ASSOCIATE EDITORS

Arjun S. Chanmugam, MD, MBA
Chair, Emergency Department
The Johns Hopkins Bayview Medical Center;
Professor and Vice Chair
Department of Emergency Medicine
The Johns Hopkins School of Medicine;
Co-Director, After Care Clinic
The Johns Hopkins Hospital
Baltimore, Maryland

Carl R. Chudnofsky, MD, FACEP
Professor and Chair
Department of Emergency Medicine
Keck School of Medicine of USC
Los Angeles, California

Peter M.C. DeBlieux, MD
Professor of Clinical Medicine
Louisiana State University Health Sciences Center
New Orleans, Louisiana

Amal Mattu, MD
Professor and Vice Chair
Department of Emergency Medicine
University of Maryland School of Medicine
Baltimore, Maryland

Stuart P. Swadron, MD, FRCPC
Associate Professor
Department of Emergency Medicine,
Assistant Dean for Pre-Health Undergraduate Studies
Keck School of Medicine of USC
University of Southern California
Los Angeles, California

Michael E. Winters, MD, FACEP, FAAEM
Associate Professor of Emergency Medicine and Medicine
University of Maryland School of Medicine;
Co-Director, Combined Emergency Medicine/Internal Medicine/Critical Care Program
University of Maryland Medical Center
Baltimore, Maryland

I H E M
iranian house of emergency medicine
خانه متخصصین طب اورژانس ایران
Roberts and Hedges’
Clinical Procedures in Emergency Medicine and Acute Care
SEVENTH EDITION

EDITOR-IN-CHIEF
James R. Roberts, MD, FACEP, FAAEM, FACMT
Professor of Emergency Medicine
The Drexel University College of Medicine
Philadelphia, Pennsylvania

SENIOR EDITOR
Catherine B. Custalow, MD, PhD
Associate Professor, Retired
Department of Emergency Medicine
University of Virginia School of Medicine
Charlottesville, Virginia

ILLUSTRATION EDITOR
Todd W. Thomsen, MD
Department of Emergency Medicine
Mount Auburn Hospital
Cambridge, Massachusetts;
Instructor in Medicine
Harvard Medical School
Boston, Massachusetts

I H E M
iranian house of emergency medicine
خانه متخصصین طب اورژانس ایران

ihem.ir
وبسایت:
t.me/ihemc
کانال تلگرام:
ایمیل:
info.ihem@yahoo.com
William J. Brady, MD, FACEP, FAAEM
Professor
Department of Emergency Medicine,
Professor
Department of Internal Medicine,
The David A. Harrison Distinguished Educator
University of Virginia School of Medicine
Charlottesville, Virginia

G. Richard Braen, MD
Professor and Founding Chairman
Department of Emergency Medicine
University at Buffalo;
Associate Dean of Graduate Medical Education
Department of Emergency Medicine
University at Buffalo
Buffalo, New York

Christine Butts, MD
Clinical Associate Professor,
Director
Division of Emergency Ultrasound
Section of Emergency Medicine
Louisiana State University Health Sciences Center
New Orleans, Louisiana

Christine M. Carr, MD, FACEP, CPE
Professor
Division of Emergency Medicine
Department of Medicine;
Department of Public Health Sciences
Medical University of South Carolina
Charleston, South Carolina

Theodore C. Chan, MD, FACEP, FAAEM
Professor and Chair
Department of Emergency Medicine
UC San Diego Health
University of California San Diego School of Medicine
San Diego, California

Mikaela L. Chilstrom, MD
Assistant Professor of Emergency Medicine
Emergency Medicine
LAC + USC Medical Center
Los Angeles, California

Arielle S. Chudnofsky, MD
PGY 3 Emergency Medicine Resident
Drexel University College of Medicine
Philadelphia, Pennsylvania

Carl R. Chudnofsky, MD, FACEP
Professor and Chair
Department of Emergency Medicine
Keck School of Medicine of USC
Los Angeles, California

Ilene Claudius, MD
Associate Professor
Department of Emergency Medicine
USC, Keck School of Medicine
Los Angeles, California

Wendy C. Coates, MD
Professor of Clinical Medicine
David Geffen School of Medicine at University of California, Los Angeles;
Senior Faculty/Education Specialist
Department of Emergency Medicine
Harbor-UCLA Medical Center
Los Angeles, California

E.C. Coffey, MD
Assistant Clinical Professor of Emergency Medicine
University of Texas Health Science Center at San Antonio
San Antonio, Texas

Alessandra Conforto, MD
Associate Professor of Clinical Emergency Medicine
Keck School of Medicine of USC
LAC + USC Medical Center
Los Angeles, California

Joseph D'Orazio, MD
Assistant Professor of Emergency Medicine,
Lewis Katz School of Medicine at Temple University
Director, Division of Medical Toxicology
Department of Emergency Medicine
Temple University
Philadelphia, Pennsylvania

Jonathan E. Davis, MD, FACEP, FAAEM
Professor & Academic Chair
Department of Emergency Medicine
Georgetown University School of Medicine & MedStar
Washington Hospital Center
Washington, D.C.

Anthony J. Dean, MD
Professor of Emergency Medicine and of Emergency Medicine in Radiology,
Director
Division of Emergency Ultrasonography
Department of Emergency Medicine
University of Pennsylvania
Philadelphia, Pennsylvania

Kenneth Deitch, DO
Attending Physician
Department of Emergency Medicine
Albert Einstein Medical Center
Philadelphia, Pennsylvania;
Attending Physician
The Lankenau Medical Center
Department of Emergency Medicine
Wynnewood, Pennsylvania

William R. Dennis, MD
Emergency Physician
St James Healthcare
Butte, Montana
CONTRIBUTORS

Valerie A. Dobiesz, MD, MPH
Core Faculty
Harvard Humanitarian Initiative,
Director of External Programs: STRATUS Center for
Medical Simulation,
Director of Women’s Leadership Initiative
Brigham & Women’s Hospital
Boston, Massachusetts

Brian E. Driver, MD
Research Director, Department of Emergency Medicine
Hennepin County Medical Center;
Assistant Professor of Emergency Medicine
University of Minnesota Medical School
Minneapolis, Minnesota

Erick Eiting, MD, MPH
Assistant Professor of Emergency Medicine
Ronald O. Perelman Department of Emergency Medicine
New York University School of Medicine
New York, New York

Timothy B. Erickson, MD
Chief, Division of Medical Toxicology
Brigham and Women’s Hospital, Department of
Emergency Medicine
Boston, Massachusetts;
Faculty, Harvard Humanitarian Initiative
Harvard Medical School
Cambridge, Massachusetts

Brian D. Euerle, MD
Associate Professor
Department of Emergency Medicine
University of Maryland School of Medicine
Baltimore, Maryland

Michael T. Fitch, MD, PhD
Professor and Vice Chair for Academic Affairs
Department of Emergency Medicine
Wake Forest School of Medicine, Winston-Salem
Winston-Salem, North Carolina

Molly Furin, MD, MS
EMS Faculty
Division of EMS and Disaster Medicine
Einstein Medical Center Philadelphia
Philadelphia, Pennsylvania

Robert T. Gerhardt, MD, MPH, FACEP
Adjunct Professor of Emergency Medicine
UTHSC-San Antonio
San Antonio, Texas;
Texas Medical Director
Prehospital and EMS Operations
Victoria Emergency Associates, LLC
Austin, Texas

John C. Greenwood, MD
Assistant Professor
Emergency Medicine
Perelman School of Medicine at the University of
Pennsylvania;
Assistant Professor
Anesthesiology & Critical Care
Perelman School of Medicine at the University of
Pennsylvania
Philadelphia, Pennsylvania

Neeraj Gupta, MD, FACEP FAAEM
Co-Director
Division of Emergency Ultrasound
Einstein Healthcare Network
Philadelphia, Pennsylvania

Richard A. Harrigan, MD, FACEP, FAAEM
Professor
Department of Emergency Medicine
Temple University School of Medicine
Philadelphia, Pennsylvania

Randy B. Hebert, MD
Assistant Clinical Professor Emergency Medicine
University of Illinois at Chicago;
Advocate
Illinois Masonic Medical Center
Chicago, Illinois

Christopher P. Holstege, MD
Professor
Departments of Emergency Medicine and Pediatrics,
Chief, Division of Medical Toxicology
University of Virginia School of Medicine;
Director
Blue Ridge Poison Center
University of Virginia Health System
Charlottesville, Virginia

Paul Jhun, MD
Assistant Professor of Emergency Medicine
University of California, San Francisco
San Francisco, California

Russell F. Jones, MD
Assistant Professor
Department of Emergency Medicine
University of California, Davis
Sacramento, California

Colin G. Kaide, MD
Associate Professor of Emergency Medicine
Department of Emergency Medicine
The Wexner Medical Center
The Ohio State University
Columbus, Ohio
Bonnie L. Kaplan, MD  
Assistant Professor,  
Associate Residency Program Director  
Department of Emergency Medicine  
University of Colorado  
Aurora, Colorado

Eric D. Katz, MD  
Executive Chair and Professor  
Department of Emergency Medicine  
University of Arizona College of Medicine;  
Chairman  
Department of Emergency Medicine  
Maricopa Integrated Health Center  
Phoenix, Arizona

John J. Kelly, DO, FACEP, FAAEM, FCPP  
Associate Chair  
Department of Emergency Medicine  
Albert Einstein Medical Center;  
Professor of Emergency Medicine  
Jefferson Medical College  
Philadelphia, Pennsylvania

John Kiel, MD  
Emergency Medicine Resident  
University of Buffalo  
Buffalo, New York

Hyung T. Kim, MD  
Health Science Associate Professor of Clinical  
Emergency Medicine  
David Geffen School of Medicine at University of  
California Los Angeles  
Los Angeles, California

Thomas D. Kirsch, MD, MPH  
Professor and Director  
National Center for Disaster Medicine and Public  
Health  
Uniformed Services University of the Health Sciences  
Bethesda, Maryland

Anne Klinke, MD, MS, FACEP  
EMS Faculty  
Department of Emergency Medicine  
Einstein Medical Center Philadelphia  
Philadelphia, Pennsylvania

Kevin J. Knoop, MD, MS  
Clinical Professor of Military and Emergency Medicine  
Uniformed Services University of the Health Sciences  
Bethesda, Maryland;  
American Red Cross Volunteer Staff Physician  
Naval Medical Center  
Portsmouth, Virginia;  
Staff Physician  
Riverside Regional Medical Center  
Newport News Virginia

J. Michael Kowalski, DO  
Attending Physician  
Department for Emergency Medicine  
Division of Medical Toxicology  
Einstein Medical Center  
Philadelphia, Pennsylvania

Alex Koyfman, MD  
Assistant Professor and Attending Physician  
Department of Emergency Medicine  
UT Southwestern Medical Center/Parkland Memorial  
Hospital  
Dallas, Texas

Diann M. Krywko, MD, FACEP  
Associate Professor,  
Director of Personal and Professional Wellness and  
Development  
Department of Medicine  
Division of Emergency Medicine  
Medical University of South Carolina  
Charleston, South Carolina

Richard L. Lammers, MD  
Professor of Emergency Medicine  
Department of Emergency Medicine,  
Assistant Dean for Simulation  
Western Michigan University Homer Stryker MD  
School of Medicine;  
Kalamazoo, Michigan

Chaiya Laoteppitaks, MD, FAAEM, FACEP  
Assistant Program Director  
Department of Emergency Medicine  
Sidney Kimmel Medical College  
Thomas Jefferson University  
Philadelphia, Pennsylvania

Glenn W. Laub, MD  
Chairman  
Department of Cardiothoracic Surgery  
Drexel University College of Medicine  
Hahnemann University Hospital  
Philadelphia, Pennsylvania

David C. Lee, MD  
Chair  
Department of Emergency Medicine  
Wellspan York Hospital  
York, Pennsylvania

Shan W. Liu

Brit Long, MD  
Physician  
San Antonio Military Medical Center, Fort Sam Houston  
Houston, Texas

Haney A. Mallemat, MD  
Assistant Professor  
Emergency Medicine and Critical Care  
University of Maryland Medical Center  
Baltimore, Maryland
CONTRIBUTORS

David E. Manthey, MD
Professor
Department of Emergency Medicine
Wake Forest School of Medicine, Winston-Salem
Winston-Salem, North Carolina

Asa M. Margolis, DO, MPH, MS
Assistant Professor
Division of Special Operations
Department of Emergency Medicine
Johns Hopkins Medical Institutions
Baltimore, Maryland

Anthony S. Mazzeo, MD
Chair
Department of Emergency Medicine
Mercy Catholic Medical Center;
Vice-Chair & Clinical Associate Professor
Department of Emergency Medicine
Drexel University College of Medicine
Philadelphia, Pennsylvania

Daniel McCollum, MD
Assistant Program Director
Department of Emergency Medicine and Hospitalist Services
Medical College of Georgia at Augusta University
Augusta, Georgia

Douglas L. McGee, DO
Chief Academic Officer
Einstein Healthcare Network;
Attending Physician
Department of Emergency Medicine
Albert Einstein Medical Center
Philadelphia, Pennsylvania

Jillian L. McGrath, MD
Assistant Professor
Department of Emergency Medicine
The Ohio State University Wexner Medical Center
Columbus, Ohio

Christopher R. McNeil, MD
Assistant Professor, Residency Program Director
Center for Emergency Medicine
University of Texas School of Medicine;
Assistant Professor, Residency Program Director
Center for Emergency Medicine
University Hospital
San Antonio, Texas

Andrew Miller, MD
Instructor
Department of Pediatrics
Harvard Medical School;
Staff Physician
Division of Emergency Medicine
Boston Children's Hospital
Boston, Massachusetts

Bohdan M. Minczak, MS, MD, PhD, FACEP/FAAEM
EMS Division Head, Faculty
Department of Emergency Medicine
Drexel University College of Medicine
Hahnemann University Hospital
Philadelphia, Pennsylvania

Aimee Moulin, MD
Associate Professor
Department of Emergency Medicine
University of California Davis Medical Center
Sacramento, California

David W. Munter, MD, MBA
Associate Clinical Professor
Department of Emergency Medicine
Eastern Virginia Medical School
Norfolk, Virginia

Joshua Nagler, MD
Department of Pediatrics
Harvard Medical School;
Fellowship Director
Division of Emergency Medicine
Boston Children's Hospital
Boston, Massachusetts

Robin M. Naples, MD
Associate Professor
Department of Emergency Medicine
University of Massachusetts Medical School
Worcester, Massachusetts

Mark J. Neavyn, MD
Assistant Professor
Department of Emergency Medicine
University of Massachusetts Medical School
Worcester, Massachusetts

Ryan Overberger, DO, MS
Clinical Assistant Professor
Sidney Kimmel Medical College
Department of Emergency Medicine
Einstein Medical Center Philadelphia
Philadelphia, Pennsylvania

Christopher R. Peabody, MD, MPH
Assistant Clinical Professor of Emergency Medicine
University of California, San Francisco
San Francisco, California

Margarita E. Pena, MD
Clinical Faculty
Department of Emergency Medicine
St. John Hospital and Medical Center
Detroit, Michigan

Phillip R. Peterson, MD
Attending Physician
Mercy Healthy System;
Clinical Associate Professor Drexel College of Medicine
Philadelphia, Pennsylvania
CONTRIBUTORS xiii

James A. Pfaff, MD
Staff Physician
San Antonio Uniformed Services Health Education Consortium;
Emergency Medicine Residency
Joint Base San Antonio-Ft. Sam Houston
San Antonio, Texas

Heather M. Prendergast, MD, MS, MPH
Professor,
Vice Chair Academic Affairs,
Assistant Dean Clinical Affairs
College of Medicine
University of Illinois Chicago Emergency Medicine
Chicago, Illinois

Robert F. Reardon, MD
Assistant Chief
Department of Emergency Medicine
Hennepin County Medical Center;
Professor of Emergency Medicine
University of Minnesota Medical School
Minneapolis, Minnesota

Salim R. Rezaie, MD
Associate Clinical Professor of Emergency Medicine/
Internal Medicine
Emergency Medicine/Internal Medicine
University of Texas Health Science Center at San
Antonio
San Antonio, Texas

Emanuel P. Rivers, MD
Clinical Professor
Emergency Medicine and Surgery
Wayne State University;
Vice Chairman and Research Director
Emergency Medicine
Henry Ford Hospital;
Senior Staff Attending
Emergency Medicine and Surgical Critical Care
Henry Ford Hospital
Detroit, Michigan

Ralph J. Riviello, MD, MS, FACEP
Professor and Vice Chair of Clinical Operations
Department of Emergency Medicine
Drexel University College of Medicine;
Medical Director
Philadelphia Sexual Assault Response Center
Philadelphia, Pennsylvania

James R. Roberts, MD, FACEP, FAAEM, FACMT
Professor of Emergency Medicine,
Senior Consultant
Division of Toxicology
The Drexel University College of Medicine;
Director
Division of Medical Toxicology
Mercy Catholic Medical Center
Philadelphia, Pennsylvania

Emily Rose, MD
Assistant Professor of Clinical Emergency Medicine
Department of Emergency Medicine
LA County + USC Medical Center
Keck School of Medicine of the University of Southern
California
Los Angeles, California

Brent E. Ruoff, MD
Chief
Division of Emergency Medicine
Washington University School of Medicine
St. Louis, Missouri

Carolyn Joy Sachs, MD, MPH
Clinical Professor of Medicine and Emergency Medicine
Ronald Reagan UCLA Emergency Medicine Department
Los Angeles, California

Leonard E. Samuels, MD
Assistant Professor
Department of Emergency Medicine
Drexel University College of Medicine
Philadelphia, Pennsylvania

Stewart O. Sanford, MD
Attending Physician
Department of Emergency Medicine
Albert Einstein Medical Center
Philadelphia, Pennsylvania

Jairo I. Santanilla, MD
Clinical Assistant Professor of Medicine
Department of Medicine, Section of Emergency
Medicine, Section of Pulmonary/Critical Care
Medicine
Louisiana State University Health Sciences Center;
Department of Pulmonary/Critical Care Medicine
Ochsner Medical Center
New Orleans, Louisiana

Genevieve Santillanes, MD
Associate Professor
Department of Emergency Medicine
Keck School of Medicine, University of Southern
California
Los Angeles, California

Richard B. Schwartz, MD
Chairman and Professor
Department of Emergency Medicine and Hospitalist
Services
Medical College of Georgia at Augusta University
Augusta, Georgia

David J. Scordino, MD
Emergency Physician
Anchorage, Alaska
Lovita E. Scrimshaw, DO  
Emergency Medicine Resident, Clinical Instructor  
Department of Emergency Medicine  
Western Michigan University School of Medicine  
Kalamazoo, Michigan

Michael A. Silverman, MD  
Chairman  
Department of Emergency Medicine  
Virginia Hospital Center  
Arlington, Virginia

Peter E. Sokolove, MD  
Professor and Chair  
Department of Emergency Medicine  
University of California San Francisco School of Medicine  
San Francisco, California

Ryan W. Spangler, MD  
Clinical Assistant Professor  
Department of Emergency Medicine  
University of Maryland School of Medicine; Attending Physician  
Mercy Medical Center  
Baltimore, Maryland

Daniel B. Stone, MD  
Clinical Assistant Professor  
Department of Medicine  
Florida International University, Herbert Wertheim College of Medicine  
Miami, Florida; Medical Director  
Department of Emergency Medicine  
Coral Springs Medical Center  
Coral Springs, Florida

Semhar Z. Tewelde, MD  
Assistant Professor  
Department of Emergency Medicine  
University of Maryland Medical System  
Baltimore, Maryland

Daniel Thomas, MD  
Emergency Medicine Resident  
University of Illinois at Chicago  
Chicago, Illinois

Laura R. Thompson, MD  
Assistant Professor of Emergency Medicine  
Department of Emergency Medicine  
The Wexner Medical Center  
The Ohio State University  
Columbus, Ohio

Jacob W. Ufberg, MD  
Professor  
Department of Emergency Medicine  
Temple University School of Medicine; Residency Director  
Department of Emergency Medicine  
Temple University Hospital  
Philadelphia, Pennsylvania

Veronica Vasquez, MD  
Assistant Professor  
Department of Emergency Medicine  
University of Southern California  
Los Angeles, California

Matthew Veltkamp, MD  
Resident Physician  
Maricopa Integrated Health Systems  
Phoenix, Arizona

Samreen Vora, MD  
HealthPartners Regions Specialty Clinics; Assistant Professor  
Department of Emergency Medicine  
Loyola University Chicago  
Chicago, Illinois

Jonathan G. Wagner, MD  
Assistant Professor of Clinical Emergency Medicine  
Department of Emergency Medicine  
Keck School of Medicine of USC  
LAC + USC Medical Center  
Los Angeles, California

Malinda Wheeler, RN, NP, SANE  
President  
Forensic Nurse Specialists, Inc.  
Long Beach, California

Michael E. Winters, MD, FACEP, FAAEM  
Associate Professor of Emergency Medicine and Medicine  
University of Maryland School of Medicine; Co-Director, Combined Emergency Medicine/Internal Medicine/Critical Care Program  
University of Maryland Medical Center  
Baltimore, Maryland

Scott H. Witt, MD  
Clinical Assistant Professor  
Associate Program Director  
Department of Emergency Medicine  
Greenville Health System  
Greenville, South Carolina

Jason Younga, MD  
Staff Emergency Physician  
Department of Emergency Medicine  
St. Anthony Medical Center  
Denver, Colorado

Richard D. Zane
The seventh edition of *Roberts and Hedges’ Clinical Procedures in Emergency Medicine and Acute Care* continues the book’s original concept of providing complete, very detailed, and up-to-date descriptions of many common, and some uncommon, procedures encountered during emergency medicine and acute care practice. The novice may find the discussions and figures devoted to the many procedures somewhat daunting or overwhelming at first; but it is hoped that most will eventually appreciate the simple discussion and complex verbiage contained in the text. The goal is to describe clinical procedures—from simple Steri-Strip application, to loop drainage of an abscess, to tracheal intubation and mechanical ventilation, to skull trephination—as though each were the nascent clinician’s first exposure to the concept, but with a depth and attention to detail that the seasoned operator would also deem helpful.

In previous editions it was difficult to find figures or photographs that conveyed the details or elucidated the vagaries to the extent one might want. The newly added color photographs, mostly digital quality, and a cornucopia of additional figures were a much needed update and morphed this edition into an obvious improvement over previous iterations. To make the text more user friendly, procedure boxes have been created, comprising a mini-atlas that allows the clinician to see the entire procedure at a glance. One can even bring the text to the bedside, viewing a single page of sequential images, the quintessential teaching tool for house staff and students. Many of the photographs were taken by me over 42 years of emergency department shifts or created or supplied by Todd W. Thomsen, MD. Some illustrations were borrowed from other sources, such as the wonderful text by Catherine B. Custalow, MD, PhD. This edition has more than 3500 images, half of which are new. More than 70 percent of the new images are the result of the artistic genius of graphics editor Dr. Thomsen. Frank Netter, watch out for Dr. Thomsen; he is rapidly attaining your status and may have already surpassed it in emergency medicine parlance. No doubt Dr. Thomsen has found his calling, blending amazing original art and electronic and digital prowess with equally impressive clinical medicine expertise. Please see Expert Consult for some bonus procedure videos from Todd.

The addition of the ultrasound-guided sections, presented in easily found and readily deciphered boxes, is the result of a gargantuan effort from our new ultrasound editor, Christine Butts, MD, an ultrasonographer extraordinaire. Another achievement of this edition is the inclusion of additional video procedures. Only wished for in other texts, many sections now reference online content that allows the reader to view videos of the procedures actually being performed. “See one, do one, teach one” has taken on new meaning with this text. This edition is now available electronically on such devices as the Kindle and iPad and is still fully searchable online at expert-consult.com.

There are, of course, many ways to approach any patient or any procedure, so this text is not a dictum and is not truly authoritative. This book does not attempt to define standard of care. The clinician should remember these caveats if involved in a medicolegal scenario. It contains practical hints and successful tactics gleaned from the literature and by years of practice, adroitly described by skilled clinicians. As with previous editions, this version also significantly incorporates the personal opinions and experience of the authors and editors. But this text is simply a clinical guide, not a legal document. Do not reference this book if you testify in court, for either the defense or the plaintiff. Today’s dogma too often becomes tomorrow’s heresy, and physician hubris is worse than incompetence. Simply stated, emergency medicine and acute care, and the human body too often defy the written word, personal opinion, or local custom and humble even the venerable and the universally praised gray-haired professor.

Many new authors have been added, as well as a number of new concepts and approaches. All procedures have been tweaked. As an example one will not find the novel loop abscess drainage technique, sedation techniques, or ENT and ophthalmology techniques so nicely described elsewhere.

My personal thanks are hereby conveyed to those who contributed to previous editions. The updated chapters often merely refine or further manipulate the scholarly work of others who originally assisted us. The current contributors include an enviable blend of friends and colleagues, some former students of mine, up-and-coming rising stars in their own right, and my prior mentors and role models—all are accomplished physicians and leaders in their own milieu. Many of the editors’ names are well known to anyone who reads the literature or attends a continuing medical education activity. All of the associate editors portray and embody the pinnacle of emergency medicine and acute care excellence. Most of the contributors, and all the associate editors, probably know more than I know, and most are likely infinitely more capable and facile with procedures. All are capable of writing a text themselves, and some have already done so; however, some are now enlightened and eschew that primal urge because they now know how difficult it is to write even a single chapter. My able and erudite associate editors, all from prestigious academic teaching programs in emergency medicine, are Arjun S. Channugam, MD, MBA; Carl R. Chudnoffsky, MD, FACEP; Peter M.C. DeBlieux, MD; Amal Mattu, MD; Stuart P. Swadron, MD, FRCP, and recently added Michael Winters, MD, FACEP, FAAEM. They provided the bulk of the original editing, but senior editor, Dr. Custalow, read every single word and reviewed every table and figure. Dr. Custalow is a more tenacious editor than the proverbial honey badger in regard to dealing with details, grammar, organization, and style. In the end, my personal bias may be evident, but Dr. Custalow was the fire and fuel for the book’s framework. As already stated, Dr. Thomsen made the text come to life with images. I would also like to thank Robert Orman, MD, and Scott D. Weingart, MD, FCCM, who served as video editors on the previous edition.
If any of our editing changed, altered, or misinterpreted the original thoughts of the contributors (and I know in some instances it must have), we apologize; but hard decisions had to be made, and waffling was rarely an option. Our book simply tells you what to do and how and when to do it, but no book can always fit every individual situation. We attempted to squarely address such omnipresent vague topics as prophylactic antibiotics, local customs, and variations in style, and accepted the fact that not all foreign bodies or tendon lacerations will be identified in the heat of the moment by even the most skilled clinician. The prescient and sagacious clinician knows that the ability to practice medicine from a book is limited, and one learns best from past experiences; and, for certain, the most instructive past experience is one that was not always textbook perfect.

James R. Roberts, MD, FACEP, FAAEM, FACMT
Every shift in the emergency department has the potential to take us into the great unknown. Frequently a patient’s presentation challenges our capabilities, and an uncommon procedure may be the only answer. Throughout my training and career, this book you are now holding has been my unwavering companion.

On a recent night shift, I encountered a patient needing two separate procedures, both well within my capabilities as an emergency physician. Yet, I found myself asking, “Do I remember exactly how to do the procedure? What is the best approach? Where are my landmarks?” I was almost positive I could recall the technique correctly, but before breaking skin, I decided a review of *Roberts and Hedges* was in order. Taking this step dispels any doubt and gives me confidence to get the job done well.

Whether it’s tapping a septic elbow, checking a compartment pressure, or performing a lateral canthotomy, the refrain is always the same: “How, exactly, do I do this?” becomes “Right, oh yes, I’ve got this down!”

As we move further into the world of online and on-demand education, some may say that textbooks are a vestige of the past, but I disagree in this instance. Even after decades in print, *Roberts and Hedges* continues to shine as the guiding light of clinical emergency medicine.

Rob Orman, MD
June 1, 2017
Gargantuan efforts, clairvoyant and perceptive suggestions, and decidedly prescient and sagacious contributions of many individuals have brought this work to fruition. Not the least of whom were the individual authors who toiled over tedious manuscripts and answered countless queries about the vagaries and vicissitudes of seemingly straightforward clinical procedures.

This book’s current edition was championed by Elsevier’s Senior Content Strategist, Kate Dimock. All of the initially submitted work was culled, corrected, and collated by the very able Senior Content Development Specialist, Jennifer Ehlers. Senior Project Manager, Rachel McMullen, also toiled over every aspect of the text. My sincere gratitude to them is warmly extended with this acknowledgment. If any reader is contemplating developing their own textbook, snag this team of publishing aficionados if you can.

Of course, the entire work was infused with vim and vigor from Catherine B. Custalow, MD, PhD, and every image was created, beautified, or otherwise superbly orchestrated by Todd W. Thomsen, MD. The final editing of Arjun S. Chanmugam, MD, MBA; Carl R. Chudnofsky, MD, FACEP; Peter M.C. DeBlieux, MD; Amal Mattu, MD; and Stuart P. Swadron, MD, FRCPC, and Michael Winters, MD, FACEP, FAAEM, completed the task. One might think that these folks have a lot of free time on their hands or, more likely, they burned gallons of midnight oil for the project. It has been an honor to be associated with such icons in the field. The contributing authors are certainly also to be congratulated on a stellar performance.

Thank you all for accomplishing a goal that was once thought, even by me, to be nothing more than a seemingly good idea, but a task too difficult to even contemplate, let alone wantonly attempt. It is hard to believe that the first edition of this book was done in 1985 on an IBM electronic typewriter, before email, scanning, fax, or Microsoft Word. We have come a long way for sure.

James R. Roberts, MD, FACEP, FAAEM, FACMT
SECTION I  Vital Signs and Patient Monitoring Techniques  

Chapter 1  Vital Signs Measurement  
Jillian L. McGrath and Daniel J. Bachmann

Chapter 2  Devices for Assessing Oxygenation and Ventilation  
Andrew Miller and Joshua Nagler

SECTION II  Respiratory Procedures  

Chapter 3  Basic Airway Management and Decision Making  
Brian E. Driver and Robert F. Reardon

Chapter 4  Tracheal Intubation  
Brian E. Driver and Robert F. Reardon

Chapter 5  Pharmacologic Adjuncts to Intubation  
Richard B. Schwartz and Daniel McCollum

Chapter 6  Cricothyrotomy and Percutaneous Translaryngeal Ventilation  
Randy B. Hebert and Daniel Thomas

Chapter 7  Tracheostomy Care  
John C. Greenwood and Michael E. Winters

Chapter 8  Mechanical Ventilation  
Jairo I. Santanilla

Chapter 9  Thoracentesis  
Barbara K. Blok

Chapter 10  Tube Thoracostomy  
Asa M. Margolis and Thomas D. Kirsch

SECTION III  Cardiac Procedures  

Chapter 11  Techniques for Supraventricular Tachycardias  
Bohdan M. Minczak and Glenn W. Laub

Chapter 12  Defibrillation and Cardioversion  
Bohdan M. Minczak and Glenn W. Laub

Chapter 13  Assessment of Implantable Devices  
James A. Pfaff and Robert T. Gerhardt

Chapter 14  Basic Electrocardiographic Techniques  
William J. Brady, Richard A. Harrigan, and Theodore C. Chan

Chapter 15  Emergency Cardiac Pacing  
Edward S. Bessman

Chapter 16  Pericardiocentesis  
Haney A. Mallemat and Semhar Z. Tewelde

Chapter 17  Cardiopulmonary Resuscitation and Artificial Perfusion During Cardiac Arrest  
Benjamin S. Abella and Lance B. Becker

Chapter 18  Resuscitative Thoracotomy  
Russell F. Jones and Emanuel P. Rivers

SECTION IV  Vascular Techniques and Volume Support  

Chapter 19  Pediatric Vascular Access and Blood Sampling Techniques  
Genevieve Santillanes and Ilene Claudius

Chapter 20  Arterial Puncture and Cannulation  
Erick Eiting and Hyung T. Kim

Chapter 21  Peripheral Intravenous Access  
Bonnie L. Kaplan, Shan W. Liu, and Richard D. Zane

Chapter 22  Central Venous Catheterization and Central Venous Pressure Monitoring  
Salim R. Rezaie, E.C. Coffey, and Christopher R. McNeil

Chapter 23  Venous Cutdown  
Veronica Vasquez and Pablo F. Aguiler

Chapter 24  Indwelling Vascular Access Devices: Emergency Access and Management  
Scott H. Witt, Christine M. Carr, and Diann M. Krywko
Chapter 54  Compartment Syndrome Evaluation  1125  
Chaiya Laoteppitaks

SECTION IX  Genitourinary, Obstetric, and Gynecologic Procedures  1141

Chapter 55  Urologic Procedures  1141  
Jonathan E. Davis and Michael A. Silverman

Chapter 56  Emergency Childbirth  1186  
Samreen Vora and Valerie A. Dobiesz

Chapter 57  Gynecologic Procedures  1211  
G. Richard Braen and John Kiel

Chapter 58  Examination of the Sexual Assault Victim  1225  
Carolyn Joy Sachs and Malinda Wheeler

SECTION X  Neurologic Procedures  1243

Chapter 59  Management of Increased Intracranial Pressure and Intracranial Shunts  1243  
Alessandra Conforto and Jonathan G. Wagner

Chapter 60  Spinal Puncture and Cerebrospinal Fluid Examination  1258  
Brian D. Euerle

Chapter 61  Special Neurologic Tests and Procedures  1281  
Mikaela L. Chilstrom

SECTION XI  Ophthalmologic, Otolaryngologic, and Dental Procedures  1295

Chapter 62  Ophthalmologic Procedures  1295  
Kevin J. Knoop and William R. Dennis

Chapter 63  Otolaryngologic Procedures  1338  
Ralph J. Riviello

Chapter 64  Emergency Dental Procedures  1384  
Kip R. Benko

SECTION XII  Special Procedures  1405

Chapter 65  Procedures Pertaining to Hypothermia and Hyperthermia  1405  
Heather M. Prendergast and Timothy B. Erickson

Chapter 66  Ultrasound  1434  
Christine Butts

Chapter 67  Bedside Laboratory and Microbiologic Procedures  1442  
Anthony J. Dean and David C. Lee

Chapter 68  Standard Precautions and Infectious Exposure Management  1470  
Peter E. Sokolove and Aimee Moulin

Chapter 69  Physical and Chemical Restraint  1481  
J. Michael Kowalski

Chapter 70  Noncardiac Implantable Devices  1499  
Paul Jhun and Christopher R. Peabody

Chapter 71  Radiation in Pregnancy and Clinical Issues of Radiocontrast Agents  1504  
Phillip R. Peterson

Chapter 72  Procedures in the Setting of Anticoagulation  1520  
Joseph D’Orazio

Appendix  Commonly Used Formulas and Calculations  1529  
Matthew Veltkamp, Eric D. Katz, and Brent E. Ruoff
Video 1.1.  
Ankle-brachial index.

Video 3.1.  
Nasopharyngeal and oropharyngeal airways. (Courtesy Jared Novack and Ernest Wang.)

Video 3.2.  
Oxygen delivery. (Courtesy Jared Novack and Ernest Wang.)

Video 3.3.  
Intubation confirmation. (Courtesy Jared Novack and Ernest Wang.)

Video 3.4.  
Pentax AWS in patient with laryngeal mass. (Courtesy James DuCanto.)

Video 3.5.  
Rapid-sequence airway to rapid-sequence intubation. (Courtesy James DuCanto.)

Video 3.6.  
Delayed sequence intubation. (Courtesy Scott D. Weingart.)

Video 3.7.  
Noninvasive positive pressure ventilation—CPAP and BiPAP. (Courtesy Jared Novack and Ernest Wang.)

Video 3.8.  
CPAP preoxygenation. (Courtesy Scott D. Weingart.)

Video 3.9.  
Boussignac CPAP. (Courtesy James DuCanto.)

Video 3.10.  
Double lumen tube placement. (Courtesy Lance Carter and Adam Petersen.)

Video 3.11.  
Bronchial blocker placement. (Courtesy Lance Carter and Adam Petersen.)

Video 4.1.  
Endotracheal intubation with continuous oxygenation. (Courtesy Alfred Sacchetti.)

Video 4.2.  
Rapid sequence endotracheal intubation. (Courtesy Alfred Sacchetti.)

Video 4.3  
Standard endotracheal intubation. (Courtesy Jared Novack and Ernest Wang.)

Video 4.4  
Skills of direct laryngoscopy. (Courtesy Scott D. Weingart.)

Video 4.5  
Intubation through AirQ. (Courtesy James DuCanto.)

Video 4.6  
Video-assisted intubation. (Courtesy Larry B. Mellick.)

Video 4.7  
Glidescope intubation. (Courtesy Mel Herbert.)

Video 4.8  
Insertion of cookgas AirQ SGA. (Courtesy Daniel Cook.)

Video 4.9  
Storz videoscope endotracheal intubation. (Courtesy Larry B. Mellick.)

Video 4.10  
Difficult airways with video laryngoscope. (Courtesy Larry B. Mellick.)

Video 4.11  
Retrograde intubation (Cadaveric). (Courtesy Siamak Moayedi and Dan Lemkin.)

Video 4.12  
Retrograde intubation. (Courtesy William H. Rosenblatt.)

Video 4.13  
Storz C-MAC intubation with Bougie. (Courtesy Larry B. Mellick.)

Video 4.14  
Awake intubation. (Courtesy Scott D. Weingart.)

Video 4.15  
Video-assisted endotracheal intubation with curved pocket Bougie. (Courtesy Fayaz Gulamani.)

Video 4.16  
Intubating around a King LT. (Courtesy James DuCanto.)

Video 4.17  
King vision. (Courtesy James DuCanto.)
Video 4.18  
Levitan FPS scope through laryngeal airway and with DL. (Courtesy James DuCanto.)

Video 4.19  
Glidescope and Shikani stylet. (Courtesy James DuCanto.)

Video 4.20  
Fiberoptic bronch with aintree through King LT. (Courtesy James DuCanto.)

Video 4.21  
McGrath intubation. (Courtesy James DuCanto.)

Video 6.1.  
Surgical cricothyrotomy. (Courtesy Siamak Moayedi and Ernest Wang.)

Video 6.2.  
Bougie-aided cricothyrotomy. (Courtesy Darren Braude.)

Video 10.1  
Needle thoracostomy. (Courtesy Jared Novack and Ernest Wang.)

Video 10.2  
Tube thoracostomy—standard technique. (Courtesy Siamak Moayedi and Dan Lemkin.)

Video 10.3  
Pigtail thoracostomy. (Courtesy Alfred Sacchetti.)

Video 10.4  
Tube thoracostomy—Seldinger technique. (Courtesy Joe Mayerle and Scott A. Joing.)

Video 10.5  
Chest tube thoracostomy hemothorax. (Courtesy Anna Bargren and Andrew Pendley.)

Video 10.6  
Securing a chest tube. (Courtesy Gary Marks.)

Video 10.7  
Tru-Close chest tube. (Courtesy Larry B. Mellick.)

Video 10.8  
Finger-bougie-endotracheal tube thoracostomy. (Courtesy Cliff Reid.)

Video 12.1  
Defibrillation and cardioversion. (Courtesy Jared Novack and Ernest Wang.)

Video 12.2  
Electrical cardioversion for atrial flutter. (Courtesy Larry B. Mellick.)

Video 12.3  
Electrical cardioversion—emergent. (Courtesy Larry B. Mellick.)

Video 15.1  
Transvenous pacemaker placement. (Courtesy Jared Novack and Ernest Wang.)

Video 15.2  
Transvenous pacemaker insertion. (Courtesy Alfred Sacchetti.)

Video 15.3  
Transcutaneous pacing. (Courtesy Jared Novack and Ernest Wang.)

Video 16.1  
Tamponade and pericardiocentesis. (Courtesy Adam Bystrzycki.)

Video 16.2  
Focused cardiac ultrasound: evaluation of pericardial effusion. (Courtesy Joshua Rempell and Michael Stone.)

Video 17.1  
LUCAS chest compression system. (Courtesy Larry B. Mellick.)

Video 17.2  
ECMO. (Courtesy Zachary Shinar and Joe Bellezo.)

Video 18.1  
Resuscitative thoracotomy. (Courtesy Siamak Moayedi and San Lemkin.)

Video 19.1  
Pediatric intravenous insertion. (Courtesy Alfred Sacchetti.)

Video 19.2  
Umbilical vein catheter. (Courtesy Mel Herbert.)

Video 20.1  
Arterial line placement. (Courtesy Chrissa Strumpe and Jared Novack.)

Video 20.2  
Arterial line insertion—arrow kit. (Courtesy Lance Carter and Adam Petersen.)
Video 20.3  
Radial arterial line insertion. (Courtesy James Bryant.)

Video 20.4  
Femoral arterial line insertion—ultrasound guided.  
(Courtesy James Bryant.)

Video 21.1  
Ultrasound-guided deep brachial IV. (Courtesy Gary Marks.)

Video 21.2  
Rapid infusion catheter. (Courtesy Tim Leeuwenburg.)

Video 22.1  
Central line kit—introduction. (Courtesy Siamak Moayedi and Dan Lemkin.)

Video 22.2  
Central line insertion—internal jugular approach. (Courtesy Jared Novack and Ernest Wang.)

Video 22.3  
Central line insertion—subclavian. (Courtesy Siamak Moayedi and Dan Lemkin.)

Video 22.4  
Supraventricular line. (Courtesy Mel Herbert.)

Video 22.5  
Central line placement—subclavian. (Courtesy Scott D. Weingart.)

Video 22.6  
Setting up the pressure set for CVP and A-lines. (Courtesy Scott D. Weingart.)

Video 22.7  
Artery or vein confirmation. (Courtesy Scott D. Weingart.)

Video 22.8  
Central line sterility. (Courtesy Scott D. Weingart.)

Video 23.1  
Venous cutdown. (Courtesy Jared Novack and Ernest Wang.)

Video 24.1  
How to access an indwelling vascular port. (Courtesy Alfred Sacchetti.)

Video 24.2  
Dialysis graft oversew. (Courtesy Alfred Sacchetti.)

Video 24.3  
Repair of bleeding dialysis shunt. (Courtesy Alfred Sacchetti.)

Video 25.1  
Intraosseous needle placement during CPR. (Courtesy Larry B. Mellick.)

Video 25.2  
Intraosseous needle placement—mistakes to avoid.  
(Courtesy Larry B. Mellick.)

Video 25.3  
Intraosseous needle placement—pediatric. (Courtesy Ernest Wang.)

Video 25.4  
Intraosseous needle insertion. (Courtesy Jared Novack and Ernest Wang.)

Video 25.5  
Intraosseous needle—humeral. (Courtesy Mel Herbert.)

Video 27.1  
Pleur-evac autotransfusion. (Courtesy Scott D. Weingart.)

Video 29.1  
Hematoma block. (Courtesy Larry B. Mellick.)

Video 31.1  
Ultrasound-guided nerve blocks in emergency care.  
(Courtesy Mike Stone.)

Video 31.2  
Wrist blocks—median, radial, and ulnar nerves. (Courtesy Daniel Gromis and Anton J. Fakhouri.)

Video 31.3  
Digital nerve block of the thumb. (Courtesy Daniel Gromis and Anton J. Fakhouri.)

Video 31.4  
Ankle nerve blocks. (Courtesy Gary Marks.)

Video 31.5  
Fascia iliaca block—pediatric femur fracture. (Courtesy Alfred Sacchetti.)

Video 31.6  
Using ultrasound to find the brachial plexus in the interscalene space. (Courtesy Jack Vander Beek.)

Video 31.7  
Ultrasound-guided median nerve block. (Courtesy Mike Stone.)
Video 31.8
Ultrasound-guided radial nerve block. (Courtesy Mike Stone.)

Video 31.9
Ultrasound-guided distal sciatic nerve block. (Courtesy Mike Stone.)

Video 31.10
Ultrasound-guided tibial nerve block. (Courtesy Mike Stone.)

Video 31.11
Ultrasound-guided ulnar nerve block. (Courtesy Mike Stone.)

Video 31.12
Ultrasound-guided axillary brachial plexus nerve block. (Courtesy Mike Stone.)

Video 31.13
Ultrasound-guided infraclavicular brachial plexus nerve block. (Courtesy Mike Stone.)

Video 31.14
Ultrasound-guided interscalene brachial plexus nerve block. (Courtesy Mike Stone.)

Video 32.1
Bier block. (Courtesy Alfred Sacchetti.)

Video 33.1
Procedural sedation with ketamine. (Courtesy Larry B. Mellick.)

Video 34.1
Equipment. (Courtesy Michelle Lin.)

Video 34.2
Anesthesia. (Courtesy Michelle Lin.)

Video 34.3
Wound irrigation. (Courtesy Michelle Lin.)

Video 34.4
Starting the sterile procedure. (Courtesy Michelle Lin.)

Video 35.1
Steri-Strips. (Courtesy Michelle Lin.)

Video 35.2
Dermabond. (Courtesy Michelle Lin.)

Video 35.3
Staples. (Courtesy Michelle Lin.)

Video 35.4
Buried sutures (subcutaneous). (Courtesy Michelle Lin.)

Video 35.5
Simple interrupted sutures. (Courtesy Michelle Lin.)

Video 35.6
Vertical mattress sutures. (Courtesy Michelle Lin.)

Video 35.7
Horizontal mattress sutures. (Courtesy Michelle Lin.)

Video 35.8
Running horizontal mattress sutures. (Courtesy Alfred Sacchetti.)

Video 35.9
Corner sutures. (Courtesy Michelle Lin.)

Video 35.10
Ingrown toenail removal. (Courtesy John Zangmeister.)

Video 36.1
Fish hook removal. (Courtesy Larry B. Mellick.)

Video 36.2
Nail gun injury. (Courtesy Larry B. Mellick.)

Video 36.3
Removal of zipper for penile entrapment. (Courtesy Mel Herbert.)

Video 36.4
Managing IUD presentations. (Courtesy Gary Marks.)

Video 37.1
Incision and drainage. (Courtesy Jared Novack and Ernest Wang.)

Video 37.2
Loop drainage technique for cutaneous abscess. (Courtesy Robert Orman.)

Video 37.3
Subungual hematoma—trephination with battery-powered cautery. (Courtesy Larry B. Mellick.)

Video 37.4
Paronychia incision and drainage. (Courtesy Larry B. Mellick.)

Video 37.5
Subungual hematoma—trephination using needle. (Courtesy John Zangmeister.)
Video 40.1
Nasogastric/orogastric tube placement. (Courtesy Lance Carter.)

Video 40.2
Nasogastric tube insertion. (Courtesy Jared Novack and Ernest Wang.)

Video 40.3
Nasogastric intubation. (Courtesy Whit Fisher.)

Video 40.4
Transabdominal feeding tube replacement. (Courtesy Whit Fisher.)

Video 40.5
MIC-KEY gastrostomy feeding tube (Halyard Health, Alpharetta, GA) placement. (Courtesy Larry B. Mellick.)

Video 41.1
Blakemore tube placement. (Courtesy Tammar Taddei.)

Video 42.1
Gastric lavage. (Courtesy Joseph Maddry.)

Video 43.1
Paracentesis. (Courtesy Alfred Sacchetti.)

Video 43.2
Paracentesis—simulation. (Courtesy Jared Novack and Ernest Wang.)

Video 46.1
Cervical collar. (Courtesy Benjamin H. Slovis, Avital Porat, Neil Singh, and Kevin Chason.)

Video 46.2

Video 46.3

Video 46.4
Traction splint. (Courtesy Benjamin H. Slovis, Avital Porat, Neil Singh, William Fleischman, and Kevin Chason.)

Video 47.1
Field amputation: introduction. (Courtesy Steve Zils and Ronald Pirrallo.)

Video 47.2
Field amputation: upper extremity. (Courtesy Steven Zils, Panna Codner, and Ronald Pirrallo.)

Video 47.3
Field amputation: lower extremity. (Courtesy Steven Zils, Panna Codner, and Ronald Pirrallo.)

Video 48.1
Extensor tendon repair. (Courtesy Mel Herbert.)

Video 48.2
Boxer’s fracture. (Courtesy Larry B. Mellick.)

Video 48.3
Salter Harris II radius fracture reduction. (Courtesy Larry B. Mellick.)

Video 49.1
Hip dislocation reduction—standard technique. (Courtesy Larry B. Mellick.)

Video 49.2
Hip dislocation reduction—Whistler technique. (Courtesy George Douros.)

Video 49.3
Hip dislocation—Captain Morgan technique. (Courtesy Alfred Sacchetti.)

Video 49.4
Scapular manipulation. (Courtesy Neil Cunningham and Gerard Fennessy.)

Video 49.5
Zero position technique. (Courtesy Neil Cunningham and Gerard Fennessy.)

Video 49.6
Can’t adduct—troubleshooting positioning. (Courtesy Neil Cunningham and Gerard Fennessy.)

Video 49.7
Difficult dislocation—using sedation for spasm. (Courtesy Neil Cunningham and Gerard Fennessy.)

Video 49.8
Shoulder dislocation reduction—Kocher’s technique. (Courtesy Neil Cunningham and Gerard Fennessy.)

Video 49.9
Shoulder dislocation reduction—Cunningham technique. (Courtesy Neil Cunningham and Gerard Fennessy.)

Video 49.10
Shoulder dislocation reduction—Spaso technique. (Courtesy George Douros.)
Video 49.11
Anterior shoulder dislocation reduction—external rotation technique. (Courtesy Daniel Gromis.)

Video 49.12
Reduction of luxatio erecta. (Courtesy Mel Herbert.)

Video 49.13
Shoulder dislocation reduction—using ultrasound to guide intraarticular lidocaine injections. (Courtesy Michael Stone.)

Video 49.14
Shoulder dislocation reduction. (Courtesy Larry B. Mellick.)

Video 49.15
Ankle dislocation reduction. (Courtesy Mel Herbert.)

Video 49.16
Finger dislocation reduction and metacarpal block. (Courtesy Larry B. Mellick.)

Video 49.17
Finger dislocation reduction. (Courtesy Larry B. Mellick.)

Video 49.18
Posterior elbow dislocation reduction. (Courtesy Larry B. Mellick.)

Video 49.19
Posterior elbow dislocations—reduction with prone and supine patient positioning. (Courtesy Daniel Gromis and Anton J. Fakhouri.)

Video 49.20
Elbow reduction. (Courtesy Mel Herbert.)

Video 49.21
Nursemaids elbow reduction. (Courtesy Larry B. Mellick.)

Video 49.22
Patella dislocation reduction. (Courtesy Larry B. Mellick.)

Video 50.1
Posterior lower leg splint with stirrup (below knee and the U slab). (Courtesy Robert Orman.)

Video 50.2
Thumb spica splint. (Courtesy Robert Orman.)

Video 50.3
Posterior lower leg split (no stirrup) using plaster. (Courtesy Robert Orman.)

Video 50.4
Sugar tong splint. (Courtesy Robert Orman.)

Video 50.5
Ulnar gutter splint. (Courtesy Robert Orman.)

Video 50.6
Mallet finger—examination and splinting technique. (Courtesy Daniel Gromis and Anton J. Fakhouri.)

Video 51.1
Ingrown toenail management. (Courtesy Larry B. Mellick.)

Video 51.2
Nail removal for onychomycosis. (Courtesy Larry B. Mellick.)

Video 51.3
Toenail removal. (Courtesy Mel Herbert.)

Video 52.1
Techniques for shoulder injections/aspirations. (Courtesy Daniel Gromis and Anton J. Fakhouri.)

Video 52.2
Subacromial bursa injection. (Courtesy Larry B. Mellick.)

Video 53.1
Techniques for wrist joint injections/aspirations. (Courtesy Daniel Gromis and Anton J. Fakhouri.)

Video 53.2
Wrist arthrocentesis. (Courtesy Siamak Moayedi and Dan Lemkin.)

Video 53.3
Elbow arthrocentesis. (Courtesy Larry B. Mellick.)

Video 53.4
Olecranon bursa aspiration. (Courtesy Larry B. Mellick.)

Video 53.5
Subacromial bursa injection. (Courtesy Larry B. Mellick.)

Video 53.6
Knee arthrocentesis—medial approach. (Courtesy Siamak Moayedi and Dan Lemkin.)

Video 53.7
Knee arthrocentesis. (Courtesy Jared Novack and Ernest Wang.)

Video 53.8
Ankle arthrocentesis. (Courtesy George Douros.)

Video 53.9
Metatarsophalangeal joint aspiration. (Courtesy Larry B. Mellick.)

Video 53.10
Gout and metatarsophalangeal joint arthrocentesis. (Courtesy Larry B. Mellick.)
**Video 54.1**
Measuring Compartment Pressures (Stryker Monitor).
(Courtesy Daniel Gromis, Anton J. Fakhouri, and Gary Marks.)

**Video 54.2**
Compartment Pressure Measurement (Stryker). (Courtesy Gary Marks.)

**Video 55.1**
Dorsal slit. (Courtesy Carlo Astini.)

**Video 55.2**
Paraphimosis reduction. (Courtesy Jared Novack and Ernest Wang.)

**Video 55.3**
Percutaneous suprapubic cystostomy. (Courtesy Siamak Moayedi and Dan Lemkin.)

**Video 56.1**
Cesarean section. (Courtesy Najeeb Layyous.)

**Video 56.2**
Perimortem C-section simulation model. (Courtesy Jason Wagner; narration by Christopher Sampson, MD.)

**Video 60.1**
Adult lumbar puncture. (Courtesy Larry B. Mellick.)

**Video 60.2**
Pediatric lumbar puncture: septic workup part I. (Courtesy Alfred Sacchetti.)

**Video 61.1**
Dix-Hallpike test and Epley maneuver. (Courtesy Larry B. Mellick.)

**Video 61.2**
Epley maneuver. (Courtesy Felipe Teran.)

**Video 61.3**
HINTS exam. (Courtesy Scott D. Weingart.)

**Video 61.4**
Tensilon test for myasthenia gravis. (Courtesy Randy Kardon and Thomas A. Oetting.)

**Video 62.1**
Introduction to the eye exam. (Courtesy David K. Duong.)

**Video 62.2**
Video visual acuity testing. (Courtesy David K. Duong.)

**Video 62.3**
Video Morgan lens insertion. (Courtesy Alfred Sacchetti.)

**Video 62.4**
Video tonometry. (Courtesy David K. Duong.)

**Video 62.5**
Slit lamp examination. (Courtesy David K. Duong.)

**Video 62.6**
Video lateral canthotomy. (Courtesy Siamak Moayedi and Daniel Lemkin.)

**Video 63.1**
Emergent management of posterior epistaxis. (Courtesy Jared Novack and Ernest Wang.)

**Video 63.2**
Anterior epistaxis management. (Courtesy Jared Novack and Ernest Wang.)

**Video 63.3**
Ear canal foreign body removal using cyanoacrylate. (Courtesy Tim Young and Melanie M. Randall.)

**Video 63.4**
Ear laceration repair. (Courtesy Mel Herbert.)

**Video 63.5**
Ear foreign body—cockroach emergency. (Courtesy Larry B. Mellick.)

**Video 63.6**
Nasal foreign body removal techniques. (Courtesy Larry B. Mellick.)

**Video 63.7**
Nasal foreign body removal—Katz extractor. (Courtesy Larry B. Mellick.)

**Video 63.8**
Mandibular dislocation reduction—part 1. (Courtesy Larry B. Mellick.)

**Video 63.9**
Mandibular dislocation reduction—part 2. (Courtesy Larry B. Mellick.)

**Video 63.10**
Reduction of spontaneous mandibular dislocation with masseteric massage. (Courtesy Daniel Gromis.)

**Video 64.1**
Reimplantation of avulsed tooth. (Courtesy Larry B. Mellick.)

**Video 64.2**
Tongue laceration repair. (Courtesy Larry B. Mellick.)
Video 64.3
Video dry socket. (Courtesy Larry B. Mellick.)

Video 66.1
Introduction to ultrasound for procedure guidance. (Courtesy Bret Nelson.)

Video 66.2
Internal jugular central line placement—ultrasound guided. (Courtesy Michael Mallin and Matt Dawson.)

Video 66.3
Lumbar puncture—ultrasound guided. (Courtesy Michael Mallin and Matt Dawson.)

Video 66.4
Peripheral IV placement—ultrasound guided. (Courtesy Michael Mallin and Matt Dawson.)

Video 66.5
Paracentesis—ultrasound guided. (Courtesy Michael Mallin and Matt Dawson.)

Video 66.6
Thoracentesis—ultrasound guided, quick reference (2 minutes). (Courtesy Michael Mallin and Matt Dawson.)

Video 66.7
Ultrasound guidance for thoracentesis: extended reference (10 minutes). (Courtesy Phillips Perera.)

Video 66.8
Radial arterial line placement—ultrasound guided. (Courtesy Michael Mallin and Matt Dawson.)

Video 66.9
Pericardiocentesis—ultrasound guided. (Courtesy Michael Mallin and Matt Dawson.)

Video 66.10
DVT ultrasound. (Courtesy Gary Marks.)

Video 66.11
Introduction to the FAST exam. (Courtesy Michael Stone.)

Video 66.12
Pulmonary ultrasound. (Courtesy Michael Stone.)

Video 66.13
Point-of-care ultrasound for the detection of abdominal aortic aneurysm. (Courtesy Heidi Harbison Kimberly.)

Video 66.14
First trimester pelvic ultrasonography. (Courtesy Michael Stone.)

Video 66.15
Ultrasound physics and knobology. (Courtesy Michael Stone.)

Video 69.1
4-Point restraint. (Courtesy Gary Marks.)

SUPPLEMENTARY VIDEOS
Arthrocentesis: shoulder. (Courtesy Todd W. Thomsen.)
Basics of wound management. (Courtesy Todd W. Thomsen.)
Chest tube placement. (Courtesy Todd W. Thomsen.)
Compartment syndrome evaluation. (Courtesy Todd W. Thomsen.)
Digital nerve block. (Courtesy Todd W. Thomsen.)
Digital nerve block. (Courtesy Todd W. Thomsen.)
Dislocation reduction of the ankle joint. (Courtesy Todd W. Thomsen.)
Dislocation reduction of the elbow joint. (Courtesy Todd W. Thomsen.)
Dislocation reduction of the hip joint. (Courtesy Todd W. Thomsen.)
Dislocation reduction of the patella. (Courtesy Todd W. Thomsen.)
Dislocation reduction of the shoulder joint. (Courtesy Todd W. Thomsen.)
Irrigation of the eye. (Courtesy Todd W. Thomsen.)
Measurement of intraocular pressure: tono-pen technique. (Courtesy Todd W. Thomsen.)
Nasogastric intubation. (Courtesy Todd W. Thomsen.)
Orotracheal intubation. (Courtesy Todd W. Thomsen.)
Tube thoracostomy. (Courtesy Todd W. Thomsen.)
Urethral catheterization: male. (Courtesy Todd W. Thomsen.)
CHAPTER 1
Vital Signs and Patient Monitoring Techniques

Vital Signs Measurement
Jillian L. McGrath and Daniel J. Bachmann

Measuring the temperature, pulse, respiratory rate (RR), blood pressure, and pulse oximetry is generally recommended for all emergency department (ED) patients, in addition to assessment of pain in the appropriate patient population. For very minor problems or for some fast-track patients (e.g., suture removal), a full set of vital signs may not be required, and this is best decided on a case-by-case basis rather than by strict protocol. Vital signs may indicate the severity of illness and also dictate the urgency of intervention. Although a single set of abnormal values suggests pathology, findings on triage or the initial vital signs may be spurious and simply related to stress, anxiety, pain, or fear. It would be incorrect and not standard of care to attribute initial triage blood pressure, RR, or pulse rate to specific pathology or to retrospectively assume that diagnostic or treatment interventions should have been initiated based solely on these readings. The greatest utility of vital signs is in their continued observation and trends over time. Deteriorating vital signs are an important indicator of a compromised physiologic condition, and improving values provide reassurance that the patient may be responding to therapy. When a patient undergoes treatment over an extended period, it is essential that the vital signs be repeated as appropriate to the clinical scenario, particularly those that were previously abnormal. In some clinical circumstances, it is advisable to monitor the vital signs continuously.1

Vital signs should be measured and recorded at intervals as dictated by clinical judgment, the patient’s clinical state, or after any significant change in these parameters. Adhering to strict protocols or disease categories is not useful or productive. An abnormal vital sign may constitute the patient’s entire complaint, as in a febrile infant, or it may be the only indication of the potential for serious illness, as in a patient with resting tachycardia.2

Emergency medical service (EMS) personnel begin assessment of the patient’s status and vital signs in the prehospital setting. Surges of epinephrine and norepinephrine commonly occur during transport by the EMS, and these catecholamines are known to alter vital signs and lead to increases greater than 10% in the heart rate.3 Vagal influences may also influence EMS-derived vital signs. Prehospital vital signs should always be interpreted with the entire clinical scenario in perspective.

Blood pressure and pulse are frequently evaluated together as a measure of blood volume. Capillary refill is discussed as an assessment of overall perfusion, circulatory volume, and blood pressure. Although body temperature is usually the last vital sign measured during resuscitation, it has special importance for patients suffering from thermal regulatory failure. With these considerations in mind, the current chapter is organized according to the priorities of patient resuscitation and evaluation. Assessment of pain as a vital sign is gaining acceptance and is discussed briefly at the end of this chapter.

NORMAL VALUES

The range of normal resting vital signs for specific age groups must be appreciated by the clinician to enable identification of abnormal values and their clinical significance. The normal ranges for vital signs are also influenced by gender, race, pregnancy, and residence in an industrialized nation. These ranges have not been validated in ED patients, who have many reasons for abnormalities in vital signs, including anxiety, pain, and altered physiology as a result of their disease states. Ranges of normal vital signs, commonly quoted as normal or abnormal in other settings, serve only as a guide and not an absolute criterion for diagnosis, treatment, further observation, or intervention in the ED.

Published vital sign norms for children are not as well accepted as those for adult patients. Table 1.1 and Table 1.2 report heart rate and RRs by age grouping and percentile for children from birth to 18 years of age. This data represents a large cross-sectional study using 6 months of nurse-documented heart rates and RRs from the electronic records of 14,014 children on general medical and surgical wards at two tertiary-care children’s hospitals. Up to 54% of heart rate observations and up to 40% of RR observations in this sample were outside textbook reference ranges.4 During the newborn period, normal arterial blood pressure rises rapidly. Values for pulse and respiration in children older than 3 years reflect an average of male and female values for 0- to 1-, 3-, 9-, and 16-year-old populations. The values for blood pressure reflect an average of male and female values for the 1- to 6-month-old and the 3-, 9-, and 16-year-old populations.5 Other studies have assessed the reference values for RR in children demonstrating the same variability of pediatric “normal” vital sign ranges.6–12
BACKGROUND

Early recognition of vital signs dates to the fourth century BC, when Herophilus first described sphygmology, or palpation of the pulse in terms of size, frequency, force, and rhythm. Chinese clinicians (second century BC) timed the pulse by the RR of the examiner in the belief that four pulsations per respiration was normal for adults. The study of pulses was greatly influenced by Galen, who expanded the subject into a rather complex and obscure art form and wrote 18 books on the subject.4

Blood pressure was first measured directly in 1733 by Hales, who recorded arterial pressure in a mare by cannulation with a brass pipe and a blood-filled glass column.5 Frank used large-bore catheters connected to a rubber membrane in a 1903 manometer.6 The invention of inflatable cuff manometers (Riva-Rocci, 1896) and discovery of the arterial phase sounds (Korotkoff, 1905) allowed the development of indirect measurement of blood pressure.5,6

Clinical thermometry was introduced by Sanctorius in 1625, with early thermometers being filled with alcohol. Mercury column thermometers were introduced by Fahrenheit in 1714. Although their routine use was supported by Boerhaave, thermometry was not established as routine clinical practice until the 1870s.7
For the adult population, normal blood pressure values have been better established. Although systolic blood pressure increases with age, normotensive or normal systolic blood pressure is defined as 90 to 140 mm Hg, and normotensive or normal diastolic blood pressure is defined as 60 to 90 mm Hg. The recent literature suggests defining an “optimal” blood pressure as 115/75 mm Hg because values at or below this level have been associated with minimal vascular mortality.13 It has been suggested that the definition of hypertension be further expanded to integrate a global cardiovascular risk assessment.14,15 Although most patients have similar blood pressure in both arms, Pesola and colleagues found that 18% of their hypertensive population16 and 15% of their normotensive population had a difference of greater than 10 mm Hg in systolic blood pressure between arms.17 Within the adult population, optimal definitions for normal systolic blood pressure probably vary with age, and particular differentiation should be made in regard to geriatric patients in the emergency setting. The recent literature suggests redefining values representative of hypotension in the elderly, especially in the setting of trauma. Systolic blood pressure readings ranging from approximately 90 to 120 mm Hg have been associated with occult hypoperfusion and increased mortality in geriatric trauma patients.18-20

In 1928 the New York Heart Association, by consensus, established the normal limits for the resting heart rate as 60 beats/min and 100 beats/min. More recent data indicate that 45 beats/min and 95 beats/min may better define the heart rate limits of normal sinus rhythm in adults of all ages. Spodick and colleagues recommended that the operational definition for the limits of the resting heart rate in adults should be 50 beats/min and 90 beats/min.21,22 This view is widely supported among cardiologists,23-25 but these ranges have not been validated in the ED setting. There is currently no consensus on what constitutes a normal adult RR; however, an RR range of 12 to 24 breaths/min is generally accepted in the existing literature as the norm for adults.26-27

Pregnancy results in alterations in the normal adult values of pulse and blood pressure. Pregnancy is characterized by significant increases in minute ventilation, and is thought to be due to the combined facilitatory effects of progesterone and estrogen on central and peripheral chemoreflex drives to breathe.28 The resting pulse rate increases throughout pregnancy from 10% to 15% over baseline values. The norms for systolic and diastolic blood pressure are dependent on patient positioning. When a pregnant patient is sitting or standing, systolic and diastolic blood pressure are dependent on patient position. When a pregnant patient is sitting or standing, systolic pressure is essentially unchanged. Diastolic pressure declines until approximately 28 weeks’ gestation, at which time it begins to rise to nonpregnant levels. When a pregnant patient is in the lateral decubitus position, both systolic and diastolic pressure decline until the 28th week and then begin to rise to nonpregnant levels (Table 1.3).29

TABLE 1.1 Pediatric Heart Rate Limits Based on Age Group and Percentile

<table>
<thead>
<tr>
<th>AGE GROUP</th>
<th>5TH PERCENTILE</th>
<th>50TH PERCENTILE</th>
<th>95TH PERCENTILE</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–3 months</td>
<td>113</td>
<td>140</td>
<td>171</td>
</tr>
<tr>
<td>3–6 months</td>
<td>108</td>
<td>135</td>
<td>167</td>
</tr>
<tr>
<td>6–9 months</td>
<td>104</td>
<td>131</td>
<td>163</td>
</tr>
<tr>
<td>9–12 months</td>
<td>101</td>
<td>128</td>
<td>160</td>
</tr>
<tr>
<td>12–18 months</td>
<td>97</td>
<td>124</td>
<td>157</td>
</tr>
<tr>
<td>18–24 months</td>
<td>92</td>
<td>120</td>
<td>154</td>
</tr>
<tr>
<td>2–3 years</td>
<td>87</td>
<td>115</td>
<td>150</td>
</tr>
<tr>
<td>3–4 years</td>
<td>82</td>
<td>111</td>
<td>146</td>
</tr>
<tr>
<td>4–6 years</td>
<td>77</td>
<td>106</td>
<td>142</td>
</tr>
<tr>
<td>6–8 years</td>
<td>71</td>
<td>100</td>
<td>137</td>
</tr>
<tr>
<td>8–12 years</td>
<td>66</td>
<td>94</td>
<td>129</td>
</tr>
<tr>
<td>12–15 years</td>
<td>61</td>
<td>87</td>
<td>121</td>
</tr>
<tr>
<td>15–18 years</td>
<td>57</td>
<td>82</td>
<td>115</td>
</tr>
</tbody>
</table>


TABLE 1.2 Pediatric Respiratory Rate Limits Based on Age Group and Percentile

<table>
<thead>
<tr>
<th>AGE GROUP</th>
<th>5TH PERCENTILE</th>
<th>50TH PERCENTILE</th>
<th>95TH PERCENTILE</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–3 months</td>
<td>27</td>
<td>41</td>
<td>62</td>
</tr>
<tr>
<td>3–6 months</td>
<td>25</td>
<td>38</td>
<td>58</td>
</tr>
<tr>
<td>6–9 months</td>
<td>23</td>
<td>35</td>
<td>54</td>
</tr>
<tr>
<td>9–12 months</td>
<td>22</td>
<td>33</td>
<td>51</td>
</tr>
<tr>
<td>12–18 months</td>
<td>21</td>
<td>31</td>
<td>48</td>
</tr>
<tr>
<td>18–24 months</td>
<td>20</td>
<td>29</td>
<td>45</td>
</tr>
<tr>
<td>2–3 years</td>
<td>18</td>
<td>27</td>
<td>42</td>
</tr>
<tr>
<td>3–4 years</td>
<td>18</td>
<td>25</td>
<td>40</td>
</tr>
<tr>
<td>4–6 years</td>
<td>17</td>
<td>24</td>
<td>37</td>
</tr>
<tr>
<td>6–8 years</td>
<td>16</td>
<td>23</td>
<td>35</td>
</tr>
<tr>
<td>8–12 years</td>
<td>15</td>
<td>21</td>
<td>31</td>
</tr>
<tr>
<td>12–15 years</td>
<td>13</td>
<td>19</td>
<td>28</td>
</tr>
<tr>
<td>15–18 years</td>
<td>13</td>
<td>18</td>
<td>26</td>
</tr>
</tbody>
</table>


RESPIRATION

Breathing is initiated and primarily controlled in the medullary respiratory center of the brainstem. The respiratory center is modulated by the pneumotaxic center, which limits the length of the inspiratory signal and greatly influences the RR and tidal volume of respiration. An abnormality in respiration may be
a primary complaint or a manifestation of other systemic diseases. Increased RRs may be seen in patients with a variety of pulmonary or cardiac diseases, and acidosis, anemia, temperature, stress, and drugs (such as stimulants and salicylates) can significantly alter the RR in the absence of cardiopulmonary dysfunction.

### Indications and Contraindications

The only contraindications to careful measurement of RR are the scenarios of respiratory distress, apnea, and upper airway obstruction, which require immediate therapeutic intervention. RR and respiratory effort should be assessed as soon as patient care demands allow.

The respiratory status of both adults and children plays a crucial role in determining the overall assessment of illness. Although it is a sensitive yet nonspecific indicator of respiratory dysfunction, the RR can also predict nonpulmonary morbidity. Several prehospital and hospital-based illness or injury severity scores feature the RR as a cardinal value. A prehospital RR of less than 10 breaths/min or greater than 29 breaths/min is associated with major injury in 73% of children.\(^3\) Using tachypnea alone as a predictor of pulmonary pathology, infants with an RR higher than 60 breaths/min are found to be hypoxic 80% of the time.\(^3\) Pediatric studies have linked abnormal RRs to in-hospital mortality and the level of care required in the ED.\(^3\) In a retrospective study exploring predictors of critical care admission for adult ED patients who were initially triaged as having low to moderate acuity, an abnormal RR at the first nursing assessment increased the odds of critical care admission by a factor of 1.66.\(^3\) An RR higher than 25 breaths/min in prehospital trauma patients was associated with increased mortality.\(^2\) Pre-arrest respiratory insufficiency (RR >36 breaths/min or pulse oximetry <90%) was an independent predictor of mortality (odds ratio [OR], 4.2) in patients with EMS-witnessed cardiac arrest.\(^7\) Although some studies have associated abnormal RRs in adult ED patients with increased mortality,\(^1\) a 2011 large prospective cohort study of adult patients found that an initial abnormal RR on triage in the ED was not an independent predictor of hospital mortality.\(^4\)

### Procedure

To measure RR (inspirations per minute), count the respirations when the patient is unaware that his or her breathing is being observed. Count for a full minute to most accurately determine the RR. The frequency of breathing is less regular than the pulse, and inaccurate measurement is more likely to occur if the count is taken for a shorter interval. It is common to measure respirations over 15 seconds and multiply by 4, but this can significantly alter the true RR per minute. An infant’s RR can easily be determined by observing or palpating the excursion of the chest or the abdominal wall.\(^4\) Infants should be observed for grunting respirations, which are produced by expiration against a partly closed glottis (an attempt to maintain positive airway pressure).

### Interpretation

#### Respiratory Rate

The reproducibility of RR measurements may be limited by significant interobserver variability.\(^5\) Clinicians should recognize this inherent variability and interpret the RR with caution. Rates obtained by nurses versus medical students varied significantly, as did those obtained by medical students versus residents and attending clinicians.\(^4\) Interobserver variability may account for a difference of up to 6 breaths/min, and variability in the same observer may account for up to 5 breaths/min.\(^3\) A study comparing RRs obtained by triage nurses with an electronic monitor found that neither provided an accurate measurement of the RR in the ED, suggesting that new clinical strategies for obtaining this vital sign may be necessary.\(^4\) RR is an independent risk marker for in-hospital mortality in community-acquired pneumonia and should be measured when patients are admitted to the hospital with pneumonia and other acute conditions.\(^6\)

Current texts vary considerably in their definition of a normal RR and cite published values that range from 8 to 24 breaths/min. In a study that specifically investigated normal RRs in an ED (afebrile ambulatory patients without respiratory complaints), females had a mean RR of 20.9 breaths/min and males had a mean RR of 19.4 breaths/min. The researchers concluded that a normal RR in the adult patient population was 16 to 24 breaths/min.\(^4\) Other studies have provided additional information on normal resting and sleeping RRs in children younger than 7 years.\(^3\) RRs obtained with a stethoscope were higher than those obtained by observation (mean difference, 2.6 breaths/min in awake and 1.8 breaths/min in asleep children). Smoothed percentile curves demonstrated a larger dispersion at birth (5th percentile, 34 breaths/min; 95th percentile, 68 breaths/min), whereas dispersion was less at 36 months of age (5th percentile, 18 breaths/min; 95th percentile, 30 breaths/min).

The RR will generally increase in the presence of fever. It is often difficult to determine whether tachypnea is a primary finding or is simply associated with hyperpyrexia. A study of children younger than 2 years in whom pneumonia was subsequently diagnosed found that age-appropriate limits for resting tachypnea in the presence of fever could be defined. A sensitivity of 74% and specificity of 77% for pneumonia were found when children 6 months of age had an RR higher than 59 breaths/min, when those aged 6 to 11 months had an RR higher than 52 breaths/min, and when those 1 to 2 years old had an RR higher than 42 breaths/min.\(^4\) Even in the face of physiologic compensation for fever, interpretation of the RR alone can help predict the presence of pulmonary disease.

#### Respiratory Pattern and Amplitude

Hyperventilation and hypoventilation can result from an extensive variety of disorders and may be related to pulmonary
Observe the respiratory patterns carefully in children. In infants, it is essential to distinguish normal periodic breathing from apnea. By definition, periodic breathing consists of three or more respiratory pauses longer than 3 seconds in duration with less than 20 seconds between pauses. There is no associated bradycardia or cyanosis. This contrasts with apnea and is a particular problem in preterm infants. Apnea is defined as a respiratory pause longer than 20 seconds. It may be associated with bradycardia and hypoxia. Periodic breathing and apnea are believed to be disorders on a continuum, both stemming from abnormal physiologic control of respiration. Periodic breathing is considered a benign disorder. Infants with symptomatic apneic episodes that result in apparent life-threatening events are thought to be at increased risk for sudden infant death syndrome.

Recognition of subtle tachypnea can be difficult in the emergency setting, although it can be the solitary indicator of disease. Another instance of pathology that can confuse routine measurement of the RR is diaphragmatic breathing or retractions. The variability in counting respiratory effort versus effective respirations is not generally appreciated in a single recorded value.

or extrapulmonary pathology. Abnormal respiratory patterns can be characteristic of metabolic or central nervous system pathologic conditions (Fig. 1.1) and may aid in the differential diagnosis. Kussmaul respirations describe the hyperventilation pattern seen in diabetic patients with ketoacidosis. Decreased RR is commonly seen with opiate toxicity. Hyperpnea, or a normal RR but clinically significant hyperventilation secondary to increased tidal volume, may be seen with salicylate poisoning.

Figure 1.1 Abnormal respiratory patterns. CVA, Cerebrovascular accident; RR, respiratory rate; VT, tidal volume. (Modified from D’Urbano J: Breathing Patterns, Breath Sounds, 2011. Available at http://www.BreathSounds.org.)
PULSE

Examine the pulse to establish the cardiac rate and regularity of the rhythm. Though rarely diagnostic, peripheral pulses may yield clues about cardiac disease, such as aortic insufficiency, and information about the integrity of the peripheral vascular supply. Doppler ultrasound has utility in locating a pulse, assessing fetal heart tones beyond the first trimester of pregnancy, evaluating peripheral lower extremity vascular insufficiency with an ankle-brachial index, and assessing blood pressure in infants or in patients with low-flow states.

Physiology

Blood flowing into the aorta with each cardiac cycle initiates a pressure wave. Blood flows through the vasculature at approximately 0.5 m/sec, but pressure waves in the aorta move at 3 to 5 m/sec. Therefore palpated peripheral pulses represent pressure waves, not blood flow.

Indications and Contraindications

Assessment of blood flow by palpation of the pulse can be used to gauge the presence of cardiac contractility and not just the electrical rhythm. Caution should be taken to not overgeneralize the presence or strength of a pulse when predicting blood pressure. The necessity for repeated pulse evaluations is dictated by the clinical complaint and the status of the patient. Continuous monitoring is not routine but may be helpful when the clinical situation predicts significant variability in heart rate, as in the setting of sepsis. An association between absence of a radial pulse or absence of both radial and femoral pulses and hypotension has been demonstrated in hypovolemic trauma patients. The variability in individual response prohibits the use of this parameter as an absolute gauge of blood pressure.

No contraindications exist to assessment of the pulse rate. Keep in mind a few cautionary notes about examination of the carotid pulse. Avoid concurrent bilateral carotid artery palpation because this maneuver could theoretically endanger cerebral blood flow. Massage of the carotid sinus, found at the bifurcation of the external and internal carotid arteries at the level of the mandibular angle, may result in reflex slowing of the heart rate. To avoid inadvertent carotid sinus massage, palpate the carotid pulse at or below the level of the thyroid cartilage. In adults with atherosclerotic disease, there is a rare risk of precipitating a cerebrovascular event by vigorous palpation of the carotid artery. Minimize this risk by prior auscultation of the carotid artery. If a bruit is present, gently palpate the carotid pulse while avoiding vigorous palpation, or use a Doppler ultrasound probe to assess carotid flow instead.

Procedure

Depending on the clinical scenario, pulses are palpable at numerous sites, although for convenience the radial pulse at the wrist is routinely used. Use the tips of the first and second fingers to palpate the pulse. The two advantages of this technique are that (1) the fingertips are quite sensitive, thereby enabling the pulse to be located easily and counted, and (2) the examiner’s own pulse may be erroneously counted if the thumb is used. Pulses are easily palpated at the carotid, brachial, femoral, posterior tibial, and dorsalis pedis arteries. Palpate the pulse at the brachial artery to appreciate its contour and amplitude. Locate the pulse at the medial aspect of the elbow and note that it is more easily palpated when the elbow is held slightly flexed. Determine the pulse rate by counting for 1 minute, particularly if any abnormality is present. Common convention in the acute care setting is to count a regular pulse for 15 seconds and multiply the resulting number by 4 to determine the beats per minute.

In neonates, use direct heart auscultation and umbilical palpation as the methods of choice to determine the heart rate. Instantaneous changes in newborn heart rates are best indicated to the resuscitation team by the clinician tapping out each heartbeat. In unstable children, palpate the central arteries, particularly the femoral and brachial pulses, instead of the more peripheral arteries. In a comparison of four methods of determining the heart rate in infants, listening at the apex of the heart was found to be more accurate than palpation of the brachial, carotid, or femoral pulses. Of the sites for palpation of the heart rate, the femoral artery has proved most valuable, especially in hypotensive infants.

Interpretation

Pulse Rate

Consider the individual’s physiology when interpreting the pulse. In infants and children, interpret the pulse rate with reference to age (see Table 1.1). Pulse varies with respiration: it increases with inspiration and slows with expiration. This is known as sinus dysrhythmia and is physiologic.

Although bradycardia is most commonly defined as a heart rate lower than 60 beats/min in adults, a well-conditioned athlete may have a normal resting heart rate of 30 to 40 beats/min. As discussed earlier, a redefinition of bradycardia to less than 45 beats/min and tachycardia to greater than 95 beats/min has been proposed based on a normal healthy population. Such definitions include 95% of the population and do not address any given individual’s normal baseline rate.

Consider whether a patient’s abnormal pulse rate is a primary or secondary condition. Examine the entire set of vital signs when attempting to discern the cause of the abnormal rate. For example, hyperthermia causes sinus tachycardia. Drug fever, typhoid fever, and central neurogenic fever are considerations when no corresponding tachycardia is found in a patient with elevated body temperature. Hypothermia, with its reduced metabolic demands, may be associated with bradycardia. Some disease states are defined by their effect on heart rate, such as thyrotoxicosis with tachycardia or myxedema coma with bradycardia.

Consider the medications that the patient may be taking or the presence of a mechanical pacemaker. Digitalis compounds, β-blockers, and antidysrhythmics may alter the normal heart rate and the ability of this vital sign to respond to a new physiologic stress. These cardioactive medications may cause the abnormality in the patient’s heart rate. Nonprescription drugs can be equally significant in their effect on heart rate. Sympathomimetic drugs such as cocaine and methamphetamine increase heart rate, as do anticholinergic drugs.

Heart Rhythm

In addition to determining the pulse rate, obtain information about the regularity of the pulse by palpation. An irregular pulse suggests atrial fibrillation or flutter with variable block, and accurate assessment of the pulse should be carried out by
auscultation of the apical cardiac sounds. The apical pulse is frequently greater than the peripheral pulse because of inadequate filling time and stroke volume, with resultant nontransmitted beats. A greater pulse deficit generally reflects more severe disease.60

Pulse Amplitude and Contour
Accurate examination and description of pulse amplitude and contour can provide additional clinical information and aid in decision making. Superimposition of one pathophysiologic state on another may modify the pulse. For example, sepsis may result in variable pulse amplitudes, depending on the stage in the development of the disease. Early in sepsis, cardiac output increases and vascular resistance decreases, causing bounding pulses. In advanced sepsis or septic shock, falling cardiac output and increased vascular resistance are seen, and pulses are diminished.61 Definable age-related changes in pulse amplitude and contour can be identified. Such changes are due to an increase in arterial stiffness, resulting in increased pulse wave velocity and progressively earlier wave reflection. This leads to increased pulse amplitude in the elderly at all commonly measured sites (carotid, femoral, and radial).61 In addition to these age-related changes, pulse wave analysis may be useful in determining arterial stiffness and the likelihood of atherosclerotic disease in a vascular laboratory setting.62 If present globally, weak pulses can be a significant finding in hypotensive patients, or an indication of limb ischemia if isolated to one extremity. Bounding pulses can be seen with a widened pulse pressure and are discussed later in the section on blood pressure. Routine measurement of pulse amplitude is not reproducible by simple palpation and requires instrumentation not available in EDs.

Pulses During Cardiopulmonary Resuscitation
Palpated femoral pulses during chest compression may represent either forward arterial blood flow or “to-and-fro” movement of blood from the right side of the heart to the venous system. A carotid pulse is preferred when assessing the adequacy of chest compressions during cardiopulmonary resuscitation (see Chapter 17).

ARTERIAL BLOOD PRESSURE
Systolic blood pressure changes with each heartbeat. Changes in arterial blood pressure over time may indicate success of treatment or worsening of the patient’s overall condition. An abrupt reduction in a patient’s arterial blood pressure usually indicates the need for immediate intervention or reconsideration of therapy. The current section discusses indirect blood pressure monitoring; intraarterial techniques are considered elsewhere. Discussion of the specific use of the Doppler device for measurement of pulse and blood pressure and for measurement of orthostatic blood pressure and changes in pulse follow this section. Despite an association between the absence of hypotension and the presence of a radial pulse or between hypotension and the absence of both radial and femoral pulses in the setting of trauma, the variability in individual responses prohibits the use of this parameter as an absolute gauge of blood pressure.45

Physiology
Arterial blood pressure indicates the overall state of hemodynamic interaction between cardiac output and peripheral vascular resistance. Arterial blood pressure is the lateral pressure or force exerted by blood on the vessel wall. It indirectly measures perfusion, and blood flow equals the change in pressure divided by resistance. Because peripheral vascular resistance varies, a normal blood pressure does not confirm adequate perfusion.63 Mean arterial blood pressure (MAP) can be estimated by adding one third of the pulse pressure (i.e., the difference between systolic and diastolic blood pressure) to diastolic pressure or by using the following measure:63,65

\[
\text{MAP} = \frac{\text{Diastolic pressure} \times 2 + \text{Systolic pressure}}{3}
\]

Many modern bedside telemetry monitors automatically incorporate MAP measurements into the blood pressure systolic and diastolic pressure readings.

Indications and Contraindications
Patients with minor ambulatory complaints unrelated to the cardiovascular system may not necessarily need their blood pressure measured in the ED, and those with hemodynamic instability need frequent monitoring of blood pressure. In children, there is a significant amount of variability regarding standard situations that require measurement of blood pressure. In general, the younger the patient, the less likely blood pressure will be measured.66,67 In newborns, infants, and even toddlers, capillary refill is sometimes substituted for standard blood pressure measurement, although viewing these tests as equivalent can lead to significant errors.

In low-flow states, Doppler measurement of blood pressure may be obtained rapidly. Repeated measurements will provide an evaluation of the adequacy of resuscitation in patients whose blood pressure cannot be auscultated by standard techniques and in those in whom intraarterial blood pressure measurements are either contraindicated or technically unobtainable.68,69 Placing a catheter for direct intraarterial measurement of blood pressure may be performed safely in the ED, but is not standard of care and has a higher risk for complications. In particular, direct measurement of arterial pressure during pulseless electrical rhythms may help discriminate between a severe shock state and otherwise nonresuscitable status.70,71 Alternative noninvasive devices for continuous blood pressure measurement (CBPM) have been introduced clinically, with varying success. One common method of CBPM uses finger cuffs equipped with infrared (IR) photoplethysmography and sophisticated technology for quantification of finger blood pressure levels. Finapres (Ohmeda, Madison, WI) was the first commercial product using this technique, and several newer products are on the market today. A number of commercial systems use an alternative method of arterial applanation tonometry to measure CBPM. Further study is needed for the validation of devices using these techniques.72,73 Relative contraindications to specific extremity blood pressure measurement include an arteriovenous fistula, ipsilateral mastectomy, axillary lymphadenopathy, lymphedema, and circumferential burns over the intended site of cuff application.

Equipment
Two types of blood pressure monitoring equipment are currently available and used in EDs: cuff type and noninvasive waveform analysis.
Cuff Type
The equipment required for indirect blood pressure measurement includes a sphygmomanometer (cuff with an inflatable bladder, inflating bulb, controlled exhaust for deflation, and manometer) and a stethoscope, Doppler device (for auscultation), or oscillometric device.74-76 A common practice in the prehospital and interhospital transport setting is to forego auscultatory blood pressure measurements with a stethoscope and instead obtain systolic values only by palpation of the first Korotkoff sound. This practice, though sometimes the only feasible method of obtaining any value in a noisy environment, poses a significant potential for error.

According to the American Heart Association guidelines, the sphygmomanometer cuff should be an appropriate size for the patient to ensure an accurate reading. The width of the bladder should be at least 40% of the distance of the limb’s midpoint (i.e., from the acromion process to the lateral epicondyle). This published figure of the ideal width, when studied in a validation review, may be higher, up to approximately 50%.77 The length of the bladder should be 80% of the midarm circumference or twice the recommended width.88 Discrepancies in matching upper arm size with cuff size have been demonstrated to produce significant errors in critically ill populations when compared with invasive intraarterial blood pressure measurements.78 The availability of appropriately sized cuffs appears to be a pervasive problem, especially as approximately 80% of patients do not fit the standard 12-cm large cuffs.79 In one study, 90% of aneroid devices had only one size of cuff available.80 A second study phase from this group showed no marked improvement in agreement of oscillatory and invasive measurements, despite using the correct cuff size.81

The manometers in common use are either aneroid, digital, or mercury gravity column, though the mercury type is much less common in modern use. All three types of manometers are convenient for bedside use, although the mercury gravity column must be placed vertically to ensure accurate measurements. An aneroid manometer uses a metal bellows that elongates with the application of pressure. This elongation is mechanically amplified and transmits the motion to the indicator needle.

Manometers require annual servicing. Mercury columns may require the addition of mercury to bring the edge of the meniscus to the zero mark. The air vent or filter at the top of the mercury column should also be checked for clogging. An aneroid manometer should be calibrated against a mercury column at least yearly. If the aneroid indicator is not at zero at rest, the device should not be used.82 Digital manometers may not be validated for all patient groups and could give inaccurate readings.

Automatic sphygmomanometers may improve physiologic monitoring with their alarm and self-cycling capabilities. They offer indirect arterial blood pressure measurement with little pain and without the risks associated with invasive arterial lines.83 Accuracy of measurements does not suffer during rapid cycling, and the potential for vascular injury from nearly continuous arterial compression dictates that most automated blood pressure units will revert back to less frequent (i.e., every 15 to 20 minutes) cycling as a safety precaution. Oscillometric blood pressure monitors detect motion of the blood pressure cuff transmitted from the underlying artery. A sudden increase in the amplitude of arterial oscillations occurs with systolic pressure and MAP, and an abrupt decrease occurs with diastolic pressure.84

There appears to be less variability with the oscillometric blood pressure method than with the auscultatory method in children. These results are not generalizable to the neonatal population, and errors are commonly encountered even when exhaustive measures are taken to control the environment.

In adult patients, numerous studies have focused on the reliability of auscultatory versus automated blood pressure measurements. Mercury column versus Dinamap readings showed increased auscultatory disparity when systolic blood pressure was greater than 140 mm Hg, the range at which accuracy should be most rigorously sought to correctly identify hypertension. In general, automated blood pressure devices yield higher systolic and lower diastolic blood pressure.95 The range of error in automated devices was, on average, 4.0 to 8.6 mm Hg.86,87 Unfortunately, these studies represent populations without critical illness and do not reflect the accuracy of readings at the extremes of hypertension and hypotension, thus making generalization to an ED population difficult.77,88 Like most durable medical equipment, sphygmomanometers are subject to deterioration with use. Automatic sphygmomanometer validation is a form of calibration and deserves quality control and traceability to ensure accurate results.88

Procedure
Obtain indirect blood pressure measurements at the patient’s bedside by palpation, auscultation, Doppler, or oscillometric methods. The technique is straightforward and accurate when the equipment is well maintained, calibrated, and used by clinicians who follow accepted standards. The patient may be lying or sitting, as long as the site of measurement is at the level of the right atrium and the arm is supported.68,76 Unless the arm is kept perpendicular to the body with the elbow resting on a desk, measurements will be 9 to 14 mm Hg higher, regardless of body position.77,90 Allowing the arm to be parallel to the body when supine and supporting the arm perpendicular to the body when measuring blood pressure may create a pseudo-drop in blood pressure. These changes are thought to be dependent on the mechanical properties of the arteries themselves and not associated with hydrostatic pressure alone.70

To palpate arterial blood pressure, inflate the cuff to 30 mm Hg above the level at which the palpable pulse disappears. Once properly inflated, palpate directly over the artery with as little pressure as possible.92 Another operative study looking at the combination of palpated systolic blood pressure and observed visual return of continuous pulse oximetry reported an underestimation of 10 to 20 mm Hg.87 The same technique is used with the Doppler device, and the palpated pulse is replaced with the Doppler auditory signal. Measurement of arterial pressure by palpation and Doppler yields only estimates of systolic blood pressure. The Doppler method is preferred when determining blood pressure in infants.88 When auscultating blood pressure at the brachial artery, apply the blood pressure cuff approximately 2.5 cm above the antecubital fossa with the center of the bladder over the artery. Apply the bell of the stethoscope directly over the brachial artery with as little pressure as possible.92 Systolic arterial blood pressure and...
pressure is defined as the first appearance of faint, clear, tapping sounds that gradually increase in intensity (Korotkoff phase I). Diastolic blood pressure is defined as the point at which the sounds disappear (Korotkoff phase V). In children, phase IV defines diastolic blood pressure (Fig. 1.2). Phase IV is marked by a distinct, abrupt muffling of sound when a soft, blowing quality is heard.

It is best to auscultate over the brachial artery because of accepted standardization of the measured values. Alternative sites include the radial, popliteal, posterior tibial, or dorsalis pedis arteries, although any fully compressible extremity artery may be used. Studies evaluating direct and indirect blood pressure measurements have demonstrated good correlation between these methods.

Occasionally it may not be feasible to obtain proximal upper extremity blood pressure measurements because of patient access issues, particularly those encountered in the prehospital setting. Forearm measurements may be obtained more easily, though correlation to standard proximal upper extremity values has been controversial. An earlier study showed fair correlation to a brachial cuff measurement (within 20 mm Hg in 86% of systolic measurements and 94% of diastolic measurements). A subsequent study demonstrated that forearm measurements cannot routinely replace upper arm measurements to determine blood pressure (the potential variance from an upper arm measurement is ± 19 mm Hg for systolic pressure). Alternatively, noninvasive finger blood pressure measurements have shown promise when compared with standard upper extremity readings. The overall discrepancy in an ED study was 0.1 mm Hg with a standard deviation of ± 5.02 mm Hg when comparing finger blood pressure and invasive MAP via radial artery cannulation.

Novel noninvasive continuous finger cuff technology offers the benefit of uninterrupted monitoring and has the advantage over invasive techniques of being safer and immediately available. Noninvasive finger cuff measurements have shown reasonable correlation, even in critically ill populations in the ED. Wrist blood pressure has been shown to have good average accuracy in the surgical environment when compared with oscillometric devices. Patient comfort is reported to be greater with these devices. The typically stated contraindications to the acquisition of upper arm blood pressure (limitation after mastectomy, etc.) may not apply. The accuracy of the palpatory, Doppler, and oscillometric methods has also been investigated. When phase I and V Korotkoff sounds are used, indirect methods typically underestimate systolic and diastolic pressure by several millimeters of mercury. During shock, the palpatory and auscultatory methods underestimate simultaneous direct arterial pressure measurements. The flush method, in which the return of color after deflation of the cuff is used to estimate blood pressure in infants, may underestimate systolic blood pressure by up to 40 mm Hg. This method is unreliable and not recommended.

Complications

Complications of indirect blood pressure measurements are minimal when the proper procedure is followed. Inadvertent prolonged application of an inflated blood pressure cuff may result in falsely elevated diastolic pressure and ischemia distal to the site of application. Invasive blood pressure monitoring is associated with a number of potential problems (see Chapter 20).

Interpretation

Normal blood pressure increases with decreasing distance from the heart and aorta. Blood pressure tends to increase with age and is generally higher in males. Individual factors that influence blood pressure include body posture, emotional or painful stimuli, environmental influences, vasoactive foods or medications, and the state of muscular and cerebral activity. Exercise and sustained isometric muscular contraction increase blood pressure in proportion to the strength of the contraction. A normal diurnal pattern of blood pressure consists of an increase throughout the day with a significant, rapid decline during early, deep sleep.

Normal lower limits of systolic blood pressure in infants and children can be estimated by adding 2 times the age (in years) to 70 mm Hg. The 50th percentile for a child’s systolic arterial blood pressure from 1 to 10 years of age can be estimated by adding 2 times the age (in years) to 90 mm Hg. Children older than 2 years are considered hypotensive when systolic blood pressure is less than 80 mm Hg. Children are able to maintain MAP until very late during shock. The finding of a normal blood pressure in a child with signs of poor perfusion should not dissuade the clinician from appropriate treatment. Most adults are considered hypotensive if systolic blood pressure is lower than 90 mm Hg, but some individuals normally exhibit a systolic pressure in that range. In the elderly, the presence of normotension within defined or published limits may not be reassuring. When considering systolic blood pressure cutoffs.
for trauma patients, 85 mm Hg for patients aged 18 to 35 years, 96 mm Hg for patients aged 36 to 64 years, and 117 mm Hg for those older than 65 years have been proposed as new standards for hypotension.\textsuperscript{10} When accompanied by signs of shock, immediate treatment is indicated. In patients with shock, blood flow cannot be reliably inferred from heart rate and blood pressure values.\textsuperscript{109,110}

**Hypertension**

Adults are hypertensive if either systolic or diastolic pressure consistently exceeds 140 or 90 mm Hg, respectively.\textsuperscript{111} A meta-analysis showed strong correlation of blood pressure to vascular (and overall) mortality down to at least 115/75 mm Hg.\textsuperscript{112} Some authors have suggested altering the blood pressure definitions to include an “optimal” blood pressure of 115/75 mm Hg.\textsuperscript{113} Other authors have suggested incorporating blood pressure into a global cardiovascular risk assessment that includes other associated risk factors.\textsuperscript{14,15} The applicability of population norms for hypertension in a stressful emergency situation is controversial. One should not make diagnostic or therapeutic decisions based solely on an abnormal initial measurement. Patients with hypertension require repeated measurements to assess whether therapy is required in the ED. Because sustained hypertension may be seen in more than one third of initially hypertensive ED patients, careful evaluation and follow-up are required.\textsuperscript{114} Unfortunately, elevated blood pressure readings are still almost uniformly ignored or unrecognized in the emergency setting, particularly in children.\textsuperscript{115} The phenomenon of *white coat hypertension* (WCH) is defined as a persistent elevation in blood pressure in the clinical setting only. The prevalence of WCH is between 20% and 94%, depending on the frequency of reassessment in the clinical setting.\textsuperscript{116} It is unclear whether patients who have isolated hypertension in the clinical setting (WCH) are at increased risk for the development of hypertension and subsequent end-organ damage.\textsuperscript{117}

**Measurement Errors**

Erroneous blood pressure measurements may result from several factors.\textsuperscript{118} Falsely low blood pressure may be caused by using an overly wide cuff, by placing excessive pressure on the head of the stethoscope, or by rapid cuff deflation.\textsuperscript{119,120} Falsely high blood pressure may be caused by the use of an overly narrow cuff, anxiety, pain, tobacco use, exertion, an unsupported arm, or slow inflation of the cuff.\textsuperscript{121} There appears to be a statistically significant difference in the error rate associated with patients weighing more than 95 kg, whether from obesity or as a result of muscular upper arms from body building.\textsuperscript{122,123}

Of note, 41% of adults observed at the University of Pittsburgh required non–standard-sized cuffs, and the use of small cuffs was associated with a mean error of 8.5 and 4.6 mm Hg in systolic and diastolic pressure, respectively.\textsuperscript{120} Other studies have confirmed relatively high rates of inappropriately diagnosed hypertension in obese patients based on erroneous cuff size.\textsuperscript{122} Other specific study populations in this area have been critically ill patients, in whom disparate cuff size can lead to significant inaccuracies based on arm circumference.\textsuperscript{124}

Hypotensive patients have unreliable Korotkoff sounds, but Doppler measurements are well correlated with direct arterial systolic pressure measurements in these patients.\textsuperscript{125} An auscultatory gap can be appreciated in hypertensive patients and may mislead the clinician. It is heard during the latter part of phase I and should not be confused with diastolic readings. Auscultation until the manometer reading approaches zero should prevent misinterpretation. In patients with aortic insufficiency or hyperthyroidism, in those who have just finished exercising, and in children younger than 5 years, measurement of diastolic blood pressure should occur at Korotkoff phase IV. Extremes of blood pressure, both hypotension and hypertension, have been found to be factors contributing to measurement errors in critically ill pediatric patients. Predictably, falsely high readings for noninvasive versus invasive measurements have been obtained in hypotensive patients and falsely low values in hypertensive states.\textsuperscript{125}

Irregular heart rates may also interfere with accurate determination of blood pressure. Take a second or third reading, with 2 minutes of deflation between recordings, and obtain an average when premature contractions or atrial fibrillation are present.

Hemiplegic patients may exhibit different blood pressures in the affected and unaffected arms.\textsuperscript{126} A flaccid extremity tends to yield lower systolic and diastolic pressure, whereas a spastic extremity tends to yield higher values than the extremity with normal motor tone. Although these differences are generally small, it is preferable to monitor blood pressure in the unaffected limb.

Numerous errors may occur in the accurate measurement of blood pressure. The only way to combat these errors is to first be cognizant of practices contributing to them. Unfortunately, few nurses can identify causes of potentially erroneous readings. In a study examining nurses’ ability to obtain accurate readings, proper technique in determining systolic blood pressure could be identified 61% of the time; diastolic blood pressure, 71% of the time; and an auscultatory gap, 54% of the time. Nurses were able to correctly identify faulty equipment 58% of the time, assess cuff size 57% of the time, determine appropriate inflation pressure 29% of the time, note the appropriate deflation rate 62% of the time, and determine correct arm positioning 14% of the time.\textsuperscript{127}

There is an increasing number of patients with heart failure, those receiving bridging measures to transplantation, or those treated with long-term circulatory augmentation devices in the form of left ventricular assist devices (LVADs); therefore it is useful to understand the difficulty in interpreting blood pressure measurements in these patients. All types of VADs fit into two categories: (1) pulsatile and (2) nonpulsatile. Pulse and blood pressure readings in patients with pulsatile VADs (Thoratec, HeartMate XVE, Novacor, C-Pulse) are comparable to values in the general non-VAD population. Nonpulsatile VADs (HeartAssist 5, Incor and Excor, Jarvik 2000, VentriAssist, MTTHeart LVAD, HVAD/MVAD, DuraHeart, DeBakey LVAD, HeartMateII/III) function by either centrifugal or axial blood flow, and this has a significant impact on the ability to detect pulses.\textsuperscript{128} Typically, these patients appear to be well-perfused with adequate skin warmth and capillary refill even though pulses may be absent. Blood pressure readings can be obtained with these nonpulsatile flow devices, and diastolic blood pressure, pulse pressure values, and MAP vary significantly depending on the speed of the pump.\textsuperscript{129}

**Pulse Pressure**

The difference between systolic and diastolic pressure is termed pulse pressure. For example, if the blood pressure is 120/80 mm Hg, the pulse pressure is 40 mm Hg. Increased pulse pressure (i.e., ≥60 mm Hg) is commonly observed with anemia, exercise, hyperthyroidism, arteriovenous fistula, aortic
regurgitation, increased intracranial pressure, and patent ductus arteriosus. A narrowed pulse pressure (≤20 mm Hg) may be a manifestation of hypovolemia, increased peripheral vascular resistance (as seen in early septic shock), or decreased stroke volume. A narrowed pulse pressure is classically noted in aortic stenosis and pericardial tamponade. Traditional vital sign measurements, such as systolic blood pressure and oxygen saturation, often fail to predict mortality or indicate the need for life-saving interventions or reductions in central blood volume until after the onset of cardiovascular collapse. There is evidence that an early indicator of reduced central blood volume in the presence of stable vital signs is the reduction in pulse pressure.130

**Differential Brachial Artery Pressure**

The presence of a systolic blood pressure difference of 10 to 20 mm Hg between the arms suggests a normal condition. If greater, it may indicate advanced atherosclerosis, coarctation of the aorta proximal to the left subclavian artery, type A aortic dissection, aortic arch syndromes, or other vascular processes preferentially affecting one extremity. The utility of upper extremity bilateral blood pressure measurements has recently come into question. One study found a 10-mm Hg systolic or diastolic difference in 53% of patients in the emergency setting and a 20-mm Hg or higher difference in 19% of patients.131 Although these numbers have not generally been found to be of this magnitude in metaanalyses,132 the unique setting of the study in the ED makes correlation particularly salient for the emergency physician. The reliability of peripheral pulse deficits in diagnosing or excluding type A aortic dissection is a frequently cited reason for evaluating blood pressure in both arms. In a metaanalysis, Teece and Hogg noted that the successful use of the absence of a clinical pulse deficit to exclude thoracic dissection in patients with chest pain was just 31%, and the authors concluded that peripheral pulse deficits are far too insensitive to warrant their use as a means of excluding thoracic aortic dissection in patients with chest pain.133 Given that many in the general population have significant differences in blood pressure in each arm, the diagnostic value of this frequently cited indication for obtaining bilateral brachial blood pressure is unproven. Essentially, most patients with a type A aortic dissection will not have a measurable blood pressure discrepancy between the arms, and most of those who do have such a finding will not have dissection.

Brachial pressure differences did not appear to be linked to age, gender, race, MAP, cardiovascular risk, or final discharge diagnosis. Smaller interarm differences have been reported in the ED setting (18% in hypertensive patients and 15% in normotensive patients when a cutoff greater than 10 mm Hg was used).16,17 Though not tested, a method proposed to minimize these differences in the ED is to take simultaneous blood pressure readings from both the left and right extremities with two calibrated automated blood pressure units.134

**Pulse-Pressure Variation**

Fluid resuscitation is an integral piece of the management of patients with circulatory failure, though administration of the proper amount and rate of parenteral fluids can be challenging. Fluid responsiveness, or the ability of the left ventricle to increase stroke volume in response to fluid administration, is an emerging concept that helps to address this challenge.135 This concept is based on the physiology of the Frank–Starling curve and the knowledge that pulse pressure (systolic pressure minus diastolic pressure) is directly proportional to stroke volume. The variation in pulse pressure seen with the respiratory cycle, or pulse-pressure variation, reflects the magnitude of respiratory change on stroke volume. This is best demonstrated by the influence of mechanical ventilation on right ventricular preload. Studies have shown that a pulse-pressure variation of greater than 13% is highly predictive of fluid responsiveness in mechanically ventilated patients.136

Though many methods exist to assess fluid responsiveness, the standard method is done with passive leg raising (PLR). This “self-volume challenge” increases preload through translocation of venous blood from the lower extremities to the thorax. A patient who exhibits a rise of more than 10% in their aortic blood flow (measured with esophageal Doppler) or cardiac index (measured with thermodilution) is considered a “fluid responder,” which is indicative of the need for further fluid administration.137 This concept is becoming increasingly useful in optimizing the fluid management of critically ill patients.

**Pulsus Paradoxus**

Normal respiration briefly decreases systolic blood pressure by approximately 10 mm Hg during inspiration. Pulsus paradoxus occurs when there is a greater than 12-mm Hg decrease in systolic blood pressure during inspiration. Pulsus paradoxus may occur in patients with chronic obstructive pulmonary disease, pneumothorax, severe asthma, or pericardial tamponade.104

To measure a paradoxical pulse, have the patient lie comfortably in the supine position at a 30- to 45-degree angle and breathe normally in an unlabored fashion (which are unusual conditions in a patient suspected of having cardiac tamponade, severe asthma, chronic obstructive pulmonary disease, or pneumothorax).135 Inflate the blood pressure cuff well above systolic pressure and slowly deflate it until the systolic sounds that are synchronous with expiration are first heard (Fig. 1.3). Initially, the arterial pulse will be heard only during expiration and will disappear during inspiration. Deflate the cuff further until arterial sounds are heard throughout the respiratory cycle. Palpation at the radial or femoral arteries may yield complete disappearance during inspiration. When present, this technique is a quick bedside confirmation of the possibility of severe tamponade.

An alternative method for determination of pulsus paradoxus is by visually observing the loss of the pulse oximetry waveform and then its reappearance.138 The plethysmographic method has been validated in intensive care unit settings.

If the difference between inspiratory and expiratory pressure is greater than 12 mm Hg, the paradoxical pulse is abnormally wide.137,138 Most patients with proven tamponade have a difference of 20 to 30 mm Hg or greater during the respiratory cycle.137-139 This may not be true of patients with very narrow pulse pressures (typical of advanced tamponade), who have a “deceptively small” paradoxical pulse of 5 to 15 mm Hg.139-141

Pulsus paradoxus has been correlated with the level of impairment of cardiac output by tamponade. In an uninjured patient with pericardial effusion, a pulsus paradoxus greater than 25 mm Hg (in the absence of relative hypotension) is both sensitive and specific for moderate or severe versus mild tamponade.137,142 An echocardiographic study found that an abnormal pulsus paradoxus had a sensitivity of 79%, a specificity of 40%, a positive predictive value of 81%, and a negative predictive value of 40% for right ventricular diastolic
The shock index (SI) is not standard of care in the ED, but a number of clinical scenarios have been studied in which the SI can be used as a predictor of severe illness or injury. An SI above 0.85 to 0.90 suggests acute illness in medical patients and a marked increase in the potential for gross hemodynamic instability in trauma patients. The SI has been studied for use in a variety of clinical scenarios from severe pneumonia to first-trimester risk for ectopic pregnancy to sepsis. It has been found to be a valid gauge of the severity of illness. Some studies have found that the initial pulse rate alone had nearly the same predictive power as the SI for the severity of illness. Although the SI appears to correlate with the left ventricular stroke work index, it has little correlation with systemic oxygen transport in patients with hemorrhagic and septic shock.

**DOPPLER ULTRASOUND FOR EVALUATION OF PULSE AND BLOOD PRESSURE**

**Principles of Doppler Ultrasound**

Doppler ultrasound is based on the Doppler phenomenon. The frequency of sound waves varies depending on the speed of the sound transmitter in relation to the sound receiver. Doppler devices transmit a sound wave that is reflected by flowing erythrocytes, and the shift in frequency is detected. Frequency shift can be detected only for blood flow greater than 6 cm/sec.

**Indications and Contraindications**

Doppler ultrasound is commonly used in the ED for the measurement of blood pressure in low-flow states, evaluation of lower extremity peripheral perfusion, and assessment of fetal heart sounds after the first trimester of pregnancy. Doppler’s sensitivity allows detection of systolic blood pressure down to 30 mm Hg in the evaluation of a patient in shock. In a patient with peripheral vascular disease in whom there is concern about the adequacy of peripheral perfusion, the ankle-brachial index provides a rapid, reproducible, and standardized assessment. Fetal heart sounds provide a baseline assessment of any pregnant patient with 12 weeks’ gestation or longer in the setting of abdominal trauma or fetal distress as a result of a complication of pregnancy. The use of Doppler ultrasound for the evaluation of deep venous thrombosis is a valuable tool, and specific training and experience are required to attain proficiency. Discussion of this topic is beyond the scope of this chapter.

**Equipment**

A nondirectional Doppler device has a probe that houses two piezoelectric crystals. One crystal transmits the signal and the other receives it. Reflected signals are converted to an electrical signal and fed to an output that transforms them to an audible sound.

Probes with a frequency of 2 to 5 MHz are best for detecting fetal heart sounds. Frequencies of 5 to 10 MHz are appropriate for limb arteries and veins. The probes should be monitored periodically for electrical damage and integrity of the crystals. The sphygmomanometers used in conjunction with the Doppler device should be calibrated periodically, as described in the section on evaluation of blood pressure.

**Procedure**

Place the Doppler probe against the skin with an acoustic gel used as an interface. The gel ensures optimal transmission and
reception of the ultrasound signal and protects the crystals. In an emergency, water-soluble lubricant (e.g., Surgilube or K-Y Jelly) may be substituted for commercial acoustic gel. Angle the probe at 45 degrees along the length of the vessel to optimize frequency shifts and signal amplitude.

To evaluate peripheral perfusion, place a sphygmomanometer cuff proximal to where the arterial pulse is being evaluated and inflate it. Place the probe over the arterial pulse and slowly deflate the cuff. The pressure at which flow is first heard is the systolic pressure under the cuff, and not the pressure at the level of the Doppler probe.

In the evaluation of peripheral vascular disease, one may determine the ankle-brachial index. It is standard for this procedure to be performed in a formal vascular laboratory, and an approximation of pressures can be determined in the ED (Fig. 1.4 and Video 1.1). Usually, only the ankle-brachial index is considered for ED purposes. Examine both brachial arteries at the medial aspect of the antecubital fossa. Angle the probe until the most satisfactory signal is obtained. Inflate the cuff and slowly deflate it until the systolic pulse is heard.

Repeat the procedure for the posterior tibial and dorsalis pedis arteries of both lower extremities. This procedure may be done with oscillometric devices and lacks sensitivity in identifying disease.153

In evaluating fetal heart tones, because of variable positioning of the fetus, an examination of several locations and angles over the uterus must be performed in search for the optimal signal. It is best to begin in the mid-suprapubic area and then explore the uterus via angulation of the probe. Once tones are located, move the probe along the abdomen to reach a position closer to the origin of the sound. Distinguish fetal heart tones from placental flow by differentiating the quality of the fetal heart tones, which will not match the maternal pulse. The placental flow and maternal pulse should be identical.

**Interpretation**

As noted earlier, in low-flow states Doppler ultrasound can detect blood pressure as low as 30 mm Hg. Calculate the ankle-brachial index of each limb by dividing the higher systolic pressure of the posterior tibial or the dorsalis pedis artery of the limb by the higher of the systolic pressures in the brachial arteries. In normal individuals, the index should be greater than 1.0 (Fig. 1.5). Patients with mild to moderate claudication have values between 0.4 and 0.9. Values lower than 0.4 indicate severe impairment and are consistent with critical limb ischemia.154 When the lower extremity has been amputated or injured, brachial-brachial indices can be used (i.e., comparison of systolic blood pressure in the injured or diseased upper extremity with the other extremity). Patients with ankle-brachial index values of 0.9 or lower have increased cardiovascular morbidity and mortality.155 One study of 323 penetrating extremity wounds found that an ankle-brachial index (or brachial-brachial index) lower than 0.9 was 72.5% sensitive and 100% specific for vascular injuries.156 Segmental lower extremity pressure measurements may help identify the level of the obstruction (Table 1.4).157 Obese patients, diabetic patients, or those with calcified vessels that are not compressible may have abnormally high systolic pressure (e.g., 250 to 300 mm Hg) and indices that do not accurately reflect flow. Normal fetal heart tones should be between 120 and 140 beats/min. Fetal heart tones may be heard as early as the 12th week of gestation.

### VITAL SIGN DETERMINATION OF VOLUME STATUS

Many techniques have been advocated to assess volume status. Unfortunately, most procedures lack a database against which to judge their reliability. Recommended methods include evaluation of skin color; skin turgor; skin temperature; supine, serial, and orthostatic vital signs; neck vein status; transcutaneous oximetry; and hemodynamic monitoring (e.g., monitoring of central venous pressure). Serial vital sign measurements have been used for assessing blood loss, but they do not reliably detect small degrees of blood loss.154,158,159 Up to 15% of the total blood volume can be lost with minimal hemodynamic changes or any alteration in supine vital signs.158,160 A decrease in pulse pressure occurs with acute blood loss,161 but the patient’s baseline blood pressure values are often unknown. Clinical examination of neck veins adds useful information and is less precise than measurement of central venous pressure. Most
Vital Signs Measurement

13

negative orthostatic test, and its value for estimating volume status is probably overstated. In patients with an acute loss of less than 20% of total blood volume, orthostatic vital signs have been shown to lack both sensitivity and specificity.161

Physiologic Response to Hypovolemia

Acute blood loss or severe hypovolemia related to dehydration decreases venous return.162 This can be seen with acute blood loss (usually greater than 20% of blood volume), severe burns, or prolonged vomiting or diarrhea that depletes body fluids. As a result, cardiac output falls and clinical manifestations of shock ensue. Several compensatory mechanisms are initiated by acute hypovolemia (Box 1.1). The dominant compensatory mechanism in shock is a reduction in arterial prejoulsion, which greatly increases peripheral vascular resistance; (2) constriction of venous capacitance vessels, which increases venous return to the heart; and (3) an increase in the heart rate and force of contraction, helping maintain cardiac output despite significant loss of volume.160

The value of sympathetic reflex compensation is illustrated by the fact that 30% to 40% of blood volume can be lost before death occurs. When sympathetic reflexes are absent, a loss of only 15% to 20% of blood volume may cause death.160 Increased sympathetic nerve activity results in the commonly recognized physical signs of shock, including pallor, cool clammy

clinicians use skin color, temperature, and moisture as a reflection of skin perfusion and sympathetic tone. This is not an accurate guide to circulatory volume because the vasomotor tone of the skin is affected by numerous diseases, as well as by emotional and environmental factors. Capillary refill has been advocated as a noninvasive test for hypovolemia, and it has been found to be inaccurate in adults (see the following discussion regarding its use in children).162 The ideal test for determining volume status would rapidly and accurately detect 5% or greater depletion of volume with a noninvasive technique. At present, no such test exists.

ORTHOSTATIC VITAL SIGNS MEASUREMENT

Orthostatic vital signs have historically been used to evaluate patients with fluid loss, hemorrhage, syncope, or autonomic dysfunction. They are also used to assess the patient’s response to therapy. The clinician is often concerned with accurate detection of acute blood loss or volume depletion. When the clinical syndrome of shock exists, assessment of a deficit in blood volume poses little difficulty. It is preferable that loss of volume be detected before loss of physiologic compensation and clinical shock occur.

Although orthostatic testing is commonly cited as a method to detect hypovolemia, it is frequently misleading and has less clinical value than often touted. The medical literature is inconsistent regarding values representative of a positive or negative orthostatic test, and its value for estimating volume status is probably overstated. In patients with an acute loss of less than 20% of total blood volume, orthostatic vital signs have been shown to lack both sensitivity and specificity.161

Figure 1.5 A, Typical pressures in a normal individual. Findings, based on resting pressure, show no evidence of occlusive disease of the large- or medium-sized arteries. Normal findings are as follows: (1) ankle-to-brachial pressure index of 1.0 or higher; (2) all segmental pressure gradients lower than 30 mm Hg; and (3) upper thigh pressure at least 40 mm Hg above brachial pressure. *Systolic pressure in mm Hg. B, Typical pressures in a patient with obstruction of the right popliteal or tibial arteries. Significant findings are as follows: (1) ankle-to-brachial pressure index less than 0.9 in the right leg; (2) abnormally high gradient from the ankle to below the knee and again from below to above the knee in the right leg; and (3) upper thigh pressure 50 mm Hg higher than brachial pressure, consistent with normal flow at the aorta-iliac level. Findings are suggestive of right popliteal occlusion or right anterior and posterior tibial occlusion, or both. (From Doppler evaluation of peripheral arterial disease: a clinical handbook, ed 5, Fredericksburg, VA, Sonicaid, Inc. Reproduced with permission.)
stimulates the sympathetic nervous system after arterial pressure loss. This causes arteriolar vasoconstriction, conservation of the release of angiotensin and antidiuretic hormone (vasopressin). Ankle-brachial index.

<table>
<thead>
<tr>
<th>TABLE 1.4 Use of Segmental Lower Extremity Pressure to Identify the Level of Obstruction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td>Normal</td>
</tr>
<tr>
<td>Right</td>
</tr>
<tr>
<td>Left</td>
</tr>
<tr>
<td>Arm</td>
</tr>
<tr>
<td>ABI</td>
</tr>
<tr>
<td>Obstruction at the Right Popliteal Artery</td>
</tr>
<tr>
<td>Right</td>
</tr>
<tr>
<td>Left</td>
</tr>
<tr>
<td>Arm</td>
</tr>
<tr>
<td>ABI</td>
</tr>
<tr>
<td>Obstruction at the Abdominal Aorta or Bilateral Iliac Obstruction</td>
</tr>
<tr>
<td>Right</td>
</tr>
<tr>
<td>Left</td>
</tr>
<tr>
<td>Arm</td>
</tr>
<tr>
<td>ABI</td>
</tr>
</tbody>
</table>

Typical pressures in a normal individual. The findings, based on resting pressures, show no evidence of occlusive disease of the large- or medium-sized arteries. Normal findings are as follows: (1) ABI of 1.0 or higher; (2) all segmental pressure gradients lower than 30 mm Hg; and (3) upper thigh pressure at least 40 mm Hg above brachial pressure.

Typical pressures in a patient with obstruction of the right popliteal or tibial arteries. Significant findings are as follows: (1) ABI less than 0.9 in the right leg; (2) abnormally high gradient from the ankle to below the knee and again from below to above the knee in the right leg; and (3) upper thigh pressure 50 mm Hg higher than brachial pressure, consistent with normal flow at the aorta-iliac level. The findings are suggestive of right popliteal occlusion or right anterior and posterior tibial occlusion, or both.

Typical pressures in a patient with obstruction of the abdominal aorta or bilateral iliac obstruction. Significant findings are as follows: (1) ABI of less than 0.9 in both legs; (2) all segmental gradients lower than 30 mm Hg; and (3) both upper thigh pressures relatively low with respect to brachial pressure. The findings are suggestive of severe aortoiliac occlusive disease.

ABI, Ankle-brachial index.

The central nervous system response to ischemia further stimulates the sympathetic nervous system after arterial pressure falls below 50 mm Hg. Subsequent compensatory mechanisms that work to restore blood volume to a normal level include the release of angiotensin and antidiuretic hormone (vasopressin). This causes arteriolar vasoconstriction, conservation of salt and water by the kidneys, and a shift in fluid from the interstitium to the intravascular space.

Several investigators have examined the changes in blood pressure and pulse that occur in supine patients with blood loss. Collectively, these studies have shown variable individual hemodynamic responses to acute blood loss of up to 1 L. The frequent inability to detect significant loss of volume with supine vital signs, and the observation that syncope frequently develops in patients with acute volume loss on rising, led to the investigation of the use of orthostatic vital signs to detect occult hypovolemia.

**Physiologic Response to Changes in Posture**

When an individual assumes the upright posture, complex homeostatic mechanisms compensate for the effects of gravity on the circulation to maintain cerebral perfusion with minimal change in vital signs. These responses include (1) baroreceptor-mediated arteriolar vasoconstriction; (2) venous constriction and increased muscle tone in the legs and the abdomen to augment venous return; (3) sympathetic-mediated inotropic and chronotropic effects on the heart; and (4) activation of the renin-angiotensin-aldosterone system. These compensatory mechanisms preserve cerebral perfusion in the upright position with minimal changes in vital signs. When a normal individual stands, the pulse increases by an average of 13 beats/min, systolic blood pressure falls slightly or does not change, and diastolic pressure rises slightly or does not change. In patients with vasodepressor syncope, the normal compensatory reflexes that preserve cerebral perfusion with changes in posture are altered. The normally increased sympathetic tone on standing is paradoxically inhibited, and an exaggerated enhancement of parasympathetic activity (bradycardia) occurs and can lead to syncope.

Few data exist regarding the true effect of acute blood loss on postural vital signs, and this parameter varies greatly among individuals experiencing hypovolemia. One study of 23 young adult volunteers from whom 500 to 1200 mL of blood was withdrawn found no reliable change in postural blood pressure, and a consistent postural increase in pulse of 35% to 40% was noted after 500-mL blood loss. Of the six individuals from whom approximately 1000 mL of blood was withdrawn, only two were able to tolerate standing, and each had a postural increase in pulse of greater than 30 beats/min. The other four individuals experienced severe symptoms on standing, followed by marked bradycardia and syncope if they were not allowed to lie down.

Following phlebotomy of 450 to 1000 mL of blood from healthy volunteers, the criterion of an increase in pulse of 30

**BOX 1.1 Homeostatic Mechanisms in Hemorrhagic Shock**

- **Sympathetic reflex compensation**
  - Arteriolar vasoconstriction
  - Venous capacitance vasoconstriction
  - Increased inotropic and chronotropic cardiac activity
  - Central nervous system ischemic response

- **Selective increase in cerebral and coronary perfusion by means of local autoregulation**

- **Increased oxygen unloading in tissues**

- **Restoration of blood volume**
  - Renin-angiotensin-aldosterone axis activation
  - Antidiuretic hormone secretion
  - Transcapillary refill

- **Increased thirst resulting in increased fluid intake**

- **Increased erythropoiesis**
beats/min or the presence of severe symptoms (syncope or near-syncpe) during a supine-to-standing test accurately distinguished between 1000-mL blood loss and no blood loss. The sensitivity and specificity of using the aforementioned criteria for detecting 1000-mL blood loss (Box 1.2) were both 98%, for an accuracy of 96% (2% false-negative results and 2% false-positive results). The investigators were unable to consistently detect blood loss of 500 mL with these criteria.161 In a similar study, the change in heart rate with postural changes after 500-mL phlebotomy was more discriminatory for blood loss than were changes in blood pressure or a change in the bioimpedance-based stroke index. Of note, these authors found that a change in heart rate of 30 beats/min or greater was 13.2% sensitive and 99.5% specific for 500-mL blood loss and that a change in heart rate of 20 beats/min or greater was 44.7% sensitive and 95.4% specific. The finding of a significant rise in pulse, though insensitive for 500-mL blood loss, was relatively specific in these healthy adult blood donors.162 In the most recent metaanalysis of orthostatic vital signs (1999), the authors concluded that a large postural pulse change (>30 beats/min) or severe postural dizziness precluding the completion of vital sign measurements is required to clinically diagnose hypovolemia secondary to acute blood loss. The analysis demonstrated that orthostatic vital signs are often absent after moderate amounts of blood loss, thus significantly limiting the test’s sensitivity (22%) in this scenario.168 A consensus statement defined orthostatic hypotension as a reduction in systolic blood pressure of at least 20 mm Hg or a reduction in diastolic blood pressure of at least 10 mm Hg within 3 minutes of standing and reinforces the theory that this is a physical finding rather than a disease process.164

Variables Affecting Orthostatic Vital Signs

Many conditions affect the compensatory mechanisms that allow patients to assume the upright posture. Most of the conditions that affect postural blood pressure regulation involve a pathologic condition that affects the sympathetic nervous system or the use of medications that alter normal cardiovascular compensatory functions. Orthostatic hypotension caused by autonomic insufficiency is not usually accompanied by tachycardia, and the orthostatic hypotension produced by acute volume depletion is commonly accompanied by a pronounced reflex tachycardia. Even in normal individuals, passive tilting generates a high incidence of orthostatic syncope.160

Because of decreased vasomotor tone, limited chronotropic response, and other factors, the elderly have a higher incidence of orthostatic hypotension leading to syncope and fall-related injuries.170 Carotid sinus hypersensitivity may play a greater role than orthostasis in geriatric syncope.171 Note that drugs that antagonize the normal autonomic compensatory mechanisms can also produce orthostatic changes. These changes can be severe enough to produce frank syncope, especially in the elderly. A study of patients with syncope seen in an ED found that orthostatic hypotension was considered to be the cause of the syncope in 24%, and that the largest proportion of these cases were drug related. Patients with orthostatic hypotension as the cause of syncope were older, had more comorbid conditions, and were found to be more frequently taking antihypertensive medications.172 Advanced patient age is an independent risk factor for orthostatic blood pressure changes and does not appear to have a direct correlation to the presence of chronic cardiovascular disease, disability, or body mass index.173 A study assessing vital sign changes prior to ward cardiac arrest found significant differences between elderly and nonelderly patients in the predictive value of the Modified Early Warning Score, noting more accurate detection of cardiac arrest by vital sign changes in nonelderly patients.174

Patients with hypertension may also have abnormal vasomotor responses to tilt testing and demonstrate more instability.165 Chronic anemia patients, who exhibit compensated blood volume, seem to have the same postural response as normal individuals.175 Ethanol ingestion exaggerates postural pulse changes and mimics the hemodynamic changes seen with acute blood loss.176

The utility of orthostatic vital signs in children has been questioned. Healthy adolescents had changes in heart rate of 21.5 ± 21.2 beats/min and variable changes in systolic blood pressure (+19 to −17 mm Hg) after 2 minutes of standing.178 A study comparing mildly dehydrated children with normal children found a significant difference in the orthostatic rise in pulse, and no difference in orthostatic blood pressure. The investigators concluded that an orthostatic increase in pulse of greater than 25 beats/min constitutes a positive tilt test and less than 20 beats/min constitutes a negative test (sensitivity of 75%, specificity of 95%, and predictive value of 92% when using near-syncpe or an increase in heart rate greater than 25 beats/min).179

Another complicating factor in interpreting orthostatic vital signs is the development of paradoxical bradycardia in the presence of blood loss. Bradycardia in the face of hemorrhage has generally been considered a preterminal finding of irreversible shock, and bradycardia has been documented in hypovolemic, conscious trauma patients as well. It has been reported that when orthostatic syncope occurs, it is accompanied by hypotension and often bradycardia.180,161 Many central nervous system factors can contribute to vagally mediated syncope in ED patients with acute traumatic blood loss, including pain, the sight of blood, anxiety, and nausea. This paradoxical bradycardia may be more frequently associated with rapid and massive bleeding. Patients with more gradual blood loss tend
to have a more typical tachycardic response. When the patient’s clinical findings are consistent with loss of volume or shock, the clinician should not allow the absence of tachycardia to change the assessment.

Indications and Contraindications

When the volume status of a patient is assessed with the use of orthostatic vital signs, several points should be remembered. Many factors influence orthostatic blood pressure, including age, preexisting medical conditions, medications, and autonomic dysfunction. It must be emphasized that data relating the effect of blood loss to orthostatic vital signs are limited to phlebotomized healthy volunteers. Great care must be used when extrapolating these data to patients with anemia, dehydration, or painful trauma. The clinician must consider the patient’s clinical condition when interpreting orthostatic vital signs in the evaluation of a patient with potential volume depletion.

Orthostatic vital signs can be considered an adjunct for the evaluation of any patient with known or suspected loss of blood volume or a history of syncope. Contraindications to orthostatic measures include supine hypotension, the clinical syndrome of shock, severely altered mental status, the setting of possible spinal injuries, and lower extremity or pelvic fractures.

The use of medications that block the normal vasomotor and chronotropic response to orthostatic tests is also a contraindication to using this test for the assessment of volume status. When the patient’s volume status is believed to be adequate and the clinician seeks to determine whether specific medications may have affected the patient’s ability to respond to postural changes, the test may be useful. In these situations, the primary finding may be the sensation of near-syncope with little or no change in vital signs.

Orthostatic vital signs are often used to assess a patient’s response to therapy. In patients receiving intravenous rehydration therapy, serial orthostatic vital signs are widely used to judge the end point of therapy before release. In one study, individual orthostatic vital sign response to saline infusion in women with hyperemesis gravidarum was associated with other measures of rehydration, including weight gain and decreased urine specific gravity. Although the individual improvement in orthostatic vital signs in response to rehydration was of clinical value, the initial orthostatic vital signs were considered insufficient as the sole indicator of clinical dehydration in this population.

Technique

To obtain orthostatic vital signs, record the blood pressure and pulse after the patient has been in the supine position for 2 to 3 minutes (see Box 1.2). Allow the patient to rest quietly and do not perform any painful or invasive procedures during the test.

Next, ask the patient to stand and be prepared to assist if severe symptoms or syncope develop. A supine-to-standing test is more accurate than a supine-to-sitting one. If severe symptoms develop on standing, defined as syncope or extreme dizziness requiring the patient to lie down, the test is considered positive and should be terminated. If not symptomatic, allow the patient to stand for 1 minute and then record the blood pressure and pulse. A 1-minute interval resulted in the greatest difference between control and 1000-mL phlebotomy groups in one study.

A number of studies have been conducted on normotensive, normovolemic patients to assess end points for orthostatic vital sign parameters. Such studies have included sitting-to-lying methods and varying rates of postural changes, including lying times of 5 to 10 minutes and standing times of 0 to 2 minutes. Arm position may affect postural changes in blood pressure and should be held constant to accurately assess orthostatic change. Complications include syncope with resulting falls and injuries.

Interpretation

The most sensitive criteria for orthostasis are tachycardia or symptoms of cerebral hypoperfusion (e.g., near-syncope). Although changes in blood pressure may be seen, they are too variable to be an indicator of loss of blood volume. Specific population-based thresholds for changes in pulse rate and blood pressure have some value in identifying patients at high risk for significant loss of blood volume, but great individual variability limits the use of this technique as a screening test. That is, a loss of 500 mL, and occasionally more, may be associated with a negative orthostatic vital sign assessment. The use of serial measurements to ascertain the response to therapy in patients considered to be at risk for loss of volume appears to have clinical utility.

In the setting of suspected blood loss, if the patient has a rise in pulse of 30 beats/min or manifests severe symptoms and if other complicating factors have been excluded, blood loss is highly likely (2% false-positive rate). The presence of a negative test indicates only that acute blood loss of 1000 mL is unlikely (2% false-negative rate) and that blood loss of 500 mL cannot be excluded (43% to 87% false-negative rate). Orthostatic changes in the SI were no more sensitive than established tilt test criteria in discriminating normal individuals from those with moderate acute blood loss (450 mL).

Criteria for significant orthostatic changes in blood pressure cannot be definitively set for the following reasons: (1) large variability in postural blood pressure has been found in the adult ED population; (2) the results of studies using passive tilt tables cannot be extrapolated to the bedside use of orthostatic vital signs; (3) studies using healthy patients with acute blood loss may not reflect the orthostatic changes seen in the elderly, those with chronic bleeding, dehydration, other medical problems, or in association with certain medications; and (4) many studies of orthostatic changes did not use a criterion standard in their determinations.

In summary, orthostatic vital sign testing is common in the ED, but this procedure has limited proven value. The clinical interpretation of orthostatic changes in blood pressure and pulse varies widely. Although this intervention may occasionally yield information not obtained through other means, the authors do not consider orthostatic blood pressure testing a standard of care in the evaluation of ED patients.

CAPILLARY REFILL

The capillary refill test is a measurement of the interval of time from the release of pressure on the nail bed or soft tissue sufficient to blanch the nail bed or superficial soft tissue, until the return to normal coloration. Delayed capillary refill is an indication of reduced skin turgor, often as a result of volume depletion or limited perfusion. Measurement of capillary refill time (CRT)
appears to be somewhat accurate in children, and its accuracy in assessing dehydration and reduced perfusion in adults is highly suspect. Skin elasticity is the characteristic that allows skin to spring back to its original shape after it has been deformed, and the speed of refilling the capillary bed after compression is responsible for the return of color to the skin.

**Indications and Contraindications**

CRT should not be determined from a dependent extremity, from a recently burned or injured extremity, or at the site of an infection or acute injury. Because CRT can be used without additional equipment and takes only a few seconds to perform, it can be a useful bedside assessment of perfusion and dehydration when used in conjunction with other objective signs of the adequacy of perfusion. It should not be considered accurate as a stand-alone tool. Capillary refill is not an appropriate alternative to measuring blood pressure in pediatric patients.

**Procedure**

The preferred sites for determining CRT are the nail bed, the thenar surface of the palm, and the heel. Alternative sites may have different CRTs, and the current standards are best developed for capillary refill determined at the nail bed. Regardless of the site chosen, position the extremity at approximately the level of the right atrium. The minimum pressure necessary to produce blanching yields the most reproducible values. Release the nail bed and begin timing with a stopwatch or simply by counting out “one-thousand-one, one-thousand-two” for an approximation of the interval. Stop the clock when the nail bed becomes pink again. Interobserver reliability has been shown to be moderate, with kappa values of less than 0.5 in the evaluation of both adults and children.

**Interpretation**

The normal CRT increases with age and is slightly longer in female patients. It is further increased by degrees of dehydration or hypoperfusion. Hypothermia, hyponatremia, congestive heart failure, malnutrition, and edema all increase CRT. Environmental conditions such as ambient air temperature can falsely alter capillary refill. Fever alone did not appear to prolong or shorten CRT in children, and a study of healthy adults found a 5% decrease in CRT for each degree Celsius rise in patient temperature.

The main difficulty in interpreting CRT is that normal values in healthy patients fall into a wide range. In 30 normal infants 2 to 24 months of age, mean CRT was 0.8 ± 0.3 second. Measurements obtained from the nail bed were more reproducible than those from the heel. Combined results from four studies evaluating capillary refill revealed a pooled sensitivity of 0.60 (95% confidence interval [CI], 0.29 to 0.91) and a specificity of 0.85 (95% CI, 0.72 to 0.98) for detecting 5% dehydration in children. The presence of a 2-second or longer delay in CRT when combined with any two or more of the findings of absent tears, dry mucous membranes, or ill general appearance predicted clinical dehydration (>5% deficit in body weight) in children (1 month to 5 years of age) with 87% sensitivity and 82% specificity. A 2015 systematic review including 21 studies suggests applying a cutoff of 3 seconds or more to define abnormal CRT in infants and children over 7 days of age.

Frequent monitoring of capillary refill may be useful in assessing responses to rapid fluid resuscitation in children. A normal CRT of 2 seconds or less has been shown to correlate with superior vena cava oxygen saturation of 70% or higher in critically ill children. The role of serial CRT measurements for assessing the response to rehydration in adults is unknown, and it does not appear to be useful for assessing acute loss of blood volume. Lima and colleagues studied the prognostic value of the subjective assessment of peripheral perfusion in critically ill patients following initial resuscitation. When an abnormal CRT was defined as greater than 4.5 seconds, coupled with extremity coolness, these parameters identified patients who had been hemodynamically stabilized and continued to have more severe organ dysfunction and higher lactate levels. In adults, CRT was found to be less sensitive and less specific than orthostatic vital signs in detecting 450-mL blood loss during blood donation. However, a 2014 study showed CRT measured as a continuous variable in a cohort of admitted medical patients, was associated with an increased 1-day and 7-day all-cause mortality.

In summary, assessment of CRT is a common technique used in the ED, and, in reality, this procedure has limited proven value, clinical interpretation is very subjective, and use varies widely. Although this intervention may occasionally yield information not obtained through other means, predominantly in children, the authors do not consider such testing an absolute standard of care in the evaluation of ED patients.

**TEMPERATURE**

Detection of abnormal body temperature may facilitate proper diagnosis and evaluation of patients’ complaints in the ED. An inability to maintain normal body temperature is indicative of a vast number of potentially serious disorders, including infections, neoplasms, shock, toxic reactions, and environmental exposures. Fever in neutropenic, immunocompromised, or intravenous drug–abusing patients may be more reliable than laboratory tests or clinician assessment in diagnosing serious illness. Infants are particularly sensitive to thermal stress and may demonstrate lower body temperatures during critical illness. In one study, although higher rates of serious bacterial infection were found in neonates who had documented fever (OR, 3.23) on admission, only 8.4% of neonates with historical fever were later diagnosed with serious bacterial infection. Pretriage or home assessment of body temperature is fraught with difficulty and unreliability. Whether taken by the oral, rectal, or tympanic routes, reports of fever at home are very difficult to interpret in the clinical setting. Some studies report rates as low as 13% in frequency when measuring temperature in the critically ill or injured, and temperature and Glasgow Coma Scale were the least frequently documented physiologic observations (less frequent than RR, oxygen saturation, heart rate, and systolic blood pressure) across ED, critical care, and medical/surgical unit settings despite temperature representing a key sign of systemic inflammatory response syndrome.

**Physiology**

Under normal conditions, the temperature of deep central body tissues (i.e., core temperature) remains at 37°C ± 0.6°C (98.6°F ± 1.08°F). Core body temperature can be maintained...
within a narrow range, whereas environmental temperature varies from as much as 13°C to 60°C (55°F to 140°F), and surface temperature rises and falls with environmental and other influences. Mean oral temperature is 36.8°C ± 0.4°C (99.2°F ± 0.7°F). Maintenance of normal body temperature requires a balance of heat production and heat loss. Heat loss occurs by radiation, conduction, and evaporation by approximately 60%, 18%, and 22%, respectively. Heat loss is increased by wind, water, and lack of insulation (e.g., clothing). Sweating, vasodilation, and decreased heat production serve to decrease temperature. Piloerection, vasoconstriction, and increased heat production serve to increase body temperature. Heat production is increased by shivering, fat catabolism, and increased thyroid hormone production.

Temperature is controlled by feedback mechanisms operating through the hypothalamus. Heat-sensitive neurons in this area increase their rate of firing during experimental heating. Receptors in the skin, spinal cord, abdominal viscera, and central veins primarily detect cold and provide feedback to the hypothalamus that signals an increase in heat production. Stimuli that change core body temperature result in reflex changes in mechanisms that increase either heat loss or heat production.

**Indications and Contraindications**

Clinicians generally measure body temperature to determine whether it is outside the normal range and as an indication of pathologic conditions that can affect core body temperature. Measurement of actual core body temperature requires the placement of invasive monitors, such as an esophageal or pulmonary artery (PA) probe. Clinicians commonly use estimates of core body temperatures to conveniently and safely assess abnormalities in core temperature. All noncore body sites and methods have inherent limitations in accuracy, and clinicians have come to accept these shortcomings in assessing most patients.

Measurement of oral temperature requires a cooperative adult or child. Patients who are uncooperative, hemodynamically unstable, septic, or in respiratory distress (with an RR > 20 beats/min) require a method of measuring temperature other than the traditional oral route. This group includes children younger than 5 years and patients who are intubated. Ingestion of hot or cold beverages has shown they can alter oral temperature readings for 5 to 30 minutes, and can falsely elevate a normal temperature or mask a fever.

Special techniques of measuring core body temperature may be indicated in certain patients (e.g., those with profound hypothermia, frostbite, or hyperthermia). Measurement of core body temperature is indicated in these individuals because it accurately measures the effects of treatment. This is the group of patients who will benefit the most from continuous temperature measurements.

**Measurement Sites**

**Core Body Temperature**

The following sites accurately reflect core body temperature and changes in it: the distal third of the esophagus, the tympanic membrane (TM) (with a direct thermistor in contact with the anterior inferior quadrant of the TM), and the PA. Other sites may represent core body temperature under certain conditions; for example, (1) the rectum when the temperature is obtained at least 8 cm from the anus with an indwelling thermistor and the body temperature is relatively constant, and (2) the bladder when measured with an indwelling thermistor.

Rectal temperature is often considered the criterion standard for body temperature in ambulatory patients and it is often used routinely in children younger than 3 years. Advantages include accuracy, sensitivity, and availability. One intensive care unit study found that rectal probe temperatures demonstrated limited variability or bias when compared with PA temperatures. Disadvantages include longer intervals for measurement, safety concerns, and inconvenience. Neutropenia and recent rectal surgery represent relative contraindications to measurement of rectal temperature. Thermistor probes (i.e., small thermocouples with instantaneous readouts) for esophageal and vascular temperature measurement provide continuous temperature readouts when attached to a potentiometer. Thermistor probes are available for measurement of esophageal, bladder, and rectal temperature with appropriate monitors.

**Peripheral Body Sites Approximating Core Temperature**

A body temperature measurement by IR radiation can be detected from the ear, including the auditory canal and TM; it is easy to use, hygienic, convenient, and quick. It can be used as a general screening technique, particularly in cases when temperature is not of great importance, such as in minor trauma. Though clearly superior to axillary temperature readings, controversy remains regarding the sensitivity and specificity of IR TM readings in the ED. In a cohort of ICU patients, agreement of tympanic with PA temperature was inferior to that of urinary temperature. Compared with PA temperatures, Δ (limits of agreement) were 0.36°C (–0.56°C, 1.28°C), –0.05°C (–0.69°C, 0.59°C), and 0.30°C (–0.42°C, 1.01°C) for tympanic, urinary, and axillary temperatures, respectively. It is possible that alterations of regional blood flow accompanying critical illness (TMs may behave as an extension of the skin or the mucous membrane in the critically ill) and the peripheral vasoconstriction that occurs with inotropes and some forms of shock may occur in the TM, making such measurements less accurate in this population.

More work is being done in the pediatric population, with an overall sensitivity of between 50% and 80% and a specificity of 85%. The lower sensitivities are found in newborns and infants younger than 3 years. Systematic reviews of pediatric studies have pooled data suggesting 65% sensitivity, which is unacceptable in the clinical setting. A 2010 study found that neither TM nor skin thermometers could reliably predict rectal temperature, and it concurred that these methods could not replace rectal temperature measurement as the “gold standard” for detecting fever in the pediatric population. Adult studies, though generally more favorable in recommending TM temperatures, have shown gaps in reliability as well. However, a 2011 systematic review suggested that TM and oral thermometry provides an accurate measure of core temperature in critically ill adults with fever. Temporal artery scanning to detect fever is increasingly being examined. In general, these devices show better sensitivity in detecting fever in infants than TM thermistors do (66% versus 49% sensitivity) and may be useful in excluding fever, defined as a rectal temperature higher than 38.3°C if the temporal artery readings are lower than 37.7°C. The point can be argued that a sensitivity of 60% in any population lacks the required sensitivity
to be useful clinically. A theoretical disadvantage of TM temperatures might be a falsely elevated estimate of the core temperature in the presence of otitis media. In one study, TM thermometers accurately reflected oral temperatures in children with otitis media.\(^{330}\)

In a group of pediatric patients receiving anesthesia, measurements taken at the temporal artery were similar to nasopharyngeal thermometer recordings, and the axillary temperature was lower than the temporal artery and nasopharyngeal temperatures.\(^{231}\)

Prehospital providers who might wish to measure IR TM temperature at low ambient temperatures should be aware that below 24.6°C, the TM readings will greatly underestimate core temperatures.\(^{332}\) EMS personnel should also be aware that in a cohort of exhausted marathon runners, rectal and IR TM temperatures have only moderate correlation.\(^{333}\) External thermometry may be unreliable in the setting of heat-related illness or hypothermia; if these diagnoses are clinically suspected and the IR TM temperature does not confirm an abnormal temperature, a measurement by rectal or esophageal probes should be obtained for diagnosis and during treatment.\(^{234}\)

Axillary and tactile temperature assessments have been demonstrated to be unreliable and insensitive. They should not be used as screening methods for core temperature abnormalities in the ED.\(^{235-237}\) Single-use Tempa-DOT thermometers show progressive temperature dot darkening with increasing temperature, and these thermometers have been adopted by many EDs. These temperature devices have been shown to have 100% sensitivity and approximately 80% specificity for identification of fever when temperature was obtained orally in both adults and children.\(^{238}\) The most rudimentary method for temperature measurement, parental assessment by tactile touch, is associated with a measured fever approximately 75% of the time.\(^{239}\) Clinician estimation of fever is almost identical (70%).\(^{240}\)

**Procedure**

Begin the temperature measurement by selecting the body site. Consider the accuracy of using a certain site to reflect core temperature, the sensitivity of the site to changes in temperature, the convenience, the time required, the safety, and the availability of the site.\(^{241}\) Insert the temperature probe and allow the probe to equilibrate with the temperature of the local body tissues.

Proper placement of the temperature probes significantly influences the results of oral, rectal, esophageal, and vascular temperatures.\(^{242-245}\) Obtain sublingual oral temperatures in either the right or the left posterior sublingual pocket with the mouth closed.\(^{242}\) Though anxiety-producing for parents, biting and breaking a mercury thermometer is generally inconsequential with regard to ingestion of either glass or mercury. Obtain a rectal temperature with the patient in the left or right lateral decubitus position and advance the probe gently to a depth of 3 to 5 cm to ensure accurate, atraumatic results.\(^{246}\) Complications associated with rectal temperature measurements are extremely rare and include rectal perforation, pneumoperitoneum, bacteremia, dysrhythmias, and syncope.\(^{331}\) Falsely low supranormal rectal temperature measurements may be seen during shock.\(^{247}\) Rectal temperature may also lag behind changes in core temperature. An esophageal catheter or PA probe can be placed for measurement of core body temperature, and these techniques are not typically used in the ED.

Though not commonly used, measurement of the temperature of a freshly voided urine specimen can validate measurement of temperature at other body sites.\(^{248}\) Urinary bladder temperature measurement has been demonstrated to be similar to PA catheter temperature.\(^{249}\)

Digital electronic probes are commonly used for the measurement of oral temperature in ambulatory patients.\(^{250}\) Electronic methods of temperature measurement are based on the thermocouple principle. Modern electronic thermometers signal when extrapolation of the temperature-time curve has occurred. Current in vitro standards call for an accuracy of ± 0.1°C (± 0.18°F) over the range of 37°C to 39°C (98.6°F to 102.2°F). Disadvantages include factors that affect clinical accuracy and sensitivity. Disposable single-use oral thermometers are now available and are as reliable as mercury or TM thermometers.\(^{251}\) In the pediatric population, pacifier thermometers record supralingual readings in infants. The average time needed to record a reading with a pacifier thermometer is 3 minutes and 23 seconds, thus making its application in emergency medicine limited, although its sensitivity (72%) and specificity (98%) rival that of alternative methods.\(^{352}\) Various IR ear thermometers are available commercially with varying operating temperature ranges, features, and reported accuracy.\(^{253}\) Complications associated with axillary, oral, ear IR, and liquid crystal thermometers are rare or unreported. TM perforation and pain have been reported as complications of placement of the thermistor probe in the auditory canal.

**Interpretation**

Normal values for body temperature are affected by the following variables: (1) site and methods used for measurement; (2) perfusion; (3) environmental exposure; (4) pregnancy, (5) activity level; and (6) time of day. Clinicians must interpret body temperature with knowledge of the range of normal values at the intended site of measurement. Although core body temperature remains nearly constant (37.0°C ± 0.6°C or 98.6°F ± 0.1°F), surface temperature rises and falls with changes in ambient temperature, exercise, and time of day. The definition of fever varies by the site of measurement and is defined by a temperature greater than 2 standard deviations above the mean. Fever has been defined as an oral temperature of 37.8°C or higher (100.0°F),\(^{246}\) a rectal temperature of 38.0°C or higher (100.4°F),\(^{254}\) or an IR ear temperature of 37.6°C or higher (99.6°F).\(^{246}\) Based on measurement of temperatures in normal, healthy infants, it is recommended that fever be defined as a rectal temperature of 38°C or higher in infants younger than 30 days, 38.1°C or higher in infants 30 to 60 days (1 to 2 months), and 38.2°C or higher in infants 60 to 90 days old (2 to 3 months).\(^{255}\) Hypothermia has been defined as a core body temperature lower than 35°C (<95°F), and hyperthermia has been defined as a core body temperature higher than 41°C (>105.8°F) with accompanying symptoms and signs.\(^{256}\) A useful nomogram and formulas for conversion of centigrade to Fahrenheit are provided in Fig. 1.6.

Temperature probes that depend on transfer of heat energy from local tissues to the probe require a period of equilibration and reliable tissue contact at the intended body site. Acceptable equilibration times for mercury-in-glass thermometers in oral, rectal, and axillary sites are 7, 3, and 10 minutes, respectively. Used in a predictive mode, electronic digital thermometers generally require 30 seconds for oral or rectal temperature
For unadjusted ear temperature using the Thermoscan Pro-1 (Thermoscan, Inc., San Diego, CA), temperature obtained with a properly positioned pulmonary artery, esophageal, or tympanic membrane thermistor.

Before taking an oral temperature the examiner should inquire about these features and possibly delay taking the temperature. A 2.7°C (4.9°F) reduction in oral temperature measurement was found when the probe was placed under the tip of the tongue instead of under the posterior sublingual pocket. Given the extrapolation that occurs with rapidly reading thermocouple devices and IR detectors, it is not surprising that the sensitivity of these devices for detection of fever is only 86% to 88%. Many clinicians have adopted the adage that when an elevated temperature is suspected or crucial in decision making and not evident with an oral thermocouple probe or IR TM thermometer, measurement with a mercury-in-glass thermometer is indicated.

Infrequently, ED patients require constant monitoring of temperature (e.g., in cases of profound hypothermia or hyperthermia). This can usually be performed by using a bladder or esophageal probe attached to a potentiometer. Patients with indwelling central venous or pulmonary arterial catheters may have electronic thermistors inserted into the central circulation to measure core body temperature. As noted earlier, rectal temperature measurements are less desirable for monitoring patients undergoing rapid changes in core temperature. Periodic IR TM temperature monitoring may represent one useful option in a hypothermic patient.

Interpretation of ear IR temperatures requires knowledge of the mode of thermometer operation and ambient temperature. Occlusion of the ear canal by cerumen may produce a falsely low reading. Most IR ear thermometers appear to be moderately sensitive for fever. If these devices are used, the clinician must be aware of the potential for a falsely low temperature. When in doubt, the measurement should be repeated with a more standard method.

Patients, parents, and caregivers often misinterpret the significance of a fever, and the term “fever phobia” has been coined to describe the ubiquitous and unsubstantiated fear that fever, by itself, is harmful or has diagnostic or prognostic significance. Some of the highest fevers are the result of benign viral infections. Fever has been demonstrated to increase the body’s immune response, making aggressive fever reduction counterproductive. Most individuals, especially children, feel better when a high fever is lowered. Uninitiated medical personnel may contribute to the incidence of fever phobia. Routine administration of antipyretics in the ED for any fever, regardless of the degree of discomfort, has fueled unwarranted concern and may result in potentially harmful interventions, unnecessary testing and treatments, avoidable side effects of

equilibration. The predictive mode uses temperature changes versus time to predict an equilibration temperature.

Normal ranges and suggested febrile thresholds for common body sites and methods should be considered in the interpretation of temperature values (Table 1.5). Interpretation of temperature measurements during clinical assessment must consider the use of antipyretics, level of activity, pregnancy, environmental exposure, and patient age. Body temperature is increased during sustained exercise, pregnancy, and the luteal phase of the menstrual cycle. Temperature also increases in the late afternoon because of diurnal variation. Body temperature is generally reduced with advanced age, and age may have an impact on the magnitude of fever. Axillary temperatures have a low sensitivity but a high specificity for fever. Axillary temperatures should not be used to screen for fever.

Oral temperature measurements are affected by the ingestion of hot or cold liquids, tachypnea, and cold ambient air. Before taking an oral temperature the examiner should inquire about these features and possibly delay taking the temperature. A 2.7°C (4.9°F) reduction in oral temperature measurement was found when the probe was placed under the tip of the tongue instead of under the posterior sublingual pocket. Given the extrapolation that occurs with rapidly reading thermocouple devices and IR detectors, it is not surprising that the sensitivity of these devices for detection of fever is only 86% to 88%. Many clinicians have adopted the adage that when an elevated temperature is suspected or crucial in decision making and not evident with an oral thermocouple probe or IR TM thermometer, measurement with a mercury-in-glass thermometer is indicated.

When rapid changes in body temperature occur, oral and TM temperature measurements appear to be more reliable than rectal temperature. In 20 adults examined during open-heart surgery, oral temperatures showed a better correlation with blood temperature during rapid cooling and rewarming. Additionally, esophageal temperature measurements showed a faster response rate compared to temperature measured in the bladder when cold saline infusion was used to induce mild hypothermia in cardiac arrest patients.

Infrequently, ED patients require constant monitoring of temperature (e.g., in cases of profound hypothermia or hyperthermia). This can usually be performed by using a bladder or esophageal probe attached to a potentiometer. Patients with indwelling central venous or pulmonary arterial catheters may have electronic thermistors inserted into the central circulation to measure core body temperature. As noted earlier, rectal temperature measurements are less desirable for monitoring patients undergoing rapid changes in core temperature. Periodic IR TM temperature monitoring may represent one useful option in a hypothermic patient.

Interpretation of ear IR temperatures requires knowledge of the mode of thermometer operation and ambient temperature. Occlusion of the ear canal by cerumen may produce a falsely low reading. Most IR ear thermometers appear to be moderately sensitive for fever. If these devices are used, the clinician must be aware of the potential for a falsely low temperature. When in doubt, the measurement should be repeated with a more standard method.

Patients, parents, and caregivers often misinterpret the significance of a fever, and the term “fever phobia” has been coined to describe the ubiquitous and unsubstantiated fear that fever, by itself, is harmful or has diagnostic or prognostic significance. Some of the highest fevers are the result of benign viral infections. Fever has been demonstrated to increase the body’s immune response, making aggressive fever reduction counterproductive. Most individuals, especially children, feel better when a high fever is lowered. Uninitiated medical personnel may contribute to the incidence of fever phobia. Routine administration of antipyretics in the ED for any fever, regardless of the degree of discomfort, has fueled unwarranted concern and may result in potentially harmful interventions, unnecessary testing and treatments, avoidable side effects of

| TABLE 1.5 Normal Ranges and Suggested Febrile Thresholds for Human Body Temperature (in Healthy Resting Patients) |
|---|---|---|
| BODY SITE | TYPE OF THERMOMETER | NORMAL RANGE (°C) | FEVER (°C) |
| Core* | Electronic | 36.4–37.9 | 38.0 |
| Oral | Mercury in glass, electronic | 35.5–37.7 | 37.8 |
| Rectal | Mercury in glass, electronic | 36.6–37.9 | 38.0 |
| Ear | Infrared emission | 35.7–37.5 | 37.6* |

*Temperature obtained with a properly positioned pulmonary artery, esophageal, or tympanic membrane thermometer.

*For unadjusted ear temperature using the Thermoscan Pro-1 (Thermoscan, Inc., San Diego, CA).
medications, and additional patient anxiety. Mandatory medical evaluation for any fever, the use of alcohol sponge baths, waking a sleeping child to administer antipyretics, alternating ibuprofen with acetaminophen, around-the-clock use of antipyretics, and misconceptions about fever causing brain damage are some of the myths that are still extant in some EDs and throughout the general population. An ED protocol stating that any child be afebrile, demonstrate a significant decrease in hyperpyrexia, or even have a temperature reading repeated before discharge has no scientific validity.

**PAIN AS A VITAL SIGN**

**Background**

One of the goals of modern medicine has been to understand, accurately assess, and adequately treat pain. A paradigm shift has occurred in which pain is viewed as a mechanism to provide a mechanical warning of actual or potential damage to cells and tissues in a specific area. An individual's pain should certainly be addressed early in the ED encounter, but reflex interventions aimed at immediately relieving all a patient's pain are inappropriate and potentially counterproductive in the long run. Importantly, the concept of using pain as "the fifth vital sign" can be problematic, and this concept has not met with universal agreement. Literal interpretation of this concept can be fraught with problems, and because of the subjective nature of pain, measurement and interpretation are, by nature, imprecise. An overemphasis on rigorously quantifying pain and aggressively treating a specific number on the pain scale has the potential to expose patients to unneeded opioids, result in unrealistic expectations by patients, and lead to opioid dependence or addiction.

Pain can cause sympathetically mediated changes in vital signs, but standard vital signs (pulse, respiration, blood pressure) do not meaningfully correlate with the level of perceived pain. It is a common error to relate changes in vital signs, or lack thereof, as an indication of a patient's true level of pain. A 2006 retrospective study of patients with confirmed painful diagnoses showed no clinically significant associations between self-reported triage pain scores and heart rate, blood pressure, or RR. Contentsions that the judicious use of analgesics will obscure a clinical condition or otherwise adversely affect clinical care are antiquated and unscientific. Adequate pain control may facilitate the overall ED encounter for both the patient and clinician.

Initiatives mandating documentation of pain as the fifth vital sign have not been associated with improvement in pain management. A prospective, multicenter study evaluating the current state of ED pain management practices concluded that there is high pain intensity, as demonstrated by intense pain on arrival (median rating, 8 of 10), and suboptimal pain management practices. It also showed assessment of pain occurring in 83% of patients, 40% of patients receiving no analgesics, lengthy delays in analgesic administration, and a large proportion of patients reporting moderate to severe pain at discharge (74%). Assuming weight-based dosing for acute pain management, analgesic regimens did not conform to the recommended regimens in a study of ED patients. Mandating the recording of a triage pain score has been shown to improve time to initial analgesic treatment. Acute or chronic pain is the most common chief complaint of more than 50% of ED patients, with the figure approaching 75% in some studies. Reliably quantifying pain should be the goal of ED clinicians and is an appropriate step in the triage process.

**Procedure/Interpretation**

The Joint Commission goals mandate that all hospitals develop comprehensive programs for the measurement, treatment, and documentation of pain. There is no perfect measurement tool suitable for the wide variety of patients and clinical scenarios experienced in EDs. Pain is a complex, subjective, multifaceted, personal experience that is difficult to easily quantify in all individuals. The perfect pain assessment tool for the ED setting would be simple and rapid to administer while providing a precise, reliable, and valid measure of pain regardless of a patient's age, cultural background, and cognitive or physical impairments. Currently no tool meets all the criteria necessary to satisfy this definition. The primary focus should be to allow patients to report and rate their pain from a personal viewpoint, as it is widely accepted that health professionals cannot rate pain intensity as accurately as patients themselves. A patient's self-report of the pain is considered the gold standard for the initial assessment of pain and tracking of a response to interventions, although such reporting does not mandate specific interventions.

Multiple unidimensional acute pain measurement instruments have been published and many have been independently validated. Common instruments include the verbal rating scale, numerical rating scale, visual analog scale (VAS), and graphic rating scales. The most used pain scale in the ED is the 1 to 10 VAS. The VAS uses a 10-cm line bounded on each end by perpendicular stops and descriptors. Zero equates to no pain, and 10 equates to the worst pain ever experienced. The initial score is not as important as change during treatment. Generally, a change in VAS scores has been shown to be valid and reliable if self-completion is appropriate. There is wide variability in this technique, and at least a 13-mm to 30-mm (1.3-cm to 3-cm) change on the scale is required to validate clinically relevant worsening or relief. Low completion rates may be seen in patients with visual or cognitive impairment. Graphic rating scales are useful for patients with limited cognitive and expressive ability, especially children. They may also be helpful to overcome language or cultural differences. A systematic review recently evaluated a variety of graphic scales in pediatric patients and supported use of the Faces Pain Scale (FPS), the Faces Pain Scale-Revised (FPS-R), the Oucher Pain Scale, and the Wong-Baker Faces Pain Rating Scale (WBFPRS). In an ED survey, the WBFPRS was the most common (81.7% of responding facilities) pain measurement tool used for pediatric pain assessment. Scale ratings may overestimate or underestimate pain in certain patients and do not mandate a specific pain reduction approach. This is better driven by clinician evaluation and consideration of the entire scenario. Such ratings rarely correlate with the seriousness of a patient's medical condition. It is not uncommon for patients to rate their pain a 12 on a scale of 1 to 10 in an attempt to emphasize their personal experience, anxiety, or fear of receiving inadequate analgesia. It may be a patient's perception that a higher rating will expedite treatment, prompt higher doses of medications, or otherwise engender more compassionate care. A common erroneous tactic or subterfuge of litigation proceedings is to incriminate a clinician's diagnosis, treatment, or disposition based on the patient's rendition of the pain scale.
Overall Goal of Pain Relief

The goal of pain management in the ED is to adequately relieve or control pain without compromising diagnosis, treatment plans, or the safety of the patient and population. Although the theoretically ideal objective is to totally relieve pain, this goal is difficult to consistently achieve in the complex ED milieu, and should not be the standard of care. Pain relief is best accomplished by combining clinician experience, real-time clinical judgment, repeated evaluation, and discussion with the patient and family. Concerns about safety and the potential for abuse of addictive pharmaceuticals often weigh into the decision-making process for emergency physicians, and universal prescriptive reporting services can be used to assist in patient care and safe prescribing practices.278,279 Strict adherence to protocols, patient self-reporting on any pain scale, or reliance on a dogmatic approach to relief of a patient’s pain can be fraught with peril. It is best to treat the patient, not the pain scale.

In summary, although the routine use of patient-reported pain scales may facilitate the administration of analgesics and be somewhat useful in trending response to analgesia, experienced clinicians and the authors recognize their limitations in an ED clinical setting. With certainty, one cannot conclude that any specific pain scale has any diagnostic or prognostic value. Pain scales are best used to raise awareness that pain—and prudent efforts to relieve it—is addressed in the ED.

Automatic sphygmomanometer.


SPIROMETRY

For patients with acute exacerbations of asthma and chronic obstructive pulmonary disease (COPD), accurately estimating the severity of airflow obstruction is a critical component of their care. A focused history plus physical examination is the cornerstone of this assessment in the practice of emergency medicine. Wide variation exists in the ability to accurately diagnose airway obstruction, and a significant proportion of patients with marked airflow obstruction present without dyspnea. This blunted perception of disease severity may be a contributor to fatal and near-fatal asthma attacks.1,2 Similarly, patients presenting with acute exacerbations of asthma may experience subjective resolution of their symptoms after therapy, even when severe airflow obstruction is still present. Given these difficulties in recognizing airflow obstruction, objective measurement provides valuable information.

Spirometry is measurement of the volume of air exhaled during forced expiration. It can be interpreted as a function of time to determine the flow rate. Spirometry gives the most complete picture of lung mechanics and is the centerpiece of pulmonary function testing. Many parameters can be derived from a spirogram, the most useful of which are forced vital capacity (FVC), which is the total volume exhaled during a forced expiratory maneuver, and forced expiratory volume in 1 second (FEV1), which is the average flow rate during the first second of the forced expiratory maneuver (Fig. 2.1).

The advent of small handheld devices allows convenient spirometric evaluation in the emergency department (ED). The most common objective measurement of respiratory mechanics used in the ED is peak expiratory flow rate (PEFR). PEFR is the maximum flow of gas achieved during a forced expiratory maneuver. It correlates well with standard spirometry and has been studied extensively in the ED and outpatient setting.3–5

Indications

Evaluation of Acute Asthma Attacks

Currently, no standards exist for the measurement of pulmonary function parameters in ED patients, and practices vary widely. Most patients with asthma exacerbations can be evaluated, treated, and given a disposition with no further pulmonary function testing other than PEFR if quantitative assessment is deemed prudent. Several consensus guidelines recommend obtaining an objective measure of airflow obstruction in all patients seen in the ED with an acute exacerbation of asthma.6–8 Other guidelines have proposed that the decision to measure PEFR in patients with acute asthma should be individualized.9 It is reasonable that mild and easily reversible disease be evaluated and treated according to clinical judgment. If any pulmonary function parameters are to be used, their use should be optimized, including measurements at arrival, after initial treatment, and periodically thereafter.6–8

Evaluation of Exacerbations of COPD

PEFR and spirometry testing can yield objective data on airflow obstruction during the ED evaluation of COPD exacerbations. Though used by some ED practitioners, consensus guidelines do not recommend routine use of these tests in the acute setting.10,11

Differentiating Causes of Dyspnea

PEFR has been studied for its ability to differentiate between COPD and congestive heart failure (CHF).12,13 Insufficient data exist to recommend its routine use for this purpose in the ED.

Evaluation of Neuromuscular and Chest Wall Disease

Diseases of the chest wall and neuromuscular system can cause respiratory compromise. Though not commonly done in the ED, pulmonary function testing and assessment of negative inspiratory force can quantify the degree of impairment and help determine the level of admission needed.

Contraindications

Need for Immediate Intervention

Patients with severe respiratory compromise should receive aggressive therapy without delaying care for pulmonary function testing. Although portable pulmonary function testing may guide management, providing immediate intervention for those in distress should be the priority. Formal pulmonary function testing has limited value for acute exacerbations, and such assessments are most predictive when patients are at their baseline functional status.

Conditions That May Be Worsened by Increased Intrathoracic Pressure

Significant elevations in intrathoracic pressure will develop in patients performing a forced expiratory maneuver. Pneumothorax and pneumomediastinum may be exacerbated by the forced expiratory maneuver and aneurysms of the aorta or the cerebral vasculature could theoretically rupture with increased pressures. The presence of these conditions should be considered a relative contraindication to pulmonary function testing.

Equipment

Spirometers can be divided into two categories. Volume spirometers measure the amount of gas exhaled as a function of time. These devices tend to be cumbersome and are not ideally suited to the ED. Flow spirometers measure the flow of gas past a certain point and use that information to extrapolate volume and time data. These machines are smaller, simpler to use, and more portable. Flow spirometers determine gas flow by measuring the difference in pressure between two points in a tube (pneumotachograph), cooling of a heated wire (hot wire anemometer), or revolutions of a rotating vane. Most handheld spirometers also measure PEFR.

The most commonly used device to measure PEFR is the “mini-Wright” peak-flow flow meter (Fig. 2.2). These meters provide accurate and reproducible measurements of PEFR.14 Variation exists between types and brands of peak-flow flow
PEFR measurements are very sensitive to technique and patient effort. Even a small decrease in effort can lead to considerable degradation of results. In addition, any deviation from technique can lead to inaccurate results and a faulty estimation of the degree of airway obstruction. Therefore emergency physicians should directly observe and guide their patients when performing this testing.

Interpretation

Obstructive diseases are characterized by a disproportionate decrease in airflow (FEV₁) in relation to the volume of gas exhaled (FVC). A decreased FEV₁/FVC ratio with preservation of FVC indicates the presence of airflow obstruction. Restrictive diseases decrease total lung capacity and therefore decrease FVC to a greater degree than FEV₁. Decreased FVC with a normal or increased FEV₁/FVC ratio is indicative of restriction. It is useful to consider the FEV₁/FVC ratio when attempting to determine whether a patient has airflow obstruction. In patients with an established diagnosis of obstructive disease, FEV₁ is the test that best reflects changes in lung function. Typical values are shown in Table 2.1. These values are dependent on age, gender, ethnicity, and height and can be predicted from mathematical equations. Isolated measurements of PEFR are not reliable in making the diagnosis of asthma because of significant variation between individuals. It is appropriate to use PEFR to monitor the degree of airflow obstruction in known asthmatics.
Although measures of airflow obstruction are not standalone tests, when considered along with other clinical factors, they can guide decisions regarding the disposition of patients with acute asthma exacerbations. The highest of three PEFR or FEV₁ measurements should be used and, whenever possible, compared with the patient’s personal best. In circumstances in which previous best values are unknown or thought to be inaccurate, comparison with predicted values is appropriate. References are available for predicted PEFR values for adults by race and ethnicity. Values for children are presented in Table 2.2 and 2.3. The National Asthma Education and Prevention Program has used the results of FEV₁ and PEFR testing to classify the severity of asthma exacerbations (Table 2.4).

Multiple guidelines and articles have advocated specific cutoff values for PEFR and FEV₁ to guide decisions on disposition. There is variation across these guidelines and no consensus that absolute cutoffs should exist. Spirometric values including PEFR and FEV₁ should be viewed as additional data points to be considered, along with other clinical variables such as response to inhaled bronchodilators, in determining the disposition of asthmatics seen in the ED. Poor response to treatment or failure to recover PEFR or FEV₁ to greater than 70% of personal best are examples of factors that might favor admission rather than outpatient management of acute asthma exacerbations.

**NONINVASIVE OXYGENATION MONITORING: PULSE OXIMETRY**

Pulse oximetry, or noninvasive measurement of the percentage of hemoglobin bound to O₂, provides real-time estimates of arterial saturation in the range of 80% to 100% and gives early warning of diminished capillary perfusion while avoiding the discomfort and risks associated with arterial puncture. Pulse oximetry has become the standard of care in a wide variety of clinical settings.

**Technology**

Oximetry is based on the Beer-Lambert law, that states the concentration of an unknown solute dissolved in a solvent can be determined by light absorption. Pulse oximetry combines the principles of optical plethysmography and spectrophotometry. The probe, set into a reusable clip or a disposable patch, is made up of two photodiodes, which produce red light at 660 nm and infrared (IR) light at 900 to 940 nm, and a photodetector, which is placed across a pulsatile vascular bed. Pulse oximeter sensors are typically placed on the finger, with the light centered over the nailbed, not the fat pad. However, other locations such as the toes and earlobes are used in some settings such as pediatrics. These particular wavelengths are used because the absorption characteristics of oxyhemoglobin and reduced hemoglobin are quite different at the two wavelengths. The majority of the light is absorbed by connective tissue, skin, bones, and venous blood. The amount of light absorbed by these substances is constant with time and does not vary during the cardiac cycle. A small increase in arterial blood occurs with each heartbeat, thereby resulting in an increase in light absorption (Fig. 2.4). By comparing the ratio of pulsatile and baseline absorption at these two wavelengths, the ratio of oxyhemoglobin to reduced hemoglobin is calculated.

Because the pulse oximeter uses only two wavelengths of light, it can distinguish only two substances. Pulse oximeters measure “functional saturation”: the concentration of oxyhemoglobin divided by the concentrations of oxyhemoglobin plus reduced hemoglobin. The disadvantage of functional saturation is that the denominator does not include other hemoglobin species that may be present, such as carboxyhemoglobin and methemoglobin. The advantage of using only two wavelengths in the oximeter is that the cost, size, and weight of the device are reduced. Alternative devices, such as the CO-oximeter, use four or more wavelengths to measure “fractional saturation,” and are able to quantify additional hemoglobin species.

**Physiology**

Arterial O₂ saturation (SO₂) measures the large reservoir of O₂ carried by hemoglobin, 20 mL of O₂/100 mL of blood, although not all O₂, will be available for use by the tissues. The majority of arterial blood contains 6 to 7 mL of O₂, with the remainder bound to hemoglobin. A drop in arterial O₂ saturation is the most sensitive indicator of impending hypoxia, and is used to determine the need for oxygen therapy. Arterial O₂ saturation is determined by pulse oximetry as the percentage of arterial hemoglobin that is saturated with O₂. Although the pulse oximeter produces a reliable estimate of arterial O₂ saturation, it is limited by the presence of other substances in the light path that absorb light (Fig. 2.4). The absorption by tissue, skin, bones, and venous blood is significant, and must be subtracted from the measured absorption to determine arterial O₂ saturation.

![Figure 2.4 Factors influencing light absorption through a pulsatile vascular bed. (From McGough EK, Boysen PG: Benefits and limitations of pulse oximetry in the ICU. J Crit Illness 4:23, 1989.)](image)
TABLE 2.2  Predicted Peak Expiratory Flow Rate in Males 8-20 Years of Age

<table>
<thead>
<tr>
<th>AGE (YEAR)</th>
<th>ETHNICITY</th>
<th>Height (Inches)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Caucasian</td>
<td>160 178 197 217</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>143 163 184 205</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>152 174 197 221</td>
</tr>
<tr>
<td>9</td>
<td>Caucasian</td>
<td>— 184 203 223</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— 165 186 207</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— 177 200 224</td>
</tr>
<tr>
<td>10</td>
<td>Caucasian</td>
<td>— — 210 230</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — 190 211</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — 205 229</td>
</tr>
<tr>
<td>11</td>
<td>Caucasian</td>
<td>— — — 239 260</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — 217 240</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — 235 260</td>
</tr>
<tr>
<td>12</td>
<td>Caucasian</td>
<td>— — — — 271 292</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — 248 271</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — 268 294</td>
</tr>
<tr>
<td>13</td>
<td>Caucasian</td>
<td>— — — — — 304 326</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — — 282 306</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — — 304 331</td>
</tr>
<tr>
<td>14</td>
<td>Caucasian</td>
<td>— — — — — — 340 363</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — — — 319 344</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — — — 342 370</td>
</tr>
<tr>
<td>15</td>
<td>Caucasian</td>
<td>— — — — — — — 378 402</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — — — — 358 384</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — — — — 384 412</td>
</tr>
<tr>
<td>16</td>
<td>Caucasian</td>
<td>— — — — — — — — 419 443</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — — — — — 401 428</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — — — — — 427 457</td>
</tr>
<tr>
<td>17</td>
<td>Caucasian</td>
<td>— — — — — — — — — 462 487</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — — — — — — 447 475</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — — — — — — 474 504</td>
</tr>
<tr>
<td>18</td>
<td>Caucasian</td>
<td>— — — — — — — — — — 482 507</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — — — — — — — 469 496</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — — — — — — — 492 523</td>
</tr>
<tr>
<td>19</td>
<td>Caucasian</td>
<td>— — — — — — — — — — — 504 529</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>— — — — — — — — — — — 492 520</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>— — — — — — — — — — — 513 543</td>
</tr>
<tr>
<td>20</td>
<td>Caucasian</td>
<td>— — — — — — — — — — — — 527 552</td>
</tr>
</tbody>
</table>

### TABLE 2.3 Predicted Peak Expiratory Flow Rate in Females 8-18 Years of Age

<table>
<thead>
<tr>
<th>AGE (YEAR)</th>
<th>ETHNICITY</th>
<th>46</th>
<th>48</th>
<th>50</th>
<th>52</th>
<th>54</th>
<th>56</th>
<th>58</th>
<th>60</th>
<th>62</th>
<th>64</th>
<th>66</th>
<th>70</th>
<th>72</th>
<th>74</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>Caucasian</td>
<td>162</td>
<td>175</td>
<td>190</td>
<td>204</td>
<td>220</td>
<td>232</td>
<td>244</td>
<td>256</td>
<td>268</td>
<td>280</td>
<td>292</td>
<td>304</td>
<td>316</td>
<td>328</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>166</td>
<td>180</td>
<td>195</td>
<td>211</td>
<td>227</td>
<td>244</td>
<td>260</td>
<td>276</td>
<td>292</td>
<td>308</td>
<td>324</td>
<td>340</td>
<td>356</td>
<td>372</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>164</td>
<td>180</td>
<td>197</td>
<td>214</td>
<td>233</td>
<td>252</td>
<td>272</td>
<td>292</td>
<td>312</td>
<td>332</td>
<td>352</td>
<td>374</td>
<td>396</td>
<td>418</td>
</tr>
<tr>
<td>9</td>
<td>Caucasian</td>
<td>195</td>
<td>209</td>
<td>223</td>
<td>239</td>
<td>255</td>
<td>271</td>
<td>287</td>
<td>303</td>
<td>320</td>
<td>338</td>
<td>356</td>
<td>375</td>
<td>395</td>
<td>416</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>190</td>
<td>205</td>
<td>221</td>
<td>237</td>
<td>254</td>
<td>271</td>
<td>287</td>
<td>303</td>
<td>320</td>
<td>338</td>
<td>356</td>
<td>375</td>
<td>395</td>
<td>416</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>195</td>
<td>212</td>
<td>229</td>
<td>248</td>
<td>267</td>
<td>286</td>
<td>306</td>
<td>325</td>
<td>345</td>
<td>365</td>
<td>386</td>
<td>408</td>
<td>430</td>
<td>452</td>
</tr>
<tr>
<td>10</td>
<td>Caucasian</td>
<td>226</td>
<td>241</td>
<td>256</td>
<td>272</td>
<td>288</td>
<td>305</td>
<td>324</td>
<td>343</td>
<td>362</td>
<td>382</td>
<td>403</td>
<td>425</td>
<td>448</td>
<td>471</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>225</td>
<td>243</td>
<td>261</td>
<td>280</td>
<td>300</td>
<td>320</td>
<td>341</td>
<td>363</td>
<td>385</td>
<td>408</td>
<td>432</td>
<td>458</td>
<td>485</td>
<td>513</td>
</tr>
<tr>
<td>11</td>
<td>Caucasian</td>
<td>256</td>
<td>271</td>
<td>287</td>
<td>303</td>
<td>320</td>
<td>338</td>
<td>356</td>
<td>375</td>
<td>395</td>
<td>415</td>
<td>436</td>
<td>458</td>
<td>481</td>
<td>504</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>240</td>
<td>257</td>
<td>273</td>
<td>291</td>
<td>309</td>
<td>328</td>
<td>348</td>
<td>368</td>
<td>389</td>
<td>410</td>
<td>432</td>
<td>455</td>
<td>478</td>
<td>502</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>255</td>
<td>273</td>
<td>292</td>
<td>312</td>
<td>332</td>
<td>353</td>
<td>374</td>
<td>395</td>
<td>416</td>
<td>438</td>
<td>461</td>
<td>484</td>
<td>509</td>
<td>534</td>
</tr>
<tr>
<td>12</td>
<td>Caucasian</td>
<td>284</td>
<td>300</td>
<td>317</td>
<td>334</td>
<td>351</td>
<td>369</td>
<td>387</td>
<td>406</td>
<td>426</td>
<td>447</td>
<td>468</td>
<td>489</td>
<td>512</td>
<td>535</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>266</td>
<td>283</td>
<td>301</td>
<td>319</td>
<td>337</td>
<td>356</td>
<td>375</td>
<td>395</td>
<td>416</td>
<td>438</td>
<td>460</td>
<td>482</td>
<td>505</td>
<td>528</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>283</td>
<td>302</td>
<td>322</td>
<td>342</td>
<td>363</td>
<td>385</td>
<td>407</td>
<td>429</td>
<td>452</td>
<td>475</td>
<td>498</td>
<td>522</td>
<td>546</td>
<td>570</td>
</tr>
<tr>
<td>13</td>
<td>Caucasian</td>
<td>311</td>
<td>328</td>
<td>345</td>
<td>362</td>
<td>380</td>
<td>399</td>
<td>418</td>
<td>438</td>
<td>459</td>
<td>480</td>
<td>502</td>
<td>524</td>
<td>547</td>
<td>570</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>293</td>
<td>310</td>
<td>329</td>
<td>347</td>
<td>366</td>
<td>386</td>
<td>407</td>
<td>428</td>
<td>450</td>
<td>472</td>
<td>494</td>
<td>516</td>
<td>539</td>
<td>562</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>311</td>
<td>331</td>
<td>351</td>
<td>372</td>
<td>393</td>
<td>416</td>
<td>439</td>
<td>463</td>
<td>488</td>
<td>512</td>
<td>537</td>
<td>563</td>
<td>590</td>
<td>617</td>
</tr>
<tr>
<td>14</td>
<td>Caucasian</td>
<td>337</td>
<td>354</td>
<td>371</td>
<td>389</td>
<td>408</td>
<td>428</td>
<td>449</td>
<td>470</td>
<td>492</td>
<td>514</td>
<td>537</td>
<td>561</td>
<td>585</td>
<td>610</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>320</td>
<td>338</td>
<td>357</td>
<td>376</td>
<td>396</td>
<td>417</td>
<td>439</td>
<td>462</td>
<td>485</td>
<td>508</td>
<td>531</td>
<td>555</td>
<td>580</td>
<td>605</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>338</td>
<td>358</td>
<td>379</td>
<td>401</td>
<td>423</td>
<td>446</td>
<td>471</td>
<td>495</td>
<td>520</td>
<td>545</td>
<td>571</td>
<td>597</td>
<td>624</td>
<td>651</td>
</tr>
<tr>
<td>15</td>
<td>Caucasian</td>
<td>344</td>
<td>361</td>
<td>378</td>
<td>397</td>
<td>415</td>
<td>435</td>
<td>456</td>
<td>478</td>
<td>501</td>
<td>524</td>
<td>547</td>
<td>571</td>
<td>596</td>
<td>621</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>330</td>
<td>348</td>
<td>367</td>
<td>386</td>
<td>406</td>
<td>426</td>
<td>447</td>
<td>468</td>
<td>491</td>
<td>513</td>
<td>535</td>
<td>558</td>
<td>581</td>
<td>605</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>343</td>
<td>363</td>
<td>384</td>
<td>406</td>
<td>428</td>
<td>451</td>
<td>475</td>
<td>499</td>
<td>524</td>
<td>549</td>
<td>574</td>
<td>599</td>
<td>625</td>
<td>651</td>
</tr>
<tr>
<td>16</td>
<td>Caucasian</td>
<td>366</td>
<td>383</td>
<td>402</td>
<td>420</td>
<td>440</td>
<td>460</td>
<td>481</td>
<td>503</td>
<td>525</td>
<td>547</td>
<td>569</td>
<td>592</td>
<td>616</td>
<td>641</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>358</td>
<td>377</td>
<td>396</td>
<td>416</td>
<td>436</td>
<td>457</td>
<td>478</td>
<td>500</td>
<td>522</td>
<td>543</td>
<td>565</td>
<td>587</td>
<td>609</td>
<td>634</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>367</td>
<td>388</td>
<td>408</td>
<td>429</td>
<td>450</td>
<td>472</td>
<td>494</td>
<td>517</td>
<td>540</td>
<td>562</td>
<td>585</td>
<td>608</td>
<td>631</td>
<td>656</td>
</tr>
<tr>
<td>17</td>
<td>Caucasian</td>
<td>369</td>
<td>386</td>
<td>405</td>
<td>423</td>
<td>443</td>
<td>463</td>
<td>483</td>
<td>504</td>
<td>526</td>
<td>547</td>
<td>568</td>
<td>590</td>
<td>612</td>
<td>637</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>368</td>
<td>386</td>
<td>406</td>
<td>426</td>
<td>446</td>
<td>467</td>
<td>488</td>
<td>509</td>
<td>531</td>
<td>552</td>
<td>573</td>
<td>595</td>
<td>616</td>
<td>640</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>370</td>
<td>391</td>
<td>412</td>
<td>435</td>
<td>458</td>
<td>482</td>
<td>506</td>
<td>530</td>
<td>553</td>
<td>577</td>
<td>599</td>
<td>623</td>
<td>648</td>
<td>673</td>
</tr>
<tr>
<td>18</td>
<td>Caucasian</td>
<td>370</td>
<td>387</td>
<td>406</td>
<td>424</td>
<td>444</td>
<td>465</td>
<td>487</td>
<td>509</td>
<td>532</td>
<td>554</td>
<td>576</td>
<td>598</td>
<td>621</td>
<td>645</td>
</tr>
<tr>
<td></td>
<td>African American</td>
<td>378</td>
<td>396</td>
<td>416</td>
<td>438</td>
<td>459</td>
<td>482</td>
<td>505</td>
<td>528</td>
<td>551</td>
<td>574</td>
<td>596</td>
<td>619</td>
<td>642</td>
<td>666</td>
</tr>
<tr>
<td></td>
<td>Mexican American</td>
<td>371</td>
<td>392</td>
<td>413</td>
<td>436</td>
<td>459</td>
<td>482</td>
<td>506</td>
<td>529</td>
<td>553</td>
<td>577</td>
<td>599</td>
<td>623</td>
<td>647</td>
<td>672</td>
</tr>
</tbody>
</table>

and arterial O₂ partial pressure (PaO₂) measures only the relatively small amount of O₂ dissolved in plasma, approximately 0.3 mL of O₂/100 mL of blood. SaO₂ correlates well with PaO₂, but the relationship is nonlinear and is described by the oxyhemoglobin dissociation curve (Fig. 2.5). In hypoxic patients, small changes in SaO₂ represent large changes in PaO₂ because these SaO₂ values fall on the steep portion of the curve. Conversely, measurements of SaO₂ are relatively insensitive in detecting significant changes in PaO₂ at high levels of oxygenation because these SaO₂ values fall on the plateau portion of the curve.

Currently available pulse oximeters are accurate and precise when saturation ranges from 70% to 100%. This range is satisfactory because for most patients an O₂ saturation of 80% is as much an urgent warning as is one lower than 70%. Testing of pulse oximeters has shown that at 75% saturation, bias is scattered uniformly between underestimation and overestimation.

**Clinical Utility**

Pulse oximetry peripheral oxygen saturation (SpO₂) offers an advantage in assessing the adequacy of oxygenation over arterial blood gas analysis by providing continuous measurements. Direct measurement of SaO₂ requires blood gas values coupled with knowledge of the actual hemoglobin levels in a patient’s blood. SaO₂ measurement is estimated with pulse oximetry. In this chapter we equate SaO₂ and SpO₂.

There are limited data on the clinical efficacy of pulse oximetry monitoring in the ED. Anesthesia studies have demonstrated that continuous monitoring of saturation can improve detection of hypoxemia and related events, though this practice has not been shown to impact major clinical outcomes such as mortality or transfer to the intensive care unit (ICU). Nonetheless, there may be benefit to use in emergency medicine, particularly in those with primary cardiopulmonary disease or critically ill patients. Rapid recognition of adverse physiologic events should allow prompt initiation of therapeutic interventions.

**Box 2.1 Clinical Applications of Pulse Oximetry**

Assessing the adequacy of preoxygenation before ET intubation  
Monitoring oxygenation during emergency airway management  
Monitoring the ventilator and changes in FiO₂  
Providing an early indicator of ventilator dysfunction  
Assisting in routine weaning from O₂ therapy  
Monitoring patients in acute respiratory distress  
Monitoring during procedural sedation and analgesia  
Monitoring during interhospital and intrahospital transport

**Indications**

Recommended uses for pulse oximetry fall into two broad categories: (1) as a real-time indicator of hypoxemia, continuous oximetry monitoring can be used as a warning system because many adverse patient events are associated with arterial desaturation, and (2) as an end point for titration of therapeutic interventions to avoid hypoxia (Box 2.1).

Pulse oximetry can also be used to assess peripheral perfusion and evaluate for possible ischemia in the extremities. Although clinical experience validates its use, minimal data are available for such utilization in the ED. Vascular surgeons will use a pulse oximetry probe on a finger or toe to assess the results of vascular surgery on the arm or leg. Peripheral artery occlusion from peripheral artery disease may be suggested by comparison of pulse oximetry readings in the extremities. Decreased peripheral oxygenation may be detected in patients with compartment syndrome, traumatic arterial injury, and external compression of the proximal circulation (Fig. 2.6).

**Procedure**

The location for the probe is determined by the clinical situation and the types of probes available (Fig. 2.7). A reusable clip-on probe works well on digits that are easily accessible. Other sites include the earlobe, the nasal bridge, the septum, the forehead/temporal artery, and the foot or palm of an infant. More central locations may provide better readings in cold ambient temperature or during movement. Tape and splints can be used to secure oximetry probes and minimize motion.

The computer analyzes the incoming data to identify the arteriolar pulsation and displays this parameter as beats per minute. Newer devices also display a pulse plethysmograph (Fig. 2.8). Simultaneously, O₂ saturation is displayed on a beat-to-beat basis. Some machines have hard-copy capability and can provide paper documentation of the patient’s status.

**Interpretation**

Patients with normal physiologic gas exchange have an O₂ saturation between 97% and 100%. When SaO₂ falls below 95%, hypoxemia may be present, which may be baseline for some patients with cardiac or lung disease. O₂ saturation below
Although pulse oximetry represents a significant advance in noninvasive monitoring of oxygenation, clinicians must recognize and understand its limitations. Pulse oximetry measures only O2 saturation. In contrast to arterial blood gas determination, pulse oximetry provides no direct information on pH or the arterial partial pressure of CO2 (PaCO2). Nonetheless, a room-air SaO2 value of 97% or higher strongly rules against hypoxemia and moderate to severe hypercapnia. A low SaO2 reading should be heeded as an important clinical warning sign. Pulse oximetry may be affected by numerous extrinsic factors, and a decline in O2 saturation with serial measurements should always prompt an evaluation of respiratory status and adequacy of circulation.

Figure 2.6 Pulse oximetry can be used to assess the distal circulation after vascular surgery for trauma and to initially evaluate other causes of decreased peripheral perfusion. A, This patient had a markedly swollen and ischemic finger from a tight ring. B, Although the need for immediate removal of the ring is clinically obvious, a pulse oximetry probe confirmed ischemia with an O2 saturation of 61%. The uninvolved fingers registered 99%. Following ring removal the saturation returned to normal, thus suggesting that fasciotomy need not be performed. C, When a discharged EpiPen caused a pale finger, injection of phentolamine was considered. D, When pulse oximetry demonstrated a saturation of 96% (97% to 98% in the other fingers), injection was not performed and the circulation spontaneously normalized over a period of 30 minutes.

Figure 2.7 Pulse oximeter sensors. A, Reusable adult sensor, attached to finger. B, Single-use adult sensor, attached to finger. C, Single-use pediatric sensor, attached to toe. Note that the light source is centered over the nail, not the fat pad. See text for various parameters that affect pulse oximetry readings.
room-air SaO2 value of 96% or less has been demonstrated to be 100% sensitive and 54% specific for detecting hypoxemia (PaO2 <70 mm Hg), and 100% sensitive and 31% specific for detecting hypercapnia (PaCO2 >50 mm Hg). Using a cutoff value of 92% or less for room-air SaO2 is more accurate in identifying hypoxemia in patients with COPD.

Pulse oximetry is not a substitute for monitoring ventilation because of the variable lag time between the onset of hypventilation or apnea and a change in O2 saturation. During procedural sedation, monitoring of ventilation is a more desirable goal for prevention of hypoxemia and hypercapnia than simple pulse oximetry (see section on Procedural Sedation and Analgesia under Carbon Dioxide Monitoring later in this chapter). Hypoventilation and the resultant hypercapnia may precede a decrease in hemoglobin O2 saturation by many minutes. Supplemental O2 may mask hypoventilation by delaying the eventual O2 desaturation that pulse oximetry is designed to monitor and recognize. Other limitations of pulse oximetry are summarized in Box 2.2.

Sources of Interference

Effects of Dyshemoglobinemias
In patients with methemoglobinemia or elevated carboxyhemoglobin levels, pulse oximetry does not accurately depict quantitative changes in hemoglobin O2 saturation. Carboxyhemoglobin results in falsely elevated SaO2 estimates of hemoglobin O2 saturation. Low quantities of methemoglobin will reduce pulse oximetry readings by approximately half the actual methemoglobin percentage. Large quantities of methemoglobin (>10%) can result in a stable pulse oximetry reading of 85% regardless of the actual SaO2. Because pulse oximetry will variably underestimate the percentage of abnormal hemoglobin, a CO-oximeter or blood gas sample for quantitative analysis is required for confirmation of these conditions.

Fetal Hemoglobin
Full-term newborns can have up to 75% of total hemoglobin in the form of fetal hemoglobin and up to 5% in the form of carboxyhemoglobin. Although fetal hemoglobin interferes with the spectrophotometric method, pulse oximetry will remain accurate. A CO-oximeter may erroneously interpret the carboxyhemoglobin level as elevated and the oxyhemoglobin level as artificially reduced. Therefore when fetal hemoglobin levels are high, CO-oximetry readings should not be used to confirm pulse oximetry readings.

Low Perfusion
To function properly, pulse oximeters require a pulsating vascular bed. Vasodilation related to hypotension, hypothermia, or the administration of vasoconstricting drugs may reduce the pulsatile component to less than 0.2% of the total signal. At this level, the true signal cannot be distinguished from background noise. Modern pulse oximeters have signal strength detectors and will display a message indicating an inadequate pulse signal when perfusion is compromised. Changing the location of the sensor to an area with higher perfusion, such as an earlobe or the forehead, may improve pulse oximetry readings.

Intravenous Dyes
A number of dyes and pigments interfere with the accuracy of pulse oximetry. Methylene blue, the treatment of methemoglobinemia, absorbs light at 660 nm, which is similar to the absorption of reduced hemoglobin, and can significantly decrease pulse oximeter saturation readings to as low as 1%. Low readings can also be seen with other intravenous dyes such as indigo carmine, indocyanine green, and fluorescein, although the rapid clearance of these agents minimizes this phenomenon.
Bilirubin
Hyperbilirubinemia does not affect the accuracy of pulse oximetry, but it may have an effect on absorption at the lower wavelengths used by CO-oximeters and result in a discrepancy between pulse oximeter and CO-oximeter readings.

Skin Pigmentation
The accuracy of pulse oximeters is somewhat reduced by darkly pigmented skin. This effect is probably due to a shift in the light-emitting diode’s output spectrum as the light output is increased. This effect is small and results in only a slight decrease in accuracy. Placing the probe on an area of lighter pigmentation, such as the fifth finger or an earlobe, is suggested to minimize this effect.

Nail Polish
Data suggest that some nail polishes may affect the accuracy of pulse oximetry.28,29 Turning the probe sideways on the finger may improve accuracy, although removing any nail polish with acetone is the favored solution. The accuracy of SaO₂ readings in the setting of synthetic nails is unknown. If a poor signal is obtained through a synthetic nail, either the nail should be removed or an alternative site should be used for placement.

High Saturation
Because the O₂ dissociation curve plateaus at saturation levels greater than 90%, a large increase in PaO₂ results in only a small increase in saturation. An error of a few percentage points could represent a large error in PaO₂. This is inconsequential for most adult patients and is of extreme importance for neonates, who are at risk for retinopathy caused by hyperoxemia.

Venous Pulsations
The increased venous pulsations resulting from right heart failure and tricuspid regurgitation can interfere with accurate readings and lead to artificially lower O₂ saturation because the pulse oximeter interprets any pulsatile measurement as arterial. Placing the probe on a site above the heart may improve accuracy. Some pulse oximeters have the capability of synchronizing pulsations at the probe site to electrocardiographic (ECG) signals, thus enhancing the signal-to-noise ratio.

Anemia
Because pulse oximetry depends on light absorption by hemoglobin, it becomes less accurate and less reliable in conditions of severe anemia.

Ambient Light
Because the pulse oximeter’s photodetector is nonspecific, high-intensity ambient light can produce interference. Surgical, fluorescent, and heating lamps are common sources of light interference. This problem can be corrected by wrapping the probe with a light barrier, such as a dark cloth or other opaque material.

Motion
Motion of the probe can produce considerable artifact and inaccurate readings. Correlating a pulse oximetry signal with an ECG waveform or using alternative probe sites, such as the ear or toe, may reduce motion artifact. Newer-generation pulse oximetry signal-processing technology has greatly reduced the effect of motion artifact.

Probe Site
The finger is the most common probe site used for adult pulse oximetry. If the finger is inaccessible or unsuitable, other probe sites, such as the earlobe, nose, and forehead (using reflectance instead of transmittance), may be used. It should be noted, though, that forehead and nasal bridge probes may be less accurate than finger and ear probes. In infants and small children, an adhesive sensor unit is preferred and frequently placed on the fingers and toes. Probes can also be secured in place over the palm, heel, or lateral aspect of the foot with a gauze or wrap.

Electrocautery
Electrical interference from devices such as electrocautery can impair the accuracy of pulse oximetry. Such interference can be reduced by increasing the distance between the surgical site and the probe.

Conclusions
Pulse oximetry is a widely available technology that provides an easy, noninvasive, and generally reliable method of monitoring oxygenation. Because measurements are continuous, pulse oximetry allows earlier detection of hypoxic episodes than intermittent arterial blood gas analysis does. Frequent measurements can lead to earlier corrective measures and prevention of adverse consequences.

Future Directions
New devices that focus on regional detection of O₂ saturation, specifically cerebral oximetry, are being studied for various clinical applications relevant to emergency medicine. This technology uses near-IR spectroscopy to obtain tissue level O₂ saturations. Data are emerging to support cerebral oximetry monitoring to assess quality of cardiopulmonary resuscitation (CPR) and to predict return of spontaneous circulation for these patients.11–13 Early studies have also suggested a role for cerebral oximetry in monitoring patients with altered mental status, including pharmacologically induced sedation.16–17 Currently there are no clear guidelines on when this technology should be used in an ED setting, and its growing experience may direct future applications in end-organ perfusion monitoring.

CO₂ MONITORING
Capnography is a noninvasive measurement of the partial pressure of CO₂ in an exhaled breath. Measurement of CO₂ at the airway can be displayed as a function of time (CO₂ concentration over time) or as an exhaled tidal volume (CO₂ concentration over volume). This chapter focuses on the use of time-based capnography because it can be adapted for use in both intubated and nonintubated patients, and therefore is the only form of CO₂ monitoring used by the emergency medical service (EMS) and the predominant form used in the ED.

The relationship of CO₂ concentration to time is graphically represented by the CO₂ waveform or capnogram (Fig. 2.9). The maximum CO₂ concentration at the end of each tidal breath is the end-tidal CO₂ pressure (PETCO₂). Changes in the shape of the capnogram are diagnostic of disease conditions,
ETCO\textsubscript{2} as displays a number and a waveform (i.e., the capnogram). This graph. CO\textsubscript{2} pressure at molecules absorb IR radiation at a very specific wavelength close to exponential relationship to the CO\textsubscript{2} concentration in vascular system); and metabolism (how effectively CO\textsubscript{2} is being produced by cellular metabolism). Information about all of the following: ventilation (how breathing patients. Pulse oximetry provides real-time feedback on oxygenation and ventilation are distinct physiologic functions that are assessed in both intubated and spontaneously breathing patients. Pulse oximetry provides real-time feedback about oxygenation, and capnography provides breath-to-breath information about all of the following: ventilation (how effectively CO\textsubscript{2} is being eliminated by the pulmonary system); perfusion (how effectively CO\textsubscript{2} is being transported through the vascular system); and metabolism (how effectively CO\textsubscript{2} is being produced by cellular metabolism).

Terminology

The ancient Greeks believed there was a combustion engine inside the body that gave off smoke (\textit{capnos} in Greek) in the form of a breath. A capnometer is a CO\textsubscript{2} monitor that displays a number (i.e., PETCO\textsubscript{2}). A capnograph is a CO\textsubscript{2} monitor that displays a number and a waveform (i.e., the capnogram).

Technology

Capnography became a routine part of anesthesia practice in Europe in the 1970s and in the United States in the 1980s. Capnography was incorporated into the American Heart Association (AHA) guidelines in 2000 and the American College of Emergency Physicians (ACEP) guidelines in 2001 and has become the standard of care for verification of endotracheal (ET) tube placement.

Most capnography technology is built on IR radiation techniques. These techniques are based on the fact that CO\textsubscript{2} molecules absorb IR radiation at a very specific wavelength (4.26 \textmu m), with the amount of radiation absorbed having a close to exponential relationship to the CO\textsubscript{2} concentration present in the breath sample. Detecting these changes in IR radiation levels by using appropriate photodetectors sensitive in this spectral region allows the CO\textsubscript{2} concentration in the gas sample to be calculated.

CO\textsubscript{2} monitors measure gas concentration or partial pressure by using one of two configurations, depending on the location of the sensor: mainstream or sidestream (Fig. 2.10). Mainstream devices measure CO\textsubscript{2} directly from the airway, with the sensor located at the proximal end of the ET tube. Sidestream devices measure CO\textsubscript{2} by aspirating a small sample from the exhaled breath through tubing and a sensor located inside the monitor. Mainstream systems are configured only for intubated patients because the sensor is located on the ET tube. Sidestream systems do not require an ET tube because the sensor is located inside the monitor and may therefore be used in either intubated or nonintubated patients. The airway interface for intubated patients is an airway adapter placed on the hub of the ET tube. For spontaneously breathing patients, a nasal-oral cannula is used. This allows concomitant CO\textsubscript{2} sampling and delivery of low-flow O\textsubscript{2} to the patient.

Sidestream systems can be high-flow (requiring 150 mL/min of CO\textsubscript{2} in the breath sample to obtain an accurate reading) or low-flow (requiring 50 mL/min of CO\textsubscript{2}). Low-flow sidestream systems have a lower occlusion rate (from moisture or patient secretions) and are more accurate in patients with low tidal volumes (neonates, infants, and patients with hypoventilation and low–tidal volume breathing).

CO\textsubscript{2} monitors can be either quantitative or qualitative (Fig. 2.11). Quantitative devices measure the precise PETCO\textsubscript{2} as either a number (capnometry) or a number and a waveform (capnography). Qualitative devices measure the range in which PETCO\textsubscript{2} falls (e.g., 0 to 10 mm Hg, >35 mm Hg) as opposed to a precise value (e.g., 38 mm Hg). The most commonly used qualitative device is the colorimetric PETCO\textsubscript{2} detector, which consists of a piece of specially treated litmus paper that turns color when exposed to CO\textsubscript{2}. Its primary use is for verification of ET tube position. If the tube is in the trachea, the resultant exhalation of CO\textsubscript{2} will change the color of the litmus paper; if the tube is in the esophagus with no CO\textsubscript{2} in the breath, no change in color will take place.
CHAPTER 2 Devices for Assessing Oxygenation and Ventilation

with PETCO₂ accurately reflecting PaCO₂. Patients with obstructive lung disease will have a more rounded ascending phase and an upward slope in the alveolar plateau (Fig. 2.12). In patients with abnormal lung function from V/Q mismatch, the gradient will widen, depending on the severity of the lung disease, and PETCO₂ will be useful only for trend ventilatory status over time and not as a spot check because it may not correlate with PaCO₂.

Indications for Intubated Patients

- Verification of ET tube placement
- Continuous monitoring of tube location during transport
- Gauging the effectiveness of resuscitation and prognosis during cardiac arrest
- Titrating PETCO₂ levels in patients with suspected increases in intracranial pressure
- Determining prognosis in patients after trauma
- Determining the adequacy of ventilation

Verification of ET Tube Placement

Unrecognized misplaced intubation (UMI) is placement of an ET tube in a location other than the trachea that is not recognized by the clinician. This life-threatening condition has been extensively documented in the EMS literature, with reported UMI rates most commonly between 7% to 10%.

After intubation, the presence of a waveform with all four phases indicates that the ET tube is through the vocal cords. A flatline waveform following intubation indicates esophageal placement except in selected conditions, including obstruction of the ET tube, complete airway obstruction distal to the tube, tracheal placement with inadequate pulmonary blood flow as a result of poor chest compressions, or prolonged cardiac arrest with no circulating CO₂ because of cessation of cellular metabolism.

The accuracy of PETCO₂ in confirming the tracheal location of an ET tube varies according to the type of CO₂ technology used. In patients who are not in cardiac arrest, qualitative colorimetric PETCO₂ and quantitative capnography studies have demonstrated 100% sensitivity and specificity for tracheal placement. In marked contrast, the use of clinical signs for verification has been shown to be unreliable. Fogging or condensation of the tube, chest wall movement, and auscultation

**Physiology**

The capnogram, which corresponds to a single tidal breath, consists of four phases (ascending phase, alveolar plateau, inspiratory limb, dead space ventilation) (see Fig. 2.9). Each phase has conventionally been approximated as a straight line. Phase I (dead space ventilation, A to B) represents the beginning of exhalation in which dead space is cleared from the upper airway. Phase II (ascending phase, B to C) represents the rapid rise in CO₂ concentration in the breath stream as CO₂ from the alveoli reaches the upper airway. Phase III (alveolar plateau, C to D) represents the CO₂ concentration reaching a uniform level in the entire breath stream (from alveolus to nose) and concludes with a point of maximum CO₂ pressure (PETCO₂). This is the number that appears on the monitor display. Phase IV (D to E) represents the inspiratory cycle in which the CO₂ concentration drops to zero as atmospheric air enters the airway.

A normal capnogram, for patients of all ages, is characterized by a specific set of elements: it includes the four distinct phases just described, the CO₂ concentration starts at zero and returns to zero (i.e., there is no rebreathing of CO₂), a maximum CO₂ concentration is reached with each breath (i.e., PETCO₂), the amplitude is dependent on PETCO₂, the width is dependent on the expiratory time, and there is a characteristic shape for all subjects with normal lung function.

Patients with normal lung function, irrespective of age, will have a characteristic rectangular- or trapezoidal-shaped capnogram and a narrow PETCO₂-PaCO₂ gradient (0 to 5 mm Hg),

![Figure 2.11 CO₂ monitors. A, Quantitative monitor. A capnography waveform (arrow) is displayed, as is a capnometry numerical reading (37). B, Qualitative device. This simple colorimetric detector is used to verify endotracheal tube position and changes color when exposed to CO₂.](image)

![Figure 2.12 Capnogram shape in A, normal subjects and B, patients with chronic obstructive pulmonary disease (COPD). (From Krauss B, Deykin A, Lam A, et al: Capnogram shape in obstructive lung disease, Anesth Analg 100:884, 2005.)](image)
for breath sounds have all been demonstrated to incorrectly identify tube location in some cases, including esophageal intubations.

Although the accuracy of PETCO₂ in verifying ET tube placement is 100% in patients with spontaneous circulation or low-perfusion states, sensitivity for tracheal placement in cardiac arrest patients ranges from 62% to 100%, depending on the type of CO₂ monitoring used and the duration of the arrest. Data are limited regarding specificity of capnography to detect esophageal intubations in cardiac arrest. When a waveform is present in an intubated patient in cardiac arrest, the ET tube can be assumed to be in the trachea. Absence of a waveform may result from esophageal intubation or a correctly placed ET tube in a patient with insufficient pulmonary blood flow.

Colorimetric studies have shown variable sensitivity because the exhaled CO₂ concentration can fall below the detection threshold. It is particularly important when evaluating PETCO₂ studies to distinguish those involving qualitative colorimetric detection from those using capnography.

**Monitoring Tube Position During Transport**

UMI (as a result of either initial misplacement of the ET tube or subsequent dislodgment during transport) can have catastrophic consequences and is largely preventable. Capnography has been demonstrated to help prehospital providers recognize ET tube dislodgement leading to more rapid correction of tube position. Continuous monitoring of tube position during transport (prehospital to hospital, interhospital, or intrahospital) is essential for patient safety. PETCO₂ confirmation of initial ET tube placement with continuous monitoring of tube position is an accepted standard of care by the American Society of Anesthesiologists and is recommended by other national organizations.

**Gauging the Effectiveness of CPR**

Studies using animal models in the 1980s demonstrated that PETCO₂ levels reflect cardiac output during CPR and can be used as a noninvasive measure of cardiac output. A landmark study in 1988 demonstrated this principle in humans (Fig. 2.13). During cardiac arrest, when alveolar ventilation and metabolism are essentially constant, PETCO₂ reflects the degree of pulmonary blood flow. Therefore PETCO₂ can be used as a gauge of the effectiveness of cardiac compressions. Effective cardiac compression leads to higher cardiac output, and the resultant increase in perfusion corresponds to a rise in PETCO₂ from baseline. The 2010 AHA Consensus statement on CPR Quality, endorsed by the ACEP, recommends monitoring end-tidal CO₂ (ETCO₂) during CPR to improve chest compression performance if ETCO₂ is <10 mm Hg and, by expert opinion, titrating to a goal ETCO₂ of >20 mm Hg.

**Indicator of ROSC**

A peak in PETCO₂ is the earliest sign of return of spontaneous circulation (ROSC) and may occur before palpable or measurable hemodynamic signs (pulse or blood pressure). When the heart is restarted, the dramatic increase in cardiac output and the resulting increase in perfusion lead to a rapid increase in PETCO₂ from baseline as the CO₂ that has built up in the blood during cardiac arrest is effectively transported to the lungs and exhaled.

The AHA guidelines emphasize the importance of continuing chest compressions without interruption until a perfusing rhythm is reestablished. Experimental evidence indicates that interruptions in chest compressions are followed by sustained periods during which flow gradually returns to pre-interruption levels. Capnography provides a means of monitoring the patient’s physiological response to ongoing resuscitative efforts with minimal interruption to compressions for the purpose of checking for pulses. Reestablishment of a perfusing rhythm will be immediately accompanied by a dramatic increase in PETCO₂.

**Assessing Prognosis After Initiation of Cardiac Arrest Resuscitation**

Multiple factors can affect the recorded ETCO₂ values during cardiac arrest resuscitations. PETCO₂ can be used as a prognostic indicator of ROSC and survival in adult cardiac arrest patients. In multiple studies, PETCO₂ levels of 10 mm Hg or lower measured 20 minutes after the initiation of advanced cardiac life support accurately predicted death in patients with cardiac arrest. This prognostic value of measuring PETCO₂ has been demonstrated in both animal and human studies.

**Identifying the Cause of Cardiac Arrest**

Though not generally used clinically, differences in PETCO₂ may be seen depending on the cause of the cardiac arrest. Animal studies reported higher PETCO₂ values at the onset of cardiac arrest caused by primary asphyxia than after arrest caused by ventricular fibrillation. Prehospital cardiac arrest studies have demonstrated higher PETCO₂ results in patients with cardiac arrest secondary to respiratory or asphyxia causes than for those with primary cardiac etiologies.

**Titrating ETCO₂ in Patients With Suspected Increased Intracranial Pressure**

PETCO₂ monitoring has been shown to play a role in controlled ventilation in patients with head injury and suspected increased intracranial pressure. CO₂ levels affect blood flow to the brain, with high CO₂ levels resulting in cerebral vasodilation and
low CO₂ levels resulting in cerebral vasoconstriction. Sustained hyperventilation (PETCO₂ ≥50 mm Hg) is detrimental to patients with increased intracranial pressure because it results in increased cerebral blood flow and potential worsening of intracranial pressure.

Sustained hyperventilation is also detrimental and associated with worse neurologic outcome in severely brain-injured patients. Unless a patient is actively herniating, ventilation with CO₂ monitoring to achieve normocapnia is recommended. The benefit of continuous PETCO₂ monitoring for this indication has been demonstrated in many prehospital-based studies. Severe head-injury patients monitored with continuous PETCO₂ have better controlled ventilation, linked to improved outcome in traumatic brain injury patients.

PETCO₂ monitoring has also demonstrated prognostic value in determining outcome in trauma victims requiring prehospital ET intubation. Specifically, lower PETCO₂ levels 20 minutes post intubation are associated with poor survival.

**Indications for Capnography in Spontaneously Breathing Patients**

In spontaneously breathing, nonintubated patients, capnography can be used in a variety of clinical indications, including:
- Rapid assessment of critically ill, injured, or seizing patients through assessment of the airway, breathing, and circulation (ABCs)
- Assessment and triage of victims of chemical terrorism and mass casualty
- Gauging the severity and response to treatment in patients with acute respiratory distress
- Determining the adequacy of ventilation in patients with altered mental status
- Detecting metabolic acidosis (including diabetic ketoacidosis and gastroenteritis)

**Assessment of Critically Ill, Injured, or Seizing Patients**

The ABCs of critically ill or injured patients can be assessed rapidly by using the capnogram and PETCO₂. The presence of a normal waveform denotes a patent airway and spontaneous breathing. Normal PETCO₂ (35 to 45 mm Hg) signifies adequate perfusion.

Capnography can be used to assess and triage critically ill or injured patients and actively seizing patients. Unlike pulse oximetry, capnography is not affected by motion artifact and provides reliable readings in low-perfusion states.

Capnography is a reliable, accurate monitoring modality for actively seizing patients. Capnographic data (respiratory rate [RR], PETCO₂, and capnogram) can be used to distinguish among:
- Seizing patients with apnea (flatline waveform, no PETCO₂ readings, and no chest wall movement)
- Seizing patients with ineffective ventilation (small waveforms, low PETCO₂)
- Seizing patients with effective ventilation (normal waveform, normal PETCO₂)

**Assessment and Triage of Victims of Chemical Terrorism and Mass Casualty**

EDs and EMS systems have focused on training to identify and effectively manage mass casualty and chemical terrorism events. Capnography can serve as a noninvasive assessment tool to quickly identify the common life-threatening complications of chemical terrorism. It can rapidly detect the common airway, respiratory, and central nervous system adverse events associated with nerve agents, including apnea, upper airway obstruction, laryngospasm, bronchospasm, respiratory failure, seizures, and coma (Table 2.5).

**Gauging Severity and Response to Treatment of Patients in Acute Respiratory Distress**

Capnography provides dynamic monitoring of ventilatory status in patients with acute respiratory distress from any cause, including asthma, bronchiolitis, CHF, COPD, croup, and cystic fibrosis. By measuring PETCO₂ and RR with each breath, capnography provides immediate information on the clinical status of the patient. RR is measured directly from the airway (nose and mouth) with an oral-nasal cannula and provides a more reliable reading than does impedance respiratory monitoring. In upper airway obstruction and laryngospasm, impedance monitoring detects chest wall movement, interprets this as valid breathing, and displays an RR even though the patient is not ventilating. In contrast, capnography will detect absence of air movement and therefore shows a flatline waveform.

PETCO₂ trends can be assessed rapidly, especially in tachypneic patients. For example, a patient with an RR of 30 breaths/min will generate 150 PETCO₂ readings in 5 minutes. This provides sufficient information to determine whether the patient’s ventilation is worsening despite treatment (increasing PETCO₂), stabilizing (stable PETCO₂), or improving (decreasing PETCO₂) (Fig. 2.14).

**Procedural Sedation and Analgesia**

Pulse oximetry is the standard technique for monitoring procedural sedation in the ED, but capnography is more likely...
### TABLE 2.5 Capnographic Identification of Life-Threatening Complications of Nerve Agents

<table>
<thead>
<tr>
<th>AGENT</th>
<th>EFFECTS</th>
<th>CAPNOGRAPHY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nerve Gas</td>
<td>Seizures, diaphragmatic weakening or paralysis, hypoventilation, respiratory depression, apnea, loss of consciousness/coma</td>
<td>• Accurate readings during seizure activity (RR, PETCO2, capnogram)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Earliest indicator of respiratory compromise</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Direct measure of ventilatory status</td>
</tr>
<tr>
<td>Nerve Gas</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Tabun</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Sarin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Soman</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• VX</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vesicants</td>
<td>Airway edema, upper airway obstruction, bronchospasm</td>
<td>• Rapid identification of upper airway obstruction</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Rapid identification of bronchospasm</td>
</tr>
<tr>
<td>Vesicants</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Mustard gas</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Lewisite</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Choking Agents</td>
<td>Rapid, progressive, noncardiogenic pulmonary edema and acute lung injury, bronchospasm, laryngospasm</td>
<td>• Earliest indicator of respiratory compromise</td>
</tr>
<tr>
<td>Choking Agents</td>
<td></td>
<td>• Rapid identification of bronchospasm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Rapid identification of laryngospasm</td>
</tr>
<tr>
<td>Choking Agents</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Chlorine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Phosgene</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Diphosgene</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Chloropicrin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Ricin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cyanide</td>
<td>Sudden loss of consciousness/coma, seizures, metabolic acidosis with tachypnea, apnea</td>
<td>• Direct measure of ventilatory status</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Accurate readings during seizure activity</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Earliest indicator of respiratory compromise</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Noninvasive identification of metabolic acidosis</td>
</tr>
<tr>
<td>Incapacitating Agents</td>
<td>Laryngospasm, bronchospasm, respiratory failure</td>
<td>• Rapid identification of laryngospasm</td>
</tr>
<tr>
<td>Incapacitating Agents</td>
<td></td>
<td>• Rapid identification of bronchospasm</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Earliest indicator of respiratory compromise</td>
</tr>
<tr>
<td>Incapacitating Agents</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Lacrimators</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Mace)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Capsaicin</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*PETCO₂*, End-tidal carbon dioxide pressure; *RR*, respiratory rate.

Capnography may be used as an adjunct to pulse oximetry and clinical assessment to detect hypoventilation and apnea earlier than pulse oximetry and/or clinical assessment alone in patients undergoing procedural sedation and analgesia in the ED.

Capnography is the earliest indicator of airway or respiratory compromise and will show an abnormally high or low PETCO2 well before pulse oximetry detects a failing oxyhemoglobin saturation, especially in patients receiving supplemental oxygen (Fig. 2.15). Capnography provides a non–impedance-based RR directly from the airway, which is more accurate than impedance-based respiratory monitoring, especially in patients with obstructive apnea or laryngospasm.

Both central and obstructive apnea can be detected almost instantaneously by capnography (Table 2.6). Loss of the capnogram, in conjunction with no chest wall movement and no breath sounds on auscultation, confirms the diagnosis of central apnea. Obstructive apnea is characterized by loss of the capnogram with continued chest wall movement and absent breath sounds. The response of the capnogram to airway alignment maneuvers can further distinguish upper airway obstruction from laryngospasm.

Capnography is more sensitive than clinical assessment of ventilation in detecting apnea along with other respiratory complications. This leads to more awareness of respiratory complications associated with procedural sedation, which is followed by appropriate and timely interventions.

Because the amplitude of the capnogram is determined by PETCO2 and the width is determined by the expiratory time, changes in either of these parameters affect the shape of the capnogram. Hyperventilation (increased RR, decreased PETCO2) results in a low-amplitude and narrow capnogram. Classic hypoventilation (decreased RR, increased PETCO2) results in a high-amplitude and wide capnogram (see Table 2.6). Acute bronchospasm results in a capnogram with a curved ascending phase and an upsloping alveolar plateau (Fig. 2.12). A PETCO2 reading higher than 70 mm Hg in patients without chronic ventilation problems indicates respiratory failure.

Two types of drug-induced hypoventilation occur during procedural sedation and analgesia (see Table 2.6). Bradypneic hypoventilation (type 1), commonly seen with opioids, is characterized by increased PETCO2 and increased PaCO2. RR is depressed proportionally greater than tidal volume, resulting in bradypnea, an increase in expiratory time, and a rise in PETCO2, graphically represented by a high-amplitude, wide capnogram (see Table 2.6).

Bradypneic hypoventilation follows a predictable course, with PETCO2 increasing progressively until respiratory failure and apnea occur. Although there is no absolute threshold at which apnea occurs, patients with acute increases in PETCO2 to above 80 mm Hg are at significant risk.

Hypopneic hypoventilation (type 2), commonly seen with sedative-hypnotic drugs, is characterized by normal or decreased PETCO2 and increased PaCO2 because airway dead space remains constant (e.g., 150 mL in the normal adult lung) and tidal volume decreases. Tidal volume is depressed proportionally greater than RR, thereby resulting in low-tidal volume breathing and leading to an increase in the fraction of airway dead space (dead space volume/tidal volume). As tidal volume decreases, the airway dead space fraction increases, resulting in an increase in the PaCO2–PETCO2 gradient. Even though PaCO2 is increasing, PETCO2 may remain normal or decrease, graphically represented by a low-amplitude capnogram.

Hypopneic hypoventilation follows a variable course. Three possibilities exist: (1) ventilation may remain stable with the low–tidal volume breathing resolving over time as drug levels in the central nervous system decrease following redistribution, (2) hypoventilation may progress to periodic breathing with intermittent apneic pauses (which may resolve spontaneously or progress to central apnea), or (3) hypoventilation may progress directly to central apnea.

The low–tidal volume breathing that characterizes hypopneic hypoventilation increases dead space ventilation as a result of inhibition of the normal compensatory mechanisms by drug effects. Minute ventilation, which normally increases to compensate for an increase in dead space, does not change or may decrease. As minute ventilation decreases, arterial oxygenation decreases. PETCO2 may initially be high (bradypneic hypoventilation) or low (hypopneic hypoventilation) without significant changes in oxygenation, particularly if supplemental O2 is given. Therefore a drug-induced increase or decrease in PETCO2 does not necessarily lead to O2 desaturation and may not require intervention.

Determining the Adequacy of Ventilation in Patients With Altered Mental Status

Patients with altered mental status, including those with alcohol intoxication, intentional or unintentional drug overdose, patients requiring chemical restraint, and postictal patients (especially those treated with benzodiazepines), may have impaired ventilatory function. Capnography can differentiate between patients with effective ventilation and those with ineffective ventilation, as well as provide continuous monitoring of ventilatory trends over time to identify patients at risk for worsening respiratory depression.

Detection of Metabolic Acidosis

In addition to its established uses for assessment of ventilation and perfusion, capnography is a valuable tool for assessing metabolic status by providing information on how effectively CO2 is being produced by cellular metabolism.

Studies have shown that PETCO2 and serum bicarbonate (HCO3) are well correlated in patients with diabetes and gastroenteritis. PETCO2 can be used as an indicator of metabolic
### TABLE 2.6 Capnographic Airway Assessment for Procedural Sedation and Analgesia

<table>
<thead>
<tr>
<th>DIAGNOSIS</th>
<th>WAVEFORM</th>
<th>FEATURES</th>
<th>INTERVENTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td><img src="image" alt="Normal Waveform" /></td>
<td>Spo₂, PETCO₂, Waveform, RR</td>
<td>Normal, Normal, Normal, Normal, Normal</td>
</tr>
<tr>
<td>Hyperventilation</td>
<td><img src="image" alt="Hyperventilation Waveform" /></td>
<td>Spo₂, PETCO₂, Waveform, RR</td>
<td>Normal, Decreased amplitude and width ↑</td>
</tr>
<tr>
<td>Bradypneic hypoventilation (type 1)</td>
<td><img src="image" alt="Bradypneic Hypoventilation Waveform" /></td>
<td>Spo₂, PETCO₂, Waveform, RR</td>
<td>Normal, ↑, Increased amplitude and width ↓↓↓</td>
</tr>
<tr>
<td>Hypopneic hypoventilation (type 2)</td>
<td><img src="image" alt="Hypopneic Hypoventilation Waveform" /></td>
<td>Spo₂, PETCO₂, Waveform, RR</td>
<td>Normal, Decreased amplitude ↓</td>
</tr>
<tr>
<td>Hypopneic hypoventilation with periodic breathing</td>
<td><img src="image" alt="Hypopneic Hypoventilation with Periodic Breathing Waveform" /></td>
<td>Spo₂, PETCO₂, Waveform, RR</td>
<td>Normal or ↓, Decreased amplitude ↓, Apneic pauses</td>
</tr>
</tbody>
</table>

*Continued*
### TABLE 2.6 Capnographic Airway Assessment for Procedural Sedation and Analgesia—cont’d

<table>
<thead>
<tr>
<th>DIAGNOSIS</th>
<th>WAVEFORM</th>
<th>FEATURES</th>
<th>INTERVENTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Physiologic variability</td>
<td><img src="waveform1.png" alt="Waveform" /></td>
<td>(\text{SpO}\text{2}), (\text{PETCO}_2), Waveform RR</td>
<td>No intervention required</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Normal</td>
<td>Continue sedation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Normal</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Varying&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Normal</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Curved</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Normal, ↑, or ↓&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Wheezing</td>
<td></td>
</tr>
<tr>
<td>Partial airway obstruction</td>
<td><img src="waveform2.png" alt="Waveform" /></td>
<td>(\text{SpO}\text{2}), (\text{PETCO}_2), Waveform RR</td>
<td>Full airway patency restored with airway alignment</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Normal</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Variable</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Noisy breathing and/or inspiratory stridor</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>PETCO&lt;sub&gt;2&lt;/sub&gt; Normal, ↑, or ↓&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Reassess patient</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Waveform Curved</td>
<td>Establish IV access</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RR Normal, ↑, or ↓&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Supplemental (\text{O}_2) (as needed)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Other</td>
<td>Cease drug administration</td>
</tr>
<tr>
<td>Partial laryngospasm</td>
<td><img src="waveform3.png" alt="Waveform" /></td>
<td>(\text{PETCO}_2), Waveform RR</td>
<td>Reassess patient</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PETCO&lt;sub&gt;2&lt;/sub&gt; Normal, ↑, or ↓&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Establish IV access</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Waveform Normal</td>
<td>Supplemental (\text{O}_2) (as needed)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RR Variable</td>
<td>Cease drug administration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PETCO&lt;sub&gt;2&lt;/sub&gt; Normal or ↓&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Waveform Zero</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>RR Zero</td>
<td></td>
</tr>
<tr>
<td>Apnea</td>
<td><img src="waveform4.png" alt="Waveform" /></td>
<td>(\text{PETCO}_2), Waveform RR</td>
<td>Airway patency restored with airway alignment</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PETCO&lt;sub&gt;2&lt;/sub&gt; Normal or ↓&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Airway not patent with airway alignment</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Waveform Zero</td>
<td>No waveform</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RR Zero</td>
<td>Positive pressure ventilation</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Other</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Normal or ↓&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Zero</td>
<td></td>
</tr>
<tr>
<td>Complete airway obstruction</td>
<td><img src="waveform5.png" alt="Waveform" /></td>
<td>(\text{PETCO}_2), Waveform RR</td>
<td>Reassess patient</td>
</tr>
<tr>
<td></td>
<td></td>
<td>PETCO&lt;sub&gt;2&lt;/sub&gt; Normal or ↓&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Establish IV access</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Waveform Zero</td>
<td>Supplemental (\text{O}_2) (as needed)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>RR Zero</td>
<td>Cease drug administration</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Other</td>
<td></td>
</tr>
<tr>
<td>Complete laryngospasm</td>
<td><img src="waveform6.png" alt="Waveform" /></td>
<td>(\text{PETCO}_2), Waveform RR</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>PETCO&lt;sub&gt;2&lt;/sub&gt; Normal or ↓&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Waveform Zero</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>RR Zero</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Other</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup>Varying waveform amplitude and width.

<sup>b</sup>Depending on the duration and severity of bronchospasm.

<sup>c</sup>Depending on the duration of the episode.

\(\text{PETCO}_2\) End-tidal carbon dioxide pressure; \(\text{RR}\), respiratory rate; \(\text{SpO}_2\), oxygen saturation as measured by pulse oximetry.
acidosis in these patients (Fig. 2.16).64–66 As the patient becomes acidotic (i.e., $\text{HCO}_3^-$ decreases), a compensatory respiratory alkalosis develops with an increase in minute ventilation and a resultant decrease in PETCO$_2$. By increasing minute ventilation, these patients are able to lower arterial CO$_2$ tension to help correct the underlying acidemia. The more acidotic, the lower $\text{HCO}_3^-$, the higher the RR, and the lower the PETCO$_2$.

PETCO$_2$ can be used to distinguish diabetics in ketoacidosis (metabolic acidosis, compensatory tachypnea, low PETCO$_2$) from those who are not (non-acidotic, normal RR, normal PETCO$_2$).64,65 Capnography could be utilized as a potential triage tool to determine the need for intravenous access and rapid assessment by a physician versus delayed physician assessment and potentially starting oral rehydration for patients with concerns for diabetic- or dehydration-related metabolic acidosis.

**Limitations**

Significant technical problems have historically restricted the effective clinical use of capnography. Such problems include interference with the sensor by condensed water and patient secretions in both mainstream and high-flow sidestream devices, cross-sensitivity with anesthetic gases in conventional CO$_2$ sensors, lack of ruggedness for intrahospital and interhospital transport, and power consumption issues related to portable battery operation time. These issues have largely been resolved in the newer-generation capnography monitors.

Problems with accuracy continue to affect high-flow sidestream systems. When the tidal volume of the patient drops below the flow rate of the system (e.g., neonates, infants, hypventilating patients with low–tidal volume breathing), the monitor will entrain room air, thereby falsely diluting PETCO$_2$ and slurring the ascending phase of the waveform.

Early capnography airway interfaces (i.e., nasal cannula) had difficulty providing consistent measurements in mouth-breathing patients and those who alternated between mouth and nose breathing. The newer oral-nasal interface has addressed these problems.

Capnography is most effective when assessing a pure ventilation, perfusion, or metabolism problem. Capnographic findings in patients with mixed ventilation, perfusion, or metabolism problems are difficult to interpret. For example, in patients with complex pathophysiology, a ventilation problem may elevate PETCO$_2$, and a perfusion problem may simultaneously lower PETCO$_2$. Absolute values and even trends over time may be difficult to interpret in these situations.

Although capnography in patients in cardiac arrest is 100% specific for tracheal placement of the ET tube, its sensitivity for esophageal placement is uncertain.

**CONCLUSION**

Capnography is a versatile noninvasive diagnostic modality for monitoring ventilation, perfusion, and metabolic status in both intubated and nonintubated patients. Clinical applications include verification and continuous monitoring of ET tube placement; determination of the efficacy of CPR in cardiac arrest; ventilatory monitoring of head-injured patients; assessment of vital signs in patients who are critically ill, injured, seizing, or undergoing procedural sedation or have altered mental status; evaluation of patients in acute respiratory distress; and detection of metabolic acidosis.
REFERENCES


49. Lah K, Krizmaric M, Grmec S: The dynamic pattern of end-tidal carbon dioxide during cardiopulmonary resuscitation: difference between asphyxial
cardiac arrest and ventricular fibrillation/pulseless ventricular tachycardia
of early traumatic brain injury care: prospective multicenter study in Austria.
Basic Airway Management and Decision Making

Brian E. Driver and Robert F. Reardon

Basic airway procedures are often overlooked in favor of more exciting intubation devices and techniques, but basic procedures are critically important and often lifesaving. Establishment of a patent airway, oxygenation, and bag-mask ventilation (BMV) remain the cornerstones of good emergency airway management (Videos 3.1–3.11). These techniques can be used quickly and in any setting. They allow practitioners to keep apneic patients alive until a definitive airway can be established.

Extraglottic devices, such as laryngeal mask airways (LMAs) and the King Laryngeal Tube (LT), are also important for the initial resuscitation of apneic patients and for rescue ventilation when intubation fails.

This chapter describes basic airway skills, including opening the airway, oxygen (O₂) therapy, BMV, and extraglottic airway (EGA) devices. These are the skills that providers can rely on when other airway techniques are difficult or impossible. Mastery of these skills will help providers manage difficult, anxiety-provoking emergency airways.

THE CHALLENGE OF EMERGENCY AIRWAY MANAGEMENT

Although other specialists are sometimes available, most emergency airways are managed by emergency medicine providers. Airway management in the emergency department (ED) is unique and significantly different from airway management in the controlled setting of an operating room. Additionally, conventional airway management tools may be ineffective in the uncontrolled emergency environment. Major challenges include hypoxia; shock; full stomach, and the presence of emesis, blood, or excessive secretions in the airway. Many patients are uncooperative and combative, making it impossible to properly examine the airway before choosing an intubation technique. Medical history, allergies, and even the current diagnosis are often unknown before emergency airway management begins. Time constraints, lack of patient cooperation, and risk for emesis and aspiration limit the use of some techniques, such as awake intubation. In trauma patients, the risk for cervical spine injury limits optimal head and neck positioning for BMV and laryngoscopy. All these factors increase the risk for complications from emergency airway management, and approximately 0.5% to 1% of all emergency airways require a surgical approach. The increasing use of video laryngoscopy may reduce the incidence of emergency surgical airways.

BASIC AIRWAY MANAGEMENT TECHNIQUES

Opening the Airway

The first concern in the management of a critically ill patient is patency of the airway. Upper airway obstruction commonly occurs when patients are unconscious or sedated. It can also be due to injury to the mandible or muscles that support the hypopharynx. In these situations, the tongue moves posteriorly into the upper airway when the patient is in a supine position. Upper airway obstruction caused by the tongue can be relieved by positioning maneuvers of the head, neck, and jaw; the use of nasopharyngeal or oropharyngeal airways; or the application of noninvasive positive pressure ventilation (NPPV).

Manual Airway Maneuvers

Airway obstruction in unconscious patients is often due to posterior displacement of the tongue (Fig. 3.1A). Research in patients with obstructive sleep apnea using NPPV supports the concept that the airway collapses like a flexible tube. The chin-lift and the
The Triple Airway Maneuver

The “triple airway maneuver” is described by some authors as a valuable method for maintaining a patent upper airway.2,12 The most common description of this maneuver is head tilt, jaw thrust, and mouth opening.2,12 Evidence demonstrates that the upper airway is more patent when the mouth is closed.23–25 Although the triple maneuver is commonly mentioned in the anesthesia literature as a valuable technique, no studies exist to support the assertion that this technique is more effective than the head-tilt/chin lift or jaw-thrust maneuvers.

Patient Positioning

The best way to position a patient’s head and neck for opening the upper airway is to mimic how patients position themselves when they are short of breath, with the neck flexed relative to the torso and with atlanto-occipital extension.2 This is known as the sniffing position and was described by Magill almost 100 years ago.26 In normal-sized supine adults this is accomplished by elevating the head approximately 10 cm while tilting the head back, so that the plane of the patient’s face tilts slightly toward the provider at the head of the bed (see Chapter 4, Fig. 4.8).2,27–29 Morbidly obese patients require much more head elevation to achieve the proper sniffing position. This can be accomplished by building a ramp of towels and pillows under the upper torso, head, and neck or by using a T roop Elevation Pillow (Mercury Medical, Clearwater, FL) or similar device (Fig. 3.2).30–33 Rather than elevating the head by a standard height, the goal of head elevation is to achieve horizontal alignment of the external auditory meatus with the sternum. This is the best position for opening the upper airway in morbidly obese patients.32–35

Importantly, the addition of NPPV may relieve airway obstruction when simple manual positioning maneuvers fail. Meier and colleagues showed that adding NPPV to the chin-lift and jaw-thrust maneuvers decreased stridor and improved the nasal fiberoptic view of the glottic opening in anesthetized children. The use of NPPV for patients with upper airway obstruction should not be considered a definitive solution.

The Jaw-Thrust Maneuver

The jaw-thrust maneuver is the most important technique used to open the upper airway. To perform the jaw-thrust maneuver, place the tips of the middle or index fingers behind the angle of the mandible (see Fig. 3.1C). Lift the mandible toward the ceiling until the lower incisors are anterior to the upper incisors. This maneuver can be performed in combination with the head-tilt/chin-lift maneuver or with the neck in the neutral position during in-line stabilization.

The Head-Tilt/Chin-Lift Maneuver

To perform the head-tilt/chin-lift maneuver, place the tips of the index and middle fingers beneath the patient’s chin (see Fig. 3.1B). Lift the chin cephalad and toward the ceiling. The upper part of the neck will naturally extend when the head tilts backward during this maneuver. Apply digital pressure on only the bony prominence of the chin and not on the soft tissues of the submandibular region. The final step in this maneuver is to use the thumb to open the patient’s mouth while the head is tilted and the neck is extended.
with a towel or pillow to move the head and neck into a more neutral position even in the setting of cervical spine trauma.

In young children, the sniffing position is often achieved without lifting the head because the occiput of a child is relatively large, so the lower cervical spine is normally flexed when the child is lying supine on a flat surface.

Airway management is usually easiest when patients are in the supine position, but the lateral position may be best for patients who are actively vomiting and those with excessive upper airway bleeding or secretions. Some evidence suggests that rotating patients to the lateral position may not prevent aspiration. Patients with suspected cervical spine injury should have their head immobilized with manual in-line stabilization if they need to be rolled to the lateral position. Airway management maneuvers will be more difficult when patients are in the lateral position.

**Foreign Body Airway Obstruction**

Awake patients with partial airway obstruction can usually clear a foreign body on their own. Intervention is required when the patient is not moving air or has altered mental status. Some patients with upper airway obstruction can be ventilated and oxygenated with aggressive high-pressure BMV, so always try this if standard BMV fails. Massive aspiration of vomitus is often a fatal event because of inability of the patient and clinician to adequately clear the airway.

**Abdominal Thrusts (Heimlich Maneuver), Chest Thrusts, and Back Blows (Slaps)**

The International Consensus Conference on Cardiopulmonary Resuscitation and Emergency Cardiopulmonary Care evaluated the evidence for different techniques to clear foreign body airway obstruction. They found good evidence for the use of chest thrusts, abdominal thrusts, and back blows or slaps. Insufficient evidence exists to determine which technique is the best and which should be used first.

The technique of subdiaphragmatic abdominal thrusts to relieve a completely obstructed airway was popularized by Dr. Henry Heimlich and is commonly referred to as the Heimlich maneuver. The technique is most effective when a solid food bolus is obstructing the larynx. In a conscious patient, stand behind the upright patient. Circle the arms around the patient's midsection with the radial side of a clenched fist placed on the abdomen, midway between the umbilicus and xiphoid. Then grasp the fist with the opposite hand and deliver an inward and upward thrust to the abdomen (Fig. 3.3A). A successful maneuver will cause the obstructing agent to be expelled from the patient's airway by the force of air exiting the lungs. Abdominal thrusts are relatively contraindicated in pregnant patients and those with procterabdomens. Potential risks associated with abdominal thrusts include stomach rupture, esophageal perforation, and mesenteric laceration, compelling the rescuer to weigh the risks and benefits of this maneuver. Use a sternal hand position for pregnant patients (Fig. 3.3B).

If a choking patient loses consciousness, use chest compressions in an attempt to expel the obstructing agent (Fig. 3.3C). The theory is the same as the Heimlich maneuver, with high intrathoracic pressure created to push the obstruction out of the airway. Some data suggest that chest compressions may generate higher peak airway pressure than the Heimlich maneuver. After 30 seconds of chest compressions, remove the obstructing object if you see it, attempt 2 breaths, and then continue cardiopulmonary resuscitation (CPR; 30 compressions to 2 breaths). Every time you open the airway to give breaths, look for the object and remove it if possible, and then continue CPR if necessary.

Back blows are recommended for infants and small children with a foreign body obstructing the airway. Some authors have argued that back blows may be dangerous and may drive foreign bodies deeper into the airway, but there is no convincing evidence of this phenomenon. Anecdotal evidence suggests that back blows are effective. No convincing data, however, indicate that back blows are more or less effective than abdominal or chest thrusts. Back blows may produce a more pronounced increase in airway pressure, but over a shorter period than with the other techniques. The AHA guidelines suggest back blows in the head-down position (Fig. 3.3D) and head-down chest thrusts in infants and small children with foreign body airway obstruction (Fig. 3.3E). The AHA does not recommend abdominal thrusts in infants because they may be at higher risk for iatrogenic injury. From a practical standpoint, back blows should be delivered with the patient in a head-down position, which is more easily accomplished in infants than in larger children. It is recommended that suction be performed on newborns rather than giving them back blows or abdominal thrusts.

Any patient with a complete airway obstruction may benefit from chest compressions, abdominal thrusts, or back blows. It is important to realize that more than one technique is often required to clear obstruction of the airway by a foreign body, so multiple techniques should be applied in a rapid sequence until the obstruction is relieved. Perform a finger sweep of the patient's mouth only if a solid object is seen in the airway.

Perform CPR on all unconscious patients with airway obstruction. Aggressive high-pressure BMV should be attempted, as this may distend the trachea enough to allow air to bypass the obstruction. In cases in which obstructive foreign bodies cannot be removed under direct visualization...
and aggressive positive pressure ventilation has failed, practitioners with advanced airway skills and proper equipment can try to push a subglottic foreign body beyond the carina, usually into the right main stem bronchus.

**Suctioning**

Patient positioning and airway-opening maneuvers are often inadequate to achieve complete airway patency. Ongoing hemorrhage, vomitus, and particulate debris frequently require suctioning. Several types of suctioning tips are available. A large-bore dental-type suction tip is the most effective in clearing vomitus from the upper airway because it is less likely to become obstructed by particulate matter. The tonsil tip (Yankauer) suction device can be used to clear hemorrhage and secretions. Its rounded tip is less traumatic to soft tissues, but the tonsil tip device is not large enough to effectively suction vomitus.

A large-bore dental-type tip device, such as the HI-D Big Stick (SSCOR Inc, Sun Valley, CA) suction tip, should be readily available at the bedside during all emergency airway management (Fig. 3.4). The large-bore tip allows rapid clearing of vomitus, blood, and secretions.
A limiting feature of many suction catheters is the diameter of the tubing. Vomitus may obstruct the standard 1/4-inch-diameter catheter. A 5/8- or 3/4-inch-diameter suction tube (Kuriyama Tubing, 5/16-inch inner diameter, 0.44-inch outer diameter, clear; www.grainger.com) has been shown to significantly decrease suction time for viscous and particulate material (see Fig. 3.4).52

Keep suctioning equipment connected and ready to operate. Everyone participating in emergency airway management should know how to use it. Interposition of a suction trap close to the suction device prevents clogging of the tubing with particulate debris.

No specific contraindications to airway suctioning exist. Complications of suctioning may be avoided by anticipating problems and providing appropriate care before and during suctioning maneuvers. Nasal suction is seldom required, except in infants, because most adult airway obstruction occurs in the mouth and oropharynx.

Avoid prolonged suctioning because it may lead to significant hypoxia, especially in children. Do not exceed 15 seconds for suctioning intervals and administer supplemental O2 before and after suctioning. Naigow and Powasner7 found that suctioning consistently induced hypoxia in dogs and that it was best avoided by hyperventilation with high-concentration O2 before and after suctioning. Hypoxia was also prevented in children by preoxygenation prior to endotracheal (ET) suctioning attempts.54

When feasible, perform suctioning under direct vision or with the aid of the laryngoscope. Forcing a suction tip blindly into the posterior pharynx can injure tissue or convert a partial obstruction to a complete obstruction.

**Artificial Airway Placement**

The simplest and most widely available artificial airways are the oropharyngeal and nasopharyngeal airways (Fig. 3.5). Both are intended to prevent the tongue from obstructing the airway by creating a passage for air between the base of the tongue and the posterior pharyngeal wall. The oral airway may also prevent teeth clenching. In cases of severe upper airway edema, such as angioedema, these devices may not function properly or be able to bypass the obstruction. The oropharyngeal airway may be inserted by either of two procedures. One approach is to insert the airway in an inverted position along the patient’s hard palate (Fig. 3.5, step 2). When it is well into the patient’s mouth, rotate the airway 180 degrees and advance it to its final position along the patient’s tongue, with the distal end of the artificial airway lying in the hypopharynx (Fig. 3.5, step 3). No rotation is necessary when the airway is placed in this manner. This technique may be less traumatic, but it takes longer.

The nasopharyngeal airway is very easy to place. It may be easiest to place it on the patient’s right naris so that the bevel is facing the septum on insertion. Be sure to lubricate the device before insertion (Fig. 3.5, step 6). Some clinicians insert a nasopharyngeal airway to dilate the nasal passages for 20 to
Oropharyngeal Airway Insertion

For oropharyngeal airway insertion, first measure. An airway of correct size will extend from the corner of the mouth to the earlobe or the angle of the mandible.

When the airway is well into the mouth, rotate it 180°, with the distal end of the airway lying in the hypopharynx. It may help to pull the jaw forward during passage.

Alternatively, open the mouth widely and use a tongue blade to displace the tongue inferiorly, and advance the airway into the oropharynx. No rotation is required with this method.

NASOPHARYNGEAL AIRWAY INSERTION

For nasopharyngeal airways, a device of correct size will extend from the tip of the nose to the earlobe.

Advance the airway into the nostril and direct it along the floor of the nasal passage in the direction of the occiput. Do not advance in a cephalad direction!

Advance the airway fully until the flared external tip of the device is at the nasal orifice.

Generously lubricate the airway prior to insertion.

Open the patient’s mouth with your thumb and index finger, then insert the airway in an inverted position along the patient’s hard palate.

30 minutes before nasotracheal intubation. Simply advance it into the nostril and direct it along the floor of the nasal passage in the direction of the occiput, not cephalad (Fig. 3.5, step 7). Advance it fully until the flared external tip of the airway is located at the nasal orifice (Fig. 3.5, step 8).

Both oropharyngeal and nasopharyngeal airways are available in multiple sizes. To find the correct size of either device, estimate its size by measuring along the side of the patient's face before insertion. An oropharyngeal airway of the correct size will extend from the corner of the mouth to the tip of the earlobe (see Fig. 3.5, step 1); a nasopharyngeal airway of the correct size will extend from the tip of the nose to the tip of the earlobe (see Fig. 3.5, step 5).

Both oropharyngeal and nasopharyngeal airways provide airway patency similar to that achieved with the head-tilt/chin-lift maneuver. The nasal airway is better tolerated by
semiconscious patients and is less likely to induce emesis in those with an intact gag reflex.

**Complications**

The nasopharyngeal airway may cause epistaxis and may be dangerous in patients with significant facial and basilar skull fractures. Semiconscious patients with nasopharyngeal airways may deteriorate and require intubation, so they should be monitored closely.

The oropharyngeal airway may induce vomiting when placed in patients with an intact gag reflex. It may also cause airway obstruction if the tongue is pushed against the posterior pharyngeal wall during insertion. The oropharyngeal airway should not be used as a definitive airway.

**OXYGEN THERAPY**

Adequate O₂ delivery depends on the inspired partial pressure of O₂, alveolar ventilation, pulmonary gas exchange, oxygen-carrying capacity of blood, and cardiac output. The easiest factor to manipulate is the partial pressure of inspired O₂, which is accomplished by increasing the fraction of inspired oxygen (FiO₂) with supplemental O₂.

**Indications and Contraindications**

Resuscitate all patients in cardiac or respiratory arrest with 100% O₂. The most certain indication for supplemental O₂ is the presence of arterial hypoxemia, defined as an arterial oxygen partial pressure (PaO₂) lower than 60 mm Hg or arterial oxygen saturation (SaO₂) less than 90%. Normal individuals will begin to experience memory loss at a PaO₂ of 45 mm Hg, and loss of consciousness occurs at a PaO₂ of 30 mm Hg. Chronically hypoxic patients can adapt and function with a PaO₂ of 50 mm Hg or lower.

When tissue hypoxia is present or suspected, give O₂ therapy. Shock states resulting from hemorrhage, vasodilatory states, low cardiac output, and obstructive lesions can all lead to tissue hypoxia and benefit from supplemental O₂. Whatever the cause of the shock state, administration of O₂ is indicated until the situation can be thoroughly evaluated and cause-specific therapy instituted. It is reasonable to administer O₂ to hypotensive patients and those with severe trauma until tissue hypoxia can definitively be excluded.

Respiratory distress without documented arterial hypoxemia is a common indication for O₂ administration, although no evidence exists to support this practice. Unless ET intubation is planned, respiratory distress without hypoxemia should generally not be considered an indication for oxygen supplementation.

Oxygen therapy is often recommended for acute myocardial infarction, but there is no difference in outcomes between patients receiving O₂ and those receiving room air after myocardial infarction. A randomized trial of room air versus O₂ supplementation for patients with acute ST-elevation myocardial infarction demonstrated that patients who received O₂ had larger infarction size, as assessed by peak myocardial enzymes and cardiac magnetic resonance imaging at 6 months. The AHA recommends O₂ in myocardial infarction only for patients with hypoxemia, signs of heart failure, shock, or respiratory distress.

Although O₂ is routinely administered to acute stroke patients, there is no convincing evidence that this practice is beneficial without documented hypoxia, and it is not recommended by current guidelines.

Administer 100% O₂ to patients with carbon monoxide poisoning. The half-life of carboxyhemoglobin is 4 to 5 hours in a patient breathing room air but can be decreased to approximately 1 hour by the administration of 100% O₂ by non-rebreather face mask at atmospheric pressure.

There are no contraindications to O₂ therapy when a definite indication exists. The risks associated with hypoxemia are grave and undeniable. Never withhold oxygen therapy from a hypoxic patient for fear of complications or clinical deterioration. Carbon dioxide retention is not a contraindication to O₂ therapy. Rather, it demands that the clinician administer O₂ carefully and recognize the potential for respiratory acidosis and clinical deterioration. Although the mechanism for the development of respiratory acidosis in patients with chronic obstructive pulmonary disease (COPD) who are administered O₂ is debated, its occurrence is not. Use caution when administering supplemental O₂ to hypoxic patients with arterial carbon dioxide pressure higher than 40 mm Hg, but do not withhold it.

**Oxygen Administration During Cardiac Arrest and Neonatal Resuscitation**

The AHA guidelines for cardiopulmonary resuscitation and emergency cardiovascular care address the potential harm of oxygen therapy and hyperoxemia following cardiac arrest and during neonatal resuscitation, and provide recommendations for best use. Recommendations from the guidelines are summarized in Box 3.1. Although it is still prudent to administer oxygen in the prehospital and ED setting, additional research may alter these recommendations. As a general guideline, fear of oxygen toxicity should not prevent the use of O₂ when there is an indication, but use the minimum concentration of O₂ necessary to achieve the therapeutic goals.

**Oxygen Delivery Devices**

A common misconception is that oxygen delivery devices can be cleanly separated into low-flow and high-flow categories. Almost any oxygen delivery device can be used across a wide range of flow rates. In fact, it is the source oxygen flow rate, not the device applied, that is the primary driver of the FiO₂ received by the patient.

Delivery of oxygen at low-flow rates provides gas flow that is less than the patient's inspiratory flow rate. The difference between the patient's inspiratory flow and the flow delivered by the device is met by a variable amount of room air being drawn into the system. Patients with normal respiratory rates and tidal volumes will require less outside air than those in respiratory distress, and therefore patients not in respiratory distress typically receive a higher FiO₂ than patients in respiratory distress, assuming equivalent supplemental oxygen flow rates. As a patient's inspiratory flow changes, so will the FiO₂ that they receive from a low-flow device.

Delivery of high-flow oxygen, with rates that match or exceed the inspiratory flow of the patient (generally >30 L/min), provides a significantly higher FiO₂ than low-flow. High-flow oxygen can achieve FiO₂ values of more than 90%.

The prongs of a nasal cannula deliver a constant flow of O₂ that accumulates in the nasopharynx and provides a reservoir of oxygen-enriched air for inspiration. The FiO₂ delivered by
**BOX 3.1 Recommendations for Oxygen Administration During Adult and Neonate Resuscitation: Excerpts From the Guidelines of the American Heart Association**

**OXYGEN SUPPLEMENTATION DURING CARDIAC ARREST**
The goals of cardiopulmonary resuscitation are to restore energy to the heart so it can resume normal function, and ensure adequate energy supply to the brain during resuscitation. Oxygen is vital to these goals, and during cardiac arrest blood flow is the major limiting factor to adequate oxygen delivery to the heart and brain. Thus, 100% FiO₂ should be administered during cardiac arrest for adults, children, and neonates to maximize oxygen delivery to vital organs.10,141,218

**OVERVIEW OF POST–CARDIAC ARREST SUPPLEMENTAL OXYGEN FOR ADULTS AND CHILDREN**43,239 Although 100% oxygen may have been used during initial resuscitation, providers should titrate inspired oxygen to the lowest level required to achieve an arterial oxygen saturation of ≥94%, to avoid potential oxygen toxicity. It is recognized that titration of inspired oxygen may not be possible immediately after out-of-hospital cardiac arrest until the patient is transported to the emergency department or, in the case of in-hospital arrest, the intensive care unit. The optimal FiO₂ during the immediate period after cardiac arrest is still debated. The beneficial effect of high FiO₂ on systemic oxygen delivery should be balanced with the deleterious effect of generating oxygen-derived free radicals during the reperfusion phase. Animal data suggests that ventilations with 100% oxygen (generating PaO₂ >350 mm Hg at 15 to 60 minutes after return of spontaneous circulation [ROSC]) increase brain lipid peroxidation, increase metabolic dysfunctions, increase neurologic degeneration, and worsen short-term functional outcome when compared with ventilation with room air or an inspired oxygen fraction titrated to a pulse oximeter reading between 94% and 96%. Data from human studies is mixed, with no clear evidence of harm or benefit from hyperoxia. There is no physiologic reason to expect that a PaO₂ >350 mm Hg is necessary or beneficial after cardiac arrest, and very well may be harmful.

Once the circulation is restored, it is reasonable to use the highest available oxygen concentration until the oxyhemoglobin saturation or partial pressure of oxygen can be measured. As an arterial oxyhemoglobin saturation of 100% may correspond to a PaO₂ anywhere between ~80 mm Hg and 500 mm Hg, in general it is appropriate to wean FiO₂ when saturation is 100%. If the oxyhemoglobin saturation is 100%, it is reasonable to reduce the oxygen supplementation, provided that the oxyhemoglobin saturation can be maintained at ≥94%. The goal oxyhemoglobin saturation is 94% to 99%. Avoiding hypoxia is more important than avoiding hyperoxia.

**ASSESSMENT OF OXYGEN NEED AND ADMINISTRATION OF OXYGEN IN THE NEONATE IMMEDIATELY AFTER BIRTH**258,260 Oxyhemoglobin saturation may normally remain in the 70% to 80% range for several minutes following birth, thus resulting in the appearance of cyanosis during that time. Clinical assessment of skin color is a very poor indicator of oxyhemoglobin saturation during the immediate neonatal period and that lack of cyanosis appears to be a very poor indicator of the state of oxygenation of an uncompromised baby following birth.

Optimal management of oxygen during neonatal resuscitation becomes particularly important because of the evidence that either insufficient or excessive oxygenation can be harmful to the newborn infant. Hypoxia and ischemia are known to result in injury to multiple organs. Conversely there is growing experimental evidence, as well as evidence from studies of babies receiving resuscitation, that adverse outcomes may result from even brief exposure to excessive oxygen during and following resuscitation. Two metaanalyses of several randomized controlled trials comparing neonatal resuscitation initiated with room air versus 100% oxygen showed increased survival when resuscitation was initiated with air.

It is recommended that the goal in babies being resuscitated at birth, whether born at term or preterm, should be an oxygen saturation value in the interquartile range of preductal saturations measured in healthy term babies following vaginal birth at sea level (see later). These targets may be achieved by initiating resuscitation with air or blended oxygen and titrating the oxygen concentration to achieve an SpO₂ in the target range using pulse oximetry. If the baby is bradycardic (heart rate <60 beats per minute) after 90 seconds of resuscitation with a lower concentration of oxygen, oxygen concentration should be increased to 100% until recovery of a normal heart rate.

**ASSESSMENT OF OXYGEN NEED AND ADMINISTRATION OF OXYGEN IN THE NEONATE**

<table>
<thead>
<tr>
<th>Targeted Preductal SpO₂ After Birth</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 min</td>
</tr>
<tr>
<td>2 min</td>
</tr>
<tr>
<td>3 min</td>
</tr>
<tr>
<td>4 min</td>
</tr>
<tr>
<td>5 min</td>
</tr>
<tr>
<td>10 min</td>
</tr>
<tr>
<td>60–65%</td>
</tr>
<tr>
<td>65–70%</td>
</tr>
<tr>
<td>70–75%</td>
</tr>
<tr>
<td>75–80%</td>
</tr>
<tr>
<td>80–85%</td>
</tr>
<tr>
<td>85–95%</td>
</tr>
</tbody>
</table>

nasal cannulas is determined by many factors, including the respiratory rate, tidal volume, pharyngeal geometry, and O₂ flow. Most importantly, at a constant O₂ flow rate, FiO₂ varies inversely with the respiratory rate.71 Despite this limitation, nasal cannulas are very comfortable for patients and are the most common O₂ delivery device. They can be used with higher flow rates for brief periods of time, but are uncomfortable to use in this manner and cause nasal dryness and irritation. Nasal cannulas are generally set to 2 to 4 L/min, which provides approximately 30% to 35% FiO₂.71 Although it may seem intuitive, patients using a nasal cannula should be reminded that they should not smoke while oxygen is being delivered (Fig. 3.6).

Simple masks receive a constant flow of O₂ from the O₂ source and have multiple vent holes. During inspiration the oxygen-enriched air that has accumulated in the mask, along with room air entrained through the vent holes, is inhaled. During expiration, 200 mL (the approximate volume of the mask) of exhaled gas is deposited in the mask, with the rest exiting through the vent holes. The continuous flow of O₂ then partially washes out the mask before the next inspiration. The mask itself provides the reservoir of oxygen-rich gas for
The Venturi mask is a titratable oxygen delivery device that is widely available, though it is used more frequently in the inpatient and intensive care unit setting than in the ED (Fig. 3.8). It is important to understand that although Venturi masks are often referred to as high flow systems they cannot provide FiO₂ above 35%. The primary advantage of the Venturi mask system is that FiO₂ can be precisely controlled between 21% and 100%.
and 35% for patients who may not tolerate a higher or imprecise FiO₂. Room air is pulled into the system through entrainment ports and mixes with the O₂ provided from the O₂ source. The proportion of entrained air—and therefore FiO₂—is constant and determined by the velocity of the O₂ jet and the size of the entrainment ports. Because the total gas flow (O₂ plus air through the entrainment ports) meets or exceeds the patient’s inspiratory flow rate, no additional entrainment of air occurs around the mask, thereby minimizing changes in FiO₂ as the patient’s respiratory pattern changes. Venturi masks are packaged with multiple inserts, each with a different size orifice for O₂ inflow. FiO₂ is determined by selecting the appropriate colored insert and O₂ flow rate according to the manufacturer’s instructions. All Venturi mask settings and inserts provide a total gas flow of 30 L/min, which matches the inspiratory flow rate of a resting adult. However, a patient in respiratory distress may have an inspiratory flow rate of 50 to 100 L/min. If the patient’s inspiratory flow rate exceeds the total gas flow delivered by the mask, additional air will be entrained around the mask, and inspired FiO₂ will decrease. This is especially true with masks rated above 35% FiO₂, which generally can only provide high FiO₂ if minimal room air is entrained around the edges of the mask. Caution should be used with masks rated above 35% in patients with respiratory distress because FiO₂ may be significantly reduced with high inspiratory flow rates.

High-flow nasal cannula oxygen (HFNC) is the delivery of heated, humidified oxygen at high flow rates (up to approximately 60 L/min) through wide-bore nasal cannulas. HFNC is hypothesized to deliver a higher FiO₂ than face mask oxygen because the nasopharynx acts as a natural reservoir that refills with oxygen after each breath, though at high flow rates face mask oxygen probably also fills the nasopharynx. HFNC requires three components: wide-bore nasal prongs, a humidifier, and a gas delivery device to control flow. Whereas HFNC requires additional equipment and a brief set-up for each patient, it delivers very high FiO₂ and is better tolerated by patients than face mask oxygen and NPPV. At flow rates of 15 L/min FiO₂ is 70% to 80%, and at flow rates of 45 L/min FiO₂ is 90% or higher. These FiO₂ values remain relatively constant, even when the patient’s mouth is open. Commercially available HFNC systems use flow rates of 5 to 40 L/min and are capable of delivering an FiO₂ of close to 100%. High-flow oxygen by nasal cannula is not well tolerated unless it is humidified, so commercially available systems (Vapotherm [Exeter, NH], Fisher and Paykel Nasal High Flow [Auckland, New Zealand], AquinOx [Smiths Medical, St. Paul, MN]) deliver oxygen with nearly 100% humidity. HFNC additionally provides low levels of positive airway pressure in the range of 1 cm H₂O to 3 cm H₂O, which increases as the flow rate increases. HFNC devices are popular in neonatal and pediatric intensive care units and are commonly used for respiratory support after extubation and for management of respiratory disease in neonates.

HFNC has been demonstrated to increase tidal volumes and increase end-expiratory lung volumes after cardiac surgery, and was associated with lower rates of escalation to more invasive modes of ventilation in two ED studies. A landmark trial that randomized patients with acute hypoxic respiratory failure without hypercapnia to receive HFNC, face mask oxygen, or NPPV found similar intubation rates but lower 90-day mortality in the HFNC group. Though the results of this trial are not necessarily generalizable to ED patients, it does demonstrate that HFNC provides similar oxygenation support for patients with severe hypoxemia as compared to NPPV. This makes HFNC a useful tool for patients who do not need immediate tracheal intubation and may be susceptible to lung injury from mechanical ventilator–induced barotrauma.

**Procedure**

In selecting the proper delivery device, consider the clinical condition of the patient and the amount of O₂ needed. Venturi systems should generally be used for patients who need precise control of FiO₂, such as COPD patients with chronic respiratory acidosis. Nasal cannulas and face masks set at lower flow rates are appropriate for patients who need supplemental O₂ but do not require precise control of FiO₂. Patients with significant hypoxemia, end-organ dysfunction, or respiratory distress require a higher FiO₂ delivery system.

Frequent clinical assessment and blood oxygen saturation (SpO₂) monitoring are needed in all patients receiving O₂ therapy. Equilibration of SaO₂ after changes in supplemental O₂ occurs within 5 minutes. FiO₂ should be titrated to achieve therapeutic goals while minimizing the risk for complications. An SaO₂ of 90% to 95% (Pao₂ = 60–80 mm Hg) is an appropriate target for most patients receiving supplemental O₂. Increases above these levels do not add appreciably to the O₂ content of blood and are unlikely to confer an additional benefit. One may exceed these parameters in patients with shock and end-organ dysfunction, but the added risk and small potential benefit should be considered on an individual basis.

An initial FiO₂ of 24% to 28% delivered by Venturi mask is indicated for patients with hypoxemia and chronic respiratory acidosis. Periodic blood gas analysis or capnography is imperative for those at risk for respiratory acidosis. In patients with COPD-associated hypercapnia, an SaO₂ of 90% (Pao₂ = 60 mm Hg) should be the goal of O₂ therapy. Mechanical ventilation should be considered when oxygenation goals cannot be achieved without progressive respiratory acidosis.

**Preoxygenation Prior to Endotracheal Intubation**

Preoxygenation prior to ET intubation is one of the most important aspects of emergency airway management, and is a different concept than supplemental oxygenation. Whereas the goal of supplemental oxygenation is to maintain normoxemia, the goal of preoxygenation is to replace all nitrogen in the lungs with oxygen, thereby creating a reservoir of oxygen available to the body during the intubation process. Preoxygenation and oxygen therapy during apnea are discussed fully in Chapter 4.

**Complications of Oxygen Therapy**

Worsening of CO₂ retention leading to progressive respiratory acidosis and obtundation in patients with COPD is the complication most likely to be seen in the ED. This phenomenon is well documented and was first described by Barach in 1937. It has been attributed to several mechanisms, including loss of hypoxic respiratory drive, ventilation-perfusion (V/Q) mismatch, and decreased hemoglobin affinity for CO₂ (Haldane
effect). This avoidable complication is best prevented by administering O₂ to chronic CO₂ retainers only when there is an indication, administering it at the smallest effective dose, and carefully monitoring clinical, capnographic, and arterial blood gas parameters.

Exposing the brain and lung to excessive concentrations of O₂ can lead to toxicity and, in severe cases, can cause acute respiratory distress syndrome (ARDS). Injury to the pulmonary parenchyma occurs as a result of the formation of reactive oxygen species. Oxygen toxicity is of special concern in premature neonates, in whom prolonged hyperoxemia can lead to retinopathy. No data describe what concentration or duration of exposure to O₂ leads to toxicity, but presumably both these factors and individual patient characteristics determine the likelihood of toxicity. The benefits of O₂ therapy in the ED usually outweigh the risk for O₂ toxicity. Fear of toxicity should not prevent the use of O₂ when there is an indication but should encourage the clinician to use the minimum concentration of O₂ necessary to achieve therapeutic goals. High concentrations of O₂ are well tolerated over short periods and may be lifesaving.

In patients receiving high concentrations of supplemental O₂, nitrogen in the alveoli is largely replaced by O₂. If this O₂ is then absorbed into the blood faster than it can be replaced, the volume of the alveoli will decrease and absorptive atelectasis can occur. Airway obstruction potentiates this problem by preventing the rapid replacement of absorbed gas.

**Indications and Contraindications**

BMV is the most common initial technique for ventilation of apneic patients and for rescue ventilation after failed intubation. Many authors note that BMV is relatively contraindicated in patients with a full stomach, those in cardiac arrest, and those undergoing rapid-sequence intubation. These patients have a high risk for stomach inflation and subsequent aspiration. Unfortunately, these are the patients for whom ED providers most commonly use BMV. In ED situations, the need for ventilation and oxygenation always takes priority over potential aspiration.

The only contraindication to attempting BMV is when application of a face mask is impossible (Fig. 3.9). It is often impossible to achieve an effective face mask seal on patients with significant deforming facial trauma and those with thick beards. An intermediate ventilation device, such as an LMA, is a better choice for initial ventilation in such patients.

**BMV Technique**

Achieving adequate ventilation with a bag-mask device requires an open upper airway and a good mask seal. Overly aggressive BMV causes stomach inflation and increases the risk for aspiration. The goal is to achieve adequate gas exchange while keeping peak airway pressure low. Squeezing the bag forcefully creates high peak airway pressure and is more likely to inflate the stomach. Several studies have shown that increased tidal volume is associated with higher peak airway pressure and increased gastric inflation. Decreased inspiratory time (faster bag squeeze) increases peak airway pressure and gastric inflation. Therefore the best method of BMV is to provide a tidal volume of approximately 500 mL delivered over 1 to 1.5 seconds.

**BAG-MASK VENTILATION**

BMV is the single most important technique for emergency airway management. Bag-mask devices are widely available and are standard equipment in all patient care settings. Although the bag-mask method of ventilation appears to be simple, it can be difficult to perform correctly. Good BMV skills are a prerequisite to more advanced methods of emergency airway management. Manually opening the airway, properly positioning the head and neck, placing an oropharyngeal airway device, and achieving a tight face mask seal are the keys to good BMV.

**Review Box 3.2** Bag-mask ventilation: indications, contraindications, complications, and equipment.
Using a ventilator (instead of a resuscitation bag) to provide the proper tidal volume and inspiratory time is an alternative to using a bag-valve device. Effective ventilation and oxygenation should be judged by chest rise, breath sounds, SpO2, and capnography.

A variety of mask configurations are available to facilitate a tight seal. The most common mask used in ED situations is a transparent disposable plastic mask with a high-volume, low-pressure cuff. This type of mask eliminates the need for an anatomically formed mask and can be used for a wide variety of patients with different facial features. Various mask sizes are available.

For a single rescuer, only one hand can be used to achieve the seal because the other must squeeze the bag. The rescuer must apply pressure anteriorly while simultaneously lifting the jaw forward. The thumb and index finger provide anterior pressure while the fifth and fourth fingers lift the jaw. The C-E clamp technique is often the most effective if an assistant is not available: the thumb and index finger form a “C” to provide anterior pressure over the mask, whereas the third, fourth, and fifth fingers form an “E” to lift the jaw (Fig. 3.10, steps 1 and 2). Generally, well-fitting intact dentures should be left in place to help ensure a better seal with the mask.

In the ED setting it is best to hold the face mask with two hands and have an assistant squeeze the bag. If face mask ventilation is difficult, the most experienced provider should hold the face mask while the less experienced provider squeezes the bag.

There are two different methods for two-handed face mask control. The traditional technique is the double C-E method, where the thumb and index finger of both hands encircle the top of the mask (Fig. 3.10, step 3) and the third, fourth, and fifth fingers of both hands form an “E” to lift both sides of the mandible to meet the mask (Fig. 3.10, step 4). The problem with the double C-E technique is that it is difficult to perform a good jaw thrust with the hands in this position.

The best two-handed method is to hold the mask in place with the thenar eminence of both hands (Fig. 3.10, step 5) and use the long fingers under the angle of the mandible to perform a jaw thrust, while also pressing the mask firmly against the face (Fig. 3.10, step 6). This technique allows the operator to perform a good jaw lift (create mandibular protrusion or an “underbite”) and create a good mask seal with the strongest muscles of the hands (Fig. 3.11). This method is best for patients with difficult mask ventilation, and it also allows inexperienced providers and those with small hands to do a better job with face mask ventilation. In addition, it is important to remember to use oropharyngeal or nasopharyngeal airways (or both) whenever face mask ventilation is difficult.

All bag-mask devices should be attached to a supplemental O2 source (with a flow rate of 15 L/min or higher) to avoid hypoxia. A significant problem with the bag-mask method is the low percentage of O2 achieved with some reservoirs. The amount of O2 delivered is dependent on the ventilatory rate, the volumes delivered during each breath, the O2 flow rate into the ventilating bag, the filling time for reservoir bags, and the type of reservoir used. For adults, a bag-mask device should have an inspiratory valve, a 1500-mL bag reservoir, and one-way exhalation port to provide adequate oxygenation during use. If the bag-mask system does not have a one-way exhalation port, room air will be entrained and the FiO2 will be diluted significantly.

Pediatric bag-mask devices should have a minimum volume of 450 mL. Pediatric and larger bags may be used to ventilate infants with the proper mask size, but care must be maintained to administer only the volume necessary to effectively ventilate the infant. Avoid pop-off valves because the airway pressure required for ventilation under emergency conditions can exceed the pressure of the valve.

Previously it had been contended that mask ventilation could be made more difficult after administration of paralytic agents. However, many studies demonstrate that BMV after paralysis is generally easier and provides larger tidal volumes. It is generally not necessary to test whether the patient can be successfully mask-ventilated before administration of paralytic agents, however it is prudent to assess for markers of difficult BMV (see Complications later).

BMV may be the best method of prehospital airway support in trauma patients and children. Murray and colleagues performed a large retrospective study suggesting that patients with severe head injury had a higher risk for mortality if they were intubated in the prehospital setting. In the same year, Gausche and colleagues reported that neurologic outcomes and ultimate survival rates after prehospital pediatric resuscitation with BMV by emergency medical service (EMS) providers were as good as those with tracheal intubation.

Complications

The main complications of BMV are gastric inflation and inability to ventilate. Langeron and colleagues performed a large prospective study of adults undergoing general anesthesia and reported a 5% incidence of difficult mask ventilation. The incidence is obviously much higher in the emergency setting. Risk factors for difficult BMV include presence of a beard, severe facial trauma, obesity, lack of teeth, age older than 55 years, history of snoring, short thyromental distance, limited mandibular protrusion, and decreased pulmonary compliance (severe asthma, COPD, ARDS, term pregnancy).
If a good technique is used, some gastric distension will generally occur. Minor gastric distention should not be considered substandard in the setting of prolonged BMV.

**Cricoid Pressure (Sellick Maneuver)**

In 1961, Sellick described the use of cricoid pressure to prevent regurgitation during anesthesia, and this technique has since become known as Sellick maneuver, though more properly termed cricoid pressure. The purpose of this technique is to apply external force to the anterior cricoid ring to push the trachea posteriorly and compress the esophagus against the cervical vertebrae. In theory, cricoid pressure compresses the distensible upper esophagus but not the airway because the cricoid ring is fairly rigid.

There is no good evidence that cricoid pressure prevents esophageal regurgitation, though it can prevent gastric...
inflation during BMV, thereby reducing the risk for subsequent regurgitation and vomiting. However, cricoid pressure has been demonstrated to reduce tidal volumes, increase peak inspiratory pressure, and prevent adequate air exchange when applied during BMV. Cricoid pressure also decreases successful insertion of and intubation through LMAs. 

BMV can produce gastric inflation, especially if high volume and high pressure are used. To avoid gastric inflation, ventilate with a small volume (500 mL or 6 to 8 mL/kg) and avoid high peak pressure by using a long inspiratory time (1 second). In addition, most airway experts recommend applying cricoid pressure during BMV to further decrease the risk for gastric inflation. It should be noted that the routine use of cricoid pressure during BMV of patients in cardiac arrest is not recommended by the AHA guidelines for cardiopulmonary resuscitation. However, during laryngoscopy and tracheal intubation (discussed in Chapter 4), it is reasonable to release or relax cricoid pressure during insertion of an LMA or if ventilation with the LMA is difficult. It may also be reasonable to release cricoid pressure during laryngoscopy and tracheal intubation (discussed in Chapter 4).

Some authors believe that improper technique is to blame for the many reported failures of cricoid pressure. The proper technique for applying cricoid pressure is to place the thumb and middle finger on either side of the cricoid cartilage, with the index finger in the center anteriorly. Apply 30 N (6.7 lb) of force to the cricoid cartilage in the posterior direction. As a reference, approximately 40 N of digital force on the bridge of the nose will usually cause pain.

**EXTRAGLOTTIC AIRWAY DEVICES**

EGAs are devices that are blindly placed above or posterior to the larynx to allow rapid ventilation and oxygenation. They are good rescue devices for patients who are difficult or impossible to ventilate and oxygenate with a face mask, especially morbidly obese patients and those with large beards or significant facial trauma. In the emergency setting, EGAs can provide temporary rescue ventilation until tracheal intubation or a surgical airway can be performed. It is important to have at least one of these devices immediately available when managing emergency airways. EGAs can be divided into two groups, LMAs and retroglottic devices.

**Laryngeal Mask Airways**

The LMA consists of a hollow shaft or airway tube connected to an oval inflatable masklike cuff designed to sit in the hypopharynx facing the glottis, with the tip at the esophageal inlet. The cuff mask is designed to form a seal around the glottis when the device is placed properly. Some LMAs (LMA Fastrach, Teleflex, Buckinghamshire, UK and LMA Proseal, Teleflex) have handles that allow the operator to increase seal pressure by lifting the entire device toward the ceiling. The first LMA, the LMA Classic (Teleflex) became available in 1988. Since then it has been used more than 200 million times and has been described in more than 2500 academic papers. LMAs are widely considered to be essential adjuncts for rescue ventilation and difficult intubation. can be inserted in less than 30 seconds, and provide effective ventilation in more than 98% of patients. LMAs are primary rescue adjuncts in the difficult airway guidelines put forth by the American Society of Anesthesiologists and the Difficult Airway Society in Europe. Advanced Cardiac Life Support guidelines suggest that the LMA is a reasonable alternative to face mask ventilation. Pediatric Advanced Life Support guidelines acknowledge the LMA as a backup device for difficult pediatric airways. In anesthesia practice, LMAs are now used for a large percentage of cases in which an ET tube may have been used in the past.

This chapter describes how to insert an intubating LMA (ILMA; LMA Fastrach, Teleflex) and use it for rescue ventilation. Intubation through the Fastrach is described in Chapter 4. The insertion and use of the nonintubating LMA Unique (Teleflex) (the disposable version of the LMA Classic) and LMA Supreme (Teleflex) will also be described because they are cheap, disposable, well tested, and widely available.

**Indications**

Both ILMAs and nonintubating LMAs can be used in the “cannot-intubate/cannot-ventilate” scenario with high success rates, and are very useful when face mask ventilation is difficult because of a beard, massive facial trauma, or obesity (Fig. 3.12).

The ILMA should be the LMA of choice for emergency use. In the cannot-intubate/cannot-ventilate scenario, adequate ventilation with the ILMA is possible in nearly all cases, and is more successful than the LMA. In this scenario ventilation with the ILMA is probably superior to bag-mask ventilation. The ILMA can also be used as a primary ventilation and intubation device for patients with difficult airways, and is a reasonable alternative to BMV for reoxygenation after a failed tracheal intubation attempt. Tracheal intubation through the ILMA can be accomplished by using a blind technique, with a laryngoscope, or under endoscopic guidance (see Chapter 4 for tracheal intubation through the ILMA). Studies of difficult airway management with the ILMA show that almost all patients can be adequately ventilated with the ILMA and 94% to 99%...
## Laryngeal Mask Airway Insertion

### Indications
- Failed rapid-sequence intubation
- Difficult bag-mask ventilation
- Difficult intubation
- Facial trauma
- Obesity
- Primary airway in cardiac arrest or use by emergency medical services

### Contraindications
- Limited mouth opening (<2 cm)
- High airway pressure
- Inadequate paralysis or sedation

### Complications
- Inability to ventilate (rare)
- Inability to intubate
- Aspiration (rare)

### Equipment
- Intubating LMA (ILMA/“Fastrach”)
- LMA unique
- 60-mL syringe

### Review Box 3.3
Laryngeal mask airways: indications, contraindications, complications, and equipment.

---

**Figure 3.12**

A, This morbidly obese patient was found asystolic by the emergency medical service. She could not be intubated in the field with multiple attempts and did not survive bag-mask resuscitation. This patient is a candidate for a laryngeal mask airway (LMA). B–D, The LMA is inserted by depressing the jaw, introduced, advanced, inflated, and attached to a resuscitation bag. Postmortem ventilation with the LMA was very easy. In retrospect, the LMA device should have been the first airway adjunct chosen by the emergency medical service (and in the emergency department), totally bypassing attempts with other methods likely to fail.

---

can be intubated through the device. In addition, for difficult airway management the ILMA is technically easier to use than the LMA. In the emergency setting, where obtaining a definitive airway (i.e., tracheal intubation) is the eventual goal, it is more practical to use an ILMA. In addition, the LMA Fastrach (Teleflex) is the most widely used and well-studied ILMA and is easier to insert than the LMA Classic (Teleflex). Finally, when the head is in the neutral position, the LMA Fastrach (Teleflex) is more likely to allow successful ventilation than the LMA Classic (Teleflex) during in-line stabilization of the cervical spine.

ILMAs are especially useful in patients with difficult bag-mask ventilation caused by a beard, severe facial trauma, or obesity because none of these factors inhibit ILMA placement. When brisk bleeding above the glottis makes ventilation and intubation difficult, the ILMA can reduce aspiration of blood and facilitate blind or fiberoptic intubation. In patients requiring urgent cricothyrotomy or percutaneous needle insertion into the trachea, the ILMA can be used to counteract anterior neck pressure. In this capacity, the ILMA provides temporary ventilation and stabilizes the cervical spine during the surgical airway procedure.

Newer LMAs (and some ILMAs) feature a channel posterior to the airway that allows an orogastric tube to be placed into the stomach while the device is used for oxygenation and ventilation. This is particularly useful in patients with gastric distension after BMV, or as a primary device (instead of BMV) in neonates, infants, and children, because they are especially prone to gastric distension with face mask ventilation.

The LMA Classic (Teleflex) (or single-use LMA Unique, Teleflex) is the most extensively tested LMA for children. It may provide a more secure and reliable means of ventilation than bag-mask ventilation. The LMA allows adequate ventilation in 98% of adults with known difficult airways and in 90% to 95% of those with unexpectedly difficult airways. It is also useful as a rescue device in difficult pediatric airways.
Two descriptive studies and 86 case reports describe use of the LMA for difficult pediatric airways. In these reports, ventilation was adequate with the LMA in nearly all pediatric patients. Case series and case reports also suggest that the LMA can provide an effective rescue airway during neonatal resuscitation if BMV and ET intubation fail.

Contraindications
The ILMA and LMA are contraindicated in patients with less than 2 cm of mouth opening because they require 2 cm of space between the upper and the lower incisors to be inserted. Any LMA is relatively contraindicated in awake patients, especially those with a full stomach, because insertion of an LMA in an awake patient will cause coughing, gagging, or emesis. If an LMA is inserted when the patient is awake and the stomach is full, there is a high likelihood of emesis and aspiration. In the ED, an LMA should be used only if the patient is unconscious or after a paralytic agent has been given. Once an LMA is inserted and ventilation is established, the patient should not be allowed to wake up or gag. Consider giving a long-acting paralytic agent or multiple doses of succinylcholine after an LMA is placed and ventilation is adequate.

Although several studies have shown that the ILMA is safe and effective for ventilation and intubation during in-line cervical spine stabilization, some evidence shows that the ILMA causes posterior pressure on the midportion of the cervical spine. The clinical importance of cervical spine pressure caused by the ILMA is unknown, and the device is generally considered safe in patients with an unstable cervical spine injury. Providers should be aware of this concern and make every effort to stabilize the ILMA in these situations.

Types of LMAs
Several manufacturers now make LMAs as the patent on the LMA Classic (Teleflex) expired in 2003. It is important to recognize that all LMAs are not the same.

There are four popular ILMAs. The LMA Fastrach (Teleflex; www.lmaco.com) has the most clinical use and research of all the ILMAs (Fig. 3.13A). Advantages include a handle that makes placement easier and allows the operator to lift up to improve the seal against the laryngeal inlet if needed. There is no gastric port to allow for gastric decompression. Blind intubation rates are significantly higher with the LMA Fastrach (Teleflex) compared to the Cookgas air-Q (CookGas LLC, St Louis, MO; www.cookgas.com); no adequate research has yet been performed comparing blind intubation rates to other devices. The Air-Q intubating laryngeal airway (CookGas LLC) has four different models and is popular with some clinicians (Fig. 3.13B). The i-Gel LMA (Intersurgical, Berkshire, UK; www.i-gel.com) features a thermoplastic elastomer cuff that does not need inflation, so it may be easier to insert than other LMAs (Fig. 3.13C). A gastric channel is present to allow for gastric decompression using an orogastric tube. The Ambu AuraGain (Ambu, Ballerup, Denmark; www.ambu.com) is a newer ILMA with gastric access that seems promising, though little research has been performed on this device to date (Fig. 3.13D).

The nonintubating LMA Supreme (Teleflex) is the latest offering from the Laryngeal Mask Company (Fig. 3.13E); it has a new mask shape that may allow a better mask seal compared to prior models (LMA Classic, Teleflex and LMA Unique, Teleflex) and a gastric evacuation channel. It features a firm curved shape and a bite block handle, so ease of insertion may be similar to that of the Fastrach (Teleflex), and it is available in all sizes from neonate to large adult. However, it does not facilitate tracheal intubation. The LMA Protector (Teleflex) is an ILMA that has recently been released, which is similar to the LMA Supreme (Teleflex) (with gastric access) and will allow intubation using a flexible endoscope.

Procedure
The insertion and use of the LMA Fastrach (Teleflex) and LMA Classic (Teleflex) will be described. Because insertion of all ILMAs and nonintubating LMAs follow similar steps, the procedure for the LMA Fastrach (Teleflex) will be illustrative of steps for ILMAs and LMAs that have a rigid body or handle, and the LMA Classic (Teleflex) will be illustrative for ILMAs and LMAs that are not rigid and do not utilize a handle.

Intubating LMAs
The first step is to select the appropriate size of ILMA, based on manufacturer’s recommendations. The Fastrach ILMA (Teleflex) is available in three sizes: size 3 for children weighing 30 kg to 50 kg, size 4 for small adults weighing 50 kg to 70 kg, and size 5 for adults weighing 70 kg to 100 kg (Table 3.1). When there is doubt about which size is appropriate, it is probably better to use the larger size.

After choosing the correct ILMA, completely deflate the cuff while pushing it posteriorly so that it assumes a smooth wedge shape without any wrinkles (Fig. 3.14, step 1). Place a small amount of water-based lubricant onto the posterior surface of the ILMA just before insertion (Fig. 3.14, step 2). Open the patient’s mouth and position the posterior mask tip so that it is flat against the hard palate, immediately posterior to the upper incisors (Fig. 3.14, step 3). Advance the airway straight into the mouth along the hard palate without rotation until the curved part of the airway tube is in contact with the patient’s chin. Then rotate the ILMA completely into the hypopharynx by advancing it along its curved axis. Keep the posterior of the mask firmly applied to the soft palate and posterior pharynx until firm resistance is felt (Fig. 3.14, step 4). Cricoid pressure impedes proper placement of the ILMA, so briefly release

<table>
<thead>
<tr>
<th>TABLE 3.1</th>
<th>Laryngeal Mask Airway, Disposable Laryngeal Mask Airway, and Intubating Laryngeal Mask Airway Size Recommendations Based on Weighta</th>
</tr>
</thead>
<tbody>
<tr>
<td>WEIGHT (kg)</td>
<td>LMA</td>
</tr>
<tr>
<td>&lt;5</td>
<td>1</td>
</tr>
<tr>
<td>5–10</td>
<td>1.5</td>
</tr>
<tr>
<td>10–20</td>
<td>2</td>
</tr>
<tr>
<td>20–30</td>
<td>2.5</td>
</tr>
<tr>
<td>30–50</td>
<td>3</td>
</tr>
<tr>
<td>50–70</td>
<td>4</td>
</tr>
<tr>
<td>70–100</td>
<td>5</td>
</tr>
<tr>
<td>&gt;100</td>
<td>6</td>
</tr>
</tbody>
</table>

aNote that only a standard LMA is available for patients less than 30 kg. ILMA, Intubating laryngeal mask airway; LMA, laryngeal mask airway.
glottic opening allows optimal ventilation and facilitates tracheal intubation. Before adjusting the ILMA, consider the patient’s position and degree of relaxation because both may affect ILMA function. The ILMA works best in the neutral or sniffing position; cervical extension may interfere with proper placement. The patient should not react to ILMA placement with coughing or gagging because this may interfere with proper placement.

Have a single operator perform the adjustment maneuvers by gripping the ILMA handle with one hand, in a “frying-pan” grip, and providing bag ventilation with the other hand. After each adjustment maneuver, assess the quality of bag ventilation and mask seal. Easy bag ventilation, good chest rise, and absence of an audible mask leak are indications of good ILMA alignment with the glottis (Fig. 3.14, step 6).

If optimal ILMA placement is not accomplished initially, adjusting maneuvers can be attempted. The purpose of adjusting maneuvers is to align the aperture of the ILMA with the glottic opening. Proper positioning of the ILMA aperture with the cricoid pressure while the device is rotated into its final position, wedged into the proximal esophagus. After insertion, the airway tube should emerge from the mouth directed somewhat caudally. Without holding the tube or handle, inflate the mask cuff (Fig. 3.14, step 5). The entire device will normally slide backward a bit when the cuff is inflated. Frequently, only half the maximum cuff volume is sufficient to obtain a good mask seal. Do not overinflate the cuff because this may make the seal worse. See the instruction manual for maximum cuff volumes. Attach a bag and ventilate the patient while using chest rise, breath sounds, and capnography to confirm adequate gas exchange. If bagging is easy and ventilation is good, the aperture of the ILMA is probably aligned correctly over the vocal cords.

To adjust the position of the ILMA, first gently pull the handle toward you without rotation along the ILMA’s curvature. Next, gently push the handle toward the patient’s feet without rotating it. Finally, try the Chandy maneuver, which consists
**Intubating Laryngeal Mask Airway Insertion**

1. Completely deflate the cuff while pushing it posteriorly, so that it assumes a smooth wedge shape without any wrinkles.

2. Arrange a small amount of water-based lubricant onto the posterior surface of the ILMA just before insertion.

3. Place the head and neck in a slightly elevated position with minimal extension. Open the mouth widely and place the posterior surface of the device against the hard palate, immediately posterior to the upper incisors.

4. Advance the ILMA straight into the mouth until the curved part of the airway tube contacts the chin. Then, rotate the ILMA into the hypopharynx until firm resistance is felt. Release cricoid pressure during this step.

5. Let go of the handle and inflate the cuff. Initially inflate the cuff with only half of the maximum volume, and increase inflation as needed. Do not overinflate the cuff. (See product manual for maximum volumes.)

6. Attach a bag and ventilate the patient. Use chest rise, breath sounds, and capnography to confirm adequate gas exchange. If bagging is easy and ventilation is good, the LMA is probably correctly aligned over the glottis.

7. If adjustment is needed, try the Chandy maneuver. First, gently rotate the ILMA farther into the hypopharynx.

8. Next, lift the handle upwards, toward the ceiling above the patient’s feet. This maneuver aligns the mask with the glottis and may provide for better ventilation.

9. If these maneuvers fail, the epiglottis may be folded down over the glottis (asterisk). Perform the “up-down” maneuver, by first rotating the ILMA out of the hypopharynx along its curvature about 5–6 cm.

10. Next, slide the ILMA back into position while pressing it against the posterior pharynx. (Note, the cuff should remain inflated during this maneuver.)

**Figure 3.14** Intubating laryngeal mask airway (ILMA or “Fastrach”; Teleflex) insertion.
of gently rotating the ILMA farther into the hypopharynx and then lifting the handle toward the ceiling above the patient's feet (Fig. 3.14, steps 7 and 8). If these simple maneuvers do not result in adequate ventilation, consider the "up-down maneuver" (Fig. 3.14, steps 9 and 10). This technique is used to correct downfolding of the epiglottis, which is common with insertion of the ILMA and may interfere with ventilation or intubation. The up-down maneuver is accomplished by rotating the ILMA out of the hypopharynx along its curvature approximately 5 cm to 6 cm while the cuff remains inflated, and then sliding it back into position while pressing it against the posterior pharynx. Do not use excessive force when placing or adjusting the ILMA.

If adjusting maneuvers do not result in adequate ventilation, it is likely that the wrong size ILMA has been used. Incorrect ILMA size is more likely to be a problem if the device is too small; attempting insertion of a larger ILMA is a reasonable first approach. If another ILMA size is not available, external anterior neck manipulation or downward pressure may bring the glottis and ILMA cuff into proper alignment. If the size of the ILMA is not in question, consider completely removing and carefully reinserting the device (see Chapter 4 for intubation through the ILMA and ILMA removal).

Nonintubating LMAs
The following steps describe the use of the LMA Classic (Teleflex) (or single-use LMA Unique, Teleflex), though these instructions will be similar for other LMAs that do not have a handle similar to the LMA Fastrach (Teleflex) or LMA Supreme (Teleflex).

The first step is to select the appropriate size LMA. The LMA is available in a wide range of sizes, from size 1 for neonates weighing less than 5 kg to size 6 for adults weighing more than 100 kg. The disposable version is available in sizes 1 through 5, but not size 6. After selecting the proper size, completely deflate the LMA cuff while pushing it posteriorly so that it forms a smooth wedge shape without any wrinkles (Fig. 3.15, step 1). Place a small amount of water-based lubricant onto the posterior surface of the LMA just before insertion (Fig. 3.15, step 2). The best patient position for insertion of the LMA is the sniffing position, with the neck flexed and the head extended. The LMA may be inserted via two different techniques, depending on access to the patient. The most common method is the index finger insertion technique. This is accomplished by holding the LMA like a pen, with the index finger at the junction of the airway tube and the cuff (Fig. 3.15, step 3). Have an assistant open the patient's mouth and insert the LMA with the posterior tip pressed against the hard palate just behind the upper incisors (Fig. 3.15, step 4). Under direct vision, use the index finger to slide the LMA along the hard palate and into the oropharynx (Fig. 3.15, step 5). As the LMA is inserted farther, extend the index finger and push the posterior cuff along the soft palate and posterior pharynx. Exert counterpressure on the back of the patient's head during insertion. Continue to push the LMA into the hypopharynx until resistance is felt. Use the other hand to hold the proximal end of the LMA tube while removing your index finger from the patient's mouth (Fig. 3.15, step 6).

An alternative method is the thumb insertion technique. Use this technique when you have limited access to the patient from behind (see www.lmana.com for details). Hold the LMA with your thumb at the junction of the cuff and the airway tube. Place the mask against the hard palate under direct vision, as with the index finger technique. Use the thumb to push the LMA into the mouth along the palate and posterior pharynx. Hold the end of the airway tube with the other hand while removing your thumb from the patient's mouth.

After the LMA is fully inserted, let go of the proximal end of the airway tube and inflate the cuff enough to achieve a good seal over the glottis (Fig. 3.15, step 7). This may require only half the maximum cuff volume. Be careful to not overinflate the LMA cuff (see the product packaging for maximal cuff volumes). Attach a bag and ventilate the patient, with chest rise, breath sounds, and capnography used to confirm adequate gas exchange (Fig. 3.15, step 8). If bagging is easy and ventilation is good, the aperture of the LMA is probably aligned correctly over the glottic opening. Proper positioning of the LMA aperture with the glottic opening allows optimal ventilation.

Several tips or techniques should be considered if LMA ventilation is inadequate. The best way to ensure proper ventilation is to optimize the insertion technique by carefully following the aforementioned directions. Position the patient's head and neck properly and ensure that the patient is deeply anesthetized or paralyzed. Listen for an audible cuff leak to make sure that a good mask seal has been achieved. Adjust the cuff volume if necessary to improve the mask seal and ensure optimal ventilation. Simply adding more air to the cuff will not necessarily improve the seal of the mask with the glottis. Cuff overinflation may cause a leak, but deflation and repositioning may improve the seal.

Sometimes adjusting the patient's head and neck position is easier than trying to change the position of the LMA. Move the patient into a better sniffing position or into the chin-to-chest position to see whether this improves the LMA cuff seal. If these positions do not help or are not possible, try a jaw-thrust or a chin-lift maneuver. Apply anterior neck pressure to help manipulate the glottis into improved contact with the LMA mask. This technique can be used in combination with any of the maneuvers just discussed.

If mask seal and ventilation are still not optimal after simple repositioning maneuvers, withdraw, advance, or rotate the LMA cuff. Another alternative is to completely remove and reinsert the LMA while paying careful attention to the details just described. If unsuccessful, change the size of the LMA. A larger LMA will usually improve ventilation even if it is more difficult to insert. It is much more common to need to increase the LMA size than to decrease it. Finally, consider using the ILMA, placing a King LT, or performing a surgical airway when ventilation with the LMA is not adequate.

Aftercare
If the LMA or ILMA will remain in place without tracheal intubation, either one can be secured like an ET tube. Removal of the ILMA after tracheal intubation is easy, but more difficult than insertion of the device (see Chapter 4).

Complications
The most important complications associated with using the LMA are aspiration of gastric contents and hypoxia. The LMA does not protect against aspiration and may actually cause vomiting if the patient gags during placement of the device. In fasted anesthetized patients, the incidence of aspiration is very low, approximately 2 per 10,000 cases. There are many descriptive studies and case reports of the use of an LMA for difficult airways with no mention of significant aspiration. Although the risk for aspiration is surely higher than 2 per
Retroglottic Airway Devices

King LT

The King LT (King Airway-LTS-D EMS, King Systems, Noblesville, IN; www.kingsystems.com) is a retroglottic device that functions similar to the esophageal-tracheal Combitube (see later). Like the Combitube, the LT is designed to isolate the glottic opening between an oropharyngeal cuff and an esophageal cuff (Review Box 3.4). Unlike the Combitube, the King LT has only one airway lumen and a simplified cuff system, so both cuffs can be inflated from a single port. The literature regarding the King LT is confusing because many versions of the device have been clinically tested during the last decade. The latest disposable versions are the LT-D and...
Retroglottic Airway Devices

**Indications**
- Primary airway in the emergency medical services setting
- Primary airway in cardiac arrest
- Failed rapid-sequence intubation
- Difficult bag-mask ventilation
- Difficult intubation
- Facial trauma
- Obesity

**Contraindications**
- Limited mouth opening
- High airway pressure
- Inadequate paralysis or sedation

**Complications**
- Inability to ventilate
- Aspiration (rare)

Specifically for the King LT device:
- Tracheal placement
- Tongue edema

**Equipment**

Review Box 3.4 Retroglottic airway devices: indications, contraindications, complications, and equipment.

The LTS-D. The LTS-D has an 18-Fr gastric suction port at the tip. The modern LTS-D has been available since 2004 and is also called the LTS II in some literature. The King LT is designed for blind placement and has a large proximal cuff and small distal cuff. Unlike the Combitube, the tip of the King LT is designed to be placed in the esophagus only. The shape of the King LT and the size of the tip in previous versions made it unlikely to be placed into the trachea. However, the latest design of the LTS-D has a narrower tip and in one study had a 10% incidence of tracheal placement. Interestingly, most patients with tracheal placement of the LTS-D were still able to be adequately ventilated. Popularity of the LTS-D has grown rapidly in EMS systems, and it is now widely used by EMS agencies in the United States. Several studies have shown that the LTS-D has a high rate of successful ventilation in the operating room setting. In addition, there are several case reports of the LT being used as a rescue device for the cannot-intubate/cannot-ventilate scenario and when placement of an LMA failed. Some data suggest that the LTS-D may be useful in neonates and small infants when direct laryngoscopy fails, though the smallest LT-D currently available in the United States is size 2 (patients 12 kg to 25 kg), and the smallest LTS-D is size 3 (patients 4 to 5 feet tall). In the EMS setting, the LT-D has a high success rate (95%) when used for ventilation of out-of-hospital cardiac arrest. One study found a higher first-attempt success rate for paramedics using the King LT in cardiac arrest compared to ET intubation. In addition, Frascone and colleagues found that the rate of successful insertion and ventilation with the LTS-D is essentially equivalent to that of standard ET intubation in the hands of paramedics.

Indications and Contraindications
In the ED, the King LT is a good choice as a primary airway in patients who are unresponsive or in cardiac arrest, especially in the uncontrolled prehospital environment. The King LT can be used in any emergency airway setting for rescue ventilation after failed BMV or failed intubation. In cases of failed intubation with an unexpectedly difficult airway, the King LT may be used to provide adequate ventilation and allow time for other methods of intubation or a controlled surgical airway.

Because the King LT is a supraglottic airway and is designed to be placed blindly, it is relatively contraindicated in patients with obstruction of the upper airway by a foreign body, and should not be used in patients with an intact gag reflex.

Placement of the King LT
The first step is to choose the proper size King LT. The LTS-D is available only in adolescent and adult sizes in the United States. Size 3 is yellow and designed for patients 4 to 5 feet in height, size 4 is red and designed for patients 5 to 6 feet in height, and size 5 is purple and designed for patients taller than 6 feet. Several pediatric sizes are available in Europe, but not in the United States. The smallest LT-D available in the United States can be placed in patients weighing as little as 12 kg (approximately 18 months old).

After determining the appropriate size King LT, check the cuffs and then completely deflate them before placement. Lubricate the device with a water-based lubricant. The best patient position for insertion of the King LT is the sniffing position, but it can be placed with the head in the neutral position if necessary. Hold the LT at the connector with the dominant hand and hold the mouth open by grasping the chin from the nondominant hand. Introduce the tip of the device into the corner of the mouth while rotating the tube 45 to 90 degrees so that the blue orientation line on the tube is touching the corner of the mouth. Pass the tip of the device into the mouth and under the tongue. As the tip passes under the base of the tongue, rotate the tube back to the midline so that the blue orientation line faces the ceiling. Without exerting force,
advance the King LT until the connector is aligned with the teeth. Inflate the cuffs with the minimum volume necessary to create a good seal (see the product brochure for maximum cuff volumes). While ventilating with a bag-valve system, gently withdraw the King LT until ventilation becomes easy and free flowing. Confirm placement with chest rise, breath sounds, and capnography (Fig. 3.16).

Complications
It is hard to assess the complication rate of the King LT because the device has been modified several times in the last decade and there is no organized surveillance of out-of-hospital airway devices. The current LT-D and LTS-D devices have been available since 2004. The LTS-D is referred to as the LTS II in some studies. The most serious complication is tracheal placement, which occurred in 10% of cases in one study and is probably significantly underappreciated and underreported. Another complication that is not uncommon and certainly underreported is tongue edema. There is one case report of massive tongue edema occurring 3 hours after placement of the King LT, and mild tongue edema is relatively common and not reported in the literature. Fig. 3.17 depicts probable pharyngeal or esophageal perforation, with massive subcutaneous neck and face emphysema following prehospital placement of a King LT. Because an autopsy was denied, the exact injury was never confirmed and may have been related to other interventions during resuscitation.

There is some concern that the large oropharyngeal balloon of the King LT might compress the carotid arteries and be detrimental in patients undergoing CPR, but currently this is only suggested in animal models; no high-quality human research has been conducted to date, though one small study that reviewed computed tomography scans of the neck for 17 patients with EGAs in place found no evidence of mechanical compression of the carotid arteries on image review.

Combitube and EasyTube
The esophageal-tracheal Combitube (Nellcor, Pleasanton, CA; www.nellcor.com) is a retroglottic airway device designed as a rescue device for difficult and emergency airways and can be placed blindly and rapidly. The Combitube (Fig. 3.18) has two parallel lumens, a small distal cuff, and a large proximal cuff. When it is placed blindly the tip will end up in the esophagus in approximately 95% of cases and in the trachea in approximately 5%. The longer lumens are used for ventilation when the tip is in the esophagus. It is perforated at the level of the pharynx and occluded at the distal end. The shorter lumen or tube is used for ventilation when the tip is in the trachea. It is open at the distal end, like a standard ET tube. The large proximal cuff or balloon is designed to occlude the pharynx by filling the space between the base of the tongue and the soft palate. The small distal cuff serves as a seal in either the esophagus or the trachea.

The Combitube provides adequate ventilation in approximately 95% of patients when placed by prehospital providers, and in nearly 100% of patients when placed by physicians. It compares favorably with the ET tube with respect to ventilation and oxygenation in cardiac arrest situations. In unconscious patients, the Combitube may provide protection from aspiration.

Indications and contraindications for the Combitube are similar to the King LT airway. Because the King is easier to use and proven to be an effective and reliable primary and rescue airway device, the Combitube is used less frequently than previously.

Placement of the Combitube
The Combitube is available in two sizes. The manufacturer recommends the smaller 37-Fr device for patients from 4 feet to 5 feet 6 inches tall and the larger 41-Fr device for patients taller than 5 feet 6 inches. Studies suggest that the smaller 37-Fr Combitube can be used safely in patients up to approximately 6 feet tall. The larger 41-Fr device is appropriate for patients taller than 6 feet.

To insert the Combitube, hold the device in the dominant hand and gently advance it caudally into the pharynx while grasping the tongue and jaw between the thumb and index finger of the nondominant hand. Pass the tube blindly along the tongue to a depth that places the printed rings on the proximal end of the tube between the patient's teeth and the alveolar ridge. If resistance is met in the hypopharynx, remove...
the tube and bend it between the balloons for several seconds to facilitate insertion. After insertion, fill the pharyngeal balloon with 100 mL of air and the distal cuff with 10 mL to 15 mL of air. The large pharyngeal balloon serves to securely seat the Combitube in the oropharynx and creates a closed system in the case of esophageal placement. Because approximately 95% of placements are esophageal, begin ventilation through the longer (blue) airway tube. Use chest rise, breath sounds, and capnography, without gastric inflation, to confirm proper placement. When the distal tip is in the esophagus, ventilation occurs through the vent holes between the distal and proximal cuffs (white arrows). If the tip of the Combitube is in the trachea, ventilation cannot be accomplished via the long (blue) airway tube. It is essential to recognize this quickly, and use the short (white) tube for ventilations.

Complications
Inappropriate balloon inflation and incorrect Combitube placement can lead to air leaks during ventilation. The most common placement error is an improper insertion angle. Use a more caudal, longitudinal direction of insertion as opposed to an anteroposterior direction of insertion. The Combitube must also be maintained in the true midline position during insertion to avoid blind pockets in the supraglottic area, which can prevent passage of the tube.

CONCLUSION
Good basic airway skills and the familiarity with and availability of proven rescue devices are the keys to emergency airway management. There are many techniques and devices that can be used to manage emergency airways. In difficult situations, providers will probably have the best success when basic skills are performed excellently.


Tracheal Intubation

Brian E. Driver and Robert F. Reardon

Intubation is often the pivotal procedure in the emergency management of critically ill patients. There are several new devices that can improve the likelihood of successful intubation, and it is important to put the intubation procedure in perspective, understanding that intubation is just one part of airway management. The primary objective of airway management is to maintain adequate ventilation and oxygenation, and intubation only allows ventilation and oxygenation after the procedure is completed, not during the procedure. It is critical to have good basic airway management skills (see Chapter 3). It is also important to use intubation techniques that have a high likelihood of first pass success, to have the wisdom to recognize when a given approach has failed, and to quickly move to a different technique. Use of an emergency airway algorithm can help providers make difficult decisions in a timely manner (see later section on Emergency Airway Algorithm).

This chapter describes nearly every possible means of tracheal intubation, with emphasis on widely used techniques. The most common means of intubation in the emergency setting is rapid-sequence intubation (RSI), and this approach must be considered very carefully. If a difficult intubation is anticipated, awake intubation may be preferred. Although flexible endoscopic devices are commonly used, with good topical anesthesia, nearly any intubating technique can be used for awake intubation. Because many difficult airways cannot be predicted, it is essential to have a well-defined backup plan and appropriate resources. The intubating laryngeal mask airway (LMA Fastrach, Teleflex, Inc, Wayne, PA) is an ideal backup device for failed RSI because it provides ventilation and oxygenation and facilitates tracheal intubation in a high percentage of patients with failed RSI.

**GENERAL APPROACH TO EMERGENCY INTUBATION**

Preplanning is the key to successful emergency airway management. Providers should follow a clear, preconceived, practiced airway algorithm that uses readily available and familiar equipment and techniques. When encountering a difficult airway, it is more important to be comfortable with a few well-proven devices and techniques than to try new or unfamiliar devices. It is important to know when to abandon one approach and move on to the next. No single approach is mandated. The best technique is the one chosen by the clinician at the bedside based on one’s individual experience and expertise given the specific clinical scenario.

A critical aspect of preparation is making sure that all essential equipment required to perform the airway maneuvers is immediately available and within easy access. This may be accomplished by wall-mounting essential equipment in the emergency department (ED) resuscitation room. Alternatively, equipment can be placed in dedicated adult and pediatric airway carts or tackle boxes in an open, organized, and labeled manner, with the carts and boxes checked and stocked regularly (Fig. 4.1). Essential equipment that is seldom used but is potentially lifesaving should be clearly identified and placed in an easily accessible location such as a dedicated difficult airway cart.

The importance of this concept cannot be overstated. Technical expertise cannot substitute for the lack of essential equipment. In airway management, failure has ominous consequences. Mental, physical, and equipment preparation maximizes the chance of success.

**AIRWAY ANATOMY**

An understanding of airway anatomy and its terminology is requisite for any discussion of airway management procedures (Fig. 4.2). The following terms are used frequently in this chapter:

- **Pharynx**: the upper part of the throat posterior to the nasal cavity, mouth, and larynx
  1. **Nasopharynx**: base of the skull to the soft palate
  2. **Oropharynx**: soft palate to the epiglottis
  3. **Hypopharynx**: epiglottis to the cricoid ring (posteriorly), including the piriform sinus/recess/fossa

- **Piriform sinus/recess/fossa**: the pockets on both sides of the laryngeal inlet separated from the larynx by the aryepiglottic folds

- **Larynx**: the anterior structures of the throat (commonly called the voice box) from the tip of the epiglottis to the inferior border of the cricoid cartilage, including the laryngeal inlet
  1. **Laryngeal inlet**: the opening to the larynx bounded anterosuperiorly by the epiglottis, laterally by the aryepiglottic folds, and posteriorly by the arytenoid cartilage

- **Arytenoid/posterior cartilage**: the posterior aspect of the laryngeal inlet separating the glottis (anterior) from the esophagus (posterior)
  1. **Corniculate cartilage**: the medial portion of the arytenoid/posterior cartilage
  2. **Cuneiform cartilage**: the lateral prominence of the arytenoid/posterior cartilage
  3. **Interarytenoid notch**: the notch between the posterior cartilage

- **Glottis**: the vocal apparatus, including the true and false cords and the glottic opening
- **Vallecula**: the space between the base of the tongue and the epiglottis

- **Hyoepiglottic ligament**: anterior midline ligament connecting the epiglottis to the hyoid bone

**PREPARATION**

Intubation is best accomplished with two operators, one to perform the intubation and the other to handle equipment, help with positioning, observe the patient and monitor, and keep track of time. Unfortunately, the ideal scenario and adequate time for preparation are not always available to the clinician, who has to make calculated adjustments based on the situation at hand. Before intubating, it is preferable to take the following steps in chronologic order: (1) attach the necessary monitoring devices and administer oxygen, (2) establish intravenous
Figure 4.1 A, Adult airway cart. Equipment and materials are visible, labeled, and accessible. B, Labeling is especially important because it lets you know what is missing. (Concept of Dr. Ernest Ruiz, Department of Emergency Medicine, Hennepin County Medical Center, Minneapolis.)

Figure 4.2 A, Anatomy of the upper airway. B, View of the larynx, epiglottis, and vocal cords seen with a laryngoscope. The key to a successful intubation is identification of the vocal cords. (A, Netter illustration used with permission of Elsevier, Inc. All rights reserved.)
access, (3) draw up essential medications and label them if time permits, (4) confirm that the intubating equipment is available and functioning, (5) reassess oxygenation and maximize preoxygenation, (6) position the patient correctly, and (7) make sure that all team members are aware of the primary procedural approach and the most likely backup plan. In the haste of the moment, it is a common error to forget to preoxygenate or to position the patient optimally. Simple omissions, such as failing to restrain the patient’s hands or remove the patient’s dentures or misplacing the suction tip, can seriously hamper the success of the procedure. Utilize universal precautions by wearing gloves, a gown, and eye and mouth protection.

The concept of using checklists to decrease medical errors and improve patient care has grown since a landmark article demonstrated decreased complications and mortality in surgical patients when checklists were utilized.7 Timely and successful tracheal intubation, while minimizing hypoxemia and other complications, is paramount when caring for ill and injured patients who require intubation. Pre-intubation checklists can reduce cognitive load for the intubating physician by creating a framework for approaching all emergency intubations. Checklists prompt clinicians to verify that all necessary equipment is available and functioning, to perform a standardized airway assessment, to execute optimal preoxygenation, and to develop an airway plan with patient-specific backups. The use of pre-intubation checklists has been shown to reduce peri-intubation complications in trauma patients.8 Example checklists are seen in Box 4.1.

**BOX 4.1 Preintubation Checklists**

**PRE-ARRIVAL**
1. Bag mask ventilation setup with oxygen running at >15 L/min
2. Suction connected and running
3. Laryngoscope functioning and ready
4. Ready an endotracheal tube: check cuff, insert stylet, and have a “straight to cuff” shape with a 35-degree distal bend
5. Back-up devices such as laryngeal mask airway or King LT airway available and ready
6. Cricothyrotomy set located
7. End-tidal CO2 detector ready

**PRE-INTRUBATION**
1. Assess the airway: open mouth, examine neck mobility, palpate anterior neck
2. Decide best approach: awake, sedated, or RSI
3. Communicate intubation medication orders to nurses, including post-intubation medications
4. Optimally position patient
5. Preoxygenate patient (usually with face mask oxygen at 60 L/min)
6. Apply nasal cannula at 10–15 L/min in preparation for apneic oxygenation
7. Discuss airway plan with entire team
8. Ensure functioning pulse oximeter
9. Ensure patient intravenous catheter
10. Ensure assistants are ready (nurses, respiratory therapists)

RSI, Rapid-sequence intubation.

**PREOXYGENATION**

Preoxygenation is one of the most important aspects of emergency airway management. The goal of preoxygenation is to replace all the nitrogen in the lungs with oxygen prior to the start of intubation attempts. This allows the lungs to act as an oxygen reservoir during the apneic period of RSI. This provides the intubator with additional time before the onset of hypoxemia, and significantly increases the chance for successful intubation on the first attempt. It is not enough to achieve a peripheral oxygen saturation (SPO2) value of 100% prior to intubation, because an SPO2 of 100% does not necessarily correspond with denitrogenation of the lungs; furthermore, partial pressure of arterial oxygen (PaO2) at 100% SpO2 can range from approximately 100 mm Hg to 600 mm Hg. Failure to preoxygenate before RSI is often a critical factor when a straightforward emergency airway becomes an unexpected airway problem. Those at greatest risk for rapid desaturation include obese, pregnant, critically ill, and pediatric patients; these populations will benefit most from optimal preoxygenation.

Preoxygenate by providing the maximal fraction of inspired oxygen (FiO2) with a simple face mask or non-rebreather mask for 3 to 5 minutes before intubation.9,10 The type of mask is less important than the oxygen flow rate; at very high flow rates the FiO2 for any device usually exceeds 90%. At lower flow rates (< 30 L/min) the FiO2 will not be high enough for adequate preoxygenation,12 so the oxygen flow rate should be at least 30 L/min. When using a standard oxygen flow meter this requires turning it up as high as possible, beyond the marked maximum of 15 L/min, to the “flush” rate. The flush rate is usually marked on each flowmeter and is typically greater than 40 L/min. Oxygen flowmeters that can measure up to 70 L/min, with flush rates up to 90 L/min, are available (see Fig. 3.7 in Chapter 3); however, a flush rate greater than 40 L/min is probably sufficient for maximal preoxygenation. Providing high flow oxygen washes out expired CO2, fills the dead space of the nasopharynx and upper airway with oxygen, compensates for any leak between the mask and the patient to avoid entrainment of room air during inspiration, and may provide low levels (1 to 2 cm H2O) of positive airway pressure.13 The importance of very high flow oxygen administration during preoxygenation cannot be overemphasized. If possible, instruct the patient to exhale maximally before beginning preoxygenation.14

Alternatively, if it is not possible to perform preoxygenation for 3 to 5 minutes prior to intubation, instruct the patient to take eight vital capacity breaths while delivering very high flow oxygen, to provide nearly the same result.9,11 Many critically ill patients will not be able to take vital capacity breaths; therefore with time permitting, the preferred method is 3 to 5 minutes of tidal breathing.

Unlike face masks, bag-mask ventilation (BMV) requires proper equipment and good technique to achieve adequate preoxygenation. Bags without one-way valves for inhalation and exhalation will not function properly during spontaneous ventilation, and will provide only room air.16 Furthermore, unless flow rates are very high (>40 L/min), the mask must be sealed perfectly to the patient's face (see Chapter 3 for proper technique).17 If the seal is imperfect, room air will be entrained and FiO2 will be close to that of room air. This is analogous to holding a mask above a patient's face, which likewise provides oxygen content near that of room air. In spontaneously breathing patients, face masks (with a very
high flow rate) are the preferred oxygen delivery device for preoxygenation.

To augment oxygen delivery and prepare for apneic oxygenation (see next section), apply a nasal cannula (at 15 L/min) to the patient during preoxygenation, simultaneously with other preoxygenation efforts.18 High flow nasal cannula (see Chapter 3 for details), if available, may also improve preoxygenation,20 though this has never been studied in combination with a high-flow face mask.

The preferred position for preoxygenation is head elevation of 20 to 25 degrees. This position minimizes atelectasis, decreases the pressure of the abdominal contents against the diaphragm, and allows the patient to continue taking deep breaths. In both obese20,21 and non-obese adults22,23 this position has been demonstrated to be advantageous for preoxygenation. For patients with spinal immobilization, the bed can be placed in 25 degrees of reverse Trendelenburg (head up) to achieve the same effect.24

If SpO2 cannot be increased above 93% to 95% after optimal preoxygenation, the addition of positive pressure using non-invasive positive pressure ventilation (NPPV) or mask ventilation with a positive end-expiratory pressure valve may improve oxygenation prior to intubation attempts.25

Sometimes patients who will benefit the most from preoxygenation are uncooperative because of delirium from hypoxia, hypercapnia, or other factors. Application of a face mask or NPPV may be difficult or impossible. These patients may benefit from careful sedation without suppression of respirations, allowing for oxygenation with a face mask or NPPV for 2 to 3 minutes before administration of a paralytic agent (also known as delayed sequence intubation). Ketamine (1 to 1.5 mg/kg by slow intravenous push) has been suggested for this technique.25 Weingart and colleagues performed sedation for preoxygenation in 62 adults, and demonstrated an increase in SpO2 for most patients without any adverse events.26 Because of the small sample size and because these patients were managed by experts in airway management and sedation, this technique is not necessarily generalizable to everyday clinical practice. If a patient is sedated for preoxygenation, the clinician should be vigilant for respiratory depression, apnea, and airway obstruction, and have all airway equipment available in case emergency control of the airway or breathing becomes necessary. In many cases, it may be safer to restrain the patient without sedation to facilitate preoxygenation.

### Apneic Oxygenation During Intubation

Another method to delay desaturation during RSI is nasopharyngeal oxygen insufflation without ventilation, termed apneic oxygenation. Even in the absence of ventilation, oxygen is able to travel down the tracheobronchial tree to the alveoli and diffuse into the bloodstream, where it is consumed and converted into carbon dioxide. Because oxygen diffuses across the alveoli much more readily than carbon dioxide, because oxygen and carbon dioxide have differences in gas solubility in blood, and because of the high affinity of hemoglobin for oxygen, more oxygen leaves the alveoli than carbon dioxide enters. This creates a pressure gradient that causes oxygen to travel from the nasopharynx to the alveoli and into the bloodstream.22

Studies have shown that providing oxygen therapy during apnea is much more beneficial than one might anticipate.27-29 Multiple studies conducted in the operating room, in normal and morbidly obese patients, have shown that nasopharyngeal oxygen insufflation results in a significant delay in desaturation after the onset of apnea, with many subjects never developing hypoxemia even after six minutes of apnea.27-30

A 2016 randomized trial in an intensive care unit (ICU) found no difference in the oxygen saturation nadir between patients who received and did not receive apneic oxygenation during intubation.32 Because ED patients are often intubated within minutes of arrival despite limited history and sometimes limited preoxygenation, and because, contrary to ICU patients, ED patients generally have not been on supplemental oxygen for the several hours preceding intubation, the results of this study should not be generalized to ED care.

Perform apneic oxygenation with every tracheal intubation to decrease the chance of severe hypoxemia. Place a standard nasal cannula beneath the main preoxygenation device (face mask or bag-valve mask). If the patient is awake, limit the flow rate to 5 to 15 L/min during the preoxygenation phase because higher flow rates can be uncomfortable. If the patient is comatose or unresponsive, set the nasal cannula to 15 L/min or higher when initially placed. When the preoxygenation device is removed for intubation, keep the nasal cannula in place. During intubation attempts, set the nasal cannula to at least 15 L/min. It may be beneficial to turn the oxygen flowmeter up as high as possible because higher flow rates have been shown to provide higher FiO2.13 If there is nasal obstruction, place a nasopharyngeal airway in one or both nares to facilitate oxygen delivery to the posterior nasopharynx. To optimize gas flow past the upper airway, position the patient for tracheal intubation, and perform maneuvers to ensure upper airway patency (i.e., jaw thrust, head tilt/chin lift). Because conventional nasal cannula oxygen delivery is not humidified, apneic oxygenation at high flow rates will cause some desiccation of the nasopharynx, but this should not cause significant harm because of the short duration of this oxygen supplementation.

High-flow nasal cannula systems may be an even better method of apneic oxygenation, and have been shown in an ICU study to be superior to simple nasal cannula, though both groups had low rates of hypoxemia.19

### ASSESSING FOR A DIFFICULT AIRWAY

When trying to predict whether there will be difficulty during emergency intubation, it is important to understand that most of the literature on prediction of difficult laryngoscopy does not apply very well in the emergency setting. A study by Levitan and colleagues in 2004 showed that two-thirds of patients who were intubated in their ED via RSI could not be assessed with the most common difficult airway prediction tests (Mallampati scoring, measurement of thyromental distance, and neck mobility testing) because of altered mental status or cervical spine immobilization.34 Even in the best circumstances, only approximately half the cases of difficult laryngoscopy can be predicted.35 Many factors such as Mallampati scoring and measurement of the thyromental distance have not been found to accurately predict difficult laryngoscopy, especially in the emergency setting.34-37 Only obvious anatomic and pathologic abnormalities and a history of difficult intubation are accurate predictors of difficult laryngoscopy.14 The American Society of Anesthesiology Difficult Airway Guidelines state that “in patients with no gross upper airway pathology or anatomic anomaly, there is insufficient published evidence to evaluate
the effect of a physical examination on predicting the presence of a difficult airway. This does not mean that emergency providers should ignore factors that are known to be associated with difficult laryngoscopy, but these must be placed in perspective.

No examination finding alone can predict difficult laryngoscopy, and a combination of multiple factors makes difficulty more likely. The classic predictors of difficult intubation include a history of previous difficult intubation, prominent upper incisors, limited ability to extend at the atlanto-occipital joint, poor visibility of pharyngeal structures when the patient extends the tongue (Mallampati classification or the tongue-pharyngeal ratio) (Fig. 4.3A), limited ability to open the mouth (suggested by a space less than three fingerbreadths between the upper and lower incisors), short thyromental distance (< 6 cm from the thyroid notch to the chin with the neck in extension) (see Fig. 4.3B), and a limited direct laryngoscopic view of the laryngeal inlet (Fig. 4.4). A relatively new test is the upper lip bite test, which has been shown in some studies to be more accurate and specific than older tests. It is essentially a test of anterior mandibular mobility, and the less mobility the more difficult it is to intubate the patient. Upper lip bite criteria are as follows: class I, the lower incisors can bite the upper lip above the vermilion line; class II, the lower incisors can bite the upper lip below the vermilion line; and class III, the lower incisors cannot bite the upper lip.

Many of these predictors cannot be assessed in the emergency setting. Some of the key predictors are apparent simply by observing the external appearance of the patient's head and neck. Patients with neck tumors, thermal or chemical burns, traumatic injuries involving the face and anterior aspect of the neck, angioedema, infection of pharyngeal and laryngeal soft tissues, or previous operations in or around the airway suggest a difficult intubation because distorted anatomy or secretions may compromise visualization of the vocal cords. Facial or skull fractures may further limit airway options by precluding nasotracheal (NT) intubation. Patients with ankylosing arthritis or developmental abnormalities such as a hypoplastic mandible or the large tongue of Down's syndrome are difficult to intubate because neck rigidity and problems of tongue displacement can obscure visualization of the glottis.

Besides these obvious congenital and pathologic conditions, the presence of a short, thick neck is one of the more common predictors of a difficult airway. Such individuals are easily identifiable by observing the head and neck in profile. Obesity alone may not be an independent predictor of difficult intubation, but obese patients with large-circumference necks are likely to be difficult to intubate. Facial hair can complicate a difficult airway by rendering BMV ineffective because of the lack of a good mask seal. One patient type that does not immediately stand out as a difficult intubation, but can be surprisingly so, is a patient with an unusually long mandibulothyroid distance (the thyroid prominence appearing low in the neck) and a short mandibular ramus. Visualization of the larynx is difficult because of the distance to the larynx and the relative hypopharyngeal location of the tongue.

Knowledge of the poor performance of difficult airway predictors should not make emergency providers more cavalier about using RSI; rather, it should create more concern. The solution is not to avoid RSI because lack of paralysis makes every intubation more difficult. Even though predicting a difficult airway is challenging, and even though a comprehensive airway assessment may not be possible because of patient behavior or clinical circumstances, performing an assessment of the airway is important and serves as a cognitive forcing strategy that guides the physician to plan and prepare for the airway intervention. It is essential to appreciate the critical importance of having a clear backup plan when intubation

aggressive management in some situations or to irreversible hypoxic injury as a result of hesitation in others. Deciding who requires a definitive airway and who needs only supportive measures is a formidable task for even the most skilled clinician.

The following parameters should be assessed before the decision is made to establish a definitive airway:

- Adequacy of current ventilation
- Potential for hypoxia
- Airway patency
- Need for neuromuscular blockade (uncooperative, full stomach, teeth clenching)
- Cervical spine stability
- Safety of the technique and skill of the operator

Consideration of these factors should guide the clinician in deciding if tracheal intubation is necessary, and in selecting the optimal technique. Choosing the initial approach is often straightforward. Difficulty arises precipitously when the initial approach fails. Time becomes critical as the risk for irreversible hypoxic injury and cardiac arrest rises. Anxiety then increases the potential for error. Forethought and practice are invaluable when managing these situations.

**Direct Laryngoscopy**

**Indications**
- Routine emergency intubation
- Difficult airways

**Contraindications**
- Limited mouth opening
- Upper airway distortion or swelling
- Kyphosis (extreme curvature of the upper back)
- Copious blood or secretions

**Complications**
- Hypoxic brain injury
- Cardiac arrest
- Aspiration
- Upper airway trauma
- Dental trauma

**Equipment**
- Laryngoscope
- Bag-valve-mask
- Endotracheal tube with stylet
- Bougie
- Yankauer suction
- Nasal and oral airways
- 12-mL syringe
- PETCO2 detector
- Tape
- Nasal and oral airways
- Endotracheal tube with stylet
- Bougie

**Review Box 4.1** Direct laryngoscopy: indications, contraindications, complications, and equipment.

with RSI fails. This situation mandates the need for a preconceived algorithm that uses proven rescue techniques applicable to a broad range of clinical scenarios, such as the bougie (flexible intubating stylet) and the intubating LMA Fastrach (Teleflex). The value of the bougie is indisputable, and it is clear that using the intubating LMA Fastrach after failed RSI has decreased the frequency of failed airways and the need for surgical intervention. Because RSI is the “go-to” method for emergency intubation, providers must be prepared to perform a surgical airway when laryngoscopy, BMV, and backup devices fail.

**EMERGENCY AIRWAY DECISION MAKING**

The airway provider must have many tools readily available to deal with an acutely compromised airway. It is important to be proficient in a number of different techniques and to tailor their use to the needs of the individual patient. Rescuers should practice potential scenarios before facing patients with a compromised airway. Failure to do so may lead to unnecessarily aggressive management in some situations or to irreversible hypoxic injury as a result of hesitation in others. Deciding who requires a definitive airway and who needs only supportive measures is a formidable task for even the most skilled clinician.

The following parameters should be assessed before the decision is made to establish a definitive airway:

- Adequacy of current ventilation
- Potential for hypoxia
- Airway patency
- Need for neuromuscular blockade (uncooperative, full stomach, teeth clenching)
- Cervical spine stability
- Safety of the technique and skill of the operator

Consideration of these factors should guide the clinician in deciding if tracheal intubation is necessary, and in selecting the optimal technique. Choosing the initial approach is often straightforward. Difficulty arises precipitously when the initial approach fails. Time becomes critical as the risk for irreversible hypoxic injury and cardiac arrest rises. Anxiety then increases the potential for error. Forethought and practice are invaluable when managing these situations.
Clinicians who perform emergency intubation, especially RSI, must understand that oxygenation and ventilation, not tracheal intubation, are paramount when caring for critically ill patients. Patients who are hypoxic on arrival and those who develop hypoxia after a failed first intubation attempt need good BMV to restore oxygenation and keep them stable enough for further intubation attempts. The importance of BMV skills cannot be overstated (see Chapter 3), and mastery of this skill alleviates much of the anxiety associated with difficult emergency airways and improves the chance for successful RSI. It is critically important to have an extraglottic airway (EGA) device immediately available during every RSI in the event that BMV is difficult or impossible. All clinicians who perform emergency intubation should be prepared to perform a surgical airway when intubation methods and backup ventilation techniques fail.

### Decision to Perform Rapid-Sequence Intubation (RSI)

RSI in anesthesia has evolved since the introduction of succinylcholine in 1951. RSI was initially used as an abbreviation for rapid-sequence induction but is now synonymous with rapid-sequence intubation. Initially, the main purpose of RSI was to decrease the risk for aspiration in patients with full stomachs who needed emergency intubation. RSI has now become the most common method of emergency airway management because paralysis facilitates optimal intubating conditions in critically ill patients.

Emergency providers should be very careful to not use RSI in a cavalier manner. When giving a paralytic agent, the provider takes complete responsibility for airway maintenance, ventilation, and oxygenation of the patient. Consider one of many awake intubation (i.e., intubation without paralysis) options in patients with known or anticipated difficult airways (Box 4.2). In the emergency setting it is useful to think of difficult airways as situations in which our usual methods of intubation and backup ventilation/oxygenation techniques fail. The goal should be to avoid RSI in patients who cannot be easily intubated via the common techniques (direct or video laryngoscopy) and cannot be ventilated with a bag-mask device. Risk factors for difficult or impossible BMV have been well studied and include the presence of a beard, obesity, lack of teeth, age older than 55 years, a history of snoring, short thyromental distance, and limited mandibular protrusion (see Box 3.2 in Chapter 3). It is also prudent to consider whether an EGA device will be difficult to place because this is often the primary backup plan. RSI is contraindicated in patients who cannot be orally intubated and it should usually be avoided in patients with laryngotracheal abnormalities caused by tumors, infection, edema, or a history of cervical radiation therapy.

If the clinician decides that RSI is not appropriate for a patient, there are many options that can be performed without paralysis (see Box 4.2). Flexible endoscopic intubation is the go-to procedure for most anesthesiologists and is described later in the chapter. Direct and video laryngoscopy can also be performed without paralysis. If excellent topical anesthesia is achieved, some patients can be intubated without any sedation.

### Ideal Versus Emergency Technique

The intubation plan that would be best in the ideal/elective situation is often not the best plan in the emergency setting. Consider the patient with rapidly increasing upper airway swelling due to angioedema or anaphylaxis, causing impending complete airway obstruction. Because of predicted difficulty with direct and video laryngoscopy, some providers would not consider this patient a candidate for RSI. Other intubation strategies might have a higher chance of success if time was not a factor. The patient will likely develop complete airway obstruction, critical hypoxia, and death in the time required to set up and perform endoscopic nasal intubation. In this situation the emergency provider is “forced to act” to complete timely tracheal intubation. The best course for this patient is likely RSI with modern video laryngoscopy equipment and a good backup plan, such as the LMA Fastrach (Teleflex), followed by a cricothyrotomy. There are many other scenarios in which the emergency airway plan is much different than the ideal/elective airway plan, such as patients with severe trauma who have multiple different life-threats. In these patients, intubation often needs to be expedited so that other life threats can be addressed in a timely manner.

### Failed Airways

A failed airway should be differentiated from a known or anticipated difficult airway. A patient is considered to have a failed airway in the following situations: (1) inability to maintain oxygenation by BMV or EGA device; (2) failure of three or more intubation attempts by an experienced operator; (3) failure of the first attempt in a “forced to act” situation.

If a patient has a failed airway, and oxygenation can be maintained, the clinician should attempt intubation by another method (e.g., flexible endoscopy, intubating LMA [ILMA], video laryngoscopy, surgical airway, and others); if a more experienced provider is not already present then one should be called for. This could be an anesthesiologist, emergency physician, paramedic, or any other clinician with significant airway expertise. If at any point there is failure of oxygenation, an EGA device should be placed while preparing for a cricothyrotomy. If the EGA device cannot oxygenate, a cricothyrotomy should be performed.

### Emergency Airway Algorithm

The realization that one cannot predict all cases of failed BMV and failed laryngoscopy mandates the need for a simple pre-conceived algorithm that uses proven rescue techniques that are applicable to a broad range of clinical scenarios, such as the bougie and the LMA Fastrach (Teleflex). The value of the bougie is indisputable, and it is clear that use of the LMA
Fastrach after failed RSI has decreased the frequency of failed airways and the need for surgical airways.\textsuperscript{1,2,3,51–58} One of the most important concepts to appreciate when using RSI is that of optimal laryngoscopy to maximize first-pass success.\textsuperscript{31,66} Preparation, preoxygenation, proper patient positioning, anterior neck maneuvers, and good laryngoscopy skills are all important components of optimal laryngoscopy. Just as important as optimal laryngoscopy is the ability to recognize when laryngoscopy (or any technique) has failed and when it is time to move on to a different approach. Patient safety during RSI depends on the provider’s ability to maintain ventilation and oxygenation if the first attempt at intubation fails.\textsuperscript{51} Critical decisions about how to maintain ventilation and oxygenation and when to use a different intubation approach are much easier when using a simple preconceived algorithm. The algorithm presented here summarizes the general approach used in the Department of Emergency Medicine at Hennepin County Medical Center (Fig. 4.5). This algorithm is presented as an example. Individual providers and institutions should determine their own algorithms based on the availability of skills and resources. There are many similarities between this algorithm and those put forth by the American Society of Anesthesiologists and the Difficult Airway Society; however, our algorithm is simpler and more applicable to emergency airway management.\textsuperscript{39,67} Most published airway algorithms are not ideal for emergency airway management because they do not account for the conditions that are commonly encountered: patients with full stomachs who are critically ill and often uncooperative, and intubations that cannot be canceled if the airway is too difficult. Many algorithms resemble wish lists of equipment and skills that are simply not available to many emergency airway providers. Our algorithm is based on the concept that oxygenation, not intubation, is the key.\textsuperscript{1} It stresses well-proven concepts, procedures, and devices, and it is modeled after a simple algorithm developed by Combes and colleagues that was validated in a large prospective study.\textsuperscript{5,53,58}

**Figure 4.5** Emergency airway management algorithm used at Hennepin County Medical Center. The end point of the algorithm is successful tracheal intubation. This algorithm is presented as an example. Individual and institutions should formulate their own algorithms based on technical skills and the availability of resources. ILMA, Intubating laryngeal mask airway.

**DIRECT LARYNGOSCOPY (DL)**

Despite the proliferation of approaches and devices designed to secure a definitive airway, DL remains the mainstay of tracheal intubation. DL is a crucial skill even in the era of video laryngoscopy, and is less prone to problems such as device failure or blood and secretions covering the video lens.\textsuperscript{68} Visual confirmation of the tube going through the vocal cords is usually possible.

**Indications and Contraindications**

DL is indicated in any clinical situation in which a definitive emergency airway is necessary, including routine and difficult airways. Relative contraindications to DL include limited mouth opening, upper airway distortion or swelling, severe kyphosis, or copious blood or secretions.

**Equipment**

**Laryngoscope**

There are two basic blade designs for DL, curved (Macintosh) and straight (Miller) (Fig. 4.6). Each comes in various adult and pediatric blade sizes. Slight variations in laryngoscopic technique follow from the choice of blade design, and it is often a matter of personal preference. The tip of the straight blade goes under the epiglottis and lifts it directly, whereas the curved blade fits into the vallecula and indirectly lifts the epiglottis via engagement of the hyoepiglottic ligament to expose the larynx.

Each blade type has advantages and disadvantages. The straight blade is often a better choice in pediatric patients, in patients with an anterior larynx or a long floppy epiglottis, and in individuals whose larynx is fixed by scar tissue. It is less effective, however, in patients with prominent upper teeth, and it is more likely to damage dentition. Use of the straight blade is also more often associated with laryngospasm because it stimulates the superior laryngeal nerve, which innervates the undersurface of the epiglottis. A straight blade may inadvertently be advanced into the esophagus and initially reveal unfamiliar anatomy until it is withdrawn. The blade has a lightbulb at the tip, which may slightly hamper vision. The wider, curved blades are helpful in keeping the tongue retracted from the field of vision and allowing more room for passing the tube through the oropharynx, and they are generally preferred for uncomplicated adult intubations. Aside from
patient considerations, some clinicians prefer the curved blade because they find that it requires less forearm strength than the straight blade.

The illumination provided by the laryngoscope can make a big difference in the ability to visualize the laryngeal inlet. The importance of these factors is underappreciated, as demonstrated by Levitan in a survey of the Macintosh blades used in 17 Philadelphia EDs. It was found that only 24% of all blades provided the brightness necessary for fine inspection. This finding was largely explained by the fact that the majority of EDs used the A-Mac (American) as opposed to the clearly superior brightness design of the G-Mac (German) or the intermediate brightness of the E-Mac (English).

**Tracheal Tubes**

The standard adult endotracheal (ET) tube measures approximately 30 cm in length. Tube size is typically printed prominently on the tube and is based on the internal diameter (ID) and measured in millimeters. The range is 2.0 to 10.0 mm in increments of 0.5 mm. The outer tube diameter is 2 to 4 mm larger than the ID. Tubes are also imprinted with a scale in centimeters that indicates the distance from a tube’s distal tip.

Adult men can generally accept a 7.5- to 9.0-mm orotracheal tube, and women can usually be intubated with a 7.0- to 8.0-mm tube. Larger tubes are theoretically desirable because airway resistance increases as tube size decreases, but in practice, a 7.5-mm tube is adequate for almost all patients. In emergency intubations, particularly if a difficult intubation is anticipated, many clinicians choose a smaller tube and change to a larger tube later if necessary. Though generally an acceptable practice, this should be avoided in burn patients because swelling may prohibit subsequent tube placement. For nasal intubation, a slightly smaller (by 0.5 to 1.0 mm) tube may be easier to advance through the nasal passages.

Correct tube size is important in the pediatric population. It is especially important when using an uncuffed tube because a good seal is needed between the ET tube and the upper part of the trachea (Table 4.1). As tube size is based on the ID, a cuffed tube should generally be a half size (0.5 mm) smaller than an uncuffed tube. The smaller ID of an appropriately sized, small cuffed tube could theoretically make it more prone to plugging from secretions. Cuffed tubes are available down to 3 mm ID, although indications for these tubes in neonates and infants are rare. A cuffed tube is used in children with decreased lung compliance who may require prolonged mechanical ventilation. In a child, the smallest airway diameter is at the cricoid ring rather than at the vocal cords, as in adults. Hence, a tube may pass the cords but go no farther. If this should occur, the next smaller size tube should be passed. The American Heart Association (AHA) states that both cuffed and uncuffed tubes are acceptable for infants and children who are tracheally intubated. If a cuffed tube is placed, careful attention must be paid to cuff pressures.

In children 2 years or older, the following formula is a highly accurate method for determining correct uncuffed and cuffed ET tube size:

Uncuffed tube size (mm) = [Age (yr)/4] + 4

Cuffed tube size (mm) = [Age (yr)/4] + 3

For most clinical situations, using the width of the nail of the patient’s little (fifth) finger as a guide is sufficiently accurate and has been shown to be more precise than finger diameter

<table>
<thead>
<tr>
<th>AGE</th>
<th>CUFFED ETT SIZE (INTERNAL DIAMETER, mm)</th>
<th>EQUIVALENT TRACHEOTOMY TUBE SIZE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preterm</td>
<td>2.5</td>
<td>00</td>
</tr>
<tr>
<td>Term</td>
<td>3.0</td>
<td>00</td>
</tr>
<tr>
<td>6 months</td>
<td>3.0–3.5</td>
<td>00–0</td>
</tr>
<tr>
<td>1–2 years</td>
<td>4.0</td>
<td>0–1</td>
</tr>
<tr>
<td>3–4 years</td>
<td>4.5</td>
<td>1–2</td>
</tr>
<tr>
<td>5–6 years</td>
<td>5.0</td>
<td>2</td>
</tr>
<tr>
<td>10 years</td>
<td>6.0</td>
<td>3</td>
</tr>
<tr>
<td>12 years</td>
<td>7.0</td>
<td>4</td>
</tr>
<tr>
<td>14 years</td>
<td>See adult sizing</td>
<td></td>
</tr>
<tr>
<td>Adults</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>7.0–8.0</td>
<td>5</td>
</tr>
<tr>
<td>Male</td>
<td>7.5–9.0</td>
<td>6</td>
</tr>
</tbody>
</table>

(Fig. 4.7) In order to reduce cognitive load during emergency situations, it may be best to use a length-based tool to determine the appropriate size ET tube, tube depth, and other resuscitation measures.

A standard tracheal tube uses a high-volume, low-pressure cuff to avoid pressure necrosis of the tracheal lining. A clinical test for determining correct cuff inflation is to slowly inject air until no air leak is audible while the patient is receiving bag-tube ventilation. This usually occurs with 5 to 8 mL of air if the proper size tracheal tube has been selected. Many clinicians use the tension of the pilot balloon as a guide to cuff inflation. Slight compressibility with gentle external pressure indicates adequate inflation for most clinical situations. For long-term use, measure and maintain cuff pressure at 20 to 25 mm Hg. Capillary blood flow is compromised in the tracheal mucosa when cuff pressure exceeds 30 mm Hg. In emergency situations, simply inflate the balloon with 10 mL of air and adjust it when the patient’s condition has stabilized.
Interest in design of the tip of the tracheal tube has grown as the Seldinger technique is increasingly being applied to intubation. When a tracheal tube is passed over a smaller-caliber introducer (Seldinger technique), regardless of whether it is a tracheal tube introducer or an endoscope, there is a reasonable chance that the tube will get hung up on the laryngeal soft tissue. A tracheal tube that has been designed to overcome this problem has a bevel oriented posteriorly and a flexible tip that decreases the distance between the tube and whatever it is being passed over (Fig. 4.8).

Check the ET tube cuff for leaks by inflating the pilot balloon before attempting intubation. Prepare the tube for placement by passing a malleable stylet down the tube to increase its stiffness and enhance control of the tip of the tube. Do not extend the stylet beyond the eyelet of the tube. Bend the tube and stylet to create a “straight-to-cuff” shape with a 35-degree distal bend. Lubricate the tip and cuff of the tube with viscous lidocaine or a water-soluble gel.

**Optimal Patient Positioning for DL**

The sniffing position, with the patient's head extended on the neck and the neck flexed relative to the torso, has traditionally been considered the best head position for DL. This position aligns the oral, pharyngeal, and laryngeal axes (Fig. 4.9A–C).

---

**Figure 4.8** Comparison of the standard tracheal tube tip with the Parker Flex-Tip Tracheal Tube (Parker Medical, Engelwood, CO). Note: When a standard tube is inserted in the normal fashion, the bevel is oriented vertically and toward the patient's left. The Parker tube tip bevel faces posteriorly and may avoid getting caught on laryngeal structures. The flexible tip of the Parker tube also provides a closer fit on an introducer.

**Figure 4.9** Head positioning for tracheal intubation. A, Neutral position. B, Head elevated. C, “Sniffing” position with a flexed neck and extended head. Note that flexing the neck while extending the head lines up the various axes and allows direct laryngoscopy. D, Morbidly obese patients are best intubated in a ramped position with elevation of the upper part of the back, neck, and head; the ideal position aligns the external auditory canal and the sternum.
Horton and colleagues described the ideal sniffing position for normal patients as neck flexion of 35 degrees and atlanto-occipital extension such that the plane of the face is −15 degrees to the horizontal position. In supine patients, neck flexion is achieved by head elevation. Depending on the size and shape of the patient, the amount of head elevation may differ significantly, and the end point should be horizontal alignment of the external auditory meatus with the sternum. In normal-size adults it is usually possible to achieve the sniffing position with 7 to 10 cm of head elevation. Morbidly obese patients require much more head elevation to achieve the proper sniffing position. In these patients, aligning the external auditory meatus with the sternum requires elevation of the head and neck, as well as the upper part of the back (see Fig. 4.9). This can be accomplished by building a ramp of towels and pillows under the upper torso, head, and neck or by using a Troop Elevation Pillow (Mercury Medical, Clearwater, FL) or similar device. Alternatively, elevating the head to a 25-degree back-up position (keeping the patient supine while placing the bed in 25-degree reverse Trendelenburg) may achieve the same purpose.

Two studies have shown that elevating the head (flexing the neck) beyond the sniffing position often improves visualization of the glottis. Because the amount of head elevation needed for optimal laryngoscopy varies depending on individual patient anatomy, it is important to make laryngoscopy a dynamic procedure. This is best accomplished by putting your right hand behind the patient’s head to lift, flex, and extend the head as needed to bring the glottis into view. Optimal positioning of the head and neck is not possible in trauma patients who require in-line stabilization of the cervical spine. This is one of the aspects that makes trauma airways so challenging and makes other maneuvers, such as external laryngeal manipulation (ELM), even more important in these patients. It should also be noted that some patients, especially those who are obese, are in neck extension when lying supine because of upper dorsal fat deposition. If this is noted, the head can be raised until the head and neck are in neutral position.

**Procedure and Technique of DL**

**Learning DL**

Existing data suggests that novices who learn tracheal intubation skills with video laryngoscopy have higher success rates at DL compared to novices who learn tracheal intubation skills with DL. When utilizing DL alone, it is estimated that at least 50 intubations need to be performed to achieve greater than 90% success. Using a video laryngoscopy system that allows both direct and video laryngoscopy (video systems with a Macintosh blade) may be ideal for learning DL skills.

**Adults**

Place the patient in the supine position with the head at the level of the lower part of the intubator’s sternum (Fig. 4.10, step 2). To maintain the best mechanical advantage, keep your back straight and do not hunch over the patient. Bend only at the knees (Fig. 4.11). Keep the left elbow relatively close to the body and flex it slightly to provide better support. In a severely dyspneic patient who cannot tolerate lying down, perform DL with the patient seated semi-erect and the clinician on a stepstool behind the patient.

Grasp the laryngoscope in the left hand with the back end of the blade pressed into the hypothenar aspect of your hand.

**Infants and Children**

It is helpful to appreciate the anatomic differences between children and adults when intubating pediatric patients (Fig. 4.13 and Table 4.2). Children’s proportionately larger heads naturally place them in the sniffing position, so a towel under the occiput is rarely necessary (Fig. 4.14). The large head of newborns can result in a posterior positioning of the larynx that prevents visualization of the vocal cords. A small towel under the infant’s shoulders should correct this problem. This is not necessary in all children; rather, the goal of any positioning maneuvers should be alignment of the tragus with the anterior shoulder.

The head may also be floppy and may benefit from stabilization by an assistant. The child’s increased tongue-to-oropharynx ratio and shorter neck hinder forward displacement of the tongue and, when coupled with a U-shaped epiglottis, can make visualization of the glottis difficult. DL in infants and young children is generally best performed with a straight blade: Miller size 0 for premature infants, size 1 for normal-sized infants, and size 2 for older children. The infant’s larynx lies
Direct Laryngoscopy

1. Check all equipment, including the light on the laryngoscope and the cuff on the endotracheal tube.
   Ensure that suction and difficult airway devices are within reach.

2. Place patient in the sniffing position, elevate the bed so that the patient’s head is at the level of the lower part of your sternum, and preoxygenate.

3. Hold laryngoscope with your left hand.
   Open patient’s mouth with your right hand and introduce the laryngoscope into the right side of the patient’s mouth.

4. Lift in the direction of the laryngoscope handle.
   Push the tongue to the left side of the mouth, slowly advance the blade, and progressively identify the base of the tongue, the epiglottis, and the posterior cartilages.

5. Place the Macintosh blade in the vallecula, or the Miller blade under the epiglottis (E), and visualize the vocal cords (VC) and arytenoid cartilages (A).
   Do not take your eyes off of the cords once they are identified.

6. Instruct an assistant to retract the right cheek for better visualization. Pass the tube on the right side of the patient’s mouth. Do not allow the tube to obstruct your view of the vocal cords during advancement.

7. Remove the stylet and inflate the pilot balloon.

8. Under direct visualization, pass the tube 3–4 cm beyond the vocal cords.

9. Confirm proper placement with end-tidal CO₂ detection, auscultation, and a chest radiograph.

Figure 4.10 Direct laryngoscopy.
Figure 4.11 Proper positioning of the clinician, patient, and assistant for tracheal intubation. The following points are demonstrated: the difficult airway cart is adjacent to the patient, the suction device is at the head of the bed, the patient is in the “sniffing position” with the occiput elevated and the clinician’s right hand ready for additional adjustment if necessary, the bed is elevated and the clinician is at the appropriate distance from the patient, and the laryngoscope handle is angled at 45 degrees.

Figure 4.12 Common problems encountered when using a laryngoscope. A, The laryngoscope blade is under the middle of the tongue, with the sides of the tongue hanging down and obscuring the glottis. B, The tongue is not pushed far enough to the left and is obscuring the glottis. C, Correct blade position with the tongue elevated and to the left. D, Use of the curved (Macintosh) laryngoscope blade. E, Use of the straight (Miller) blade.
Cricoid Pressure

There is some good evidence that cricoid pressure (Sellick’s maneuver) helps prevent gastric inflation during BMV, though cricoid pressure during BMV reduces tidal volume, increases peak inspiratory pressure, and prevents good air exchange. There are many reports of significant regurgitation and aspiration regardless of the application of cricoid pressure. Cricoid pressure also decreases successful insertion of and intubation through LMAs. Despite the lack of evidence, many experts believe that Sellick’s maneuver is critical during RSI. Because aspiration has dire consequences and because cricoid pressure has traditionally been considered integral to patient safety during emergency airway management, it is reasonable to apply cricoid pressure as long as it does not interfere with ventilation and intubation. There is significant evidence that it can interfere with ventilation and intubation, so it is best to apply cricoid pressure on a case-by-case basis with a full understanding of the benefits and drawbacks of cricoid pressure. If cricoid pressure is utilized, it should be released immediately if there is any difficulty either intubating or ventilating a patient in an emergency setting. Routine use of cricoid pressure during BMV of patients in cardiac arrest is not currently recommended in the AHA Guidelines for Cardiopulmonary Resuscitation and Emergency Cardiovascular Care. It is reasonable to release or relax cricoid pressure during insertion of an LMA, during intubation with an ILMA, or if ventilation with the LMA is difficult. Some authors believe that improper technique is to blame for the many reported failures of Sellick’s maneuver. The proper technique for applying Sellick’s maneuver is to place the thumb and middle finger on either side of the cricoid cartilage and the index finger in the center anteriorly. As a reference, approximately 40 N of digital force on the bridge of the nose will usually cause pain. ELM, Bimanual Laryngoscopy, and BURP

ELM is the application of pressure on the thyroid cartilage in an attempt to improve the view of the larynx during laryngoscopy. The differences between cricoid pressure, ELM, and backward, upward, rightward pressure (BURP) are often misunderstood. Cricoid pressure is the application of pressure at the anterior cricoid ring to displace it posteriorly to attempt to occlude the esophagus, with the intent of preventing regurgitation and aspiration; cricoid pressure is not intended to improve visualization during laryngoscopy. ELM is the application of pressure on the thyroid cartilage during laryngoscopy to help optimize visualization of the glottis. BURP is often the best combination of forces that need to be applied to the thyroid cartilage during ELM. Bimanual laryngoscopy refers to use of the right hand to perform ELM.

Cricoid Pressure

There is some good evidence that cricoid pressure (Sellick’s maneuver) helps prevent gastric inflation during BMV, though cricoid pressure during BMV reduces tidal volume, increases peak inspiratory pressure, and prevents good air exchange. There are many reports of significant regurgitation and aspiration regardless of the application of cricoid pressure. Cricoid pressure also decreases successful insertion of and intubation through LMAs. Despite the lack of evidence, many experts believe that Sellick’s maneuver is critical during RSI. Because aspiration has dire consequences and because cricoid pressure has traditionally been considered integral to patient safety during emergency airway management, it is reasonable to apply cricoid pressure as long as it does not interfere with ventilation and intubation. There is significant evidence that it can interfere with ventilation and intubation, so it is best to apply cricoid pressure on a case-by-case basis with a full understanding of the benefits and drawbacks of cricoid pressure. If cricoid pressure is utilized, it should be released immediately if there is any difficulty either intubating or ventilating a patient in an emergency setting. Routine use of cricoid pressure during BMV of patients in cardiac arrest is not currently recommended in the AHA Guidelines for Cardiopulmonary Resuscitation and Emergency Cardiovascular Care. It is reasonable to release or relax cricoid pressure during insertion of an LMA, during intubation with an ILMA, or if ventilation with the LMA is difficult. Some authors believe that improper technique is to blame for the many reported failures of Sellick’s maneuver. The proper technique for applying Sellick’s maneuver is to place the thumb and middle finger on either side of the cricoid cartilage and the index finger in the center anteriorly. Apply 30 N (6.7 lb.) of force to the cricoid cartilage in the posterior direction. As a reference, approximately 40 N of digital force on the bridge of the nose will usually cause pain. ELM, Bimanual Laryngoscopy, and BURP

ELM is the application of pressure on the thyroid cartilage in an attempt to improve the view of the larynx during laryngoscopy. Multiple studies have shown that ELM performed by the laryngoscopist (bimanual laryngoscopy) is superior to having an assistant apply anterior neck pressure. Bimanual laryngoscopy is best because the direction and amount of force that will optimize laryngeal exposure is variable. BURP is sometimes optimal, and it often worsens the laryngoscopic view, so it is best to move the larynx in a variety of directions.
### TABLE 4.2 Comparison of the Airway in Adults and Children

<table>
<thead>
<tr>
<th>COMPARISON</th>
<th>CHILD</th>
<th>ADULT</th>
<th>CLINICAL CONSEQUENCES OR ADJUSTMENTS FOR CHILDREN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Head</td>
<td>Proportionately larger (up to approximately age 10 yr)</td>
<td>Proportionately smaller</td>
<td>A child is naturally in the sniffing position when supine. Do not place a towel under the occiput; a child may benefit from elevation of the shoulders. The large head may be “floppy” and require the assistant to hold the head still during intubation.</td>
</tr>
<tr>
<td>Teeth</td>
<td>Easily knocked out</td>
<td>Stable unless decay or trauma is a factor</td>
<td>Teeth may be knocked out and aspirated or forced into trachea.</td>
</tr>
<tr>
<td>Tonsils or adenoids</td>
<td>Large and friable</td>
<td>Generally not a problem</td>
<td>Nasotracheal intubation in a child may cause excessive bleeding and is not recommended. Adenoid or tonsil tissue may plug the endotracheal tube or cause airway obstruction from aspiration.</td>
</tr>
<tr>
<td>Tongue</td>
<td>Relatively larger</td>
<td>Relatively smaller</td>
<td>The tongue is difficult to displace anteriorly in a child. Consider using a straight blade.</td>
</tr>
<tr>
<td>Larynx</td>
<td>Opposite C2-C3</td>
<td>Opposite C4-C6</td>
<td>A more superiorly located larynx or an “anterior” larynx is more difficult to visualize. Consider using a straight blade.</td>
</tr>
<tr>
<td>Epiglottis</td>
<td>U shaped, shorter, stiffer</td>
<td>Flatter, more flexible</td>
<td>The epiglottis is more difficult to manipulate in a child; it may fold down and obstruct the view with use of a curved blade. Consider using a straight blade.</td>
</tr>
<tr>
<td>Vocal cords</td>
<td>Concave upward; anterior attachment of the cords lower than posterior, thereby creating a slant</td>
<td>Horizontal</td>
<td>A concave shape does not affect intubation, but it may affect ventilation. For partial airway obstruction or to break laryngospasm, consider positive pressure ventilation with a jaw-lift maneuver to open the arytenoids. The anterior superior slant of the vocal cords may cause the endotracheal tube to hang up on the anterior commissure as it passes into the larynx. Rotate the tube 90 degrees counterclockwise. Overextension of the neck may cause partial airway obstruction as a result of airway collapse.</td>
</tr>
<tr>
<td>Length of the trachea</td>
<td>Relatively shorter</td>
<td>Relatively longer</td>
<td>A short trachea increases the likelihood of main stem bronchus intubation. Follow the formula for correct depth of placement (cm depth = 0.5 × age [yr] + 12) measured from the corner of the mouth. The double black line on the endotracheal tube should pass just beyond the cords.</td>
</tr>
<tr>
<td>Airway diameter</td>
<td>Relatively smaller; smallest diameter at the cricoid ring</td>
<td>Relatively larger; smallest diameter between the vocal cords</td>
<td>Laryngoscope-induced trauma, edema, and foreign material will significantly alter the diameter of the airway. Be gentle. Extremes of flexion or extension may kink the airway. If trouble with bag-valve-mask ventilation occurs, reassess the degree of head flexion or extension. Cricoid pressure may cause complete airway obstruction. The endotracheal tube may pass through the cords but be too large to pass through the cricoid ring. If unable to pass into the trachea, use the next smaller tube.</td>
</tr>
<tr>
<td>Residual lung capacity</td>
<td>Relatively smaller</td>
<td>Relatively larger</td>
<td>A child becomes hypoxic more quickly than an adult does. Closely monitor O2 saturation and avoid prolonged periods without ventilation.</td>
</tr>
</tbody>
</table>

To determine the optimal ELM. The best way to quickly apply a variety of different forces to the larynx to determine the optimal ELM is by manipulation of the larynx with the laryngoscopist’s right hand (Fig. 4.15).\(^{113}\)

Operator-directed posterior displacement of the larynx during laryngoscopy was described by Brunnings in 1912.\(^{113}\) After Sellick described the use of cricoid pressure to avoid regurgitation in 1961, it became common to have an assistant apply anterior neck pressure during laryngoscopy.\(^{7}\) In 1993, Knill reported that having an assistant apply BURP to the cricoid or thyroid cartilage improved visualization of the glottis during two cases of difficult laryngoscopy.\(^{119}\) In 1993, Takahata and coworkers performed a prospective study of 630 intubations and found that BURP produced better laryngeal exposure than...
Figure 4.15 External laryngeal manipulation (ELM) (or bimanual laryngoscopy). A, ELM step 1. The laryngoscopist optimizes view of the larynx by reaching around to the patient's neck with the right hand and manipulating the thyroid cartilage while performing laryngoscopy. B, ELM step 2. The assistant's hand replaces the laryngoscopist's hand on the anterior aspect of the neck and maintains the position of the larynx while the laryngoscopist places the tracheal tube.

Figure 4.16 The epiglottis is elevated and visualization of the vocal cords is improved by applying pressure on the hyoepiglottic ligament and external laryngeal manipulation. (Courtesy Richard M. Levitan, MD, Airway Cam Technologies, Inc., Wayne, PA. Used with permission.)

Figure 4.17 Note the assistant manipulating the anterior part of the neck and retracting the cheek for better visualization.

just backward pressure in patients with difficult laryngoscopy.\textsuperscript{140} A 2005 prospective crossover trial by Snider and colleagues found no benefit with routine application of the BURP maneuver.\textsuperscript{141}

Studies by Benumof, Levitan, and colleagues have demonstrated that it is best to apply pressure on the thyroid cartilage (not the cricoid cartilage) and suggested that ELM should be applied by the laryngoscopist's right hand, not by an assistant.\textsuperscript{115,137,142} They also found that the direction of force required for optimal ELM was not always upward and rightward and that the amount of backward pressure was variable.\textsuperscript{115} In a 1996 study of 181 patients, Benumof and Cooper found that external manipulation was optimal when applied to the thyroid cartilage in 88\% of patients and to the cricoid cartilage in only 11\% of patients.\textsuperscript{115} In a 2006 cadaveric study of 1530 laryngoscopies by 104 laryngoscopists, Levitan and associates found that bimanual laryngoscopy (ELM performed by the laryngoscopist) consistently improved laryngeal exposure by novice intubators.\textsuperscript{142}

Bimanual laryngoscopy with ELM should be performed whenever the laryngeal view is not optimal after good laryngoscopic technique. To perform this procedure, the intubator applies posterior pressure on the thyroid cartilage. The force vector (right or left, upwards or downwards, and amount of posterior pressure) will vary patient to patient, and the intubator should find the force vector that provides the best laryngeal view. Once this is established, an assistant applies the same force vector to the thyroid cartilage as the intubator removes pressure to free the hand in order to pass the tracheal tube. The assistant holds pressure while the intubator completes tracheal intubation.

**Passing the Tube**

Once the vocal cords have been visualized, the final step is to pass the tube through the vocal cords and into the trachea
under direct vision. It is best to use a malleable stylet for all emergency intubations. The best stylet shape is straight with a 35-degree hockey-stick bend at the proximal cuff (“straight-to-cuff”). In a 2006 study, Levitan and colleagues showed that stylet bend angles greater than 35 degrees made ET tube passage more difficult. Hold the tube in your right hand and introduce it from the right side of the patient’s mouth. Lateral retraction of the cheek by an assistant may greatly aid overall visualization (see Fig. 4.17). Advance the tube toward the patient’s larynx below the line of sight with the bend facing upward. When advanced in this manner, the tube does not obstruct the view of the larynx until the last possible moment before the tube enters the larynx. If the patient is not chemically paralyzed, pass the tube during inspiration, when the vocal cords are maximally open. It enters the trachea when the cuff disappears through the vocal cords. Advance the tube 3 to 4 cm beyond this point. It is not enough to see the tube approach the cords; watch the tube pass through the vocal cords to ensure tracheal placement. Directly observing the tube pass through the cords is the best way to immediately confirm correct placement. If part of the glottis is visualized and it is difficult to pass the tube, consider using a bougie (tracheal tube introducer).

**Tracheal Tube Introducer (Bougie)**

If DL does not bring the vocal cords fully into view, a tracheal tube introducer may be used to facilitate intubation. This adjunct is a long, thin, semirigid introducer that, with the aid of a laryngoscope, is passed through the laryngeal inlet and over which an ET tube is advanced through the cords and into the trachea. The technique, originally described more than 60 years ago by Macintosh, was recommended for patients in whom visualizing the vocal cords was difficult. It has also been shown to be effective when the laryngeal inlet cannot be visualized at all. It is the most common airway adjunct used in British EDs for complicated intubations. Its efficacy has been demonstrated prospectively during difficult intubations in the operating room, as a pivotal component of a difficult airway algorithm in the operating room, and when compared with conventional laryngoscopy in the ED. A variety of tracheal tube introducers are available today (Fig. 4.18). The original adjunct was called the gum elastic bougie, or simply “the bougie,” and is currently available in a reusable form for both adult and pediatric patients (Eschmann Tracheal Tube Introducer, Portex Sims, Kent, UK). The adult size comes in two forms: a 60-cm (15-Fr) version with a short, 40-degree hockey-stick curve at the end, and a straight one that is 70 cm. The adult version can accommodate a 5.5-mm ET tube. The pediatric version is 70 cm (10-Fr) and straight and can accommodate a 4.0-mm tube. A polyethylene introducer designed for single use is also available and comes only in the 60-cm version (Flextrach ET Tube Guide, Greenfield Medical Sourcing, Austin, TX). A variation of this concept is the FROVA Introducer (Cook Critical Care, Bloomington, IN), a plastic introducer with a similar profile to the others except that it has a hollow lumen through which the patient can be ventilated when an accompanying adapter is attached.

Consider using a tracheal tube introducer when a difficult airway is anticipated; it can also be helpful in all intubations when visualization of the laryngeal inlet is limited. A trauma patient with cervical spine precautions is a typical example. The presence of blood and vomitus rarely prevents placement of the bougie into the trachea. Its safety record is impressive despite decades of use, and reports of complications are rare.

Shaping the introducer may not be necessary in many cases, but with difficult laryngeal views, create a 60-degree bend in the distal introducer (see Fig. 4.18D). Ideally, tracheal tube introducer-assisted intubation is a two-person procedure (Fig. 4.19). As laryngoscopy begins, the assistant has both a styletted ET tube and bougie prepared and available. The intubator performs laryngoscopy in the normal fashion to obtain the best possible view of the larynx. If the cords are in full view, proceed with intubation using a styletted ET tube. If the view is suboptimal, an assistant can pass the tracheal tube introducer to the operator for placement anterior to the arytenoids and into the larynx. If only the epiglottis is visible, place the introducer, with a 60-degree distal bend, just under the epiglottis and direct it anteriorly. With the laryngoscope still in place and the introducer stabilized by the operator, the assistant slides the ET tube over the introducer. Pass the tube through the larynx. Just before entering the larynx, rotate the tube 90 degrees counterclockwise to avoid having the tip of the ET tube get caught on the laryngeal structures (Fig. 4.20). Withdraw the laryngoscope and confirm proper tube placement. While securing the ET tube, ask the assistant to remove the introducer.

There are a number of findings that confirm successful introducer placement. If any portion of the arytenoids is visible and the introducer was seen to pass anterior to them without resistance, the introducer is in the airway. Unlike seeing an ET tube “go through the cords” when in fact the laryngeal inlet may have been momentarily obscured by the tube or balloon, the smaller-caliber introducer does not obscure the view of the glottis and thus avoids this potential pitfall. In addition to better visual confirmation, successful passage is indicated, up to 90% of the time, by feeling clicks produced by the angled tip of the introducer as it strikes against the tracheal rings. An assistant will also usually feel confirmatory movement in the airway if the anterior aspect of the neck is palpated. If there is still any question whether the introducer is in the airway, gently advance it at least 40 cm, at which point resistance should be felt as the introducer passes the carina and stops inside a main bronchus. If the bougie does not stop when advanced approximately 40 cm, the introducer is most likely in the esophagus. Withdraw it and reattempt placement. Several technical points should be emphasized. The first is that it is important to create a curve in the distal portion of

![Figure 4.18](image-url) Several types of tracheal tube introducers. The classic introducer, the gum elastic bougie, A, is reusable and comes in curved- and straight-tipped adult forms and a straight pediatric form. The straight bougies, B, are 70 cm long and the curved-tipped bougies are 60 cm long. The blue introducer, C, (Flextrach ET Tube Guide, Greenfield Med., Austin, TX) is polyethylene and designed for single use and comes only in a curved-tipped adult form (60 cm). D, Create a 60-degree bend in the distal portion of the introducer if the laryngeal inlet cannot be seen on laryngoscopy.
counterclockwise, and advance it again (see Fig. 4.20). Fourth, although there may be some benefit to lubricating the distal end of the introducer, in emergency intubations, lubricating the full length of the introducer makes it slippery and hard to handle without conferring any obvious advantage. Lubricating the ET tube, conversely, remains critical for smooth passage through the vocal cords. Fifth, a common error when first using the tracheal tube introducer is to remove the laryngoscope before passing the tube over the introducer. This often results in difficulty placing the tube because it is displaced posteriorly by the weight of the pharyngeal soft tissues and gets hung up on the laryngeal structures. Reinsert the laryngoscope. Pull the tube back 2 cm to disengage the soft tissue. Rotate the tube 90 degrees counterclockwise and then readvance
it. Sixth, in instances in which it is difficult to get the introducer sufficiently anterior to access the laryngeal inlet, make sure that the introducer lines up with the operator's line of vision. If the introducer enters the mouth at a significant angle above this line, most often when the clinician is too close to the patient, it may be deflected posteriorly by the lip or intraoral structures and escape the attention of the operator. This creates the impression that the introducer is “too floppy.”

In the prehospital setting, where assistance might not be available, the laryngoscope should be removed to mount the ET tube onto the introducer. Once the tube is on the introducer, reinsert the laryngoscope and advance the introducer through the glottic opening. Advance the ET tube while rotating it 90 degrees counterclockwise to ensure successful passage into the trachea. Mounting the tube onto the introducer to insert them as a unit is not advised because it is often difficult to direct the introducer into the laryngeal inlet as it moves within the ET tube.

**Laryngospasm**

If the patient is not paralyzed, laryngospasm, or persistent contraction of the adductor muscles of the vocal cords, may prevent passage of the tube. Pretreatment with topical lidocaine may decrease the likelihood of laryngospasm, though this is not routinely performed. After laryngospasm is noted, one option is to spray lidocaine (2% or 4%) directly onto the vocal cords. An infrequent but effective means of achieving tracheal anesthesia is transtracheal puncture and injection of 3 to 4 mL of lidocaine through the cricothyroid membrane. Laryngospasm is usually brief and often followed by a gasp. Be ready to pass the tube at this moment. Occasionally, the spasm is prolonged and needs to be disrupted with sustained anterior traction applied at the angles of the mandible, as in the jaw-thrust maneuver. Do not force the tube at any time because it could cause permanent damage to the vocal cords. Consider using a smaller tube. Prolonged, intense spasm may ultimately require muscle relaxation with a paralyzing drug (see Chapter 5). Pediatric patients are far more prone to laryngospasm than adults. In a child, if vocal cord spasm prevents passage of the tube, a chest-thrust maneuver may momentarily open the passage and permit intubation.

**Positioning and Securing the Tube**

Secure the ET tube in a position that minimizes both the chance of inadvertent, main stem endobronchial intubation and the risk for extubation. The tip should lie in the midtrachea with room to accommodate neck movement. Because tube movement with both neck flexion and extension averages 2 cm, the desired range of tip location is between 3 and 7 cm above the carina.

The average tracheal length is between 10 and 13 cm. On a radiograph, the tip of the tube should ideally be 5 ± 2 cm above the carina when the head and neck are in a neutral position. On a portable radiograph, the adult carina overlies the fifth, sixth, or seventh thoracic vertebral body. If the carina is not visible, it can be assumed that the tip of the tube is properly positioned if it is aligned with the third or fourth thoracic vertebra. In children, the carina is more cephalad than in adults, and it is consistently situated between T3 and T5. In children, T1 is the reference point for the tip of the ET tube.

Estimate the proper depth of tube placement before radiographic confirmation by using the following formulas, in which length represents the distance from the tip of the tube to the upper incisors in children and from the upper incisors or the corner of the mouth in adults:

- **Children:** Tracheal tube depth (cm) = \[\text{age (yr)} / 2\] + 12
- **Adults:** Tracheal tube depth (cm) = 21 cm
- **Med:** Tracheal tube depth (cm) = 23 cm

In adults, this method has been shown to be more reliable than auscultation in determining the correct depth of placement. One can anticipate that tall male patients will often require deeper placement, to 24 or 25 cm, and that short women will often require a shallower placement of 19 or 20 cm.

Inflate the cuff to the point of minimal air leak with positive pressure ventilation. In an emergency intubation, inflate with 10 mL of air and adjust the inflation volume after the patient is stabilized.

After placement of the tracheal tube, auscultate both lungs under positive pressure ventilation. Take care to auscultate posterolaterally because auscultation anteriorly can reveal sounds that mimic breath sounds and arise from the stomach. With the tube in place and the cuff inflated, secure the tube in place. Attach commercial ET tube holders, adhesive tape, or umbilical (nonadhesive cloth) tape securely to the tube and around the patient's head (Figs. 4.21 and 4.22). Position the tube at the corner of the mouth, where the tongue is less likely to expel it. This position is also more comfortable for the patient and allows suctioning. A bite block or oral airway to prevent crimping of the ET tube or damage from biting is commonly incorporated into the system used to secure the tube.

Unintentional extubation can have disastrous consequences, particularly if the patient was difficult to intubate initially. Secure the ET tube immediately after correct placement has been confirmed. Orotracheal intubation is associated with a higher rate of unplanned extubation than NT intubation. During transport, moving the intubated patient, or obtaining radiographs, designate one person to tend to the ET tube to avoid unplanned extubation. Inadequate sedation is another risk factor for unplanned extubation. If long-acting paralytics have not been administered, consider sedation or physical restraints to prevent self-extubation by an agitated or confused patient.

**Confirmation of Tracheal Tube Placement**

**Clinical Assessment**

Confirm tracheal placement clinically by seeing the tube pass through the vocal cords (Table 4.3). If any question remains, apply posterior pressure on the ET tube while the laryngoscope is still in place and expose the tube by altering the angle as it passes between the cords. Absent or diminished breath sounds, any sound or vocalization, increased abdominal size, and gurgling sounds during ventilation are clinical signs of esophageal placement. If the patient can moan or groan, the tube is not in the trachea! Critically, esophageal placement is not always obvious. One may hear “normal” breath sounds if only the midline of the thorax is auscultated. The presence of condensation of the ET tube as a means of confirming tracheal placement may also be misleading. Blinded observers noted condensation of the ET tube during ventilation in 23 of 27 esophageal intubations in an animal model. One way to clinically assess tracheal placement after several ventilations...
if after noting asymmetric breath sounds the tube is withdrawn a defined distance beyond the point at which equal breath sounds are first heard: 2 cm in children younger than 5 years and 3 cm in older children.

**Esophageal Detector Device**

An aspiration technique used to determine ET tube location was first described by Wee in 1988.164 The technique takes advantage of the difference in tracheal and esophageal resistance to collapse during aspiration to locate the tip of the tracheal tube. After intubation, attach a large syringe (Positube esophageal...
Figure 4.22 A, A commercial disposable tube holder is ideal and preferred to secure an endotracheal (ET) tube without the use of messy tape. B, A plastic disposable ET holder firmly secures the ET tube with a small clamp. C, When positioning a patient for transfer to another bed or for a chest radiograph, ensure the integrity of the ET tube by placing the right hand firmly against the right side of the face while holding the tube securely with the same hand. The other hand immobilizes the neck. (B, Courtesy Laerdal Medical, Wappingers Falls, NY.)

### TABLE 4.3 Assessing Proper Tube Placement

<table>
<thead>
<tr>
<th>TEST</th>
<th>INTERPRETATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Observe the tube pass through the vocal cords</td>
<td>Accurate way to ensure placement; if in doubt, look again after intubation</td>
</tr>
<tr>
<td>End-tidal CO₂ measurements</td>
<td>Reliable if a good persistent waveform is present; can be misleading with nasotracheal intubation if the tip is curled supraglottically: it will give a positive CO₂ reading</td>
</tr>
<tr>
<td>Auscultation of breath sounds over the chest</td>
<td>May be misleading, especially if only the midline is examined; listen in both axillae</td>
</tr>
<tr>
<td>Auscultation over the stomach</td>
<td>Gurgling indicates esophageal placement</td>
</tr>
<tr>
<td>Condensation (fog) forms inside the tube with each breath</td>
<td>Not reliable to confirm tracheal placement</td>
</tr>
<tr>
<td>Observe the chest rise with positive pressure and fall with release</td>
<td>Generally reliable if good chest rise is present; may be absent in patients with a small tidal volume or severe bronchospasm</td>
</tr>
<tr>
<td>Feel air exiting from the end of the tube after inflation</td>
<td>Reliable</td>
</tr>
<tr>
<td>Air remains in the lung after the end of the tube is occluded and exits when the occlusion is removed</td>
<td>Reliable, but one may “ventilate” a closed area of the esophagus</td>
</tr>
<tr>
<td>Ask the patient to speak; listen for moaning or other sounds</td>
<td>If the tube is in proper place, no sound is possible</td>
</tr>
<tr>
<td>Chest radiograph</td>
<td>Cannot always differentiate between tracheal and esophageal placement. If known to be in the trachea by other measures, radiography can assess for proper depth of tube insertion.</td>
</tr>
<tr>
<td>Aspiration technique</td>
<td>Tracheal location is confirmed (assuming patient ET tube) if 30–40 mL of air is aspirated without resistance; probable esophageal location if unable to aspirate the syringe easily or delayed bulb refill occurs; can be misleading with nasotracheal intubation if the tube has curled supraglottically</td>
</tr>
<tr>
<td>Fiberoptic bronchoscope</td>
<td>Reliable if tracheal rings are seen down the endotracheal tube</td>
</tr>
<tr>
<td>Lighted stylet down the endotracheal tube</td>
<td>Reliable if transillumination seen in the low midline portion of the neck</td>
</tr>
<tr>
<td>Ultrasound detection of tracheal tube location</td>
<td>Appears to be reliable but not extensively studied</td>
</tr>
</tbody>
</table>
detector, Flotec, Indianapolis, IN) to the end of the ET tube and withdraw the plunger of the syringe. If the tube is placed in the trachea correctly, the plunger will pull back without resistance as air is aspirated from the lungs. If the tracheal tube is in the esophagus, resistance is felt when the plunger is withdrawn because the pliable walls of the esophagus collapse under the negative pressure and occlude the end of the tube. Another device that uses the same principle as syringe aspiration is the self-inflating bulb (e.g., Ellick device).

In the initial study conducted in an operating room, tube placement was identified correctly in 99 of 100 cases (51 esophageal, 48 tracheal). The result was considered equivocal in the remaining case. That tube was removed and found to be nearly totally occluded with purulent secretions. Slight resistance was noted in one patient with right main stem intubation; the resistance decreased when the tube was pulled back. Before use, always check the esophageal detector device for air leaks. If any connections are loose, the leak may allow the syringe to be withdrawn easily, thereby mimicking tracheal location of the tube.

When using the aspiration technique, apply constant, slow aspiration to avoid occlusion of the tube from tracheal mucosa drawn up under the high negative pressure. If the tracheal tube is placed correctly, 30 to 40 mL of air can be aspirated without resistance. If air was initially aspirated and some resistance is then encountered, the tracheal tube should be pulled back between 0.5 and 1.0 cm and rotated 45 degrees. This takes the tube out of the bronchus if it has been placed too deeply and changes the orientation of the bevel if the tube has been temporarily occluded with tracheal mucosa. Air is easily aspirated if the tube was in the trachea, but repositioning it will make no difference if the tube was in the esophagus. The syringe aspiration technique can be used before or after ventilation of the patient. Inflation of the tube cuff has no effect on the reliability of the test. This device is reliable, rapid, inexpensive, and easy to use.

A squeeze-bulb aspirator is an alternative to the syringe technique. Attach the bulb to the ET tube and squeeze; if the tube is in the esophagus, it is accompanied by a flatus-like sound followed by absent or markedly delayed refilling. Insufflation of a tube in the trachea is silent with instantaneous refill. An early study with the Ellick evacuator bulb device reported that 82% of esophageal intubations were identified. A later study using a slightly different bulb device (Respirronics, Murrysville, PA) found that all 45 esophageal intubations were detected. The device is cheap, easy to use, and operated single-handedly in less than 5 seconds. The bulb should not be used in freezing temperatures because of loss of elasticity. Confusion may occur if the esophageal tube is tested more than once because subsequent inflations may be silent. With repeated assessments, false-positive refilling of the bulb may occur as a result of instillation of air during the first attempt. This observation has led to a recommendation that the bulb be compressed before it is attached to the ET tube. Delayed, though complete refilling of the bulb may occur with bronchial tube placement or placement in the more pliable pediatric airway. The bulb suction modification of the aspiration technique has not been studied as thoroughly as the syringe technique.

A significant number of false positives occur with esophageal detection devices (the tube is correctly placed in the trachea, but the device suggests that it is in the esophagus). These patients are almost uniformly obese. Endoscopic evaluation found that the tracheal wall was invaginated into the ET tube because of the negative pressure. In such circumstances, if the intubation was felt to be successful, visually re-confirm that the ET tube is through the cords before removing the ET tube. Alternatively, if the patient has a perfusing rhythm and an expired CO₂ device is available, it should be used. To date, there has been one reported case of unrecognized esophageal intubation undetected by the syringe aspiration technique. In this case there was marked gastric distention from forceful BMV. The esophageal detection device is not reliable in confirming tracheal tube depth and position after intubation because easy aspiration of air will occur if the tip of the tube is located supraglottically (expired CO₂ will also be misleading).

**End-Tidal CO₂**

End-Tidal CO₂ detection is probably the best technique, apart from visualizing the tube pass through the cords, to confirm tracheal placement of the ET tube. A high level of CO₂ in exhaled air is the physiologic basis for capnography and the principle on which end-tidal CO₂ pressure (PETCO₂) detectors were developed (see Chapter 2). Continuous waveform capnography is recommended by the AHA guidelines as the most reliable method of confirming and monitoring correct placement of an ET tube (class I, level of evidence [LOE]). This recommendation is based on multiple studies showing 100% accuracy of waveform capnography for detecting correct ET tube placement. Continuous waveform capnography is accurate even in cardiac arrest. Patients with prolonged cardiac arrest will still have a typical square waveform but a low PETCO₂ value.

When waveform capnography is not available, emergency providers may have to rely on colorimetric CO₂ indicators, which correspond to CO₂ levels flowing through the device when placed on the tracheal tube adapter (see Fig. 4.10, step 10). The typical device displays opposite colors (e.g., yellow and purple) to indicate low levels of CO₂ in esophageal gas versus the high levels of CO₂ exhaled from the respiratory tree. Handheld quantitative or semiquantitative electronic CO₂ monitors are also available. Eventually, all prehospital defibril-lators will have advanced monitoring capability that includes capnography.

A multicenter study of a colorimetric device demonstrated an overall sensitivity of 80% and a specificity of 96%. In patients with spontaneous circulation and an inflated tracheal tube cuff, the sensitivity and specificity were 100%. The poor sensitivity (69%) noted in patients in cardiac arrest was due to the fact that low exhaled CO₂ levels were seen with both very-low-flow states and esophageal intubation. The device must be used with caution in cardiac arrest victims. CO₂ levels return to normal after return of spontaneous circulation. Colorimetric changes may be difficult to discern in situations with reduced lighting, and secretions can interfere with the change in color. Regardless of the monitoring device, patients should be ventilated for a minimum of six breaths before taking a reading. Recent ingestion of carbonated beverages can result in spuriously high CO₂ levels with esophageal intubation. Colorimetric changes do not exclude supraglottic positioning of the tip of the ET tube. Adequate ventilation and oxygenation may be achieved in the supraglottic position, but there is still a risk for aspiration in the absence of a protected airway and the potential for further tube dislodgement. Glottic positioning may be difficult to detect clinically. The only signs may be
persistent cuff leak or diminished chest rise with ventilation. Radiographic evidence or direct visualization confirms the diagnosis.\textsuperscript{172}

**Ultrasound Detection of Tracheal Tube Location**

Some data suggest that transtracheal ultrasound may play a future role in confirming tracheal location of the ET tube location after intubation. All studies are small, and sensitivity ranges from 96% to 100% for confirming tracheal placement, whereas specificity (detecting esophageal intubation) ranges from 88% to 100%.\textsuperscript{175–178} If the ET tube is the in the trachea, acoustic shadowing is seen posterior to the anterior tracheal rings only. If the ET is in the esophagus, the esophagus is opened by the ET tube and shadowing is seen posterior to the anterior esophageal wall (as well as the trachea). This method relies on the esophagus being located in the paratracheal position; if the esophagus is directly posterior to the trachea, then detecting esophageal intubation is very difficult.

Sonographic sliding signs can also be used immediately after tracheal intubation is confirmed by waveform capnography to evaluate for main stem intubation prior to obtaining a chest radiograph.\textsuperscript{179,180} Assuming there is no other underlying lung pathology, absence of sliding on the left after intubation indicates probable right main stem intubation. The ET tube can be withdrawn 2 cm, and sliding signs reassessed.

**Comparison of Detector Devices**

In the setting of spontaneous circulation, both syringe aspiration and PETCO2 detection are highly reliable means of excluding esophageal intubation. An animal study comparing these techniques with clinical assessment and measuring the speed and accuracy of determining tube placement demonstrated that both the syringe esophageal detector device and PETCO2 detection were highly accurate, approaching 100%.\textsuperscript{181} The esophageal detector device was more rapid, with determination in 13.8 seconds versus 31.5 seconds for PETCO2 detection. The detector device remained accurate when air was insufflated into the esophagus for 1 minute, thus simulating unrecognized esophageal placement. Clinical assessment alone yielded an alarming 30% rate of failure to identify esophageal intubation. In the setting of cardiac arrest, the aspiration method is more reliable than colorimetric CO2 detection, although waveform capnography remains reliable even in low flow states.

An unequivocal method for determining tracheal tube location uses the endoscope. Passage of the scope through the tube with visualization of the tracheal rings confirms ET placement and position within the trachea. Placement of a lighted stylet below the tracheal tube and successful transtracheal illumination can also be used to determine correct ET placement.\textsuperscript{182}

At present, there is no perfect device for confirmation of ET tube placement in all situations. Be aware of the limitations of each device; it is ideal to use multiple means of confirmation to ensure tracheal placement of the ET tube.

**Complications of Intubation**

Failure to achieve adequate ventilation and oxygenation is the most serious complication of tracheal intubation. The potential for hypoxia exists just before intubation as more conservative oxygenation methods are attempted and then fail, during difficult intubation when ventilation is halted for an attempt at intubation, and after intubation when esophageal intubation goes undetected. Because irreversible cerebral anoxia occurs within minutes, conservative airway management maneuvers should be limited to 2 to 3 minutes; failure to achieve adequate oxygenation should lead to a quick decision to intubate.

As a guide, limit intubation attempts to the amount of time that a single deep breath can be held by the patient. This is especially important in a child because the functional residual capacity of a child’s lungs is less than that of an adult. Historically, the maximum recommended duration of an intubation attempt in an apneic patient has been 30 seconds, followed by a period of BMV before intubation is attempted again. Longer attempts at intubation are permissible when guided by accurate data from an oxygen saturation monitor because oxygen saturation may remain in the normal range for much longer in patients who have been preoxygenated. As a general rule, intubation attempts may continue if oxygen saturation is above 90% and should be interrupted for BMV when oxygen saturation drops below 90%.

Assessment of tube location is the top priority immediately after placement. The best assurance of tracheal placement is to see the tube pass through the vocal cords. Techniques to assess tube placement were discussed earlier. If esophageal intubation is discovered, removal of the tube may be followed by emesis. Apply cricoid pressure during tube removal and maintain it until the intubation is successful. Keep a large-bore suction tip catheter readily available should vomiting occur. Alternatively, leave the first tube in the esophagus to serve as a temporary gastric-venting device and as a guide to intubation until tracheal intubation is achieved.

Though seldom associated with serious complications, unrecognized placement of the tip of the ET tube in the right main stem bronchus may cause hypoxia, atelectasis, pneumothorax, and unilateral pulmonary edema.\textsuperscript{183} Obtain a chest radiograph soon after intubation to confirm tube positioning. Endobronchial intubation was clinically unrecognized without a chest film in 7% of prehospital intubations in one study.\textsuperscript{184} Persistent asymmetric breath sounds after correct tube positioning suggests unilateral pulmonary pathology (e.g., main stem bronchus obstruction, pneumothorax, hemothorax).

Prolonged efforts to intubate can also cause cardiac decompensation. Pharyngeal stimulation can produce profound bradycardia or asystole, thereby confirming the need for an assistant to monitor cardiac rhythm throughout the intubation. Keep atropine available to reverse the vagal-induced bradycardia that may occur secondary to suctioning or laryngoscopy. Prolonged pharyngeal stimulation may also result in laryngospasm, bronchospasm, and apnea. Hypotension requiring intervention in the peri-intubation period occurs in approximately 1.5% of intubations.\textsuperscript{64} If a patient is profoundly hypovolemic and does not need immediate intubation, it may be best to resuscitate with intravenous fluids prior to intubation to avoid worsening hypotension.

Generally, dentures are removed for intubation but kept in place for BMV. Check for loose or missing teeth before and after orotracheal intubation. Look for any avulsed teeth not found in the oral cavity on the postintubation chest film. Dental injury occurs in approximately 0.5% of intubations.\textsuperscript{64} Laceration of the mucosa of the lips, especially the lower lip, may occur. Tracheal or bronchial injuries are rare, but serious and usually occur in infants and the elderly as a result of their decreased tissue elasticity.\textsuperscript{185}

Emesis with aspiration of gastric contents is another serious complication that can occur during intubation. Avoiding BMV when possible may help reduce the risk of this adverse event. Case reports of adult respiratory distress syndrome and chronic
lumbar disease are thought to be due to aspiration of activated charcoal.185,186 In patients who are obtunded or who are at high risk for seizures or vomiting, consider tracheal intubation before the administration of activated charcoal.

There are ongoing concerns that DL may cause or worsen spinal cord injury in a patient with an unstable cervical spine injury. These concerns are essentially theoretical, with no credible data to prove or disprove a true effect. Many anesthesiologists prefer awake endoscopic intubation in this setting, but no data support one approach over another. A cadaveric study of intubation during fluoroscopy showed that DL with in-line immobilization in the setting of complete C4-5 ligamentous instability did not result in clinically significant movement.187 The greatest degree of motion occurs at the atlanto-occipital junction and decreases with each sequential interspace, and studies of cervical spine instability at these higher levels have not been performed.188 It can also be argued that cadaveric studies do not accurately depict the trauma setting, and yet this model is probably going to remain the best one available. Unless new information emerges regarding the risks of orotracheal intubation with DL, it appears to be a safe approach when performed in conjunction with in-line immobilization. The cervical collar should be removed for intubation, and cervical immobilization held manually by an assistant.

Intubation can be complicated by a persistent air leak. This is generally caused by failure of either the cuff or the pilot balloon or by positioning the cuff balloon between the vocal cords. If the cuff balloon is leaking, replace the tracheal tube (see later section on Changing Tracheal Tubes). If the pilot balloon is determined to be leaking, this can usually be remedied without changing the tube.189 An incompetent one-way balloon valve can be fixed by placing a stopcock in the inflating valve. Reinflate the cuff and shut off the stopcock to solve the problem. If the leak involves the pilot balloon or if the distal inflation tube has been inadvertently severed, cut off the defective part and slide a 20-gauge catheter into the inflation tube. Then connect the stopcock to the catheter, inflate the cuff, and close the stopcock.

Tracheal stricture used to be a significant late complication of long-term intubation with low-volume, high-pressure cuffs. The use of high-volume, low-pressure cuffs has markedly decreased the incidence of this complication.190 Tubes with high-pressure cuffs are obsolete and should be avoided.

**Conclusion**

DL is the most common means of securing a definitive airway. With adequate preparation and emphasis on preoxygenation and positioning, it is usually successful. RSI has rendered patients much more amenable to DL for emergency airway management (see Chapter 5). Once the patient is paralyzed, it becomes the clinician’s supreme responsibility to ventilate, oxygenate, and protect the patient’s airway. Mastery of DL fulfills part of this obligation. Being prepared for failure and having a successful backup plan fulfills the remainder of this responsibility. The remainder of this chapter is devoted to adjuncts and alternatives to DL.

**VIDEO AND OPTICAL LARYNGOSCOPES**

Video and optical laryngoscopes continue to transform airway management. These devices are all somewhat similar in that they use video, fiberoptics, or mirrors to visualize the larynx. They provide a better view of the glottis with less effort and have a shorter learning curve than DL. Only the Macintosh shaped video laryngoscopes are designed to sweep the tongue aside and allow either a direct or video view of the larynx. The other devices are made to look around the curve of the tongue rather than lifting it or pushing it aside, so they are more angulated. The drawback of more angulated devices is that getting an excellent view of the vocal cords does not always correlate with easy intubation191,192 because it can be difficult to manipulate the ET tube around the sharp curve of the blade. Adding a tube channel to the blade obviates the need to manipulate the ET tube around the sharp curve but may add other complexities.193 Because expense is a major impediment to the widespread use of video laryngoscopy, the relatively inexpensive Airtraq device (see Category 3 later) may be a good choice in settings in which difficult intubation is rare and funding is limited. Video laryngoscopy is associated with higher first pass success rates compared to DL, especially in patients with known or suspected difficult airways. Some experts recommend that video laryngoscopy be used for all emergency intubations.194

Video and optical laryngoscopes can be divided into three broad categories:

1. **Video laryngoscopes with standard Macintosh blades (Storz C-MAC, GlideScope Titanium MAC, McGrath MAC, Venner A.P. Advance Mac Blade).**
2. **Video laryngoscopes with angulated blades (GlideScope, McGrath Series 5, Storz D-Blade).**
3. **Video or optical laryngoscopes with a tube channel (Airtraq, Pentax AWS, KingVISION).** (See the following sections for pictures and manufacturers of each device.)

**Video Laryngoscopes With Standard Macintosh Blades**

These devices have blades that are exactly the same or very similar to a standard English or German Macintosh direct laryngoscope. They all have a digital camera adjacent to the light source a few centimeters proximal to the tip. Each of these devices can be used for either conventional DL or video laryngoscopy. The Storz C-MAC (Karl Storz, Tuttingen, Germany) is the only device in this group that has been sufficiently tested (Fig. 4.23).194-202 The laryngeal view obtained on the video monitor is better than the direct view in most patients.193,197 The improved laryngeal view provided by the camera is especially helpful in morbidly obese patients and in trauma patients who require cervical spine immobilization.194,201

Unlike other video laryngoscopy systems, those with Macintosh blades still allow direct visualization of airway structures even if blood or secretions obscure the camera view, or if there are any other camera malfunctions.68 This may help avoid the need to change equipment during the intubation process. The use of a video laryngoscopy system that still allows the ability to perform DL may be ideal for learning DL skills, as novices who learn intubation on a video laryngoscope system have higher success rates when performing DL.87

**Background**

The Storz Video Macintosh Laryngoscope was introduced into clinical practice by Kaplan, Ward, and Berci, and in 2002.200 The latest version of this device, the Storz C-MAC, is smaller,
more portable, and cheaper. The C-MAC consists of a one-piece blade, a cable, and a monitor. It can record digital video or still images onto a Secure Digital memory card. This section will discuss the use of the Storz C-MAC because it is the most used and most tested video laryngoscopy system with a Macintosh blade. The procedure will be similar for other video laryngoscopy systems with a Macintosh blade. The GlideScope Titanium MAC (Verathon Medical, Bothell, WA), McGrath MAC (Medtronic, Minneapolis, MN), and A.P. Advance Mac Blade (Venner Medical, Singapore) appear similar in design, but they are all newer and less tested.

**Indications**

The C-MAC can be used for almost any orotracheal intubation attempt in an emergency setting. If laryngoscopy is not predicted to be difficult, the operator can intubate using DL, with video laryngoscopy immediately available if the airway proves to be difficult or if there is difficulty passing the tracheal tube introducer or ET tube. However, many experts feel that all emergency intubations should be performed with video laryngoscopy, though DL skills remain crucial. The C-MAC allows both direct and video laryngoscopy.

The C-MAC can be used when DL is difficult. In 2006, Kaplan and coauthors published a prospective multicenter trial in which the DL view was compared with the video view during 865 intubations with the Storz Video Macintosh system. They reported 7 cases in which the direct view of the larynx was easy and the video view was difficult and 100 cases in which the direct view was difficult and the video view was easy. In 2009, Jungbauer and colleagues studied 200 patients with Mallampati class III or IV anatomy. They randomized patients to intubation with DL or with the Storz Video Macintosh system. The success rate with DL was 92% and the success rate with the Storz Video Macintosh system was 99%. Other studies also demonstrate that the C-MAC is superior to DL for difficult airways. The C-MAC is also superior to DL after a failed first attempt, though sometimes DL is used as a rescue device when video laryngoscopy fails.

The Storz C-MAC may be a good option for routine intubations at academic institutions that are teaching novice intubators. By viewing the video screen, the senior physician can teach laryngoscopy skills in real time while the learner performs DL. In 2008, Howard-Quijano and coauthors published a prospective, randomized, crossover study of 37 novice intubators (medical students and residents) who performed 222 intubations with the Storz Video Macintosh system. The novice intubators performed only DL and were not allowed to see the video monitor in any case. intubations were randomized so that either the instructors could see the video monitor or it was covered by a drape. Novice intubators were given 2 minutes to intubate before the instructor took over. When instructors were allowed to see the video monitor,

**Video and Optical Laryngoscopy**

**Indications**
- Routine emergency intubation
- Teaching of direct laryngoscopy skills to novices
- Difficult airways
- Failed direct laryngoscopy
- Morbid obesity
- Trauma patients with cervical spine immobilization

**Contraindications**
- Limited mouth opening
- Severe kyphosis
- Copious blood or secretions

**Complications**
- Dental trauma
- Oropharyngeal trauma

**Equipment**

- Storz C-MAC
- Pentax AWS
- McGrath Series 5
- Airtraq

Ancillary equipment (endotracheal tubes, etc) not depicted

*Review Box 4.2* Video laryngoscopy: indications, contraindications, complications, and equipment.
the novice intubators had a 69% success rate and a 3% rate of esophageal intubation. When instructors were blinded to the video monitor, the novice intubators had a 55% success rate and a 17% rate of esophageal intubation.

Two studies compared the Storz C-MAC with the GlideScope and McGrath systems. A 2009 study of 450 patients with normal airways by van Zundert and colleagues showed that those intubated with the Storz C-MAC were intubated more quickly and had a higher first-pass success rate than did those intubated with the GlideScope or McGrath MAC.192 Another study compared the C-MAC with the GlideScope and McGrath in 150 morbidly obese patients and found that those intubated with the C-MAC had significantly fewer intubation attempts and a shorter intubation time.201 The C-MAC is more successful in trauma patients with cervical spine immobilization.206,207

Contraindications
Contraindications to use of the C-MAC are rare. Like other video laryngoscopes, the lens on the video camera is susceptible to obscuration by secretions or blood. Unlike other video devices, the C-MAC facilitates direct visualization of anatomic structures.

Procedure
Much of the procedure for laryngoscopy with the C-MAC is similar to that for DL. When the tip of the Storz C-MAC is advanced into the vallecula, it gives an 80-degree-wide field of view and offers a panoramic view of the glottis. Moving slowly and using a progressive visualization technique to find the tongue, epiglottis, posterior cartilages, and vocal cords (as with DL) is the best approach to performing laryngoscopy with the C-MAC. Keep in mind that although the tip of the C-MAC is usually placed in the vallecula, it may help to directly lift the epiglottis with the tip of the blade if laryngoscopy is difficult; this is known as the straight-blade technique.208

Complications
A complication that may be encountered is blind passage of the ET tube into the mouth while fixating on the video monitor. This can result in damage to oral and pharyngeal structures. It is less common with the C-MAC than with the GlideScope and McGrath MAC.

Video Laryngoscopes With Angulated Blades
The GlideScope (Verathon, Bothell, WA), McGrath Series 5 (Teleflex, Buckinghamshire, UK), and Storz D-Blade (Karl Storz, Tuttlingen, Germany) video laryngoscopes are similarly shaped devices that have sharply angulated, unchanneled, and narrow-flanged blades.191,192,209,210 All these devices have a distal angulation of approximately 60 degrees and a digital camera a few centimeters proximal to the tip. These blades do not allow laryngoscopy by direct visualization. They are designed to follow the natural curvature of the upper airway and look around the tongue rather than displace it. Excellent visualization of the glottis is nearly always achieved when the distal tip of the blade is in or near the vallecula. Because the point of reference is from the location of the camera near the distal tip of these devices, they reliably provide a panoramic view of the larynx that cannot be achieved with DL. Both the GlideScope and the McGrath allow easy visualization of the larynx, even with inexperienced providers or difficult airway anatomy.204,202,209,211–216 Maneuvering the ET tube around the severe angle of the blade and into the trachea is more difficult compared to systems with a Macintosh blade, and this is where problems can occur.195 In one large trial involving inexperienced GlideScope operators, the device was abandoned in 3.7% of intubations despite the fact that an excellent view of the glottis was achieved in most of these cases.209

Background
In 2001, the original GlideScope video laryngoscope became the first commercially available video laryngoscope. The first report of its use was in a case report of the management of a difficult airway.177 Three GlideScope models are now available: the Titanium (Fig. 4.24), which is the newest version that features reusable titanium blades or disposable plastic blades of a similar shape; the AVL, which has both single-use and reusable plastic blades to cover a reusable video baton with a mountable monitor; and the Ranger, which has a small portable monitor and the choice of a reusable blade or a video baton with disposable blades (Ranger Single Use). All GlideScope models have a lens antifog mechanism and digital recording capability. Though this section will discuss hyperangulated blades, the Titanium system also offers blades with a Macintosh shape.

The McGrath portable video laryngoscope is another device with similar characteristics but is more compact. It became available in 2005, and the current model is called the McGrath Series 5 (Fig. 4.25). The shape of the blade is very similar to that of the GlideScope, but it has a small video monitor (1.7 inches diagonally) on the end of the handle. It also has an adjustable-length, detachable metal blade (camera stick) that is covered by a disposable plastic blade during use, so no part of the handle or metal blade makes contact with the patient. It has no recording capability.

The Storz D-Blade became available in 2010 and also has a blade shape similar to that of the GlideScope. It is an optional blade for use with the Storz C-MAC system and has the same functionality as the other C-MAC blades but does not allow the option of direct visualization of the larynx. It was designed specifically for difficult intubations.

The GlideScope and its use are discussed as representative of this class of indirect video-assisted laryngoscopes with thin, sharply angulated and unchanneled blades.

Indications
Like the C-MAC, video laryngoscopy systems with hyperangulated blades can also be used for routine intubations and...
have higher success rates than DL.218,219 The GlideScope, McGrath Series 5, and Storz D-Blade may be especially useful when DL is difficult or fails. In 2010, Noppens and coauthors reported a series of 61 patients with failed DL, all with Cormack-Lehane grade III and IV views, who subsequently had good laryngoscopic views and 95% were successfully intubated with a McGrath video laryngoscope.220 In 2011, Cavus and colleagues published a very similar series of 20 patients with grade III and IV views who had failed DL but subsequently had good laryngoscopic views and were all successfully intubated with the Storz D-Blade.221 There are many case reports of easy intubations with the GlideScope and McGrath video laryngoscopes after failed DL.217,222 In addition, several case reports show that the GlideScope and McGrath can facilitate successful awake intubation in patients with known or suspected difficult airways.223–225

A significant advantage of the more angulated video laryngoscopes is that they can provide a good view of the larynx when the neck is in the neutral position. Not surprisingly, there are several small studies and case reports demonstrating that the GlideScope and McGrath provide good laryngoscopic views and a high rate of successful intubation in patients with cervical spine immobilization.226–228 There is also a case report of a patient with severe ankylosing spondylitis who was easily intubated on the first attempt with the McGrath.229

There may be patients in whom the larynx is so anterior that devices with a Macintosh blade are unable to adequately visualize the airway structures; hyperangulated blades may be especially useful in this situation.

Finally, the GlideScope is useful for routine intubations by inexperienced or novice intubators, because it takes more than 50 intubations with DL to become proficient.89 In a 2009 study by Nouruzi-Sedeh and coworkers, 20 novices each intubated five patients with DL and five patients with the GlideScope. They had a 51% success rate with DL and a 93% success rate with the GlideScope.214

**Contraindications**

Contraindications to using angulated video laryngoscopes are not well described in the literature. As with any video or optical device, blood or secretions on the lens may decrease visualization of the larynx. Fogging was a problem with older endoscopic devices, but this problem is rare with the newer video devices. Limited mouth opening (<2 cm) can make insertion of these devices more challenging, though they require less mouth opening than devices with Macintosh blades.191

The biggest contraindication is probably lack of experience or absence of a rigid stylet or tracheal tube introducer to allow the operator to maneuver the ET tube around the sharp angle of the blade.191

**Procedure and Technique**

Grasp the video laryngoscope in the left hand and place it in the patient’s mouth with the scissor technique. Under direct vision, not by viewing the screen, advance the blade through the oropharynx along the midline of the tongue (Fig. 4.26, step 4). Then look up at the monitor while continuing to advance the blade down the midline of the tongue, progressively identifying the base of the tongue and the epiglottis. With a gentle lifting motion, place the tip of the blade in the vallecula. Elevate the epiglottis and expose the laryngeal inlet (see Fig. 4.26, step 5). If the glottis is not well visualized, tilt the handle back slightly to enhance exposure. If needed, manipulate the neck externally to enhance visualization. If more exposure is required, place the tip of the blade under the epiglottis and gently lift and tilt back. While attempting to optimize the laryngeal view, be careful to not place the blade too close to the laryngeal inlet because it may tip the larynx anteriorly and inferiorly, thus making it more difficult to access the laryngeal inlet and pass the tube through it.

Use a rigid steel GlideRite (Verathon Medical) stylet, which has the same 60-degree curve as the blade of the GlideScope, McGrath, and D-Blade. Alternatively, a malleable stylet with a 60-degree distal bend or a bougie may be acceptable, but these devices may fail if tube passage is challenging. Pass the ET tube through the oropharynx under direct visualization until it passes under the curve of the blade, and then look for it on the monitor (see Fig. 4.26, step 5). If needed, manipulate the neck externally to enhance exposure. If more exposure is required, place the tip of the blade under the epiglottis and gently lift and tilt back. While attempting to optimize the laryngeal view, be careful to not place the blade too close to the laryngeal inlet because it may tip the larynx anteriorly and inferiorly, thus making it more difficult to access the laryngeal inlet and pass the tube through it.

Use a rigid steel GlideRite (Verathon Medical) stylet, which has the same 60-degree curve as the blade of the GlideScope, McGrath, and D-Blade. Alternatively, a malleable stylet with a 60-degree distal bend or a bougie may be acceptable, but these devices may fail if tube passage is challenging. Pass the ET tube through the oropharynx under direct vision until it passes under the curve of the blade, and then look for it on the monitor (see Fig. 4.26, step 6). Decrease the chance of soft tissue injury by carefully passing the ET tube along the side of the GlideScope.230 When the tip of the ET tube comes into view, direct it into the glottis and advance it to the appropriate depth (see Fig. 4.26, step 7). If a malleable stylet or bougie is used, it may be difficult to get the tube to go anterior enough. If a malleable stylet must be used, try introducing the tube from the right side of the patient and rotating it 90 degrees and vertically into a midline position behind the tongue. This will help the tip of the stylet maintain its shape as it passes through the oropharynx. Another option is to use a Parker Flex-It (Parker Medical, Engelwood, CO) intubation stylet to direct the tip of the ET tube anteriorly.209 It is important to realize that the fixed steel (GlideRite) stylet needs to be withdrawn as the ET tube is passed through the vocal cords because the tip of the stylet will be essentially pointing toward the ceiling as it approaches the laryngeal inlet (see Fig. 4.26, step 9). The large knob on the proximal end of the GlideRite stylet is designed to be pushed up by the tip of the intubator’s thumb, thus making ET tube advancement and stylet withdrawal a one-handed procedure.

**Complications**

Several relatively minor complications have been reported with use of the GlideScope. There are case reports of injury to the soft palate and tonsillar pillars, as well as two cases of puncture.
Video Laryngoscopy (GlideScope)

1. Prepare the endotracheal tube. Depending on the system that you are using, insert a rigid stylet or preload the tube onto the device.

2. Check the rest of your equipment, including the light source and video monitor.

3. Preoxygenate and premedicate the patient as clinically indicated.

4. Place the blade into the mouth under direct visualization (don't look at the video monitor yet). Keep the blade in the midline of the tongue throughout the procedure.

5. Slowly advance the blade while watching the video monitor. Progressively identify the tongue and epiglottis.

6. Under direct visualization (don't look at the monitor), pass the styleted tube through the mouth and into the posterior pharynx.

7. Place the blade in the vallecula or under the epiglottis, gently lift, and identify the vocal cords.

8. Direct the tube through the vocal cords under direct visualization on the video monitor.

9. While the operator firmly secures the endotracheal tube, an assistant removes the stylet as the tube is passed through the vocal cords.

NOTE: This image sequence depicts video laryngoscopy using the GlideScope. The procedure sequence may vary with the use of other devices. Refer to text for additional information. Be familiar with the equipment used at your institution prior to use.

Figure 4.26 Video laryngoscopy (GlideScope, Verathon Medical, Bothell, WA). Note that there are times when the operator does not look at the screen but rather uses direct vision (steps 4 and 6).
of the right palatopharyngeal arch, one requiring surgical repair.230-232

Summary
Video-assisted laryngoscopy is a major advancement in visualization of the laryngeal inlet. Hyperangulated blades are very promising because they provide an improved laryngeal view and have a shorter learning curve than DL. The majority of intubation failures are due to an inability to pass the tube through the larynx despite excellent glottic views.239 Using a fixed steel stylet (GlideRite stylet) helps, and does not eliminate this problem.239 Advancements need to be made in achieving tracheal intubation once the glottis is seen. The addition of a tube channel obviates the need to maneuver the ET tube around the sharp angle of the blade; channeled devices are described in the next section.

Video and Optical Laryngoscopes With a Tube Channel
Four devices allow indirect video or optical visualization of the larynx and also provide a channeled blade to guide the ET tube through the vocal cords. The Airtraq (Fig. 4.27) (Prodol Meditec S.A., Vizcaya, Spain) and Pentax AirWay Scope (Fig. 4.28) (AWS) (Pentax, Tokyo) are the only devices in this group that have been sufficiently tested. The King Vision aBlade (Ambu, Ballerup, Denmark) and Venner A.P. Advance Difficult Airway Blade (DAB) (Venner Medical, Singapore) are newer and less tested. The ET tube is inserted into the channel before the device is placed in the airway, and the channel guides the ET tube into the trachea once the glottis is visualized. This design helps alleviate the most common problem associated with video and optical laryngoscopy: difficulty placing the ET tube despite an excellent view of the glottis.

All the channel-guided devices have the same curvature as the normal upper airway. They allow visualization of the glottis by looking around the tongue instead of trying to straighten the airway and push the tongue out of the way. They consistently provide a better view of the glottis and may lead to less airway trauma and hemodynamic stimulation than occurs with DL.231-235 The concept of a laryngoscope shaped like the natural curvature of the upper airway and containing a channel to guide the ET tube is not new, but the addition of optics and video just recently made this concept a reality.235-238 The advantage of these devices is that they provide an excellent view of the glottis with little need for maneuvering and they eliminate the requirement for the hand-eye coordination needed to pass the ET tube around an acute curve.240-243

Indications
Difficult intubation with DL is an indication for using a channel-guided optical or video device. The Airtraq and Pentax AWS have a high rate of success in patients with known or predicted difficult airways.213,233,234 In 2009, a study by Asai and colleagues reported 270 patients with difficult airways by DL (grade III and IV views) and found that 268 of these patients (99.3%) had good glottic visualization and were easily intubated with the Pentax AWS.234 In a 2009 trial of patients with expected difficult airways, Malik and coworkers randomized 75 patients to DL, the Pentax AWS, or the GlideScope.213 The rate of successful intubation was 84% with DL, 96% with the GlideScope, and 100% with the Pentax AWS.

The Airtraq and Pentax AWS are particularly useful for difficult airways because of cervical spine immobilization and morbid obesity.242-245 In 2007, Ndolo and coauthors published a study of 106 morbidly obese patients who were randomized to intubation with DL or the Airtraq.246 All patients randomized to the Airtraq were successfully intubated, and six patients in the DL group required rescue intubation with the Airtraq. In 2007, Maharaj published a study of 40 patients with cervical spine immobilization who were randomized to DL or the Airtraq.244 All patients randomized to the Airtraq group were intubated on the first attempt and were intubated faster and with less difficulty than the DL group.

In 2008, Enomoto and associates studied 203 patients with manual in-line neck stabilization who required intubation.243 They randomized these patients to intubation with DL or the Pentax AWS. The intubation success rate with DL was 89% versus 100% with the Pentax AWS. A similar study by Liu...
and coworkers in 2009 included 70 patients with manual in-line stabilization of the cervical spine and randomized them to intubation with the Pentax AWS or the GlideScope. They reported a 100% rate of successful intubation with the Pentax AWS and 89% with the GlideScope. In addition, two studies used fluoroscopy to compare cervical spine motion during intubation with DL versus intubation with the Pentax AWS. Both studies found that intubation with the Pentax AWS caused less movement of the upper cervical spine than DL.

The Airtraq is generally used during RSI but may also be used for awake intubation of patients with difficult airways. When compared with DL, both the AirTraq and Pentax AWS consistently provide better visualization of the glottis and have a higher rate of successful intubation. Two small studies and one large trial have shown that the Airtraq has a very high rate of success after failed DL. In 2011, Amathieu and coauthors reported the results of a large prospective trial in which they used an algorithm that specified use of the Airtraq as a backup for all cases of failed RSI with DL. They had 28 patients who could not be intubated with DL after multiple attempts, including the use of a bougie. Intubation with the Airtraq was successful in 27 of 28 of these patients (97%), and the other patient was intubated with an ILMA.

**Contraindications**
The greatest drawback of channeled video and optical devices is that copious amounts of blood or secretions can obscure the view. Because they do not allow a direct line of sight to the larynx, visualization and intubation are dependent on the video or optical image. If blood or fluid covers the tip of the lens, the image is obscured. This problem can be minimized by aggressively suctioning the hypopharynx before placing the device in the mouth.

Inability to open the mouth or severely limited mouth opening is a contraindication to using the channel-guided devices just described. If the patient’s mouth opening is at least 2 cm, these devices can succeed in cases in which DL would be impossible. The normal adult-size Airtraq requires 18 mm of mouth opening, and the pediatric and infant sizes require 12.5 mm. The Pentax AWS requires approximately 20 mm of mouth opening.

**Procedure**
Before beginning the procedure check that you are using the correct size device, ensure that it is functioning properly, and choose the correct size ET tube for the device and the patient. Insert the ET tube into the channel of the device and advance it to the end of the channel while ensuring it is not advanced so far that it obscures the video or optical lens. Insert the tip of the blade into the mouth vertically so that the handle of the device is pointed toward the patient’s feet. Rotate it into the pharynx and hypopharynx along the midline of the tongue until the tip is in the vallecula. If the glottis is not visualized immediately, try backing the blade out 1 to 2 cm and lift the device gently in the direction of the handle (toward the ceiling). A common mistake is to insert the blade too deep initially, which offers a narrower view of the glottis and may make intubation difficult. When the vocal cords are well visualized in the center of the video or optical image, advance the ET tube through the cords by sliding it forward within the channel. If the ET tube does not go through the cords, pull it back and realize that you need to adjust the position of the entire device to change the trajectory of the tube. If the tube tends to go posterior to the vocal cords, lift the device toward the ceiling and tip the handle back slightly (while avoiding contact with the upper teeth), and then advance the tube again. Alternatively, pass a bougie through the ET tube (or in the channel without an ET tube) and direct the curved tip up and through the vocal cords.

When using the Pentax AWS, it is important to know that the ET tube leaves the channel in line with the tip of the blade (more anterior than the Airtraq). Therefore when the tip of the AWS blade is in the vallecula, the tube often strikes the epiglottis when it is advanced. To avoid this problem, advance the tip of the AWS blade posterior to the epiglottis (as with straight-blade laryngoscopy), and lift the epiglottis out of the way before advancing the ET tube.

**Aftercare**
A common mistake when using laryngoscopes with a tube channel is to quickly remove the laryngoscope after the ET tube is advanced into the trachea. Because the ET tube is already inside the trachea, it is best to attach a resuscitation bag and ventilate the patient. After giving several breaths, stabilizing oxygenation, and confirming tracheal placement with capnography, slowly and carefully remove the device while providing ongoing ventilation. It is appropriate to leave the laryngoscope in place for a few minutes after intubation to allow visual reconfirmation of proper tracheal placement, particularly if there is a problem with ventilation or oxygenation immediately following intubation. To remove the tube from the channel of these devices, hold the end of the ET tube firmly in the right hand and carefully wiggle the device to the left. When the tube slides out of the channel, slowly rotate the laryngoscope out of the patient’s mouth. Reconfirm tracheal placement with capnography and adjust the depth of the ET tube as needed.

**Complications**
No significant complications have been reported with the tube channel video and optical devices. Because the blades of these devices are rigid and cannot be directly visualized after entering the mouth, they have potential for soft tissue damage. Complications of other video laryngoscopes are primarily a result of blind ET tube placement through the mouth and pharynx, and the tube channel devices eliminate this problem.

**Intubating Laryngeal Mask Airways**
ILMAs are unique devices because they are easy to insert, provide excellent ventilation and oxygenation (better than BMV), and also provide a reliable means of tracheal intubation. Many clinicians who have used the Fastrach ILMA in an emergency setting insist on having this device in their difficult airway tool kit. It is impossible to overstate the value of this device for failed emergency RSI, and unlike other rescue devices, a large volume of data support its use in general and in the setting of failed RSI. It has become the primary rescue device for failed RSI in many EDs.
Intubating Laryngeal Mask Airway

**Indications**
- Failed rapid-sequence intubation
- Cannot-intubate/cannot-ventilate scenario
- Difficult mask ventilation
- Expected difficult intubation
- Refractory hypoxemia despite preoxygenation

**Contraindications**
- Unable to open mouth
- Awake patient

**Complications**
- Laryngeal or esophageal injury from blind intubation
- Aspiration (rare)

**Equipment**
- Intubating LMA (ILMA) Fastrach
- Large-volume syringe
- ILMA stabilizer rod
- ILMA endotracheal tube
- Standard endotracheal tube (optional)
- 10-mL syringe

**Types of ILMAs**

There are four popular ILMAs currently available (see Chapter 3, Fig. 3.12). The LMA Fastrach (Teleflex; http://www.lmaco.com) has, by far, been the most used and most studied ILMA. It has excellent blind intubation rates, much higher than the Air-Q (Cookgas LLC, St. Louis, MO) or i-Gel (Intersurgical, Berkshire, UK) and there are numerous investigations detailing its success as a rescue device in difficult and failed airways. A unique feature of the Fastrach is the metal handle that makes insertion easier and enables lifting of the device to create a better seal against the glottis. There are also newer ILMA devices that show promise: the Air-Q, i-Gel, and Auragain (Ambu, Ballerup, Denmark). Whereas the smallest Fastrach is size 3 (for children 30 to 50 kg), the other ILMAs are available in all sizes, including neonates less than 5 kg. The LMA Protector (Teleflex; http://www.lmaco.com) is an ILMA that will soon be released, and is similar to the LMA Supreme (with gastric access) but will allow intubation using a flexible endoscope.

An important feature of newer ILMAs is gastric access channels, allowing placement of an orogastric tube through the gastric port and into the stomach with the ILMA in place. This is particularly useful in patients with distended stomachs after prolonged BMV, and in children who are more prone to gastric insufflation even with shorter BMV times. The Air-Q is available in four versions, one of which has a gastric port (Air-Q Blocker). The i-Gel and Auragain both have gastric ports. The LMA Fastrach does not have gastric access.

**Indications**

ILMAs are indicated as an alternative to BMV or as a conduit for intubation of difficult airways. In the cannot-intubate/cannot-ventilate scenario, it is a reliable rescue device. In this situation, adequate ventilation with the ILMA is possible in nearly all cases. The ILMA is more successful for ventilation and intubation of difficult airways than the LMA Classic. For inexperienced providers, ventilation with the ILMA is usually superior to face mask ventilation.

The ILMA can also be used as a primary ventilation and intubation device in patients with difficult airways. Studies of difficult airway management with the ILMA show that almost all patients can be adequately ventilated with the ILMA and 94% to 99% can be intubated through the device. Patients who are difficult to intubate by DL are often easy to intubate with the ILMA because many anatomic factors that cause difficult DL do not affect placement or function of the LMA Fastrach.

The LMA Fastrach is especially useful in patients with difficult face mask ventilation because of a beard, severe facial trauma, or obesity because none of these factors inhibit Fastrach placement. When brisk bleeding above the glottis makes ventilation and intubation difficult, the Fastrach can prevent aspiration of blood and facilitate blind or endoscopic intubation. In patients requiring urgent cricothyrotomy or percutaneous needle insertion into the trachea, the ILMA can be used to counteract anterior neck pressure. In this capacity, the Fastrach provides temporary ventilation and aids in stabilization of the cervical spine during the surgical airway procedure.

*References 2, 53, 55, 57, 128, 252-254, 258, 267, 268.*
Contraindications

One limitation of the LMA Fastrach is that it cannot be used in infants and small children because the smallest size, a No. 3, is not suitable for patients smaller than 30 kg. Other ILMAs (or nonintubating LMAs) should be used in patients less than 30 kg. Other ILMAs require flexible endoscopic guidance for reliable tracheal intubation.

LMAs are contraindicated in patients with less than 2 cm of mouth opening. They are unlikely to be successful in patients with grossly distorted supraglottic anatomy from disease processes or posttrauma scarring. They are also relatively contraindicated in awake patients because of the high risk for emesis when the gag and airway reflexes are intact.

Intubation through the LMA Fastrach

The majority of intubations through the ILMA are performed blindly by using either the designated LMA ET tube or a standard ET tube. Regardless of the tube used, it is critical that the ILMA be optimally adjusted before attempting blind or flexible endoscopic intubation through the device.

The LMA ET tube, also known as the LMA Fastrach, is designed specifically for the ILMA. There are two versions of the LMA ET tube: a reusable and a single-use disposable. The reusable version is made of silicone and the single-use version is made of polyvinyl chloride (PVC). The specialized LMA ET tubes are soft and straight and have a midline-beveled tip. These features are designed to allow the LMA ET tubes to emerge from the ILMA mask at an acute angle and to minimize potential injury to the vocal cords and esophagus. The drawback of the specialized Fastrach ET tubes is that they have low-volume high-pressure cuffs, which could potentially cause ischemic damage to the trachea, and there are no clinical data on how long these tubes can remain in place. Use of a standard ET tube for intubation through the LMA Fastrach has been well studied and is the current practice in many EDs. Another option is to use a Parker Flex-Tip ET tube (Parker Medical) (see Fig. 4.8).

Procedure and Technique for Blind Intubation

The LMA Fastrach is the only device that enables reliable tracheal intubation without the use of a flexible endoscope. If another ILMA is used, tracheal intubation has much lower success rates and, in general, blind intubation should not be attempted.

Before intubation through the LMA Fastrach, make sure that the patient is ventilating optimally through the device. Determine this by manually ventilating the patient while holding the ILMA handle with a “frying-pan” grip. If any resistance is felt, adjust the handle by slight rotation in the sagittal plane and then lift the entire device toward the ceiling (see the “Chandy maneuver,” Fig. 4.29, step 3).

Before inserting the ET tube, lubricate it generously. Advance the ET tube with the curve opposite that of the LMA Fastrach curve (Fig. 4.29, step 5). When the tube has advanced to 15 cm, the tip will start to emerge from the LMA Fastrach mask. Just before advancing the tube, use the frying-pan grip and apply a slight anterior lift (not a tilt) to further align the aperture of the ILMA with the glottis (see second part of the Chandy maneuver, Fig. 4.29, step 6). Do not use a levering action. While holding the handle in this position, gently pass the tracheal tube to approximately 16.5 cm. In this position, the ET tube will push the epiglottic elevating bar up and may now come in contact with the larynx or esophagus. If cricoid pressure is being applied, decrease it because it may interfere with passage of the ET tube through the glottis. If no resistance is encountered, advance the tube into the trachea until the tracheal tube adapter comes in contact with the proximal end of the ILMA tube (see Fig. 4.29, step 7). Do not use force when advancing the tube.

If the ET tube does not pass into the trachea easily, withdraw the ET tube to the 15-cm mark and readjust the position of the LMA Fastrach. If the tube meets resistance at approximately 17 cm, this may indicate a fully down-folded epiglottis or impaction of the tip of the tube against the anterior laryngeal wall. Rotating the tube may overcome impaction of the tip. To correct a down-folded epiglottis, remove the ET tube and perform the “up-down maneuver” by rotating the ILMA outward 5 to 6 cm without deflating the mask and then sliding it back into the hypopharynx (see Chapter 3, Fig. 3.13, steps 9 and 10). If these maneuvers are unsuccessful, it is likely that the wrong size LMA Fastrach is being used. Consider using a flexible endoscope to guide intubation (also see the “Insertion Technique and Maneuvers Guide,” on the company website: http://www.lmana.com).

Once the LMA ET tube has passed into the trachea, inflate the tube cuff and attempt to ventilate the patient. Check for proper tube placement with a PETCO2 detector. If the tube is in the trachea, deflate the cuff of the ILMA. There is no rush to remove the ILMA; it can remain in place for an hour or longer if more pressing patient care issues need to be addressed first.

Using a Standard ET Tube

Use of a standard ET tube is not recommended by the manufacturer, but it is a well-studied and common practice. The manufacturer warns that using a standard ET tube may be associated with a greater likelihood of laryngeal trauma, but there are no reports of such trauma in the literature. The only report of significant trauma with blind intubation through the LMA Fastrach is a case report of esophageal perforation and subsequent death caused by use of the specialized LMA ET tube. A laboratory study showed that a standard PVC ET tube exerts 7 to 10 times more pressure on distal structures than the silicone LMA ET tube, and the clinical relevance of this finding is unknown. Some experts suggest warming the tip of a standard PVC ET tube (to soften it) before insertion through the LMA Fastrach.

Because of the potential for injury with blind intubation through the ILMA, use endoscopic guidance if there is any difficulty. If using a standard ET tube, insert the ET tube with its curvature opposite the curvature of the LMA Fastrach tube (see Fig. 4.29, step 5). This allows the ET tube to exit the Fastrach at a less acute angle and then to advance into the trachea more easily.

If intubation through the LMA Fastrach fails, the device can still be used to provide ventilation and oxygenation during more invasive airway procedures, such as retrograde intubation or cricothyrotomy.

Flexible Endoscopic Intubation through LMAs

A flexible bronchoscope can be used to verify the position of the larynx either before or during intubation. When intubating
Endotracheal Intubation With the ILMA (Fastrach)

1. Insert the ILMA by sliding the posterior surface of the mask along the palate and posterior pharynx until firm resistance is felt (see Chapter 3).

2. Inflate the ILMA cuff and begin ventilations through the device as you prepare for intubation.

3. CHANDY MANEUVER
   Step 1
   Rotate handle in the sagittal plane to move the cuff forward against the larynx.
   This maneuver provides a good seal and optimal ventilations.

4. Insert the LMA Fastrach ET tube to the 15-cm mark.
   At this point, the tip of the tube will begin to emerge from the LMA.

5. Correct
   If using a standard ET tube, align the curve so that it is opposite of the LMA curve.
   This allows the tube to exit the mask at an angle more conducive to tracheal entry.

6. CHANDY MANEUVER
   Step 2
   Prior to passing the tracheal tube, lift the handle anteriorly (not tilted) to better align the cuff and larynx for smooth ET tube passage.

7. If no resistance is felt, advance the tube into the trachea until the adapter comes into contact with the ILMA airway tube.
   Do not use force.

8. Inflate the ET tube cuff, ventilate, and check tube position with PETCO₂.
   If the tube is properly positioned, deflate the LMA cuff.

9. To remove the LMA, deflate the LMA cuff and remove the adapter from the tube.
   Rotate the LMA out of the mouth while using the stabilizer rod to keep the tube in place.

10. Remove the stabilizing rod when the mask is clear of the mouth, and complete removal of the LMA.
    Reattach the adapter and resume ventilation.

Figure 4.29 Endotracheal (ET) intubation with the intubating laryngeal mask airway (ILMA) (Fastrach, Teleflex, Buckinghamshire, UK).
through the ILMA over an endoscope, a standard ET tube is sufficient and there is no reason to use the specialized LMA ET tube.

To use a flexible endoscope to intubate through the LMA Fastrach, first advance the ET tube through the ILMA Fastrach to 15 cm. At a 15-cm depth the view through the endoscope should show the glottis beyond the epiglottic elevating bar. Advance the ET tube 1.5 cm before advancing the endoscope. This protects the camera elements from being damaged by the epiglottic elevating bar. The view at 16.5 cm should show the vocal cords and trachea. Advance the endoscope into the trachea and then pass the ET tube over the endoscope. If the vocal cords are not visualized immediately, see the LMA Insertion Technique and Maneuvers Guide, described in Chapter 3.

To use a flexible endoscope to intubate through other LMAs, the procedure is similar. First, place the ET tube approximately halfway to the opening in the laryngeal mask bowl. At this point, it may be useful to connect a swivel adapter to the ET tube adapter (see Fig. 4.29) that will allow ongoing ventilation with the ability to place the flexible endoscope through the other port. If this step is performed, the cuff of the ET tube should be inflated during endoscopic intubation to minimize the air leak during ventilation. After the ET tube is placed halfway to the opening of the laryngeal mask bowl, the endoscope is introduced into the ET tube and advanced until the laryngeal inlet is visualized. Advance the endoscope into the trachea and then pass the ET tube over the endoscope. If the laryngeal inlet is not immediately visualized, the laryngeal mask should be repositioned before attempting intubation.

For children, the most common flexible bronchoscope has an outer diameter of 3.6 to 3.8 mm, which enables intubation of a 4.5 ET tube or larger. This allows intubation through an ILMA for children as small as 2 to 3 years old. Children younger than this should not be intubated through an ILMA, unless a smaller endoscope is used.

**ILMA Removal After Intubation**

To remove the ILMA, first deflate the cuff, if present (see Fig. 4.29, step 8), and be careful to not deflate the ET tube cuff. Start by removing the ET tube adapter. Then hold the proximal end of the ET tube in place while rotating the ILMA out of the hypopharynx. As the ILMA passes over the ET tube and out of the mouth, hold the ET tube in place with a stabilizer rod (see Fig. 4.29, step 9), without advancing the tube farther. If a stabilizer rod is not available, a second ET tube can also be used to aid in ILMA removal. When the ET tube pilot balloon comes into contact with the stabilizer rod, remove the stabilizer rod to allow the pilot balloon to travel through the ILMA tube. Then reattach the ET adapter and resume ventilation. Adjust the depth of the ET tube as needed.

**Intubation through the LMA Classic**

The recommended technique for tracheal intubation through the LMA Classic uses an endoscope and has a high success rate but requires a smaller ET tube and some adjustments. Blind intubation through a nonintubating LMA (LMA Classic or LMA Unique) has a poor success rate and is not recommended. Using a tracheal tube introducer as a guide through the LMA is unlikely to be successful and is not recommended.

After ensuring that the LMA is ventilating properly, place a well-lubricated ET tube into the LMA tube and advance it to a depth of 24 cm (No. 5 LMA), so that the tip of the ET tube has just passed the fenestrations. Pass a lubricated endoscope through the ET tube and advance it through the vocal cords. If the epiglottis is deflected downward, manipulate the tip of the endoscope under the epiglottis until the vocal cords come into view. Pass the ET tube over the endoscope and into the trachea, and then inflate the ET tube cuff and ventilate the patient. Check for correct ET tube placement with a PETCO₂ detector. Cricoid pressure may impede placement and intubation through the LMA. Release cricoid pressure, if necessary, to accomplish these procedures.

**Complications When Intubating through LMAs**

Although most patients can be safely intubated blindly through the ILMA, there is a small chance of injury to the larynx or esophagus, especially with multiple blind attempts. There is one case report of a death caused by esophageal perforation during blind intubation through the ILMA. Consider using endoscopic guidance if blind intubation is difficult or if the clinician is inexperienced. Intubation through a nonintubating LMA is not recommended unless facilitated by an endoscope.

**Summary**

LMAs provide an excellent means of oxygenation and ventilation when face mask ventilation is difficult or impossible. In addition, ILMAs are particularly useful when managing patients who are difficult to intubate with DL. By facilitating the management of difficult or failed ventilation and difficult or failed intubation, ILMAs are an indispensable component of the difficult airway algorithm.

**FLEXIBLE ENDOSCOPIC INTUBATION**

Flexible endoscopic intubation is the most common technique used by anesthesiologists for known difficult airways. Physicians who perform endoscopic intubations daily have a success rate of nearly 100% when using this technique for difficult intubations. In the ED, with more difficult intubating conditions and less experienced fiberoscists, the success rate is 50% to 90%, 286–290 The most common reason cited for failure of flexible endoscopic intubation in the operating room is clinician inexperience. In the ED, failure is most often attributed to poor visibility from blood, vomitus, and other secretions. 288–290 Mlinik and colleagues found that successful ED endoscopic intubations averaged 2 minutes whereas failures averaged 8 minutes. They recommended consideration of alternative approaches if intubation attempts take more than 3 minutes. Flexible endoscopic intubation is often the best method for intubating awake patients with a known difficult airway. It can be accomplished via the nasal or oral route and is better tolerated than DL. It usually provides excellent visualization of the airway and permits evaluation of the airway before placement of the tube. The expense of the equipment, its fragility, and the length of time required to both achieve and maintain technical proficiency are drawbacks. Because intubation with a flexible endoscope is performed much less commonly than other methods, clinicians are encouraged to learn and maintain
Flexible Fiberoptic Intubation

Indications
- Known or suspected difficult airway
- Distorted airway anatomy
- Swelling
- Abscess/infection
- Morbid obesity
- Trauma
- Tumors/previous radiation therapy

Contraindications
- Nasal approach
  - Severe midface trauma
  - Coagulopathy
- Relative
  - Active airway bleeding
  - Vomiting

Complications
- Hypoxia from prolonged intubation attempts
- Vomiting
- Laryngospasm
- Soft tissue trauma

Equipment
- Flexible fiberoptic bronchoscope (FOB)
- Endotracheal tube

Review Box 4.4 Flexible endoscopic intubation: indications, contraindications, complications, and equipment.

this skill when performing bedside nasopharyngoscopy for patients with severe sore throat, new hoarseness, suspected foreign body, possible laryngeal injury after neck trauma, and in other cases when upper airway edema, obstruction, or pathology is suspected. All clinicians who perform airway management should also perform nasopharyngoscopy with laryngoscopy on a regular basis. Expense has become less of an issue in recent years with the introduction of disposable endoscopes like the Ambu aScope (Ambu, Ballerup, Denmark).

Endoscopes are graded according to their external diameter, measured in millimeters. Endoscopes specifically designed for ET intubations are available from several companies (Pentax, Olympus, Machida, Storz, and Fujinon). A practical size for an intubating scope is approximately 4 to 5 mm. Although it is physically possible to pass an ET tube sized 0.5 mm larger over an endoscope, the fit is tight. As a rule, the ET tube should be approximately 1 mm larger than the intubating scope. The size of the working channel, the port that allows suction, administration of oxygen, and passage of fluid or catheters, are also important when evaluating endoscopes. A working channel of approximately 2 mm is desirable to allow adequate suction of secretions, though working channels are not strictly necessary for endoscopic intubation.

Older fiberoptic systems required the intubator to look into an eyepiece when performing intubation. Newer flexible endoscopic systems plug into a video monitor, enabling assistants and learners to visualize the airway anatomy; newer systems enable the intubator to maintain a more comfortable position while holding the endoscope and sheath properly.

Indications and Contraindications
Patients with known or suspected difficult airways are good candidates for awake or semi-awake endoscopic intubation. Patients with distorted airway anatomy, including swelling of the mouth or tongue, upper airway abscess or infection, morbid obesity, cervical spine injury, trismus, and penetrating and blunt neck trauma, are all good candidates for awake endoscopic intubation. Patients with laryngeal tumors, especially those with a history of radiation therapy encompassing the cervical region, may be impossible to intubate by any other nonsurgical method. An endoscope can also be helpful when assessing and intubating patients with airway obstruction from presumed foreign body aspiration. Flexible endoscopic intubation is best used as the initial approach to tracheal intubation, and it may also be used as a rescue device when other methods fail.

Flexible endoscopic intubation can also be performed through an ILMA or LMA after difficult ventilation or failed intubation.

Contraindications to the nasal approach are severe midface trauma and coagulopathy. Patients who are likely to receive thrombolytics should also be excluded. Although there are no clear contraindications to endoscopic orotracheal intubation, active airway bleeding, excessive oral sections, and vomiting are relative contraindications because successful endoscopic intubation is rarely achieved in these settings. Because of the time that must be expended preparing for and performing flexible endoscopic intubation, patients with impending airway closure from a dynamic process that is causing severe upper airway obstruction or swelling should have their airway secured by other means. Hypoxia despite good attempts at oxygenation is another relative contraindication, especially if the intubator is inexperienced in flexible endoscopic intubation.

Procedure and Technique

Preparation
Proper preparation of the upper airway is crucial for successful awake or semi-awake endoscopic intubation. If time permits,
administration of a drying agent such as glycopyrrolate 10 to 20 minutes before the procedure will reduce oral secretions and increase effectiveness of topical anesthesia. Topical anesthesia should be applied to the posterior oropharynx, hypopharynx, and larynx for all endoscopic intubations. If nasal intubation is planned, administer topical anesthesia to the selected naris; application of a vasoconstrictor such as oxymetazoline (Afrin, Bayer, Whippany, NJ) or phenylephrine is recommended to increase the caliber of the nasal passage and reduce the chance of epistaxis.

Deliver local anesthetic to the upper airway by one of several methods. Application of 4% or 5% lidocaine cream by “buttering” the base of the tongue is an effective technique to anesthetize the posterior tongue, vallecula, epiglottis, and laryngeal structures. The tongue is held in protrusion as the base of the tongue is buttered with 4% or 5% lidocaine cream using a tongue depressor. Because the patient is unable to swallow with a protruded tongue, the ointment warms, liquefies, and moves posteriorly and inferiorly to anesthetize the laryngeal structures. It is contended that the ointment also penetrates the mucosa to anesthetize the glossopharyngeal and superior laryngeal nerves. Nebulized lidocaine (4 to 6 mL of a 4% solution) can be used to anesthetize the entire upper airway if time permits. Lidocaine (3 mL of a 4% solution) can be injected percutaneously through the cricothyroid membrane via a 20-gauge needle, thereby providing anesthesia to the larynx and trachea. A flexible atomizer device that attaches to a standard syringe (Fig. 4.30) (LMA MADgic, Teleflex) can be used to accurately apply atomized 4% aqueous lidocaine to the posterior oropharynx and upper laryngeal structures; tracheal anesthesia can be accomplished by having the patient inspire as the medication is sprayed. Some laryngeal and tracheal anesthesia can be achieved by oral spray with a laryngeal tracheal anesthetic set, but this is probably less effective than other methods. Finally, 4% aqueous lidocaine can be sprayed through the working channel of the endoscopic scope during the procedure via the “spray as you go” technique.

Nasal anesthesia can be accomplished by several methods. A flexible atomizer (Fig. 4.31) can be passed to the posterior nasopharynx, and then 4% aqueous lidocaine is sprayed as the patient sniffs. This is repeated several times as the flexible atomizer is slowly withdrawn. Nebulized 4% lidocaine can anesthetize the nasal passage, assuming the patient is not mouth breathing. Viscous lidocaine gel can be injected with a syringe into the nasal passage as the patient sniffs; absorption into the mucous membranes is probably enhanced if a nasal airway is placed after application. This has the added benefit of further dilating the nasal passage. Some clinicians use topical cocaine 4% to anesthetize the nose because of excellent tissue penetration.

The maximum dose of lidocaine for airway anesthesia is approximately 4 mg/kg (approximately 250–300 mg in an adult). Sedation for endoscopic intubation can be accomplished with ketamine, etomidate, propofol, fentanyl, alfentanil, or midazolam (see Chapter 5). The goal of sedation is to preserve spontaneous respiration, but limit patient movement and reaction to the procedure. A combination of good topical anesthesia and mild sedation allows the best chance for successful intubation. If excellent topical anesthesia is achieved, parenteral sedation is sometimes unnecessary.

Figure 4.30 This malleable atomizer (LMA MADgic device; Teleflex, Buckinghamshire, UK) allows topical anesthesia to be applied accurately to multiple areas.

Figure 4.31 Attaching a standard swivel adapter (A) to the end of the ET tube during endoscopic intubation through the intubating laryngeal mask airway (ILMA) allows the patient to be ventilated during the entire procedure (B). BVM, Bag-valve-mask; ETT, endotracheal tube; FO, flexible fiberoptic scope; SC, stopcock.
The optimal position of the neck is extension, as opposed to the slight cervical flexion desired when using DL. Extension allows better visualization of the glottis by elevating the epiglottis off the posterior pharyngeal wall. This is especially pertinent in a comatose patient who lacks the muscle tone necessary to maintain an open airway. If problems arise with the tongue and soft tissues falling back and obscuring the view of the endoscope, apply a jaw-lift maneuver or grasp the tongue and pull it forward and away from the soft palate and posterior pharyngeal wall. This also moves the epiglottis away from the posterior pharyngeal wall and facilitates exposure of the cords. Extend the head to accomplish the same objective.

Endoscopic intubation may be performed with the patient in the upright, semi-upright, or supine position. Stand facing the patient or at the patient’s head, depending on personal preference. The upright and semi-upright positions help keep pharyngeal soft tissue from obstructing the airway. The upright position may be more familiar to emergency clinicians who are skilled at diagnostic nasopharyngoscopy.

The greatest impediment to successful endoscopic intubation is an inability to visualize the larynx because blood or secretions have covered the optical element and cannot be removed. Suction actively just before introduction of the endoscope. Application of a small amount of soap to the lens may decrease fogging. If the endoscope has a working channel, suction minor secretions through the suction port during the procedure. The camera can also be cleaned by pressing the lens against a moist mucosal surface or having the patient swallow. Whereas some advocate oxygen insufflation through the suction port to blow away secretions, defog the tip, and increase the inspired oxygen content, there are case reports of gastric insufflation causing gastric rupture, as well as pulmonary barotrauma. If oxygen insufflation is used to clear secretions, it should be used for a few seconds at a time. Supplemental oxygen can be administered separately, through the nose or mouth.

Once the scope has entered the trachea, difficulty advancing the ET tube may be encountered. The tip of the tube most commonly catches on the right arytenoid cartilage or vocal cord. To fix this, withdraw the tube 2 cm, rotate it counterclockwise 90 degrees, and readvance the tube to remedy the problem (see Fig. 4.20).

**Nasal Approach**

The nasal approach is technically easier than the oral approach because the angle of insertion allows better visualization of the larynx with minimal manipulation of the endoscope. Patient cooperation is also less critical with this approach. In an unconscious patient, the tip of the scope is less likely to impinge on the base of the tongue with a nasal approach. In some patients the mouth is not accessible at all because of trismus, swelling, or trauma (e.g., angioedema caused by an angiotensin-converting enzyme [ACE] inhibitor), so the nasal route is the only option for endoscopic intubation (Fig. 4.32).

If conditions permit, choose the most patent nostril. In a cooperative patient, determine this by simply occluding each nostril and asking the patient to identify the nostril that is easiest to breathe through. Identify the most patent nostril by direct vision or by gently inserting a gloved finger that is lubricated with viscous lidocaine into the nostrils. If time is not an issue, an effective method to dilate the nasal cavity and administer an anesthetic is to pass a lidocaine gel–lubricated nasopharyngeal airway (nasal trumpet) into the selected nostril. Leave this airway in place for several minutes, and introduce progressively larger trumpets.

First, place a well-lubricated ET tube in the nostril to a depth of approximately 10 cm before passing the scope through it. Alternatively, mount the ET tube over the scope and first pass the scope through the nostril (Fig. 4.33). Either sequence is acceptable. The advantage of first passing the tracheal tube through the nose is that it avoids the possibility of secretions covering the scope and positions the scope just above the laryngeal inlet. One disadvantage is that it may cause epistaxis, and, in some patients, the tube may not pass easily into the nasopharynx. However, if the tube will not pass into the nasopharynx it is best to discover this at the beginning of the procedure rather than after the trachea has been intubated with the endoscope.

At an insertion depth of 10 cm, the ET tube should have advanced around the bend into the nasopharynx. If negotiating this bend is difficult, place a well-lubricated endoscope through the tube and into the oropharynx to serve as a guide for the ET tube. Once the tracheal tube is in the oropharynx, perform thorough oropharyngeal suctioning before introducing the scope into the ET tube. Advance the endoscope toward the larynx. The epiglottis and vocal cords are seen with little or no manipulation of the tip of the endoscope in 90% of patients. Advance the scope and keep the cords in view by making frequent minor adjustments of the tip of the scope.

In a comatose or obtunded patient, the tongue and other soft tissues may obscure the view of the larynx. This can be alleviated by asking an assistant to pull the tongue forward or to apply a chin- or jaw-lift maneuver. Advance the scope through the larynx to the carina and pass the ET tube over the firmly held endoscope into the trachea. Remember that in adults, the average distance from the naris to the epiglottis is 16 to 17 cm. If the scope has been advanced much beyond this distance and the glottis is still not seen, the scope is probably in the esophagus. If the scope meets resistance at approximately this same level and only a pink blur is visible, the tip of the scope is probably in a piriform sinus. Transillumination of the soft tissues may confirm this and indicate the necessary corrective maneuvers.

**Oral Approach**

Oral endoscopic intubation is indicated when nasal intubation is contraindicated, most commonly because of severe midface trauma or clinician inexperience. Skilled endoscopists often find the oral approach just as easy as the nasal approach. For the less experienced, the oral approach may be more difficult because the path of the scope is less defined by the surrounding soft tissue and the tip of the scope is more likely to impinge on the base of the tongue or vallecula. Keeping the scope in the midline and elevating the soft tissue by pulling the tongue forward or applying the jaw-lift maneuver will minimize this difficulty. Because the oropharyngeal axis is not as well aligned with the larynx as the nasopharyngeal axis, more scope manipulation is required when using the oral approach.

Difficulty with the oral approach can be minimized by using an oral intubating airway (Fig. 4.34). This adjunct resembles an oropharyngeal airway but is longer and has a cylindrical passage through which the endoscope and tracheal tube are passed. The tip of this airway lies just above the laryngeal inlet and ensures midline positioning. Make sure that the patient is either adequately anesthetized or obtunded before the oral airway is placed to minimize gagging or emesis. The ET tube
Figure 4.32 A, This patient with life-threatening angiotensin-converting enzyme inhibitor-induced angioedema is in severe distress, and oral intubation is impossible. This can be a lethal condition. B, Endoscopic nasotracheal intubation is a good choice, but it is very difficult, if not impossible, in a struggling patient. Ketamine anesthesia is ideal, does not depress respirations, and allows easy administration of supplemental oxygen. A less ideal option is blind nasotracheal intubation. A surgical airway may be required. Note: In this case the tracheal tube was premounted on the scope before the scope was passed into the patient’s nose. Alternatively, the tube may be first passed approximately 10 cm through the nose, and then the scope passed through the tube and into the trachea.

Figure 4.33 Flexible endoscopic intubation. Note that the tracheal tube is first premounted on the scope. The fiberoptic scope enters the trachea and then serves as a guide over which the tracheal tube is passed. Larger image, The nasal approach. Inset, Use of an oral intubating airway via the oral approach. (Courtesy Department of Emergency Medicine, Hennepin County Medical Center, Minneapolis.)

Figure 4.34 Examples of oral intubating airways. The Williams Airway Intubator (SunMed, Grand Rapids, MI) (left) cradles the endotracheal (ET) tube in an open, curved guide, whereas the Ovassapian Fiberoptic Intubating Airway (Teleflex, Buckinghamshire, UK) (right) positions the ET tube on the posterior surface of the intubation airway.

can be placed to the tip of the oral airway before the oral airway is inserted, and then the scope passed through the ET tube. Alternatively, place a well-lubricated endoscope, premounted with an ET tube, through the well-positioned oral intubating airway. An assistant holds the oral airway and can make minor adjustments to place the tip of the oral airway just above the laryngeal inlet. This is made easier if an endoscopic system with a video screen is used, so that the intubator and assistant can both observe the camera image. Once the airway structures are visualized, intubate the trachea with the scope (see Fig. 4.33, inset). Advance the ET tube over
the scope and into the trachea. This may require the same counterclockwise maneuver as described with the nasal approach. After successful intubation, the oral airway can be left in place as a bite block or may be removed over the ET tube after removal of the tube adapter. Some oral intubating airways can be removed from the mouth without disconnecting the ET tube adapter.

An alternative, albeit less practical, approach to the traditional oral endoscopic intubation for an anticipated difficult airway requires two clinicians. One performs DL, places the tip of the endoscope under the epiglottis, and blindly advances it while the second clinician, holding the body of the scope, directs the tip of the scope through the cords (Fig. 4.35). Most patients who could be intubated with this technique can probably be intubated using video laryngoscopy with a well-curved bougie.

**Endoscope Technique**

Flexible endoscopic intubation will be more successful if proper technique is maintained. The scope body and endoscope controls are held in one hand while the other hand stabilizes the sheath as it enters the ET tube or patient. The intubator directs motion in one plane with the endoscope controls; to move perpendicular to the plane under direct control the tip of the scope must be rotated. This is best accomplished by keeping the endoscopic sheath taut between the scope body and the point of entry into the patient, and performing rotation of the scope body to affect rotation at the tip of the scope. If the sheath is not held taut, rotation of the scope body will not cause rotation at the tip because of excessive slack in the sheath. The intubator can attempt to rotate the tip by twisting the sheath with the stabilizing hand, but this is technically more difficult and usually less effective. Newer endoscope systems with video monitors make it easier to hold the endoscope and sheath properly (Fig. 4.36).

**Complications**

Complications of endoscopic orotracheal intubation include hypoxia from prolonged intubation attempts, emesis, and laryngospasm. Oxygen saturation monitoring should alert the clinician to hypoxia. Most complications seen with endoscopically guided NT intubation are associated with passing the ET tube through the nasopharynx. Epistaxis is the most common, followed by other nasopharyngeal injuries. A rare but potentially significant complication may result if, on blind advancement of the endoscope through the ET tube, the tip of the scope exits through Murphy’s eye (the distal side port of the ET tube). Attempts at passing the ET tube through the larynx will fail because the tip of the tube, now extending off the midline, will catch on the laryngeal structures.

**Summary**

The primary advantages of endoscopic intubation are the ability to visualize upper airway abnormalities, to negotiate difficult airway anatomy, and to carefully perform tracheal intubation under visual guidance. Endoscopic intubation is noninvasive and well tolerated if meticulous attention is paid to topical anesthesia. Its major limitation is poor visibility
because of blood and secretions. If drying agents and suction cannot mitigate excessive secretions or blood, it is best to avoid endoscopic intubation. Procedural time can be another limitation. Endoscopic intubation requires more practice than many other methods of airway management, and considerable experience should be obtained before using the endoscope in an airway emergency. Endoscopic intubation is more likely to be successful if used early in the management of a difficult airway rather than as a last resort after repeated failure with DL.

OPTICAL AND VIDEO INTUBATING STYLETS

A class of devices that incorporates optics or video into a semirigid metal or metal-reinforced stylet was introduced in the late 1990s. Intubating stylets are different from other video laryngoscopy devices in two important ways. First, they do not create potential space for tube passage like devices with blades. Second, they intubate the trachea directly. Intubating stylets remain popular with some anesthesiologists, but are rarely used in emergency airway management with so many video laryngoscopy devices available.64

Intubating stylets can be used in conjunction with DL or as stand-alone devices. They require laryngoscopic assistance in 8% to 20% of cases and in general are more successful when used in this way.102,103 Visibility can be hampered by blood and secretions, but less so than with flexible endoscopy.104 These devices are more intuitive and require less practice than flexible endoscopy. The Clarus Video System, Shikani Optical Stylet and Levitan FPS Scope (Clarus Medical, Minneapolis) are examples of these malleable optical stylets (Fig. 4.37). The Bonfils Retromolar Intubation Fiberscope (Karl Storz Endoscopy, Tuttingen, Germany) is rigid but otherwise structurally and functionally similar to the Shikani. The distal end of the Bonfils has a fixed curve of 40 degrees, whereas the other scopes are malleable up to 120 degrees. The intubator can look through an eyepiece or a video screen, depending on the model used.

Indications and Contraindications

Semirigid fiberoptic stylets are useful when the glottis cannot be seen readily, and may be a useful adjunct for any difficult airway, especially if mouth opening is limited. A relative contraindication is the presence of significant blood or oral secretions.

Procedure and Technique

The semirigid fiberoptic stylet can be used with or without DL. In either case, place the patient in the sniffing position. Load an ET tube onto a lightly lubricated stylet so that the ET tube extends 1 to 2 cm beyond the end of the stylet. If the device is used alone, create an accentuated curve of between 70 and 80 degrees at the proximal aspect of the cuff of the tube so that it can negotiate the oropharynx. Suction the oropharynx well before attempting intubation. Grasp the mandible with the left hand. Next, raise the jaw to lift the tongue and epiglottis off the posterior hypopharyngeal wall. Ask an assistant to apply a jaw-thrust maneuver or grasp the tongue with gauze and retract it anteriorly. Place the device into the mouth, and while following the curve of the tongue, bring it up under the epiglottis with fiberoptic or video guidance. When the laryngeal inlet is visualized, direct the tip into the larynx. Advance the ET tube as you withdraw the stylet. If resistance is met while advancing the tube, the tip may be catching on the anterior larynx or trachea. Rotate the tube clockwise 120 degrees at the proximal end, which will result in a 90-degree rotation at the tip. With the bevel now anterior, allow the tube to advance without catching. It is important to remember that the direction of rotation of the ET tube is clockwise if resistance is encountered after going through the cords. For resistance encountered before the cords, such as occurs with tracheal tube introducers, fiberoptic scopes, and NT intubation, rotate the tube counterclockwise.

Semirigid optical stylets can be used with DL, either primarily or after encountering difficulty while using the device alone. With this approach, make the angle of the distal stylet less acute, at approximately 35 degrees, and introduce the device only after obtaining maximal visualization with the laryngoscope. If the epiglottis can be seen, advance the tip of the stylet just underneath it via direct vision. Be careful to not embed the tip of the ET tube in the supraglottic soft tissue because it will obscure visibility. At this point, guide the stylet–ET tube unit into the glottis and advance the tube off the stylet.

There have been no reported complications with semirigid fiberoptic stylets other than the failures or prolonged attempts usually related to poor visibility from blood and secretions.102

Summary

Semirigid fiberoptic stylets combine the features of direct and indirect laryngoscopy and provide a valuable tool when approaching a difficult airway. They can be used alone or in conjunction with a laryngoscope.

OROTRACHEAL INTUBATION WITH A KING LARYNGEAL TUBE (LT) OR COMBITUBE IN PLACE

King LT (Ambu, Ballerup, Denmark) airways are used extensively in the prehospital setting because they can be placed quickly and provide reliable oxygenation and ventilation in the vast majority of patients. It is common for patients to arrive to the ED with a King LT (or Combitube [Covidien/
Medtronic, Minneapolis, MN) in place; almost all will need to have it replaced with an ET tube at some point. Intubation through the King LT devices is technically possible with fiberscopes or a bougie, but many who have tried this maneuver in clinical practice have been disappointed. The reason is that the aperture does not necessarily directly align with the glottis, even when the device is ventilating properly. In one study the glottis was visualized in only 51% of patients when a fiberscope was passed through the King LT. The clinician has the choice of whether to remove the functioning retroglottic airway before attempting intubation, or to leave the device in place during intubation. An advantage of leaving the device in place is the maintenance of oxygenation and ventilation until just before the attempt begins, and the ability to reinflate the balloons and resume ventilation immediately after a failed attempt, without having to replace the laryngeal tube.

DL with the King LT in place is difficult. In contrast, video laryngoscopy usually allows excellent visualization of the glottis and placement of an ET tube while the King LT remains in place (with the balloons deflated), and is a reliable method of securing the airway with a laryngeal tube in place. This approach allows providers to avoid the risk of removing a functional airway in a patient who may be difficult to intubate (Fig. 4.38).

To perform this procedure, use a video laryngoscope to obtain a view of the large proximal balloon. Deflate the balloons. The glottis will usually come immediately into view. If it does not, optimize the view using the laryngoscope; small adjustments (rotation or slight retraction) of the laryngeal tube may also be needed. Pass a bougie into the airway, and then have an assistant place an ET tube onto the bougie; intubate the trachea as described previously. Inflate the ET tube and confirm placement. After the tube is known to be in the trachea and patient is stable, remove the laryngeal tube.

**Figure 4.38** Tracheal intubation around a King LT (Ambu, Ballerup, Denmark) airway using video laryngoscopy. A, The laryngoscope is placed in the mouth and advanced until the large balloon comes into view. B, The balloon is deflated. Usually the glottis will immediately appear. If it does not, optimize the view by ensuring the laryngoscope blade is in the vallecula and engaged in the midline hyoepiglottic ligament. C, Pass a bougie into the trachea. D, The tube is passed over the bougie. After confirmation that the endotracheal tube is in the trachea, the King LT airway can be removed. If there is difficulty during the procedure, or if intubation is unsuccessful, inflate the King LT balloons and resume ventilation.

**BLIND NASOTRACHEAL (NT) INTUBATION**

Blind NT intubation can be one of the more technically demanding airway approaches, with the outcome being heavily dependent on the skill and experience of the clinician and a certain amount of luck. The primary advantage of the blind technique is that it minimizes neck movement and does not require mouth opening. In extenuating circumstances, it can be accomplished without an intravenous line.

NT intubation is technically more difficult than oral intubation, and it has definite advantages. Blind NT intubation is possible with the patient in the sitting position, a distinct advantage when intubating a patient with congestive heart failure who cannot tolerate lying flat. In fact, patients in respiratory distress are the easiest to intubate blindly because their air hunger results in increased abduction of the vocal cords, which facilitates entry of the tube into the trachea. Danzl and Thomas reported a success rate of 92% in a large series of ED patients, but success rates are highly dependent on clinician skill, and success is probably significantly lower in current practice because this technique is used less frequently than in decades past.

An NT tube has advantages that extend beyond the immediate difficulties of airway control. The patient cannot bite the tube or manipulate it with the tongue. Oral injuries may be cared for without interference from the tube. An NT tube is more easily stabilized and generally easier to care for than an orotracheal tube. It is better tolerated by the patient, permits easier movement in bed, and produces less reflex salivation than an orotracheal tube.

**Indications and Contraindications**

Patients requiring airway control who have spontaneous respirations can be considered for blind NT intubation. The typical patient is one with an anticipated difficult airway and persistently low oxygen saturation despite preoxygenation. Patients with severe chronic obstructive pulmonary disease (COPD) or asthma who have high airway pressures and may be difficult to ventilate with a face mask are another group to consider for NT intubation. Some other common examples of a difficult airway to consider for NT intubation are patients with short thick necks, trismus, neck immobility, and oral injuries. Other conditions that preclude successful orotracheal intubation include severe arthritis, fixed deformities of the cervical spine, or ACE inhibitor-induced angioedema.

Apnea is the major contraindication to blind NT intubation because attempts to place the tube without respirations as a guide are futile. Avoid nasal intubation in patients with severe nasal or midface trauma. In the presence of a basilar skull fracture, an NT tube may inadvertently enter the cranial cavity. Nasal intubation is also relatively contraindicated if the patient is taking anticoagulants, has a known coagulopathy, or has recently been or soon will be administered thrombolytics. Blind NT intubation should be avoided in patients with expanding neck hematomas and oropharyngeal trauma. If the patient has known abnormal glottic anatomy that would impede blind tube passage, other methods will probably be more successful. Patient combativeness, if not controlled with sedation, makes blind intubation difficult. Inability to open the mouth (such as a wired jaw) is a relative contraindication because emesis may be induced and it may be impossible to clear the vomitus; endoscopic guidance may be preferred to reduce the...
risk of emesis. Exercise judgment in each individual case and be prepared to use neuromuscular blocking agents or to bypass the upper airway with a surgical technique if such a complication develops.

Procedure and Technique

Place the patient in the sniffing position with the proximal part of the neck slightly flexed and the head extended on the neck. Like endoscopic NT intubation, patient preparation is important. Apply phenylephrine drops, oxymetazoline (Afrin) spray, or 4% cocaine spray to both nares to dilate the nasal passages and reduce the risk of epistaxis. Topical anesthesia of the nares, oropharynx, hypopharynx, and larynx with lidocaine spray (4%) is also indicated if time permits (as described previously). If available, cocaine is ideal because it is both a vasoconstrictor and an anesthetic, but caution is necessary in hypertensive patients. Choose the most patent nostril. In a cooperative patient, simply occlude each nostril and ask the patient which naris is easier to breathe through. The most patent nostril can also be identified by direct vision or by gently inserting a gloved finger lubricated with viscous lidocaine into the nares. If time permits, pass a nasal airway first and allow it to remain in place to physically dilate the passage.

After preparation of the nostril, insert a well-lubricated 7.0- or 7.5-mm ET tube along the floor of the nasal cavity. Do not direct the tube cephalad, as one might expect from the external nasal anatomy, but rather direct it straight backward toward the occiput and along the nasal floor. Twist the tube gently to bypass any soft tissue obstruction in the nasal cavity. It is sometimes recommended that the tube’s bevel be oriented toward the septum to avoid injury to the inferior turbinate. At 6 to 7 cm, one usually feels a “give” as the tube passes the nasal choana and negotiates the abrupt 90-degree curve required to enter the nasopharynx. This is the most painful and traumatic part of the procedure and must be done gently. If resistance persists despite continued gentle pressure and twisting of the tube, pass a suction catheter or endoscope down the tube and into the oropharynx to allow successful passage of the tube over the catheter. If this attempt fails, try the other nostril. To avoid this difficulty from the outset, use a controllable-tip tracheal tube (Endotrol, Mallinckrodt Medical, Inc., St. Louis). The tube allows you to increase the flexion of the tube, thereby facilitating passage beyond this tight curve. One study found that the Endotrol tube enhanced first-attempt success with blind NT intubation. A study of paramedic-performed blind NT intubation reported success rates of 58% with standard ET tubes versus 72% with ET tubes with a directional controllable tip.

As the tube advances through the oropharynx and hypopharynx, it approaches the vocal cords, and breath sounds from the tube typically become louder. Foggling of the tube may also occur. At the point of maximal breath sounds, the tube is lying immediately in front of the laryngeal inlet. The tube is most easily advanced into the trachea during inspiration, when the vocal cords are maximally open. As the patient begins to breathe in, advance the tube in one smooth motion. If a cough reflex is present, the patient usually coughs and becomes stridulous during this maneuver, which suggests successful tracheal intubation. The absence of such a response should alert the clinician to probable esophageal passage. Remember that in adults, the average distance from the naris to the epiglottis is 16 to 17 cm; if the scope is advanced past this distance and is not in the trachea, it is probably in the esophagus. If there is a delay in advancing the tube, consider adding oxygen to the end of the tube or providing supplemental oxygen to the mouth or other nares to increase the inspired oxygen concentration. Once the tube is in the trachea, vocalization should cease. Persistent vocalizations suggest esophageal intubation. Breath sounds coming from the tube and tube fogging are other signs of correct ET tube placement. Reflex swallowing during blind NT intubation may direct the tube posteriorly toward the esophagus. If this occurs, direct conscious patients to stick out their tongue to inhibit swallowing and prevent consequent movement of the larynx. Application of laryngeal pressure may also help avoid esophageal passage.

After intubation, auscultate over both lungs while applying positive pressure ventilation. If only one lung is being ventilated, withdraw the tube until breath sounds are heard bilaterally. The optimum distance from the external nares to the tip of the tube is approximately 28 cm in males and 26 cm in females. After verification of tracheal placement, inflate the cuff and secure the tube.

Technical Difficulties

The NT tube may slide smoothly through the hypopharynx and into the trachea on the first pass. Unfortunately, such is not always the case; in an operating room series, the first attempt was successful in less than 50% of instances. When the initial pass is unsuccessful, there are four potential locations of the tip of the tube: anterior to the epiglottis in the vallecula; anterior to the epiglottis in the piriform sinus, or in the esophagus.

Observation and palpation of the soft tissues of the neck during attempted passage of the NT tube are helpful in finding the misplaced tube. Before reattempting placement, withdraw the tube slightly. Do not remove it from the nose because this will create additional trauma to the nasal soft tissues. If cervical spine injury is suspected, avoid any cervical maneuver that moves the neck significantly. To move the ET tube in relation to the laryngeal inlet, it is important to know how the tip of the ET tube responds to head movements. Flexion of the neck will move the tip posteriorly; extension of the neck will move the tip anteriorly; rotation of the neck to the right and left will move the tip of the tube contralateral to the direction of rotation. Based on the suspected location of the tip of the tube, the following methods will help achieve success when difficulties with tube placement are encountered.

Anterior to the Epiglottis

Difficulty advancing the tube beyond 15 cm or palpation of the tip of the tube anteriorly at the level of the hyoid bone suggests an impasse anterior to the epiglottis in the vallecula. Withdraw the tube 2 cm, decrease the degree of neck extension, and readvance the tube.

Arytenoid Cartilage and Vocal Cord

Contrary to the classic teaching, studies have demonstrated a propensity for an NT tube, when placed through the right nares, to lie posteriorly and to the right as it approaches the larynx. It is not surprising that the most common obstacles to advancement of the NT tube are the right arytenoid and the vocal cords. No data are available on common obstacles encountered if the tube is placed in the left nares. If the tube appears to be hanging up on firm cartilaginous tissue, withdraw
An alternative method is to tilt the patient’s head toward the side of the misplacement and then reattempt placement.323

**Esophageal Placement**

Esophageal placement is indicated by a smooth passage of the tube and the loss of breath sounds through the tube. The larynx may be seen or felt to elevate as the tube passes under it. Assisted ventilation will usually produce gurgling sounds when the epigastrium is auscultated. Withdraw the tube until breath sounds are clearly heard, and reattempt passage while applying pressure to the cricoid. Increase extension of the head on the neck during placement. If attempts continue to result in esophageal misplacement, the following maneuver may result in successful tracheal intubation. From the precise point at which breath sounds are lost, withdraw the ET tube 1 cm and inflate the cuff with 15 mL of air, which results in elevation of the tube off the posterior pharyngeal wall, and can help angle the tube toward the larynx. Advance the tube 2 cm. Continued breath sounds indicate a probable intralaryngeal location. At this point, deflate the cuff and advance the ET tube into the trachea (Fig. 4.41). This technique may be particularly useful in patients with cervical spine injury because it requires no manipulation of the head or neck.324 This maneuver, when used on the first pass in 20 patients in the operating room, was successful in 75% of cases.316 This maneuver can also be combined with the passage of a suction catheter after inflation of the ET tube cuff. The suction catheter will often pass easily into the trachea; the cuff can be deflated and the tube advanced into the trachea.

If a controllable-tip ET tube (Endotrol) is used, flex the tip anteriorly to help avoid esophageal placement.313 Remember that the tip is very responsive to pulling on the ring. A common mistake is exerting too much force on the ring, which can result in the tube curling up before the larynx and preventing advancement. There has been a case report of an Endotrol tube “kinking” at the point of sharpest curvature and causing difficulty with suctioning but no problems with ventilation.325 Another device that allows flexion of the distal ET tube is the Parker Flex-Tip stylet (Parker Medical).

**Laryngospasm**

Laryngospasm is a common problem that arises when NT intubation is attempted. It is usually transient. Withdraw the tube slightly and wait for the patient’s first gasp to advance the tube. This is frequently successful because the vocal cords are widely abducted during inhalation. Assess laryngeal anesthesia, and if topical and nebulized lidocaine has already been administered without success, consider transcricothyroid anesthesia (e.g., 2 mL of 4% lidocaine).326 Occasionally, a jaw lift is necessary to break prolonged spasm. Another option is to use a smaller tube.

**Complications**

Epistaxis is the most common complication of blind NT intubation. Severe epistaxis was encountered in only 5 of 300 cases reported by Danzl and Thomas.309 Tintinalli and Claffey327 reported severe bleeding in 1 of 71 patients and less serious bleeding in 12 others. Bleeding is not usually a problem unless it provokes vomiting or aspiration, which is a serious potential problem in obtunded patients with trismus or a decreased gag reflex. Other immediate complications include turbinate fracture, intracranial placement through a basilar skull fracture,
retropharyngeal laceration or dissection, and delayed or unsuccessful placement. Sinusitis in patients with an NT tube is common and can be an unrecognized cause of sepsis. Rare but potentially fatal delayed complications include mediastinitis after retropharyngeal abscess and massive pneumocephalus.

Because most complications occur during tube advancement through the nasal passage and proximal nasopharynx, the complications of blind NT intubation and placement under direct vision are largely the same. Retropharyngeal laceration and esophageal intubation are more of a threat with blind placement techniques because they are more likely to go unrecognized.

Summary

Blind NT intubation is being used less frequently than in the past because clinicians are increasingly becoming comfortable performing oral intubation in patients with potential cervical spine injuries. In addition, emergency physicians frequently use paralytics to facilitate orotracheal intubation. Nevertheless, blind NT intubation remains an effective and potentially lifesaving approach to a difficult airway.

DIGITAL INTUBATION

Digital intubation uses the index and middle fingers to blindly direct the ET tube into the larynx. It is particularly well suited to the prehospital situation, such as when a trapped victim cannot be positioned for intubation. A prehospital series of 66 digitally intubated patients demonstrated an 89% success rate.

Indications and Contraindications

Digital intubation is indicated in a deeply comatose patient whose larynx cannot be visualized and who has a contraindication to NT intubation. Digital intubation could be particularly helpful in situations with poor lighting, abnormal patient positioning (e.g., in an extrication situation), or when standard intubation equipment is unavailable or malfunctioning, and no other approaches are possible. Advantages include speed and ease of placement, immunity to constraints visualizing the larynx, and little neck movement. Contraindications are primarily precautions to protect the clinician. Digital intubation should not be attempted in any patient with a significant risk of biting. This includes calm, awake patients and agitated patients.

Procedure and Technique

Place the patient’s head and neck in the neutral position. Stand at the patient’s right side, facing the patient. Introduce your left index and middle fingers into the right angle of the patient’s mouth and slide them along the surface of the tongue until the epiglottis is felt. The tip of the epiglottis should be palpated 8 to 10 cm from the corner of the mouth in average adults. Use of a stylet in the tube is optional, but the largest reported series had good success without a stylet. For a clinician with short fingers or a patient with an anterior larynx, a stylet is advantageous. If a stylet is used, place it in the tube and bend it into the form of an open J with the distal end terminating in a gentle hook. Introduce a lubricated tube from the patient’s left side between the tongue and the rescuer’s two fingers (Fig. 4.42). Cradle the tube between two fingers and guide the tip beneath the epiglottis. Apply gentle anterior pressure to direct the tube into the larynx. If the clinician has sufficiently long fingers, place them posterior to the arytenoids to act as a “backstop” for the tube, to both avoid esophageal passage and assist in laryngeal placement. If a stylet has been used, withdraw it at this time while simultaneously advancing the tube. An alternative to using a stylet for directing the tube anteriorly is to select an ET tube with a controllable tip (Endotrol, Mallinckrodt Medical, Inc., St. Louis).

A variation of the technique of digital intubation has been described for intubating a newborn. In this technique, only...
Although the most recent experience with digital intubation in adults has been prehospital use, there is no reason why it should be confined to this setting. The majority of moribund ED patients who defy orotracheal intubation are never given a trial of digital intubation. This omission likely deprives some patients of expeditious airway management.

**RETROGRADE INTUBATION**

Retrograde orotracheal intubation is a technique of guided ET intubation that involves the use of a wire or catheter placed percutaneously through the cricothyroid membrane or high trachea and exiting through the mouth or nose. An ET tube is then passed over this guide and advanced through the vocal cords into the trachea. Introduced by Butler and Cirillo in 1960, the technique has undergone several recent modifications that have enhanced its value as a means of establishing a definitive airway when more conventional techniques have failed.

### Indications and Contraindications

Retrograde intubation is indicated when definitive airway control is required and less invasive methods have failed. Indications include trismus, ankylosis of the jaw or cervical spine, upper airway masses or swelling, unstable cervical spine injuries, and maxillofacial trauma. It can be used to convert transtracheal needle ventilation (see Chapter 6) into a definitive airway. It was used successfully in a 1-month-old with developmental abnormalities. It can be particularly helpful in trauma patients with airway bleeding that prevents visualization of the glottis.

Contraindications to retrograde intubation include the availability of a less invasive means of airway control and inability to open the mouth. If rapid control of the airway is needed, consider other alternate strategies including a cricothyrotomy because retrograde intubation takes several minutes to set up and perform. A relative contraindication is an apneic patient who cannot be effectively ventilated with a bag-valve-mask device. In this setting, it is advisable to first establish transtracheal needle ventilation (see Chapter 6) before attempting retrograde intubation or proceed directly to cricothyrotomy.

### Equipment

Materials include (1) local anesthetic and skin preparation material, (2) an 18-gauge needle, (3) a 60-cm epidural catheter needle combination or an 80-cm (0.88-mm diameter) spring guidewire (J tip preferred), (4) a hemostat, (5) long forceps (e.g., Magill) for grasping the wire in the pharynx, (6) an ET tube of appropriate size, (7) a syringe for the tube cuff, and (8) material for securing the tube. A prepackaged alternative is the Cook Retrograde Intubation Set (Cook Critical Care, Bloomington, IN), which also contains a sheath.

### Procedure and Technique

Locate the three important anatomic landmarks by palpation: hyoid bone, thyroid cartilage, and cricoid cartilage. Prepare the skin overlying the cricothyroid membrane and anesthetize it. Maintain a cephalad orientation of the needle bevel, and
result in successful passage through the larynx. When satisfied that the tube has entered the trachea, stabilize the tube and pull the guidewire out through the mouth. Then advance the tube farther into the trachea.

The classic method of retrograde intubation, as just described, has undergone modifications that facilitate passage of the ET tube through the glottis. A significant advance has been the addition of a plastic sheath that is passed antegrade over the wire until it meets resistance where the wire penetrates the laryngeal mucosa (see Fig. 4.42). This sheath needs to be stiff enough to effectively guide an ET tube, yet small enough to easily pass through the vocal cords without impinging on any supraglottic or glottic structures.

If a sheath is available, after grasping the wire from the mouth, thread the plastic sheath over the wire until it comes to rest against the anterior laryngeal wall. Withdraw the wire from the mouth and advance the sheath. Once the sheath is well within the trachea, pass the ET tube over the sheath. If any resistance at the arytenoids or vocal cords is encountered, pull the tube back 1 to 2 cm and rotate it 90 degrees counterclockwise. One advantage of the antegrade sheath is that it lies freely in the larynx, allowing more posterior passage through the widest distance between the cords. In contrast, the wire pulls the ET tube anteriorly toward the narrow commissure of the vocal cords and is more likely to result in impingement of the tube on the cords. In addition, use of the sheath permits unrestricted advancement of the ET tube, but a wire entering the larynx 1.0 to 1.5 cm below the vocal cords prevents the tube from advancing more than this distance before removal of the wire.

If no sheath is available, consider placing the needle inferiorly in the trachea, thereby increasing the distance that the ET tube can be advanced before being stopped by the wire. This will decrease the likelihood of dislodging the tip of the ET tube when the guidewire is withdrawn.

---

**Review Box 4.5** Retrograde intubation: indications, contraindications, complications, and equipment.
Complications

The complications of retrograde intubation are largely related to puncture of the cricothyroid membrane (see Chapter 6). The potential for hemorrhage is minimized by taking care to puncture the cricothyroid membrane in its lower half to avoid the cricothyroid artery. Subcutaneous emphysema may occur, but it is of no clinical significance because no air is insufflated during this technique. A small incidence of soft tissue infection is reported with translaryngeal needle procedures, and ensuring that the wire is withdrawn from the mouth rather than the neck can minimize this problem. The final complication, failure to achieve intubation, has been mitigated by addition of the antegrade sheath over the wire.

Summary

Retrograde intubation is an underused technique for achieving tracheal intubation in a patient who cannot be intubated by less aggressive means. It is more invasive than fiberoptic intubation but requires less skill. Retrograde intubation usually takes several minutes to complete, and the patient can undergo BMV throughout much of the procedure. Recent modifications in the technique guarantee this method a prominent place in
the management of difficult airways, particularly when active bleeding compromises the airway.

**CHANGING TRACHEAL TUBES**

A tracheal tube with a leaking cuff is a vexing problem, especially if the original intubation was difficult. A method of replacing the tube without losing control of the tracheal lumen is preferred. This can be achieved by passing a guide down the defective tube, withdrawing the tube while leaving the guide in place, and introducing a new tube over the guide and into the trachea (Fig. 4.45).

A number of different guides have been described (e.g., simple nasogastric tubes, 18-Fr Salem sump tubes, feeding tubes), and they are poor substitutes for a bougie. Tracheal

---

**Retrograde Intubation**

1. Place a saline-filled needle through the cricothyroid membrane.
   - Orient the needle bevel cephalad.
   - Aspirated air bubbles indicate tracheal entry.

2. Advance a J-wire cephalad through the needle.
   - Grasp the wire in the pharynx and pull it out, long enough to advance the sheath.

3. A) Grasp the wire at the neck with a clamp to keep it taut.
   - B) Pass the antegrade sheath over the wire into the trachea while keeping both ends of the wire taut.

4. Pull the jaw forward, remove the guidewire, and advance the sheath into the trachea.

5. Advance the tracheal tube over the sheath into the trachea, again while pulling the jaw forward.

6. Remove the sheath.

---

**Figure 4.45** Retrograde intubation using a guidewire and antegrade sheath. (Courtesy Department of Emergency Medicine, Hennepin County Medical Center, Minneapolis.)
Preventing Unplanned Extubation

Unplanned extubation is undesirable and can be medically disastrous if unrecognized or if the medical condition requires immediate reintubation. Not all extubations can be prevented. They most often occur when patients suddenly pull out their own tube or during transport. It is not uncommon for a deceased patient to arrive at the medical examiner with a tube in an incorrect position that was dislodged during transit. Unplanned extubation is best prevented with the use of physical and chemical restraints, aggressive sedation of intubated patients, and careful attention to the integrity of the ET tube during procedures, such as positioning for postresuscitation chest radiographs or transfer to another bed.

**CONCLUSION**

Emergency airway management in critically ill or injured patients with acute airway compromise is one of the greatest challenges for clinicians. Although the definition of a difficult airway will change as our ability to visualize the laryngeal inlet continues to improve, the challenge of emergency airways will persist. Mastery of the basics of airway management and DL, competency in a number of guided or indirect intubation techniques, and the ability to perform a surgical airway are all necessary skills for emergency airway management. Mastery of technique, advanced preparation of equipment, and experience in clinical decision-making are essential. Scenario visualization and advanced simulation models can provide an excellent means of practicing the difficult decision making and technical maneuvers necessary for effective emergency airway management.

See Videos 4.1 to 4.21 for airway management procedures using various devices.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Endotracheal (ET) intubation in the emergency setting presents a challenge distinct from that associated with intubation of fasted, premedicated patients in the operating room (OR). Patients in the emergency department (ED) are frequently uncooperative and unstable and may have medical problems or anatomic abnormalities that are completely unknown to the treating clinician. It is challenging that within a matter of minutes and with scant data the clinician must assess and control the airway while diagnosing and managing the patient’s other life-threatening problems.

In 1979, Tarylle and colleagues reported that complications occurred in more than half the patients intubated in a university hospital ED. They called for improved house officer training in ET intubation, including “more liberal use of the procedures and agents used in the OR, including sedatives and muscle relaxers.” Since this report, the use of neuromuscular blockade and agents used in the OR, including “more liberal use of the procedures and agents used in the OR, including sedatives and muscle relaxers.” Since this report, the use of neuromuscular blockade and rapid-sequence intubation (RSI) have become the standard for emergency medicine practice.5,6

In addition to RSI, emergency physicians now use airway devices such as videoscopes and flexible fiberoptic bronchoscopes to manage difficult and complex airways. In high-risk patients it is increasingly common for awake intubation techniques to be utilized. This allows for the safe management of an emergent airway without the risk of completely eliminating the patient’s airway reflexes. A related approach that has been described as delayed-sequence intubation has emerged in recent years. This technique may be considered as procedural sedation to allow preoxygenation prior to intubation. Both awake intubation and delayed-sequence intubation techniques have their own unique pharmacologic considerations. Clinicians must concentrate not only on the manual skills of airway management, but also on selection of the appropriate drugs to achieve specific objectives. These objectives include: (1) immediate airway control, including induction of unconsciousness and muscle paralysis; (2) analgesia and sedation in awake patients; (3) minimization of the adverse physiologic effects of intubation, including systemic and intracranial hypertension; and (4) prevention of harm during the postintubation period, including inadequate sedation, hemodynamic instability, or oversedation.

This chapter reviews the mechanisms and strategic use of the drugs that are currently available to facilitate intubation in the ED.

OVERVIEW OF RSI

The sequential process for quickly intubating a patient in an emergency situation is referred to as rapid-sequence intubation. The steps in performing RSI are often described by the six “P’s”: preparation, preoxygenation, pretreatment, paralysis and induction, placement of the tube, and postintubation management. This sequential technique of rapidly inducing unconsciousness (induction) combined with muscular paralysis to create optimal conditions for intubation has gained broad acceptance among ED clinicians. Many patients do not afford the clinician the time or opportunity to comply with the ideal scenario of tracheal intubation described in this chapter. RSI, as described in this chapter, is the ideal method of emergency airway management for intubations that are not anticipated to be difficult. Consider awake techniques of intubation in high-risk patients with airways that are anticipated to be difficult.

Preparation occurs before and during preoxygenation. Assess the airway to determine the likelihood of a difficult intubation. Simultaneously, establish an intravenous (IV) line and connect the patient to cardiac, pulse oximetry, and end-tidal CO2 monitors when available. Assemble all necessary drugs and equipment for oral intubation and the desired backup equipment for airway management.

Begin RSI preoxygenation as soon as possible by administering 100% oxygen. The intent is to displace nitrogen from the lungs and replace it with an oxygen reserve that will last several minutes. Under optimal conditions, breathing 100% oxygen for 3 minutes has been demonstrated to maintain acceptable oxygen saturation for up to 8 minutes in previously healthy apneic individuals. Another method is to give four vital capacity (maximal) breaths of 100% oxygen from a face mask, which can also maintain acceptable saturation for 6 minutes. Comparable results may not be extrapolated to the ED setting because of differences in the underlying health and cooperation of the patient population.

A recent advance in preoxygenation of patients prior to intubation is the Nasal Oxygenation During Efforts Securing A Tube (NO DESAT) technique. In the NO DESAT technique a nasal cannula is placed underneath the non-rebreather face mask. In awake patients, the nasal cannula can be comfortably set to 5 L/min of oxygen. After induction agents are given, this can be safely increased to 15 L/min of oxygen. The nasal cannula may be left on throughout the attempt to intubate, as it will not interfere with the ability to place an orotracheal tube. This has the potential to prolong the time to desaturation, but clinical studies of this technique’s application in the ED are currently lacking.

Pretreatment consists of the administration of medications to mitigate the potential untoward responses to intubation. Pretreatment during RSI usually occurs 2 to 5 minutes before induction of unconsciousness or muscular paralysis. Although preoxygenation should be maintained for as long as practical before beginning intubation, the ideal situation and circumstances are not always present, and clinical judgment is the deciding factor for this portion of RSI. The clinical utility of routine pretreatment to improve patient-oriented outcomes in ED RSI has been challenged.9

Paralysis and induction involve the induction of a state of unconsciousness with a sedative agent, followed immediately by muscle paralysis. A protocol for ED-based RSI is summarized in Box 5.1.

ET intubation and RSI have also expanded beyond the ED into the prehospital setting. Prehospital RSI protocols use a sedative plus a paralytic for patients not in cardiac arrest, with success rates as high as 92% to 98%.10-14 Without a full complement of medications, prehospital intubation becomes significantly more difficult, and success rates drop to approximately 60%.11 Rates of misplaced ET tubes and complications by
Rapid-Sequence Intubation: The 6 “P’s”

1 Preparation

Preparation occurs prior to and during oxygenation. Assess the airway for difficulty. Establish an IV line, place the patient on the monitors, and assemble all required medications and equipment.

2 Preoxygenation

Begin RSI preoxygenation as soon as possible by placing the patient on 100% oxygen. Oxygen will displace nitrogen from the lungs and provide an oxygen reserve that will last several minutes.

3 Pretreatment

Consider pretreating with medications such as lidocaine, atropine, and/or fentanyl (depending on the clinical scenario) 2 to 3 minutes prior to induction and paralysis.

4 Paralysis and induction

Administer a sedative agent to induce loss of consciousness (induction). After induction, administer a paralytic agent to achieve muscle relaxation, which greatly facilitates intubation.

5 Placement of the tube

Once the patient is sedated and paralyzed, place the endotracheal tube. Intubation techniques are reviewed in Chapter 2.

6 Postintubation management

Confirm proper tube position with an end tidal CO₂ detector, auscultation, and chest radiograph. Assess for adequate tissue oxygenation and response to previously administered drugs.

Figure 5.1 The 6 “P’s” of rapid-sequence intubation. IV, Intravenous; RSI, rapid-sequence intubation.
paramedics may be much higher than previously reported. Studies indicate that outcomes may be worse for patients with traumatic brain injury intubated in the prehospital setting than in the ED. For these reasons many prehospital systems have moved away from the use of RSI.

The technique for proper ET tube placement is discussed in Chapter 4. Postintubation monitoring should assess for proper tube placement, adequate tissue oxygenation, and response to previously administered drugs. After laryngoscopy, ensure ongoing analgesic and anxiolytic therapy.

### Rapid-Sequence Intubation Protocol

1. Preoxygenate (denitrogenate) the lungs by providing 100% oxygen by mask. Apply nasal cannula at high flow at this time if using nasal cannula for apneic oxygenation.
2. Assemble the equipment required:
   - Bag-valve-mask device connected to an oxygen delivery system.
   - Suction with a Yankauer tip.
   - ET tube with an intact cuff, stylet, syringe, and tape.
   - Laryngoscope and blades, in working order.
   - Backup airway equipment.
3. Check to be sure that a functioning, secure intravenous line is in place.
4. Continuously monitor cardiac rhythm and oxygen saturation.
5. Premedicate as appropriate:
   - Fentanyl: 2 to 3 µg/kg given at a rate of 1 to 2 µg/kg per min intravenously.
   - Atropine: 0.01 mg/kg by intravenous push for children or adolescents.
   - Lidocaine: 1.5 to 2 mg/kg intravenously over a period of 30 to 60 seconds.
6. Induce anesthesia with one of the following agents administered intravenously: ketamine, etomidate, fentanyl, midazolam, or propofol.
7. Give succinylcholine, 1.5 mg/kg by intravenous push (use 2 mg/kg for infants and small children) or rocuronium 1.2 mg/kg.
8. Apnea, jaw relaxation, and/or decreased resistance to bag-mask ventilation (use only when oxygenation before rapid-sequence intubation cannot be optimized by spontaneous ventilation) indicates that the patient is sufficiently relaxed to proceed with intubation.
9. Perform ET intubation. If unable to intubate during the first 20-second attempt, stop and ventilate the patient with the bag-mask device for 30 to 60 seconds. Monitor pulse oximetry readings as a guide.
10. Treat bradycardia occurring during intubation with atropine, 0.5 mg by intravenous push (smaller dose for children; see item 5).
11. Once intubation is completed, inflate the cuff and confirm ET tube placement by auscultating for bilateral breath sounds and checking the pulse oximetry and capnography readings. Ultrasound for lung sliding on both sides may be a useful adjunct.
12. Secure the ET tube.

**ET, Endotracheal.**

### Pretreatment: Preventing the Complications of Intubation

Numerous reports have highlighted the physiologic responses to tracheal intubation and attempted to define their immediate or long-term adverse effects and to offer interventions to ameliorate potential organ injury. It is certain that intubation and adjunctive medications have the potential to alter reflexes, intracranial pressure (ICP), blood pressure, and pulse rate, and may induce disturbances in cardiac rhythms, but the actual clinical consequences of these commonly observed changes are largely unknown. Clinical experience suggests that most transient alterations in physiology occurring with ED intubation produce no specific or readily documented long-term sequelae, or are often consequences that cannot easily be monitored or prevented. Prudent clinicians are aware of the potential adverse effects of intubation and are cognizant of potential methods to minimize them. Careful monitoring of the postintubation condition will guide specific interventions.

Overzealous attempts to suppress the physiologic responses that normally accompany airway manipulation may be counterproductive. It would be desirable to provide airway control under the best of circumstances and with the least amount of injury to the patient, but the ideal approach to the physiologic responses to intubation is simply unknown. Most information has been extrapolated from experimental animal models or from the anesthesia experience and similar issues may not apply to the milieu of the ED experience. It is critical to prevent hypotension and hypoxia during intubation, particularly in those with neurologic injury. Patients that theoretically would benefit the most from pretreatment medications are those that may be least able to tolerate any delay in obtaining a definitive airway. The following discussion serves as a general clinical guide to alterations in the physiologic response to intubation.

### The Pressor Response

In addition to the ubiquitous sinus tachycardia, a number of dysrhythmias have been reported after intubation. They are primarily ventricular in origin and include ectopic beats, bigeminy, and short runs of ventricular tachycardia. No studies have established a direct relationship between the response and subsequent clinical deterioration in a large patient population. It is also unclear whether attenuation of the pressor response prevents dysrhythmias or electrocardiographic evidence of ischemia. Ideally, it would be desirable to avoid sudden increases in blood pressure in unstable patients with acute cardiac or vascular disease. Unfortunately, it is unclear if outcomes are improved by attempting to mitigate the pressor response.

Multiple medications have been evaluated to attempt to reduce the pressor response. Lidocaine has been the most extensively evaluated, but it has not been shown to improve outcomes. A dose of 1.5 to 2 mg/kg may slightly reduce the heart rate or blood pressure increase caused by intubation. One small trial demonstrated that nebulized tetracaine reduced increases in heart rate during intubation, but this is not commonly done in emergency practice. Fentanyl dosed at 2 to 5 µg/kg is likely to be more effective at blunting the pressor response than lidocaine. Fentanyl will be discussed in more detail later, in relation to its possible benefit in preventing increases in ICP.
Lidocaine and fentanyl are the drugs with the largest evidence base supporting their use to decrease the pressor response. Other drugs, including thiopental, sodium nitroprusside, labetalol, nitroglycerin, verapamil, nifedipine, clonidine, fentanyl, sufentanil, etomidate, and magnesium, have shown variable responses. In light of the uncertainty of the benefit of any premedication in improving patient outcomes, these other medications cannot be recommended for routine use. In patients at very high risk for harm by transient increases in heart rate or hypertension (such as those experiencing hypertensive emergency or having an active myocardial infarction), it would be reasonable to consider fentanyl or lidocaine. Given the lack of patient-oriented outcomes in the available literature, it is also very reasonable to not premedicate these patients and proceed with intubation without delay.

**Hypotension**

A more pressing concern for the majority of patients intubated in the ED is the importance of avoiding hypotension. Post-intubation hemodynamic instability occurs in over 10% of emergency intubations. Unlike the uncertainty regarding patient harm as a result of transient hypertension, transient hypotension is associated with poor patient outcomes. This is of even higher concern in patients with known or suspected traumatic brain injury.

Avoid hypotension by initiating adequate resuscitation prior to attempts at intubation. If the patient’s clinical condition permits, pursue appropriate volume resuscitation with blood in actively bleeding patients and fluids in hypovolemic patients prior to intubation. Even relatively hemodynamically stable drugs, such as etomidate, may contribute to hypotension in critically ill patients by reducing the endogenous catecholamine response. The physiologic changes that contribute to hypotension following intubation include both the reduced venous return from positive pressure ventilation and the effects of medications given during intubation. It is a challenge to balance the need for airway control and the need to adequately resuscitate prior to intubation.

Once hypotension has occurred during or after intubation, the fastest way to correct this is with the use of vasopressors. If time permits clinically, it is advisable to start vasopressors prior to attempting to intubate if a patient is already hypotensive. Due to the necessary delays in setting up a vasopressor through a pump, there has been recent increased attention to the use of push-dose vasopressors to treat peri-intubation hypotension (Box 5.2 and Fig. 5.2). Further details about how to mix a dilute solution of vasopressor for this purpose will be provided later. It is certainly reasonable to administer temporary vasopressors for this purpose through a peripheral IV line to avoid further delays in intubation to place a central line. Although currently we lack patient-oriented data on whether push-dose pressors improve outcomes in hypotensive patients, the clear harm of even transient hypotension makes the use of these agents a reasonable approach until additional data is generated.

**Intracranial Hypertension**

Stimulation of the respiratory tract by maneuvers such as laryngoscopy, tracheal intubation, and ET suctioning is commonly associated with a brief rise in ICP. The exact mechanism responsible for this rise is unknown. One potential mechanism is the coughing and gagging that frequently follow manipulation of the upper airway and subsequent transmission of intrathoracic pressure to the cerebral circulation. An alternative explanation is the release of catecholamines that accompanies laryngoscopy, which causes a rise in mean arterial pressure and cerebral perfusion pressure. A small rise in ICP has been reported after the administration of succinylcholine. The value of pretreatment with defasciculating doses of neuromuscular blockers (NMBs) to prevent rises in ICP is unknown.

Although the exact significance of a transient rise in ICP is not known, it is possible that it may be detrimental in patients with head trauma or intracranial hypertension. This theoretical harm comes from a possible reduction in cerebral blood flow if a rise in intracranial hypertension is not compensated for by a rise in systemic blood pressure. A number of drugs, including lidocaine, succinylcholine, and the majority of anesthesia induction agents, have been studied to determine whether their use prevents this response. Many of the existing clinical data are not particularly relevant to the ED setting because they are derived from patients in various stages of general anesthesia, during which a wide variety of drug combinations and doses are utilized.

Good evidence suggests that deep general anesthesia prevents the rise in ICP associated with intubation. Depending on the drug used, anesthesia may compromise cardiovascular performance and critically reduce cerebral blood flow. The ideal anesthetic agents to facilitate intubation of patients with acute intracranial pathology may be those that have minimal effects on cardiovascular performance, such as etomidate or fentanyl. Etomidate has been demonstrated to prevent changes in both cerebral perfusion pressure and ICP after tracheal intubation of patients with space-occupying intracranial lesions.

Fentanyl is perhaps the agent with the best evidence supporting its use in prevention of increased ICP during intubation. The dose is 2 to 5 μg/kg, and is higher than dosages typically used for other indications of fentanyl. To be effective, this dose must be given well before intubation, preferably at least 3 minutes prior to intubation.

Ketamine is traditionally contraindicated in patients with head injuries, but this has come into question. Whereas there may be minimal increases in ICP with ketamine, this effect is
offset by a rise in systemic blood pressure. This appears to preserve cerebral blood flow and may in fact be cerebroprotective.41 There are no current studies that have demonstrated poor patient outcomes due to the use of ketamine in critically ill patients.42

**Atropine for Prevention of Bradycardia**

Bradycardia is not infrequent during intubation, especially in young children and neonates. Traditionally, premedication with atropine has been advocated to reduce the incidence of bradycardia.43 A dose of 0.01 mg/kg given intravenously is the standard dose for premedication. While there may be a slight decrease in the incidence of bradycardia with this medication, patient-oriented outcomes have not been proven to benefit.40,43 It is not unreasonable to use atropine only in response to bradycardia, as opposed to using it as a premedication.

**Bronchospasm**

Patients with reactive airway disease may have further bronchospasm during the process of intubation. Lidocaine has traditionally been used to decrease the incidence of bronchospasm in asthma patients who require intubation. A dose of 1.5 to 3 mg/kg given intravenously has been advocated.39 This medication has not been proven to reduce the incidence of bronchospasm, but nebulized albuterol has been shown to be of benefit.44

---

**Figure 5.2** How to make push-dose epinephrine (see Box 5.2).
At the present time, the clinical consequences of intubation-induced physiologic changes are not thoroughly understood. The role of drugs in preventing these changes is equally unclear. Despite this lack of data, it may be intuitively reasonable to attempt to protect patients at theoretical risk. The approach outlined in Box 5.3 is recommended.

**INDUCTION AGENTS**

After premedication, a sedative agent is used to induce loss of consciousness. A number of diverse drugs are available in the ED to induce unconsciousness before intubation, including barbiturates, benzodiazepines, etomidate, ketamine, opiates, and propofol. The choice of a particular induction agent depends on the experience and training of the clinician, the patient’s clinical status, drug characteristics, and institutional protocols (Box 5.4). Considerable evidence indicates that the selected sedative agent influences the quality of intubation conditions and the rapidity of their attainment. These effects persist even when paralytic agents are used. Commonly used drugs and their doses are summarized in Table 5.1.

**Etomidate**

Etomidate is an ultrashort-acting nonbarbiturate hypnotic agent that has been used successfully as an anesthesia induction agent in Europe since the mid-1970s and in the United States since 1983. A significant benefit of etomidate in the emergency setting is its lack of cardio depressant effects. Several case series have now demonstrated its safety and effective use in ED RSI. Extensive experience with this agent now exists in both pediatric and adult patients, and it is an agent of choice for most ED intubations.

Etomidate is a carboxylated imidazole that is both water and lipid soluble. The drug reaches peak brain concentrations within 1 minute of IV infusion and induces unconsciousness within 30 seconds of administration. Its effects last less than 10 minutes after a single bolus dose. Redistribution of the drug is quite rapid, which accounts for its short duration of action. Etomidate is rapidly hydrolyzed in the liver and plasma and forms an inactive metabolite excreted primarily in urine.

---

**BOX 5.3** Sample Protocol for Intubation of a Head-Injured Adult Patient

1. Preoxygenate with 100% O₂ for 2 to 3 minutes.
2. Administer lidocaine, 1.5 to 2 mg/kg intravenously.
3. Sedate with fentanyl, 3 to 5 µg/kg.
4. Induce anesthesia with etomidate, 0.3 mg/kg.
5. Paralyze with succinylcholine, 1.5 mg/kg or rocuronium 1.2 mg/kg.
6. Perform intubation.
7. Maintain postintubation analgesia and sedation.
8. Maintain paralysis if indicated (vecuronium, 0.1 mg/kg).

*The benefit of this traditional protocol is unproved but can be supported if contraindications do not exist.*

**BOX 5.4** Recommendations for Sedation of Patients Undergoing Rapid-Sequence Intubation in Specific Circumstances

The most appropriate medications for sedation before rapid-sequence intubation are based on evaluation of the clinical scenario, and no specific recommendations are appropriate for all circumstances. Different situations, too complex to list here, lend themselves to the use of certain agents (see text). There are no specific standards that must be followed, and the medical literature can be confusing, contradictory, or inadequate. The following conditions and sedation agent recommendations may guide the clinician. Note that appropriate paralytic drugs should also be used.

**HEAD INJURY OR POTENTIALLY ELEVATED INTRACRANIAL PRESSURE**

Pretreatment with the various medications described in the text are appropriate but of unproven value. Adequate cerebral perfusion pressure should be maintained to prevent secondary brain injury. Etomidate is suggested for induction of these patients, but ketamine is also likely to be safe in this population. For hypotensive patients, etomidate or ketamine may be used.

**STATUS EPILEPTICUS**

Midazolam or thiopental may be used for induction. Reduced doses should be used in the unusual circumstance of seizure with hypotension. Midazolam may be used for induction in those with adequate blood pressures. Ketamine and propofol may also be used for induction as they may have some antiepileptic properties.

---

**HEMODYNAMICALLY STABLE PATIENT WITH SEVERE BRONCHOSPASM**

Induction with ketamine or propofol is suggested. Etomidate and midazolam are acceptable alternatives. In hemodynamically unstable patients with severe bronchospasm, ketamine is suggested. Thiopental should not be used in these patients because it provokes release of histamine and can induce or exacerbate bronchospasm.

**PATIENTS WITH CARDIOVASCULAR COMPROMISE**

Etomidate is suggested because of the hemodynamic stability that it provides. If the patient is in shock or severely hypotensive, ketamine and/or etomidate are suggested. If etomidate is used in a patient with sepsis who has associated hypotension refractory to treatment with fluid resuscitation and a vasopressor, a single dose of hydrocortisone (100 mg intravenously) may be given (value unproven).

**FOR AN “AWAKE LOOK” BEFORE INTUBATION OR DELAYED-SEQUENCE INTUBATION**

Ketamine is suggested, but it may not be appropriate when these patients have cardiovascular disease or hypertension.

TABLE 5.1 Recommended Anesthetic Dosing for Rapid-Sequence Intubation and Clinical Considerations

<table>
<thead>
<tr>
<th>DRUG*</th>
<th>DOSE</th>
<th>PREFERRED IN</th>
<th>AVOID IN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thiopental</td>
<td>3–5 mg/kg IV</td>
<td>↑ ICP, SE</td>
<td>Hypotension, RAD</td>
</tr>
<tr>
<td>Methohexital</td>
<td>1–1.5 mg/kg IV</td>
<td>↑ ICP, SE</td>
<td>Hypotension, RAD</td>
</tr>
<tr>
<td>Fentanyl</td>
<td>5–15 µg/kg IV</td>
<td>SE</td>
<td>Hypotension</td>
</tr>
<tr>
<td>Midazolam</td>
<td>0.1–0.3 mg/kg IV</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ketamine</td>
<td>1–2 mg/kg IV</td>
<td>RAD, Hypotension</td>
<td></td>
</tr>
<tr>
<td>Etomidate</td>
<td>0.3 mg/kg IV</td>
<td>Hypotension</td>
<td>Sepsis (unclear)</td>
</tr>
<tr>
<td>Propofol</td>
<td>2 mg/kg IV</td>
<td>↑ ICP, SE</td>
<td>Hypotension</td>
</tr>
</tbody>
</table>

*Any of these can drugs be used before the administration of a neuromuscular blocking agent to induce anesthesia (see text).

ICP, Intracranial pressure; IV, intravenous; RAD, reactive airway disease; SE, status epilepticus.

The recommended dose is 0.3 mg/kg via rapid IV bolus. There is virtually no accumulation of the drug, and anesthesia may be maintained through repeated doses; however, etomidate should not be used as an infusion or in repeated bolus doses for maintenance of sedation after intubation in the ED.

Etomidate acts on the central nervous system (CNS) by stimulating γ-aminobutyric acid (GABA) receptors and depressing the reticular activating system. It produces electroencephalographic changes similar to those produced by barbiturates stimulating the reticular activating system. It produces electroencephalographic changes similar to those produced by barbiturates. Because etomidate has no analgesic activity, it is suggested for induction of patients requiring RSI. The most common immediate side effects of etomidate are pain at the site of injection, nausea, vomiting, and rather common myoclonic jerks. Pain on injection is reported in up to two thirds of patients, although the clinical significance of this transient discomfort immediately prior to intubation is unclear. Use of a large vein, simultaneous saline infusion, and opioid premedication can reduce the discomfort in appropriate situations. Myoclonic activity has been reported in approximately one third of cases and is believed to be caused by disinhibition of subcortical activity rather than CNS stimulation and does not represent seizure activity.

Ketamine

Unique among anesthetic agents, ketamine produces a dissociative anesthesia characterized by excellent analgesia and amnesia despite the appearance of wakefulness. As a drug that is potent and relatively safe with a rapid onset and brief duration of action, ketamine fits the profile of an agent that could be used sometimes dramatic effect can be avoided through the use of NMBs and is rarely seen in the ED, where paralytic agents are regularly used with RSI. No treatment of myoclonus is necessary, and it is of no clinical significance. If persistent, an IV benzodiazepine may be administered. Etomidate need not be avoided in patients with seizure disorders, status epilepticus, head injury, or stroke.

Some degree of altered adrenal function has been demonstrated even after a single dose of etomidate. The true clinical effect is unknown, and because the alteration in adrenal function appears to persist for 12 to 24 hours, there is theoretical concern about potential clinical consequences. Etomidate is a reversible inhibitor of 11β-hydroxylase, the enzyme that converts 11-deoxycortisol to cortisol. Although cortisol levels do not fall below the normal physiologic range, even a single induction dose of etomidate causes a measurable decrease in the level of circulating cortisol that occurs in response to the administration of exogenous adrenocorticotropic hormone (ACTH).

A substudy of the Corticosteroid Therapy of Septic Shock (CORTICUS) trial evaluating the use of hydrocortisone in septic shock provided further evidence about the safety of the use of etomidate in patients with sepsis. There was a statistically significant increase in mortality (42.7% versus 30.5%) for those that received etomidate. Unfortunately, this effect was not diminished by the administration of hydrocortisone. In a separate study that compared etomidate with ketamine, a multicenter randomized trial of critically ill patients requiring emergency intubation found no significant difference in organ failure score, 28-day mortality, or intubating conditions between patients given etomidate for induction and those given ketamine. No serious, drug-related adverse events were reported with either medication. Even though adrenal insufficiency occurred at a higher rate in the etomidate group (86%), it also developed in approximately 48% of patients receiving ketamine. A Cochrane metaanalysis that reviewed eight studies found no conclusive evidence that a single dose of etomidate negatively impacted patient-oriented outcomes.

Clinicians performing postintubation ACTH stimulation testing should be aware that the results may be affected by prior administration of etomidate. The empirical administration of glucocorticoids for the first 24 hours after a dose of etomidate has been given to patients with sepsis is not supported by outcome studies.

The debate over the use of etomidate as an induction agent in those with known sepsis is unlikely to be definitively answered in the near future. Despite numerous studies with conflicting results, there is no clear evidence that patients are harmed by a single dose of this drug. It is not always immediately clear on arrival in the ED if patients who require intubation have sepsis. It is the recommendation of the authors that ketamine be considered in patients requiring intubation who have known sepsis. While the use of etomidate is not a violation of the standard of care, the favorable pharmacologic properties of ketamine make it an ideal induction agent for those with known sepsis.
effectively to facilitate intubation. Ketamine has been advocated for pharmacologic control of undifferentiated agitated and violent patients with excited delirium. It does possess some pharmacologic properties that theoretically limit its use in selected circumstances.

Ketamine is a water- and lipid-soluble drug that rapidly penetrates the CNS. Like the barbiturates, ketamine accumulates rapidly and then undergoes redistribution with subsequent degradation in the liver. The recommended dose of ketamine before intubation is 1 to 2 mg/kg administered intravenously over a 1-minute period. Anesthesia occurs within 1 minute of completing the infusion and lasts 5 to 10 minutes. A smaller additional dose (0.5 to 1 mg/kg) may be given 5 minutes after the initial dose if needed to maintain anesthesia. The intramuscular dose for intubation has not been well studied, and a suggested dose is 4 to 5 mg/kg. Onset of action occurs within 2 to 3 minutes. Because of its good vascularity, the anterior thigh muscle is theoretically the preferred site for administration. Unlike other anesthetic agents that depress the reticular activating system, ketamine acts by interrupting association pathways between the thalamocortical and limbic systems. Characteristically, the eyes remain open and patients exhibit spontaneous, though not purposeful, movements. Increases in blood pressure, heart rate, cardiac output, and myocardial oxygen consumption are seen and are most likely mediated through the CNS. In vitro studies indicate that ketamine is a myocardial depressant, and the CNS-mediated pressor effects generally mask the direct cardiac effects, making it potentially useful in patients with hemorrhagic shock or hypotension. Respirations are initially rapid and shallow after ketamine administration, but they soon return to normal.

Other features of ketamine anesthesia include increased skeletal muscle tone, preservation of the laryngeal and pharyngeal reflexes, hypersalivation, and relaxation of bronchial smooth muscle. Discussions exist regarding the use of ketamine in patients with head injury or potentially elevated ICP. Ketamine can potentially cause a rise in ICP through sympathetic stimulation, theoretically exacerbating the condition of patients with elevated ICP. Recent literature questions the degree to which ketamine raises ICP, with some studies actually showing decreased ICP with the administration of ketamine. A 2009 trial demonstrated the safety of ketamine compared to etomidate in critically ill patients. Ketamine may benefit patients with a neurologic injury by increasing cerebral perfusion.

Other studies suggest that ketamine does not interfere with cerebral metabolism or increase cerebral oxygen consumption and does not reduce regional glucose metabolism. Ketamine can also offset any decrease in mean arterial pressure caused by fentanyl, a drug commonly used as part of RSI in patients with a head injury. For similar reasons, some have advocated the addition of ketamine if propofol is to be used for induction. The resultant “ketofol” mixture is less likely to cause hypotension than propofol monotherapy. A prospective randomized controlled trial is underway to provide additional information about the performance of ketofol as an induction agent.

Overall, evidence suggesting that ketamine elevates ICP to any significant clinical extent is weak, and evidence that harm might ensue is weaker. Based on current evidence it is reasonable to conclude that ketamine is an appropriate induction agent for RSI in patients with suspected ICP elevation and normal blood pressure or hypotension. In patients with hypertension and suspected ICP elevation, ketamine should be used with caution because of its tendency to further increase blood pressure. Patients with severe hypertension may be better served by using etomidate or propofol for induction.

The most promising use of ketamine as an intubation adjunct has been in the setting of acute bronchospastic disease. Ketamine relaxes bronchial smooth muscle either directly through enhancement of sympathomimetic effects or indirectly through inhibition of vagal effects. Ketamine also increases bronchial secretions and may decrease the incidence of mucous plugging that is commonly seen in decompensating asthmatic patients. Clinical reports have demonstrated a reduction in airway resistance and an increase in pulmonary compliance within minutes of ketamine administration. Bronchospastic patients struggling to breathe and unable to tolerate oxygen masks or bronchodilators because of hypoxic encephalopathy will continue to breathe deeply and rapidly with ketamine anesthesia, thereby allowing the maximum delivery of oxygen before a more elective intubation (Fig. 5.3).

A potential side effect that has raised some concern about the use of ketamine for RSI is its tendency to produce postanesthesia emergence reactions, a characteristic that it shares with the structurally similar drug phencyclidine. The reemergence phenomenon, such as disturbing dreams as patients emerge from ketamine-induced anesthesia, is much less of a concern when the drug is used for RSI. In fact, there are no convincing data indicating that, when used for RSI, ketamine produces unpleasant reemergence reactions that are significant or common enough to limit its use for this purpose. One study found that although dreams occurred frequently following sedative doses of ketamine, they were generally pleasant and the frequency of reemergence phenomena and delirium was markedly reduced by the concomitant use of a benzodiazepine. Rarely, reactions may be marked and distressing, with symptoms including floating sensations, dizziness, blurred vision, out-of-body experiences, and vivid dreams or nightmares. The true incidence of emergence reactions following RSI is unknown.

Figure 5.3 This asthmatic is diaphoretic, confused, and agitated and cannot tolerate inhaled bronchodilators. He is about to suffer respiratory arrest and cannot be preoxygenated before intubation. Pulse oximetry shows an oxygen saturation of 82% to 84%. Ketamine is an ideal agent under these circumstances. Keeping him on the stretcher was difficult but within 60 seconds following intravenous administration of ketamine (100 mg), he stopped fighting but kept breathing rapidly. A non-rebreathing oxygen mask was tolerated and oxygen saturation rose to 98%, after which he was electively intubated under controlled preoxygenated conditions. A ketamine infusion (1 mg/kg per hr) was maintained for a few hours.
Propofol does not prolong the QT interval. Although the manufacturer lists egg or soybean allergy as a contraindication to the use of propofol, significant allergic reactions to the newer preparation of the drug are extremely rare. Evidence from 2016 suggests that there is no connection between propofol and egg, soy, or peanut allergy.79

Propofol suppresses sympathetic activity, thereby causing myocardial depression and peripheral vasodilatation, particularly in the elderly or hypovolemic patients and when administered simultaneously with opioids. Hypotension can be minimized with appropriate fluid loading or reversal with pulse dosing IV ephedrine. Propofol reduces cerebral blood flow and may cause mild CNS excitatory activity (e.g., myoclonus, tremors, hiccups) during induction. Propofol has been used in the treatment of status epilepticus,80 but more data is needed on its safety as an induction agent for this purpose. Pain on injection occurs commonly, even when the drug is infused slowly.78 Pretreating the infusing vein with 3 mL of 1% lidocaine (30 mg) injected over a 30-second period or choosing a large antecubital vein will ameliorate this pain. It is unclear if this transient pain is meaningful in patients being induced for intubation due to the likely amnesia caused by drug administration.

Propofol is a reasonable induction agent for RSI in the ED, especially if the patient is hypertensive or in status epilepticus. The safety of etomidate and ketamine are the main challenges to the routine use of propofol for intubation. Whereas propofol has an excellent record for stable patients in the OR and for procedural sedation,40 the common side effect of hypotension precludes its use in many critically ill patients. Its short duration of action and ease of titration make it a more attractive agent for postintubation sedation than as a routine agent for induction.

**Benzodiazepines (Midazolam)**

The benzodiazepines are a class of drugs characterized by anxiolytic, hypnotic, sedative, anticonvulsant, muscle relaxant, and amnestic effects. Several of these properties make the benzodiazepines appealing adjuvant agents for intubation, particularly when used in combination with opioids. It is important to remember that benzodiazepines do not have analgesic effects. Although they may produce excellent sedation and impair the patient's memory of an unpleasant experience, they will not prevent the pain associated with intubation. The high risk of hypotension in critically ill patients who receive both benzodiazepines and opiates limit this drug class's use for induction in the ED.

Midazolam has replaced diazepam as a preoperative sedative agent, even in elderly patients.81,82 When compared with diazepam, the primary advantages of midazolam include a twofold increase in potency, shorter half-life, and decreased potential for cardiorespiratory depression. Midazolam is water soluble in an acid medium and does not require suspension in propylene glycol like other benzodiazepines. It is rarely associated with phlebitis and can be given intramuscularly when a very rapid onset of action is not required. At physiologic pH, midazolam is lipophilic and rapidly accumulates in the CNS, with onset of sedation occurring in as little as 1 to 2 minutes. Outside the CNS it accumulates in fatty tissue and extensively binds to plasma, which accounts for the paucity of non-CNS side effects. Its half-life of elimination is 1 to 4 hours and is dependent on release of the drug from adipose tissue and protein-binding sites. The period of sedation after a single IV

and clinical experience suggests that it is not an issue for RSI in the ED. Such reactions are less common in children than in adults and may be suppressed with benzodiazepines. Both diazepam and lorazepam appear to be useful in adults and the latter is more effective, most likely because of its enhanced amnestic effect. Midazolam is effective in adult patients at doses of 0.07 mg/kg7 and may be the preferred agent because it has potent amnestic effects and a short duration of action. Studies in children have failed to show a reduction in the rate of emergence reactions in those treated with both ketamine and midazolam.71,72

Despite preservation of pharyngeal and laryngeal reflexes in patients sedated with ketamine, aspiration can still occur.73 Ketamine does not relax skeletal muscles, and production of the desired intubating conditions requires the simultaneous administration of a paralytic agent, thereby removing all upper airway reflexes. Despite the possibility of aspiration, the use of ketamine is associated with very few airway complications. A study of over 1000 uses of ketamine for pediatric sedation revealed no cases of aspiration.74

Ketamine’s excellent safety profile and preservation of respiratory drive make it an excellent agent for awake intubations or delayed-sequence intubation.7 This will be discussed further in later sections.

**Propofol**

Propofol is a popular drug among anesthesiologists for OR-based induction and is ideal for ED use. Multiple reports have demonstrated the safety and efficacy of propofol for ED procedural sedation.75,76 Although some ED clinicians now routinely intubate with propofol, its role as an adjunct to intubation in the ED is undergoing evolution.77 Propofol is an alkylphenol sedative-hypnotic used for induction and maintenance of general anesthesia. The drug has no analgesic activity and it exerts a powerful amnestic effect. Propofol tends to decrease vomiting through an unknown antiemetic effect. It produces dose-dependent depression of consciousness ranging from light sedation to coma. Propofol is a highly lipophilic, water-insoluble compound that undergoes rapid uptake by vascular tissues, including the brain, followed soon afterward by redistribution to muscle and fat. The drug is metabolized by the liver and excreted in urine.78 After an induction dose of 2 mg/kg intravenously, unconsciousness occurs within 1 minute and lasts for 5 to 10 minutes. A smaller dose (1 to 1.5 mg/kg) is recommended in the elderly and when simultaneously administering other CNS depressants. Because propofol has a short duration of action and patients rapidly regain consciousness, repeat boluses are not a practical way to maintain a desired level of sedation after intubation has been completed.78 Therefore a slow drip infusion of 50 to 200 µg/kg per min IV titrated to effect is preferred.

Propofol reduces airway resistance, making it a useful induction agent for patients with bronchospasm. Its neuroinhibitory effects make propofol a possible induction agent for patients with intracranial pathology, provided that they are hemodynamically stable. The propofol-induced decrease in mean arterial pressure, generally approximately 10 mm Hg, can reduce cerebral perfusion pressure, thereby theoretically exacerbating CNS injury. As even transient hypotension has been associated with poor outcomes in critically ill patients, propofol should not be used if the patient is hypotensive or at high risk for hypotension.80
The cardiac index and coronary artery blood flow are not generally affected. In the prehospital setting, hypotension with midazolam was found to be dose related, and it should be used cautiously in patients with hypovolemia or traumatic brain injury. Respiratory depression may occur even at standard doses, but it most often follows rapid administration of an excessive dose. Respiratory depression is also more likely to occur in debilitated or elderly patients and in those simultaneously receiving opioids. The effects of midazolam are rapidly reversed by administration of the benzodiazepine antagonist flumazenil. A study of various induction agents for ET intubation suggested that the use of midazolam alone for RSI may be associated with suboptimal intubating conditions and increased difficulty. Other induction agents may be preferred over midazolam alone during RSI.

Whereas once a common agent for induction, midazolam cannot be recommended as a routine induction agent for intubation in the ED. At the doses required for consistent sedation, hypotension is very common. One study showed that 19% of those intubated with the drug had hypotension. The association of hypotension with poor outcomes precludes the routine use of this medication. A possible exception to this is for patients in status epilepticus, and additional research is needed in this area. Midazolam is a common agent for post-intubation sedation, and it is more appropriate for this use than as an induction agent.

**Opioids (Fentanyl)**

Although any of several opioids administered intravenously could be used to induce unconsciousness, fentanyl has significant advantages over other opioid agents. A synthetic opioid, it has the potential for adverse effects with midazolam is similar to that with other benzodiazepines. A small increase in heart rate is seen frequently, as is a small decrease in systolic blood pressure. Changes in blood pressure may be exaggerated in the presence of hypovolemia. An ED-based study reported a mean 10% decrease in systolic blood pressure, with 19% of intubated patients having systolic blood pressure lower than 90 mm Hg. The cardiac index and coronary artery blood flow are not generally affected. In the prehospital setting, hypotension with midazolam was found to be dose related, and it should be used cautiously in patients with hypovolemia or traumatic brain injury. Respiratory depression may occur even at standard doses, but it most often follows rapid administration of an excessive dose. Respiratory depression is also more likely to occur in debilitated or elderly patients and in those simultaneously receiving opioids. The effects of midazolam are rapidly reversed by administration of the benzodiazepine antagonist flumazenil. A study of various induction agents for ET intubation suggested that the use of midazolam alone for RSI may be associated with suboptimal intubating conditions and increased difficulty. Other induction agents may be preferred over midazolam alone during RSI.

Whereas once a common agent for induction, midazolam cannot be recommended as a routine induction agent for intubation in the ED. At the doses required for consistent sedation, hypotension is very common. One study showed that 19% of those intubated with the drug had hypotension. The association of hypotension with poor outcomes precludes the routine use of this medication. A possible exception to this is for patients in status epilepticus, and additional research is needed in this area. Midazolam is a common agent for post-intubation sedation, and it is more appropriate for this use than as an induction agent.

**Opioids (Fentanyl)**

Although any of several opioids administered intravenously could be used to induce unconsciousness, fentanyl has significant advantages over other opioid agents. A synthetic opioid, it has been widely used since its introduction in 1968. Its favorable pharmacologic properties include rapid serum clearance, high potency, and minimal release of histamine. Fentanyl quickly crosses the blood-brain barrier and produces analgesia in as little as 1 to 2 minutes. Serum levels decline rapidly from peak concentrations because of extensive tissue uptake. Unlike morphine, the brain concentration of fentanyl falls in conjunction with the serum level. The duration of analgesic action is 30 to 40 minutes, although at high doses a second peak of activity may be seen several hours later because of release of the bound drug from tissue stores. Fentanyl is approximately 50 to 100 times as potent as morphine. This unique combination of potency and short half-life permits the administration of numerous small doses that can be titrated to the desired clinical effect. Similar to other opioids, fentanyl is competitively reversed with naloxone. Related agents include remifentanil and sufentanil, but these synthetic opioids are not generally available in the ED setting.

The relative safety of fentanyl permits considerable latitude in dosing. When used as a primary anesthetic agent for major surgical procedures, doses ranging from 50 to 100 µg/kg produce minimal side effects. Comparatively tiny doses produce sedation, and 3 to 5 µg/kg, given at a rate of 1 to 2 µg/kg per min, is generally an effective analgesic dose. More rapid administration will cause greater depression of the level of consciousness. Mostert and colleagues reported successful awake intubation in 99 of 103 patients who were administered an average cumulative dose of 3.7 µg/kg. Most of these patients were able to follow commands, and many recalled events surrounding the intubation. A small percentage could not be intubated, even after receiving 500 µg of fentanyl.

Larger doses, perhaps up to 25 µg/kg, may be needed to produce ideal intubating conditions, although if given rapidly, 10 µg/kg is usually adequate. Even this lower dose is more likely to produce unconsciousness than a lesser depth of sedation and it may cause a longer period of unresponsiveness than is desirable. It is preferable to use a low dose of fentanyl (2 to 3 µg/kg) for analgesia combined with a paralytic agent (e.g., succinylcholine) to produce adequate muscle relaxation and a sedative (e.g., midazolam) to reduce anxiety and produce amnesia for the event.

Unlike other opioids, fentanyl causes little or no release of histamine, and its use is seldom associated with emesis or hypotension. It is probably safer than morphine and hydroxypropone in hypovolemic patients. Fentanyl also has significantly fewer emetic effects than other opioids do. Adverse effects that have been reported with fentanyl are few and primarily follow the rapid IV infusion of very large doses. Like other opioids, fentanyl may cause rigidity of the skeletal musculature, including the chest wall and diaphragm. Rigidity occurs with doses in excess of 15 µg/kg, but it has also been reported with doses as low as 10 µg/kg and may also be related to rapid administration. The muscular rigidity may be prevented or treated with standard doses of succinylcholine or naloxone. This rigidity is generally not a major concern during RSI due to the use of paralytics, but may be observed during procedural sedations. The most common significant complication with fentanyl is respiratory depression, and it generally occurs when fentanyl is given in combination with other CNS depressants or in excessive amounts.

Whereas fentanyl appears to be a safe agent for use in the ED, its main limitation as an induction agent is the variable patient response to the drug. Unlike ketamine and etomidate, the effect of a standard dose of fentanyl is unpredictable. This is especially true in patients that are not opioid naive. As mentioned previously, a small percentage of patients did not have adequate sedation despite doses of 500 µg, limiting its use as a sole induction agent. The main use of fentanyl in current practice is as a pretreatment drug to prevent increased ICP or elevated blood pressure.

**NEUROMUSCULAR BLOCKING AGENTS**

NMBs are used to achieve muscle relaxation for intubation. They permit complete airway control and greatly simplify visualization of the vocal cords. This is particularly important when intubation must be performed quickly under less than
ideally circumstances. Sedatives may provide some muscle relaxation, but this requires rapid administration of large doses, which poses a risk for depression of the cardiovascular system. The combination of a paralytic agent and a sedative or analgesic agent is generally superior to the use of either agent alone. A 1999 report showed an 18% failure-to-intubate rate with either agent alone. A 1999 report showed an 18% failure-to-intubate rate with a sedative alone versus a 0% failure rate for sedatives plus paralytics.107 Procedural complication rates such as significant airway trauma and aspiration were also markedly higher in the group receiving sedation alone.

The only absolute contraindication to the use of NMBs is the inability to manage the airway once the patient becomes apneic. Though not absolutely contraindicated, it is considered inhumane to paralyze and intubate an alert patient. A sedative or an analgesic agent should always be administered simultaneously if the patient is able to perceive pain.

NMBs are classified as either depolarizing or nondepolarizing. Depolarizing agents mimic the action of acetylcholine (ACh) and produce sustained depolarization of the neuromuscular junction, during which time muscle contractions cannot occur. Nondepolarizing agents competitively block the action of ACh at the neuromuscular junction and prevent depolarization and therefore muscle contractions. Commonly used NMBs and their dosages and characteristics are listed in Table 5.2.

### Succinylcholine

The standard depolarizing agent currently in use is succinylcholine.102 It has a chemical structure similar to that of ACh and depolarizes the postjunctural neuromuscular membrane. Administration is followed by a brief period of muscle fasciculations that correspond to the initial membrane depolarization and muscle fiber activation. Unlike ACh, which is released in minute amounts and hydrolyzed in milliseconds, succinylcholine requires several minutes for breakdown to occur. During this time the neuromuscular junction remains depolarized, but the muscles relax and will not contract again until the neuromuscular end plate and adjacent sarcoplasmic reticulum return to the resting state and are again depolarized. Relaxation proceeds from the small, distal, rapidly moving muscles to the proximal, slowly moving muscles. The diaphragm is one of the last muscles to relax.

Succinylcholine is rapidly degraded in serum by the enzyme pseudocholinesterase, and the duration of action of a single dose is 5 to 10 minutes.108 Relaxation may be maintained by repeated IV injections. Prolonged or repeated use of the drug enhances its vagal stimulatory effects, thereby resulting in bradycardia, hypotension, and other muscarinic effects. These effects may be seen even at normal doses, particularly in children.109 For this reason, atropine pretreatment at a dose of 0.02 mg/kg has been recommended by some experts for small children and for adults receiving multiple doses, although the need and optimal dose are still in question.110 Repeated dosing with succinylcholine may produce a desensitization blockade and create a scenario in which the neuromuscular membrane returns to the resting state and becomes resistant to further depolarization by succinylcholine.109,110 In general, there is little need for repeated doses of succinylcholine if appropriately dosed the first time. If paralysis in excess of 5 to 10 minutes is desired, longer-acting, nondepolarizing agents such as rocuronium should be used.

The recommended dose of succinylcholine is 1 to 1.5 mg/kg given intravenously. Dosages at the upper end of this range are suggested to guarantee complete relaxation and avoid the need for repeated dosing. Dosage calculations should also be based on actual, not lean body mass because of alterations in both volume of distribution and pseudocholinesterase activity.107 Neonates and infants require a slightly higher dose of succinylcholine (2 mg/kg intravenously) as a result of their higher volume of distribution.107,108 It is crucial that succinylcholine be administered as a rapid bolus because slow administration may lead to incomplete relaxation. Use of a rapid 20-mL to 30-mL saline flush after IV administration may enhance its desired effect.

There are a number of potential adverse effects of the use of succinylcholine, ranging from minor to life-threatening. Muscle fasciculations accompany the initial depolarization of the neuromuscular membrane. Fasciculations are most prominent in muscular patients and create deep, aching muscle pain that may last for days.109 It is unclear whether any regimen will totally prevent the succinylcholine-induced fasciculations (seen in 75% to 100% of patients) and myalgias (seen in 10% to 83% of patients), with varying effects reported after numerous interventions.110 Interestingly, higher doses of succinylcholine may be associated with less myalgia. Traditionally, fasciculations have been prevented by the preadministration of a defasciculating dose (0.01 mg/kg) of pancuronium or vecuronium. The evidence available does not suggest that succinylcholine worsens outcomes in at-risk patients, nor does any evidence suggest that defasciculation improves outcomes in at-risk patients.17

A major consideration in the use of succinylcholine is its propensity to cause hyperkalemia. This electrolyte disturbance is believed to occur secondary to asynchronous depolarization of muscle cells and resulting cellular injury. Elevation in serum potassium occurs in normal patients after standard doses but is typically clinically inconsequential, with increases of less than 0.5 mmol/L (mEq/L).111 Increases in potassium are not prevented with defasciculating doses of nondepolarizing agents. Marked hyperkalemia is associated with increased extrajunctional muscle ACh receptors, which develop in patients with prolonged diseases of the neuromuscular system. Susceptibility may occur within as few as 5 to 7 days and persists indefinitely. In these cases, the hyperkalemic response may be as much as 5 mmol/L (mEq/L). Such conditions include late severe burns,112 major muscle trauma,113 spinal cord injury, muscular dystrophy,
multiple sclerosis, and other upper motor neuron diseases such as amyotrophic lateral sclerosis. These large elevations occur only in patients who have had significant tissue injury or muscle denervation for several days or weeks before the use of succinylcholine. Importantly, succinylcholine is not contraindicated in the initial management of patients with acute injuries, including burns, major crush injuries, and spinal cord injuries. Succinylcholine is also not contraindicated in normokalemic patients with renal failure because the magnitude of the rise in serum potassium is the same as in patients with normally functioning kidneys. A retrospective review of the use of succinylcholine in 38 operative cases with moderate pre-RSI hyperkalemia (5.6 to 7.6 mmol/L) suggested that the risk for hyperkalemia-related complications may be lower than feared. In a review of more than 41,000 intubations, 38 patients had hyperkalemia with a mean serum potassium level of 5.9 mmol/L, so Schow and colleagues concluded that it is safe to administer succinylcholine to patients with a potassium level of 5.5 to 6.0 mEq/L.

Succinylcholine is best avoided (if other equally effective pharmacologic options such as rocuronium exist) in the setting of known or suspected preexisting hyperkalemia (e.g., renal failure patients not receiving regular dialysis or demonstrating a wide QRS complex). There is a higher risk of undiagnosed myopathies in pediatric patients, but succinylcholine remains a safe drug for the majority of pediatric intubations.

Malignant hyperthermia is a rare complication with an autosomal dominant inheritance pattern that is triggered by multiple anesthetic agents, including succinylcholine. Most provocative agents, such as halothane, are not used in the ED setting, and it is extremely rare for the ED physician to encounter this complication. It occurs in approximately 1 in 15,000 children and 1 in 50,000 adults. The clinical syndrome consists of high fever, tachypnea, tachycardia, cardiac arrhythmias, hypoxia, acidosis, myoglobinuria, and impaired coagulation. Unabated muscle contractions mediated by abnormal calcium channels are the physiologic basis for this condition. Treatment includes aggressive cooling measures, volume replacement, and correction of hypoxia as well as acid-base and electrolyte abnormalities. Dantrolene sodium, a direct-acting skeletal muscle relaxant, is thought to be effective in reducing the muscle hypermetabolism that causes the dramatic hyperpyrexia. An associated abnormal response to succinylcholine is isolated masseter spasm, but it can occur in isolation or portend the subsequent development of malignant hyperthermia. Though rare, it has been reported in the emergency medicine literature. In this condition, forcible sustained contraction of the masseter muscles occurs and prevents mouth opening and oral intubation. Management is controversial, with recommendations ranging from use of a bag-valve-mask, securing a surgical airway until the contraction abates, to attempting to suppress the contractions through administration of a nondepolarizing NMB. A case report of an ED patient with masseter spasm during RSI described successful management with the use of a nondepolarizing NMB (vecuronium). Unlike routine cases in the OR, the vast majority of ED patients require definitive airway management and cannot be allowed to simply have the succinylcholine wear off. With the limited evidence available on the management of this rare condition, it is the author’s recommendation that rocuronium be used to treat masseter spasm attributed to succinylcholine.

Prolonged paralysis after the administration of succinylcholine may occur in clinical conditions that result in decreased pseudocholinesterase levels and subsequent decreased metabolism of succinylcholine. Physiologic states associated with this condition include hepatic disease, anemia, renal failure, organophosphate poisoning, pregnancy, chronic cocaine use, advanced age, bronchogenic carcinoma, and connective tissue disorders. Patients with these conditions experience a twofold to threefold increase in the duration of apnea. Patients with cocaine intoxication may also experience prolonged muscle relaxation because cocaine is competitively metabolized by the cholinesterases. An inherited deficiency of pseudocholinesterase is also present in approximately 0.03% of the population and can lead to prolonged paralysis from the administration of succinylcholine. The prolonged duration of paralysis for these patients is likely to be of little clinical significance for the majority of ED patients, as the vast majority of patients intubated in the ED are not ready for extubation shortly afterwards and are unlikely to be harmed by slightly prolonged paralysis.

Succinylcholine can also result in an increase in ICP, but the magnitude and significance of the increase in ICP remains controversial. Several investigators have reported increases in the range of 5 to 10 mm Hg, but other researchers have shown no increase. There is no evidence of neurologic deterioration associated with these transient elevations in ICP. Mechanisms that have been proposed to explain the elevated ICP include: (1) a direct effect of fasciculations, (2) an increase in cortical electrical activity with a resultant increase in cerebral blood flow and blood volume, and (3) sympathetic postganglionic stimulation. Limited studies have been performed to evaluate the significance of this rise in ICP in a brain-injured human patient population. These studies have shown no significant change in electroencephalographic activity or ICP with succinylcholine, but the small size of the studies limits the ability to draw conclusions about clinical outcomes.

Questions concerning the safety of succinylcholine in the setting of acute intracranial pathology do not have clear answers. The drug has been used widely and successfully in this setting and its continued use is supported by this experience. The very real risk for airway compromise and secondary cerebral insult because of hypoxia from delayed or failed intubation must always be weighed against the theoretical harmful effects. Succinylcholine, despite its many potential side effects, is currently the most frequently used agent for neuromuscular blockade with RSI because of its rapid onset and offset and reliable muscle relaxation characteristics.

**Nondepolarizing Agents**

Nondepolarizing agents act in a competitive manner to block the effects of ACh at the neuromuscular junction. Drugs in this class include pancuronium, atracurium, vecuronium, mivacurium, and rocuronium. These drugs, particularly the intermediate-acting agents, have fewer side effects than succinylcholine and they have the potential for reversal. They generally have a longer onset and duration of action than succinylcholine, thus making them less attractive choices for RSI because of the delay in muscle relaxation. In most instances, succinylcholine remains the agent of choice to facilitate emergency intubation, and nondepolarizing agents are indicated to maintain paralysis after intubation. Knowledge of appropriate nondepolarizing NMBs is important for situations when succinylcholine may be contraindicated.
Because nondepolarizing agents act competitively at neuromuscular junction receptors, increasing the concentration of ACh may reverse their effects. Cholinesterase inhibitors such as neostigmine or edrophonium may be used but will not be effective until some spontaneous signs of reversal are seen. The concept of reversal is of limited clinical importance in the ED with rare exceptions, such as performance of a neurologic examination on a previously paralyzed patient. When reversal is required, neostigmine, 0.02 to 0.04 mg/kg, is given by slow IV push. An additional dose of 0.01 to 0.02 mg/kg may be given in 5 minutes if reversal is incomplete, but the total dose should not exceed 5 mg in adults. Atropine, 0.01 mg/kg (with a minimum dose of 0.1 mg in children and a maximum dose of 1 mg in adults), should be given concurrently with neostigmine to block its systemic cholinergic effects.129-131

Another potential reversing agent, sugammadex, binds directly to the aminosteroid NMB rather than to cholinesterases.132 For reversal of shallow or profound neuromuscular blockade, 2 mg/kg or 4 mg/kg intravenously, respectively, is recommended. The most concerning risk with this medication is anaphylaxis.133 Unfortunately, no studies other than case reports currently address its use in emergency situations. It is approved for use in Europe, but was only recently approved for use in the United States and is not widely available yet.

A study by Lee and colleagues134 demonstrated that rocuronium with sugammadex administration leads to faster recovery than succinylcholine after neuromuscular blockade. Whereas the rocuronium/sugammadex arm recovered faster than the succinylcholine arm, sugammadex still took several minutes to reverse many patients. This may be unacceptable in a “cannot-intubate/cannot-oxygenate” situation. A case report of sugammadex use in a cannot-intubate/cannot-oxygenate situation showed that despite partial reversal of paralysis, a tracheostomy was still required.135 The availability of this reversal agent for use in the ED does not change the need to be prepared for a surgical airway if required. The high cost of this medication will also make it cost prohibitive for many EDs.136

**Long-Acting Agents: Pancuronium**

Pancuronium is an aminosteroid that is primarily excreted in urine within 1 hour of IV administration.137 Classified as a long-acting agent, its onset and duration of action are dose related. After a typical IV dose of 0.1 mg/kg, paralysis occurs within 2 to 5 minutes and lasts approximately 60 minutes. Paralysis may be maintained safely by repeated bolus or drip infusion. Because the effects of the drug are cumulative, repeating the original dose significantly lengthens the duration of paralysis.

Relatively few adverse effects are associated with the use of pancuronium. Many patients experience an increase in heart rate, blood pressure, and cardiac output because of the vagolytic effects of the drug. Ventricular tachycardia and severe hypertension have been reported but are quite rare.137,138 Pancuronium may cause release of histamines, resulting in bronchospasm or anaphylactic reactions.139 Prolonged paralysis may occur, primarily in patients with myasthenia gravis or with significant impairment in renal function. One consensus panel recommended pancuronium for maintaining paralysis, except in patients with cardiac disease or hemodynamic instability, for whom they recommended vecuronium.125 This may be appropriate for an ED patient requiring prolonged paralysis.

**Intermediate-Acting Agents: Vecuronium, Atracurium, Mivacurium, and Rocuronium**

Vecuronium and atracurium are intermediate-acting agents with an onset of action of approximately 3 minutes and a duration of action of 30 minutes. Mivacurium has an onset of action of 2 to 3 minutes and a duration of action of 15 to 20 minutes. Rocuronium has an onset of action of 1 to 1.5 minutes and a duration of action of 20 to 75 minutes (longer in geriatric patients). These drugs have minimal cardiovascular effects, cause little release of histamine (with the exception of mivacurium),130 and lack cumulative effects.136

The recommended doses of vecuronium, atracurium, mivacurium, and rocuronium are listed in Table 5.2. Use of larger doses hastens the onset of action but greatly prolongs the period of paralysis. For example, vecuronium at a dose of 0.25 mg/kg intravenously will cause paralysis in as little as 1 minute, but the period of paralysis will last 1 to 3 hours.131,132 Because a rapid onset of action comparable to that of succinylcholine is achieved at high doses of intermediate-acting agents, they may be used as the sole agents to facilitate intubation, particularly if a long period of paralysis is desired after intubation. The paralysis induced by vecuronium or atracurium may be maintained by repeated boluses or drip infusion. Unlike both pancuronium and succinylcholine, these agents have no side effects specifically related to repeat dosing in the ED. A repeated dose of 0.01 to 0.02 mg/kg of vecuronium will extend the period of paralysis by 12 to 15 minutes.

Rocuronium, a structural analogue of vecuronium, is emerging as a desirable alternative agent for RSI when succinylcholine is contraindicated. At doses of 0.6 to 1.2 mg/kg, rocuronium consistently provides good to excellent intubating conditions within 1 minute of administration. Its duration of action is dose dependent and ranges from 20 to 75 minutes.135,136 Smaller anesthesia and ED-based studies have demonstrated its clinical utility and safety in RSI protocols.130,143,145 Rocuronium creates appropriate intubating conditions faster than vecuronium,146 making rocuronium the nondepolarizing agent of choice for RSI in the ED.

A 2015 Cochrane analysis including 37 studies compared rocuronium to succinylcholine for RSI. It revealed slightly superior intubating conditions with succinylcholine when compared to rocuronium. Once limited to studies that only utilized 1.2 mg/kg rocuronium, this difference disappeared. It is the authors’ recommendation that 1.2 mg/kg of rocuronium be used as the routine dose of this medication.

There is insufficient information to clearly demonstrate whether succinylcholine or rocuronium is the superior paralytic for RSI in the ED. Succinylcholine has a much shorter duration, and also has a much longer list of contraindications. EDs frequently receive undifferentiated patients in respiratory failure requiring immediate airway management, often before a full medical history can be obtained or initial potassium level has been assessed. As mentioned previously, it is far from clear how harmful the transient elevation in potassium is for patients requiring RSI.137 Rocuronium has much cleaner pharmacology and a much longer duration of action. It is unclear what role sugammadex will play as a potential reversal agent for rocuronium, or how a patient in respiratory failure would be best managed after initially failing RSI.142 Both agents have their merits and emergency physicians should become comfortable using both agents.
THE “SEDATED LOOK” EVALUATION OF THE AIRWAY BEFORE RSI

In selected stable patients, conditions may exist that preclude the immediate use of RSI, and the more prudent approach would be to assess the airway and intubation needs or potential complications before using paralytics. Examples are patients with angiotensin-converting enzyme inhibitor–induced angioedema or smoke inhalation, where clinical issues of RSI and intubation can be assessed by directly visualizing the larynx. This approach, referred to as a “sedated look” or “awake look,” is used when the clinician suspects a difficult intubation. This approach allows the clinician to verify that the laryngeal structures are indeed visible and accessible before committing to paralysis. This technique allows the patient to maintain respiratory drive during analysis of the airway. This approach is distinct from the practice of intubation with sedation alone or nonparalytic RSI, in which the patient receives a full induction dose of a sedative agent but no neuromuscular blocking agent. This older practice may create a vulnerable and compromised patient in whom intubating conditions are then problematic.

Traditionally in the OR, topical anesthesia (e.g., nebulized 4% lidocaine) along with moderate sedation has been used to allow a view into the airway while enabling the patient to maintain respiratory drive and protect airway reflexes. Although more research is needed to determine which medications are best for sedated looks in the ED, ketamine may be ideal in this circumstance by allowing the patient to maintain respiratory drive while providing analgesia, amnesia, and sedation. Ketamine’s analgesic properties allow it to be used as the sole agent in patients with a bloody traumatized airway, for which topical anesthesia is unlikely to work effectively. Other options are discussed in the following section.

AWAKE INTUBATION

An alternative to induction of unconsciousness in patients requiring intubation is the use of local anesthetic and sedative agents in conscious patients. The availability of relatively effective and safe induction agents makes this a less attractive alternative than in the past, but these techniques may be desirable in specific patients, such as for fiberoptic intubation or nonparalytic RSI, in which the patient receives a full induction dose of a sedative agent but no neuromuscular blocking agent. This approach allows the clinician to verify that the laryngeal structures are indeed visible and accessible before committing to paralysis. This approach allows the clinician to verify that the laryngeal structures are indeed visible and accessible before committing to paralysis. This approach is distinct from the practice of intubation with sedation alone or nonparalytic RSI, in which the patient receives a full induction dose of a sedative agent but no neuromuscular blocking agent. This older practice may create a vulnerable and compromised patient in whom intubating conditions are then problematic.

Additionally, in the OR, topical anesthesia (e.g., nebulized 4% lidocaine) along with moderate sedation has been used to allow a view into the airway while enabling the patient to maintain respiratory drive and protect airway reflexes. Although more research is needed to determine which medications are best for sedated looks in the ED, ketamine may be ideal in this circumstance by allowing the patient to maintain respiratory drive while providing analgesia, amnesia, and sedation. Ketamine’s analgesic properties allow it to be used as the sole agent in patients with a bloody traumatized airway, for which topical anesthesia is unlikely to work effectively. Other options are discussed in the following section.

Local or topical anesthesia techniques may be used in patients who are awake, either in place of or as a supplement to IV analgesia or sedation. They are particularly useful as adjuncts to nasotracheal and fiberoptic intubation but do not generally provide the degree of analgesia or relaxation desirable for traditional laryngoscopy. In addition, the time required to achieve good topical anesthesia may limit the usefulness of these techniques in emergency situations. Topical anesthesia may be achieved by direct application, by cricothyroid membrane puncture, or by inhalation of a nebulized anesthetic.

Direct Application of Topical Anesthetics

Achieving anesthesia of the oral and pharyngeal mucosa is a relatively simple procedure that involves the use of commonly available agents such as 4% lidocaine or a combination such as 14% benzocaine, 2% butamben, and 2% tetracaine (Cetacaine, Cetylite Inc.). Achieving anesthesia of the hypopharynx is more difficult because optimal results require application of the anesthetic to the epiglottis and vocal cords. If time allows, administer an anticholinergic agent, such as glycopyrrolate, at a dose of 0.2 mg IV to help reduce airway secretions. Give this medication 15 minutes prior to intubation for maximal effect. It is important to attempt to dry the airway with suction and gauze, as glycopyrrolate will only reduce additional secretions and will not remove existing moisture. A dry airway will make topical anesthetics more effective.

Begin this procedure by spraying the tongue and pharynx with a topical agent. Atomization devices that attach to standard syringes (e.g., Mucosal Atomization Device [MAD]; Wolfe Tory Medical, Inc., Salt Lake City, UT) can provide effective drug dispersal without a forceful spray (Fig. 5.4). The more forceful pressurized canister sprays commonly provoke a cough reflex. After allowing at least 2 to 3 minutes to achieve numbing of the tongue and pharynx, spray the epiglottis and vocal cords with the MAD device. A malleable extension tube allows the tip of the MAD to pass around the base of the tongue, thereby permitting direct spraying of the epiglottis and vocal cords. This is generally well tolerated. An alternative method is to visualize the epiglottis and vocal cords with a laryngoscope and directly spray them with the anesthetic agent. The use of a laryngoscope to visualize the vocal cords, however, is much more stimulating to the patient and often not well tolerated. Another alternative is percutaneous injection of an anesthetic agent into the trachea at the level of the cricothyroid membrane.

Figure 5.4 The Mucosal Atomization Device. (Courtesy Wolfe Tory Medical, Inc., Salt Lake City, UT)
Parkes and colleagues demonstrated that the plasma levels of lidocaine following nebulized lidocaine for awake intubation were detectable, but far below the toxic threshold. Further evidence of the effectiveness and safety of this procedure has been found through its use in decreasing the pain of placement of nasogastric tubes.

### Sedation for Awake Intubation

Many patients can be intubated while awake with adequate topical anesthesia, but anxiolysis and mild to moderate sedation may be helpful for selected patients. The use of propofol in low doses (0.2 to 0.3 mg/kg) may be helpful, but exercise caution because propofol can cause hypotension in critically ill patients despite using a low dosage.

A new sedative agent, dexmedetomidine (an α₂-adrenoreceptor agonist), has been described for use in awake intubation. Dexmedetomidine produces sedation and anxiolysis with minimal respiratory depression. Patients become sleepy but, if stimulated, can easily be aroused and are generally cooperative. These properties make it seem like an ideal agent for awake intubation, but its use is limited in emergencies by a requisite 10-minute loading dose followed by a maintenance infusion. Future studies will be needed before this medication can be recommended for use in the ED.

The sedative that is most conducive to awake intubation is ketamine. It preserves spontaneous respirations, does not generally cause hypotension, and can safely provide a deep level of sedation. Ketamine's tendency to cause increased airway secretions may be partially corrected for with glycopyrrolate or atropine, but there is rather limited data on the need to use an anticholinergic. Ketamine may be used at subdissociative doses (< 0.3 mg/kg) if adequate topical anesthesia has been obtained. It may alternatively be used at dissociative doses (> 1 mg/kg) but theoretically can cause more side effects. An attempt to appropriately anesthetize the airway is important as ketamine preserves airway reflexes. Ketamine's safety profile makes it a good choice for sedation in awake intubations. The clinician should be prepared for possible laryngospasm during the use of a scope with ketamine. Whereas no cases of this have been reported, preparedness for transition to a surgical airway or traditional RSI is essential.

### Delayed-Sequence Intubation

Some patients may be unable to tolerate adequate preoxygenation prior to an initial attempt to intubate. Delayed-sequence intubation has been described as procedural sedation for the procedure of preoxygenation. Ketamine is given at 1 mg/kg IV to provide sedation. This allows the patient to tolerate preoxygenation with any appropriate combination of high-flow nasal cannula, bag-valve-mask (preferably with positive end-expiratory pressure valve), non-rebreather mask, or noninvasive ventilator. Once adequately preoxygenated, the patient's airway may be managed by intubation with a direct laryngoscope, video laryngoscope, or fiberoptic scope.

### Conclusion

There have been many major advances in the pharmacologic adjuncts to intubation in the ED. There has been an increasing de-emphasis on the importance of premedications prior to intubation. The importance of avoiding hypotension and hypoxia has caused far less focus on the use of lidocaine, fentanyl, or other medications prior to intubation. Ketamine has now joined etomidate as a first-line induction agent, with many

---

**Figure 5.5** Cricothyroid membrane puncture. Prepare the skin with an antiseptic and then puncture the cricothyroid membrane in the midline. Advance the needle until air can be aspirated and then rapidly inject 2 mL of 4% lidocaine. Alternatively, 3 to 4 mL of 2% lidocaine can be used.

**Cricothyroid Membrane Puncture**

Direct application of topical anesthetics to the subglottic region can also be achieved through cricothyroid membrane puncture (Fig. 5.5). In this procedure, identify the cricothyroid membrane immediately below the thyroid cartilage. After antiseptic skin preparation, puncture the overlying tissue and membrane with a 22-gauge needle in the midline and just above the superior border of the cricoid cartilage. Take care to maintain the needle in the midline at all times to avoid injury to the recurrent laryngeal nerves. Advance the needle until air can be aspirated, which indicates placement of the needle in the trachea. Inject a 2-mL volume of 4% lidocaine rapidly. If the 4% concentration is not available, use 3 to 4 mL of 1% to 2% lidocaine. Typically, this will precipitate a cough and distribute the anesthetic over the upper part of the trachea, vocal cords, and epiglottis. Whereas more invasive than the other described techniques, intratracheal injection may provide superior local anesthesia, but there is limited evidence comparing different techniques.

**Nebulized Anesthesia**

Nebulized anesthesia is a simple and painless technique that can be used to facilitate awake intubation when the patient's condition is stable enough to permit a several-minute delay. Deliver the anesthetic via a standard nebulizer and face mask connected to an oxygen source that delivers 4 to 8 L/min. It is critical that the oxygen source not be set to the high-flow level because propofol can cause hypotension in critically ill patients despite using a low dosage.

Deliver the anesthetic via a standard nebulizer and face mask connected to an oxygen source that delivers 4 to 8 L/min. It is critical that the oxygen source not be set to the high-flow level because propofol can cause hypotension in critically ill patients despite using a low dosage.
providers using it almost exclusively. Rocuronium has emerged as the nondepolarizing paralytic of choice in most EDs in recent years. An increased emphasis on awake intubations and the new concept of delayed-sequence intubation has further expanded the use of topical anesthetics and ketamine to assist in difficult airways. A mastery of the pharmacology of drugs used to intubate remains a critical skill for all providers that see critically ill patients.

ACKNOWLEDGMENT

The editors and authors wish to acknowledge the contribution of Laura R. Hopson and Greene Shepherd to this chapter in previous editions.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


C H A P T E R  6

Cricothyrotomy and Percutaneous Translaryngeal Ventilation

Randy B. Hebert and Daniel Thomas

Few clinical scenarios are as critical as when a patient's airway cannot be controlled with traditional endotracheal (ET) intubation. Although cricothyrotomy is rarely required, the incidence of surgical airways has decreased even further since the advent of adjunctive intubation techniques. The conditions accompanying an airway emergency are often stressful and chaotic and require the emergency department (ED) physician to be intimately familiar with this procedure.

When ET intubation has failed or is contraindicated, cricothyrotomy is often the procedure of last resort. Both surgical cricothyrotomy and needle cricothyrotomy entail puncture of the cricothyroid membrane through the overlying skin to gain access to the airway.

Surgical cricothyrotomy is a procedure in which an incision is made in the cricothyroid membrane and a tracheostomy tube or modified ET tube is placed into the airway to ventilate the patient (Video 6.1). Tracheostomy differs from cricothyrotomy in that the incision is made between two of the tracheal rings. Needle cricothyrotomy refers to insertion of a catheter via percutaneous needle puncture of the cricothyroid membrane to allow percutaneous translaryngeal ventilation (PTLV). The term transtracheal jet ventilation is often used interchangeably with PTLV in conjunction with needle cricothyrotomy, but PTLV is more accurate because the cricothyroid membrane is part of the larynx and not the trachea. PTLV is sometimes provided by bag insufflation instead of jet ventilation. The term jet ventilation usually refers to low-frequency jet ventilation with oxygen from a wall source as opposed to high-frequency jet ventilation from a dedicated jet ventilator.

ANATOMY

The central structure of importance is the cricothyroid membrane, an elastic membrane located anteriorly and midline in the neck. The membrane is bordered superiorly by the thyroid cartilage and inferiorly by the cricoid cartilage. The lateral aspects of the cricothyroid membrane are partially covered by the cricothyroid muscles, but the central triangular portion is subcutaneous, which makes it an ideal location to access the airway.

Identify the cricothyroid membrane by locating the prominent thyroid cartilage superior to it. The thyroid cartilage consists of two lateral laminae that join at an acute angle in the midline to form the laryngeal prominence. Commonly known as the Adam's apple, this structure is more pronounced in males. The internal aspect of the anterior body of the thyroid cartilage provides the attachment for the vocal ligaments. Superior to the thyroid cartilage and connecting it to the hyoid bone is the thyroid membrane, which allows passage of the superior laryngeal vessels and the internal branch of the superior laryngeal nerve through its laterally located foramina.

The cricoid cartilage forms the inferior border of the cricothyroid membrane and is the only completely circumferential cartilaginous structure of the larynx. It is composed of a broad posterior segment that tapers laterally to form a narrow anterior arch. The tracheal rings descend inferior to the cricoid cartilage.

Identify the cricothyroid membrane between the previously mentioned structures as a shallow depression measuring approximately 9 mm longitudinally and 30 mm transversely. If the depression is obscured by soft tissue swelling, estimate the location of the cricothyroid membrane at approximately 2 to 3 cm inferior to the laryngeal prominence or four fingerbreadths above the sternal notch.

The area overlying and immediately adjacent to the cricothyroid membrane is relatively avascular and free of other significant anatomic structures. The cricothyroid arteries branch from the superior thyroid arteries and may form a small anastomotic arch traversing the superior aspect of the cricothyroid membrane. The external branch of the superior laryngeal nerve runs along the lateral aspect of the larynx and innervates the cricothyroid muscles inferior to the membrane. The isthmus of the thyroid gland most often overlies the second and third tracheal rings, although an aberrant pyramidal lobe of the gland may extend just superior to the cricothyroid membrane. The anterior attachments of the vocal cord structures are protected by the thyroid cartilage (Fig. 6.1).

In children, the larynx is positioned more superiority than in adults. There is also more overlap between the thyroid cartilage and the cricoid cartilage, thus making the cricoid membrane proportionally smaller (Fig. 6.2).

SURGICAL CRICOTHYROTOMY

Indications and Contraindications

The chief indication for surgical cricothyrotomy is an inability to secure the airway with less invasive techniques in a patient with impending or ongoing hypoxia.

Surgical cricothyrotomy, like any invasive procedure, is associated with significant complications and should not be attempted until less invasive measures have failed. No simple algorithm fits all cases. When time and the clinical situation allow, it may be appropriate to attempt to intubate multiple times with traditional laryngoscopy or to try alternative intubation techniques. Emergency decisions are subject to controversy and differ on a case-by-case analysis, but alternatives to cricothyrotomy include bag-valve-mask ventilation, the gum elastic bougie, and laryngeal mask airways. At some point, further attempts at intubation become futile and the benefits of a surgical airway outweigh the risks associated with ongoing hypoxia.

When approaching a patient with a compromised airway, the clinician must have a clear potential algorithm in mind with a well-defined plan that shifts the airway approach from laryngoscopy to alternative techniques, and then to cricothyrotomy. The first step in deciding whether cricothyrotomy is indicated is anticipating a possible difficult intubation.
Cricothyrotomy

**Indications**
Inability to maintain >90% saturation between intubation attempts or after three attempts
Inability to bag-mask-valve ventilate the patient between intubation attempts or after three attempts
Multiple attempts at endotracheal intubation fail to secure the airway after failed rescue maneuvers (e.g., gum elastic bougie intubation, intubating laryngeal mask airway)

**Contraindications**
Age younger than 5–12 years (depending on the source)
Tracheal transection, fracture, or obstruction below the cricothyroid membrane

**Complications**
_Acute_
- Bleeding
- Tube malposition
- Bronchial intubation
- Laryngotracheal injury
- Tension pneumothorax
- Tube obstruction

_Late_
- Subjective voice changes
- Difficulty swallowing
- Infections
- Persistent shortness of breath
- Persistent stoma
- Subglottic or glottic stenosis

**Equipment**

**Traditional surgical cricothyrotomy**
- Tracheal hook
- Tracheostomy tube
- Tracheal dilator
- Scalpel with a No. 11 blade
- Cuffed 6.0 endotracheal tube

**Melker technique**
- 18-gauge over-the-needle catheter
- Melker airway catheter
- 12-mL syringe
- Scalpel with a No. 11 blade
- Guidewire in plastic housing

Review Box 6.1 Cricothyrotomy: indications, contraindications, complications, and equipment.
CHAPTER 6  Cricothyrotomy and Percutaneous Translaryngeal Ventilation

129

if adequate ventilation and oxygen saturation greater than 90% can be maintained.28,29 Because of the anatomic differences between children and adults, including a smaller cricothyroid membrane and a rostral, funnel-shaped, and more compliant pediatric larynx, surgical cricothyrotomy has been contraindicated in infants and young children. The exact age at which surgical cricothyrotomy can be performed is controversial and not well defined. Various textbooks list the lower age limit from 5 years30 to 10 years31 or 12 years.32 The advanced cardiac life support and pediatric advanced life support define an infant airway as age up to 1 year and a child airway as age 1 to 8 years.

Several studies in the anesthesia and emergency medicine literature have attempted to identify predictors of a difficult airway. A Mallampati score can be determined in cooperative patients who are able to sit upright. It classifies the degree that the faucial pillars, soft palate, and uvula can be visualized (Fig. 6.3). A higher score predicts a more difficult ET intubation.19 A Mallampati score can be obtained only in a limited number of ED patients requiring intubation.20 A modified LEMON (Look externally, Evaluate, Mallampati score, Obstruction, Neck mobility) score, when excluding the Mallampati score, is more easily applied to ED patients for prediction of more difficult ET intubation21 (Fig. 6.4). Additional indicators of a difficult airway include obesity, oropharyngeal edema, hemorrhage, and laryngospasm22–25 (Box 6.1). In anticipation of a failed airway, it may be reasonable to mark the cricothyroid membrane using ultrasound guidance to prepare for the possibility of a cricothyrotomy.26

Cricothyrotomy is indicated when a difficult airway becomes a “failed airway”. Various algorithms have been designed to define a failed airway. The American Society of Anesthesiologists suggests defining a failed airway as an inability to maintain oxygen saturation greater than 90%, signs of inadequate ventilation (cyanosis, absent breath sounds, hemodynamic instability) with positive pressure bag-mask ventilation, or more than three failed attempts at ET intubation or failure to intubate after 10 minutes by an experienced operator.27 As more rescue airway adjunctive devices (such as the laryngeal mask airway, gum elastic bougie, or lighted stylet) become available, it is reasonable to continue beyond three attempts at ET intubation if adequate ventilation and oxygen saturation greater than 90% can be maintained.28,29

Because of the anatomic differences between children and adults, including a smaller cricothyroid membrane and a rostral, funnel-shaped, and more compliant pediatric larynx, surgical cricothyrotomy has been contraindicated in infants and young children. The exact age at which surgical cricothyrotomy can be performed is controversial and not well defined. Various textbooks list the lower age limit from 3 years20 to 10 years21 or 12 years.32 The advanced cardiac life support and pediatric advanced life support define an infant airway as age up to 1 year and a child airway as age 1 to 8 years.
Some authors also identify tracheal transection or low tracheal obstruction (below the cricoid) as absolute contraindications to cricothyrotomy because of the need to secure the airway below the injury.\textsuperscript{31} (Box 6.2).

**Equipment**

The equipment necessary to perform a traditional surgical cricothyrotomy includes a scalpel with a No. 11 blade, a Trousseau dilator, a tracheal hook, and a tracheostomy tube or modified ET tube (see Review Box 6.1). Bent 18-gauge needles may substitute for tracheal hooks. In addition, the sterile tray may include a syringe and lidocaine with epinephrine for local anesthesia, sterile drapes or towels, antiseptic preparation solution, \(4 \times 4\)-cm sterile gauze, scissors, hemostats, and suture material. The average adult’s cricothyroid membrane is approximately 9 mm longitudinally and 30 mm horizontally. Familiarity with the dimensions of several standard tracheostomy and ET tubes is essential when selecting the appropriate size for surgical airways. Cuffed tracheostomy tubes are recommended, and they come in various sizes. Shiley tracheostomy tubes are commonly available in most EDs. The No. 4 tube has an inner diameter (ID) of 5.0 mm and an outer diameter (OD) of 9.4 mm, and the No. 6 tube has an ID of 6.4 mm and an OD of 10.8 mm. Shiley tracheostomy tubes have three parts: a cuffed outer cannula, a removable inner cannula, and a removable obturator that is solid and removed after insertion (Fig. 6.5). ET tubes are often used temporarily in place of a tracheostomy tube. With respect to ID, ET tube OD can vary with the manufacturer. As an example, the Mallinckrodt TaperGuard Evac Endotracheal Tube with IDs of 6.0 and 8.0 mm have ODs of 9.0 and 11.8 mm, respectively.\textsuperscript{33} Although a No. 11 scalpel blade is most commonly used, a No. 20 blade is recommended in some variations of the technique. Commercially available kits include the Melker Cricothyrotomy Kit (Cook Critical Care, Bloomington, IN) for percutaneous cricothyrotomy, which uses the Seldinger technique to insert a cuffed or uncuffed airway catheter.

**Procedure**

Positioning plays a critical role in success, and the ideal patient position may be impossible because of clinical parameters. For example, hypoxic patients often cannot recline. Ketamine anesthesia does not suppress the respiratory drive and may aid in patient cooperation and positioning. Ketamine is theoretically the superior agent to facilitate the procedure. When feasible, use the supine position with the neck exposed. Unless the patient has a known or suspected cervical spine injury, hyperextend the neck to more readily identify the landmarks. Surgical cricothyrotomy can safely and successfully be performed with minimal cervical spine movement.\textsuperscript{34} Preoxygenate the patient by bag-mask ventilation. Prepare the skin of the anterior aspect of the neck with antiseptic solution and create a sterile field with the use of drapes or towels. If the patient is awake or responding to pain, give a subcutaneous and translaryngeal injection of lidocaine with epinephrine as a local anesthetic. Test the integrity of the balloon on the tracheostomy or ET tube by inflating it with 10 mL of air. Wear sterile gloves and take standard precautions by wearing a mask, goggles, and gown. Some preparatory steps may be omitted depending on the urgency of the procedure.

**Traditional Technique**

The “traditional” (open) cricothyrotomy technique (Fig. 6.6) has changed little since the original description of elective cricothyroidotomy by Brantigan and Grow in 1976.\textsuperscript{36} McGill and colleagues\textsuperscript{37} described the addition of a tracheal hook for emergency cricothyrotomy in 1982. In a follow-up report in 1989, Erlandson and colleagues\textsuperscript{38} emphasized the importance of making an initial vertical skin incision and using a relatively small (No. 4 Shiley) tracheal tube. These modifications have generally been accepted and are commonly described as part of the traditional technique.\textsuperscript{39}

Prepare the tracheostomy tube by testing the balloon, removing the inner cannula, and inserting the solid white obturator. If right hand dominant, stand at the bedside on the patient’s right side. Stabilize the larynx with the nondominant hand by grasping both sides of the lateral thyroid cartilage with the thumb and middle finger. Palpate the depression over the cricothyroid membrane with the index finger. Control the larynx throughout the procedure by stabilizing it in this manner (see Fig. 6.6, step 1). In shorter or more obese patients, or in patients with neck swelling, these landmarks may be more difficult to identify by palpation alone.\textsuperscript{40,41} If the laryngeal landmarks are not easily identifiable, bedside ultrasonography...
SURGICAL CRICOXYROTOMY: TRADITIONAL TECHNIQUE

1. Extend the neck whenever possible for better access to the trachea. Immobilize the larynx with your nondominant hand and palpate the cricothyroid membrane with your index finger.

2. Make a 3- to 5-cm vertical midline incision through the skin and subcutaneous tissues. Palpate the membrane through the skin to confirm the anatomy.

3. Make a <1-cm horizontal incision through the cricothyroid membrane. Note that the skin incision is vertical, but the membrane incision is horizontal.

4. Insert the tracheal hook in the opening of the membrane, and rotate it cephalad, while grasping the inferior border of the thyroid cartilage. Ask an assistant to provide upward traction on the hook.

5. Place the tips of the Trousseau dilator into the opening in the membrane and spread in the longitudinal (vertical) plane.

6. Rotate the handle 90° until the handle is vertical or parallel to the neck.

7. Insert the tube between the blades of the dilator until the flanges rest against the skin of the neck. Keep your thumb on the obturator during tube insertion.

8. Carefully remove the Trousseau dilator and the obturator.

9. Replace the inner cannula of the tracheostomy tube and inflate the balloon.

10. Ventilate and confirm tube position by auscultation and end-tidal CO₂. Secure the tube in place.

Figure 6.6 Surgical cricothyrotomy: traditional technique. (Adapted from Custalow CB: Color atlas of emergency department procedures, Philadelphia, 2005, Saunders.)
has been shown to assist in identifying the cricothyroid membrane, as well as decreasing injuries to the larynx and trachea.

While holding the scalpel with a No. 11 blade in the dominant hand, make an approximately 2- to 3-cm vertical incision through the skin and subcutaneous tissue (see Fig. 6.6, step 2). With the index finger of the nondominant hand, palpate the cricothyroid membrane through the incision. It is important to understand that the remainder of the procedure should be performed by palpation of the anatomy, not visualization, because bleeding may obscure the field and there is no time to delay while trying to achieve hemostasis. If the cricothyroid membrane cannot be palpated, extend the initial incision superiorly and inferiorly and try to palpate again. Using the stabilizing nondominant index finger as a guide, make a horizontal incision of less than 1.0 cm in length through the cricothyroid membrane (see Fig. 6.6, step 3). Note that the skin incision is vertical but the membrane incision is horizontal. Place the nondominant index finger into the stoma momentarily and exchange the scalpel for the tracheal hook.

Using the dominant hand, place the tracheal hook into the opening in the cricothyroid membrane. Rotate the handle cephalad while grasping the inferior border of the thyroid cartilage with it. Ask an assistant to provide upward traction or provide traction yourself by passing the handle of the hook to the nondominant hand (see Fig. 6.6, step 4). Use the tracheal hook to stabilize the larynx and keep it in place throughout the remainder of the procedure.

With the dominant hand, place the tips of the Trousseau dilator into the opening in the membrane with the spreading action oriented initially in the longitudinal or vertical plane so that the handle is facing horizontal or perpendicular to the direction of the neck (see Fig. 6.6, step 5). This instrument works opposite that of most ordinary instruments, such as hemostats. Squeezing the handles opens rather than closes the blades. This can be confusing the first time you use this instrument, and it is worth practicing before you need it in an emergency. If this instrument is not available in an emergency, Mayo scissors, a hemostat, or even the blunt end of a scalpel handle can be used to dilate the incision in the cricothyroid membrane.

Dilate the incision vertically with the Trousseau dilator. Hold the handles of the Trousseau dilator with the nondominant hand and rotate the handle 90 degrees until the handle is vertical or parallel to the neck (see Fig. 6.6, step 6). If the dilator is left horizontal, the blades of the dilator may impede passage of the tracheostomy tube into the trachea. While holding the dilator with the nondominant hand, take the tube in the dominant hand and insert it between the blades of the dilator until the flanges rest against the skin of the neck (see Fig. 6.6, step 7). Keep the thumb on the obturator throughout the procedure. Carefully remove the Trousseau dilator (see Fig. 6.6, step 8). Remove the obturator and insert the inner cannula. Inflate the balloon (see Fig. 6.6, step 9). Remove the tracheal hook while being especially careful to not puncture the cuff.

If a tracheostomy tube is not available or if there is difficulty placing the tracheostomy tube into the opening in the cricothyroid membrane, try using a 6-0 cuffed ET tube cut to a shorter length. The ID/OD ratios of tracheostomy tubes are comparable to those of ET tubes. Use of a gum elastic bougie (Video 6.2) may facilitate and even hasten placement of an ET tube through the cricothyroid membrane into the trachea. The advantage of using the bougie is that you can get immediate confirmation that the device is inside the trachea because of the “washboard” vibration that the curved tip makes as it contacts the tracheal rings.

Modify the ET tube by cutting the distal end and replacing the adapter to the cut end (Fig. 6.8). Be careful to not cut the pilot balloon or inflation port. If the ET tube is shortened, it is less likely to kink once it is attached to a ventilator. Advance the ET tube only approximately 5 cm from the tip to avoid main stem intubation. Keep in mind that standard ET tubes do not have centimeter markings at the distal end. Inserting the ET tube so that the distal cuff is approximately 2 cm beyond the cricothyroid membrane usually ensures proper placement.
Confirm proper placement in the same manner as with ET tube placement: end-tidal CO₂, bilateral chest movement, and breath sounds. Secure the tracheostomy tube with a circumferential tie around the neck or with sutures (see Fig. 6.6, step 10). Verify placement with a post-procedure portable chest radiograph.

**Rapid Four-Step Technique (Brofeldt)**
Brofeldt and colleagues developed a rapid four-step technique (RFST) to decrease the amount of time required to establish an airway and reduce complications of hypoxia. It combines aspects of traditional cricothyroidotomy and ET intubation. If right handed, stand at the bedside to the patient’s left. Palpate the depression over the cricothyroid membrane with the nondominant hand (Fig. 6.9, step 1). With the dominant hand, make a single horizontal stab incision with a No. 20 scalpel blade approximately 1.5 cm in length through the skin, subcutaneous tissue, and cricothyroid membrane (see Fig. 6.9, step 2). With the scalpel blade as a guide, pick up the cricoid cartilage with the tracheal hook and provide traction in the caudal direction to stabilize the trachea (see Fig. 6.9, step 3).

**Figure 6.8** A, Modify the standard endotracheal (ET) tube for use in surgical cricothyrotomy as a temporary alternative to a tracheostomy tube. Cut the proximal end of the ET tube but be careful to not cut the balloon inflation apparatus. B, Replace the adapter on the cut end of the tube.

**Figure 6.9** Surgical cricothyrotomy: rapid four-step technique. Extension of the neck (if clinically feasible) facilitates the procedure. (Redrawn from Brofeldt BT, Panacek EA, Richards JR: An easy cricothyrotomy approach: the Rapid Four Step Technique. *Acad Emerg Med* 3:1060, 1996).
Place a No. 4 cuffed tracheostomy tube or a 6-0 cuffed ET tube through the opening (see Fig. 6.9, step 4).

Because this technique omits dilating the stoma with the Trousseau dilator, it may be more difficult to pass a tracheostomy tube. A gum elastic bougie, using the Seldinger technique, may assist in this step.51

Bair and colleagues52 modified this technique further by introducing a new device called a Bair Claw to replace the tracheal hook. The technique is similar to the four-step method except for positioning the operator at the head of the bed instead of the patient’s side and the use of a double-hook device rather than a single hook. By replacing the single hook with the double hook, they found a decrease in the incidence of cricoid ring fractures in cadavers (Fig. 6.10).

This method has been further simplified, using ultrasound to localize the cricothyroid membrane. Then, a single horizontal laceration is made, again through the skin, subcutaneous tissue, and cricothyroid membrane. The scalpel is then turned 90 degrees to allow passage of a gum elastic bougie, over which an ET tube is passed.53

Melker Percutaneous Cricothyrotomy Technique

The Melker Cricothyrotomy Kit (Cook Critical Care, Bloomington, IN) is a prepackaged commercial kit that uses the Seldinger technique to place a tracheostomy tube over a guidewire. The kit comes supplied with a 6-mL syringe, an 18-gauge needle with an overlying tetrtafluoroethylene (TFE) catheter (the TFE catheter is not included in some kits), a guidewire, a tapered dilator, and a Melker airway catheter in lieu of a tracheostomy tube. Similar to retrograde intubation or needle cricothyrotomy with PTLV, the cricothyroid membrane must be easy to identify because no initial skin incision will be made. Anatomic distortion will make locating the cricothyroid membrane with a needle more difficult.

Preparation for this technique is similar to that for the other techniques. Positioned at the patient’s left, palpate the cricothyroid membrane with the nondominant hand. With the dominant hand, attach the needle to the syringe and insert it through the cricothyroid membrane pointing caudally at a 45-degree angle relative to the skin surface (Fig. 6.11, step 1). Be careful not to advance the needle too far because this may result in perforation of the posterior aspect of the trachea. To help recognize when the trachea has been entered, place a small amount of saline in the syringe before the procedure. Apply gentle negative pressure while advancing the syringe. When the membrane is pierced and the trachea is entered, air will be aspirated into the syringe and air bubbles will appear in the saline.

When the needle is in the trachea, pull the syringe and needle back and advance the flexible TFE catheter through the distal end of the trachea to its hub. If the needle does not have an overlying catheter, leave the needle in place and remove the syringe. Thread the guidewire through the needle or the catheter (see Fig. 6.11, step 2). Once the guidewire is placed securely in the trachea, remove the needle or catheter. With a disposable No. 15 scalpel, make a small incision in the skin at the point of guidewire entry to facilitate passage of the dilator and airway catheter (see Fig. 6.11, step 3).

Place the gray-tipped dilator into the airway catheter and thread it over the wire as one unit (see Fig. 6.11, step 4). Once it is through the skin and into the trachea, advance the airway catheter to its hub until it is flush against the neck. Remove the guidewire and dilator. Confirm placement in the trachea by standard methods. Secure the kit in place with “trach tape.”

Melker kits on the market differ with respect to airway catheter ID and whether the airway catheter is cuffed. Some kits do not contain a needle with an overlying catheter.54

Complications

Surgical cricothyotomy is performed infrequently and usually under circumstances that are inherently chaotic. These patients often have confounding medical issues, with high morbidity and mortality rates. Evaluation of short- and long-term complications in this population is difficult.55

Surgical cricothyrotomy has been studied to assess the periprocedure and short-term complications that occur with significant frequency. Acute complication rates have been reported to be between 8.7%56 and 40%.38 The most frequent complications are uncontrollable bleeding and misplacement of the tube.57,58 Most bleeding is from small superficial vessels that can be controlled, but significant bleeding can also occur as a result of the procedure. The cricoid arteries branch from the superior thyroid arteries and anastomose at the anterior superior aspect of the cricothyroid membrane. The laterally running superior thyroid arteries are more often damaged when the initial incision is broad and horizontal. To prevent hemorrhage from these vessels, make the initial skin incision longitudinal as in the traditional technique, and maintain careful awareness of the landmarks.58 When making the horizontal incision in the cricothyroid membrane, avoid the cricoid artery by incising the membrane at its inferior aspect. Misplacement of the tracheostomy or ET tube during cricothyrotomy is a concern, just as esophageal intubation is a complication with ET intubation. If the opening in the cricothyroid membrane is not carefully stabilized during the procedure, the tube may inadvertently be inserted into subcutaneous tissue. This complication can be recognized by the presence of subcutaneous emphysema when attempting to ventilate the patient. It is essential to recognize this immediately to prevent the development of hypoxia and obliteration of anatomic landmarks. Failure to detect end-tidal CO2 and absence of breath sounds by auscultation should alert the physician to a misplaced tube. If suspected, remove the tube and reassess the airway. A misplaced tube can pass into any location other than through the cricothyroid membrane, but the most crucial locations are those that do not enter the airway because this will lead to hypoxia and death if not recognized.

Many other occult complications have been reported less frequently or have been described in case reports, such as main stem bronchial intubation,59 laryngotracheal injury,60 tension pneumothorax,61 and obstruction of the tracheostomy tube...
To decrease the morbidity and mortality associated with prolonged hypoxia and other factors inherent in an airway emergency, attempts have been made to determine whether any technique is superior with regard to complication rate and time needed to secure the airway. When comparing Brofeldt and colleagues' RFST with the traditional five-step technique, Davis and colleagues found an increased incidence of cricoid ring fracture when the single hook was used for caudal traction on the cricoid cartilage and concluded that the traditional technique produced a lower complication rate. A study by Holmes and colleagues in which the same two techniques were performed by inexperienced medical students and residents on human cadavers concluded that the single-hook RFST was executed significantly faster than the traditional technique. They noted that there were more complications with the RFST but that the difference in complication rates failed to reach statistical significance. Davis and colleagues revisited this comparison in a later study and replaced the single hook in the RFST with the double-hooked Bair Claw. The revised study showed that the airway could be secured faster with the

---

**MELKER PERCUTANEOUS CRICOThYROTOMY**

1. Palpate the cricothyroid membrane and advance the needle at a $45^\circ$ angle in a caudal direction. Aspirate on the saline-filled syringe as you advance; air bubbles will enter the syringe when the trachea is entered.

2. Advance the catheter over the needle and then remove the needle. Thread the guidewire through the catheter into the trachea. Once the guidewire is in place, remove the catheter.

3. Make a small incision at the point of guidewire entry to facilitate passage of the dilator and airway catheter.

4. Place the dilator into the airway catheter and thread them over the wire as a unit until it is flush with the skin. Remove the guidewire and dilator, confirm placement, and secure.

**Figure 6.11** Melker percutaneous cricothyrotomy. As with other cricothyrotomy techniques, extension of the neck (if clinically feasible) exposes the trachea and facilitates the procedure.
RFST and that the complication rate was comparable; they also observed that the Bair Claw did not cause any fractures of the cricoid cartilage. Bair and colleagues’ retrospective report of ED cricothyrotomy showed a lower complication rate with the RFST than with the traditional technique.

Consensus cannot be drawn from the literature comparing the traditional method with the percutaneous Seldinger (Melker kit) method. Some studies show no difference in time to ventilation or complication rate when the traditional technique is compared with the Seldinger technique. Some studies report that the surgical method is faster than the Seldinger method, whereas others conclude the opposite.

Many complications of cricothyrotomy are relatively minor in comparison to those caused by prolonged hypoxia. Be aware of these potential complications and prepare for them, and do not delay performing the procedure out of fear of them. Reduce complication rates by maintaining a sterile field and being familiar with the techniques and anatomy.

When deciding which surgical cricothyrotomy technique to use, consider the advantages and disadvantages of each technique, the clinical scenario, availability of equipment, and comfort and familiarity with each individual technique.

Success Rates

Success rates for first-attempt ET intubation in the ED are quite high. The reported success rate is 90% for all ED intubations and 98% for attending physician intubations. The “rescue” cricothyrotomy rate is reported to be just 0.7% according to the National Emergency Airway Registry. A study including more than 6000 trauma patients reported a 0.3% cricothyrotomy rate in those requiring airway management within the first hour after arrival. In pediatric patients, the success rate with the first attempt for all ED intubators is slightly less at 85%, with rescue cricothyrotomy performed less than 1% of the time (1 of 156 patients). As an overview, few emergency physicians have the opportunity to gain extensive experience with surgical airways, and no standards of care have been developed that define the exact role of cricothyrotomy in clinical practice. It is difficult to successfully perform an emergency surgical airway, and even with proper training and standard experience, not all attempts with this technically difficult procedure will be successful. Although the reported success rate for cricothyrotomy has been quite high (89% to 100%) in most studies, one study found only a 62.5% success rate. In a community hospital setting, the rather optimistic success rate reported from trauma centers probably cannot be duplicated. In one ED study, the incidence of failed cricothyrotomy (e.g., tube misplacement into the pretracheal space or failed attempts) was 3.6%, with earlier ED studies being in the 7.9% to 10% range. In the prehospital setting, the reported failure rates are 6% to 12% for paramedics, 0% for physicians, and 0% to 38% for air transport medics. In a cadaver model, the first-time performance of cricothyrotomy by intensive care unit clinicians, versus standard surgical cricothyrotomy with the Seldinger technique, resulted in successful tracheal placement in only 70% with the standard technique and 60% with the Seldinger technique. In an animal model, paramedics had a 90.9% success rate with a percutaneous technique and a 100% success rate with the open surgical technique.

PERCUTANEOUS TRANSLARYNGEAL VENTILATION

PTLV is a procedure in which oxygen is delivered through a 12- to 14-gauge catheter inserted through the cricothyroid membrane via needle cricothyrotomy. Needle cricothyrotomy does not differ greatly from the Seldinger technique variation of surgical cricothyrotomy. Administration of oxygen through the percutaneous translaryngeal catheter is by bag insufflation for pediatric patients younger than 5 years, and by a high-flow source for older children and adults. The methods for controlling jet ventilation have evolved over recent years. The initial use of continuous oxygen flow provided adequate oxygenation but not ventilation. The technique has evolved so that shorter bursts of oxygen are followed by a longer passive exhalation to resemble a more physiologic respiratory state.

Indications and Contraindications

The indications for and contraindications to needle cricothyrotomy with PTLV are similar to those for surgical cricothyrotomy. Indications include failed attempts at ET intubation, inability to bag-mask ventilate to an oxygen saturation greater than 90%, or airway obstruction above the level of the cricothyroid membrane. Based on the operator’s experience, needle cricothyrotomy may be relatively indicated over surgical cricothyrotomy in adult patients. Much of the otolaryngology literature supports the use of PTLV as a means of nonemergency ventilation during head and neck surgery because the smaller ventilation catheter provides a relatively unobstructed field in which to work. In an emergency airway situation, needle cricothyrotomy may successfully allow time to establish an airway via surgical cricothyrotomy or the ET route. Case reports describe PTLV to be relatively indicated over the more invasive surgical cricothyrotomy when ET intubation has failed as a result of copious oropharyngeal secretions. Providing temporary ventilation through the needle catheter may allow...
sufficient time to clear the upper airway of secretions or obstructions and give the operator more time to establish ET intubation.96,97

Surgical cricothyotomy is contraindicated in infants and young children. The contraindication arises from the fact that the cricothyroid membrane is too small to insert a tracheostomy tube and there is a significant risk for injury to surrounding structures. Needle cricothyrotomy is the preferred method of securing the airway in crash airway situations in infants and young children.98

An absolute contraindication to needle cricothyrotomy is the ability to secure the airway without difficulty through less invasive means.99 Similar to surgical cricothyrotomy, needle cricothyrotomy is contraindicated in cases of laryngotraheal transection or fracture because the airway needs to be established below this level.11 It has been suggested that in cases of complete upper airway obstruction, needle cricothyrotomy is relatively contraindicated in comparison to surgical cricothyrotomy. This concern, which is debatable, is due to the fact that the PTLV catheter theoretically does not permit adequate expiration volumes and results in hypercapnia and barotrauma.100

**Equipment**

The essential material needed for PTLV includes a needle with an overlying catheter, oxygen tubing, an oxygen source with a means of regulating the pressure, and a means to connect them together.

Commercial kits are available, but a standard 12- or 14-gauge Angiocath attached to a 3- or 5-mL syringe can be used to make the puncture through the cricothyroid membrane. The catheter can be left in place to serve as the conduit for oxygen delivery. The larger the diameter of the catheter, the greater the oxygen flow, depending on the method of oxygen delivery.101 Commercial catheters, such as wire-coiled nonkinking catheters and fenestrated catheters, are available as part of prearranged kits (Fig. 6.12). Larger-caliber, 3.0 to 4.0 mm-ID percutaneous tracheal catheter devices are also available.

---

**Review Box 6.2** Percutaneous translaryngeal ventilation: indications, contraindications, complications, and equipment.
There are two different basic means and therefore armories of equipment to choose from to deliver oxygen through the transtracheal catheter. One method uses a standard ventilation bag to supply oxygen through the catheter. This requires the constant effort of manual bag insufflation as long as the patient is being oxygenated and ventilated. Attach the bag to the adapter of a 7.0-mm ET tube and insert it into the back of a plungerless 3-mL syringe connected to the trancylaryngeal catheter (Fig. 6.13). Alternatively, attach the bag directly to the catheter with the adapter of a 3.5-mm pediatric ET tube. An inherent problem with this setup is the rigidity of the system. Although the trancylaryngeal catheter itself is flexible, there is no flexibility from the hub of the catheter to the bag. Slight movements of the bag relative to the patient may cause dislodgment of the catheter. To ameliorate this obstacle, connect standard intravenous infusion tubing directly to the trancylaryngeal catheter and attach the distal cut end of a 2.5-mm ET tube to the bag.

In an alternative method, supply oxygen can be supplied from a standard 50 psi wall source, connecting the high-pressure oxygen tubing to the wall source. Then secure the oxygen tubing to a manual on/off valve that is attached to the hub of the catheter. The on/off valve controls the inspiratory-to-expiratory ratio. This valve may be a trigger that is pushed down and released, the third arm of a three-way stopcock open to the atmosphere (Fig. 6.14), or holes placed at the end of the oxygen tubing. A pressure gauge connected to a hand-triggered jet injector may also be used to control the amount of air pressure reaching the catheter (Fig. 6.15). Commercial kits are available that contain prepackaged systems already assembled. Otherwise, assemble the apparatus in the ED. In an emergency, it is unlikely that one would be able to assemble an apparatus for PTLV from individual components in a timely manner. If a prepackaged PTLV kit is not available, prepare the appropriate components from the ED ahead of time and place them with other airway supplies for easy access.

Assemble additional material such as antiseptic preparation solution, sterile drapes, sterile gauze, and suture material or trach tape in the kit.

**Procedure**

As with the surgical cricothyrotomy technique, place the patient in the supine position with the neck exposed. Prepare the skin
of the anterior aspect of the neck. Wear appropriate protective equipment, including sterile gloves, gown, protective eyewear, and a face shield. Hyperextend the patient’s neck unless a suspected cervical spine injury prohibits it. Infiltrate the skin with local anesthetic.

Similar to the needle insertion technique used for guidewire-assisted surgical cricothyotomy, while positioned at the patient’s left, locate the cricoid membrane with the nondominant hand. Locate the thyroid cartilage and cricoid cartilage and palpate the cricoid membrane in the depression between the two while keeping in mind that this depression will be proportionately smaller in children (Fig. 6.16, step 1).

Attach a 12- to 14-gauge Angiocath to a 3- or 5-mL syringe filled with 1 to 2 mL of saline or lidocaine. Once the cricoid membrane has been located, insert the needle through the overlying skin, subcutaneous tissue, and membrane directed at a 30- to 45-degree angle caudally (see Fig. 6.16, step 2). While doing so, aspirate gently with the syringe. The cricoid membrane has been pierced and the airway entered once through step 3. While aspirating, if air bubbles are seen in the fluid or there is an increase in the ease of air aspiration (see Fig. 6.16, step 4 and 5). Hold the catheter by hand until the oxygen supply is connected and appropriate placement is confirmed (see Fig. 6.16, step 6). Make sure that the hub of the catheter is flush against the skin to avoid an air leak and then secure it with a circumferential tie around the neck.

Oxygen can be supplied to the catheter through several different conduits. With the resuscitation bag setup, manually ventilate the lungs through the catheter by squeezing and releasing the bag. When coupled with a 14-gauge Angiocath, ventilation with a resuscitation bag produces low maximal tidal volumes, approximately 100 mL per 1 second of inspiratory time in one study. Children, especially those younger than 5 years, have small total lung capacities and need smaller tidal volumes. In these cases, use the bag instead of the jet ventilator. To use this setup, control the volume of air inspired and adjust it breath by breath based on chest wall motion, pulse oximetry, and capnography. This method is not appropriate for adults because the operator cannot provide adequate tidal volumes and allow enough time for exhalation.

If using high-flow oxygen supplied from a wall source, attach the oxygen tubing to the wall source and secure the distal end of the tubing apparatus to the hub of the translaryngeal catheter. The flow rate recommended for children is 1 L/min per year of age, with titration upward in increments of 1 L/min based on chest wall movement. Most wall-mounted oxygen flowmeters have a maximum flow rate marked at 15 L/min. Pressures generated at this flow rate have been shown to be inadequate to sustain ventilation in adults. To provide the additional pressure necessary to ventilate an adult, open the oxygen flowmeter to full output. Ventilate the patient by alternating between allowing and inhibiting airflow through the catheter. This is done by occluding and then releasing the hole or holes if using a stopcock or the ENK oxygen flow modulator (Cook Medical, Bloomington, IN) or by pushing and releasing a trigger if using this type of modulator. Watch for chest wall rise and fall, and make sure to allow enough time for exhalation before the next cycle. A pressure gauge, if attached, can help guide inspiration time.

Complications

Complications reported with needle cricothyotomy are similar to those associated with surgical cricothyotomy: bleeding, misplacement of the catheter, subcutaneous emphysema, and pneumothorax. Complications more specific to needle cricothyotomy include catheter kinking and perforation of the posterior aspect of the trachea.

One would assume that a translaryngeal placed catheter would not afford any airway protection against aspiration because the diameter of the catheter is not nearly large enough to occlude the lumen of the trachea. A few studies, though, have shown a decreased rate of aspiration in dogs that were ventilated with PTLV versus control animals who were not ventilated, thus suggesting some airway protection with this mode of ventilation.

The potential complications more specific to PTLV than to ventilation through a tracheostomy tube stem from the idea that, especially in cases of complete upper airway obstruction, egress of inspired gas is limited through the relatively small translaryngeal catheter. It has been reported that PTLV inevitably causes retention of CO₂ in adults. This leads to poor ventilation despite adequate oxygenation. This assumption may be a remnant of earlier oxygenation techniques in which continuous low-flow “apneic oxygenation” was used without ventilation. Many animal studies have shown that adequate ventilation, normal blood pH, and normal arterial CO₂ partial pressure can be maintained with PTLV for 30 or even 60 minutes. Factors that seem to improve ventilation are increased expiratory time and a high-flow oxygen source. Even with partial or nearly complete oropharyngeal obstruction, adequate ventilation has been achieved. Unfortunately, none of these studies looked at ventilation for extended periods.

Barotrauma is a significant risk associated with PTLV and occurs when upper airway obstruction is preventing air from being exhaled. This causes an increase in lung volume and pressure and leads to lung injury. Lenfant and colleagues found that, using a lower respiratory rate and the ENK oxygen flow modulator versus the Manujet (Mainline Medical, Inc.,
Norcross, GA), decreases pulmonary pressure, which theoretically reduces the likelihood of barotrauma. The ENK oxygen flow modulator may also be superior to a three-way stopcock for similar reasons. The Ventrain device (Armstrong Medical Ltd, Coleraine, Northern Ireland), which allows more rapid expiration, has also been shown to have improved minute ventilation, peak airway pressures, and decreased hemodynamic depression compared to previous methods when studied in animal models. Use of a bidirectional valve or an expiratory ventilation assistance ejector device has been shown to improve ventilation dynamics and decrease the complications associated with PTLV in patients with complete upper airway obstruction (Box 6.4).

**CONCLUSION**

The majority of crash airway scenarios are controlled successfully with ET intubation. The development of airway aids such as the gum elastic bougie, laryngeal mask airway, and
fibroscopic laryngoscope has obviated the need to convert efforts to an invasive surgical technique. Situations arise when an airway cannot be secured with one of these aids or the patient cannot be ventilated via bag-mask-valve ventilation, and then a surgical approach is indicated.

Researchers have attempted to delineate which method of gaining emergency airway access is superior by comparing all techniques across the spectrum from traditional surgical cricothyrotomy to needle cricothyrotomy to commercially available kits such as the QuickTrach (Teleflex Medical, Morrisville, NC). Given the variety of equipment and techniques to choose from and the advantages and disadvantages reported about each throughout the literature, it would be difficult for an inexperienced physician to know which to choose in a critical situation. As most physicians do not perform this procedure during residency on a living patient, and only approximately 68% do so on a recently deceased patient, familiarity with equipment is paramount. Aside from being familiar with the equipment available in the ED, research shows that physicians at all levels of training have increased success rates when they are exposed to simulated situations before an actual airway emergency.

**REFERENCES**

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Watson PG: Percutaneous transtracheal jet ventilation: a safe, quick, and temporary way to provide oxygenation and ventilation when conventional methods are unsuccessful. Chest 116:1689, 1999.


INTRODUCTION

Placement of a tracheostomy tube is a common procedure in critically ill patients. Common indications for this procedure include upper airway obstruction, head or neck trauma, and prolonged respiratory failure. Approximately one fourth of patients in the intensive care unit (ICU) will require a tracheostomy tube for prolonged respiratory support or weaning from mechanical ventilation. Advances in health care allow many patients with tracheostomies to live at home or in other relatively low-technology environments such as rehabilitation facilities and nursing homes. Tracheostomy care is often provided by a variety of caregivers, including family members, home health care nurses, and patient care technicians.

Patients with tracheostomies are seen in emergency departments (EDs) for a variety of problems related to the tracheostomy. Many emergency clinicians will have minimal experience with such patients and it should be emphasized that complications or problems with tracheostomies often require specialty consultation in the ED to be fully evaluated or definitively remedied. Common complaints include difficulty breathing as a result of tube obstruction, tube displacement, or equipment failure; poor oxygensation from infection or altered patient anatomy; and bleeding. In some cases, complications can be life threatening. Emergency physicians must be knowledgeable regarding the evaluation and management of patients with tracheostomies. This chapter reviews the relevant tracheal anatomy and essential tracheostomy equipment, discusses pertinent tracheostomy care, provides a systematic approach to the evaluation and management of selected complications, and identifies high-risk patient populations.

BACKGROUND

Most tracheostomies are performed electively. For elective tracheostomies, the surgical site is between the first and second or the second and third tracheal rings. With an open, or surgical, tracheostomy, the anterior aspect of the trachea is generally left sutured to the skin until the tract matures, approximately 4 to 5 days after the procedure. In recent years, percutaneous dilational tracheostomy has become the preferred technique for many ICU patients. It can be performed at the bedside and eliminates the risks associated with transporting critically ill patients to an operating room. Postoperative complications vary and depend on the timing and insertion technique. Early postoperative complications tend to arise in the first few days to weeks. Sixteen to twenty percent of patients experience early complications, and 6% to 8% experience late complications. Although elective and emergency tracheostomies are generally performed with the same technique, the complication rate associated with emergency procedures may be higher than the rate for elective tracheostomies.

TRACHEAL ANATOMY AND PHYSIOLOGY

The lower respiratory tract begins at the vocal cords. Inferior to the vocal cords lies the cricoid cartilage, which encases the 1.5- to 2-cm subglottic space. Inferior to the cricoid cartilage is the trachea (Fig. 7.1). The typical adult trachea is 10 to 12 cm in length. The anterior and lateral walls of the trachea are supported by 18 to 22 incomplete cartilaginous rings. A fibromuscular sheet lying anterior to the esophagus completes the posterior wall. The interior diameter of the adult trachea is 12 to 25 mm, and it is lined with mucosa covered by respiratory epithelium. Blood is supplied to the trachea by branches of the inferior thyroid, innominate (brachiocephalic), bronchial, and subclavian arteries. Critically, the innominate artery (IA) lies in close proximity to the tracheostomy stoma. From its origin at the aortic arch, the IA courses between the sternum and the anterior aspect of the trachea and veers right at the sternomandibular joint. The location of the IA is important because erosion of the anterior tracheal wall can lead to life-threatening bleeding. The recurrent laryngeal nerve innervates the intrinsic laryngeal muscles and mucosa below the vocal cords. Efferent vagal fibers stimulate bronchoconstriction, mucosal secretions, and vasodilation. Efferent sympathetic fibers of the pulmonary plexus stimulate tracheal bronchodilation and vasoconstriction.

The upper airway, including the oropharynx and nasal passages, filters particulate matter, humidifies inspired air, and aids in the expectoration of secretions. These functions are reduced in patients with a tracheostomy. Placement of a tracheostomy bypasses humidification and results in the formation of thick, dry secretions. In the absence of humidification, squamous metaplasia and chronic inflammatory changes develop in the trachea. Bronchoconstriction resulting in reduced airflow can occur if the inspired air temperature is below room temperature. Normal mucociliary clearance is also impaired because of the increased viscosity of respiratory secretions, which underlies chronic illness and respiratory infections, particularly with Mycoplasma or viral pathogens.

The tracheostomy procedure weakens the anterior tracheal wall and blunts the normal cough mechanism, an important component for clearance of secretions by the trachea. Normally, the epiglottis and vocal cords close to trap air in the lungs and raise intrathoracic pressure before a cough. Patients with a tracheostomy tube are generally unable to generate sufficient pressure to initiate a strong cough and facilitate airway clearance. In addition to an impaired cough mechanism, immune responses are often blunted in patients with tracheostomies as the result of underlying illnesses, chronic lung disease, chemotherapy, or acquired immunosuppression.

EVALUATION OF TRACHEOSTOMY PATIENTS

Every ED patient with a tracheostomy who has a respiratory complaint or a complaint related to the tracheostomy should be thoroughly evaluated. Begin the primary assessment with a review of the patient’s vital signs and evaluation of the airway. Examine the tracheostomy and consider dislodgement, obstruction, and fracture of the tube. Next, assess the patient’s...
VENTILATOR SETTINGS OR TRACHEOSTOMY CARE, INCLUDING INCREASED OXYGEN USE, INCREASED SUCTIONING, EQUIPMENT FAILURE, OR CHANGES IN EQUIPMENT.

After the primary assessment and focused history, perform a thorough physical examination to differentiate between tracheostomy complications and other causes of the patient's respiratory complaint. The differential diagnosis may include pneumonia, exacerbation of chronic obstructive pulmonary disease, congestive heart failure, pulmonary embolism, pneumothorax, and acute coronary syndrome. During the secondary evaluation, evaluate the patient's stoma for signs of bleeding, infection, or skin breakdown. Make a note of the specific type and model of equipment used in the event that a replacement tube is needed.

GENERAL EQUIPMENT FOR TRACHEOSTOMY PATIENTS

Before performing any procedure on the tracheostomy tube, it is important that the emergency physician ensure that essential equipment is readily available at the patient's bedside. These items are shown in Fig. 7.2. Adequate preparation is crucial in preventing a poor outcome should complications arise. Most of this equipment can be placed in a designated airway box that can easily be accessed within the room or ED.

ROUTINE TRACHEOSTOMY MAINTENANCE

Routine tracheostomy maintenance involves (1) regular cleaning of the tube, (2) frequent stomal care, and (3) periodic monitoring of cuff pressure. There are many different types of tracheostomy tubes, and the focus of this section is on the most common, those with a removable inner cannula (Fig. 7.3A). Regular cleaning of the tracheostomy tube and inner cannula can prevent the accumulation of dried secretions. Lack of cleaning and maintenance of the inner cannula is the primary cause of tube obstruction (see Fig. 7.3B). Under normal circumstances the inner cannula should be in place. It sits snugly within the tracheostomy tube and can easily be removed without disturbing the tube itself. The inner cannula should be cleaned daily.
maximum recommended cuff pressure, the tube may have become dislodged, which requires further evaluation.

It is essential that adequate air humidification be provided to patients with a tracheostomy. Inadequate humidification can result in obstruction of the tube from thick secretions, sputum retention, keratinization or ulceration of the tracheal mucosa, and impaired gas exchange as a result of lung atelectasis. Ambulatory patients and those who require low-flow oxygen can be fitted with a heat-moisture exchanger that attaches to the external opening of the tracheostomy tube. Patients receiving long-term ventilation or high-flow oxygen require regular saline nebulizer treatments delivered by an in-line humidification system.

VENTILATING TRACHEOSTOMY PATIENTS

To properly ventilate ED patients with a tracheostomy, it is important to determine the make, model, and type of tube (Fig. 7.4). For proper ventilation, a 15-mm tube/Ambu bag or ventilator tube adapter must be present, either on the tube itself or on the end of an inner cannula that has an inflatable cuff. The inner cannula adapter will accept an Ambu bag or ventilator tubing, and the cuff will allow positive pressure ventilation. If the patient requires manual or mechanical ventilation and the tracheostomy tube is not suitable for ventilatory support, immediately replace it with a 6-0 cuffed endotracheal (ET) tube for ventilation.

TRACHEAL SUCTIONING

Tracheal suctioning is required to remove secretions or aspirated material from the upper airway in patients whose cough is impaired or in whom an artificial airway is in place. Tracheal suctioning can be performed through an ET tube, a tracheostomy tube, a minitracheostomy placed in the cricothyroid membrane, or the nasopharynx. The indications, equipment, procedure, and complications are similar for each technique.
Indications

The primary indications for tracheal suctioning are to remove secretions, enhance oxygenation, or obtain samples of lower respiratory tract secretions for diagnostic tests. In the ED, tracheal suctioning should be performed to enhance oxygenation in any tracheostomy patient in respiratory distress. In addition, tracheal suctioning should be performed when the patient has coarse rales, rhonchi, or tubular breath sounds; acute or worsening dyspnea; or arterial oxygen desaturation. It is important to emphasize that tracheal suctioning should be performed only when it is clinically indicated; frequent, routine suctioning is not recommended.18,19

There are no absolute contraindications to tracheal suctioning. Relative contraindications include severe bronchospasm, which may worsen with suctioning, and persistently elevated intracranial pressure (ICP), which is exacerbated by suctioning.20 Bronchodilators, sedatives, and paralytics may alleviate these symptoms. Tracheal suctioning should be undertaken with caution in patients with cardiovascular instability because of an increased risk for associated dysrhythmias.21

Equipment

A suction catheter and vacuum system, open or closed, is required to perform tracheal suctioning (Fig. 7.5). It is recommended that the diameter of the suction catheter be no larger than half the inner diameter of the tracheostomy tube.22,23 The size of the suction catheter in French gauge (Fr) can be calculated as follows:

\[
\text{Size (Fr)} = 2 \times (\text{Size of the tracheostomy tube} - 2)
\]

For example, a 7-0 tracheostomy tube will require a 10-Fr suction catheter because \(2 \times (7 - 2) = 10\) Fr. If the catheter is too small, it will not remove excess secretions adequately. If the catheter is too large, it can obstruct airflow during insertion and cause alveolar collapse with resultant hypoxemia.

In adults, the suction catheter should be inserted only 10 to 15 cm, depending on the length of the tracheostomy tube. The goal of suctioning is to remove secretions only from the proximal airways. Shallow suctioning occurs when the catheter is placed just beyond the hub of the tracheostomy tube to remove proximal secretions. Premature suctioning occurs when a catheter is inserted such that the distal side ports are beyond the caudal end of the tracheostomy tube. Deep suctioning occurs when the suction catheter is advanced until resistance is met. It is used for clearing excess secretions in the lower airways.19 Deep suctioning has not been shown to be more beneficial than shallow suctioning and should not be performed routinely.

**Review Box 7.1** Tracheal suctioning: indications, contraindications, complications, and equipment.
because it might damage the mucosal epithelium and lead to an increase in granulation tissue.24,25

A number of suction catheter tips have been designed to maximize removal of secretions without causing mucosal injury. Tips may have a single or multiple side ports proximal to the distal tip. Directional or Coude tip catheters are available for selective suctioning of the main stem bronchi.

A closed-system airway encases a suction catheter in a sterile sheath attached to ventilator tubing. This prevents the suction catheter from being contaminated by contact with the outside environment and allows tracheal suctioning to be performed without interrupting ventilatory support. The vacuum should be set to the lowest possible pressure to reduce atelectasis. Vacuum pressure should not exceed 80 mm Hg in infants or 150 mm Hg in adults.22

Procedure and Technique

Before tracheal suctioning, continuous pulse oximetry, cardiac monitoring, and continuous capnography, if available, should be initiated (Fig. 7.6, step 1). Awake and alert patients should be sitting upright with their head in a neutral position. For mechanically ventilated patients, the head of the bed should be elevated to 30 degrees to improve respiratory mechanics. Aseptic technique should be used throughout all suctioning procedures to prevent the introduction of bacteria. Backup airway equipment should be readily available (see Fig. 7.2).

Preoxygenate the patient for at least 30 to 60 seconds. For mechanically ventilated patients, increase the fraction of inspired oxygen (FiO2) to 100%. For nonventilated patients, provide 10 to 15 L of high-flow oxygen. Humidify the air before suctioning to reduce the viscosity of respiratory secretions. Routine instillation of normal saline has not been shown to provide regular clinical benefit and is no longer recommended.22,26

Once the patient has been adequately preoxygenated, insert the suction catheter through the inner cannula in situ (see Fig. 7.6, step 2). If the carina is irritated during deep suctioning, a vigorous cough reflex will be activated. After reaching the desired depth, withdraw the suction catheter 1 to 2 cm and then apply suction while slowly removing the catheter (see Fig. 7.6, steps 3 and 4). Gently rotate the catheter as it is withdrawn to facilitate removal of secretions. The duration of suctioning should not exceed 10 to 15 seconds.27

Monitor the patient throughout the procedure for signs of cardiac dysrhythmia, hypoxia, or a rise in end-tidal CO2. Immediately stop suctioning if any of these signs develop (see...
section on Complications of Suctioning). If marked respiratory distress is presumed to be secondary to significant tracheal obstruction, continue with expeditious suctioning to remove the obstruction (see section on Obstruction and Complications from Tube Changes).

**Complications of Suctioning**

Complications that occur during or after suctioning are relatively common and can result in significant morbidity. Most complications can be anticipated and simple maneuvers can reduce their incidence and severity.

Hypoxemia from suctioning may cause increased ICP, dysrhythmias, or even death. In neonates, hypoxia during suctioning may contribute to spontaneous intracerebral hemorrhage. A number of factors contribute to suctioning-related hypoxia, including interruption of mechanical ventilation, aspiration of air from the respiratory tract, and suctioning-related atelectasis. Use in-line suction catheters for ventilator-dependent patients to allow continuous oxygen delivery and positive pressure ventilation. Select the catheter size carefully to reduce the evacuation of airway gases during suctioning and help prevent atelectasis. Limit the duration of suctioning to 10 to 15 seconds and perform no more than three passes in succession. Measurement of arterial oxygen saturation may not be sufficient to assess hypoxia after suctioning. Oxygen consumption increases during suctioning despite insignificant changes in oxygen saturation. This increase in oxygen consumption is more prevalent in patients who possess a vigorous cough, are agitated, or resist suctioning.

Dysrhythmias associated with suctioning may be caused by hypoxia, increased myocardial oxygen consumption, vagal stimulation, hypoventilation, or catecholamine release. Vagal stimulation caused by suctioning can cause bradycardia and hypotension. Bradycardia in the setting of hypoxia potentiates ventricular dysrhythmias, including ventricular fibrillation. Nebulized or intravenous atropine is recommended for bradycardia and can be used as pretreatment in patients at risk for bradycardia, particularly infants. Digoxin enhances vagal activity and may potentiate the vagal stimulation of ET suctioning. Sympathetic stimulation may occur as a result of hypoxia, pain, or stress of the procedure. Pain medications, anxiolytics, and preparation of the patient for the procedure may blunt the sympathetic response. Suctioning should be stopped immediately if a dysrhythmia develops.

Increases in ICP during suctioning are well documented. ET suctioning can induce a strong cough reflex, which is thought to contribute to the increase in ICP by raising intrathoracic pressure, reducing cerebral perfusion pressure, and increasing systemic blood pressure. In susceptible patients, increases in ICP can lead to devastating outcomes. If there is concern that increases in ICP could harm the patient, several preventive steps should be taken before initiating ET suctioning. The most important interventions are providing adequate sedation and maximizing oxygenation. Hyperventilation before and between passes of the suction catheter can transiently lower ICP by reducing systemic CO₂. One minute before suctioning, hyperventilate the patient by increasing the respiratory rate to approximately 30 breaths/min. If not heavily sedated, most patients will cough vigorously when suctioned. Lidocaine can be instilled into the trachea to blunt the cough reflex, thereby preventing an increase in ICP. To anesthetize the trachea locally, instill 1 to 1.5 mg/kg of 2% lidocaine into the tracheal tube.

After administering the medication, prepare the patient for suctioning by ensuring that sedation and oxygenation are adequate. After 10 minutes to allow the lidocaine to produce its local effect, begin tracheal suctioning. This method has been shown to prevent increases in ICP and changes in cerebral hemodynamics.

Atelectasis can occur when airway gases are suctioned too rapidly. To reduce this complication, choose a suction catheter that is less than half the inner diameter of the tracheostomy tube and minimize the duration and suction pressure. Atelectasis can be minimized by using a closed suction system and providing positive end-expiratory pressure after suctioning. Hyperventilation should not be performed routinely to resolve suction-related atelectasis.

Mucosal injury is a common complication of tracheal suctioning. Invagination of the mucosa into the side ports of the catheter occurs during suctioning and causes the tracheal mucosa to become denuded, edematous, and predisposed to bleeding. Mucosal damage also interferes with mucociliary transport. Tracheitis can occur as a result of frequent or improperly performed suctioning.

**Minitracheostomy Suctioning Procedure**

The minitracheostomy (minitrach) was designed to improve tracheal hygiene in patients with intact cough reflexes, normal ventilatory function, and vocalization. The minitrach serves as a small port solely for suctioning secretions. Commonly, a 4-mm indwelling cuffless cannula is inserted through the cricothyroid membrane into the trachea. Patients who are suctioned through a minitrach are at lower risk for gagging and aspiration because they are able to maintain laryngeal and glottic function. The minitrach device is seldom used in children because they have smaller airway diameters.

The technique used to suction through a minitrach is the same as that for tracheal suctioning. The smaller port size may require that smaller catheters be used. Most patients with minitrachs are decannulated before discharge from the ICU and are rarely seen in EDs.

**CHANGING A TRACHEOSTOMY TUBE**

**Indications**

Maturation of the tracheostomy tract is generally completed by postoperative day 7. Most ED patients with a tracheostomy are seen after the stomal tract has matured, so routine changes of the tracheostomy tube can be done safely in the ED. Indications for exchange of the tracheostomy tube include cuff rupture or leak, leakage around the tube caused by tracheomalacia, other changes in tracheal anatomy, complete or partial tube occlusion, and conversion to an alternative tube style. There are no absolute contraindications to exchanging a tracheostomy tube in the ED as long as the stomal tract has matured. Before undertaking the exchange, consider whether further tissue trauma or hemorrhage might occur as a result of it or whether anatomic abnormalities could make the exchange difficult.

There is conflicting evidence on how frequently tracheostomy tubes should be changed. Do not perform tube changes on a predetermined schedule, but rather as the patient’s clinical condition dictates. There is some evidence that tubes left in
Changing a Tracheostomy Tube

**Indications**
- Complete or partial tube obstruction
- Mechanical failure
- Peritubal leak
- Cuff rupture
- Conversion to an alternative tube style (only with a mature stomal tract)

**Contraindications**
**Absolute:** None
**Relative:** Tracheostomy placement <7 days earlier

**Complications**
- Hemorrhage
- False passage
- Obstruction
- Prolonged procedure time
- Tract closure
- Hypoxia
- Subcutaneous emphysema
- Aspiration

**Equipment**
- Red rubber catheter (optional)
- Outer cannula
- Obturator
- Inner cannula
- Lubricating jelly
- 10-mL syringe
- Tracheostomy tube:

**Table 7.1** Common Tracheostomy Tube Sizes With Dimensions

<table>
<thead>
<tr>
<th>Portex Cuffed D.I.C. Tracheostomy Tubes</th>
<th>Shiley Tracheostomy Tubes*</th>
</tr>
</thead>
<tbody>
<tr>
<td>TUBE SIZE (mm) AND COLOR CODE</td>
<td>TUBE SIZE (JACKSON)</td>
</tr>
<tr>
<td>-----------------------------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>6.0 (orange)</td>
<td>4</td>
</tr>
<tr>
<td>7.0 (green)</td>
<td>6</td>
</tr>
<tr>
<td>8.0 (white)</td>
<td>8</td>
</tr>
<tr>
<td>9.0 (blue)</td>
<td>10</td>
</tr>
<tr>
<td>10.0 (yellow)</td>
<td>10</td>
</tr>
</tbody>
</table>

*Shiley also offers tracheostomy tubes with both distal and proximal (relative to the cuff) extended lengths for patients with large necks or other abnormal anatomy.

DIC, Disposable inner cannula.


place longer than 3 months are at higher risk for infection. Most manufacturers recommend that tracheostomy tubes be changed approximately 30 days after placement.

**Equipment**

When changing a tracheostomy tube, familiarize yourself with the resources and equipment available in the ED. Keep the equipment readily available, if not at the patient's bedside. The necessary equipment is depicted in Fig. 7.2.

Tracheostomy tubes may be made from metal or other synthetic material such as plastic. Plastic tubes are either polyvinyl chloride, which softens at body temperature, or silicon, which is naturally soft and unaffected by temperature. Metal tubes are constructed of silver or stainless steel and lack both a cuff and a 15-mm connector for attachment to a ventilator or Ambu bag.

**Sizing**

To determine the appropriate size of tube, consider the internal diameter (ID), outside diameter (OD), length, and curvature of the tube. The Chevalier Jackson sizing system indicates the length and tapering of the OD. This sizing method applies to metal tubes and most Shiley dual-cannula tracheostomy tubes. Table 7.1 lists common tracheostomy tubes and their dimensions. For dual-cannula tracheostomy tubes, the inner cannula has a 15-mm connection for a ventilator. Note that tracheostomy
tubes with the same ID can have very different ODs and lengths. Consider the pretracheal distance before selecting a replacement tube of the appropriate size for obese patients (see section on Special Populations).

Single-cannula tracheostomy tubes are sized by the ID of the tube at its smallest dimension. The size of the tube is usually stamped on the flange. Before changing a tube, have the appropriate size of tube at the bedside along with tubes that are one or two sizes smaller. As a rule of thumb, most women can accommodate a tube with an OD of 10 mm, and most men can accommodate a tube with an OD of 11 mm. Table 7.1 lists the recommended sizes of tracheostomy tubes.

### Components

Most tracheostomy tubes have three standard components: an outer cannula, an obturator, and an inner cannula (see Fig. 7.3.4). The outer cannula is the permanent portion of the tracheostomy tube and should not be removed unless complications arise or a tube change is needed. Attached to the outer cannula is a flange on either side with eyelets used to tether the tube to the patient’s neck. The obturator is a white rounded or cone-shaped object that is used to facilitate insertion of the tube. When inserted into the outer cannula, it extends several millimeters beyond the distal end of the tube.

Tracheostomy tubes can be cuffed or uncuffed. Cuffed tracheostomy tubes are used for patients on long-term mechanical ventilation and those at risk for aspiration. Cuffed tubes also prevent loss of volume during positive pressure ventilation while preventing air leaks across the vocal cords. Speech is not possible for patients with a cuffed tube. Most tubes have a high-volume, low-pressure cuff that reduces mucosal injury and the risk for tracheal erosion or stenosis. Low-volume, low-pressure cuffs may be more effective in preventing aspiration. Inflate the tracheostomy cuff and deflate it by attaching a syringe to the Luer-Lok port at the proximal end of the pilot balloon and either injecting or removing approximately 10 mL of air. Determine cuff pressure by connecting the Luer-Lok port to a handheld manometer.

Uncuffed and metal tubes are used in patients with adequate ventilatory effort who are alert and at low risk for aspiration. Depending on the size of the tracheostomy tube and how much of the tracheal diameter is filled by the tube itself, the patient may be able to speak. Air must be able to bypass the tube and be transmitted across the vocal cords. Digital occlusion of the tracheostomy tube or the use of specialized speaking valves can occlude expired air from the tracheostomy and facilitate voice production. Fenestrated tubes also allow air to be transmitted across the vocal cords. The fenestrations are generally located at the superior, posterior arch of the tube but can also be found on the inner cannula in some models.

During weaning from tracheostomy, tracheal buttons can be used to maintain patency of the stoma in patients who do not require mechanical ventilation. They can be retained permanently if decannulation is not possible. Tracheal buttons have a hollow outer cannula and a solid inner cannula and extend from the outer skin into the tracheal lumen. Tracheal buttons can become displaced into the tracheal lumen if they are not tethered correctly and may become clogged with secretions. They may also have a speaking valve, such as the Passy-Muir (Passy-Muir, Inc., Irvine, CA) or the Shiley Phonate (Mallinckrodt Medical, St. Louis, MO). These devices generally clip or twist onto the 15-mm coupling of the tracheostomy tube or inner cannula. Remove the speaking valve before changing the tracheostomy tube.

### Procedure

It is essential to ensure that all airway equipment is at the patient’s bedside before performing the tube exchange. In addition to having airway equipment ready, place the patient on continuous pulse oximetry, cardiac monitoring, and capnography, if available, to confirm tube placement. Identify the tracheostomy tube model and determine its size. Have this size and two other tubes that are one or two sizes smaller in the event that tube replacement is difficult. Inspect all equipment for proper function, including the replacement tube cuff for leaks and the obturator for ease of insertion and removal (Fig. 7.7, step 1). Coat the replacement tracheostomy tube with a water-based lubricant (see Fig. 7.7, step 2).

Place the patient in a semirectum position with the neck slightly extended to ensure proper alignment of the external stoma and the tracheostomy tract. Do not flex the neck because this may misalign the tissues and make tube replacement more difficult. Remove the old tracheostomy dressing and clean the stomal site. If awake, preoxygenate the patient by placing a non-rebreather oxygen mask over the tracheostomy site for 3 minutes and then suction the oropharynx. If the patient is ventilator dependent, increase the FiO₂ to 100% for at least 1 minute before beginning the tube exchange to ensure adequate oxygenation. If needed, provide the patient with soft restraints or anxiolytic medication to improve compliance.

The tracheostomy tube can be changed by either of two methods. If the tracheostomy tract is well matured, the tube can be exchanged with an obturator. To remove the old tracheostomy tube, first deflate the cuff completely (if present) (see Fig. 7.7, step 3). Remove the existing tube with an “out-then-down” movement while the patient exhales (see Fig. 7.7, step 4). Next, with the obturator in place, insert the new tube into the stoma at a 90-degree angle to the cervical axis (see Fig. 7.7, step 5). If two experienced providers are available, one can be responsible for deflating the cuff and removing the old tube while the other inserts the new device. Next, gently push the tube downward in a fluid, sweeping motion so the external flange is flush against the neck. If necessary, use a tracheal hook to hold the stoma open. Remove the solid obturator immediately, insert the hollow internal cannula, inflate the cuff, return the patient’s head to the neutral position, and secure the external flange (see Fig. 7.7, steps 6 to 8).

The second technique involves exchanging the tube with a modified Seldinger technique using a red rubber catheter, nasogastric tube, gum elastic bougie, or fiberoptic nasopharyngeal scope (Fig. 7.8). This technique is preferable if the tracheostomy tract is not well defined or if there is concern that tube exchange could be difficult. Using a nasopharyngeal scope can also confirm placement by way of direct visualization. To exchange tubes with this method, first premeasure the distance needed to extend the guidewire device beyond the distal tip of the old tracheostomy tube. Advance the guide to the premeasured distance, deflate the cuff, and remove the tube as described previously. Next, without the obturator in place, advance the new device over the guide until it is seated securely in the trachea. Remove the guide and secure the new tube. Weimann and Bander developed a modified Seldinger technique involving the use of an airway exchange catheter to allow jet or bag ventilation into the trachea during tube...
Remove the obturator, and insert the inner cannula. Insert the new tube (with the obturator in place) into the stoma at a 90-degree angle; push the tube downward in a fluid, sweeping motion.

Inspect the tube for proper function; check for a cuff leak. Apply a water-based lubricant to the tube.

Deflate the cuff on the old tube completely. Remove the old tube during exhalation.

Insert the new tube (with the obturator in place) into the stoma at a 90-degree angle; push the tube downward in a fluid, sweeping motion. Remove the obturator, and insert the inner cannula.

Attach the ventilator to the tracheostomy tube. Inflate the cuff.

**Figure 7.7** Changing a tracheostomy tube. Maintain the neck in the same position (slightly extended) for removal and insertion so that the tract is not lost. If the tube is already out, and the specifics of the tract are unknown, extending the neck to align the stoma with the tissue tract is the first best position for tube replacement.
inner cannula, or at the distal end of the outer cannula. In one review, 30% of ED visits for respiratory distress were due to an obstructed tube. Obstruction is caused most commonly by dried respiratory secretions and, less often, by blood, aspirated material, or granulation tissue. The obstructing object may act as a ball valve and allow air to enter but restricts expiration. Such obstruction is easily remedied in the ED by removing the inner cannula and then cleaning and replacing it.

Preparation

First, assess the patient’s airway for patency and respiratory status. Place the patient on a cardiorespiratory monitor, a pulse oximeter, and continuous capnography. Gather appropriate airway equipment at the bedside. Be sure to have a large-bore suction catheter available in case the appropriate size of catheter is inadequate. Examine the tracheostomy tube to see whether the inner cannula is obstructed or any obvious external obstruction is present.

Interventions

Administer high-flow oxygen and encourage patients who can breathe spontaneously to cough. Manually remove any obstruction seen at the external tracheal tube opening. If there is no obvious external obstruction, remove the inner cannula and suction the secretions. Inspect the inner cannula and remove any obstructing objects. If the patient’s tracheostomy does not have an inner cannula, suction the tracheostomy tube to remove obstructing plugs. Thick secretions can be loosened in a critically ill patient by instilling normal saline into the tracheostomy tube, but this is no longer recommended for routine suctioning.

If the obstruction persists, evaluate the distal tip of the tracheostomy tube. First, deflate the tracheostomy cuff and supply high-flow oxygen via face mask if the patient is breathing spontaneously or via a bag-valve-mask device if the patient is unable to breathe without assistance. Note that if the tracheostomy cuff is inflated, you will not be able to provide oxygen or ventilatory support from above the tracheostomy tube. If the patient is still in exchange. This adjunct delivers intratracheal oxygen and is helpful if hypoxemia is likely to occur.

Confirm proper placement of the tube within the trachea with one of several possible techniques. Traditionally, the patient is ventilated and correct tube placement confirmed by observation of equal chest rise and auscultation of bilateral breath sounds. Although both signs are important to confirm correct placement, quantitative waveform capnography is now a class I American Heart Association recommendation for confirmation of ET intubation. It should be used for tracheostomy tube placement, if available. Multiple studies have shown quantitative capnography to be a highly sensitive tool for confirming correct placement of airway devices. Correct tube placement can also be confirmed by direct visualization of the tracheal rings with a fiberoptic scope.

COMPLICATIONS OF TRACHEOSTOMY

Complications related to tracheostomy placement are common and sometimes life-threatening. This section addresses the emergency development of late postoperative complications (occurring more than 3 weeks after the operation). Patients with immediate and early postoperative complications are usually still hospitalized and are less likely to be seen in the ED. Late postoperative tracheostomy complications encountered in ED patients include obstruction, dislodgement, equipment failure, infection, anatomic disruption, and hemorrhage. Management of the patient varies depending on the type of complication that the patient is experiencing. Rapid recognition of the problem is paramount and can drastically affect the outcome.

Obstruction and Complications From Tube Changes

Obstruction of the tracheostomy tube is a common complication and can occur at the external opening of the tube, within the inner cannula, or at the distal end of the outer cannula. In one review, 30% of ED visits for respiratory distress were due to an obstructed tube. Obstruction is caused most commonly by dried respiratory secretions and, less often, by blood, aspirated material, or granulation tissue. The obstructing object may act as a ball valve and allow air to enter but restricts expiration. Such obstruction is easily remedied in the ED by removing the inner cannula and then cleaning and replacing it.
Dislodgement

Displacement of the tracheal tube is a serious complication that can have a disastrous outcome if not recognized and corrected quickly. Patients at highest risk for poor outcomes from a dislodged tube are those who are obese, who have recently inserted or changed tubes, who have anatomic anomalies, and who are difficult to ventilate or were difficult to intubate in the past.56–58 Evidence of tube dislodgement is usually obvious. Signs and symptoms include hypoxemia, agitation, respiratory distress, altered mental status, subcutaneous emphysema, and high airway pressure. Dislodgement can occur during patient transfers, when traction is placed on the tube, or when the tube is manipulated for bag-valve-mask ventilation or ventilator tube connection. If a tracheostomy tube is too long, it can become dislodged inferiorly, which causes the tip to either abut the mucosal wall of the trachea, or obstruct it at the level of the carina.

It is important to know when and why the tracheostomy was placed because such information will influence evaluation and management of the patient.59 If a tracheostomy tube becomes dislodged within the first 7 days after initial placement, the stoma can close rapidly and make reinsertion difficult. Blind, forceful attempts at reinsertion of the tracheostomy tube in the early postoperative period can result in the creation of a false passage and possible respiratory arrest. If accidental decannulation occurs before the tract has time to form, orotracheal intubation is the safest approach in the ED, unless the patient has had a laryngectomy (see section on Special Populations). Reinsertion of the tracheostomy tube is possible, but the patient will most likely require ET intubation to secure the airway.70 Patients with abnormal neck anatomy or other causes of a “difficult airway” may not benefit from reinsertion in the ED. In these cases, the procedure may need to be performed in the operating room. Seek emergency surgical consultation for complicated cases.

Preparation

As for all patients with tracheostomy complications, begin with a quick primary assessment and place the patient on appropriate monitoring devices. Continuous waveform capnography can be invaluable in determining whether the tracheostomy tube is placed properly. Absence of a waveform or end-tidal CO2 partial pressure less than 10 mm Hg is an indication that the airway device has become dislodged or was placed inappropriately.60 It is essential to have advanced airway equipment at the bedside, including two replacement tracheostomy tubes (each with an inner cannula). A fiberoptic nasopharyngeal scope or bronchoscope, if available, can be helpful. Position the patient with the neck in extension to maximize alignment of the stoma and trachea. Neck flexion can cause downward displacement of the tube by as much as 3 to 4 cm.16

Interventions

First, determine that the tube flanges rest snugly at the skin. Lateral neck x-ray films may reveal that the opening of the tracheostomy tube is abutting the anterior tracheal wall or obstructing the tracheal lumen. If the tracheostomy tube appears to be minimally displaced, attempt to reposition the tube by gentle manipulation. If the tracheostomy tube is completely dislodged or no waveform is present on capnography, tube replacement should be attempted.

If the tracheostomy stoma was created less than 7 days earlier, be prepared to orally intubate the patient. First, cut any flange sutures and remove the tracheostomy tube (Fig. 7.9). Stay sutures may be placed to hold the stoma open and better visualize the tracheal opening. If the patient is stable, attempt to reinsert the tracheostomy tube with the assistance of a gum elastic bougie or fiberoptic scope.45 If the patient is unstable with an obstructed tracheostomy tube, cut the flange sutures, remove the tube, and orally intubate the patient.

If the tracheostomy stoma was created 7 to 30 days earlier, remove the tube and any other cause of obstruction from the stoma. If the patient can ventilate independently, allow the patient to oxygenate and then reinsert the tube when ready. If the patient is unstable, occlude the stoma with moist gauze or other occlusive device and provide bag-valve-mask ventilation. Once the patient is stabilized, replace the tracheostomy tube. If a replacement tube is not available, check the outer diameter of the tracheostomy tube, and insert a cuffed ET tube of equal size (or at minimum, 6-0) as a substitute.

If the tracheostomy is more than 30 days old, the tube may not need to be replaced. If the patient is unstable, prepare as described for reinsertion or ET intubation. If the patient is stable and ventilating spontaneously without any signs of distress, contact the appropriate specialty care physician to discuss the need for emergency reinsertion.

Once the tube is reinserted, confirm correct placement with auscultation and waveform capnography. Tube position can also be confirmed by direct visualization with a fiberoptic scope.
**False Passage**

A false lumen can be created during replacement of a tracheal tube or during repositioning of a dislodged tube. Obese patients are at high risk for false passage because of their redundant neck tissue (see section on Special Populations). Subcutaneous air, crepitus, or distortion of anterior neck landmarks may indicate placement of the tracheostomy tube into a false passage. Absence of a waveform on capnography confirms misplacement of the tracheostomy tube. Abdominal distention after bag-valve-mask ventilation may indicate placement of the tracheal tube through a tracheoesophageal fistula (TEF). If a false passage is suspected, remove and replace the tracheostomy tube expeditiously.

**Equipment Failure**

**Fracture**

Tracheostomy tubes do not fracture frequently. When fractures do occur, they are most often located at the juncture of the flange and the tube connection.\(^61,62\) A fractured tube fragment may migrate inferiorly and obstruct the tracheal lumen. Patients may have acute respiratory complaints such as cough, dyspnea, choking, or wheezing. Prolonged retention of a foreign body can result in chronic respiratory symptoms such as wheezing, coughing, or recurrent bouts of pneumonia or bronchiectasis. To manage this problem, replace the tube if possible and consider bronchoscopy for retrieval of the tube fragment.\(^63\)

**Tracheal Cuff Complications**

Complications related to the tracheal tube cuff include perforation, which results in a poor seal and increased risk for aspiration; overinflation, which causes pressure on or impingement of the esophageal lumen; and distention of the cuff distal to the tracheal tube, which results in obstruction of the tracheal tube opening. Mucosal injury is less common since the use of low-pressure cuffs has become standard practice. Pain with ventilation or swallowing, inadequate oxygenation, or the presence of gastric secretions in the tracheostomy tube may indicate cuff problems. Verify inflation pressure with a manometer (target range, 18 to 25 mm Hg) and appropriate cuff position. Replacement of the tube is indicated if the symptoms persist.

**Infection**

Patients who require long-term tracheostomy and mechanical ventilation are at high risk for nosocomial pneumonia, ventilator-associated pneumonia (VAP), and tracheobronchitis. In general, the frequency of infection increases with the duration of mechanical ventilation, and the risk for infection is highest in the first week following intubation. Nonventilated tracheostomy patients are also at increased risk for pneumonia, tracheobronchitis, stomal infections, and other soft tissue infections.

Risk factors for systemic infection include impairment of host defenses and exposure to large numbers of bacteria that bypass the upper airway defense systems. In healthy patients, the upper respiratory tract is colonized by normal oropharyngeal flora. In tracheostomy patients, the normal flora can be replaced by virulent pathogens such as enteric gram-negative bacteria.\(^54\) The organisms most commonly cultured from tracheostomy stomas and tubes are *Pseudomonas aeruginosa*, *Acinetobacter*, and *Staphylococcus aureus*.\(^65\) Colonization rates are high in these patients, even in the absence of systemic infection. The tracheostomy tube bypasses the natural protective barriers of the upper airway. Suctioning a colonized tracheostomy tube can introduce bacteria into the lower respiratory tract. Underlying medical conditions, prolonged hospitalization, impaired host defense mechanisms, and poor nutrition all increase susceptibility to infection.

Ventilator-associated tracheobronchitis (VAT) is the result of colonization of the upper airway and can eventually progress to VAP.\(^66\) Patients with VAT have fever, production of purulent sputum, and positive respiratory cultures but no evidence of a new infiltrate on chest radiography.\(^66\) VAT is frequently caused by multidrug-resistant organisms. When choosing antimicrobial therapy, consider recent hospitalizations, previous infections, recent antibiotic use, and the hospital's antibiogram.

Tracheostomy patients on long-term ventilation are also at risk for VAP. These patients are similar to those with VAT, except that they have an infiltrate on chest radiography. A common cause of VAP is aspiration. Patients become predisposed to aspiration if they have altered neurologic function, have an abnormal swallowing mechanism, or are mechanically ventilated and kept in the supine position for a prolonged time. Some degree of aspiration occurs in 33% to 61% of all patients with tracheostomies.\(^67,68\) To reduce the risk for aspiration, keep patients requiring mechanical ventilation in a semirecumbent position. It is generally recommended that 30 to 45 degrees is adequate to reduce the risk for VAP.\(^69-71\)

Stomal infections and cellulitis are also common.\(^31\) The bacteria cultured most frequently from patients with tracheostomy-related cellulitis are *S. aureus*, *Pseudomonas* species, and *M. alba*. Consider *Candida albicans* infection in patients previously treated with antibiotics and those who have an underlying immunocompromised state. Peristomal cellulitis can usually be treated with good wound care and oral antibiotics. The most dangerous complications from cellulitis are mediastinitis, mediastinal abscess, necrotizing fasciitis, and para-tracheal abscess. Consider these complications in patients who have pain with breathing, pain with swallowing, or signs of systemic infection. Strongly consider a deep neck infection in diabetic patients.\(^72\) \(\beta\)-Hemolytic streptococci or coagulase-positive staphylococci are the cause in 90% of patients with craniocervical necrotizing fasciitis. \(^31\) Sputum samples, obtained via tracheal suctioning, should be analyzed when infectious complications are suspected. Radiologic studies, namely, a chest radiograph, cervical computed tomography (CT), and chest CT, should be ordered as indicated.

Antimicrobial treatment in the ED should cover the most common organisms for the suspected site of tracheostomy-related infection. Broad-spectrum systemic antimicrobials, along with adjunct aerosol therapy, should be given urgently to high-risk patients.\(^74\) Depending on the patient's condition, surgical evaluation may be needed.\(^41\)

**Tracheal Stenosis and Tracheomalacia**

Tracheal stenosis and tracheomalacia are late complications of a tracheostomy, often occurring weeks to months after decannulation. Its overall incidence is unknown, and clinically significant stenosis has been estimated to develop in 10% of all tracheostomy patients.\(^75\) Tracheal stenosis occurs most often at the level of the stoma and can develop proximal (suprastomal) or distal to the stoma as a result of a poorly fitting tube or...
cause of a TEF is a poorly fitted tracheostomy tube or an overinflated cuff. Injury to the esophageal mucosa from a nasogastric or orogastric tube can also cause a TEF.

Most patients with a TEF have increased secretions, recurrent pneumonia, or aspiration of gastric contents while receiving mechanical ventilation. The most frequent sign of a TEF is cough after swallowing. A persistent cuff leak or abdominal distention may also indicate a TEF. The clinician may be able to auscultate breath sounds over the lung fields and the epigastrium simultaneously, but this finding is unreliable. Bronchoscopy, barium esophagography, or mediastinal CT can aid in making the diagnosis.

Interventions

Once a TEF is diagnosed, the immediate goal should be to reduce the amount of tracheal and pulmonary soilage by inflating a cuff below the level of the fistula. If the current tracheostomy tube is not long enough to allow placement of a cuff distal to the TEF, exchange it with a longer tracheostomy tube or an ET tube. An orogastric or nasogastric tube can also be inserted to prevent gastric contents from further contaminating the respiratory tract. Early consultation with an otolaryngologist or thoracic surgeon is appropriate because the definitive treatment is surgical. Antibiotics that target gram-negative, gram-positive, and anaerobic organisms should be initiated if there is a suspicion for pneumonia. TEF is not usually an immediately life-threatening complication, and tube exchange can be performed in conjunction with the surgical team.

Bleeding

Major Bleeding

Major bleeding is one of the most feared complications of tracheostomy. A history of bleeding or minor bleeding that has stopped spontaneously cannot automatically be attributed to minor irritation or skin erosion. It may not be possible to fully evaluate bleeding from a tracheostomy site in the ED without consultation or specialized equipment such as a fiberoptic endoscope. Sources of bleeding include the thyroid vessels, anterior jugular veins, brachiocephalic (innominate) artery, carotid artery, and aortic arch. Bleeding from esophageal or gastric sources may occur if a TEF is present...
or if the patient has aspirated blood. Erosion of a major vessel from the cuff or tip of the tube is responsible for 10% of all tracheostomy hemorrhages and is a devastating complication. The IA is the vessel most commonly involved.\(^80\)

A tracheoinnominate artery fistula (TIF) is a late complication of tracheostomy that usually occurs within the first 4 weeks after insertion, but it can happen at any time.\(^12,81\) The mortality rate associated with a TIF approaches 100%.\(^82\) The anatomic proximity of the IA to the trachea places it at higher risk for fistulization if the anterior tracheal wall is injured. The vessel crosses from left to right as it moves superiorly and lies immediately anterior to the trachea at the level of the superior thoracic inlet. Risk factors for TIF include placement of the stoma below the third tracheal ring, caudal migration of the tracheal tube, and the presence of a cephalad-coursing IA.

A single episode of hemoptysis or tracheal bleeding may be the only warning sign of a TIF. Any amount of bleeding or hemoptysis exceeding 10 mL within 48 hours of insertion of the tube should be considered a “sentinel bleed” and an indication that a fatal hemorrhage may be imminent. Some patients report only a new cough or retrosternal pain.\(^83\) Presume that a history or evidence of 10 mL or more of blood is from an arterial source.

**Preparation**

When evaluating a patient with a suspected TIF, place advanced airway equipment at the bedside. Emergency surgical consultation is mandatory. Position the patient with the head elevated and the neck in slight extension. Secure adequate intravenous access, and prepare the patient to go to the operating room. In addition to the equipment noted in Fig. 7.2, have a scalpel with a No. 10 or 11 blade and a 50-mL syringe at the bedside.

**Interventions**

If the patient is stable, an attempt to visualize the bleeding site with direct visualization using a fiberoptic scope should be performed. Look for evidence of bleeding on the anterior tracheal wall at or below the sternal notch. If significant tracheal bleeding or a clot is present, first hyperinflate the tracheostomy tube cuff with the 50-mL syringe to compress the artery against the posterior sternal wall. Inflate the balloon slowly to prevent rupture of the cuff. Depending on the make and model of the tube, inflating the cuff with the entire 50 mL may not be possible. If a TIF has been caused by cuff erosion, this procedure should tamponade the bleeding. If the patient’s tracheostomy tube does not have a cuff, replace it with a cuffed ET or tracheostomy tube.

If cuff hyperinflation is unsuccessful, the TIF may be located at or beyond the distal tip of the tracheostomy tube. In this circumstance, the current tracheostomy tube must be removed and replaced with either an oral ET tube or an ET tube inserted through the stoma.

Position the cuff of the ET tube below the stoma at the level of the upper part of the sternum and hyperinflate it. If the patient continues to bleed despite this maneuver, apply digital pressure through the tracheal stoma to compress the anterior tracheal wall against the sternum. Digital pressure is considered the most reliable technique to stop hemorrhage and can provide control of bleeding during transport to the operating room (Fig. 7.11).\(^41\) Extension of the stoma with a vertical incision to the jugular notch may be necessary if the provider cannot reach the TIF through the original stoma. The clinician should be careful not to damage surrounding vasculature when making the incision.

**Minor Bleeding**

All bleeding from a tracheostomy site must be evaluated for a potentially life-threatening event. Seemingly minor or self-limited bleeding may be a harbinger of subsequent severe hemorrhage. Sentinel hemorrhages frequently precede massive bleeding. Consider endoscopic examination for complete evaluation unless a superficial bleeding site is confirmed.

Minor bleeding is most likely the result of irritated granulation tissue and is usually confined to the skin surrounding the stoma. Bloody secretions from the tracheostomy tube may represent tracheitis, bleeding running down from the skin or thyroid, or superficial tracheal ulceration from tracheal suctioning or tracheal tube pressure.

Examine the stoma site and tube first in an attempt to locate the source and quantify the volume of blood loss. If the source of bleeding is within the stoma or from within the trachea, remove the tracheostomy tube if it was placed more than 7 days before the current event. Visualize the tracheal lumen, proximal end of the trachea, and inner stoma with a nasopharyngoscope or a small pediatric laryngoscope. It is important to differentiate superficial erosions from active bleeding. Do not attempt to remove clots in the trachea because this may increase the rate of hemorrhage. Consider obtaining a basic metabolic panel, complete blood count, and coagulation studies.
to evaluate for other factors complicating bleeding, such as uremia, thrombocytopenia, or coagulopathy.

**Preparation**

Prepare patients with minor tracheostomy bleeding in the same way that you would those with major bleeding. Gather the appropriate airway equipment at the bedside. In addition to the standard equipment shown in Fig. 7.2, obtain the following:

- Sterile gauze
- Sterile saline for irrigation
- 22-gauge needle with syringe
- 1% lidocaine with 1:100,000 epinephrine
- Absorbable hemostat (e.g., Surgicel, Ethicon, Cincinnati, OH)
- Suture kit with suturing material
- Electrocauterization or chemical cauterization supplies, if available

**Interventions**

For external or stomal bleeding, begin with local irrigation to find the source of the bleeding. Most incisional or stomal bleeding can be stopped by applying direct pressure for 3 to 5 minutes. Application of absorbable hemostatic material may improve the outcome of direct pressure application. If an external bleeding site continues to ooze, consider adjunctive treatment. Options include injecting 0.5 to 1.0 mL of lidocaine with epinephrine near the source, placing a single suture for hemostasis, or using cauterization. Last, replace the tracheostomy tube. Following tube replacement, suction carefully to confirm resolution of the bleeding and to identify secondary sources of bleeding.

If stomal bleeding or intratracheal sites do not account for the bleeding, consider other causes. Placement of a nasogastric tube will help in the identification of gastrointestinal bleeding. Examine the nasopharynx and oropharynx for bleeding sources. If the patient has undergone radiation therapy, examine the area above the level of the tracheostomy stoma, where mucosal injury secondary to radiation damage may be the cause of blood in the tracheal secretions.

**TRANSESOPHAGEAL PUNCTURE FOR VOICE RESTORATION**

TRANSESOPHAGEAL PUNCTURE FOR VOICE RESTORATION

Transepophageal puncture (TEP) has become one of the most widely used and accepted techniques for voice rehabilitation. Developed in the 1980s, it can be performed as a primary or secondary procedure after laryngectomy or other pharyngeal surgeries. A puncture site is created through the anterior esophagus and posterior tracheal wall, where the TEP prosthesis is later inserted. The mucosa in segments of the pharyngeal esophagus vibrates in response to airflow, thereby creating speech.

**Complications**

Operative and immediate postoperative complications of TEP are infrequent.84 Long-term complications include stomal stenosis, aspiration of the prosthesis, fistula leakage, TEP necrosis, and swallowing impairment. Reported infectious complications associated with TEP include deep neck abscess, aspiration pneumonia, and cervical cellulitis.85

The emergency care provider should be aware of the most common complications, few of which are life threatening. Accumulation of thick or inspissated secretions or food above the TEP can cause upper airway obstruction. Prosthesis dislodgement, occlusion, or erosion secondary to infection should be considered in all patients with acute changes in voice production or decreased ability to speak.86 Esophageal edema causing dysphagia and loss of TEP speech has been reported and should be differentiated from other causes of esophageal obstruction.

When evaluating patients with a TEP, review the medical records, assess airway and esophageal patency, and determine whether the patient has experienced changes or difficulty in voice production. In stable patients, management of prosthesis complications should be referred to a specialist, most commonly an otolaryngologist.

**TRANSTRACHEAL OXYGEN DELIVERY SYSTEMS**

Low-flow oxygen is prescribed for patients who have adequate ventilatory function but chronic hypoxia. Many patients with chronic obstructive pulmonary disease, pulmonary fibrosis, sleep apnea, lung cancer, and α1-antitrypsin deficiency are candidates for outpatient use of supplemental oxygen.87–89 Traditionally, supplemental oxygen has been delivered by nasal cannula. Although nasal cannula oxygenation is easy to administer, it has several side effects, including drying of the nasal mucosa, epistaxis, ear discomfort, contact dermatitis from the oxygen tubing, and dry throat.90–92 Use of a nasal cannula is inefficient because it delivers oxygen only during inspiration and the oxygen must traverse the anatomic dead space of the nares and hypopharynx.

Transtracheal oxygen (TTO) delivery systems enhance the efficiency of oxygenation by administering oxygen directly to the lower respiratory tract. These systems reduce complications, improve patient comfort, and increase compliance.90 Oxygen is delivered through all phases of the respiratory cycle and directly into the trachea, thereby bypassing dead spaces in the upper airway.93 As a result, the required oxygen flow rate is commonly reduced by at least 50%.94 Gas mixture in the distal end of the trachea is more effective in eliminating CO2. Clinically, TTO systems usually reduce the patient’s work of breathing and exertional dyspnea.95 Physiologic benefits include reduced erythrocytosis, decreased pulmonary vascular resistance, improved cor pulmonale, improved arterial oxygen tension, and increased exercise capacity.91

TTO catheters are small tubes that deliver oxygen directly to the lumen of the trachea. The catheter is held in place by a subcutaneous tract, and is inserted into the lower part of the trachea. Low-flow oxygen (2 to 10 L/min) is supplied directly to the trachea by a narrow (7- to 11-Fr) catheter. Typically, an 11-cm catheter sits in the trachea with its tip 1 to 2 cm above the carina.89 It can have a single or multiple distal ports for oxygen flow. The catheter is held in place by a thin band or necklace through two openings in the flange.

The surgical procedure is often done in an outpatient setting with the patient under local anesthesia. Initially, a small stent is placed percutaneously into the anterior aspect of the neck; it is then replaced with a TTO catheter in 1 to 2 weeks, after
the tract matures. Dislodgement of the catheter during this time can result in closure of the tract within a matter of minutes. Once the tracheocutaneous fistula has epithelialized, the catheter may be inserted. Early catheter changes should be done in the clinician's office via a modified Seldinger technique if the integrity of the stoma is questionable. After the tract has fully matured, most patients can change their catheter at home.

Regular maintenance includes cleaning and changing the TTO catheter. One milliliter of sterile saline is instilled into the catheter, and a cleaning rod is inserted as far as possible. The cleaning rod is inserted and extracted three times to remove secretions and encrustations from the lumen of the catheter. Catheters are usually changed according to the manufacturer's recommendations, from twice daily to once every 2 weeks. The stoma should be cleaned twice each day and inspected thoroughly for signs of infection. All patients should be given supplemental oxygen by nasal cannula during catheter maintenance procedures. Adequate humidification, cleaning, and systemic hydration will help reduce the incidence of mucous blockage.

Early complications (developing within 3 weeks after the procedure) occur in approximately 30% of patients and include bleeding, infection, pneumothorax, costochondritis, and dislodgement (which can be caused by coughing). Late complications occur in approximately 40% of patients and include mucous plugging, bleeding, infection, and hemoptysis. A mucous ball is an accumulation of impregnated mucus that adheres to the outer surface of the TTO catheter tip. It can cause coughing, wheezing, and dyspnea. Life-threatening airway obstruction resulting from the formation of a large mucous ball has been reported.

Dyspnea or increased coughing may indicate that the TTO catheter is obstructed by a mucous ball, that the tube is kinked, or that the tip of the catheter is positioned cephalad to the stoma. Obstruction within the catheter tubing may cause a whistling sound from the oxygen tank humidifier. Always examine the patient for signs of subcutaneous air and catheter dislodgement.

If routine broad-spectrum antibiotics are used by the patient, Candida infections can develop at the stoma. Such infections are more common in patients receiving systemic antibiotics or long-term corticosteroids or those with diabetes. Tracheal chondritis may result from bacterial infection of the cartilage. Many patients with chondritis have a deep, indurated, nonfluctuant lump around the tract that may be tender to palpation. A 3-week course of an oral antibiotic that specifically covers S. aureus should be prescribed as treatment.

Interventions

If obstruction is suspected while the patient is in the ED, clean and replace the catheter. If the stoma tract has not healed or appears to be infected, change the catheter via a modified Seldinger technique. Use a water-soluble lubricant for the catheter change. If changing the catheter does not relieve the obstruction and the patient's airway is intact, encourage the patient to perform maneuvers that raise intrathoracic pressure to increase the force of the cough. Ask the patient to sit upright, hold a pillow to the abdomen, and cough forcefully after three deep inspirations. This maneuver may help mobilize secretions or small mucous plugs in the airway. Dyspnea and coughing should lessen with effective removal of the obstruction.

Manage minor bleeding at the catheter site with gauze packing or cautery. If significant bleeding is identified or suspected, consult a specialist on an emergency basis and manage the airway definitively as clinically indicated (see the section on Major Bleeding). Manage skin and pulmonary infections with the techniques discussed for tracheostomy care.

Stents

Tracheal stenosis and tracheomalacia are known complications of artificial airways. Management options include surgery and placement of silicone stents in the trachea. Patients who have had their tracheostomy tubes successfully decannulated may need stents if symptomatic stenosis or tracheomalacia occurs. Indications for and complications of tracheal stents are beyond the scope of this chapter, but the clinician should be aware of the possibility that these indwelling devices may be present in patients who have undergone head and neck surgery or previous tracheostomies.

SPECIAL POPULATIONS

Obese Patients

The prevalence of obesity (body mass index [BMI] ≥30) and morbid obesity (BMI ≥40) has increased dramatically over the past few decades. Obesity increases the patient's risk for cardiovascular and metabolic comorbidity, and obese patients have a higher risk for ventilatory dysfunction secondary to altered respiratory mechanics. Lung compliance, functional residual capacity, and expiratory reserve volume in an obese patient are reduced exponentially in relation to BMI. As weight increases, vital capacity and total lung capacity decrease. Ultimately, obese patients have decreased pulmonary reserve, which can cause a rapid onset of hypoxia if they become critically ill.

Morbidly obese patients are particularly at risk for life-threatening complications related to tracheostomy. Tubo obstruction and accidental dislodgement appear to be more common in obese patients. Dislodgement is specifically associated with increased rates of morbidity and mortality. Delayed recognition of tube dislodgement because of abnormal neck anatomy accounts for nearly 30% of tracheostomy-related deaths in the obese population.

Preparation

For the management of an obese patient with a tracheostomy-related complaint, it is important to have all advanced airway equipment at the bedside (see Fig. 7.2). In addition to standard equipment, attempt to obtain an extra long or adjustable-flange tracheostomy tube to span the elongated pretracheal distance that is common in morbidly obese patients. In obese patients, standard tracheostomy tubes may be too short proximal to the cuff and have a higher risk for malposition. In addition to standard monitoring, use continuous capnography, if available, to prevent delay in the recognition of tube dislodgement. Physical signs, such as reduced breath sounds and subcutaneous air, may be more difficult to recognize in obese patients. Fiberoptic bronchoscopy may be needed to confirm correct tube placement. The patient should be positioned with the neck in slight extension, and the head of the bed should be elevated to approximately 30 degrees.
Interventions
Approach an obese patient with the same protocols outlined previously in this chapter for the applicable complication. If the tube is obstructed, deflation of the tracheostomy cuff may not be sufficient to allow adequate ventilation because external compression caused by abnormal neck anatomy may occur. If tube dislodgement is suspected, ET intubation may be preferred over blind reinsertion because of the increased risk for false passage. Tube placement can be confirmed with direct visualization via fiberoptic bronchoscopy.

Laryngectomy Patients
It is important to distinguish the patient with a routine tracheostomy from the patient with a laryngectomy as this could potentially limit the ability to provide ventilation by way of the upper airway. A patient who has undergone a total laryngectomy no longer has an upper airway that is in continuity with the lungs. A patient with a partial laryngectomy will have a contiguous upper and lower airway, and interventions by way of the upper airway may be limited.

The basic principles of emergency tracheostomy care still apply to the laryngectomy patient, and it is critical to recognize that a patient with a total laryngectomy cannot be resuscitated with traditional oxygenation and ventilation techniques, such as oropharyngeal bag-valve-mask ventilation and face mask oxygenation. Interventions should be made by way of the stoma rather than the upper airway. Partial laryngectomy patients should also be ventilated through their stoma, and their mouth and nose should be kept closed to avoid air escape.

Interventions
Should a patient with a laryngectomy need positive pressure ventilation, there are a few important steps to consider prior to inserting a cuffed tracheostomy or ET tube through the stoma. First, it is important to inspect the laryngectomy stoma for any mechanical obstruction. Suction the stoma to make sure it is clear of excessive secretions or mucous plugs. Stoma buttons should be removed, whereas tracheoesophageal speaking valves may be left in place as long as they are not displaced.

External ventilation with a pediatric face mask or laryngeal mask airway can be performed. A standard cuffed-tracheostomy tube may be placed to facilitate mechanical ventilation. If inserting an ET tube, it is important that the provider avoid advancing it too far as this can lead to main stem bronchus intubation. The approximate distance between the skin of the stoma and carina is only approximately 6 cm.

Equipment
When replacing a tracheostomy tube in a child, always have at least two tracheostomy tubes available: the current size and a size smaller. Pediatric tubes generally have a much smaller diameter than adult tracheostomy tubes, and many of them do not have an inner cannula. Pediatric tracheostomy tubes rarely have inflatable cuffs, except for those used for certain special indications. Suction catheters, a bag-valve-mask device, ET tubes, resuscitation medication, and equipment appropriate for the pediatric population should be available when treating pediatric tracheostomy patients. Continuous capnography may detect certain tracheostomy complications and is recommended.

Sizing
Pediatric tracheostomy tubes share most of the same components of the tubes used in adults. The ID of the tube is stamped on the outer cannula flange, and that information should guide the clinician’s choice of replacement tubes. Recommended tube sizes according to age are listed in Table 7.2. Age guidelines can be helpful, but they may not be reliable in pediatric patients because of complex medical problems. Premature infants may be small for their age and weight, thus making estimation of tube size even more difficult.

Like ET tubes, tracheostomy tubes are sized by the tube’s ID. When the tracheostomy tube is seated correctly in a pediatric patient, it should extend at least 2 cm beyond the stoma and no closer than 1 to 2 cm above the carina. Its curvature should be such that when the tube is placed appropriately, the distal portion of the tube is concentric and collinear with the trachea. After replacing a tracheostomy tube, confirm its position by auscultating breath sounds, using capnography, and confirming the location of the distal tip with a chest radiograph.

Cuff
The general rule that cuffed ET tubes should not be used in patients younger than 6 years is under a great deal of debate and does not universally apply to tracheostomy patients. If a

<table>
<thead>
<tr>
<th>AGE</th>
<th>SIZE</th>
<th>INNER DIAMETER (mm)</th>
<th>OUTER DIAMETER (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature</td>
<td>00</td>
<td>3.1</td>
<td>4.5</td>
</tr>
<tr>
<td>Newborn–3 mo</td>
<td>0</td>
<td>3.4</td>
<td>5.0</td>
</tr>
<tr>
<td>3–10 mo</td>
<td>1</td>
<td>3.7</td>
<td>5.5</td>
</tr>
<tr>
<td>10–12 mo</td>
<td>2</td>
<td>4.1</td>
<td>6.0</td>
</tr>
<tr>
<td>13–24 mo</td>
<td>3</td>
<td>4.8</td>
<td>7.0</td>
</tr>
<tr>
<td>2–9 yr</td>
<td>4</td>
<td>5.0</td>
<td>8.5</td>
</tr>
<tr>
<td>10–11 yr</td>
<td>6</td>
<td>7.0</td>
<td>10</td>
</tr>
<tr>
<td>≥12 yr</td>
<td>8</td>
<td>8.5</td>
<td>12.0</td>
</tr>
</tbody>
</table>

patient requires high-pressure ventilation or only nocturnal ventilation or is at risk for aspiration, a cuffed tube may be appropriate. Cuffed tubes are also used in patients with a tracheal anomaly. Replacement of these tubes should ensure that the individual’s anatomic and physiologic needs are met.

Cuff pressure recommendations for pediatric patients are less than 20 cm H$_2$O. With few exceptions, low-pressure, high-volume cuffs should be used.

**Humidifiers**

Humidifiers for pediatric tracheostomy tubes attach to the external port. Some humidifiers have lithium-coated moisture exchangers. Systemic absorption of lithium is unlikely to cause clinical symptoms in adults but may be a consideration in children. Use humidifiers regularly to keep secretions loose and help prevent obstruction of the tube.

**Suctioning**

Suctioning recommendations in pediatric patients clearly support the use of a premeasured suction catheter to reduce the rate of mucosal irritation and to limit the development of granulation tissue. The premeasured technique uses an exact depth of insertion, which reduces epithelial damage if the catheter is inserted too deeply and inadequate suctioning if the catheter is not inserted deeply enough. Depth of insertion can be estimated by measuring a similar tube before inserting the suction catheter. In children with fenestrated tracheostomy tubes, suction catheters may accidentally go through the fenestrations and cause mucosal irritation. If this happens repeatedly, granulation tissue may develop at the site.

**Complications**

Pediatric tracheostomy complications are similar to those in the adult population. Their incidence is estimated to be between 5% and 49%. Late complications most likely to be seen in the ED include obstruction, dislodgement, bleeding, pneumothorax, infection, pneumomediastinum, TEF, and TIF. Chronic respiratory complications include tracheomalacia, tracheal stenosis, vocal cord paralysis, and vocal cord fusion.

Granuloma formation is common in pediatric patients. It often occurs in the tracheal lumen either at the superior margin of the tracheostomy or at the level of the tip of the tube. The development of granulation tissue is the result of persistent mucosal irritation and inflammation. Any patient with a clinically significant granuloma should be evaluated by a pediatric specialist for definitive care.

Pneumomediastinum and pneumothorax are generally thought to be early complications of tracheostomy, but they should always be considered in a pediatric tracheostomy patient. In children, the pleural apices rise higher than in adults and can even extend into the lower part of the neck. These complications can be caused by a dislodged tube positioned in a false passage or a malpositioned tube that causes an increase in intrathoracic pressure.

The most common bacterial species that colonize pediatric tracheostomies are *P. aeruginosa* and *S. aureus*. As in adults, colonization does not require treatment unless signs of acute infection are present. Suprastomal collapse of the anterior tracheal wall is very common in pediatric patients and can cause air trapping. A tracheostomy tube that places excessive pressure on the tracheal rings will cause inflammation, chondritis, and weakening of the cartilaginous rings. Significant collapse can hamper subsequent decannulation success.

Management of major bleeding in pediatric patients follows the same recommendations given for adults. The smaller stoma size may prevent the application of digital pressure to the site of hemorrhage. If the clinician is unable to reach the source of the bleeding, expeditious ET intubation followed by cuff overinflation may temporize the bleeding until the patient can be taken to the operating room.

**CONCLUSION**

A tracheostomy emergency can be one of the most high-risk scenarios in the ED. The emergency physician must be well prepared for a variety of tracheostomy complications and should approach the patient in an organized, stepwise fashion so that interventions can be initiated in a timely fashion. Often specialty consultation in the ED is required to fully assess or treat complications. Tube dislodgement and obstruction are common and associated with an extremely high mortality rate, especially in obese patients. Always ensure correct tube position, have backup airway equipment readily available, and obtain surgical support early if the patient needs to go to the operating room.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
64. Park DR: The microbiology of ventilator-associated pneumonia. Respir Care 50:742, 2005.
Mechanical Ventilation
Jairo I. Santanilla

INTRODUCTION
Mechanical ventilation (MV) is an important tool for resuscitation of critically ill patients in the emergency department (ED). It is vital that ED practitioners have a thorough understanding of the basics of MV and to know when to apply these principles and how to support patients in respiratory or cardiac failure. Hospital overcrowding has led to a delay in transfer of mechanically ventilated patients out of the ED, and ventilator management often falls on the emergency medicine physician. In addition, during nights and weekends in some facilities, the ED physician may be called on to troubleshoot or stabilize mechanically ventilated patients in the intensive care unit (ICU). The traditional view of MV as a prescription that fits virtually all patients equally should be discarded as a gross misunderstanding of pulmonary pathophysiology. Increasing evidence has shown that the mechanism of lung ventilation by MV may be as deleterious as it is helpful.1 Because patients remain in the ED while mechanically ventilated, ED clinicians should embrace the established paradigm of pulmonary-protective MV strategies as a cornerstone of care.

BASIC PHYSIOLOGY
Understanding basic pulmonary physiology is essential to understanding how to initiate MV. This ensures that the method of gas delivery meshes with the patient’s underlying physiology to avoid ventilator-induced lung injury.

Minute Volume and Alveolar Ventilation
The volume of air that moves in and out of a patient’s lungs per minute is termed minute volume (\(V_E\)). \(V_E\) is the product of tidal volume (\(V_T\)) and respiratory frequency or rate (\(f\)):

\[ V_E = V_T \times f \]

Normal \(V_E\) is 7 to 10 L/min. \(V_T\) can be further broken down into alveolar volume (\(V_A\)) and dead space volume (\(V_{DS}\)):

\[ V_T = V_A + V_{DS} \]

In healthy young persons, the anatomic dead space is accounted for by the trachea and the larger airways and is approximately 2.2 mL/kg lean body weight. In disease states, in addition to the anatomic dead space, there is a variable amount of “pathologic” dead space, which corresponds to ventilated alveoli and respiratory bronchioles that are not adequately perfused. The sum of anatomic and pathologic dead space is often referred to as physiologic dead space.

Alveolar minute ventilation (\(V_{A}\)) is the product of rate times \(V_T\) minus dead space:

\[ V_A = (V_T - V_{DS}) \times f \]

\(V_A\) and the rate of \(\text{CO}_2\) production by the body determine the partial pressure of \(\text{CO}_2\) in the alveoli (\(P_{VCO2}\)), which is approximately equal to systemic arterial \(\text{CO}_2\) tension.

Volume-Pressure Relationship
Volume and pressure are related for a given respiratory system. A given volume (\(V\)) will create a certain pressure (\(P\)) relative to the compliance (\(C\)) of the respiratory system. The respiratory system consists of the ventilator tubing, endotracheal (ET) tube, trachea, airways, lung parenchyma, chest wall, and diaphragm tension. For example, a 500-mL volume will create a certain pressure based on the compliance of the respiratory system. Increasing the volume will increase the pressure in the system. Decreasing the volume will result in lower pressure. Decreasing the compliance (i.e., making the system “stiffer”) will increase the pressure in the system. Increasing the compliance will decrease the pressure in the system.

\[ P = V/C \]

Conversely, this relationship also holds for volume. For example, a pressure of 20 cm \(\text{H}_2\text{O}\) will create a certain volume based on the compliance of the system. Increasing pressure will result in higher volume. Decreasing pressure will result in lower volume. Decreasing compliance will result in lower volume. Increasing compliance will result in higher volume.

\[ V = P \times C \]

Airway Pressures
Plateau Pressure
Plateau pressure is measured at the end of inspiration with a short breath-hold (Fig. 8.1). At this point no airflow should be occurring. This is considered a static pressure. By understanding the aforementioned volume-pressure relationship, one can easily deduce how plateau pressure is inversely related to respiratory system compliance and directly related to volume. Anything that decreases compliance will increase plateau pressure. Increasing compliance will decrease plateau pressure. Decreasing volume will decrease plateau pressure (a major tenet in lung-protective ventilation).

Peak Airway Pressure
Peak airway pressure is derived during inspiration and thus incorporates airflow. Because there is air movement during this measurement, it is considered a dynamic pressure. It reflects the dynamic compliance of the entire respiratory system and incorporates static compliance and airflow. Peak airway pressure can never be lower than plateau pressure. In addition, because its main distinction is that it incorporates airflow, it is reflective of resistance to airflow. Anything that decreases compliance or increases resistance to airflow will increase peak airway pressure. Increasing compliance or decreasing resistance to airflow will decrease peak airway pressure.

A physiologically appropriate means of detecting and monitoring bronchospasm is the peak-plateau gradient. A normal gradient is less than 4 cm \(\text{H}_2\text{O}\) pressure, and elevated values indicate increased airflow resistance. The efficacy of treatment with \(\beta_2\)-agonists, steroids, intravenous magnesium, or diuresis may be assessed by monitoring the changes in this gradient (Fig. 8.2).
Positive End-Expiratory Pressure

Positive end-expiratory pressure (PEEP) is the pressure in the airway at the end of exhalation. PEEP helps keep the large noncartilaginously supported airways and the smaller alveoli open to prevent collapse, atelectasis, and ensuing hypoxia. The ventilation required to compensate for this triad commonly worsens lung compliance and is associated with ventilator-induced lung injury. Progressive increases in PEEP result in elevations in both total lung pressure and total lung volume. For example, serial elevations in PEEP often result in increased plateau pressure and elevated functional residual capacity (FRC; i.e., lung volume).

Extrinsic PEEP

When discussing MV and PEEP, most often authors are referring to extrinsic PEEP (PEEP). This is also referred to as applied PEEP. It is the PEEP that is extrinsically applied by the ventilator. When PEEP is used without a subscript in this chapter, it refers to PEEP. The useful PEEP range is from 3 to 20 cm H₂O. PEEP is used to increase FRC and move the zero pressure point of each alveolar unit more proximally in the airway and thereby prevents early alveolar collapse. By so doing, PEEP increases the available number of alveolar units that can participate in gas exchange. The primary effect of PEEP on gas exchange is improvement in oxygenation, not removal of CO₂. CO₂ clearance is rather efficient and will be well preserved, even in hypoxic situations. By opening one alveolar unit, the tendency of adjacent units is to open as well (i.e., alveolar codependency) (Fig. 8.3). Excessive PEEP will compromise hemodynamics. There are two primary questions to ask when using PEEP to augment oxygenation: (1) What is the “optimal PEEP?” and (2) Is the current amount of PEEP compromising the patient’s hemodynamics?

PEEP is not without untoward side effects, and increased levels of PEEP can lead to lung injury and hemodynamic compromise. Increased intrathoracic pressure can result in cardiac compression and collapse, principally of the right atrium. It is imperative that the patient be adequately volume-resuscitated because preload depletion compounds this problem. Desired levels of PEEP simply may not be possible because of deleterious effects on cardiac output.

Optimal PEEP can be determined in several ways. One is to increase PEEP until there are no longer increases in partial pressure of oxygen (PO₂). However, this method may result in several untoward events. First, oxygen tension may increase steadily, but carbon dioxide pressure (PCO₂) may increase as a result of alveolar overdistention. With overdistention, alveolar pressure may exceed pulmonary arteriolar pressure and actually decrease pulmonary blood flow and clearance of CO₂. Second, alveolar overdistention may increase total intrathoracic pressure and result in diminished venous return and cardiac output. Third, decreased venous return may cause cerebral venous hypertension. The optimal PEEP for one organ system may be deleterious for another. For example, the optimal PEEP for ideal oxygenation may be the worst PEEP for cerebral venous drainage.

An alternative is to increase PEEP until a complication of PEEP occurs (e.g., elevation in PCO₂, hypotension) and then reduce PEEP if needed (inability to tolerate hypercapnia) or expand the patient’s intravascular volume to combat the decreased venous return.

Another excellent method of determining the optimal PEEP is guided by assessing changes in plateau pressure with changes
upstroke of the loop because the vast majority of the functional lung units are already open and ready to be ventilated (see Fig. 8.4). This strategy is known as the open lung model of MV.6 Irrespective of what technique is used, it is currently widely agreed that plateau pressure should not exceed 30 cm H₂O. If respiratory system compliance is so low that plateau pressure exceeds 30 cm H₂O, either PEEP or VT has to be decreased. If this is not possible because of either recalcitrant hypoxia or acidosis, rescue therapies may need to be used (see the section on Acute Lung Injury and Acute Respiratory Distress Syndrome).

Intrinsic PEEP
Intrinsic PEEP (PEEPi) is additional pressure that is generated within the airways from trapped gas that should have been exhaled but for various reasons (commonly obstruction to exhalation such as in chronic obstructive pulmonary disease [COPD]) was not. PEEPi is also referred to as auto-PEEP, dynamic hyperinflation, and breath stacking. For the remainder of this chapter it will be referred to as PEEPi.

PEEPi can cause hemodynamic instability secondary to decreased venous return, just like high levels of PEEP.7 PEEPi may be detected in two ways: (1) evaluation of the flow-time trace or (2) disconnection of the patient from the ventilator and listening for additional exhaled gas after an exhalation should have occurred.6 The flow-time trace will demonstrate that the exhalation is not yet completed before the next breath has been initiated (Fig. 8.6).8

INDICATIONS FOR MV
There are wide-ranging reasons for patients to require MV in the ED, and there are no absolute contraindications. Many time-honored indications for invasive ventilation are now identified as appropriate indications for noninvasive ventilation and are addressed later. Indications for MV range from loss of airway anatomy (edema, direct or indirect trauma, burns, infection), loss of protective airway mechanisms (intoxicants, brain injury, stroke), inability to ventilate, inability to oxygenate, or the expected clinical course. Indications for ET intubation may be separated into several categories—emergency, urgent, delayed, and elective—based on the urgency of establishing a definitive airway.
EQUIPMENT—STANDARD OPTIONS

Perhaps one of the most confusing aspects of MV is the plethora of terms and acronyms that are used. Understanding the basic terminology helps clarify this subject. The following discussion explores machine features and settings. Regardless of which ventilator is used, a limited number of standard features are common to each (Fig. 8.7).

**Set Respiratory Rate**

Most ventilators allow the clinician to set a respiratory rate. The respiratory rate and actual VT determine a patient’s minute ventilation. Patients intubated for airway protection because of trauma or toxicosis often do well with a normal minute ventilation. Initially setting the respiratory rate at 10 to 14 breaths/min and VT at 7 to 8 mL/kg ideal body weight (IBW) is usually sufficient. Adjustments can be made based on arterial blood gas (ABG) analysis, end-tidal CO₂, or venous blood gas and pulse oximetry. Patients who are septic or have severe acidosis often require higher minute ventilation. Respiratory rates can be increased, as can VT, but volumes higher than 10 mL/kg IBW should not be used because of the risk of inducing ventilator-associated lung injury. In special scenarios such as acute lung injury (ALI) and acute respiratory distress syndrome (ARDS), initial VT values should be lowered to 6 mL/kg IBW within 2 hours of intubation. Some of these specific scenarios are discussed later.

**Fraction of Inspired Oxygen**

All ventilators can deliver an adjustable fraction of inspired oxygen (FIO₂). Recommendations are to set it initially at 1.0 because the act of transitioning from negative pressure ventilation (normal physiologic breathing) to positive pressure ventilation (PPV) may unpredictably alter ventilation-perfusion (V/Q) matching. Although initially an FIO₂ of 100% is optimal, it is beneficial to quickly titrate FIO₂ down because of the theoretical risk for oxygen toxicity. Make adjustments based on ABG analysis or pulse oximetry, with a goal of keeping arterial PO₂ higher than 60 mm Hg or arterial oxygen saturation (SaO₂) at 88% to 92% to avoid potential oxygen toxicity (see Table 3.3 in Chapter 3). Such adjustments may best be accomplished in the ICU rather than the ED, after the entire clinical scenario can be analyzed and all interventions are appropriately adjusted. However, if there is a delay in transfer to the ICU, adjustments should be made in the ED.

**PEEP**

PEEP, is typically set at 5 to 8 cm H₂O. Most patients should be started at a PEEP of 5 cm H₂O, which is considered a physiologic level. It is used to offset the gradual loss of FRC in supine, mechanically ventilated patients. PEEP can be increased by 2 cm H₂O every 10 to 15 minutes as needed or tolerated by patients who remain hypoxic. The initial goal is to reduce FIO₂ to nontoxic levels. This goal is coming under increasing scrutiny as new information challenges the time frame and concept of O₂-induced lung injury at FIO₂ levels greater than 0.6. Exercise care when using PEEP levels higher than 8 cm H₂O in the setting of elevated intracranial pressure (ICP), unilateral lung processes, hypotension, hypovolemia, or pulmonary embolism. High PEEP can potentially lead to hypotension as it increases intrathoracic pressure and decreases venous return and, subsequently, cardiac output.

**Flow Rate**

The flow rate is the speed, in liters per minute, that the ventilator is delivering gas. It is found in volume-targeted modes, but not pressure-targeted modes. The flow rate is typically set at 60 L/min, which means that a set VT will be delivered at that speed. Patients wanting higher flow rates will not receive them and may display air hunger. Increase flow rates to deliver the set volume faster and shorten the inspiratory time (thereby increasing the inspiratory-to-expiratory [I/E] time ratio).

**Waveform**

The waveform determines how the ventilator delivers the flow of gas. It is traditionally set to a “decelerating waveform” in an effort to optimize recruitment because of different time constants in the lung.

**Decelerating (Ramp)**

Once the maximal inspiratory flow is reached, the rate of gas delivery immediately begins to slow in a preprogrammed fashion. When compared with the square waveform, longer time is spent in inhalation to deliver the set VT or achieve the target pressure and allow improved oxygenation. This waveform also achieves lower peak airway pressure and higher mean airway pressure.
**Square**

Once the maximal inspiratory flow rate (IFR) is achieved, gas flow is constant until the set volume is delivered. When that point is reached, gas flow is terminated. This waveform is best for patients with asthma, COPD, and head injury because gas delivered with this waveform allows a longer expiratory time (Te, thereby increasing the I/E ratio) and lower mean airway pressure. The longer Te is beneficial for patients with restrictive airway disease such as asthma or COPD. Spending a longer time in exhalation allows improved venous drainage from the brain and greater loss of trapped VT. The drawback with this waveform is increased peak airway pressure, which often requires lower VT. This can lead to inadequate alveolar recruitment in patients with ALI.

**I/E Time Ratio**

The ratio of inspiratory time (the time that it takes to take a breath) to expiratory time (the time that it takes to exhale) is automatically reported in some modes, whereas in others it is dialed in.

The normal I/E ratio in a spontaneously breathing, non-intubated patient is 1:4.\(^{11}\) Intubated patients commonly achieve I/E ratios of 1:2. Shorter ratios may lead to decreased exhalation by compromising Te. In its extreme form, inverse ratio ventilation (IRV), the normal pattern of breathing is reversed. A longer time is spent in inhalation to allow more time for oxygenation and recruitment. The decrease in Te can lead to air trapping, elevated mean airway pressure, and rising P\(_{CO_2}\). These problems lead to hypercapnia, respiratory acidosis, and PEEPi.\(^{12}\) (Fig. 8.8).

**Trigger**

The trigger is the aspect that initiates a machine-generated breath. The trigger can be set to detect either a pressure or a flow gradient. It should be set so that the patient can trigger the ventilator without great effort.

**Sensitivity**

This refers to the sensitivity of the trigger. If the trigger is set too high (not sensitive enough), the work of breathing incurred by the patient can be substantial. Some providers have been known to set the sensitivity at a high level if the patient is markedly overbreathing the set rate. This is not recommended because it causes an undue increase in the work of breathing. Many ventilators are set to a pressure trigger with a sensitivity of 1 to 3 cm\(\text{H}_2\text{O}.\)^{11} If the sensitivity is set too low (too sensitive), the ventilator can “auto-trigger” (inappropriate initiation of machine-generated breaths) because of oscillating water in the ventilator tubing, hyperdynamic heartbeats, or patient movement.

**MODES OF VENTILATION**

Once some of the standard features are understood, the next step is determining the ventilator’s target. Most ventilators can be set to achieve spontaneous breathing, volume-targeted ventilation, pressure-targeted ventilation, or some combination. In volume-targeted ventilation, the ventilator is set to reach a determined volume regardless of the pressure required to do so. Pressure-targeted modes are set to reach a determined pressure regardless of the volume generated. Dual modes combine the benefits of both strategies (Fig. 8.9).

**Spontaneous Breathing**

Spontaneously breathing patients can be supported on the ventilator by pressure support ventilation (PSV). In this mode, the ventilator provides a supplemental inspiratory pressure to each of the patient-generated breaths. The clinician sets F\(_{IO_2}\) and PEEPi. The patient dictates the respiratory rate and generates the desired flow rate. The applied pressure is turned off once the flow decreases to a predetermined percentage. VT is dictated by the pressure support given, patient effort,

---

**Figure 8.8** Effect of flow rate on inspiratory (T\(_i\)) and expiratory (T\(_e\)) time. Note that as the flow rate changes, there are corresponding alterations in the effective times for inspiration and exhalation. Deflections above the x-axis (time) indicate inspiration, and those below indicate exhalation. The tidal volume (VT) delivered for each cycle is the same, but T\(_i\) and T\(_e\) are different.

**Figure 8.9** Initial ventilator settings for standard patients. AC, Assist/control mode; F\(_{IO_2}\), fraction of inspired oxygen; IBW, ideal body weight; PEEPi, positive end-expiratory pressure; PCV, pressure-cycled ventilation; PSV, pressure support ventilation; RR, respiratory rate; SIMV, synchronized intermittent mandatory ventilation; VCV, volume-cycled ventilation; \(V_T\), tidal volume.
and compliance of the respiratory system. There is no set respiratory rate, although most modern ventilators have a backup apnea rate.

**Volume-Cycled Ventilation**

Volume-cycled ventilation (VCV) may also be termed volume-limited, volume-control, volume-assist, or volume-targeted ventilation. Volume-targeted modes are the most commonly used and the most familiar mode of MV in adults. As its name implies, “volume”—in this case VT—is the ventilator’s targeted parameter. With this target, the ventilator determines the base ventilation rate, but if the patient tries to breathe faster than the set rate, additional breaths can be initiated by the patient, known as spontaneous breaths (i.e., assist breaths). A potential downside is inappropriate hyperventilation.

**Pressure-Cycled Ventilation**

Pressure-cycled ventilation (PCV) may also be termed pressure-limited, pressure-control, pressure-assist, or pressure-targeted ventilation. As its name implies, “pressure” is the ventilator’s targeted parameter. The ventilator will generate an inspiratory pressure that has been set by the clinician. With this target, the ventilator alters gas flow to achieve and maintain a preset airway pressure for the duration of a preset inspiratory time (T). Gas flow is terminated when the preset pressure is achieved. The volume delivered is determined by the compliance of the patient’s respiratory system, airway resistance, T, and the pressure target. In addition, the clinician sets the desired PEEP, respiratory rate, FIO2, I : E ratio, and trigger mode. Pressure is maintained with a variable or intermittent flow rate for the set T,. In the setting of hypoxemia, T may be increased quite precisely to increase mean airway pressure and oxygenation. This strategy is much more difficult, if not impossible, to manipulate with VCV.

An advantage of PCV is that airway pressure is tightly managed to limit or eliminate alveolar overdistention and to reduce ventilator-induced lung injury. It should be noted that the clinician does not control waveform or peak inspiratory flow. Patients can generate their desired flow rate and thus reduce air hunger. Pressure-targeted modes, which are growing in popularity, might have better pressure distribution, improved dissemination of airway pressure, and greater distribution of ventilation.

One problem with PCV is that the volume received by the patient is potentially variable. Any change in system compliance or resistance (or both) will affect the VT generated. For example, if the patient bites on the ET tube or a mucous plug develops, the set pressure that was generating an adequate volume will no longer do so. In contrast, a sudden increase in system compliance might result in the generation of VT that may be considerably larger than desired. Instead of the traditional pressure alarm limits, one must adjust and be cognizant of VT and minute ventilation alarm settings. Uncertainties such as these have led many clinicians to favor volume-targeted strategies or dual-controlled strategies in the acute care setting.

**Modes of Ventilation Commonly Used in the ED**

Assist/control (AC) and synchronized intermittent mandatory ventilation (SIMV) are the ventilation modes most commonly used in the ED. Both are acceptable, and no data have demonstrated a better outcome with either mode. Other modes are also acceptable based on clinician preference.

**AC Ventilation**

Here, ventilator-initiated breaths, known as machine breaths (i.e., control breaths), are provided at a preset rate. Every breath is fully supported by the ventilator, regardless of whether the breath is initiated by the patient or the ventilator. The clinician sets the base ventilation rate, but if the patient tries to breathe faster than the set rate, additional breaths can be initiated by the patient, known as spontaneous breaths (i.e., assist breaths). A potential downside is inappropriate hyperventilation.

AC modes may be either volume or pressure cycled. Both assist breaths and control breaths will reach the set target, be it a set volume (in a volume-targeted mode) or a set pressure (in a pressure-targeted mode). In VCV, the spontaneous breath receives the same VT that is set for the machine breath. In PCV, the spontaneous breath receives the same pressure that is set for the machine breath. To be more specific, in the volume-targeted mode, the clinician sets VT, as well as the IFR, flow waveform, sensitivity to the patient’s respiratory effort (i.e., trigger), and the basal ventilatory rate. In the pressure-targeted mode, the clinician determines the basal ventilatory rate and how sensitive the ventilator will be to the patient’s respiratory effort and also selects pressure levels and T,. Hence, in this mode VT is not set by the ventilator but is dependent on the compliance of the lung and chest wall and airway pressure. This helps avoid pressure-induced lung injury, but a specific VT is not guaranteed.

For the patient to trigger the ventilator and initiate flow for a spontaneous breath, mean airway pressure must decrease by a preset amount below PEEP if set on a pressure trigger or flow to be generated if set on a flow trigger. The amount necessary to open the inflow valve is the sensitivity setting. Caution should be exercised to avoid auto-PEEP (also known as breath stacking) when using volume-targeted AC modes. Because each mechanically delivered breath is given at full VT, patients with a high actual respiratory rate on AC may not have sufficient time to completely exhale between breaths. This results in progressive air trapping, which leads to an increase in auto-PEEP (PEEP) (see Fig. 8.6). This is of clinical concern in patients with asthma, in whom auto-PEEP can significantly reduce cardiac output and even promote cardiovascular collapse.

**SIMV**

SIMV provides breaths at a preset rate (machine breath), similar to the AC mode. The patient can initiate an additional spontaneous breath between the mandated or preset number of ventilator-supported breaths. Such spontaneous breaths above the preset ventilatory rate are not supported by the ventilator,
and the patient receives only a spontaneous VT that reflects the depth and time spent in the patient-controlled inspiration. For each of these nonmandatory (i.e., spontaneous) breaths, the patient has a high work of breathing. SIMV is typically partnered with PSV to aid in spontaneous breathing support and to overcome the intrinsic resistance associated with MV. This mode was initially recommended by those who thought that, as a patient’s need for mechanical ventilatory support decreased, the set respiratory rate could be decreased and the patient “weaned” to PSV alone and ensuing extubation. Subsequent data have shown that this method of liberation actually increases the number of ventilator days.15 The synchronized version of intermittent MV allows the ventilator to attempt to coordinate spontaneous and machine breaths to prevent it from delivering a scheduled breath on top of a spontaneous breath or during exhalation after a spontaneous breath. This could lead to elevated mean airway pressure, alveolar overdistention, and biotrauma.16

Both volume-targeted ventilation and pressure-targeted ventilation modes can be set to either an AC or SIMV mode to achieve the desired minute ventilation. In a chemically paralyzed patient with no intrinsic respiratory drive, AC and SIMV look virtually identical. Both will reach their target (volume or pressure) at the set rate. If a patient triggers the ventilator at a rate greater than the set rate, these two strategies diverge. In AC, each breath above the set respiratory rate will result in a full mechanically supported breath to reach either the set volume or pressure target. In SIMV, the ventilator will give only the set number of breaths that the clinician has selected. Each additional breath will require the patient to generate a spontaneous VT without mechanical assistance. This patient-generated breath must overcome any resistance caused by the artificial airway and ventilator circuitry. Pressure support should be added to SIMV for patient-generated breaths to reduce any increase in the work of breathing related to the resistance imposed by the ventilator circuit and ET tube.

**Advanced Modes of MV**

**Dual-Control Modes**

Advanced modes of MV use a closed-loop ventilator logic that combines the features of volume- and pressure-targeted ventilation (Box 8.1). These modes automatically alter control variables, either breath to breath or within a breath, to ensure a minimum VT or minute ventilation.17 Detailed explanation of these modes is beyond the scope of this chapter. If these modes are encountered, one should discuss options with a respiratory therapist and critical care medicine specialist.

**Other Modes**

**High-Frequency Ventilation**

High-frequency ventilation (HFV) attempts to achieve adequate gas exchange by using asymmetric velocity profiles when combining very high respiratory rates with VT levels that are smaller than the volume of anatomic dead space. It is used more commonly in neonates and infants with neonatal respiratory failure. There has been renewed interest in using HFV in adult patients with ALI or ARDS under the rationale that the small VT may cause less ventilator-associated lung injury. More trials are necessary to determine whether HFV can improve mortality outcomes in these patients.18

---

**Box 8.1 Advanced Modes of Mechanical Ventilation**

**BREATH TO BREATH**

- Pressure-regulated volume control (PRVC)
- Auto flow
- Volume control plus (VC+)
- Adaptive pressure ventilation (APV)
- Variable-pressure control (VPC)

**WITHIN A BREATH**

- Volume-assured pressure support ventilation (VAPSV)
- Pressure augmentation

**OTHER**

- High-frequency ventilation (HFV)
- Airway pressure release ventilation (APRV)
- Bilevel
- Proportional assist ventilation

**Airway Pressure Release Ventilation and Bi-Level Ventilation**

Both these modes are proprietary names yet function in essentially the same manner. The clinician sets a pressure high, a pressure low, and a time at each level (time high and time low). Although at first glance this appears to be similar to PCV, it differs markedly in that the majority of time is spent at pressure high with brief periods at pressure low. The patient typically spends 4 to 6 seconds in time high. Pressure high may be as high as 40 cm H₂O or greater. Ventilation occurs during the release from pressure high to pressure low. Time low is typically 0.2 to 0.8 second in restrictive lung disease and 0.8 to 1.5 seconds in obstructive lung disease. It is probably prudent to start at 0.8 and titrate to meet individual patient requirements. Time low is also referred to as the release phase.19

The long time that high-level pressure is maintained achieves oxygenation, and the short release period achieves clearance of CO₂ (Fig. 8.10). The long time at high-level pressure results in substantial recruitment of alveoli from markedly different regional time constants at rather low gas flow rates. The establishment of PEEP, by the short release time enhances oxygenation. CO₂ clearance is aided by recruitment of the patient’s lung at close to total lung capacity. Elastic recoil creates large-volume gas flow during the release period.

In a paralyzed patient, airway pressure release ventilation and bilevel ventilation (APRV/Bi-Level) are identical to pressure-targeted IRV. For these reasons, some have described this mode as inverse ratio ventilation. A major difference between APRV/Bi-Level and IRV is that IRV typically requires chemical paralysis or heavy sedation. APRV/Bi-Level is a fundamentally different mode from cyclic ventilation. This mode allows the patient to spontaneously breathe during all phases of the cycle, thus making it relatively more comfortable and reducing the level of sedation or paralysis needed. This mode is enabled to succeed by having a floating valve that is responsive to the patient’s needs, regardless of the location within the respiratory cycle. The patient is allowed to breathe in or out during the pressure high phase and during the release phase. Accordingly, the sequence is called a phase cycle. There is no set Tᵢ or Tₑ.
and no readily identifiable respiratory rate in the traditional sense. During the pressure high phase, patients may exhale 50 to 200 mL or more of gas as the lung volume becomes full of gas. This is not a full exhalation, and the release of excess gas should not be counted as a breath. APRV has been used successfully for neonatal, pediatric, and adult forms of respiratory failure. It is considered an alternative open–lung model approach to MV. 19

Given the spontaneous nature of the mode, there should be virtually no need for continuous infusion of neuromuscular blocking drugs in patients placed on this mode of ventilation. This may result in a shorter length of ICU stay and a reduced incidence of prolonged neuromuscular blockade syndrome. The need for sedatives is reduced because patients are more comfortable on this spontaneous mode than on cyclic ventilation. 20 APRV/Bi-Level has gained popularity in patients with hypoxemic respiratory failure because it improves oxygenation by optimizing alveolar recruitment and V/Q matching. 21

A common mistake with this mode is setting time low too long. This essentially mimics a pressure-targeted SIMV strategy. Transport of patients on APRV with pressure high greater than 20 cm H2O should occur with the patient attached to the ventilator instead of being hand-ventilated. 22 Hand ventilation is unable to match the manner of gas delivery and pressure dynamics that the patient requires. Attempts at hand ventilation, even with an appropriately set PEEP valve, are frequently complicated by unexpected hypoxemia and hemodynamic instability.

Noninvasive Positive Pressure Ventilation

Noninvasive ventilation is defined as the provision of ventilatory assistance without an invasive artificial airway. Noninvasive ventilators consist of both negative and positive pressure ventilators. Because negative pressure ventilation is so rarely used today, our discussion is limited to PPV.

Before the 1960s, the use of negative pressure ventilation in the form of a tank ventilator (the “iron lung”) was the most common form of MV outside the anesthesia suite. It was not until the 1952 polio epidemic in Copenhagen that anesthesiologist Bjorn Ibsen showed that he could improve the survival of patients with respiratory paralysis by using invasive PPV.

Nonetheless, negative pressure or “iron lungs” were the mainstay of ventilatory support for patients with chronic respiratory failure until as late as the mid-1980s. In the early 1980s, nasal continuous positive airway pressure (CPAP) was introduced to treat obstructive sleep apnea. These tightly fitting masks proved to be an effective means of assisting ventilation, and noninvasive positive pressure ventilation (NPPV) quickly displaced traditional negative pressure ventilation as the treatment of choice for chronic respiratory failure in patients with neuromuscular and chest wall deformities. Current NPPV devices are able to provide a set respiratory rate, set VT, and set FiO2. The use of NPPV has also been integrated into the acute inpatient setting, where it is now used to treat acute respiratory failure. 21

Definitions

The current literature uses different definitions for NPPV. Although some authors use NPPV as an umbrella term that includes CPAP and bilevel positive airway pressure, more recently, authors have used the term NPPV as synonymous with bilevel and consider CPAP a separate entity. The terms NPPV and noninvasive intermittent positive-pressure ventilation (NIPPV) are often used interchangeably with bilevel.

As its name suggests, CPAP supplies continuous positive pressure via a tightly fitting face mask (Fig. 8.11). NPPV
The term CPAP is synonymous with PEEPi and the area available for gas exchange, and improves the V/Q ratio. CPAP increases alveolar recruitment and size, thereby enhancing the patient’s work of breathing. Careful application of PEEP can reduce this gradient and decrease the patient’s work of breathing. PPV also creates an increase in intrathoracic pressure. This causes preload to decrease as a result of diminished venous return and also decreases transmural pressure, which reduces afterload.

or bilevel provides an inspiratory positive airway pressure (IPAP) in addition to end-expiratory positive airway pressure (EPAP), and breaths are usually triggered by the patient (Fig. 8.12). On many such devices, backup rates may be set that deliver bilevel pressure, even if patients fail to initiate a breath.

### Rationale for Using NPPV

The most important advantage of NPPV is avoiding the complications associated with invasive MV. It has been well documented that invasive MV increases the incidence of airway and lung injury and augments the risk for nosocomial pneumonia. NPPV avoids these complications by keeping the upper airway defense mechanisms intact and allows the patient to retain the ability to eat, clear secretions, and communicate normally when NPPV is used intermittently (NIPPV). NPPV has the potential to reduce the mortality of a selected group of patients with acute respiratory failure and may shorten hospital stays, thereby reducing cost. Specific to the ED, appropriate initiation of NPPV may avoid unnecessary intubation of certain patients, hence avoiding ICU admission, reducing cost, decreasing complications, and improving mortality.

### Pathophysiologic Effects of NPPV

CPAP increases alveolar recruitment and size, thereby enhancing the area available for gas exchange, and improves the V/Q relationship. The term CPAP is synonymous with PEEPi, and bilevel EPAP. It can also negate the effects of PEEP. In patients with dynamic hyperinflation (such as asthma or COPD), an escalating PEEP increases the magnitude of the drop in airway pressure that the patient must generate to trigger a breath. This causes increased work of breathing for the patient. Careful application of PEEP can reduce this gradient and decrease the patient’s work of breathing. PPV also creates an increase in intrathoracic pressure. This causes preload to decrease as a result of diminished venous return and also decreases transmural pressure, which reduces afterload.

In NPPV, IPAP is similar to pressure support and, when combined with EPAP, further augments alveolar ventilation, thereby allowing some rest of the respiratory muscles during the inspiratory phase (Box 8.2).

### Acute Exacerbation of COPD

Numerous studies have shown that NPPV can reduce the need for intubation, length of hospital stay, and in-hospital mortality in patients with acute exacerbations of COPD. NPPV should be initiated early and along with standard medical therapy. If it is started late, after the failure of medical treatment, the benefits conferred by NPPV (hospital mortality, length of ICU stay, number of days on the ventilator, overall complications) are eliminated. Early implementation of NPPV in patients seen in the ED with an acute exacerbation of COPD and without contraindications should be considered the standard of care.

### Acute Cardiogenic Pulmonary Edema

CPAP has been shown to produce a reduction in the rate of intubation and a trend toward decreasing mortality. CPAP and NPPV both reduce the risk of intubation in the ED. An early article described an increased rate of acute myocardial infarction in patients with acute cardiogenic pulmonary edema treated with NPPV, but several subsequent trials have refuted this increased risk for myocardial infarction.

### Hypoxemic Respiratory Failure

Although some of the literature suggests that NPPV may be beneficial in the setting of acute hypoxemic respiratory failure, doubt still exists. A large multicenter trial of NPPV in patients with acute hypoxemic respiratory failure that excludes patients with cardiogenic pulmonary edema and COPD may help clarify the use of NPPV in this setting.

NPPV has been used increasingly in the ICU for hypoxemic respiratory failure. It can be considered for patients who are hemodynamically stable with single-organ (i.e., pulmonary) failure, but extremely close monitoring is required.

### Immunocompromised Patients

When faced with a severely immunosuppressed patient with acute hypoxemic respiratory failure in the ED, early initiation of NPPV may be beneficial in avoiding the serious complications of ET intubation. NPPV can keep upper airway defenses intact and minimize the risk for ventilator-associated pneumonia, which is universally fatal in these patients. NPPV has been shown to be associated with a lower rate of ET intubation, shorter ICU stay, and lower ICU mortality. In-house mortality did not differ significantly.

### Box 8.2 Indications for Initiating Noninvasive Positive Pressure Ventilation

- Exacerbation of chronic obstructive pulmonary disease
- Exacerbation of congestive heart failure and cardiogenic pulmonary edema
- Exacerbation of asthma
- Immunocompromised patients
- Hypoxemic respiratory failure
- Do-not-resuscitate/do-not-intubate advance directive

---

**Figure 8.12** Bilevel positive airway pressure (BiPAP) S/T noninvasive ventilation system (Phillips Respironics, Inc, Andover, MA). Adjustable parameters include inspiratory positive airway pressure, expiratory positive airway pressure, and breaths per minute. Both BiPAP and continuous positive airway pressure are used to support ventilation in patients with decompensated congestive heart failure, chronic obstructive pulmonary disease, pneumonia, and asthma, but neither mode has a clear benefit over the other.
“Do-Not-Intubate/Do-Not-Resuscitate” Patients

In do-not-intubate/do-not-resuscitate (DNI/DNR) patients willing to undergo NPPV, success would be measured by improved ventilation, oxygenation, and comfort. NPPV can provide support for the patient while the underlying cause of the respiratory failure is being treated. NPPV should be discontinued if it is not producing the desired response or if the patient is unable to tolerate this therapy. In these circumstances it should be a joint decision between the health care team, the patient, and the family to limit NPPV and transition toward comfort measures.

In patients who have chosen to forego any life-sustaining therapy and are receiving comfort care measures, NPPV might be used as a form of palliative care in an attempt to reduce the associated dyspnea. In this circumstance, the use of NPPV is considered successful if it alleviates the patient’s symptoms. If it causes any discomfort to the patient, it should be discontinued. This use of NPPV is controversial, and no studies have assessed the benefits of NPPV in these patients. Another use of NPPV in patients who have chosen comfort care measures is a time-limited trial to achieve the goal of survival until the arrival of family and friends. In this situation, NPPV would be used to provide life support until friends and family can achieve closure.

Even if a patient with a known DNI advance directive arrives at the ED with acute respiratory failure of reversible etiology, it can be beneficial discussing the use of NPPV with the patient and family. In this situation, communication about the expectations and goals of care is of utmost importance.

Initiation of NPPV

There is no standard approach to the initiation of NPPV. Different methods have been used in clinical trials, yet these methods have never been compared. There are two main strategies: a high-low approach and a low-high approach (Fig. 8.13).

Emphasis should be placed on the importance of close follow-up of patients in whom NPPV is started in the ED. It is important to serially assess patient response as soon as 30 minutes after the initiation of NPPV. Those who are persistently tachypneic and acidemic should be considered for intubation sooner rather than later. It is imperative to closely observe patients for deterioration. ABGs should be checked within 1 to 2 hours after initiation of NPPV to assess treatment success or failure. Patients who do not improve clinically should be considered for intubation.

Cautions With the Use of NPPV

Most studies involving NPPV have excluded patients who were hemodynamically unstable, had an altered level of consciousness, or were unable to protect their airway. This was based on the concern that a depressed sensorium would predispose the patient to aspiration. The International Consensus Conference in Intensive Care Medicine on Noninvasive Positive Pressure Ventilation in Acute Respiratory Failure held in April 2000 considered the presence of severe encephalopathy, as manifested by a Glasgow Coma Scale score lower than 10, to be a contraindication to NPPV.

Other accepted contraindications to NPPV are listed in Box 8.3.

Recently, studies have looked specifically at the use of NPPV in patients with hypercapnic coma secondary to acute respiratory failure. This was based on the observation that some DNI patients who declined intubation had successful outcomes with NPPV therapy despite their initial comatose state. Diaz and colleagues conducted an observational study and found that success rates were comparable between the comatose and noncomatose group.

Scala and coworkers performed two studies, both showing that NPPV could be used successfully in treating patients with exacerbations of COPD and hypercapnic encephalopathy. Their 2007 study showed that performance of NPPV by an experienced team led to similar short- and long-term survival, fewer nosocomial infections, and shorter durations of hospitalization than in patients who underwent MV.

Of note, these studies were conducted in the ICU or specialized respiratory care units with a nursing ratio of at least 1:3. The patients were very closely monitored by staff while they received NPPV. This high nursing-to-patient ratio may not be feasible in a busy ED, however.

The other key point in these studies was the rapid improvement in neurologic status that occurred 1 to 2 hours after the initiation of NPPV. The importance of close monitoring of patients in whom NPPV is started is crucial in identifying those who will fail this therapy.

**Box 8.3 Contraindications to Noninvasive Positive Pressure Ventilation**

- Impending cardiovascular collapse or respiratory arrest
- Severe upper gastrointestinal bleeding
- Facial surgery, trauma, or deformity limiting placement of the mask
- Upper airway obstruction
- Inability to cooperate or protect the airway, altered mental status
- Inability to clear respiratory secretions
- High risk for aspiration

---

**Figure 8.13** Initiation of noninvasive positive pressure ventilation (NPPV). EPAP, End-expiratory positive airway pressure; IPAP, inspiratory positive airway pressure; PEEP, positive end-expiratory pressure; PSV, pressure support ventilation.
High-Flow Nasal Cannula

The high-flow nasal cannula (HFNC) is a relatively new oxygen delivery system. A conventional nasal cannula uses a low-flow system and at higher flow (≥6 L/min) can cause nasal dryness, epistaxis, and patient discomfort. The HFNC system (Fig. 8.14) is a novel device that combines oxygen, pressurized air, and warm humidification to deliver tolerable flow of up to 40 L/min through a nasal cannula. FIO₂ and flow rates can be adjusted. With higher flow, less room-air entrainment occurs, and the higher flow rates match the dyspneic patient’s increased minute ventilation.

Its use has been studied more extensively for neonatal respiratory care, where ongoing studies suggest that HFNC may be as effective as nasal CPAP in preterm neonates. One study looked at the effect of HFNC on exercise performance with acute respiratory failure. Positive pressure ventilation in the emergency department, with immunosuppressed patients being targeted in the hope of avoiding intubation, may be as effective as nasal CPAP in preterm neonates. One study has investigated the use of HFNC in patients with acute respiratory failure. Anecdotally, this system has been used with some success in adults in the ICU, with immunosuppressed patients being targeted in the hope of avoiding intubation. It seems to be better tolerated than NPPV, and at higher flow it is believed to provide a certain amount of continuous positive pressure. More studies will have to be performed before this technology becomes a mainstay of treatment in patients with acute respiratory failure.

NPPV has been shown to work well in patients with reversible conditions and acts as a bridge while allowing medical therapy (e.g., bronchodilators, steroids, diuretics) to take effect, thereby potentially avoiding the need for ET intubation. Those with less reversible causes of their acute respiratory failure (ARDS, pneumonia) may be less likely to respond. With careful patient selection, NPPV can be safely initiated in the ED, give time for medical treatment to work, and potentially avoid ICU admission. Patients being considered for intubation should be evaluated for the potential use of NPPV.

Close monitoring and follow-up of patients placed on NPPV is crucial in determining whether the therapy has been successful or whether further intervention is needed. In EDs in which the staff has been adequately trained, selected patients, even those in a hypercapnic coma, can be considered for a short trial of NPPV. Extreme diligence in monitoring these patients must be used, and if improvement is not seen in 1 to 2 hours, intubation should not be delayed.

Conclusions

The short-term administration of paralyzing agents, in addition to sedation and analgesia, for mechanically ventilated patients in the ED is appropriate to improve patient-ventilator synchrony, enhance gas exchange, diminish the risk for barotrauma, decrease muscle oxygen consumption, facilitate short procedures, and prevent movement in patients with elevated ICP. Importantly, neuromuscular blocking agents (NMBAs) have no sedative, amnestic, or analgesic properties, and all patients administered such an agent must receive concomitant sedation and analgesia.

The long-term use of chemical paralysis in the ICU has greatly diminished because of prolonged recovery secondary to drug and metabolite accumulation. Cases of prolonged paralysis, also known as critical illness myopathy, postparalysis syndrome, polyneuropathy of critical illness, and acute quadriceps myopathy syndrome, lead to protracted ICU stays. In the ICU, daily discontinuation of NMBAs for a few hours or avoiding their use entirely has been recommended to potentially decrease the incidence of these conditions. When clinically feasible, it is advisable to stop administration of the NMA in the ED and reassess the need for continued paralysis.

A number of drugs are available, but the nondepolarizing agents vecuronium and pancuronium are the most commonly prescribed agents for short-term paralysis in the ED. Both are given by intermittent bolus administration as needed. Continuous infusion, though potentially useful, is rarely used or indicated in the ED. Nondepolarizing NMBAs can be reversed by neostigmine at a dose of 0.035 to 0.07 mg/kg.

Vecuronium has an intermediate duration of action with a half-life of 80 to 90 minutes, slightly longer in the elderly, but not affected by renal or hepatic failure. Minimal adverse cardiovascular side effects have been reported, and there is minimal risk for hypotension. Vecuronium has minimal adverse cardiovascular effects and is the drug of choice for patients with cardiovascular disease or hemodynamic instability. The dose for intermittent bolus administration is 0.1 mg/kg.
These patients with the use of paralytics to keep their respiratory rate and expiratory time controlled. This should be a final option and be done with the understanding that the side effects of steroids and paralytics can be quite devastating, but paralysis cannot be avoided in certain cases. In the ICU, paralysis is commonly titrated to an effect monitored by a peripheral nerve monitor applied over the ulnar or other peripheral nerve.

### BOX 8.4 Common Sedatives and Analgesics Used With Mechanical Ventilation

**SEDATIVES**
- Propofol, 5–80 µg/kg per min
- Dexmedetomidine, 0.2–1.5 µg/kg per hr
- Lorazepam, 1–5 mg IV prn; infusion at 0.5–7 mg/hr if multiple doses required
- Midazolam, 1–6 mg IV prn
- Haloperidol, 2–10 mg IV every 20–30 minutes until stable, then 25% of the loading dose q 6 hr prn
- Remifentanil infusion: 0.025–0.2 µg/kg per hr (no bolus used, provides both sedation and analgesia)

**ANALGESICS**
- Fentanyl, 25–100 µg IV q 15–30 min prn
- Hydromorphone, 0.2–1 mg IV prn
- Morphine, 2–5 mg IV prn
- Fentanyl infusion: 50–200 µg/hr if more than 2–3 boluses per hour of above agents are needed

$q$, Every; prn, as needed.

Pancuronium is long acting with a half-life of 100 to 130 minutes. Its duration of action is prolonged in patients with renal or hepatic failure. There is a moderate risk for adverse cardiovascular effects associated with pancuronium, such as tachycardia, hypertension, and increased cardiac output secondary to vagal blockade. The dose for intermittent bolus administration is 0.05 to 0.1 mg/kg.

### SEDATION

See Box 8.4 for common sedatives and analgesics. It is recommended that only the minimal amount needed to achieve comfort be used. Sedation should be targeted to a specific sedation scale (such as the Richmond Agitation Sedation Scale). Agents with a quick offset are preferable to improve spontaneous awakening and for breathing trials once the patient is a candidate for extubation. Benzodiazepines should be avoided if possible. If paralyzing drugs are used for neuromuscular blockade, it is imperative to provide aggressive sedation because pain and anxiety in paralyzed patients cannot be evaluated. It is prudent to conclude that a paralyzed ventilated patient is awake and can hear and feel unless sedated.

### SPECIFIC DISEASE PROCESSES

#### Asthma and COPD (Fig. 8.15)

As asthma and COPD treatment (β-agonists, steroids) take effect, peak pressure will begin to lower, the peak expiratory flow rate should increase, and the expiration flow time should shorten (Fig. 8.16). It is necessary to adequately sedate these patients because their hypoxemic state is a powerful stimulus to breathe rapidly. Opiates such as fentanyl and sedatives such as propofol and ketamine have gained increased roles in these patients. Occasionally, one will be required to chemically weaken

### Initial Ventilator Settings

- Volume targeted (VCV), AC, RR 8–10, Vt 6 mL/kg IBW, PEEP 5, FiO₂ 100%
- Flow rate 80 L/min, square waveform
- These strategies may worsen hypercapnia (termed permissive hypercapnia) because of a decrease in minute ventilation and is tolerated if pH >7.15
- NaHCO₃ infusion or the administration of THAM (tris[hydroxymethyl] aminomethane) may be required to keep arterial pH above 7.15 to 7.20

### After Intubation

- Adequate sedation and analgesia. Consider the use of propofol, ketamine, or opiates
- Monitor flow-time curves and pressure-time curves
- Follow peak pressure and plateau pressure
- Consider chemical weakening as a last resort (goal TOF 2–4)
- Consider heliox if available

#### Figure 8.15 Initial ventilator settings for exacerbation of asthma.

- AC, Assist/control mode; BVM, bag-valve-mask ventilation; FiO₂, fraction of inspired oxygen; IBW, ideal body weight; VT, intravenous fluid; NaHCO₃, sodium bicarbonate; PEEP, positive end-expiratory pressure; RR, respiratory rate; RSI, rapid sequence induction; THAM, tris(hydroxymethyl)aminomethane; TOF, train of four; VCV, volume-cycled ventilation; VT, tidal volume.

#### Figure 8.16 Schematic of a flow-time curve showing improvement in an obstructive process. Note that the peak expiratory flow rate (PEFR) increases and the expiratory time (Te) decreases with improvement and flow reaches zero before the next breath is given.
distribution. No blockade results in four twitches of the adductor pollicis muscle causing four supramaximal triggering stimuli; complete blockade yields no response. A common goal of blockade is to use enough of the agent to result in two twitches out of a “train of four.” A peripheral nerve monitor is not commonly used in the ED and is not a standard intervention during resuscitation and interim ED care.

Valuable information can be gained from flow-time curves and pressure-time curves. Improvement (see Fig. 8.16) or worsening of airway obstruction and air trapping (see Fig. 8.6) can often be evident in these curves before becoming clinically apparent. In ideal situations, PEEP, can be measured with an end-expiratory hold. This measurement is often inconsistent and difficult to obtain. Some authors have recommended that PEEP be applied at 80% of the measured PEEP. Because of the difficulty in accurately measuring PEEP, and the potential hazard of adding too much PEEP, others have suggested that it should be set at 50% of the measured PEEP. If PEEP becomes severe enough, it will begin to affect plateau pressure. Current recommendations for obstructive airway disease are to keep plateau pressure below 35 cm H2O, but many clinicians follow the ALI/ARDS recommendations and maintain plateau pressure at less than 30 cm H2O.

PEEP, can be used to decrease some of the negative pressure that the patient has to generate to initiate a breath. For example, a patient with a PEEP of 10 cm H2O and a set pressure trigger of ~2 cm H2O (and no applied PEEP) has to generate an alveolar pressure of 12 cm H2O to produce a breath. Adding a PEEP of 5 cm H2O means that the ventilator will trigger a breath when alveolar pressure is 3 cm H2O. The patient has to generate a decrease of only 7 cm H2O (instead of 12 cm H2O), thereby reducing the inspiratory effort required.

Although seemingly counterintuitive, decreasing the respiratory rate and VT in a critically ill asthmatic may be beneficial and result in an acceptable elevation in Pco2, termed permissive hypercapnia. Hence, it may not be advisable to meet arbitrary Pco2 levels in a ventilated asthmatic but rather concentrate on maintaining acceptable oxygenation (Pao2 > 60 mm Hg, SaO2 of 88% to 92%) while minimizing PEEP, and optimizing plateau pressure.

If cardiovascular collapse occurs in a ventilated asthmatic with either pulseless electrical activity or sudden hypotension, a first step in troubleshooting is to remove the patient from the ventilator. This is both a diagnostic and a therapeutic maneuver for air trapping. Some clinicians also advocate fluid loading and rapid and deep chest compressions while the patient is disconnected from the ventilator to expel the excess volumes of air trapped by prior aggressive ventilation (Fig. 8.17). Tension pneumothorax must also be considered (see later).

**ALI and ARDS**

Initial ventilator settings for patients with ALI and ARDS can be found in Fig. 8.18.

Efforts should be made while patient is in the ED to decrease VT down to 6 mL/kg IBW, as recent data have shown settings made in the ED are continued in the ICU. Forty-one percent of patients never received recommended settings during hospitalization. Fuller and colleagues also showed lung protective ventilation was used only 47% of the time in patients with ARDS in the ED.

A common finding in lung-protective ventilation is the occurrence of patient-ventilator dyssynchrony. This is thought to be due to the patient wanting a higher flow rate than the ventilator is providing while on a volume-targeted strategy. This occasionally leads to double or triple cycling of the ventilator. It should be noted that in this situation the patient is actually receiving a higher VT and not benefiting from lung-protective ventilation. Sedation needs to be optimized, and at times different modes, such as pressure-targeted modes, may be attempted. Temporarily weakening the patient with paralytics may be considered.
There are several areas of uncertainty with MV in patients with ALI or ARDS. Patients with traumatic brain injury, intracranial hemorrhage, fulminant hepatic failure, and elevated ICP in whom ARDS develops must be managed carefully because lung-protective ventilation may induce hypercapnia. Acutely, this may lead to cerebral vasodilation and an increase in ICP. There is little evidence to support the recommendation for any particular rescue therapy in patients with severe refractory hypoxia, such as recruitment maneuvers, high-dose albuterol, IRV, HFV, prone ventilation, and extracorporeal membrane oxygenation. In dire circumstances, these modalities may be used on the basis of clinician preference and expertise and consultation with a critical care specialist.

**COMPLICATIONS OF MV**

Though necessary to sustain life, MV is associated with a number of pathophysiologic derangements that can lead to morbidity and mortality, including pulmonary barotrauma, ventilator-associated lung injury, hemodynamic compromise, PEEP, and elevated ICP.

**Pneumothorax**

Pneumothorax that is not associated with trauma in a mechanically ventilated patient typically stems from alveolar overdistention (continuous or episodic) and leads to alveolar rupture and escape of gas into the pleural space. In patients receiving PPV, it is wise to drain the pleural space to prevent a simple pneumothorax from progressing to tension pneumothorax with hemodynamic compromise. Loculated pneumothoraces may be successfully drained percutaneously under ultrasound (US) or computed tomography (CT) guidance. Successful drainage of air space disease leads to enhanced liberation from MV. Pneumothorax or tension pneumothorax may also result from aggressive bag-valve-mask ventilation. Patients with intrinsic lung disease such as COPD or asthma are more prone to the development of pneumothorax than the average patient.

A simple pneumothorax can be drained by surgical tube thoracostomy with a small-bore tube (24 Fr), a commercially available pneumothorax kit (Arrow [Teleflex, Morrisville, NC]), or a pigtail catheter placed into the pleural space via the Seldinger technique (see Chapter 9). Each of these catheters should be placed into a chest drainage collection unit that incorporates a water seal chamber and variable suction control. Treat persistent air leaks initially with continuous suction (usually suction at 20 cm H₂O) to evacuate the pleural space and promote coaptation of the visceral and parietal pleurae. Reduce suction and place the chest tube on water seal only after resolution of any air leak. Remove the chest tube directly from the water seal if no pneumothorax is apparent on a chest film or after a test period of tube clamping and subsequent radiographic evaluation. The author favors a 4-hour period of clamping because recurrent pneumothorax is easier to treat by unclamping a tube than by placing a new one. Not all patients with a pneumothorax require invasive techniques to evacuate air from the pleural space. It is important to recognize that small pneumothoraces occurring in spontaneously breathing patients (i.e., negative pressure ventilation) may be reevaluated in 4 to 6 hours with a repeated chest radiograph and drained only if they are expanding. This option is not advised for patients who are on any form of PPV because a simple pneumothorax can rapidly become a tension pneumothorax with subsequent hypotension and death. Tension pneumothoraces may be recognized by tachycardia, hypotension, elevated peak airway pressure (if mechanically ventilated, tachypnea if not), jugular venous distention, thoracic resonance by percussion on the affected side, diminished or absent breath sounds on the affected side, and tracheal deviation away from the affected side. Because not all signs or symptoms are present in all patients, treatment should be dictated by the patient’s clinical condition. Loculated pneumothoraces or fluid collections develop in certain patients. If the collections are either single or immediately adjacent to one another and readily identified, they may be drained under US guidance at the bedside. Loculations are frequently in inaccessible areas or are difficult to image with US. CT scanning of the thorax can provide precise anatomic definition of the presence and number of loculated collections and be used as a guide for the interventional radiologist. Successful treatment involving CT-guided drainage of loculated pleural collections (air and fluid) to assist in weaning of patients from mechanical ventilator support has been reported.

**Ventilator-Induced Lung Injury**

There are several causes of ventilator-induced lung injury, including biotrauma, volutrauma, barotrauma, and atelectasis-related trauma. Biotrauma refers to the self-sustaining process of lung injury from MV that follows alveolar overdistention or rupture, alveolar hypoperfusion, and repetitive shear stress across alveolar walls. Originally, this problem was thought to be caused by too much pressure (barotrauma). Current principles hold that elevated airway pressure is a straightforward reflection of excess volume delivered to a lung that cannot accept excess gas (i.e., in volutrauma, excess volume is delivered). Lung injury is an inhomogeneous process with areas of normal lung immediately adjacent to diseased and injured segments. The healthy and compliant segments with shorter regional time constants will readily accept gas, but their neighbors with reduced compliance and longer regional time constants will not. The end result is overdistention of the compliant segments, alveolar injury, liberation of inflammatory cytokines and chemokines, activation of endothelin and arachidonic acid pathways, and the expression of adhesion molecules along the vascular endothelium. This leads to infiltration of inflammatory cells, release of destructive lysosomal enzymes, and induction of toxic oxygen metabolites. Avoiding this inflammatory cascade is an intelligent means of protecting a patient’s lungs from volume-induced lung injury. Such a notion has given rise to lung-protective ventilator strategies based on low-Vt ventilation (6 mL/kg IBW) and low plateau pressure (< 30 cm H₂O). Several studies have reported the development within hours of ventilator-induced lung injury in patients with normal lungs that were ventilated with larger Vt (12 mL/kg). Current recommendations are for all MV to be conducted with lower Vt than the once-standard 12 to 15 mL/kg. Patients with abnormal lungs (interstitial lung disease, lung resection, severe pneumonia, edema) or the presence of an ALI risk factor (sepsis, aspiration, transfusion) should be ventilated at a Vt of 6 mL/kg IBW. These patients may initially be markedly acidotic and may require starting Vt at 8 mL/kg IBW and titrating down to 6 mL/kg IBW within 2 hours. Those with normal lungs and no ALI risk factors should be started at a Vt of less than 10 mL/kg IBW.
Hemodynamic Compromise

In all circumstances, the volume of venous return exactly matches the volume of cardiac output. Any process that impedes venous return will decrease the available volume that establishes cardiac output. For patients receiving PPV, each gas delivery increases intrathoracic pressure, and exhalation decreases that pressure. Venous return principally occurs during exhalation. If the ventilator settings lead to increased intrathoracic pressure during exhalation, venous return will be reduced. Variables that can lead to this circumstance are increased PEEP, auto-PEEP, and IRV. Venous return not only depends on a relatively negative pressure within the thoracic cavity but also relies on a sufficient amount of time for flow into the thoracic vasculature and right side of the heart. Significantly high respiratory rates may compromise venous return. An additional untoward side effect of impaired venous return is cerebral venous hypertension from impeded venous drainage. Because of the absence of valves between the cerebral parenchyma and the right atrium, increased pressure on the right atrium reduces cerebral venous flow and may contribute to cerebral ischemia in patients with traumatic brain injury or stroke, especially in those with compromised systemic hemodynamics. These patients are prone to watershed infarction, and cerebral venous hypertension may increase this risk.

Intrinsic PEEP

Maneuvers directed at elimination of PEEP, result in a decrease in inspiratory time and an increase in expiratory time. Decreasing the respiratory rate and VT and increasing the IFR decreases intrathoracic pressure, and exhalation decreases that pressure. Venous return principally occurs during exhalation. If the ventilator settings lead to increased intrathoracic pressure during exhalation, venous return will be reduced. Variables that can lead to this circumstance are increased PEEP, auto-PEEP, and IRV. Venous return not only depends on a relatively negative pressure within the thoracic cavity but also relies on a sufficient amount of time for flow into the thoracic vasculature and right side of the heart. Significantly high respiratory rates may compromise venous return. An additional untoward side effect of impaired venous return is cerebral venous hypertension from impeded venous drainage. Because of the absence of valves between the cerebral parenchyma and the right atrium, increased pressure on the right atrium reduces cerebral venous flow and may contribute to cerebral ischemia in patients with traumatic brain injury or stroke, especially in those with compromised systemic hemodynamics. These patients are prone to watershed infarction, and cerebral venous hypertension may increase this risk.

Difficulty Triggering the Ventilator

To trigger a ventilator, a patient must cause either a drop in pressure or an increase in airflow at the proximal airway, depending on the type of settings used. The magnitude of change required to trigger the ventilator is adjusted by setting the sensitivity, usually in the range of −1 to −2 cm H₂O below the level of PEEP. Difficulty triggering the ventilator is often hard to detect. When it becomes obvious by physical examination that the patient is using the accessory muscles of respiration to trigger the ventilator, the problem may be severe. The condition can be detected earlier by inspecting the pressure-volume time curve on the ventilator display, if available. A large negative deflection at the beginning of inhalation suggests that ventilator sensitivity needs to be increased.

Auto-Cycling

Auto-cycling refers to a phenomenon when the ventilator set in AC mode begins to rapidly trigger without the patient initiating respiration. The cause is usually vacillations in airway pressure that the ventilator “interprets” as patient effort. Tremors, shivering, voluntary motion, convulsions, and oscillating water in the ventilator circuit are all potential causes. Auto-cycling should prompt immediate disconnection from the ventilator circuit and ventilation with a bag-valve device until the problem is resolved.

Rapid Breathing

When attempting to ventilate a patient with an obstructive process, the goal is to eliminate PEEP. Permissive hypercapnia is best achieved at low respiratory rates, but at the same time hypercapnia is a powerful stimulus to breathe. This can typically be quelled with a combination of sedation with high-dose opiates, ketamine, or propofol. Neuromuscular blockade should be considered a last resort undertaken only after careful consideration of the risk of prolonged paralysis and potential development of neuropathy in critical illness. If undertaken, the goal should be to weaken the patient sufficiently to inhibit dyssynchrony with the ventilator. Other common causes of rapid breathing include sepsis, pulmonary embolism, pregnancy, hepatic encephalopathy, intracranial hypertension, stroke or hemorrhage, and hypercapnia.

Outstripping the Ventilator and Double Cycling

Hypercapnia will develop in patients receiving low-VT ventilation for ARDS or for an obstructive process and generate an increased respiratory drive. Outstripping the ventilator refers to the patient’s effort to draw a higher VT than set while in a volume-targeted mode. This can be detected by observing the exhaled VT or by finding a negative deflection at the end of inhalation on the pressure-volume time plot. Double cycling occurs when the patient desires a larger VT than is set and continues to inspire despite the delivery of a breath. The ventilator will then provide a second breath almost immediately after the first. This is especially problematic because the actual VT delivered is twice the set volume. As with controlling rapid breathing, the solution is sedation and analgesia, particularly with opiates. Switching to a pressure-targeted mode or increasing the set VT may alleviate this issue.

Straining Over the Ventilator

Straining over the ventilator indicates that the patient on a volume-targeted mode is attempting to inhale at a flow rate in excess of the set IFR. On the pressure-volume time plot, the rise in pressure during inhalation will be concave rather than convex. Potential solutions are to raise the IFR, switch to a pressure-targeted mode or PSV, or use sedation and analgesia.

Coughing

Coughing is a common problem that arises from increased secretions, airway foreign body (ET tube), or an underlying pulmonary disease process. Coughing can lead to auto-cycling, poor patient comfort, dislodgment of the ET tube, and rarely airway injury. Accurate placement of the ET tube above the carina should be confirmed. Suctioning and provision of warmed, humidified air are often helpful. If these simple measures fail to provide relief, aerosolized lidocaine or suppression with opiates may increase patient comfort.
Equipment Failure

Whenever a patient decompensates while undergoing MV, consideration should be given to equipment failure as the cause. Interruption of the oxygen supply, erroneous settings, disconnected ventilator circuitry, and obstructed tubes are all potential culprits. Immediate action should include disconnection from the ventilator and bag ventilation with 100% O_2. The mnemonic made popular by the American Heart Association Pediatric Advanced Life Support Course is useful for recalling the causes of unexpected decompensation: DOPE (Dislodgment of the ET tube, Obstruction of the tube, Pneumothorax, and Equipment failure). Confirmation of ET tube placement, suctioning via the ET catheter, auscultation, chest radiography, and equipment troubleshooting are necessary actions.

TROUBLESHOOTING

Critically ill patients are encountered in the ED every day. Some of them required intubation in the prehospital setting, whereas others are intubated on arrival or during ED evaluation. After their airway has been secured and their condition stabilized, patients may remain in the ED because of a lack of critical care beds. In some hospitals, a dedicated intensivist will take charge of these patients, but in others, this type of coverage may not be available. Acute complications or deterioration must be handled by the emergency physician.

This section provides a framework for managing crashing, mechanically ventilated patients. The information provided will assist the practitioner in determining whether the patient’s condition is related to the underlying pathology that necessitated MV or whether it is being caused by MV itself.

Determine Hemodynamic Stability

The initial step in managing a crashing ventilated patient is to determine the patient’s hemodynamic stability. A ventilated patient is, by definition, critically ill. The patient can fall anywhere on the spectrum from being ventilated for airway protection, with normal vital signs, blood pressure, and oxygen saturation, to being ventilated and in cardiac arrest. Determining where the patient is on this spectrum, including assessment for hypotension and hypoxia, will dictate how much time that the practitioner has to implement rescue strategies. In addition, it is important to anticipate the patient’s clinical course. The approach to a patient who is intubated for hypoxia stemming from pneumonia and whose blood pressure and oxygenation gradually trend down over a period of hours or days is different from the approach to a patient who is declining over a span of minutes (Fig. 8.19).

As a general rule, the following evaluation should be performed on patients with new unexplained hypotension (systolic blood pressure [SBP] < 90 mm Hg), new unexplained hypoxia (SaO_2 < 90%), or a new marked change in vital signs (a drop in SBP by more than 20 mm Hg or a drop in SaO_2 by more than 10%) within an hour. Patients with stable SBP between 80 and 90 mm Hg and SaO_2 between 80% and 90% should be evaluated expeditiously in the hope of halting the decline. Those with SBP lower than 80 mm Hg or SaO_2 less than 80%, and those who continue to decline rapidly, should be evaluated quickly, with consideration given to entering the cardiac arrest/near arrest algorithm. These values are arbitrary demarcation points and do not take precedent over bedside clinical judgment.

Cardiac Arrest and Near Arrest Patients

Time is of the essence in a patient with cardiac arrest or near arrest. Advanced cardiac life support should be implemented quickly, and there are some key points to remember in a ventilated patient in cardiac arrest or one who becomes acutely hemodynamically unstable. The ED practitioner should develop a stepwise approach in this situation. During each step, the practitioner should “look, listen, and feel” to run through the differential diagnosis.

During stabilization of these patients it is important to keep in mind the original pathology that necessitated intubation. Sudden decompensation in an asthmatic is a common example of physician intervention worsening the scenario unless the pathophysiology of the arrest is appreciated (see Fig. 8.17). A crashing ventilated patient may simply be worsening from the primary pathology. A multitrauma patient may have an intra-thoracic or intra-abdominal catastrophe, and a septic patient may be deteriorating clinically from lack of source control.

It is also important to determine and address special circumstances that the ventilator can precipitate, the most significant of which are tension pneumothorax and severe auto-PEEP. Tension pneumothorax can lead to marked hypotension as a result of decreased cardiac output and marked hypoxia from V/Q mismatch. Auto-PEEP (also referred to as PEEP, breath stacking, or dynamic hyperinflation) is caused by trapped volume in the pulmonary system. If severe enough, it will eventually lead to increased intrathoracic pressure. This can cause hypotension and decreased cardiac output from decreased venous return and marked hypoxia from V/Q mismatch.

In critically ill ventilated patients with respiratory distress who are hemodynamically unstable, the following steps will assist the ED physician in determining the cause of the decompensation.

Step 1: Disconnect the Patient From the Ventilator

This is perhaps the easiest step to perform. It can be both diagnostic and therapeutic in a crashing ventilated patient. A quick rush of air or prolonged expiration of trapped air from the ET tube can be diagnostic of ventilator-induced auto-PEEP. A few seconds of observation can determine whether this entity is present. Return of hemodynamic stability implies that the maneuver was successful.

Patients undergoing cardiopulmonary resuscitation (CPR) should not be connected to a ventilator. The variations in intrathoracic pressure caused by CPR will trigger ventilator breaths at high rates if set on AC mode. Patients treated with inhaled nitric oxide (iNO) should not be removed from iNO abruptly, and effort should be made to quickly establish this supply through the bag-valve system. In addition, care should be taken when disconnecting patients who are on high PEEP, such as those with ARDS. When disconnecting the ventilator, it is important to address auto-PEEP because derecruitment can occur and hypoxia may be worsened. Once auto-PEEP has been ruled out, PEEP valves may be used to maintain PEEP levels and thus avoid derecruitment. PEEP valves may be problematic in a markedly hypotensive patient. They may increase intrathoracic pressure and decrease venous return.
Step 2: Breathing—Hand-Ventilate With 100% Oxygen
Ensure that 100% oxygen is being delivered and limit the respiratory rate to 8 to 10 breaths/min. Particular attention should be paid to the delivery of hand ventilation. Inadvertent rates as high as 40 breaths/min are often used in codes. Excessive rates will increase intrathoracic pressure and lead to a decrease in venous return and cardiac output. Look at both sides of the chest to determine whether there is equal chest rise. Unequal chest rise can signify main stem intubation, pneumothorax, or a mucous plug. Listen for air escaping from the mouth or nose (a sign of an air leak). Listen over the epigastric area and in both axillae. Decreased breath sounds may provide clues regarding main stem intubation, pneumothorax, or an atelectatic lung. Feel for subcutaneous crepitus (a sign of pneumothorax) and assess for difficulty in hand ventilating (a sign of low dynamic or static respiratory system compliance).

Step 3: Airway—Determine That the ET Tube Is Functioning and in the Proper Position
The ET tube functions by providing a conduit to the lower part of the trachea. The cuff creates a seal between it and the inner wall of the trachea. To determine whether the ET tube is functioning properly, pass the suction catheter and listen for an air leak (Fig. 8.20). Easy passage of the suction catheter does not guarantee that the ET tube is in the trachea because the catheter may be passing down the esophagus. If it is difficult or not possible to pass the suction catheter, the ET tube is either dislodged or obstructed. Attempt to correct this by repositioning the head in the case of a twisted or bent ET tube or inserting a bite block if the patient is biting on the tube. Dislodged or obstructed ET tubes require reintubation. Patients with dislodged ET tubes should be treated as difficult intubations because unplanned extubations are notorious for causing trauma to the glottis and for leading to vocal cord edema.

In a cardiac arrest or near-arrest patient, the best choice for determining that the ET tube is in the proper position is direct visualization of the ET tube passing through the cords. This step is often omitted in a crashing ventilated patient because of the belief that the ET tube has not migrated. Unfortunately, unrecognized ET tube migration can occur during routine care of a critically ill patient. Patients are frequently moved in and out of emergency medical service vehicles, transferred to and from stretchers for imaging studies, and turned for procedures or bathing, all of which are capable of dislodging the tube. This visualization step can be performed while providing ventilation by hand.
Other simple techniques may be used to confirm that the ET tube is in the trachea. Direct visualization of the carina with a fiberoptic scope is an option, but this device is not typically readily available to the ED practitioner. Another quick and readily available technique is to pass an intubating stylet (gum elastic bougie or Eschmann introducer). The stylet is passed gently through the ET tube. If resistance is met at 30 cm, the ET tube is most likely in the trachea. If the stylet passes beyond 35 cm without resistance, the ET tube is probably in the esophagus. If resistance is met too soon, the intubating stylet may be catching on the ET tube.

At least one of these techniques to determine proper positioning should be used early enough in the resuscitation to correct any airway issues. In addition, proper positioning should be confirmed before simply removing the ET tube and reintubation, particularly if the patient is deemed to have a difficult airway (unless it is glaringly evident that the patient is extubated).

**Step 4: Special Procedures**

If the patient is still in cardiac arrest or near arrest after being disconnected from the ventilator, ensuring proper placement of the ET tube, and hand-ventilating with 100% oxygen, a clinical decision will be required regarding needle decompression of the chest. If time permits, a focused history from the bedside nurse, respiratory therapist, or paramedic and a focused physical examination will indicate which side of the chest to decompress. In addition, depending on the urgency of the situation, bedside US and chest radiography may be used. Bedside US has been shown to exclude pneumothorax in the presence of “lung slide.” This is depicted in M-mode as the “seashore sign” (Fig. 8.21) and, in its absence, as the “stratosphere sign/bar code sign” (Fig. 8.22).

At times, the clinical situation does not allow imaging studies, and the focused history and physical examination may not be helpful. In these cases, needle decompression of both sides of the chest should be considered if other more likely causes of acute decompensation are not found. It is important to remember that chest tube placement is required in patients after needle decompression.

**Stable and Nearly Stable Patients**

If the patient is deemed stable or near stable or quickly regains stability after disconnection from the ventilator and ventilation by hand, the event should be approached in a systematic manner.
The patient should be placed on 100% oxygen during this evaluation.

**Step 1: Obtain a Focused History**
A focused history should be obtained from the practitioners most involved in the patient's care (bedside nurse, respiratory therapist, resident, paramedic). Valuable information includes the indication for intubation, difficulty of the intubation, depth of the ET tube, ventilator settings, and recent procedures or moves (central line insertion; chest tube placement; removal or transition to a water seal; thoracentesis; ET tube manipulation; transport off the stretcher; rotation for cleaning, a procedure, or chest radiography).

**Step 2: Perform a Focused Physical Examination**
Take a general survey of the patient. Observe for agitation, attempts to pull at the ET tube and lines, gasping for breath (the patient will have the mouth open and appear dyspneic), and tearing of the eyes.

**Airway**
Look at the ET tube and determine if it has migrated from its previous position. It is possible that the ET tube has migrated out of the trachea or into a main bronchus. Adjust if necessary. Listen for escaping air (an air leak) from the mouth or nose (see Fig. 8.20). This typically signifies that the tube has lost its seal with the trachea and occurs with extubation or cuff failure. Feel the pilot balloon. If it is deflated, the cuff is deflated. Add air to the pilot balloon. If this stops the air leak, make a note that air was added to the balloon. If the pilot balloon does not inflate or deflates with time, there is a defect in the pilot balloon-cuff apparatus, and the ET tube will probably need to be exchanged. Occasionally, it may be possible to repair the pilot balloon mechanism with commercially available kits. This is a good option in patients who are difficult to intubate.

Determine that the ET tube is functioning properly by passing the suction catheter. If it is difficult or not possible to pass the suction catheter, the ET tube is either dislodged or obstructed. Attempt to correct by repositioning the head in the case of a twisted or bent ET tube or inserting a bite block if the patient is biting on the tube. Dislodged or obstructed ET tubes require reintubation.

Determine that the ET tube is in proper position if at any point in the evaluation it is suspected that extubation has occurred. Any of the techniques discussed in the previous section may be used.

**Breathing**
Look at both sides of the chest to determine whether there is equal chest rise. Unequal chest rise can signify main stem intubation, pneumothorax, or a mucus plug. Look at the ventilator tubing and determine whether there is an oscillating water collection. Listen for air escaping from the mouth or nose (a sign of an air leak). Listen over the epigastric area and in both axillae. Decreased breath sounds may provide clues regarding main stem intubation, pneumothorax, or an atelectatic lung. Feel for subcutaneous crepitus (a sign of pneumothorax).

**Circulation**
Check for pulses and cycle the blood pressure cuff frequently. If the patient has an arterial line, make sure that the transducer is level. Determine the need for fluid bolus or vasopressors.

**Step 3: Assess Gas Exchange**
Hypoxia can be diagnosed with pulse oximetry if the waveform is reliable. The waveform should not be highly variable and the frequency of the waveform should match the heart rate on the cardiac monitor. In a few instances, such as carbon monoxide poisoning, pulse oximetry is not reliable. In these cases or if the pulse oximeter is not picking it up, an ABG sample should be obtained. Patients with a partial pressure of O₂ (PaO₂)/FiO₂ ratio of less than 200 should be evaluated for ARDS. Those with a ratio between 200 and 300 should be evaluated for ALI. A lung-protective strategy should be implemented in those determined to have ALI or ARDS (see Fig. 8.18). Hypoventilation cannot be identified with pulse oximetry. ABG analysis is beneficial in this event.

**Step 4: Check Respiratory Mechanics**
Determine whether peak pressure and plateau pressure have changed from their previous values. These values should be obtained on volume-targeted modes. Airway pressure is a function of volume and respiratory system compliance. The respiratory system incorporates the ventilator circuit, ET tube, trachea, bronchi, pulmonary parenchyma, and chest wall. A set volume with a set system compliance results in a specific pressure. Peak pressure is a function of volume, resistance to airflow, and respiratory system compliance. Plateau pressure is obtained during an inspiratory pause, thus eliminating airflow, and therefore reflects only respiratory system compliance. An isolated increase in peak pressure is indicative of increased resistance to airflow. An isolated increase in plateau pressure is indicative of a decrease in respiratory system compliance. Note that plateau pressure can never be higher than peak pressure and that if plateau pressure rises, so will peak pressure. It is important to keep in mind the Δ relationship (peak pressure – plateau pressure). These measurements assume a comfortable patient, and peak pressure and plateau pressure values are not reliable in a “bucking” patient.

**Step 5: Observe Ventilator Waveforms**
The two most helpful ventilator waveforms are the flow-time curve and the pressure-time curve. The flow-time curve can be used to detect air trapping. The pressure-time curve can be used to determine plateau pressure with an inspiratory hold (see Fig. 8.1).

A notching in the pressure-time curve during inspiration can signify air hunger. In this situation, the patient desires a higher flow rate than the ventilator is delivering. It is commonly seen in volume-targeted modes. Increasing the flow rate will often alleviate this phenomenon. Another solution is to change to a pressure-targeted mode. Double cycling can also be seen on ventilator waveforms. This occurs when the patient desires a higher Vt than the ventilator is set to deliver. The patient is still inspiring when the first breath has finished cycling and the ventilator immediately gives a second mechanical breath. This is frequently seen with low-Vt ventilation, which is used in patients with ARDS and status asthmaticus. It is important to recognize because the actual Vt being provided is essentially twice the set Vt. This has important ramifications for patients with ARDS and obstructive processes such as asthma and COPD, for whom the goal is lower Vt. Typically, improved sedation with emphasis on blunting the respiratory drive with opiates alleviates double cycling. Other adjustments that may prove helpful are increasing the flow rate, increasing Vt by 1 mL/
Step 6: Imaging Studies—Chest Radiograph and Bedside US
Evaluate the chest radiograph for ET tube position, main stem intubation, lung atelectasis, pneumothorax, and a worsening parenchymal process. Bedside US, if available, is typically quicker in evaluating for pneumothorax, but it will not provide information on the location of the ET tube, lung atelectasis, or parenchymal processes (see Figs. 8.21 and 8.22).

Step 7: Evaluate Sedation
Patients requiring MV often need sedation and analgesia to make the ET tube and ventilation tolerable. Some patients, such as those with drug overdoses or traumatic head injuries, may not require any sedation. Others may tolerate intubation quite well while almost fully awake. The majority of patients require some form of sedation or analgesia. Note that a plethora of non–MV-associated conditions, such as unrecognized bladder obstruction, alcohol or drug withdrawal, occult fractures, compartment syndrome, or bowel ischemia, can cause significant agitation that can mistakenly be attributed to the stress of intubation and ventilation.

Selection of agents should be based on the desired effect. If a patient appears agitated, sedative-hypnotics should be used. Such drugs include benzodiazepines, propofol, dexmedetomidine, and haloperidol. It is important to note that these agents do not provide an analgesic component. If a patient is being given adequate sedative doses and still appears agitated, consider pain as a cause. Typical opiates that can be used are fentanyl, hydromorphone, and morphine. A remifentanil infusion is ultrashort acting and can provide both sedation and analgesia. Remifentanil does not accumulate in patients with renal or hepatic insufficiency. Prompt reversal of analgesia and sedation are seen on discontinuation. Remifentanil is an alternative to fentanyl for patients requiring frequent neurologic assessment or those with multiorgan failure.

The goal of sedation and anesthesia in ventilated patients who are not being evaluated for extubation is one in which the patient will arouse with gentle stimulation but will return to a sedated state when left alone. Patients who are being sedated and require deep stimulation to get a response are oversedated.

Patients who display air hunger and have a high respiratory rate can be given a trial of opiates to relieve their symptoms. Proper sedation and analgesia are paramount in patients being treated with a strategy that allows or permits hypercapnia, such as those with status asthmaticus, and in patients being treated with lung-protective strategies, such as those with ARDS. Hypercapnia is a powerful stimulus to the respiratory drive, and opiates are often required to control respiratory rates. Patients who tend to be difficult to control (besides those with status asthmaticus and ARDS) include patients with hepatic encephalopathy or intracranial processes such as a mass effect or hemorrhage. Chemical weakening with intermittently dosed paralytics may be required if patients have undergone a good trial of sedation, analgesia, and ventilator changes and are still markedly tachypneic. Careful consideration should be given before this step because prolonged paralysis has been implicated in critical illness polyneuropathy. In addition, expert consultation should be obtained before prolonged paralysis of a neurosurgical patient. The goal in chemical paralysis in these patients is to weaken them enough to control their interaction with the ventilator. Usually, this does not require a full dose of the paralytic.

Hemodynamic instability in mechanically ventilated and sedated patients may be a result of medications because sedatives and analgesics can precipitate or worsen hypotension. As a general rule, continuous infusions should be withheld in these cases. Patients who are hypoxic and agitated but not hypotensive may benefit from improved sedation. It is possible that their pulmonary status is so tenuous that they are agitated from the hypoxia and their condition is worsened by the oxygen consumption caused by their agitation. Patients who are agitated and hypotensive may respond well to a low-dose benzodiazepine and opiate if the agitation is a precipitant of hypotension. In all these cases, it is imperative to determine whether sedation is a factor in the decompensation. Chemical paralysis should be reserved as a final option. It is important to remember that without continuous electroencephalographic recordings, seizure activity cannot be monitored if the patient is paralyzed.

SPECIAL SCENARIOS

Two special scenarios should be mentioned. One is a crashing intubated pediatric patient and the second is a patient with a tracheostomy. The approach described earlier can be used in pediatric patients, but there are a few caveats that may improve the approach. The first is to recognize that migration of the ET tube is common with small movements of the head and neck. A simple solution is to use a cervical collar for immobilization. Second, small ET tubes are often uncuffed and do not have a pilot balloon. Air leaks in this scenario should prompt the clinician to consider that the ET tube is either dislodged or too small. Finally, specialized equipment such as intubating stylets and fiberoptic scopes are typically not available in pediatric sizes.

Important questions that have ramifications in the care of a crashing ventilated patient with a tracheostomy are the following: (1) Does the patient have a laryngectomy? (2) Why does the patient have a tracheostomy? and (3) How old is the tracheostomy? These are important questions because patients with a laryngectomy cannot be intubated orally, patients with a tracheostomy secondary to anatomic considerations or difficult or failed airways may be difficult to intubate orally, and the tract in a patient with a recent tracheostomy (less than a week old) may not have matured enough to safely reintroduce a tracheostomy tube.

LIBERATION FROM THE VENTILATOR

Occasionally, patients can be considered for extubation while still in the ED. Before extubating a patient, several questions should be answered in the affirmative (Fig. 8.23). Common scenarios include resolution of exacerbations of COPD or asthma, exacerbation of congestive heart failure, and metabolism of intoxicants. Patients who were difficult intubations or required multiple attempts should have a planned extubation. It may be prudent to allow the edema from a traumatic intubation to subside.

Once these screening questions have been answered in the affirmative, one should perform an awakening trial followed
by a spontaneous breathing trial. Extubation should be considered if the patient does not fail the breathing trial.

For patients intubated for exacerbation of COPD, NPPV should be considered. For patients with COPD who failed a spontaneous breathing trial, extubating to NPPV decreases mortality, hospital length of stay, the incidence of ventilator-associated pneumonia, and the total duration of MV.83

**CONCLUSION**

Mechanically ventilated patients are typically the most critically ill patients that the ED practitioner will manage. The underlying disease process that required intubation is typically life-threatening. When patients become unstable, the physician should take a stepwise approach toward determining whether the patient is deteriorating because of the underlying disease process or because of interaction with the ventilator. It is hoped that the approach presented here will assist practitioners with a framework to evaluate and stabilize crashing ventilated patients. For further information, Wood and Winters present an up-to-date review of the care of ventilated patients in the ED and evaluation of potential problems.84

---

Figure 8.23 Liberation from ventilator. ED, Emergency department; $\text{FiO}_2$, fraction of inspired oxygen; PEEP, positive end-expiratory pressure; PSV, pressure support ventilation; RR, respiratory rate.
REFERENCES


Thoracentesis is a percutaneous procedure during which a needle is inserted into the pleural space to remove fluid for diagnostic or therapeutic purposes. When performed using real-time ultrasound (US) guidance, thoracentesis has an extremely low rate of complications.

ANATOMY

The thoracic cavity is lined by thin serous membranes: the visceral pleura, which wraps around the lungs, and the parietal pleura, which lines the inner surface of the thoracic cavities and meets the visceral pleura at the root of the lungs in the mediastinum. The space between these two linings is called the pleural space. Each pleural membrane consists of a superficial mesothelial cell layer and an underlying connective tissue layer containing systemic capillary beds. These membranes are leaky, allowing for easy movement of both fluid and protein. In the healthy state, hydrostatic pressure of the systemic capillaries allowing for the movement of fluid and protein. In the healthy state, hydrostatic pressure of the systemic capillaries produces a small net movement of fluid across both pleurae into the low-pressure pleural space.

Because the lymphatic system has the ability to remove 30 times more pleural fluid than its basal rate, it is believed that in most instances the development of a pleural effusion requires both an increase in fluid entry into the pleural space and a decrease in its removal.

ETIOLOGY OF PLEURAL EFFUSIONS

The most common causes of pleural effusion in adults are congestive heart failure (CHF), pneumonia, malignancy, pulmonary embolism (PE), and cirrhosis. The most common cause in children is pneumonia, followed by heart failure, and malignancy. Pleural effusions can be classified as either transudates or exudates. Distinguishing between transudates and exudates narrows the differential diagnosis and directs management and therapy. A comprehensive list of causes can be found in Box 9.1. The most common are discussed in the following sections.

Transudates: Overwhelming the System

Transudates are caused by either an increase in intravascular hydrostatic pressure or a decrease in intravascular oncotic pressure. This generates an increased flow of fluid into the pleural space across noninjured capillary beds. These effusions are typically straw colored and serous with very low cellular and protein content.

The most common cause of a transudate is CHF. The increased pulmonary capillary hydrostatic pressure in patients with CHF results in a net flow of fluid into the pulmonary interstitium. This fluid readily moves across the leaky visceral pleura into the pleural space. In addition, elevated systemic capillary pressures associated with CHF increase fluid flow across the pleural membranes into the pleural space and decrease lymphatic flow out of the thorax. Any process that causes compromised left ventricular outflow can result in a pleural effusion, including myocardial infarction, cardiomyopathy, and valvular disease. These effusions are typically small and bilateral with larger effusions on the right (Fig. 9.1). Highly asymmetric and large effusions should raise suspicion for another cause.

Patients with cirrhosis are frequently hypoalbuminemic, which leads to a chronic state of decreased plasma oncotic pressure. This imbalance between the hydrostatic and oncotic forces across the pleural membrane results in an effusion. In addition, experiments have shown that high volumes of ascites can stretch the diaphragm enough to allow fluid to pass through preexisting microdefects.

Similarly, in several other disease states such as the hypoalbuminemic state of patients with nephrotic syndrome, intraabdominal fluid associated with peritoneal dialysis, or retroperitoneal fluid with obstructive uropathy can cause transudative pleural effusions in patients with renal disease.

Exudates: Pathology of Tissues, Destroying the System

Exudates are caused by pleural inflammation, increased pleural membrane permeability, or lymphatic obstruction. More than 90% of exudative effusions are due to malignancy, pneumonia, PE, and gastrointestinal diseases (e.g., pancreatitis, esophageal perforation). Tuberculosis (TB) should be added to this list in patients from high–TB prevalence regions.

The primary mechanism of cancer-related pleural effusion is obstruction of pleural lymphatic outflow, either via damage to lymphatic stomata or more distal involvement of the mediastinal lymph nodes. Other mechanisms include metastases to the pleurae increasing capillary permeability and obstruction of the thoracic duct with consequent chylothorax.

A pleural effusion associated with pneumonia or a lung abscess is termed a parapneumonic effusion. In the first, or exudative, stage of parapneumonic effusion, an increase in pulmonary interstitial fluid leads to flow of sterile fluid and inflammatory cells across the visceral pleura into the pleural space. When the infection and resultant inflammation continue unchecked, there is increased capillary permeability with bacterial and additional inflammatory cell transfer to the pleural space, causing the parapneumonic effusion to become complicated. This worsening pleural space inflammation results in the deposition of fibrin on the pleural surfaces with formation of loculations, and finally, in a tough peel that encases the lung.

A pleural effusion develops in at least 20% of those with PE. Therefore when the cause of the effusion is unclear, PE should be strongly considered. The effusions associated with PE are often too small to require thoracentesis, but they have been shown to be uniformly exudative, probably resulting from ischemia- and infarction-induced increases in pulmonary capillary permeability.
Traumatic Effusions: Acute and Catastrophic Destruction of the System

Effusions can also result from significant injuries such as the following. Esophageal rupture can result from forceful vomiting, as in the case of Boerhaave’s syndrome, or from instrumentation, as in the case of endoscopy, gastroesophageal balloon tamponade tube placement, or rigid nasogastric tube placement. Hemothorax can result from traumatic injuries, cannulation of the subclavian vein or artery, PE, or aortic aneurysm rupture. Chylothorax develops from acute disruption of the thoracic duct, usually in the setting of trauma or malignancy. These effusions are usually large-volume collections that accumulate over an extremely short period and rapidly compromise both oxygenation and circulation.

CLINICAL PRESENTATION

Symptoms of small pleural effusions are usually representative of the underlying disease process and not the effusion itself. An effusion greater than 300 mL may cause symptoms on its
own, usually dyspnea, chest pain, or cough. The dyspnea from a larger pleural effusion is typically the result of impaired chest wall and diaphragm movement and not decreased oxygenation or lung function.12 Chest pain may be dull and aching from the effusion itself or pleuritic from localized irritation of the parietal pleura, which has abundant nerve fibers. Cough is usually attributed to bronchial irritation from compression of the lung parenchyma. Patients occasionally present in extremis from a large pleural effusion that is compressing the heart and causing tamponade physiology with cardiovascular compromise.13

There are numerous described physical exam findings for pleural effusion, but these may be difficult to appreciate in the emergency department (ED) environment depending on the acuity of the patient and the ambient noise (Box 9.2). Dullness to percussion and decreased tactile fremitus over the effusion have been shown to be the most useful physical exam findings on systematic review.14 Percussion over the effusion produces a characteristic dull resonance, which shifts when the patient changes position if the fluid is free flowing. Placing the flat hand over the patient’s back while the patient says the word “toy” or the phrase “blue moon” will demonstrate a reduced or absent fremitus because the fluid separates the lung from the thoracic wall and absorbs vibrations from the lung.

**RADIOLOGIC DIAGNOSIS**

**Chest Radiograph**

Because pleural fluid is denser than air-filled lung, a free-flowing effusion will first accumulate in the most dependent parts of the thoracic cavity: the subpulmonic space and the lateral costophrenic sulcus. Pleural effusions are usually visible on an upright chest radiograph if 200 to 250 mL of fluid is present. A lateral radiograph may reveal an effusion of 50 to 75 mL.

The earliest recognized sign of a pleural effusion on an upright chest radiograph is blunting of the lateral costophrenic angle, which may be seen on either the frontal or the lateral view (Fig. 9.2). With a larger free-flowing effusion, the pleural fluid appears as a meniscus that curves downward toward the mediastinum in the frontal view and appears “lowest” midway through the thoracic cavity on the lateral view (Fig. 9.3). The presence of air from pneumothorax or abscess may alter the appearance of the meniscus to more of a straight line (air-fluid level).

Occasionally, up to 1000 mL of fluid collects in the subpulmonic space and causes neither blunting of the costophrenic angle nor a meniscus appearance on the upright radiograph. This is called a **subpulmonic effusion** (Fig. 9.4). This should be

---

**BOX 9.2 Exam Findings in Pleural Effusion**

- Decreased chest wall excursion
- Bulging intercostal spaces
- Egophony at superior border of effusion
- Diminished breath sounds
- Dullness to percussion
- Decreased tactile fremitus
- Pleural friction rub

---

![Figure 9.2](image1.png) Posteroanterior upright chest radiograph indicating a pleural effusion with a bilateral meniscus at the costophrenic angles. The pleural effusion has a curvilinear upper margin concave to the lung and is higher laterally than medially.

![Figure 9.3](image2.png) Left-sided pleural effusion seen on A, posteroanterior and B, lateral radiographs. The meniscus can be visualized on both views (arrows).
suspected if the hemidiaphragm is elevated and the hemidiaphragm dome peaks more laterally than expected on the upright frontal radiograph.

Pleural effusions are challenging to identify on chest radiograph in the supine patient, and even a significant amount of fluid may not be appreciated. If the effusion is large enough, a diffuse haziness may be appreciated (Fig. 9.5). Other findings include apical capping, obliteration of the hemidiaphragm, partial opacification of a hemithorax, and a widened minor fissure.

Obtaining bilateral decubitus radiographs when a pleural effusion is seen or suspected will confirm the presence of a free-flowing effusion and allow for visualization of loculations, contained abscesses, infiltrates, or masses. With the side of the effusion down, a simple pleural effusion will follow gravity and layer between the floating lung and the chest wall (Fig. 9.6). A lateral decubitus view on the opposite side draws the fluid toward the mediastinum and allows further visualization of the lung parenchyma.

With a diseased or scarred lung, tissue adhesions can trap pleural fluid within the parietal, visceral, or interlobar surfaces. Because these adhesions anchor the fluid, loculated effusions are often described as “D-shaped” (Fig. 9.7). Fluid loculated in the fissures assumes a lenticular shape.

In the case of a massive pleural effusion, the entire hemithorax is opacified (Fig. 9.8). On such films, identification of mediastinal shift is a key to identifying the underlying disease process. In the absence of a diseased lung or mediastinum, large fluid collections push the mediastinum contralaterally. When the mediastinum is shifted toward the effusion, the lungs and main stem bronchi are diseased, obstructed, or both. When the mediastinum is fixed midline, it is likely invaded by tumor.

**Computed Tomography**

Computed tomography (CT) is more sensitive than plain films in detecting very small effusions and can readily assess the
There are definite advantages to using US for assessment of pleural effusions as well. In particular, it is noninvasive and can be performed at the bedside. US is superior to chest radiographs in diagnosing effusion and can detect effusions as small as 5 mL. Although some details can be seen only with CT, US can identify fluid loculations, separate fluid from pleural thickening, and distinguish solid from fluid pleural lesions. US can also be used to identify both pulmonic and abdominal causes of pleural effusion (Fig. 9.9). Most importantly, US-guided thoracentesis is associated with a significantly lower rate of complications.

**INDICATIONS**

Thoracentesis can be performed for either diagnostic or therapeutic purposes. With diagnostic thoracentesis, 50 mL of pleural fluid is obtained for laboratory studies to elucidate the cause of effusion. Most new effusions that measure greater than 1 mm on decubitus radiograph, CT, or US require diagnostic thoracentesis unless there is a clear clinical diagnosis (e.g., CHF) and no evidence for superimposed pleural space infection. Therapeutic thoracentesis is performed to relieve the dyspnea associated with a large pleural effusion and typically involves removal of 1.5 L of fluid.

**CONTRAINDICATIONS**

There are no absolute contraindications to thoracentesis. One relative contraindication is cutaneous infection or herpes zoster of the overlying chest wall. In this case an alternate patient position and site can typically be chosen. Preprocedure
withdrawn, a 2-inch 22-gauge needle (spinal needle for larger patients) can be used. For therapeutic procedures, using an over-the-needle thoracentesis catheter (e.g., 8-French catheter over an 18-gauge needle) avoids prolonged insertion of a needle in the pleural space while large volumes of fluid are being removed. Commercial thoracentesis catheters have built-in safety features which may include a blunt spring-loaded safety cannula that extends beyond the sharp needle tip once the pleural space is entered to protect the lung from puncture, a one-way valve that prevents air entry into the catheter during removal of the needle, and a built-in side port for drainage of fluid (Fig. 9.10). It is important to familiarize yourself with the features of the thoracentesis catheter available in your institution.

**Figure 9.7** Loculated pleural effusion. A, A posteroanterior radiograph demonstrates the D-shaped appearance of a right-sided loculated pleural effusion (arrows) in the midchest region. B, Occasionally, pleural effusions (arrows) may become loculated in the fissures.

**Figure 9.8** Massive pleural effusion A, before and B, after thoracentesis. Notice the underlying mass that is seen following thoracentesis. A computed tomography scan would have prospectively confirmed this radiograph.

coa... 

**TECHNIQUE**

Thoracentesis is usually an elective procedure. Obtain and document informed consent, according to hospital policy, before starting the procedure. Follow sterile technique throughout the entire procedure to avoid introduction of infection.

**Equipment**

The specific device used for thoracentesis depends on type of procedure, hospital availability, and provider experience. For diagnostic purposes in which a small volume of fluid is being withdrawn, a 2-inch 22-gauge needle (spinal needle for larger patients) can be used. For therapeutic procedures, using an over-the-needle thoracentesis catheter (e.g., 8-French catheter over an 18-gauge needle) avoids prolonged insertion of a needle in the pleural space while large volumes of fluid are being removed. Commercial thoracentesis catheters have built-in safety features which may include a blunt spring-loaded safety cannula that extends beyond the sharp needle tip once the pleural space is entered to protect the lung from puncture, a one-way valve that prevents air entry into the catheter during removal of the needle, and a built-in side port for drainage of fluid (Fig. 9.10). It is important to familiarize yourself with the features of the thoracentesis catheter available in your institution.

Gather all the necessary equipment before the procedure (see Review Box 9.1 for a list of recommended equipment). If the procedure is diagnostic, remember to draw and send peripheral blood to the laboratory for serum protein and lactate dehydrogenase (LDH) measurement. Keep atropine available at the bedside in case the patient has a vagal reaction during the procedure. Monitor oxygen saturation by pulse oximetry and administer supplemental oxygen as needed.
Thoracentesis

**Indications**
- Suspected pleural space infection
- New effusion without a clear clinical diagnosis
- Relief of dyspnea associated with a large effusion

**Contraindications**
- **Absolute**
  - None
- **Relative**
  - Severe clotting abnormality
  - Infection or herpes zoster at selected site

**Complications**
- Pneumothorax
- Reexpansion pulmonary edema
- Cough
- Air embolism
- Infection
- Catheter fragment in the pleural space
- Hemothorax
- Intraabdominal hemorrhage

**Equipment**
- Skin cleanser
- Gauze
- Sterile drape
- 2 10-mL syringes
- Blood gas syringe
- Blood culture bottles
- Lidocaine
- Scalpel
- Over-the-needle catheter
- Occlusive dressing
- 60-mL syringe
- Large evacuated container
- 3-way stopcock
- Occlusive tubing
- 25-gauge needle
- 60-mL syringe

**Patient Position and Site of Entry**

Fig. 9.11 illustrates various patient positions for thoracentesis. Upright positioning is desired for draining most pleural effusions. For this position, have the patient sit upright on the edge of the bed with arms extended on a bedside table or Mayo stand (Fig. 9.12). If the effusion is sufficiently large, allow the patient to lean forward slightly. Locate the height of the effusion and the level of the diaphragm using US (Fig. 9.13 and Ultrasound Box 9.1). The site of entry should be one to two intercostal spaces below the highest level of the effusion in the midscapular or posterior axillary line (Fig. 9.14). In all
cases, the entry level must be above the ninth rib as the risk for diaphragmatic or hepatic/splenic injury increases with entry below this level. Also avoid entry medial to the midscapular line, as the intercostal artery runs more centrally and tortuously when it first enters the intercostal space at the spine, increasing the risk for arterial injury and hemothorax.\textsuperscript{19–21} Greater than 10 mm of fluid thickness should be present on US at the selected site. Mark your entry site using a skin marker. If the patient is too ill to sit upright, perform the procedure with the patient in the lateral decubitus position, with the side of the effusion down and the back at the edge of the bed. Insert the needle at the posterior axillary line in this position. Alternatively, position the patient supine with the head of bed elevated as much as possible. Use the midaxillary line as the point of needle insertion for this position. For both these alternate positions, determine the fluid level and thickness by bedside US, and make sure the selected site is above the ninth rib. Take a time-out immediately before the procedure to verify the correct patient, procedure, and site.

**Anesthesia and Pleural Fluid Localization**

Once the skin has been sterilized and draped, create a skin wheal at the upper edge of the rib just below the marked entry site using 1% or 2% lidocaine and a 25-gauge needle. Use the upper edge of the rib as the needle insertion site to minimize

---

**Figure 9.11** A–C, Various patient positions for thoracentesis. Note the midscapular line when the patient is sitting upright. This anatomic line is important in that thoracentesis should not be performed medial to this marker because of the increased incidence of trauma to the neurovascular bundle.

**Figure 9.12** Preferred positioning for thoracentesis by the posterior approach is with the patient sitting upright and leaning over an adjacent Mayo stand.

**Figure 9.13** Ultrasound image of a right-sided pleural effusion. Ultrasound can be used during the procedure itself if a sterile probe cover is used. This enables the operator to view the tip of the needle entering the effusion and to thus avoid vital structures such as the lung or diaphragm. Note the superior, inferior, and medial landmarks that are used during the procedure itself.
ULTRASOUND BOX 9.1: Thoracentesis

by Christine Butts, MD

Ultrasound is an excellent tool for both identifying and directing the aspiration of pleural effusions. It can be used to both “mark the spot” for thoracentesis and directly guide the procedure.

Equipment

A 3.5- to 5-mHz transducer will reliably identify most pleural effusions. A higher-frequency transducer should be used when direct visualization of the procedure is desired.

Interpretation of Images

A normal lung is filled with air and will typically appear as a hazy gray area when viewed by ultrasound (Fig. 9.US1). Reverberation or comet tail artifacts may also be seen (Fig. 9.US2). When pleural fluid is present, it will typically be seen in the most dependent area of the thorax, typically in the recesses above the diaphragm. Pleural fluid will appear as anechoic (black) collections of varying size (Fig. 9.US3). Depending on the consistency of the fluid, it may appear heterogeneous, with areas of lighter gray representing more solid components (such as clotted blood).

Procedure and Technique

Once a pleural effusion is known or suspected to be present, scan the area over the posterior aspect of the thorax with a low-frequency transducer (Fig. 9.US4). Identifying the abdominal organs (such as the kidney, liver, and spleen) and diaphragm first will often help orient the sonographer (Fig. 9.US5). Assess the recess superior to the diaphragm for the presence and size of the effusion. If the procedure is to be performed blindly (such as with a very large effusion), mark a location several rib spaces above the diaphragm to avoid any possible intraabdominal injury. The procedure can then proceed in the usual fashion. If the procedure is to be performed under direct ultrasound guidance (such as with smaller effusions), cover the high-frequency transducer with a sterile sheath. Once the field has been prepared, use the sterile transducer again to locate the fluid pocket. At this time,

Figure 9.US2 This image demonstrates two key artifacts seen in a normal lung. As with the previous image, the pleura is recognized as a hyperechoic horizontal line deep to the ribs (at the far right and left of the image). A comet tail, or a small vertical line, can be seen extending deep to the pleura (arrow). This is a normal artifact created by the pleura. A-lines, or a series of horizontal lines extending deep to the pleura, can also be seen (arrowhead). These artifacts are created by reverberation and may be seen in a normal aerated lung.

Figure 9.US3 Large pleural effusion seen on ultrasound with a low-frequency transducer. In this longitudinal image, the thorax is seen on the left side of the image, and a large anechoic (black) fluid collection can also be seen (arrow). Within the thorax the lung can be seen as the hypochogenic (dark gray) mass on the left of the image.

Continued
190  \textit{SECTION II  RESPIRATORY PROCEDURES}

\textbf{ULTRASOUND BOX 9.1: Thoracentesis—cont’d}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure9.14}
\caption{The inferior tip of the scapula is at the seventh rib. Do not insert the thoracentesis needle below the ninth rib. (From Thomsen T, Setnik G, editors: Procedures consult—emergency medicine module, 2008 Elsevier Inc. All rights reserved.)}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure9.us4}
\caption{Placement of the ultrasound transducer over the posterior of the thorax to identify a pleural effusion.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure9.us5}
\caption{Longitudinal image of the lower part of the thorax and upper part of the abdomen demonstrating a large pleural effusion. The \textit{bright white} (hyperechoic) line arcing over the kidney (arrow) is the diaphragm, which is an important landmark in planning the location of aspiration.}
\end{figure}

Do not insert the needle inferior to the ninth rib

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure9.14}
\caption{Placement of the ultrasound transducer over the posterior of the thorax to identify a pleural effusion.}
\end{figure}

accidental trauma to the intercostal artery located along the inferior margin of each rib. Next use a 22-gauge needle to anesthetize the deeper tissues. Hold the needle perpendicular to the chest wall to avoid inadvertent trauma to the intercostal artery. With each 1 to 2 mm of needle advancement, aspirate and then infiltrate the tissues with 1 to 2 mL of anesthetic. While the aspiration-infiltration process is continued, “walk” the needle above the superior edge of the rib and advance it through the intercostal space until the pleural space is entered (Fig. 9.15). On entering the pleural space, a pop may be felt. Aspirate fluid to ensure that the pleural space has been reached. Once fluid is aspirated, withdraw the needle slightly and infiltrate a small amount of lidocaine to anesthetize the parietal pleura. Finally, grasp the needle just proximal to the skin with the thumb and index finger and withdraw it. This allows measurement of the proper depth of penetration needed during subsequent needle insertion. If no fluid is encountered (dry tap) in the setting of a free-flowing pleural effusion, the needle is too short or the site chosen is too high or too low. If air bubbles are encountered, the chosen sight is likely too high. For both of these scenarios, as long as the patient has no new symptoms to suggest pneumothorax or intraabdominal injury, reevaluate the fluid level using sterile US guidance and reattempt fluid localization.

\textbf{Needle Aspiration Technique for Diagnostic Thoracentesis}

Once you have completed anesthesia and pleural fluid localization, use a 22-gauge needle attached to a 60-mL syringe and the same technique to reenter the pleural space and aspirate 50 mL of fluid. Stabilize the needle at the skin with your thumb and index finger during fluid removal to prevent movement. Once aspiration is complete, instruct the patient to exhale fully, and then withdraw the needle.
**Over-the-Needle Catheter Insertion Technique for Therapeutic Thoracentesis**

As mentioned previously, there are numerous over-the-needle catheters available, each with varying types and degrees of safety features. Here we will discuss performing the procedure with the most basic needle-catheter. Familiarize yourself with the features present on the device at your institution prior to initiating the procedure.

Once you have completed anesthesia and pleural fluid localization, use a scalpel to pierce the skin at the needle insertion site. This will ease entry of the catheter through the skin. Attach the thoracentesis catheter to a 10-mL syringe (Fig. 9.16). Mark the depth of the pleural space that was determined from the anesthetic needle by gently grasping the shaft of the needle-catheter unit with the index finger and thumb of the nondominant hand. This stabilizes the device and controls the advance. Walk the needle-catheter unit over the rib through the anesthetized area and into the pleural space while applying gentle negative pressure with the syringe. When fluid is encountered, angle the needle-catheter unit slightly caudally. While holding the needle steady, twist the catheter to break its seal and advance it into the pleural space. Withdraw the needle and immediately cover the exposed hub of the catheter with your finger to prevent entry of air into the pleural space. Next, attach a three-way stopcock with 60-mL syringe into the catheter hub. The lever should be closed to the unused stopcock port. Aspirate fluid with the syringe for diagnostic studies, then close the lever to the patient. Now connect the high-pressure collection tubing with attached vacuum bottle to the unused stopcock port. Turn the lever closed to the syringe and fluid should begin to drain into the vacuum bottle. The syringe can now be removed. If you do not have a vacuum collection system, you can alternate syringe aspiration with expelling fluid from the syringe, through the tubing, and into a sterile collection bag. Turn the stopcock lever off to the patient when changing collection devices. Stop the procedure when 1500 mL of fluid is removed. The recommended maximum of 1500 mL is to avoid significantly negative pleural pressures, which has been associated with both symptomatic hypovolemia and the potentially fatal complication of reexpansion pulmonary edema.22,23 Larger volumes of fluid may be removed if pleural pressure is monitored, but this is not typically done in the ED setting. Once aspiration is complete, instruct the patient to exhale fully, and then withdraw the needle.

Commercial thoracentesis catheters with safety features can greatly simplify the procedure. Most have a self-sealing valve that obviates the need to cover the catheter hub with your finger as the needle is removed, and a built-in side port that allows fluid collection without a three-way stopcock (see Fig. 9.10).

**PEDIATRIC PATIENTS**

The indications for and contraindications to performing thoracentesis are much the same in children as in adults. Positioning is also similar but will probably require an assistant to help hold the patient and prevent movement. Sedation may be helpful when respiratory distress is minimal. Determine the effusion level by exam and bedside US. Make sure that the needle insertion site is not lower than the ninth rib. The smallest possible needle or needle-catheter device is recommended.

**POSTPROCEDURE EVALUATION**

Traditionally, postprocedure imaging, either x-ray or bedside US, is considered an important part of the process. Patients who are at risk for adhesions, have new symptoms after thoracentesis, require multiple needle passes, have lung disease, or are mechanically ventilated are at higher risk and should have either x-ray or bedside lung US performed after the procedure. Notably, the postprocedure radiograph rarely shows new findings that aid in identifying the cause of the effusion, such as an underlying mass or infiltrate (Fig. 9.17).
Thoracentesis

Position the patient, and then identify and mark the insertion site. The insertion site is best chosen after ultrasound evaluation.

Prepare the area with antiseptic and apply a sterile drape.

Aspirate and stop advancing once pleural fluid is obtained.

Figure 9.16 Steps in the thoracentesis procedure. (From Thomsen T, Setnik G, editors: Procedures consult—emergency medicine module, 2008, Elsevier Inc. All rights reserved.)
PLEURAL FLUID ANALYSIS

Pleural fluid should be analyzed in an organized and thoughtful manner based on clinical suspicion for a disease process. The most cost-effective approach is to perform an initial evaluation to determine whether the fluid is transudative or exudative and obtain other tests only if the fluid is an exudate. Many laboratories are comfortable performing all required tests from the initial 60-mL syringe, so dividing the specimens into individual tubes at the bedside is often unnecessary. Analyze the fluid within 4 hours when possible, but results remain accurate for much longer if the sample is refrigerated at 4°C. Exceptions include using an anticoagulant-containing tube (e.g., lavender top) for differential cell count to prevent clumping of cells, inoculating aerobic and anaerobic blood culture bottles in addition to standard fluid culture to increase the yield of identifiable organisms, and transferring samples for pleural fluid pH immediately to a blood gas syringe, placing it on ice, and analyzing it within 1 hour to prevent falsely high pH measurements. Because of differences in laboratory preference, consult your individual laboratory for its policies on specimen collection and delivery.

Visual Inspection

Identifying a cause based on pleural fluid appearance alone is inaccurate, but certain findings are suggestive. The presence of blood suggests trauma, malignancy, pulmonary infarction, or pneumonia. White or milky fluid suggests the presence of lipids, whereas purulent, malodorous fluid indicates empyema. Pleural effusion containing food particles is highly suggestive of esophageal rupture.

Distinguishing Transudate From Exudate: Light’s Criteria

The first step in evaluation of pleural fluid is categorization of the fluid as exudative or transudative. As discussed earlier, exudates are caused by pleural inflammation, increased pleural membrane permeability, or lymphatic obstruction and therefore contain high levels of LDH and protein. Light and coworkers published criteria for distinguishing transudates from exudates in 1972 based on measurements of serum and pleural fluid protein and LDH. The value for LDH was subsequently changed to accommodate variations in assay conditions. These criteria have since become known as Light’s criteria (Box 9.3). Because Light’s criteria have very high sensitivity but lower specificity, it may misclassify up to 25% of transudates. An exception to using Light’s criteria for differentiating exudates from transudates is in the setting of diuretic-treated CHF, as diuretic use has been shown to increase pleural fluid protein and LDH concentrations, thus making the transudative fluid appear exudative by Light’s criteria. Other criteria for differentiating exudate from transudate should be used in this setting (Box 9.4).

Once the fluid is classified as transudative, it requires no further fluid analysis, and therapy is directed at the underlying cause of the effusion (e.g., CHF, cirrhosis, nephrotic syndrome).

In the presence of an undiagnosed exudative effusion, however, more extensive fluid evaluation is required.

Evaluation of Exudates

For all undiagnosed exudates, pleural fluid should be sent for differential cell count, glucose, Gram stain and culture, and cytologic examination at a minimum (Table 9.1).
fl uid hematocrit if the fluid appears bloody. Clinical suspicion for an underlying disease process should guide additional fluid assessment.

**Differential Cell Count**
Exudates typically have a pleural fluid white blood cell count greater than 1000 cells/mm³, and counts may reach levels higher than 100,000 cells mm³, most commonly with parapneumonic effusions. The differential cell count can be useful in identifying the cause of an exudative effusion. A predominance of neutrophils indicates an acute process, such as pneumonia or pulmonary infarction. A predominance of lymphocytes is consistent with a more chronic pleural process, such as malignancy or TB. Eosinophil counts greater than 10% often have no clear etiology but have traditionally been associated with blood or air in the pleural space.

**Glucose**
The concentration of glucose in exudates is extremely variable and, in general, does not correlate with any specific disease process. Routine measurement of pleural fluid glucose for exudative effusions is recommended because a low glucose concentration (< 60 mg/dL) narrows the differential diagnosis to complicated parapneumonic effusion, malignancy, TB, rheumatoid effusion, and the uncommonly seen paragonimiasis or Churg-Strauss syndrome. Keep in mind, however, that these diagnoses are not excluded by a high or normal pleural fluid glucose.

**BOX 9.4 Separating Exudate From Transudate in the Diuretic-Treated CHF Patient**
The effusion is likely a transudate if one of the following is present:
- Serum protein minus pleural protein > 3.1 g/dL
- Serum albumin minus pleural albumin > 1.2 g/dL
- Pleural fluid N-terminal pro-brain natriuretic peptide > 1500 pg/mL

CHF, Congestive heart failure.

**TABLE 9.1 Exudates: Evaluation of an Undiagnosed Exudative Effusion**

<table>
<thead>
<tr>
<th>PLEURAL FLUID ASSAY</th>
<th>RESULT</th>
<th>PROBABLE DIAGNOSIS (DIFFERENTIAL DIAGNOSIS)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Standard Testing: Perform on All Exudative Effusions</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Differential Cell Count</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WBC count</td>
<td>&gt; 10,000 cells/mm³</td>
<td>Parapneumonic effusion (pancreatitis, PE, collagen vascular disease)</td>
</tr>
<tr>
<td>Neutrophils</td>
<td>&gt; 50%</td>
<td>Acute pleural process: infection, pulmonary infarction</td>
</tr>
<tr>
<td>Lymphocytes</td>
<td>&gt; 50%</td>
<td>Chronic pleural process: malignancy, chronic TB</td>
</tr>
<tr>
<td>Eosinophils</td>
<td>&gt; 10%</td>
<td>Air or blood in the pleural space (numerous other causes)</td>
</tr>
<tr>
<td>Cytology</td>
<td>Abnormal cells</td>
<td>Malignancy</td>
</tr>
<tr>
<td>Glucose</td>
<td>&lt; 60 mg/dL</td>
<td>Parapneumonic effusion, malignancy, TB, collagen vascular disease, esophageal rupture</td>
</tr>
<tr>
<td>Gram stain and Culture</td>
<td>Presence of organism</td>
<td>Pleural space infection</td>
</tr>
<tr>
<td>Pleural fluid hematocrit</td>
<td>&gt; 50% of peripheral hematocrit</td>
<td>Hemothorax (trauma, malignancy, PE)</td>
</tr>
<tr>
<td><strong>Selective Testing: Order if High Clinical Suspicion for Diagnosis</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adenosine deaminase</td>
<td>&gt; 40 IU/L</td>
<td>TB (lymphoma, empyema)</td>
</tr>
<tr>
<td>Amylase</td>
<td>Pleural fluid amylase &gt; serum amylase</td>
<td>Pancreatitis, esophageal rupture, malignancy</td>
</tr>
<tr>
<td>Cholesterol</td>
<td>&gt; 45 mg/dL</td>
<td>Exudative effusions (inaccurate alone)</td>
</tr>
<tr>
<td>Creatinine (with serum measurement)</td>
<td>Pleural fluid creatinine &gt; serum creatinine</td>
<td>Urinothorax</td>
</tr>
<tr>
<td>N-terminal pro-brain natriuretic peptide</td>
<td>≥ 1500 pg/mL</td>
<td>Congestive heart failure</td>
</tr>
<tr>
<td>pH</td>
<td>&lt; 7.2</td>
<td>Complicated parapneumonic effusion (collagen vascular disease, esophageal rupture, malignancy, TB)</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>&gt; 110 mg/dL (&gt; 50 if fasting or malnourished)</td>
<td>Chylothorax (intrathoracic TPN infusion)</td>
</tr>
</tbody>
</table>

PE, Pulmonary embolism; TB, tuberculosis; TPN, total parenteral nutrition; WBC, white blood cell.
Culture
If bacterial infection is a consideration, culture the pleural fluid by both bedside inoculation of blood culture bottles and standard laboratory culture. The addition of blood culture bottles increases the overall yield of identifiable pathogens in parapneumonic effusions by 20%, from a baseline of 40% yield with standard laboratory culture alone. As little as 2 mL of fluid can be used in each bottle without affecting the results.

Pleural Fluid Hematocrit
In general, the presence or absence of red blood cells is not useful in determining the cause of effusion because it takes only a small amount of blood to impart a blood-tinged appearance. A grossly bloody effusion is suggestive of trauma, malignancy, or pulmonary infarction. A pleural fluid hematocrit of greater than 50% of the peripheral hematocrit is indicative of hemothorax and often requires tube thoracostomy.

Cytology
Perform cytologic analysis on all undiagnosed effusions or when malignancy is suspected. The sensitivity for diagnosing pleural malignancy hovers at approximately 55%, regardless of whether small or large (> 50 mL) volumes are analyzed.

Parapneumonic Effusion
Patients with suspected parapneumonic effusions warrant rapid evaluation and outcome risk assessment based on pleural anatomy, pleural fluid bacteriology, and pleural fluid chemistry. All patients with parapneumonic effusions require at least diagnostic thoracentesis with the goal of identifying those with complicated parapneumonic effusions. Indications for tube thoracostomy or other surgical procedure include large or loculated effusions, pleural thickening on CT, aspiration of frank pus (empyema), pleural fluid pH lower than 7.20, pleural fluid glucose concentrations lower than 60 mg/dL, and positive Gram stain or culture (Box 9.5).

COMPLICATIONS

Pneumothorax
Pneumothorax is the most frequently reported complication of thoracentesis with an overall rate of approximately 6%. It can develop in even the most pristine and seemingly uncomplicated procedure and does not necessarily denote poor technique. Fortunately, thoracostomy tubes are required in only one third of cases. Procedure-related factors that appear to contribute to pneumothorax include an inexperienced operator, larger needle size, multiple needle attempts, and large-volume fluid removal. A recent large observational study showed an increased risk of pneumothorax in patients who are underweight. The risk for iatrogenic pneumothorax is significantly lower in all patient groups when thoracentesis is performed using real-time US guidance.

Pneumothorax Ex Vacuo
Pneumothorax ex vacuo is a form of pneumothorax that occurs when the lung is unable to reexpand during therapeutic thoracentesis due to the presence of an endobronchial lesion or other structural pathology. Negative pleural pressures created in this scenario cause a transient passage of air from the lung to the pleural space. It is a likely explanation for pneumothorax occurring after a seemingly uncomplicated therapeutic thoracentesis.

Cough
Cough is typically considered a minor complication that results only in patient discomfort, though it can be a harbinger of a postprocedure pneumothorax. Terminate the procedure if persistent coughing occurs.

Uncommon Serious Complications
Less common, but serious complications include hemothorax, splenic rupture, abdominal hemorrhage, unilateral pulmonary edema, infection, air embolism, and fragmentation of the catheter in the pleural space.

Hemothorax is typically caused by intercostal artery laceration and should be suspected with rapid reaccumulation of pleural fluid or by a change in the patient's vital signs during or after the procedure. Other causes include laceration of the lung, diaphragmatic, or internal mammary vessels. Hemothorax requires immediate surgical consultation, thoracostomy tube placement, and close monitoring of hemodynamics with resuscitation as needed.

Puncture of the spleen or liver through the diaphragm may result in a localized organ hematoma or, more seriously, hemoperitoneum. Clinically, this is suspected when the needle pass does not yield fluid (dry tap) and is followed by the patient's complaint of abdominal pain. If this diagnosis is suspected, appropriate resuscitation is the initial treatment, followed by abdominal CT imaging with contrast, when possible. If the patient is hemodynamically unstable, obtain bedside FAST (focused assessment with sonography) US and immediate surgical consultation.

Reexpansion pulmonary edema is a complication associated with rapid reexpansion of the lung and negative pleural pressures during thoracentesis. Symptoms include dyspnea, tachypnea, tachycardia, cough, and frothy sputum. It is believed that this problem can be avoided by discontinuing the procedure when pleural pressure is greater than –20 mm Hg, which is unlikely to occur until after 1500 mL of fluid has been acutely removed. Because pleural pressure is rarely measured in the ED, 1500 mL of fluid is considered to be the maximum amount that can be safely withdrawn at one time.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Tube thoracostomy is a procedure used to evacuate an abnormal accumulation of fluid or air from the pleural space and can be performed on an elective, urgent, or emergent basis. Air or fluid can accumulate in the pleural space as a result of spontaneous or traumatic pneumothorax, pleural fluid accumulation of blood, malignancy, infection (empyema), or lymph (chylothorax). The first modern methods to evacuate the contents of the pleural space were developed in the 19th century, but these techniques did not become widespread until 1918, when they were used to treat post-influenza empyema. Military experience demonstrated that thoracic drainage combined with antiseptics and antibiotics reduced mortality related to thoracic trauma from 62.5% during the Civil War, to 24.6% in World War I, and to 12% in World War II.¹

**PATHOPHYSIOLOGY**

The lung is surrounded by two layers, the parietal pleura, which lines the interior of the chest wall, and the visceral pleura, which covers the lungs. They are separated by a potential space that usually contains a thin layer of fluid that lubricates the pleural space. A small negative pressure within the pleural space helps keep the lung inflated and the two layers closely apposed. With inspiration, chest expansion increases negative intrathoracic pressure and leads to inflation of the lung from an influx of air. If the pleural space is disrupted, air, blood, or other fluid can accumulate between the two layers of the pleura. The result is that the pressure gradient associated with normal respiration is compromised and can lead to “collapse” of the lung. As the amount of fluid or air increases, respiratory function worsens leading to symptoms of dyspnea, often with pleuritic chest pain and anxiety. The degree of respiratory compromise depends on the volume of fluid or air in the pleural space, the patient’s age, baseline pulmonary status, and the integrity of the chest wall. In severe cases, a one-way valve type of mechanism can occur within the respiratory apparatus, leading to tension pneumothorax physiology. The hallmark of tension pneumothorax is a progressive accumulation of air in the pleural space with each breath that will result in severe respiratory dysfunction, cardiovascular compromise, and ultimately obstructive shock if untreated.

**Pneumothorax**

A pneumothorax is caused by the presence of air in the pleural space and loss of the normal negative pressure (Fig. 10.1). Air can enter the pleural space from outside of the chest wall as a result of a penetrating injury. Air can also enter internally from a ruptured lung bleb or damaged trachea. Iatrogenic injuries can occur from needle procedures such as subclavian venous cannulation, transthoracic biopsy, thoracentesis, positive pressure ventilation (PPV), or cardiopulmonary resuscitation (Fig. 10.2).

Pneumothoraces are commonly divided into “open” and “closed.” An open pneumothorax indicates that the skin and

---

**Tube Thoracostomy**

**Indications**
- Spontaneous and traumatic pneumothorax
- Hemothorax
- Empyema
- Patients with penetrating chest trauma undergoing positive pressure ventilation or long-distance transport

**Contraindications**
- **Absolute**
  - None
- **Relative**
  - Presence of multiple pleural adhesions
  - Presence of emphysematous blebs
  - Coagulopathy

**Complications**
- Infection
- Laceration of an intercostal vessel
- Pulmonary injury
- Intra-abdominal or solid organ tube placement
- Failure of reexpansion of pneumothorax
- Reexpansion pulmonary edema

**Equipment**
- Sterile drapes
- Local anesthetic
- Antiseptic
- Gauze
- Needle driver
- Forceps
- Scalpel with No. 10 blade
- Mayo scissors
- Large Kelly clamps
- Chest tube
- Occlusive dressing
- Silk suture
- Tape

---

Review Box 10.1  Tube thoracostomy: indications, contraindications, complications, and equipment.
Figure 10.1 Pneumothorax. A, Anteroposterior chest radiograph of a right-sided, seemingly small, simple pneumothorax (PTX). Note the absence of peripheral lung markings on the right side and the distinct line indicating the edge of the collapsed lung (arrow). Although this appears to be a small PTX, it produced significant dyspnea in this patient with chronic obstructive pulmonary disease and therefore required a chest tube. B, Computed tomography scan showing the extent of the collapse. Adhesions kept part of the lung expanded.

Figure 10.2 A, After successful cardiopulmonary resuscitation and intubation, this patient began to deteriorate, with a precipitous drop in blood pressure and decreasing oxygen saturation. It was believed that the cause of the initial cardiac arrest was returning. Marked subcutaneous air was noted in the scrotum and abdominal wall, but little air was noted in the chest wall tissue. The subcutaneous air from a pneumothorax and positive pressure ventilations created by the resuscitation had curiously tracked via abdominal wall tissue planes, a distinctly unusual place for air to accumulate. Usually subcutaneous air is felt around the chest tube insertion site, neck, and chest wall. B, A chest tube (arrow) quickly reversed the decompensation from the pneumothorax. C, This patient suffered respiratory arrest from heroin injected into a neck vein (note the extensive scar). After bag-mask resuscitation the respiratory depression returned and the blood pressure dropped precipitously. He was unresponsive to naloxone and was very difficult to ventilate. He had a small pneumothorax from a nick in the lung from the neck injection. The positive pressure ventilations produced a tension pneumothorax.
Spontaneous (Closed) Pneumothorax
Spontaneous pneumothorax is caused by the rupture of a subpleural lung bleb with little or no trauma and can be categorized as either primary or secondary based on the presence of underlying lung disease. Primary spontaneous pneumothorax occurs in a patient without overt lung disease. The typical patient with spontaneous pneumothorax is a tall, thin, 20- to 40-year-old male smoker (Fig. 10.3). Secondary spontaneous pneumothoraces occur in patients with underlying lung or pleural disease, including emphysema, chronic bronchitis, asthma, Marfan’s syndrome, infection, and neoplasm. The morbidity, mortality, and long-term complications associated with pneumothorax increase in patients with underlying lung disease. Whereas a primary pneumothorax may be managed by selective observation or might simply be aspirated, a secondary pneumothorax often requires a more aggressive management approach.

The sudden onset of pleuritic chest pain and dyspnea with exertion or at rest is the most common finding. More subtle manifestations may also occur with little or no pain and only mild dyspnea on excursion that the patient may ignore for days. A person with a small spontaneous pneumothorax may never seek medical attention, and the process will resolve without treatment. The signs and symptoms do not always correlate well with the size or cause of the collapsed lung. Tube thoracostomy is the most common treatment, but new trials suggest that conservative management or aspiration of first-time primary pneumothoraces results in similar outcomes as traditional tube thoracostomy, though with fewer complications, shorter hospital stay, and lower cost. Depending on the size of the pneumothorax, conservative management and aspiration are both reasonable initial interventions in clinically stable patients. However, to date no high-quality clinical trials have definitely demonstrated that aspiration is a superior treatment methodology.24 Rarely, spontaneous pneumothorax may be bilateral. In addition, in rare cases, a spontaneous pneumothorax may progress to tension pneumothorax as described previously, which can result in a potentially life-threatening condition.

Traumatic Closed Pneumothorax
Closed pneumothorax usually occurs from a rib fracture that penetrates the lung, but can also occur when an alveolus or bleb ruptures after blunt trauma. The air leak from a closed pneumothorax is generally self-limited but can, in rare cases, progress to a tension pneumothorax.

Traumatic Open Pneumothorax
An open pneumothorax occurs when the chest wall is penetrated and the negative intrapleural pressure is lost. Each breath can increase intrapleural pressure, especially if the diameter of the chest wound is greater than the diameter of the trachea. With each respiratory attempt, air moves preferentially through the chest wall opening rather than down the trachea and thus prevents meaningful ventilation of the involved lung.

Tension Pneumothorax
An open pneumothorax can occasionally manifest as a tension pneumothorax, which is a life-threatening condition that requires immediate intervention. A tension pneumothorax occurs when an injury creates a one-way “flap valve” mechanism that allows air into the pleural space with inspiration but then closes with expiration and traps the air (Fig. 10.4). The progressive accumulation of air in the pleural space leads to ipsilateral complete lung collapse and then impingement on the mediastinum with a shift of the heart toward the uninvolved side. This restricts ventricular filling and subsequently decreases cardiac output. This severe disruption in both respiratory and cardiac function can lead to hypotension and reduced ventilation.
Hemothorax

Hemothorax is an accumulation of blood in the pleural space as a result of injury to the heart, great vessels, or vessels of the lungs, mediastinum, or chest wall. Bleeding from the lung parenchyma is usually low pressure, self-limited, and ceases when a chest tube is inserted. However, bleeding from an intercostal artery, a pulmonary artery, or the internal mammary artery can be profuse and often requires surgical intervention.

Empyemas and Effusions

An empyema is an accumulation of pus in the pleural space, usually from a parapneumonic infectious effusion (Fig. 10.5). An empyema can also be caused by violation of the thoracic

Figure 10.4 Traumatic tension pneumothorax. A and B, Pathophysiology of a tension pneumothorax. During inspiration, air enters the pleural space through a one-way valve either from the outside or from the lung itself. On expiration, the injury/valve closes and traps increasing amounts of air in the pleural space. Eventually, the mediastinum shifts and cardiac filling and ultimately cardiac output are compromised. C, This elderly patient sustained a tension hemopneumothorax after slipping and falling on ice. The left hemithorax is very dark (radiolucent) because of total collapse of the left lung (large white arrow). Note the dramatic shift of the mediastinum to the right, indicative of tension. Multiple posterior rib fractures are present but are difficult to appreciate on this film (small white arrows). The air-fluid level (black arrow) indicates the presence of air in the pleural cavity in addition to fluid (blood). D, A computed tomography scan of the same patient again demonstrates the findings seen on the conventional radiograph.
To hypotension or severe dyspnea, especially in those patients with tension pneumothorax. With a spontaneous pneumothorax, 95% of patients complain of the sudden onset of sharp, pleuritic chest pain, shoulder pain, or both. Sixty percent of patients experience dyspnea, and 12% have a mild cough. Dyspnea and anxiety are more common in older patients.

Tension pneumothorax (TP) must be considered in any patient with sudden or severe respiratory or cardiac deterioration and in intubated patients who become difficult to ventilate. Signs of TP include increased airway pressure, hypotension, or elevated central venous and pulmonary artery pressure. Severe dyspnea, restlessness, agitation, and a feeling of impending doom can develop rapidly in conscious patients with tension pneumothorax. Initially these patients may be tachycardic and tachypneic but can quickly become hypotensive.

The symptoms of hemothorax can be similar to those of pneumothorax but may be accompanied by hypotension as blood accumulates in the pleural space. The onset of symptoms with effusions is usually much more gradual, with increasing shortness of breath and dyspnea on exertion occurring over a period of days to weeks.

**DIAGNOSIS**

**Symptoms**

The symptoms of patients with abnormal collections in the pleural space range widely depending on the size of the pneumothorax, the rapidity of accumulation, the age of the patient, and the presence of comorbidities, especially underlying lung disease. Specific symptoms range from mild dyspnea with exertion and pleuritic chest pain with small disruptions to hypotension or severe dyspnea, especially in those patients with tension pneumothorax. With a spontaneous pneumothorax, 95% of patients complain of the sudden onset of sharp, pleuritic chest pain, shoulder pain, or both. Sixty percent of patients experience dyspnea, and 12% have a mild cough. Dyspnea and anxiety are more common in older patients.

Tension pneumothorax (TP) must be considered in any patient with sudden or severe respiratory or cardiac deterioration and in intubated patients who become difficult to ventilate. Signs of TP include increased airway pressure, hypotension, or elevated central venous and pulmonary artery pressure. Severe dyspnea, restlessness, agitation, and a feeling of impending doom can develop rapidly in conscious patients with tension pneumothorax. Initially these patients may be tachycardic and tachypneic but can quickly become hypotensive.

The symptoms of hemothorax can be similar to those of pneumothorax but may be accompanied by hypotension as blood accumulates in the pleural space. The onset of symptoms with effusions is usually much more gradual, with increasing shortness of breath and dyspnea on exertion occurring over a period of days to weeks.

**Physical Examination**

**Unstable Patients**

During the initial phase of resuscitation (airway, breathing, circulation, disability), consider the diagnosis of pneumothorax in patients who are tachycardic, hypotensive, and dyspneic. These symptoms are not specific as similar ones do occur with pulmonary embolism, pericardial tamponade, large pleural effusion, and severe, multilobar pneumonia. Serial examination is critical as the diagnosis of tension pneumothorax by physical examination can be very subtle. Use the phrase "look, listen, and feel" as your guideline. Observe the chest wall, which may reveal asymmetric chest expansion, and the neck and forehead.
veins, which may be distended, even if the patient is hypotensive. The trachea may be deviated away from the side of the pneumothorax. When percussing the chest wall, hyperresonance on the affected side and subcutaneous emphysema may be present. Auscultation may demonstrate diminished breath sounds on the injured side. In one prospective study, the sensitivity, specificity, and diagnostic accuracy of auscultation for hemothorax and pneumothorax were 84%, 97%, and 89%, respectively. A false-negative auscultation is more likely than a false-positive one. Pulsus paradoxus may be evident. For intubated patients, an early sign of tension pneumothorax is difficulty ventilating because of increased airway pressure. Additionally, poor compliance may be noted when providing respiratory support by bagging.

In injured patients with apnea, hypotension, or cardiopulmonary arrest, diagnose and treat a tension pneumothorax by immediate needle or catheter decompression thoracentesis. In patients who may have TP, do not take the time to obtain and review a radiograph because delay can lead to increased morbidity and mortality in patients with this emergency condition. X-rays cause a delay in therapeutic intervention. Confirm the diagnosis of tension pneumothorax after needle decompression by rapid improvement in vital signs and a rush of air through the needle.

**Stable Patients**

In more stable patients (and those with smaller accumulations) the findings on physical examination are less sensitive, and a chest radiograph, ultrasound, or even a computed tomography (CT) scan is usually necessary to make a definitive diagnosis. Physical findings may include unilaterally decreased breath sounds, tachypnea, tachycardia, decreased tactile fremitus, increased resonance with percussion, or subcutaneous emphysema. Alternatively, the examination may reveal little to no abnormalities with a small pneumothorax. Patients with a pneumothorax involving less than 20% of the hemithorax will often have completely normal findings on chest examination, including equal breath sounds (Fig. 10.6). Pleural fluid collections are difficult to detect by physical examination, particularly with less than 500 mL of fluid in the pleural space. Breath sounds may be decreased and percussion of the bases may be dull.

Parapneumonic empyemas are often accompanied by fever, cough, chest pain, dyspnea, and purulent sputum (see Fig. 10.5). Physical examination may reveal diminished breath sounds, dullness on percussion, egophony, and diminished tactile fremitus on the involved side. Fever will often develop in patients with an indwelling chest tube and empyema. The pleural fluid drainage may be copious and purulent, and respiratory symptoms may worsen.

**Radiography**

**Plain Radiographs**

In stable patients, a chest radiograph is an essential tool for diagnosing a pneumothorax. In unstable patients with a potential tension pneumothorax, the diagnosis should be made clinically, but in rare cases a portable radiograph may be obtained in the resuscitation room if the patient is being carefully monitored by a clinician. As will be discussed later in this chapter, bedside ultrasound may prove advantageous as a diagnostic modality in these peri-stable patients. The best plain radiographs for detecting hemothorax or pneumothorax are traditional upright inspiratory posteroanterior and lateral chest radiographs.

Diagnostic sensitivity is not increased with an expiratory upright chest radiograph. Upright is preferable to a supine chest radiograph, particularly for a hemothorax, because even with large amounts of blood there may only be slight differences in the density of the lung fields as the blood may layer out evenly. With an upright chest radiograph, 300 to 500 mL of fluid is needed to cause blunting of the costophrenic angle (Fig. 10.7). When CT is not available, other useful views...
include a bilateral decubitus chest radiograph, with the pneumothorax expected to be seen on the side away from the table (upward) as gravity pulls the affected lung down. A pleural effusion or hemothorax, however, may be better seen on the side toward the table (downward).

On a chest radiograph, the partially collapsed lung of a pneumothorax appears as a visceral pleural line with no pulmonary markings beyond it (see Figs. 10.1, 10.3, 10.6, and 10.7). It is easy to initially mistake large blebs for a pneumothorax or to identify the scapular border, skin folds, or indwelling lines as a pneumothorax, but a CT scan quickly resolves the issue (Fig. 10.8). Other radiographic findings include hyperlucency of the affected hemithorax, a double diaphragm contour, increased visibility of the inferior cardiac border, better visualization of pericardial fat at the cardiac apex, and possibly a depressed diaphragm. If subcutaneous air is noted on the chest radiograph of a patient with blunt chest trauma, assume that the air came from an injured lung and that a pneumothorax exists until proven otherwise.

It is difficult to accurately predict the size of a pneumothorax on plain radiographs. Greater accuracy in predicting size can best be accomplished with a CT scan. With a tension pneumothorax, the chest radiograph reveals lung collapse, a depressed hemidiaphragm on the affected side, and a shift of the mediastinum and trachea to the opposite side (see Fig. 10.4). Keep in mind, with a bilateral pneumothorax, a mediastinal shift may not be seen.

Thoracic CT
The gold standard for diagnosis is a thoracic CT scan, which can even detect a pneumothorax that is not easily visible on a plain radiograph. CT scans of the chest are more sensitive than plain radiographs in detecting hemothorax and pneumothorax. They are also more accurate for estimating the size and other characteristics of a pneumothorax (see Figs. 10.1, 10.4, and 10.8). CT scans are not routine for the diagnosis of a pneumothorax, but are more useful for hemothoraces and other fluid collections. They also offer invaluable information on the cause of such abnormalities. A CT scan may be useful when the diagnosis is unclear or when looking for small amounts of pleural fluid. CT scans are particularly helpful in determining whether an empyema is loculated or draining successfully. Approximately 10% of trauma patients with normal findings on a chest radiograph will demonstrate a small hemothorax or pneumothorax. The clinical significance of a small, previously undetected occult injury is probably not great, and it has been suggested that a small pneumothorax seen only on CT may be left untreated and simply observed in otherwise stable patients. Some patients with a pneumothorax seen only on CT may also safely undergo PPV without placement of a chest tube.

Ultrasound
Ultrasound is useful in diagnosing both hemothoraces and pneumothoraces, and its use is reviewed in the Ultrasound Box.

Thoracic ultrasound is a useful tool at the bedside to rapidly assess the lung fields. This includes the ability to assess the pleural space, diagnose pleural and lung parenchymal abnormalities and guide pleural-based procedures, including thoracentesis with or without catheter placement, and evaluation of placement for tube thoracostomy. Thoracic ultrasound has several advantages over standard chest radiographic imaging, including absence of radiation, portability, real-time imaging, and the ability to perform dynamic imaging. Sensitivity of lung ultrasound in the detection of pneumothorax is higher than conventional chest radiography. Thoracic ultrasound is considered superior in the detection and characterization of pleural effusion, and provides an advantage for the guidance of bedside pleural interventions. Additionally, bedside thoracic ultrasound is helpful in differentiating between consolidation, atelectasis, and effusion, all of which can appear as an opacity
To evaluate for pneumothorax, a high-frequency transducer should be used to ensure a high degree of resolution. In a supine patient the transducer should be placed on the anterior chest wall in the midclavicular line at approximately the second to third intercostal space (Fig. 10.US1). The depth of the image should be adjusted until the ribs are seen as brightly echogenic (white) arcs with acoustic shadows behind them. The pleural line can be found just deep to the ribs and is represented as a horizontal, echogenic line (Fig. 10.US2). In a normal lung, the visceral and parietal pleural layers are directly opposed to one another (save for a thin layer of pleural fluid). When the patient breathes in and out, the layers “slide” past each other to allow the lungs to expand. When this is viewed with ultrasound, the pleural line can be seen to slide back and forth with patient respiration. This is referred to as the “slide sign.” In cases in which this interface is disrupted (such as by a pneumothorax), the sliding is lost. When these patients are evaluated with ultrasound, the pleural line will be seen as a static echogenic line that does not change with respiration. Although the slide sign carries significant sensitivity for ruling out a pneumothorax, other secondary confirmatory findings should be sought as well.

Comet-tail artifacts may be seen in a normal lung. These are hyperechoic vertical lines that extend deep to the pleural interface (Fig. 10.US3). Typically, they will be seen in small numbers in a normal lung. Patients in whom a pneumothorax is present are noted to lack comet tails because this artifact arises from the normal pleural interface, which is disrupted. The presence of comet tails may be used by the sonographer to further rule out a pneumothorax.2,3

Motion mode (M-mode) may also be used to further evaluate for the presence of a pneumothorax. M-mode is used to evaluate objects in motion and plots a linear representation of motion on screen. Objects that move toward the surface (or toward the transducer) are
or “white out” on chest radiograph. Ultrasound can also be used to evaluate chest wall pathology.

Lower-frequency probes with smaller footprints, such as the phased-array probe, should be used for the evaluation of pulmonary and pleural pathology as the probe can allow for scanning between rib interspaces. For chest wall structures, changing from the 3.5 to 5 MHz phased-array probe to the linear probe with a frequency of 7.5 to 10 MHz will provide better visualization. Additionally, the linear high-frequency probe is better suited for evaluation of the pleural line, which is discussed later.

The normal pleura appears as an echogenic line between the chest wall and the air-filled lung. Lung sliding, which represents the visceral pleura moving against the parietal pleura, is a key finding in a lung without a pneumothorax. When air separates the two pleural layers, as in a pneumothorax, the movement disappears.

There are two main types of lung artifacts caused by air-tissue interfaces that are commonly observed, A-lines and B-lines. Horizontal artifact, or A-lines, are echogenic lines between rib shadows that are observed in normal lungs. B-lines arise at the border between aerated and compressed lungs. They are often described as comet-tail and are characteristic in that they spread to the edge of the screen without fading, move synchronously with the lung during respiration, and tend to erase A-lines. Multiple B-lines are often termed lung rockets and are pathologic.

Thoracic ultrasound can be an integral part in the evaluation of pneumothorax in a variety of situations. Examples include patients with chest trauma, evaluation for postprocedural iatrogenic pneumothorax, patients presenting with undifferentiated shortness of breath, hypoxia or chest pain, and following chest tube placement to assess resolution of pneumothorax.

When evaluating for pneumothorax the patient should ideally be in the supine position for visualization of the lung apices. The probe indicator should be pointed cephalad at approximately the third-to-fourth intercostal space between the parasternal and midclavicular line. The probe is moved progressively toward the lateral chest, checking for lung sliding at different locations. Air will first move to the most superior part of the hemithorax, which in the supine patient, corresponds to the lung apices. The evaluation of pneumothorax should focus on observing for the presence of four sonographic artifact signs: lung sliding, including by M-mode sonography, B-lines, the lung pulse, and the lung point sign. The lung point sign is the most specific sonographic sign for pneumothorax and occurs when the lung intermittently comes into contact with the chest wall during inspiration. This represents the edge of the pneumothorax. The presence of a lung point is 100% specific for ruling in pneumothorax. Lung pulse describes the vertical movement of the pleural line synchronous to the cardiac rhythm and is caused by transmission of heartbeats through the lung.

Furthermore, thoracic ultrasound can be advantageous when evaluating the need for a pleural procedure and during the performance of this procedure. Blind insertion of a chest tube can result in significant complications in a patient with pleural adhesions from prior infections, pleurodesis, or pulmonary surgery. Guidance by ultrasound can help mitigate the risk of complications.

Ultrasound-guided thoracentesis, with or without subsequent placement of the drainage catheter, can be performed either by static or dynamic technique. Placement of tube thoracostomy is typically performed by static technique (or simply by landmark) after the diagnosis has been made and evaluation of the previously mentioned potential complications completed. Static ultrasound guidance identifies the intercostal spaces, pleura, and effusion, as well as the best angle of approach.

---

**ULTRASOUND BOX 10.1: Recognizing Pneumothorax—cont’d**

![Figure 10.US5](image_url) Bar code sign consistent with a pneumothorax. This M-mode image shows lack of movement of the pleura. Unlike the previous image, the solid lines can be seen to continue past the point of the pleural line. Because the presence of the pneumothorax causes the pleura to appear stationary, no movement is seen in this image.
The skin is then marked on the patient, the probe is removed, and the procedure carried out the same way as during landmark thoracentesis or tube thoracostomy. The examiner may also choose to observe diaphragmatic excursion through several cycles to avoid selecting a location where the instrumentation occurs too close to the diaphragm, potentially resulting in laceration.

Thoracic ultrasound is considered superior to chest CT in visualizing stranding and septations in complex pleural effusions. The ability to detect such findings may prove useful as they can predict difficulty with tube drainage.22 Additionally, distinguishing large bullae from a pneumothorax can be very challenging, as plain radiograph of the chest may be inadequate to make the diagnosis. Making the correct diagnosis is critical as the management approach for these two entities varies significantly. Whereas CT of the chest remains the diagnostic modality of choice for characterization of large bullae, there are case reports in the literature where thoracic ultrasound was used to correctly differentiate bullae from pneumothorax. Specifically, the comet-tailing phenomenon of the movement of the lung tissue against the pleura during respiration was noted to be present in bullous disease but absent in pneumothorax.21

INDICATIONS FOR TUBE THORACOSTOMY

Pneumothorax

Tube thoracostomy is by far the most common treatment for all types of pneumothoraces, but controversy exists over the treatment of small traumatic and spontaneous primary pneumothoraces. Treatment should address several basic issues: (1) removal of air, (2) preventing further air accumulation, (3) healing the injury that caused the initial accumulation, (4) promoting lung reexpansion, and (5) preventing or limiting recurrences.22 The American College of Chest Physicians has developed useful guidelines for the management of primary and secondary spontaneous pneumothoraces (Box 10.1).21

Chest tube placement is probably not necessary in healthy patients with small primary spontaneous or isolated small traumatic pneumothoraces in the absence of respiratory compromise or concomitant injuries, or when PPV will not be required. With no intervention, a small pneumothorax will resolve over a period of days to weeks. Supplemental oxygen will speed the process of lung reexpansion by increasing the rate of pleural air absorption. No controlled studies have been conducted to compare the various interventions for primary spontaneous pneumothorax, and there is wide variation in treatment approaches. In particular, no significant difference has been found between simple aspiration and tube drainage with regard to the immediate success rate, early failure rate, duration of hospitalization, and 1-year success and pleurodesis.24 Needle aspiration is associated with reduced analgesia requirements, lower pain scores, and a lower percentage of patients being hospitalized. Studies have shown excellent short- and long-term outcomes after video-assisted thoracoscopic treatment (VATS) of primary and secondary spontaneous pneumothorax, and rates of recurrence are lower than after other treatment modalities. However, VATS continues to have a higher complication rate than simple tube thoracostomy, with 4.6% of patients with spontaneous pneumothorax experiencing serious side effects as a result of the procedure.25,26

In patients with a simple pneumothorax, prolonged suction is rarely required, and the tube can simply be attached to a Heimlich valve or underwater seal.27–29 Treatment is different for patients with a secondary spontaneous pneumothorax because the presence of underlying disease leads to more serious compromise and more frequent complications. In patients with chronic obstructive pulmonary disease, malignancy, cystic fibrosis, pneumonia, and tuberculosis, a chest tube usually cannot be avoided.

The underlying lung pathology of a patient with a spontaneous pneumothorax is best initially evaluated with CT. Frequently, this is followed by diagnostic or therapeutic visual inspection of the lung and pleural space by fiberoptic thoracoscopy. Extensive evaluation is not, however, usually recommended for the first episode of a small primary pneumothorax. When patients have recurrent spontaneous pneumothoraces, further evaluation (CT, thoracoscopy) and evaluation for surgical treatment are indicated. Patients who have had one spontaneous pneumothorax have a 30% to 50% chance of recurrence within 2 years, and after the second pneumothorax there is a 50% to 80% chance of developing a third. Surgery may be recommended for a first pneumothorax in the following situations: life-threatening tension pneumothorax, massive air leaks with incomplete reexpansion, an air leak persisting 4 days after a second tube has been placed, associated hemothorax with complications, identifiable bullous disease, or failure of easy reexpansion in patients with cystic fibrosis.

In patients with traumatic causes, the urgency and type of treatment depend primarily on the stability of the patient; a hypotensive patient with a tension pneumothorax requires immediate decompression with a chest tube or needle thoracostomy, whereas a patient with normal vital signs and a small pneumothorax may be observed initially. Emergency needle thoracostomy is only a temporary solution for a compromised patient with a tension pneumothorax. Once done, needle thoracostomy necessitates an ipsilateral tube thoracostomy. Other factors that modify treatment include the patient’s age, the size of the pneumothorax, whether bilateral pneumothoraces are present, and whether the current episode represents a recurrence. A chest tube is usually indicated for a pneumothorax created by needle decompression.

Because of the risk for tension pneumothorax, a chest tube should be considered in all patients with a penetrating chest injury if PPV will be used or if they will be transported a long distance for definitive care. However, CT scans of trauma victims have demonstrated that many patients with small pneumothoraces that would have escaped detection with standard radiographs have safely undergone PPV without clinically evident pneumothorax developing. Close observation for signs of tension pneumothorax is necessary for patients in whom a chest tube is not placed and PPV is used.

Hemothorax

Tube thoracostomy is the treatment for hemothorax, but it is also used to monitor the amount and rapidity of blood output, which determines the need for additional interventions, including video-assisted thoracoscopy or open thoracotomy. Approximately three fourths of patients with a traumatic hemothorax can be managed by tube thoracostomy and volume replacement alone. The remaining patients will require immediate or delayed elective thoracotomy. The indications for surgery after an acute hemothorax vary somewhat between authors but are summarized
in Box 10.2. Early institution of blood replacement is recommended for patients with massive hemothoraces (>2000 mL) because they are often associated with continued hemorrhage. Autotransfusion of the shed blood is desirable whenever possible. For patients with active blood loss but stable hemodynamics, VATS can be used to locate and stop the bleeding, evacuate blood clots, and break down adhesions.

**Empyema**

Treatment of empyema depends on the severity of the infection and the patient’s underlying condition. Some patients with empyema can be treated with serial thoracenteses, but most will require continuous drainage with a chest tube (see Fig. 10.5). Thoracoscopic decortication represents definitive therapy for severe cases. Usually, diagnostic thoracentesis is performed first to assess the fluid for signs of infection. Thick pus on thoracentesis, positive Gram stain, fluid glucose level lower

**BOX 10.1 Guidelines of the American College of Chest Physicians for the Management of Primary and Secondary Spontaneous Pneumothoraces**

**PRIMARY SPONTANEOUS PNEUMOTHORAX (NO UNDERLYING LUNG DISEASE)**

A clinically stable patient must have all of the following present: respiratory rate lower than 24 breaths/min, heart rate higher than 60 beats/min or less than 120 beats/min, normal blood pressure, room-air $O_2$ saturation higher than 90%, and the ability to speak in whole sentences between breaths.

**CLINICALLY STABLE PATIENTS WITH SMALL PNEUMOTHORACES (<3-CM APEX-TO-CUPOLA DISTANCE)**

Clinically stable patients with small pneumothoraces (PTXs) should be observed in the emergency department (ED) for 3 to 6 hours and be discharged home if a repeated chest radiograph excludes progression of the PTX (good consensus). Patients should be provided with careful instructions for follow-up within 12 hours to 2 days, depending on the circumstances. A chest radiograph should be obtained at the follow-up appointment to document resolution of the PTX. Patients may be admitted for observation if they live distant from emergency services or follow-up care is considered unreliable (good consensus). Simple aspiration of the PTX or insertion of a chest tube is not appropriate for most patients (good consensus) unless the PTX enlarges. The presence of symptoms for longer than 24 hours does not alter the treatment recommendations.

**CLINICALLY STABLE PATIENTS WITH LARGE PNEUMOTHORACES (≥ 3-CM APEX-TO-CUPOLA DISTANCE)**

Clinically stable patients with large PTXs should undergo a procedure to reexpand the lung and should be hospitalized in most instances (very good consensus). The lung should be reexpanded by using a small-bore catheter (≤14 Fr) or placement of a 16- to 22-Fr chest tube (good consensus). Catheters or tubes may be attached either to a Heimlich valve (good consensus) or to a water seal device (good consensus) and may be left in place until the lung expands against the chest wall and the air leaks have resolved. If the lung fails to reexpand quickly, suction should be applied to a water seal device. Alternatively, suction may be applied immediately after placement of a chest tube in all patients managed with a water seal system (some consensus).

Reliable patients who are unwilling to undergo hospitalization may be discharged home from the ED with a small-bore catheter attached to a Heimlich valve if the lung reexpanded after the removal of pleural air (good consensus). Follow-up should be arranged within 2 days. The presence of symptoms for longer than 24 hours does not alter management recommendations.

**SECONDARY SPONTANEOUS PNEUMOTHORAX**

**Clinically Stable Patients With Small Pneumothoraces**

Clinically stable patients with small PTXs should be hospitalized (good consensus). Patients should not be managed in the ED with observation or simple aspiration without hospitalization (very good consensus). Hospitalized patients may be observed (good consensus) or treated with a chest tube (some consensus), depending on the extent of their symptoms and the course of their PTX. Some of the panel members argued against observation alone because of a report of deaths with this approach. Patients should not be referred for thoracoscopy without prior stabilization (very good consensus). The presence of symptoms for longer than 24 hours did not alter the panel members’ recommendations.

**Clinically Stable Patients With Large Pneumothoraces**

Clinically stable patients with large PTXs should undergo placement of a chest tube to reexpand the lung and should be hospitalized (very good consensus). Patients should not be referred for thoracoscopy without prior stabilization with a chest tube (very good consensus). The presence of symptoms for longer than 24 hours did not alter the panel members’ recommendations.

This is meant to be a guide, and clinical judgment should always be used.


**BOX 10.2 Indications for Surgery After Tube Thoracostomy Based on Results of the Thoracostomy**

- Massive hemothorax, >1000-mL to 1500-mL initial drainage
  - Continued bleeding
    - >300 to 500 mL in the first hour
    - >200 mL/hr for the first 3 or more hours
  - Increasing size of the hemothorax on a chest film
  - Persistent hemothorax after two functioning tubes are placed
  - Clotted hemothorax
  - Large air leak preventing effective ventilation
  - Persistent air leak after placement of a second tube or inability to fully expand the lung

This is meant to be a guide and clinical judgment should always be used.
particularly those with penetrating chest injuries. The goal is to open the pleural space quickly to allow any accumulated air to escape and decompress the chest cavity. This can be accomplished with a scalpel and forceps, as is done at the beginning stages of a thoracostomy (Fig. 10.9), or more commonly, decompression can be achieved with a large-bore needle/angiocatheter combination (minimum of 16 gauge) (Fig. 10.10). Place the catheter in the second intercostal space at the midclavicular line on the side with diminished breath sounds or on both sides if the diagnosis is unclear. If unable to obtain access to this landmark, or if unsuccessful in penetrating into the pleural space, an alternative site in the fourth or fifth intercostal space at the midaxillary line can be used. Remove the needle, but leave the angiocatheter in place to create a simple pneumothorax. Regardless of whether needle decompression is successful in improving the patient's vital signs, a standard tube thoracostomy should be immediately performed following the decompression. If the needle decompression is not effective an open thoracostomy can be started, even without the immediate availability of a chest tube, to help create an exit for the air so that the patient's respiratory and cardiovascular functions can normalize. Needle decompression causes an open pneumothorax and in most cases needs to be converted to an open thoracostomy.

CONTRAINdications

For unstable trauma patients with a pneumothorax or hemothorax, there are no absolute contraindications to tube thoracostomy. In critical patients, placement of a chest tube is often performed empirically. In stable patients, relative contraindications include anatomic problems such as the presence of multiple pleural adhesions, emphysematous blebs, or scarring. Coagulopathic patients should be evaluated for replacement of clotting factors before any invasive procedure.

TREATMENT

Treatment of a Possible Tension Pneumothorax in an Unstable Patient

Immediate decompression of the chest must be considered in all injured patients with unexplained hypotension or tachypnea, particularly those with penetrating chest injuries. The goal is to open the pleural space quickly to allow any accumulated air to escape and decompress the chest cavity. This can be accomplished with a scalpel and forceps, as is done at the beginning stages of a thoracostomy (Fig. 10.9), or more commonly, decompression can be achieved with a large-bore needle/angiocatheter combination (minimum of 16 gauge) (Fig. 10.10). Place the catheter in the second intercostal space at the midclavicular line on the side with diminished breath sounds or on both sides if the diagnosis is unclear. If unable to obtain access to this landmark, or if unsuccessful in penetrating into the pleural space, an alternative site in the fourth or fifth intercostal space at the midaxillary line can be used. Remove the needle, but leave the angiocatheter in place to create a simple pneumothorax. Regardless of whether needle decompression is successful in improving the patient's vital signs, a standard tube thoracostomy should be immediately performed following the decompression. If the needle decompression is not effective an open thoracostomy can be started, even without the immediate availability of a chest tube, to help create an exit for the air so that the patient's respiratory and cardiovascular functions can normalize. Needle decompression causes an open pneumothorax and in most cases needs to be converted to an open thoracostomy.

Prehospital Treatment

Emergency needle decompression thoracostomy may be used in the prehospital setting when a patient who is suspected of having a pneumothorax rapidly deteriorates or is initially seen in extremis. In such cases, attach the needle (or catheter) to a
Radiopaque strip along the length of the tube that is interrupted by a series of holes along the distal length of the tube. The strip allows visualization of the tube’s location on the post-procedure radiograph and ensures that the side ports are within the pleural cavity. Tube sizes vary from 12 to 42 Fr, with smaller tubes used for smaller pneumothoraces and larger (a minimum of 36 Fr) tubes for hemothorax and empyema. For pediatric patients, 14-, 16-, 20-, and 24-Fr tubes are adequate.

PROCEDURE
Conduct the procedure (Videos 10.1-10.8) as steriley as possible and wear a gown, glove, mask, and goggles. When possible, obtain consent.

Emergency Department Treatment

Equipment
The standard instruments for a tube thoracostomy tray are listed in Box 10.3 and depicted in Review Box 10.1. The most basic needs are a scalpel, a large (Kelly) clamp, and the thoracostomy (chest) tube.

Thoracostomy tubes are clear plastic tubes of various diameters that are open at both ends (Fig. 10.11). There is a radiopaque strip along the length of the tube that is interrupted by a series of holes along the distal length of the tube. The strip allows visualization of the tube’s location on the post-procedure radiograph and ensures that the side ports are within the pleural cavity. Tube sizes vary from 12 to 42 Fr, with smaller tubes used for smaller pneumothoraces and larger (a minimum of 36 Fr) tubes for hemothorax and empyema. For pediatric patients, 14-, 16-, 20-, and 24-Fr tubes are adequate.

PROCEDURE
Conduct the procedure (Videos 10.1-10.8) as steriley as possible and wear a gown, glove, mask, and goggles. When possible, obtain consent.
possible, elevate the head of the bed 30 to 60 degrees (Fig. 10.13) to lower the diaphragm and decrease the risk for injury to the diaphragm, spleen, and liver. Abduct the arm on the affected side, place it over the patient’s head, and restrain it in that position. To facilitate, explain the process to the patient in detail if conditions are appropriate and use an assistant to help hold the patient and offer comforting support. Clean the skin with a standard surgical scrub and drape the field with sterile towels.

**Anesthesia**

**Preprocedure**

Tube thoracostomy can be extremely painful, so consider giving parenteral analgesics or procedural sedation to stable patients before starting the procedure. A common problem is inadequate systemic analgesia and local anesthesia. Use generous local anesthesia, such as up to 4 mg/kg of locally injected 1% lidocaine with or without epinephrine. Slowly inject local anesthetic over the superior aspect of the rib, through the muscle, periosteum, and parietal pleura and along the entire anticipated tract of passage of the tube (Fig. 10.14). Intermittently aspirate for air or fluid with the needle to find the pleural cavity. If air or fluid is not found, change the insertion site.

**Procedure**

During the procedure, an additional injection of local anesthetic (1% lidocaine) can be administered just prior to bluntly penetrating through the tough parietal pleura with the large Kelly clamp, as this is often the most painful part of the procedure.

**Postprocedure**

Once the tube is in place, administer local anesthetic through the chest tube into the pleural space to reduce pain caused by the tube rubbing against the pleura. In a stable patient, add

---

**Tube Insertion Site**

Insert the tube over the top of the rib rather than near the bottom to avoid the neurovascular structures located on the inferior aspect of the ribs. The most common location for a chest tube is the midaxillary to anterior axillary line, usually in the fourth or fifth intercostal space (Fig. 10.12). This approach is cosmetically preferable and better tolerated than the anterior chest wall approach in the second intercostal space of the midclavicular line. The fifth intercostal space is approximately at the level of the nipple or the inferior scapular border in most patients, although the position of the female breast mass leads to variance. To avoid penetrating the abdominal cavity, choose a more superior insertion site because the external landmarks can be misleading. An important point to consider is that the diaphragm of a supine patient who is not taking a deep breath is much higher than expected.

Hold the tube next to the chest wall with the tip of the tube at the level of the clavicle to estimate the distance that the tube needs to be advanced from the incision site to the apex of the lung. Place a clamp on the tube to mark the maximum length that the tube should be inserted to prevent the tube from advancing too far. Confirm that the last drainage hole is within the pleural space at the level of the insertion site to ensure that the tube has been advanced sufficiently far. In markedly obese patients, it is common to fail to advance the tube far enough, with the last hole being left in fatty tissue rather than in the pleural space.

**Patient Preparation**

If indicated clinically, start oxygenating and monitoring the patient continuously with cardiac and pulse oximetry. When
infiltrating and aspirating until the pleura is breeched and air is infiltrated over the rib at the site of the incision. With a generous amount of local anesthetic.

Both the skin and pleura should be infiltrated to control pain. Vacuum drainage. Use parenteral analgesic agents as needed without drainage of the thorax, reinitiate standard gravity or the pleura before pushing through (Fig. 10.17). As stated few centimeters distal to the incision and rest the tip against the tough parietal pleura is met. Close the clamp and push it through which the tube will enter, followed by “tunneling” the tube under the skin and up over the next rib. This is theoretically done to prevent air leaks, but there is no good evidence to support this practice. It is more common to simply make the incision at the intended location so that there is a straight path to the pleural cavity over the appropriate rib. This avoids the problem of losing the entrance point or increasing damage to the soft tissues, both issues that are particularly problematic in obese patients. After the incision is made, insert a large Kelly clamp to push and spread the deeper tissues. Bluntly dissect a tract over the rib while avoiding the intercostal vessels and nerve on the inferior margin of each rib (Fig. 10.16; see also Fig. 10.15, step 2). Firm resistance will usually be felt when the tough parietal pleura is met. Close the clamp and push it forward with firm pressure to penetrate the pleura and enter the cavity. Considerable force may be needed. To prevent the clamp from penetrating too deeply, hold it at the midshaft a few centimeters distal to the incision and rest the tip against the pleura before pushing through (Fig. 10.17). As stated previously, penetrating the pleura is usually the most painful portion of the procedure, so consider injecting extra anesthetic or analgesic at this point. On entering the pleural cavity, a palpable pop may be felt and a rush of air or fluid may occur. With only the tips of the clamp in the pleural cavity, spread the clamp to make an adequate hole in the pleura and then withdraw it (see Fig. 10.15, step 3). Make the opening in the parietal pleura wide enough to comfortably insert both a finger and the tube, but avoid a larger pleural opening to reduce the risk of an air leak (Fig. 10.18).

Another common problem occurs at this point, particularly in obese patients: the dissected tract and pleural opening can be lost when the clamp is withdrawn. To prevent this problem, slide a sterile gloved finger over the clamp and into the pleura before withdrawing the clamp (Fig. 10.19). This is done to further define the tract, to verify that the pleura has been entered, and to be sure that no solid organs have been penetrated. Whenever possible, leave the finger in the pleural space, make a 360-degree sweep to verify the correct space, to feel for adhesions, and to ensure that the hole is not lost (see Fig. 10.15, step 4). Pass the tube over, under, or beside the finger into the pleural space, with the fingertip being used to guide the course of the tube (Fig. 10.20). This step allows the clinician to feel the tube passing into the pleural cavity, avoids subcutaneous dissection by the tube, and enhances proper direction of the tube. Pass the tube alone or hold on to a large curved clamp with the tip of the tube protruding beyond the tip of the clamp (Fig. 10.21; see also Fig. 10.15, step 5). Normally, the tube should pass with little resistance. If resistance is met, the tube may not be in the pleural cavity and instead is passing subcutaneously, is in a fissure, or is abutting against the mediastinum (Fig. 10.22). Still using the finger that remains in the pleural space, direct the tube posteriorly, medially, and superiorly until the last hole of the tube is clearly in the thorax, the marker clamp that was previously attached touches the chest wall, or resistance is felt (see Fig. 10.15, step 6). Ensure that all the holes in the tube are within the pleural space. Rotate the tube 360 degrees to reduce the likelihood of kinking.

Attach the tube to the previously assembled water seal or suction before releasing the clamp. Ask the patient to cough and look for bubbles in the water seal chamber to check for patency of the system.

In cases in which there may be broken ribs or significant resistance, some authors recommend the use of a bougie to ensure that the tract (that was formed by the Kelly clamp and the inserted finger) is not lost. After confirmation of the correct space by finger sweep, insert the bougie while the finger is in the pleural space and direct it posteriorly, medially, and superiorly until the last hole of the tube is clearly in the thorax, the marker clamp that was previously attached touches the chest wall, or resistance is felt (Fig. 10.23). Using the bougie as a guide allows the operator to identify the correct distance for advancement. Once the operator is satisfied that the bougie is in the correct place, the chest tube can then be inserted over it. As with most modified Seldinger techniques, the bougie must be secured by one hand at all times, and the tube must be advanced to its correct depth over the bougie, while the bougie remains stationary.

Confirmation of Tube Placement

There are many ways to confirm the location of the tube. Slide a finger along the tube to verify that it has entered the pleural cavity. Look for condensation on the inside of the tube and

![Image](image-url)
Position the patient, prepare the skin, and administer local anesthetic.

Use a scalpel with a No. 10 blade to make a transverse 3- to 5-cm incision through the skin and subcutaneous tissue, over the rib.

With only the clamp tips in the pleural cavity, spread the clamps to make an adequate hole in the pleura, and then withdraw it.

The opening in the pleura should be wide enough to insert a finger and the tube. Avoid making a larger opening to reduce air leak.

Alternatively, if a finger is not used as a tube guide, hold the tube in a large curved clamp, and pass it into the pleural cavity. The tube should pass with little resistance. If resistance is met, the tube may not be in the pleural cavity and may be passing subcutaneously, entering a fissure, or abutting the mediastinum.

Secure the tube to the chest with sutures.

Specific techniques to secure the tube are discussed in detail in text.

Use a large Kelly clamp to push and spread the deeper tissues, and bluntly dissect a track over the rib, while avoiding the vessels on the inferior surface of the rib.

Firm resistance will be felt when the parietal pleura is met. Close the clamp and push it forward to penetrate the pleura.

Before removing the clamp, slide a finger over it and into the pleural cavity so that the dissected tract is not lost.

Leave the finger in the pleural space, and pass the tube alongside the finger during insertion. Verify that the pleural cavity has been entered, and that no solid organs are present.

Direct the tube posteriorly, medially, and superiorly until the last hole of the tube is clearly intrathoracic or resistance is felt.

Attach the tube to the previously assembled water seal or suction system. Ask the patient to cough, and observe bubbles in the water seal chamber to assess patency of the system.

After suturing the tube, place an occlusive dressing of petrolatum-impregnated gauze at the point where the tube enters the skin.

This will help prevent air leaks.

**Figure 10.15** Tube thoracostomy. (From Custalow CB: *Color atlas of emergency department procedures*, Philadelphia, 2005, Saunders.)
listen for air movement, which is audible during respirations. Observe free flow of blood or fluid. Check to be sure that the tube rotates freely after insertion. A chest radiograph is used for definitive assessment of tube placement (Fig. 10.23). If the tube and the most proximal hole are not completely within the pleural space, and if the field has remained sterile, advance the tube. If the tube is kinked or dysfunctional or the sterile field has been lost and advancement is required, place a new tube in sterile fashion through the same tract. If the tube has been advanced too far, simply withdraw it to the correct depth.

**Securing the Tube**

Once the position of the tube has been verified with a radiograph, secure it. There are numerous methods to secure a tube. The usual one is to sew the tube to the skin with large 0 or 1-0 silk or nylon sutures. Nylon sutures are acceptable but must be tied tightly or they will slip on the surface of the tube. One common method is to use a “stay” suture, in which the same suture that closes the skin incision is used to hold the tube (Fig. 10.24, plate 1). After this suture is used to close the skin incision at the site of insertion of the tube, wrap the ends tightly and repeatedly around the chest tube and tie it securely. Tie the sutures tightly enough to indent the chest tube slightly and avoid slippage. If the skin incision is especially long, use additional simple sutures to close it completely.

Another optional suture technique can both help close the skin around the tube and completely close the incision once the chest tube is removed. To do this, place a horizontal mattress suture approximately 1 cm across the incision on either side

---

**Figure 10.16** Direct the Kelly clamp over the top of the rib to avoid the intercostal neurovascular bundle that courses along the inferior portion of the rib. Do not use excessive tunneling. (From Millikan JS, Moore EE, Steiner E: Complications of tube thoracostomy for acute trauma, *Am J Surg* 140:739, 1980.)

**Figure 10.17** A and B, Use the right hand to push the clamp through the pleura (white arrows). Firmly encircle the midportion of the clamp with the fingers of the left hand to serve as a safety stop measure to avoid penetrating too deeply.

**Figure 10.18** Right-sided subcutaneous emphysema after chest tube placement secondary to making too large a hole in the pleura with a subsequent air leak (arrows). It is usually benign and self-limited, but with positive pressure ventilation it can be problematic. Because there is no way to close the pleura, making just the right-sized hole initially is the key to avoiding this complication.
CHAPTER 10  Tube Thoracostomy

213

wrap the center section tightly around the tube (see Fig. 10.24, plate 4). Repeat this with a second piece of tape placed at a 180-degree angle to the first. In addition, securely tape the tube connections. To further secure the tube, use tape to create a loop or stalk by wrapping it around the tube. Then press the tape together for 1 to 2 cm before applying the tape to the chest wall (see Fig. 10.24, plate 5).

Drainage and Suction Systems

A basic understanding of chest tube drainage systems is necessary to prevent life-threatening complications associated with their use. All drainage systems have two essential components: a one-way valve to allow air or fluid to drain out of the pleural

Figure 10.19  After puncturing the pleural lining and spreading with the clamp, slide a gloved finger over the clamp to ensure that the pleural space has been reached (white arrow) and that no solid underlying masses are present. Then withdraw the clamp but keep the finger inside the tract. Use the finger as a guide for the chest tube to ensure entry into the pleural cavity.

Figure 10.20  Leave your finger in the pleural space and use it to guide the course of the tube (arrows).

Figure 10.21  Hold the tube with the tip of the tube protruding beyond the tip of the clamp to reduce the risk for pulmonary injury.

Figure 10.22  Subcutaneous placement of a chest tube (arrows) can occur because the tube can dissect through tissue planes with relative ease. If this tube had been directed posteriorly, the radiograph would erroneously “confirm” intrapleural placement despite the tube being subcutaneous throughout its entire course. There would be no air fogging the tube with respirations but a computed tomography scan would settle the issue of a misplaced tube.
bubbling can be detected. Vigorous bubbling does not equate with more suction.

The bottle combinations example is provided to illustrate the principles but is rarely used now. Many types of commercial, enclosed systems are available that essentially combine a two-bottle method that can be connected to suction, but with the addition of an “air leak chamber” (see Fig. 10.26C). Bubbling in this chamber indicates the presence of an air leak, usually from the drainage system itself as a result of a loose tube connection. If bubbling is present, first check the drainage system, tube, and connectors for any problems or loose connections. If the leak continues, check whether all the holes of the chest tube are within the thorax. If the bubbling continues after these steps, a continued air leak may be due to a large hole in the lung parenchyma, which is usually seen only with expiration or with coughing. A continuous air leak or a leak during inspiration indicates a larger and possibly more significant lung injury. Surgical intervention is indicated if an air leak persists for longer than 72 hours or if the lung is not completely reexpanded.

The drainage reservoir must remain below the level of the chest to prevent the fluid in the collection system from reentering the chest. Place the reservoir on the floor or hang it from the edge of the bed. Simple respirations do not generate enough negative intrathoracic pressure to pull the water in the reservoir up to the height of the chest if the reservoir is kept on the floor and the patient is either sitting or lying at standard chair or bed height. The length of the tubing should be long enough so that the reservoir can be kept below the level of the patient, but not long enough to cause it to form dependent loops of fluid or kinks. Dependent loops collect fluid and create an additional water seal that, if large enough, can require greater intrapleural pressure to drain. If these pressures become high enough (15 to 25 cm H₂O), a tension pneumothorax may result.

When the drainage system is functioning properly, the height of the fluid level in the drainage tube fluctuates with inspiration and expiration. Absence of respiratory fluctuation or a decrease in drainage may indicate that the system is blocked or that the lung is fully expanded. If the tube is blocked, the chest tube, collecting tubing, or both, can be changed or stripped to dislodge clots. Replacing the tube is a complicated process associated with risks, but the stripping procedure is controversial because of concern that the potentially high negative pressure from the procedure could damage lung tissue. For stripping, clamp the tube proximally and progressively compress it distally and then release it to allow the tube to spring open. The sudden increase in negative pressure may extract clots and fluid from a more proximal location. If the blockage is within the thorax, the tube can be cleared by forcing air or fluid back into the chest. The tube must be clamped distally and then compressed and stripped to force the contents proximally.

Perform occlusive clamping of a chest tube only with close monitoring because a tension pneumothorax can result in rare cases. Patients with chest tubes in place are best transported with a Heimlich valve or water seal only, not with the tube clamped. Avoid clamping the chest tube as a trial maneuver before removing it.

Prophylactic Antibiotics

The use of prophylactic antibiotics after placement of a chest tube in the emergency department (ED) is common, but no
To secure the tube, first close the skin incision with a "stay" suture near the tube. Tie the knot securely and leave the suture ends long for wrapping around and tying the tube. Wrap the suture tightly at least twice around the tube, enough to indent the tube slightly, and tie securely.

Another method to close the wound and secure the tube is with a horizontal mattress suture combined with a stay suture. A horizontal mattress suture is placed on either side (above and below) of the tube and held only with a surgeon's knot. The loose ends are also wrapped around the tube and tied loosely in a bow to identify the suture. This suture will be untied and used to close the skin incision after removal of the tube.

To dress the wound and reduce the risk for air leaks, an occlusive dressing should be applied. First wrap the base of the tube at the skin incision with a petroleum-impregnated dressing. A two-layer dressing of gauze sponges with a Y-shaped cut centered at the tube is shown. Place the second layer at a 90° angle to the first.

One method to further secure the tube is to use wide, split cloth tape. The distal half of a 15- to 20-cm-long wide piece of tape is longitudinally split into three pieces. The two outside pieces are placed on the skin on either side of the tube, and the center strip is wrapped around the chest tube itself. This process may be repeated with a similar piece of tape placed at a 90° angle. The tape is securely anchored to the skin (benzoin is optional, but the skin must be clean and dry), and the torn tape is wrapped around the tube. Each anchoring piece is covered by another piece of tape.

The tube can be further secured with an additional anchor system farther down on the tube. Wrap a 20- to 25-cm piece of tape or elastic, adhesive dressing around the tube and seal at least 3 cm of the tape together on the side of the tube nearest the chest wall. Spread the remaining tape against the dry skin of the chest wall and secure with additional tape.

Figure 10.24 Methods of securing a thoracostomy tube.
specific standards exist. It appears reasonable to provide short-
term antibiotic prophylaxis because analyses of many studies 
have shown that antibiotics reduce the incidence of chest 
tube–associated empyema or pneumonia. Guidelines from 
the Eastern Association of Surgeons of Trauma recommend 
the use of first-generation cephalosporins during the first 
24 hours for patients undergoing chest tube drainage for 
hemothorax.

**Tube Removal**

Chest tubes are rarely removed by emergency clinicians. The 
usual indication for removal of a chest tube is after a chest 
radiograph demonstrates complete resolution of the pneumo-
thorax and there is no evidence of an ongoing air leak. Before 
removal, discontinue suction and establish a water seal. The 
literature is not clear about the use of a chest radiograph after 
suction is discontinued and after pulling the tube. Textbooks 
continue to recommend such practice, but studies have shown 
that it is not necessary for postoperative patients.

To remove the chest tube, ask the patient to sit upright at 
approximately a 45-degree angle. Use sterile technique, clean 
the skin, and drape the insertion site. Suturing equipment 
is needed to close the wound after the tube is removed, except 
when a purse-string suture was placed at the time of insertion,
in which case only sterile scissors are needed to cut the suture. 
Keep additional equipment available to reinsert a chest tube 
if the lung collapses. Prepare a petrolatum- or antibiotic-
impregnated gauze dressing to cover the wound.

If a purse-string suture was placed previously, loosen it and 
prepare it for closing the wound. Cut the skin loop of the 
suture holding the tube to the skin and remove it from the 
skin. With the tube clamped, it has traditionally been advised 
to ask the patient to inhale fully and perform a mild Valsalva 
maneuver. Although this is theoretically beneficial in minimizing 
residual air following tube removal, it is of no proven value 
and many clinicians use no special protocols for removal of
the tube. Pull the tube out in one swift motion. Quickly tie the purse-string suture or suture the wound closed, and then cover it with an occlusive dressing. If no purse-string suture is present, once the tube is removed, suture the skin as quickly as is prudent for the physician. Observe the patient for 2 to 6 hours and obtain a chest radiograph before discharge. Any increase in symptoms requires prompt reevaluation. After 48 hours the patient may remove the dressing. Remove the sutures in 7 to 10 days.

OTHER TECHNIQUES

Small-Bore Tube Thoracostomy

Since the advent of common pleural drainage techniques in the early 20th century, physicians have preferred large-bore pleural drainage catheters (PDCs) inserted via blunt dissection for thoracic drainage and decompression. This preference for tubes greater than 16 Fr in diameter was born largely of concern for fibrin blockage, inadequate drainage capability, and the perception of frequent dislodgement of small-bore PDCs. To date, no quality randomized clinical trials have compared the use of large- versus small-bore PDCs, mainly because of the difficulty of conducting comparative studies in critically ill patients.

Despite these impediments, studies have hinted at the adequacy of small-bore PDCs for the treatment of many conditions. Review of the data available suggests that the rate of blockage and dislodgement is similar with large- and small-bore PDCs and that drainage capability is comparable in all but the most extreme circumstances. Another study has shown that large-bore PDC placement is associated with significantly more pain and slightly increased rates of empyema.

Review of the pulmonology and critical care literature reveals no consensus on the appropriate use of small-bore PDCs; however, review articles have attempted to establish rough guidelines for their use. The literature suggests that patients with malignant effusion, simple pneumothorax without penetrating injury, or a suspected air leak, as well as those with simple or complex parapneumonic effusions, are good candidates for small-bore PDCs. Patients with empyema, penetrating injury, traumatic hemotorax or complex pneumothorax, and those on mechanical ventilation require large-bore PDCs because of the potential for large-volume air leaks in such patients and the need for rapid evacuation of the pleural space.

Many protocols are available for use of catheter aspiration as the first step in treating simple pneumothoraces. In general, patients with successful aspiration are observed in the ED for 4 to 6 hours after catheter insertion, and if a repeated radiograph shows no reaccumulation of air, the catheter is removed. After 2 more hours another chest radiograph is obtained, and the patient is released if there is no recurrent pneumothorax. Patients with continued residual pneumothorax often receive a conventional chest tube.

Guidewire Technique for Catheter Aspiration

Catheters designed specifically for aspirating a pneumothorax are made of flexible, thrombosis-resistant radiopaque material with multiple distal side ports to reduce the risk of occlusion. Commercially available small-bore catheter systems are ideal for this procedure.

The catheters are placed via a standard “over-the-wire” (Seldinger) technique. The most common insertion site is the second intercostal space in the midclavicular line, but either of the standard locations (the midaxillary to anterior axillary line, usually in the fourth or fifth intercostal space, or the midclavicular line, second intercostal space) can be used. Place the patient in a semi-upright position. Clean the skin with an antiseptic solution and drape the area. Infiltrate locally with lidocaine for anesthesia. Advance the guide needle in a straight line at a 60-degree angle cephalad over the top of the rib (Figs. 10.27 and 10.28). Unless a straight tract is created, it will be difficult to advance the floppy catheter, so a tunneling approach cannot be used. When the pleural space is identified by intermittent aspiration, halt advancement of the needle. Stabilize the needle and feed a guidewire through the needle and into the pleural space. Remove the needle while stabilizing the guidewire to keep it in the pleural space. Make a small incision in the skin with a No. 11 blade at the base of the wire to allow passage of the catheter through the skin. Some systems use a dilator over the wire to open the path through the soft tissue. Thread the mini-catheter over the guidewire and into the pleural space. Remove the wire and dilator while leaving the catheter in the pleural space. Advance the catheter through the subcutaneous tissue with a twisting motion. Secure the catheter to the skin with a suture and dress the incision site. The catheter may be removed after a period of observation, or suction may be maintained for a few days. If the catheter becomes clogged with mucus or blood, inject sterile saline through the device to clear it.

To aspirate the pneumothorax, attach a three-way stopcock to the catheter and slowly aspirate air with a 60-mL syringe until resistance is felt. Gentle wall suction can also be used because a number of aspirations may be required until all the air exits. Take a chest radiograph to determine whether the lung is fully expanded. If a residual pneumothorax is present, attempt further aspirations. If the residual pneumothorax persists and air cannot be aspirated, the catheter may be kinked or blocked with soft tissue. To relieve the blockage, place the patient in the full upright position and have the patient cough or take a deep breath. Alternatively, the catheter can be twisted or rotated gently.

TUBE THORACOSTOMY IN PEDIATRIC PATIENTS

Pneumothoraces can occur in the neonatal population. They are often associated with resuscitative measures (such as mechanical ventilation) for meconium aspiration or prematurity. For the rest of the pediatric population, trauma is the most common cause. A pneumothorax will develop in approximately one third of children with thoracic trauma. As in adults, physical examination of newborns and infants with pneumothorax can be highly variable, thus necessitating the use of a chest radiograph for diagnosis. Ideally, both anteroposterior and cross-table lateral x-ray projections are used because small pneumothoraces may be seen only on the lateral view. A chest CT is more sensitive and specific.

In general, tube thoracostomy is the treatment of choice once a symptomatic pneumothorax is detected in infants. When signs of tension pneumothorax are present, immediately aspirate with a plastic catheter-over-the-needle device. Small pneumothoraces (<20% of the hemithorax) in relatively asymptomatic
The Seldinger-type catheter kit contains a pigtail catheter and all necessary equipment, including local anesthesia, introducing needle and syringe, scalpel, guidewire, and dilator.

After generous local anesthesia, advance the introducing syringe in a straight line over the top of the fifth rib until air is aspirated. Unless a straight tract is created, it will be difficult to advance the floppy catheter, and a tunneling approach cannot be used.

Advance the guidewire through the introducing needle into the pleural space, and then remove the needle. The procedural steps are analogous to initiating a central venous catheter via the Seldinger technique.

Puncture the skin at the site of wire insertion with a scalpel. Make the incision large enough to accept the dilator and pigtail catheter.

Advance the dilator over the wire to create a tract for the catheter. Remove the dilator while leaving the wire in place. Again, this is analogous to establishing a central venous line.

Advance the pigtail catheter over the wire through the dilated tract. It will assume its pigtail configuration when it is in the pleural space. A twisting motion may be needed to advance the catheter through subcutaneous tissue. Advance the catheter to the hilt and secure to suction. This catheter may be removed after a period of observation, or the suction may be maintained for a few days.

Figure 10.27 Aspiration of a pneumothorax with an Arrow 14-French Percutaneous Cavity Drainage Catheterization Kit (Teleflex, Morrisville, NC). This 23-cm pigtail multihole catheter is ideal for such purposes. Air can be aspirated from the catheter with a syringe, or the catheter can be attached to suction or a Heimlich valve. This catheter is not used for patients on a ventilator, those with continuing air leaks, or those with a hemothorax. It is ideal for stable patients who have a primary pneumothorax or a collapse that can be expected to be stable if the lung is reexpanded (such as induced by intravenous drug use, minor blunt trauma, or insertion of a central venous catheter). If used for a few days, the catheter will become clogged with mucus or blood, which may be cleared by injecting sterile saline through the device.
infants (e.g., those without other problems who do not require positive airway pressure) can be observed.

The technique of tube thoracostomy in pediatric patients is essentially the same as that in adults, but the body size and small spaces between the ribs make the procedure more difficult. The size of the tube increases with the patient's weight, starting with 8-Fr to 10-Fr catheters for premature infants (Table 10.1). Because of the risk for future breast deformities, avoid the midclavicular approach. Instead, use the anterior axillary line through the fifth intercostal space for newborns and infants.

### COMPLICATIONS

The most common complications include infection, laceration of an intercostal vessel, laceration of the lung, and intraabdominal or solid organ placement of the chest tube (Boxes 10.4 and 10.5). Local infection at the insertion site is common and often related to the emergency nature of the procedure. Subcutaneous

### Table 10.1

<table>
<thead>
<tr>
<th>WEIGHT (KG)</th>
<th>CHEST TUBE (FR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;3</td>
<td>8–10</td>
</tr>
<tr>
<td>3–5</td>
<td>10–12</td>
</tr>
<tr>
<td>6–10</td>
<td>12–16</td>
</tr>
<tr>
<td>11–15</td>
<td>17–22</td>
</tr>
<tr>
<td>16–20</td>
<td>22–26</td>
</tr>
<tr>
<td>21–30</td>
<td>26–32</td>
</tr>
<tr>
<td>&gt;30</td>
<td>32–40</td>
</tr>
</tbody>
</table>

---

**Figure 10.28** Catheter-over-the-needle technique for aspiration of a pneumothorax.
### BOX 10.4 Physical Complications of Tube Thoracostomy

**INFECTION**
- Pneumonia
- Empyema
- Local infection of the incision
- Osteomyelitis
- Necrotizing fasciitis

**PHYSIOLOGIC**
- Allergic reactions to the surgical preparation or anesthesia
- Pulmonary atelectasis
- Reexpansion pulmonary edema
- Reexpansion hypotension

**INJURIES—BLEEDING**
- Local incision hematoma
- Intercostal artery or vein laceration
- Internal mammary artery laceration (with midclavicular line placement)
- Pulmonary vein or artery injury
- Great-vessel injury

**INJURIES TO SOLID ORGANS OR NERVES**
- Lung, liver, spleen, diaphragm, stomach, colon; long thoracic nerve, intercostal nerve

**MISCELLANEOUS**
- Subcutaneous or mediastinal emphysema
- Persistent pneumothorax
- Retained hemothorax
- Recurrence of pneumothorax after removal of the chest tube

Emphysema is a frequent, usually benign complication that is self-limited and generally caused by an excessively large opening in the pleura. Subcutaneous air may be a result of the incident that caused the pneumothorax in the first place. The development of palpable subcutaneous air is another complication of chest tube placement. It is usually limited to the insertion site but can become massive with PPV in a patient with continued air leak or an occluded tube.

Intercostal arteries or veins may be lacerated, but this can be minimized by using blunt dissection and carefully directing the tube just above the rib. The tube may adequately tamponade such bleeding. If the bleeding persists, extend the incision to ligate the bleeding vessel. If the bleeding continues, consult a thoracic surgeon.

### BOX 10.5 Mechanical Complications of Tube Thoracostomy

**MECHANICAL PROBLEMS**
- Dislodgement of the chest tube from the chest wall
- Incorrect tube position
- Subcutaneous placement
- Intraabdominal placement

**AIR LEAKS**
- Leaks within the drainage system (tubing or drainage device)
- Last tube port not within the pleural space
- Leaks from a skin site

**BLOCKED DRAINAGE**
- Flow of drainage contents into the chest from elevation of the drainage bottles
- Kinked chest tube or drainage tubes
- Occlusion of the tube by clots

Failure of a pneumothorax to reexpand may be due to a mechanical air leak, but it may also indicate a bronchopleural fistula, a continued parenchymal lung leak, or a bronchial injury. Tension pneumothorax can occur if a blockage in the drainage system at any point is associated with a continued air leak from the lung. Reinsetion or placement of a second tube may be indicated if the first tube is not functioning properly. In general, if a chest tube is not functioning properly and the patient is deteriorating, remove the tube and insert another one. Manipulating the tube by pushing it deeper into the chest cavity can lead to an increased risk for infection.

A rare complication of tube thoracostomy is unilateral reexpansion pulmonary edema. The pulmonary edema ranges from mild to severe, but fatalities have been reported. The condition may occur shortly after reexpansion or may be delayed a number of hours. A common factor in these cases seems to be a prolonged period between the development of a pneumothorax and the onset of treatment, but the exact time frame is quite variable. Usually, the pneumothorax has been present for at least 3 to 4 days. Proposed mechanisms include anoxic damage to the alveolar-capillary basement membrane from prolonged pulmonary collapse, loss of surfactant, or rapid fluid shifts. Treatment is supportive, with ventilatory support occasionally required. Reintroduction of air back into the pleural space and temporary occlusion of the ipsilateral pulmonary artery are other suggested, but unproved interventions.

**REFERENCES ARE AVAILABLE AT** [www.expertconsult.com](http://www.expertconsult.com)
REFERENCES


Techniques for Supraventricular Tachycardias
Bohdan M. Minczak and Glenn W. Laub

INTRODUCTION

Patients in the emergency department frequently complain of palpitations, heart fluttering, or a rapid heartbeat, which is often coupled with weakness, chest pain, or dizziness. The physician must determine the exact rate, rhythm, origin, and cause of the tachycardia and then gain control of the heart rate (HR) by slowing or normalizing it, or by treating the underlying cause. Determining the cause, origin, and rhythm of the tachycardia is often complicated by the fact that the underlying rate may be very fast (in excess of 150 to 300 beats/min), which makes interpretation of the electrocardiogram (ECG) more difficult. Furthermore, the sources or pacemakers producing or facilitating the tachyarrhythmia may be from one or multiple locations: in the sinoatrial (SA) node, in one or more ectopic atrial foci, in the atrioventricular (AV) node, or in the ventricular free walls or septum. There may also be an abnormal conduction pathway between the atria and the ventricles. In some conditions, one or more pacemakers can be discharging simultaneously. To facilitate the diagnostic process, discrimination between atrial and ventricular electromechanical activity must be attempted. This chapter provides a framework to facilitate the decision-making process with a focus on emergency interventions for various tachyarrhythmias.

Techniques for unmasking, identifying, and treating the various forms of tachyarrhythmias are presented in Box 11.1. This chapter addresses the utility of the vagal reflex in treating and managing various pathophysiologic conditions and the use of medications and cardioversion as they apply to the treatment of various supraventricular tachycardias (SVTs). The major focus is on the evaluation and treatment of SVTs.

A more comprehensive discussion regarding the treatment of ventricular tachycardia (VT) is provided in Chapter 12.

OVERVIEW/SIGNIFICANCE: ANATOMY AND PHYSIOLOGY OF SUPRAVENTRICULAR TACHYCARDIA

Normally, the human heart beats at approximately 80 beats/min (+20 beats/min). If the HR exceeds 100 beats/min, it is called tachycardia. If it drops below 60 beats/min, it is called bradycardia. The heart’s ability to increase the rate of a normal sinus rhythm is primarily related to age: the maximum HR possible with a sinus tachycardia is approximately 220 beats/min minus age, with normal variations as high as 10 to 20 beats/min. As an example, a 60-year-old man cannot usually mount a sinus tachycardia higher than 160 beats/min in response to sepsis, exercise, fever, anxiety, or adrenergic stimulation. Faster rates would indicate a pathologic cardiac rhythm, not a physiologic response.

There are two general categories or types of tachycardias: SVT and VT. SVT describes a rapid HR that has its electro-chemical origin either in the atria or in the upper portions of the AV node. VTs originate in the ventricular free walls or interventricular septum (or both). VTs can quickly become unstable and require special consideration (Fig. 11.1F).

SVTs can be further classified as narrow-complex (QRS duration < 0.12 second; or three small boxes on the ECG) and wide-complex tachycardias (QRS duration > 0.12 second). The rhythms of these dysrhythmias can be regular or irregular. Examples of narrow-complex SVTs are sinus tachycardia (see Fig. 11.1A); atrial fibrillation (AF) (see Fig. 11.1C); atrial flutter (see Fig. 11.1D); AV nodal reentry; atrial tachycardia (see Fig. 11.1B), both ectopic and reentrant; multifocal atrial tachycardia (MAT); junctional tachycardia; and accessory pathway-mediated tachycardia. The term wide-complex tachycardia describes rhythms such as VT (see Fig. 11.1F), SVT with aberrancy (see Fig. 11.1E), or a preexcitation tachycardia facilitated by an accessory pathway between the atria and ventricles.

Tachycardias can be benign or can have significant physical effects on the patient. When the HR is 60 beats/min, approximately one cardiac cycle of contraction (systole) and relaxation (diastole) occurs per second. The excitation for cardiac contraction typically originates in the SA node, the intrinsic pacemaker of the heart. The pacemaker impulse traverses across and depolarizes the atria, which causes atrial contraction or systole. Subsequently, the depolarization reaches the AV node. On initiating depolarization of the AV node, the conduction velocity of this depolarizing impulse transiently decreases (i.e., undergoes “decremental conduction”) so that the ventricles can fill with blood from the antecedent atrial contraction. (Remember: the duration of diastole must be roughly twice the duration of systole to allow adequate ventricular filling.) The AV node also serves as a gate or selective block to prevent an excessive
AF (via the AV node), or simply producing a sinus bradycardia stimulation results in slowing of electrical activity, examples of atrial depolarization impulses to the ventricles. Hence, vagal whereas vagal output to the AV node enhances nodal blockade decreasing the depolarization rate of the “intrinsic pacemaker,” adrenal gland and cause the HR to increase. Parasympathetic epinephrine and norepinephrine, which are released from the atria and ventricles, and by circulating hormones such as nerves (i.e., the autonomic nervous system). Sympathetic input to a balance of input from the sympathetic and parasympathetic during ventricular depolarization) (Fig. 11.2).

The discharge rate of the SA node is usually modulated by a balance of input from the sympathetic and parasympathetic nerves (i.e., the autonomic nervous system). Sympathetic input to the heart is provided by the adrenergic nerves, which innervate the atria and ventricles, and by circulating hormones such as epinephrine and norepinephrine, which are released from the adrenal gland and cause the HR to increase. Parasympathetic input to the heart is provided by the vagus nerve (cranial nerve X) fibers. These nerve fibers innervate the SA and AV nodes. Vagal output to the SA node causes slowing of the HR by decreasing the depolarization rate of the “intrinsic pacemaker,” whereas vagal output to the AV node enhances nodal blockade of atrial depolarization impulses to the ventricles. Hence, vagal stimulation results in slowing of electrical activity, examples being termination of an SVT, slowing of the ventricular rate of AF (via the AV node), or simply producing a sinus bradycardia (via the SA node). Under normal physiologic circumstances, the HR is modulated to meet the metabolic needs of the body’s peripheral circulation. Changes in AV electrochemical events (i.e., rates and rhythms) are manifested as changes in the electrocardiographic intervals and waveforms.

As noted earlier, SVT rhythms can be either sinus (i.e., originating in the SA node: sinus tachycardia) or ectopic (i.e., originating in atrial myocytes above the ventricles). The rate of discharge of the SA node often varies as a result of various physiologic and pharmacologic stimuli, including fever, hypovolemia, shock, anemia, hypoxia, anxiety, pain, cocaine, and amphetamines. These conditions often require or precipitate increased blood flow and hence cardiac output (CO) to peripheral tissues. This increase in peripheral blood flow or CO is accomplished by an increase in heart rate (HR = CO × SV [stroke volume]). These are usually normal, benign physiologic responses to various stimuli or triggers. Direct treatment of the rate of discharge of the SA node is not generally necessary; however, determining and treating the cause of the sinus tachycardia usually eliminates the fast HR. Nonetheless, when single or multiple ectopic, spontaneously discharging foci develop in the atria or upper portions of the AV node, they can begin to “take over” or “override” the normal pacemaker activity in the heart (i.e., the SA node) and produce a rapid HR exceeding 100 beats/min. These foci may develop as a result of increased irritability or automaticity of atrial myocytes secondary to electrolyte abnormalities, hypoxia, pharmacologic agents, or atrial stretch caused by volumetric overload. If these foci are not treated or suppressed and the atrial depolarization rate proceeds to accelerate to rates greater than 150 beats/min (i.e., the heart is beating in excess of 2 beats/sec) and the impulses get through the AV node to the ventricles, the time for diastolic filling of the ventricles will be compromised and result in a precipitous drop in CO. This will ultimately cause a drop in CO regardless of the increase in HR. Furthermore, as CO begins to drop, mean arterial blood pressure (MAP) will decrease and cause hypoperfusion of the brain and other peripheral tissue (MAP is the product of CO times total peripheral resistance [TPR]: MAP = CO × TPR). Treatment of this tachycardia can be achieved pharmacologically by suppressing the automaticity of myocytes with medications (e.g., calcium channel blockers or β-blockers) and subsequently treating the underlying cause or causes, such as hypoxia, electrolytes, and the like. Decreasing the hemodynamic consequences of this arrhythmia requires increasing the “blocking” of these impulses from reaching the ventricles via the AV node. This can be done by enhancing vagal input to the AV node or by pharmacologic enhancement of AV blockade. Multiple rapid depolarizations of the atria, which are conducted to the ventricles, can ultimately have a bimodal type of response: a modest increase in HR will cause an increase in CO, whereas a massive increase in the atrial rate with a concomitant increase in the ventricular rate will cause a drop in CO. This can lead to an unstable patient with signs and symptoms such as confusion, altered mental status, or persistent chest pain. When the patient becomes unstable, immediate treatment is indicated.

In addition to areas of increased automaticity that can precipitate SVTs, reentry can also cause SVTs. Reentry describes a condition whereby a depolarization impulse is being propagated down a pathway in which some of the myocytes are still in the effective refractory period and a unidirectional block is present and preventing the impulse from traveling normally down this pathway. However, as the impulse travels around the area of the unidirectional block, the tissue allows the depolarization front to travel in the opposite (antidromic) direction, back to the initial point of entry into this pathway. This allows the depolarization wave front to restimulate the myocytes and

**BOX 11.1 Diagnostic and Therapeutic Approaches to Supraventricular Tachycardias**

**VAGAL MANEUVERS**
1. Carotid sinus massage
2. Pressure on the carotid sinus
3. Valsalva technique
4. Forced expiration of air against a closed glottis
5. Apneic facial exposure to cold water (“cold water diving reflex”)
6. Immersion of the face into cold water
7. Oculocardiac reflex
8. The trigemino-vagal reflex initiated by pressure on the eyeball

**PHARMACOLOGIC AGENTS**
1. Adenosine
2. Calcium channel blockers (verapamil, diltiazem)
3. β-Blockers, including esmolol
4. Digoxin
5. Amiodarone
6. Procainamide

**CARDIOVERSION**
Administration of a synchronized shock

The changes in electrochemical voltage during these events are depicted on the ECG in the usual sequential PQRST (the P wave indicates SA nodal depolarization, the PR interval denotes atrial depolarization followed by activation of the AV node, and the QRS complex summarizes electrical activity during ventricular depolarization) (Fig. 11.2).

The discharge rate of the SA node is usually modulated by a balance of input from the sympathetic and parasympathetic nerves (i.e., the autonomic nervous system). Sympathetic input to the heart is provided by the adrenergic nerves, which innervate the atria and ventricles, and by circulating hormones such as epinephrine and norepinephrine, which are released from the adrenal gland and cause the HR to increase. Parasympathetic input to the heart is provided by the vagus nerve (cranial nerve X) fibers. These nerve fibers innervate the SA and AV nodes. Vagal output to the SA node causes slowing of the HR by decreasing the depolarization rate of the “intrinsic pacemaker,” whereas vagal output to the AV node enhances nodal blockade of atrial depolarization impulses to the ventricles. Hence, vagal stimulation results in slowing of electrical activity, examples being termination of an SVT, slowing of the ventricular rate of AF (via the AV node), or simply producing a sinus bradycardia (via the SA node). Under normal physiologic circumstances, the HR is modulated to meet the metabolic needs of the body’s peripheral circulation. Changes in AV electrochemical events (i.e., rates and rhythms) are manifested as changes in the electrocardiographic intervals and waveforms.

As noted earlier, SVT rhythms can be either sinus (i.e., originating in the SA node: sinus tachycardia) or ectopic (i.e., originating in atrial myocytes above the ventricles). The rate of discharge of the SA node often varies as a result of various physiologic and pharmacologic stimuli, including fever, hypovolemia, shock, anemia, hypoxia, anxiety, pain, cocaine, and amphetamines. These conditions often require or precipitate increased blood flow and hence cardiac output (CO) to peripheral tissues. This increase in peripheral blood flow or CO is accomplished by an increase in heart rate (HR = CO × SV [stroke volume]). These are usually normal, benign physiologic responses to various stimuli or triggers. Direct treatment of the rate of discharge of the SA node is not generally necessary; however, determining and treating the cause of the sinus tachycardia usually eliminates the fast HR. Nonetheless, when single or multiple ectopic, spontaneously discharging foci develop in the atria or upper portions of the AV node, they can begin to “take over” or “override” the normal pacemaker activity in the heart (i.e., the SA node) and produce a rapid HR exceeding 100 beats/min. These foci may develop as a result of increased irritability or automaticity of atrial myocytes secondary to electrolyte abnormalities, hypoxia, pharmacologic agents, or atrial stretch caused by volumetric overload. If these foci are not treated or suppressed and the atrial depolarization rate proceeds to accelerate to rates greater than 150 beats/min (i.e., the heart is beating in excess of 2 beats/sec) and the impulses get through the AV node to the ventricles, the time for diastolic filling of the ventricles will be compromised and result in a precipitous drop in CO. This will ultimately cause a drop in CO regardless of the increase in HR. Furthermore, as CO begins to drop, mean arterial blood pressure (MAP) will decrease and cause hypoperfusion of the brain and other peripheral tissue (MAP is the product of CO times total peripheral resistance [TPR]: MAP = CO × TPR). Treatment of this tachycardia can be achieved pharmacologically by suppressing the automaticity of myocytes with medications (e.g., calcium channel blockers or β-blockers) and subsequently treating the underlying cause or causes, such as hypoxia, electrolytes, and the like. Decreasing the hemodynamic consequences of this arrhythmia requires increasing the “blocking” of these impulses from reaching the ventricles via the AV node. This can be done by enhancing vagal input to the AV node or by pharmacologic enhancement of AV blockade. Multiple rapid depolarizations of the atria, which are conducted to the ventricles, can ultimately have a bimodal type of response: a modest increase in HR will cause an increase in CO, whereas a massive increase in the atrial rate with a concomitant increase in the ventricular rate will cause a drop in CO. This can lead to an unstable patient with signs and symptoms such as confusion, altered mental status, or persistent chest pain. When the patient becomes unstable, immediate treatment is indicated.

In addition to areas of increased automaticity that can precipitate SVTs, reentry can also cause SVTs. Reentry describes a condition whereby a depolarization impulse is being propagated down a pathway in which some of the myocytes are still in the effective refractory period and a unidirectional block is present and preventing the impulse from traveling normally down this pathway. However, as the impulse travels around the area of the unidirectional block, the tissue allows the depolarization front to travel in the opposite (antidromic) direction, back to the initial point of entry into this pathway. This allows the depolarization wave front to restimulate the myocytes and
favoring reentry, and the hemodynamic consequences may be attenuated by enhancing AV nodal blockade of the ventricles (e.g., through vagal stimulation, medication), thus slowing the ventricular response to this condition. Termination of reentry can be accomplished by either pharmacologic modification

**Figure 11.1 Tachydysrhythmias.**

ECG AND MEMBRANE POTENTIAL OF VENTRICULAR CELLS

Myocardial cell

Extracellular fluid

Overshoot

+30 mV

−70 mV

1 mV

P

Q

R

S

T

ECG

Phase 0: Upstroke. Increased sodium conductance into myocytes
Phase 1: Early repolarization. Increased potassium conductance out of myocytes
Phase 2: Plateau. Calcium influx into myocytes/potassium efflux increasing
Phase 3: Repolarization. Sodium influx decreased, calcium influx decreased/potassium efflux still present
Phase 4: Steady state. Sodium, potassium, calcium conductance returns to resting membrane potential

Figure 11.2 Electrocardiographic and Membrane Potential of Ventricular Cells.

Figure 11.3 Cardiac Conduction in Supraventricular Tachycardia. A, Normal depolarization down path 1 and 2 that will “extinguish” or “cancel out” at point 3. Normal depolarization/repolarization and conductance. B, Reentrant pathway. 1: Normal conduction; 2: delayed/slowed conduction with unidirectional block; 3: normal conduction pathway.
of the myocytes to render them refractory to depolarization impulses for a longer period in a stable patient or by synchronized cardioversion to uniformly depolarize the myocytes and terminate the conditions favoring the SVT in an unstable patient.

Another situation to consider in the development and propagation of SVTs is the presence of preexcitation or an accessory pathway between the atria and the ventricles. Arrhythmias secondary to these causes can be managed with the use of appropriate pharmacologic agents to either suppress conduction through the accessory pathway or block AV nodal transmission without enhancing conduction through the accessory pathway.

To complete this discussion, we must also consider that there may be the possibility of an interventricular conduction delay (“aberrant conduction”) being present before the development of an SVT. If this is the case, the SVT may appear as a wide-complex tachycardia and can be confused with other dysrhythmias. However, an even more dangerous situation can occur if a wide-complex tachycardia of ventricular origin (VT) is present and is misdiagnosed as an SVT with aberrancy. As a result, the patient could be treated inappropriately with calcium channel blockers or β-blockers, resulting in vasodilation, loss of inotropy, and ultimately cardiac arrest. The safest course of action is to always assume that a regular wide-complex tachycardia is VT and treat with ventricular antiarrhythmia medications or electrical cardioversion. These therapies will generally result in successful cardioversion of the rhythm, regardless of whether the rhythm is actually VT or SVT with aberrancy.

The clinician must have a means of slowing down and sorting out these physiologic events so that appropriate diagnosis and treatment or intervention decisions can be made. With the application of vagal maneuvers, in some cases the activity of the atria and ventricles may be isolated enough to facilitate a correct diagnosis. An understanding of the underlying pathophysiology will guide appropriate treatment.

### Indications for Vagal Maneuvers

Vagal maneuvers are potentially useful in attempting to slow down or break an SVT. They are also indicated in settings in which slowing conduction in the SA or AV node could provide useful information (Box 11.2 and Figs. 11.4A-E and 11.5A-D). Such settings include patients with wide-complex tachycardia, in whom carotid sinus massage (CSM) aids in the distinction between SVT and VT. CSM can elucidate narrow-complex tachycardia in which the P waves are not visible, or aid in detection of suspected rate-related bundle branch block or pacemaker malfunction. After CSM, a wide-complex SVT may be converted to normal sinus rhythm, P waves may be revealed after increased AV node inhibition, or ventricular complexes may narrow as the ventricular rate slows. Because CSM slows atrial and not ventricular activity, AV dissociation may be seen more easily and is indicative of VT (see Fig. 11.4). In rapid AF or atrial flutter with a 2:1 block, either P waves or irregular ventricular activity with absent P waves may be revealed (see Figs. 11.5A and B). Sinus tachycardia may also be more apparent once P waves are unmasked by slowing the SA node (see Figs. 11.4C and D). Adenosine may be used for the same diagnostic purpose in these situations as well. In order of decreasing frequency, the electrocardiographic changes seen with CSM and vagal maneuvers are presented in Box 11.3.

### BOX 11.2 Potential Observations With Vagal Maneuvers in the Management of Tachydysrhythmias

1. Vagal maneuvers may slow the atrial rate in VT or complete heart block and may therefore demonstrate previously hidden P waves or obvious (AV) dissociation.
2. Abrupt changes in the heart rate without conversion are a result of increasing AV block.
3. Gradual slowing of the ventricular rate suggests the presence of sinus rhythm. Only rarely do vagal maneuvers decrease AV conduction in the presence of a sinus mechanism.
4. The dysrhythmias most likely to convert to sinus rhythm are PAT and paroxysmal nodal tachycardia.
5. Dysrhythmias that are associated with AV conduction defects (PAT with block, atrial flutter, and atrial fibrillation) infrequently convert to a sinus rhythm, but the ventricular rate slows. Rarely, atrial slowing will be sufficient to allow 1:1 AV conduction, which may actually increase the ventricular rate (see Fig. 11.6).

AV, Atrioventricular; PAT, paroxysmal atrial tachycardia; VT, ventricular tachycardia.

### BOX 11.3 Order of Decreasing Frequency of Electrocardiogram Changes With Vagal Maneuvers

1. Sinoatrial slowing, which occurs in approximately 75% of cases and leads to sinus arrest approximately 3% of the time.
2. Atrial conduction defects, as manifested by an increase in width of the P wave on the electrocardiogram.
3. Prolongation of the PR interval and higher degrees of atrioventricular block, which are seen in approximately 10% of cases.
5. Complete asystole, defined as sinus arrest without ventricular escape lasting longer than 3 seconds, which occurs in 4% of cases.
6. Premature ventricular contractions.

Vagal maneuvers, CSM in particular, may also be a useful aid in the diagnosis of syncope in the elderly. Some 14% to 45% of elderly patients referred for syncope are thought to have carotid sinus syndrome (CSS).2-4 CSS is defined as an asystolic pause longer than 3 seconds or a reduction in systolic blood pressure greater than 50 mm Hg in response to CSM (Fig. 11.6). Because it shares many characteristics with sick sinus syndrome, it has been suggested that both are manifestations of the same disease. CSS causes cerebral hypoperfusion, which can lead to dizziness and syncope. Analysis of patients with CSS indicates that it results from baroreflex-mediated bradycardia in 29%, hypotension in 37%, or both in 34%.3,6 Therefore syncope, chronic near-syncope, or a fall of unclear etiology in the elderly is an important indication for diagnostic CSM.3,6 Although the use of digoxin has been overshadowed by the use of other potentially less toxic agents such as calcium
channel blockers and β-blockers, the clinician can still prospectively simulate the cardioinhibitory effects of digoxin on a patient by performing vagal maneuvers. This can guide use and dosage of the medication before initiating treatment with digoxin. Significant slowing or block with CSM suggests a similar sensitivity to digoxin, and a smaller loading dose should be considered (Table 11.1).

Equipment and Setup
Before the initiation of any clinical intervention such as vagal maneuvers, administration of medication, or cardioversion for SVT, place the patient on a cardiac monitor, establish intravenous (IV) access, and infuse a slow, keep-vein-open (KVO; 60 mL/hr saline IV) solution through the IV line. Monitor the patient with a pulse oximeter and blood pressure monitor.
CSM is a bedside vagal maneuver involving digital pressure on the richly innervated carotid sinus (Fig. 11.7). The procedure is likely underused by clinicians but should be routinely considered as an initial intervention. It takes advantage of the accessible position of this baroreceptor for diagnostic and therapeutic purposes. Its main therapeutic application is for termination of SVTs caused by sudden paroxysmal atrial tachycardia. It also has diagnostic utility in the assessment of paroxysmal atrial tachycardia and other arrhythmias.

Keep antiarrhythmic medications readily available at the bedside. Keep a defibrillator/pacemaker at the bedside in anticipation of a worsening dysrhythmia. Administer oxygen for the procedure, especially if conscious sedation is anticipated. Place the patient in the Trendelenburg position if tolerated. Merely placing the patient in this position may terminate the SVT as a result of increased pressure on the carotids and maximum carotid bulb stimulation. This position may also prevent syncope if there is a significant decrease in blood pressure or HR.
tachydysrhythmias and rate-related bundle branch blocks. In addition, it can provide clues to latent digoxin toxicity, as described previously, by potentiating manifestations of the toxicity. It can also be used to sort out the differential diagnosis of syncope.

Returning to the use of CSM as a diagnostic technique for assessing digoxin toxicity, the adverse effects and toxicity from digoxin depend more on the response of the host than on the actual digoxin level. In cases of suspected digoxin toxicity, before the digoxin level is available or when it is in the normal range, CSM may be a useful diagnostic adjunct. Significant inhibition of AV node conduction associated with ventricular ectopy, especially ventricular bigeminy, should lead to suspicion of digoxin toxicity.9

### Contraindications

CSM is likely underused by clinicians but is contraindicated in the very rare patient likely to suffer neurologic or cardiovascular complications from the procedure. Patients with a carotid bruit should not undergo CSM because of the theoretical risk for carotid embolization or occlusion. A recent cerebral infarction is another contraindication because even a marginal reduction in cerebral blood flow may produce further infarction. Age, by itself, is not a contraindication to CSM. However, the elderly are more likely to have carotid artery disease and may experience transient and, very rarely, permanent neurologic or visual symptoms after CSM. Complications are thought to be due to transient cerebral ischemia or embolization of plaque, similar to a transient ischemic attack.

### Table 11.1 Ventricular Response to Carotid Sinus Massage and Other Vagal Maneuvers

<table>
<thead>
<tr>
<th>Type of Arrhythmia</th>
<th>Atrial Rate (beats/min)</th>
<th>Response to Carotid Sinus Massage and Release</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal sinus rhythm</td>
<td>60–100</td>
<td>Slowing with return to the former rate on release</td>
</tr>
<tr>
<td>Normal sinus bradycardia</td>
<td>&lt; 60</td>
<td>Slowing with return to the former rate on release</td>
</tr>
<tr>
<td>Normal sinus tachycardia</td>
<td>&gt; 100–180</td>
<td>Slowing with return to the former rate on release; appearance of diagnostic P waves</td>
</tr>
<tr>
<td>AV nodal reentry</td>
<td>150–250</td>
<td>Termination or no effect</td>
</tr>
<tr>
<td>Atrial flutter</td>
<td>250–350</td>
<td>Slowing with return to the former rate on release; increasing AV block; flutter persists</td>
</tr>
<tr>
<td>Atrial fibrillation</td>
<td>400–600</td>
<td>Slowing with persistence of a gross irregular rate on release; increasing AV block</td>
</tr>
<tr>
<td>Atrial tachycardia with block</td>
<td>150–250</td>
<td>Abrupt slowing with return to a normal sinus rhythm on release; tachycardia often persists</td>
</tr>
<tr>
<td>AV junctional rhythm</td>
<td>40–100</td>
<td>None; ± slowing</td>
</tr>
<tr>
<td>Reciprocal tachycardia using accessory (WPW) pathways</td>
<td>150–250</td>
<td>Abrupt slowing; termination or no effect; may unmask WPW</td>
</tr>
<tr>
<td>Nonparoxysmal AV junctional tachycardia</td>
<td>60–100</td>
<td>None; ± slowing</td>
</tr>
<tr>
<td>Ventricular tachycardia</td>
<td>60–100</td>
<td>None; may unmask AV dissociation</td>
</tr>
<tr>
<td>Atrial idioventricular rhythm</td>
<td>60–100</td>
<td>None</td>
</tr>
<tr>
<td>Ventricular flutter</td>
<td>60–100</td>
<td>None</td>
</tr>
<tr>
<td>Ventricular fibrillation</td>
<td>60–100</td>
<td>None</td>
</tr>
<tr>
<td>First-degree AV block</td>
<td>60–100</td>
<td>Gradual slowing caused by sinus slowing; return to the former rate on release</td>
</tr>
<tr>
<td>Second-degree AV block (I)</td>
<td>60–100</td>
<td>Sinus slows with an increase in block; return to the former rate on release</td>
</tr>
<tr>
<td>Second-degree AV block (II)</td>
<td>60–100</td>
<td>Slowing</td>
</tr>
<tr>
<td>Third-degree AV block</td>
<td>60–100</td>
<td>None</td>
</tr>
<tr>
<td>Right bundle branch block</td>
<td>60–100</td>
<td>Slowing with return to the former rate on release</td>
</tr>
<tr>
<td>Left bundle branch block</td>
<td>60–100</td>
<td>Slowing with return to the former rate on release</td>
</tr>
<tr>
<td>Digitalis toxicity–induced arrhythmias</td>
<td>Variable</td>
<td>Do not attempt CSM</td>
</tr>
</tbody>
</table>

AV, Atrioventricular; CSM, carotid sinus massage; WPW, Wolff-Parkinson-White (syndrome).

The presence of diffuse, advanced coronary atherosclerosis is associated with increased sensitivity of the carotid sinus reflex. This hypersensitivity is further augmented during an anginal attack or acute myocardial infarction. Brown and coworkers found that the degree of carotid sinus hypersensitivity was directly proportional to the severity of coronary artery disease as documented by cardiac catheterization. Patients with acute myocardial ischemia or with recent myocardial infarction are already at higher risk for VT or ventricular fibrillation (VF). A CSM-induced prolonged asystole may further predispose them to these dysrhythmias. Therefore CSM should be avoided in these patients.

Both digoxin and CSM act through a vagal mechanism to inhibit the AV node. Patients taking digoxin may experience greater inhibition of the AV node with a longer AV block as a result. Patients with apparent manifestations of digoxin toxicity or known digoxin toxicity should not undergo CSM because the AV inhibition may be profound.

Technique
This technique can be performed with or without a concomitant Valsalva maneuver. Alternatively, pressure can be applied to the abdomen by an assistant. Some clinicians prefer to place the patient supine or with the head of the bed tilted downward. Passively raising the supine patient’s legs is an additional maneuver to be used with the Valsalva maneuver (Fig. 11.8). The use of both a Valsalva maneuver and supine position/leg raise are suggested as routine techniques. Begin CSM on the patient’s right carotid bulb because some investigators have found a greater cardioinhibitory effect on this side. However, scientific agreement on this issue is not unanimous. Simultaneous bilateral CSM is absolutely contraindicated because the cerebral circulation may be severely compromised.

Before attempting CSM, first auscultate for carotid bruits on both sides of the neck (Fig. 11.9, step 2). The presence of a bruit is a contraindication to massage.

Keep the patient relaxed for two reasons. A tense platysma muscle makes palpation of the carotid sinus difficult, and an anxious patient will be less sensitive to CSM as a result of heightened sympathetic tone.

Tilt the supine patient’s head backward and slightly to the opposite side. Passively raise the legs. Palpate the carotid artery just below the angle of the mandible at the upper level of the thyroid cartilage and anterior to the sternocleidomastoid muscle (see Figs. 11.7 and 11.9, step 3). Once the pulsation is identified, use the tips of the index and middle fingers to administer CSM for 5 seconds in a posteromedial direction, aiming toward the vertebral column. Although earlier practitioners used a longer duration of massage, a shorter period minimizes the risk for complications and is adequate for diagnostic purposes in the majority of patients. Pressure on the carotid sinus may be steady or undulating in intensity; the force, however, must not occlude the carotid artery. The temporal artery may be simultaneously palpated to ensure that the carotid remains patent throughout the procedure.

If unsuccessful, repeat CSM after 1 minute. If the procedure is still unsuccessful, massage the opposite carotid sinus in a similar fashion. If not already performed, use simultaneous Valsalva maneuvers with the patient in the head-down position/leg raise to enhance carotid sinus sensitivity before the technique is abandoned (see Fig. 11.9, step 4). CSM may be repeated once antiarrhythmic medications (e.g., calcium channel blockers and β-blockers) have been given, and often the combination is more effective. However, repetition of CSM after the administration of adenosine is not thought to have any utility.

Complications
Neurologic complications of CSM are rare and usually transient. In a review of neurologic complications in elderly patients undergoing the procedure, Munro and associates found seven complications in a total of 5000 massage episodes, an incidence of 0.14%. Reported deficits included weakness in five cases and visual field loss in two others. In one case the visual field loss was permanent. Patients in this study were excluded from CSM if they had a carotid bruit, recent cerebral infarction, recent myocardial infarction, or a history of VT or VF. The duration of massage was 5 seconds. Lown and Levine described one patient with brief facial weakness during several thousand tests. Carotid emboli and hypotension have both been implicated as possible causes of the neurologic deficits. Unintentional
occlusion of the carotid artery may also be responsible for some neurologic complications.

Cardiac complications include asystole, VT, or VF. A normal pause of less than 3 seconds is part of the physiologic response to CSM; a longer pause may be diagnostic of CSS (see Fig. 11.6). In a review of reported cases of ventricular tachydysrhythmia, five cases were described. All five patients were receiving digoxin, and in several cases VT or VF followed AV block. Digoxin is associated with more prolonged AV block from CSM, which perhaps leaves these patients more vulnerable.

Valsalva Maneuver

In general, mean changes in bradycardia are greatest with the Valsalva maneuver and the diving response. During the Valsalva maneuver (i.e., exhaling against a closed glottis or bearing down as though to defecate), intrathoracic pressure increases and leads to increased arterial pressure as a result of increased afterload. It is easily done by having the patient take a deep breath, put their thumb in their mouth with closed lips, and attempt to exhale without expelling any air. This increased pressure is transferred to the peripheral vascular system. Venous return to the heart is decreased, which results in a decrease in the SVT. This is followed by increased venous pressure. All these changes in pressure lead to an initial increase in HR and carotid sinus pressure. As the maneuver is sustained, vagal tone is increased, thereby leading to a compensatory decrease in SA and AV conduction. This is the expected or desired diagnostic or therapeutic response. There is no reason not to routinely perform a Valsalva maneuver/leg raise with CSM.

Contraindications

Patients must be able to cooperate with the clinician’s commands. Dyspneic or tachyphneic patients may not be able to hold their breath for the period needed to complete the maneuver.

Technique

The patient should be supine, with a cardiac monitor in place, IV access secured, antiarrhythmics available, and defibrillation available. Ask the patient to take a deep breath and hold it, or attempt to blow it out against their thumb in their mouth.
encircled by closed lips. Instruct the patient to bear down and try to exhale without allowing air to leave the lungs. Passively raise the patient’s legs (see Fig. 11.8). Ask the patient to try and hold this position for 10 to 20 seconds. An adjunctive method is to have the patient take and hold a deep breath and try to push against the clinician’s hand with the abdomen while the clinician gently pushes on the anterior wall of the abdomen. Then perform CSM, first on the right side for 3 seconds. Perform CSM on the left side if this is not successful.

Apneic Facial Exposure to Cold (Diving Reflex, or Diving Bradycardia)

**Technique**

This technique can be viewed as a variation on the simple Valsalva maneuver. It has been found to be useful in children who may be unable to cooperate or may be incapable of performing a Valsalva maneuver. Classically, the technique consists of covering the face with a bag of crushed ice and cold water (0°C to 15°C) for 15 to 30 seconds and then observing the ECG for a break in the tachycardia. Another variation of this technique is to drip ice water into the nostril of a small child. The procedure is based on the classic diving reflex of bradycardia. Slowing the SVT to unmask the hidden, underlying rhythm is similar to the effects of CSM. Conversion of sudden atrial tachycardia to sinus rhythm should be observed in 15 to 35 seconds. The procedure is convenient and noninvasive and can be self-administered.

Berk and colleagues demonstrated in healthy volunteers that immersion of the face in cold water and the Valsalva maneuver can produce a greater vagal response than CSM. Lim and associates in 1998 and Mehta and coworkers in 1988 also found that the Valsalva maneuver was more effective than CSM for conversion of induced SVT.

Another technique that was used, but has fallen out of favor, is direct ocular pressure. There are many contraindications to this technique, such as retinal or lens surgery, glaucoma, thrombotic-related eye conditions, and penetrating or recent blunt trauma to the eye. This procedure is no longer recommended.

**Selected Pharmacologic Agents**

(Fig. 11.10 and Box 11.4)

After unsuccessful CSM, a pharmacologic approach to SVT is preferred in stable patients. In the presence of severe hypotension, chest pain, or other evidence of extremis, cardioversion...
Adenosine depresses the AV node and sinus node activity. Adenosine is indicated for the conversion of PSVT associated with or without accessory tract bypass conduction (WPW, Lown-Ganong-Levine [LGL]). The other use of adenosine is for diagnostic slowing of SVT to unmask AF, atrial flutter, or VT. The diagnostic and therapeutic effects of adenosine on tachydysrhythmias are similar to those elicited by vagal maneuvers. Adenosine's safety is derived from its short duration of action, usually approximately 10 to 12 seconds.

Adenosine should not be used in patients with a known history of second- or third-degree AV block or sick sinus syndrome unless there is a functioning internal pacer. Additionally, if the patient has a known hypersensitivity to adenosine or a history of severe reactive airway disease or active wheezing, the drug should not be used. In addition, adenosine should not be used in patients with an underlying accessory pathway (WPW, LGL) in the setting of AF. In this circumstance, the HR may increase because the enhanced AV node blockade permits conduction through the bypass tract.

**Dosage**

The initial dose recommended is a 6-mg rapid bolus administered over a period of 1 to 3 seconds. The dose should be immediately followed by a 20-mL saline flush (Fig. 11.11). If no response occurs within 1 to 2 minutes, a 12-mg dose should be administered in the same manner as the initial dose. This second dose should also be followed by a 20-mL saline bolus.

Side effects of adenosine are common and transient. Many patients experience a rather unsettling feeling, and this should be the preferred intervention. In unusual circumstances, such as rapid AF in patients with known Wolf-Parkinson-White (WPW) syndrome, certain medications should be used with caution, and cardioversion may be considered the first-line intervention.

**Adenosine**

The use of vagal maneuvers has been eclipsed in recent years by the use of adenosine, an endogenous, ultrashort-acting, vagal-stimulating purine nucleoside that is ubiquitous in body cells. Its action is to slow conduction time through the AV node and depress the AV node. If vagal maneuvers have been attempted and failed to produce the desired response, use of adenosine is an appropriate subsequent intervention.

Extracellular adenosine is cleared rapidly from the circulation by the erythrocyte and vascular endothelium system that transports adenosine intracellularly. Here, rapid metabolism via a phosphorylation or deamination cycle produces inosine or adenosine monophosphate. Adenosine produces a short-lived pharmaceutical response because it is metabolized rapidly by the described enzymatic degradation. The half-life of adenosine is less than 10 seconds, with the metabolites becoming incorporated into the high-energy phosphate pool.

**Indications and Contraindications**

Most forms of paroxysmal supraventricular tachycardia (PSVT) affect a reentry pathway involving the AV node, and adenosine

---

**Figure 11.11 Treatment of Supraventricular Tachycardia (SVT) With Adenosine.**

A, Because of its ultrashort half-life (< 10 seconds), adenosine must be given rapidly, followed by a 20-mL saline flush. B, Adenosine slows conduction through the atrioventricular node, thereby effectively converting SVT (left side of the screen) to sinus rhythm (right side of the screen).
be explained to the patient before administering the drug. Common sensations include flushing, dyspnea, and chest pain. Important drug interactions include theophylline or related methylxanthines (caffeine and theobromine), which can block adenosine receptor sites. If these medications are being taken by the patient, administer a larger dose of adenosine. If the patient is taking dipryidamole or carbamazepine, these drugs may block uptake of adenosine and potentiate its effects, so contemplate administering a smaller IV dose of adenosine (e.g., 3 mg). Adenosine is safe and effective in pregnancy. Also, if the patient has a central line or a transplanted heart, try an initial 3-mg dose.

**Procaainamide**
A time-honored antiarrhythmic, procaainamide slows conduction and decreases the automaticity and excitability of atrial, ventricular, and Purkinje tissue. It also increases refractoriness in atrial and ventricular tissue. Procaainamide prolongs the QT interval without having much effect on Purkinje fibers or ventricular tissue.

**Indications and Contraindications**
A long-established clinical application is for management of the rate of SVT, SVT with aberrant conduction (wide-complex SVT), AF or atrial flutter associated with WPW conduction, and VT. The advantage of using procaainamide is the ability to convert to the oral form when rate control is achieved. The dose of procaainamide recommended by advanced cardiac life support is usually 20 mg/min, although in urgent situations up to 50 mg/min can be used. Procaainamide is generally used in clinical situations in which time is not a factor in patient care. Long-term management in the emergency department necessitates monitoring of the plasma concentrations of procaainamide and its N-acetylprocaainamide metabolite. Hypotension and conduction disturbances (torsades de pointes, heart block, and sinus node dysfunction) are often signs of high plasma levels. Use caution in patients with a history of hypokalemia, long QT intervals, and torsades de pointes. Hematologic and rheumatologic disturbances are factors in long-term use. The endpoint of administration of the drug is when the arrhythmia is suppressed, hypotension occurs, the QT duration increases by 50% over baseline, or a maximum of 17 mg/kg of the drug has been administered (1.2 g in a 70-kg adult).

**Amiodarone**
Amiodarone has become one of the workhorses for treatment of dysrhythmias in the emergency department. It is often considered a Vaughan-Williams class III drug because it is a potassium channel blocker. However, this medication also blocks sodium and calcium channels and α- and β-adrenergic receptors. As a result of its potassium-blocking properties, amiodarone prolongs the action potential duration and increases refractoriness of the atrial and ventricular tissue, the sinus and AV nodal tissue, and Purkinje fibers. Amiodarone also blocks sodium channels in depolarized tissue. It slows depolarization in the SA node and slows conduction through the AV node. Its calcium antagonist effect is minimal.

**Indications and Contraindications**
Amiodarone is used for the control of narrow-complex supraventricular and ventricular dysrhythmias. It is useful in the management of narrow-complex tachycardias that originate from a reentry rhythm (SVT). It is effective in the conversion of stable wide-complex tachycardias and is useful in managing polymorphic VT with a normal QT interval. Amiodarone can be used for wide-complex tachycardias of undetermined origin. This drug can also be used for the management of AF and atrial flutter with aberrancy, SVT with accessory pathway conduction, and the rare adult junctional tachycardia. Another use for this medication is control of the rapid ventricular rate as a result of accessory pathway conduction in preexcited atrial arrhythmias. It is a strong second-line choice with procaainamide for hemodynamically stable VT.

Its use can precipitate heart failure, hypotension, and severe bradycardia. When used with β-blockers and calcium channel blockers, amiodarone can have the added risk of hypotension and bradycardia. Torsades de pointes has been reported after the use of amiodarone in conjunction with drugs that have increased the QT interval. In addition, amiodarone should not be used in the presence of AF with accessory pathways; there are multiple case reports of patients decompensating after receiving IV amiodarone when it was used for rapid AF plus WPW syndrome.

In the setting of possible AF with an accessory pathway, procaainamide is safer. Finally, note that amiodarone will also prolong the QT interval, so it is best to avoid this drug in patients with a preexistent long QT interval, just like procaainamide.

**Dosage**
The IV dosage is 150 mg administered over a 10-minute period. Follow this with a 1-mg/min infusion for 6 hours and then a 0.5-mg/min maintenance infusion over an 18-hour period. If a dysrhythmia is refractory or resistant, 150 mg can be repeated every 10 minutes to a maximum of 2.2 g/24 hr. The major adverse effects of amiodarone are bradycardia and hypotension.

**β-Adrenergic Blockade**
β-Blockers are very useful agents for control of the ventricular response in patients with PSVT, AF or atrial flutter, and atrial tachycardia. No β-blocker offers a distinctive advantage over another because when used clinically, they can all be titrated to a desired effect on dysrhythmias and hypertension. Examples of β-blockers are atenolol, metoprolol, propranolol, and esmolol. Recently sotalol IV has been added to the AHA 2015 Adult Tachycardia with a pulse algorithm (more information later). Separating the different drugs and their use is the various pharmacologic characteristics that control adverse reactions, speed of onset, dosage regimens, contraindications, and drug interactions.

The electrophysiologic effect of β-blockers results from inhibition of binding of catecholamine at β-receptor sites. These medications reduce the effects of circulating catecholamines and this is manifested as a decrease in HR, blood pressure, and myocardial contractility. The PR interval may be prolonged, but the QRS and QT intervals are not affected. Their actions are most noted on cells that are most stimulated by adrenergic actions. Typically, these sites are the sinus node, the Purkinje fibers, and ventricular tissue when it is stimulated by catecholamines. These medications also have various cardioprotective effects in patients suffering from acute coronary syndromes. They exert their cardioprotective effects by decreasing myocardial workload, and hence they decrease myocardial oxygen consumption and demand. β-Blockers are useful in the treatment of narrow-complex tachycardias that originate secondary to a reentry phenomenon or an automatic focus (MAT, an ectopic pacemaker, or a junctional rhythm). These drugs can also be used to control rates in patients suffering from...
AF or atrial flutter, as long as ventricular function is nominal. Some representative dosages of these β-blockers are: (1) atenolol (β1), 5 mg IV slowly over a period of 5 minutes; if no effect, repeat in 10 minutes; (2) metoprolol (β1), 5 mg IV slowly, may repeat up to 15 mg total; and (3) propranolol, 0.1 mg/kg IV by slow push and divided into three equal doses at 2- to 3-minute intervals; the total dose may be repeated in 2 minutes. The administration rate of the drug should not exceed 1 mg/min.

In general, β-blockers should not be used in patients with diabetes, lung disease, bradycardia, or heart block; in conjunction with a calcium channel blocker; in patients with hypotension; or in the presence of a vasospastic condition. β-Blockers should also not be used in patients with AF in the presence of bypass tracts, as is true for the calcium channel blockers, adenosine and amiodarone.

Propranolol

Propranolol is the representative drug of the β-adrenergic blockade agents. It is nonselective and has β1 and β2 effects on the heart, which allows it to be used to control rapid ventricular rates. Rate slowing is caused by: (1) slowing of impulse formation in the SA node, and (2) depression of myocardial contractility. The usual effects on the ECG are rate reduction and prolongation of the PR interval. The QRS and QT intervals are not affected. Because it is relatively nonselective (it has effects on both β1 and β2 receptors), its contraindications are extensive.30,42,43

Sotalol

Sotalol is a nonselective competitive β-adrenergic receptor blocker which manifests class III antiarrhythmic activity. This drug is now used to mitigate life-threatening VT and VF. Its utility is providing the clinician with the ability to control symptomatic AF or atrial flutter. Use of the drug can be considered if the Valsalva maneuver does not resolve the AF or atrial flutter.

This medication should NOT be used in patients with significant bradycardia (HR < 50 beats/min), sick sinus syndrome, prolonged QT syndrome, cardiogenic shock, uncontrolled congestive heart failure, or those with respiratory issues such as bronchospasm or asthma. Also avoid its use in higher level blocks such as a second- or third-degree heart block. It can be used cautiously if the patient has a pacemaker.

Sotalol is excreted via the kidneys, therefore the clinician needs to be aware of the patient’s creatinine clearance (> 40 mL/min) if this medication is to be considered.

Use caution when using this medication with a breastfeeding female as sotalol passes into the breast milk.

If possible, avoid using this drug in combination with calcium channel blockers, catecholamine-depleting medications, medications used to treat diabetes, β2-adrenergic agonists, and clonidine.

Some side effects of this medication are light-headedness, weakness, headache, fatigue, dizziness, and dyspnea. In addition, this drug can precipitate significant bradycardia. Due to the QT prolongation this drug produces, there is a potential proclivity for the development of torsades de pointes.

This medication blocks the β receptors, and prevents activation of the G-protein complex. This leads to a dearth in the production of cyclic adenosine monophosphate. As a result, normal calcium metabolism is derailed in the myocytes, causing a slowing of the HR and a decrease in the inotropic state of the heart, thus decreasing force of contraction and potentially SV. In addition, this medication blocks potassium channels in the myocardium and prolongs the refractoriness of the heart to another excitatory impulse, potentially quelling the tachycardia.

Dosage. The dosing of sotalol is quite simple. The American Heart Association (AHA) recommends that 100 mg IV (1.5 mg/kg) be administered over 5 minutes.

The following drugs and medications are available and can be used at the discretion of the physician, depending on the clinical scenario and expert consultation. These medications are available and still in use by many clinicians.

The aforementioned medications are 2015 AHA recommendations.

Esmolol

Esmolol is a rapid-, short-acting, β1-selective (cardioselective) β-blocker. At therapeutic doses, it inhibits β1 receptors in cardiac muscle. At higher doses, its selectivity is lost and it affects β2 receptors in the lung and vascular system. Esmolol is rapidly metabolized in erythrocytes and has a half-life of approximately 2 to 9 minutes. Its elimination half-life is approximately 9 minutes.31,35

The complicated dosing regimen of esmolol makes it more difficult to use.

Indications and Contraindications. Esmolol is indicated for the rapid conversion of SVT and rapid control of the ventricular rate in patients with non–preexcited AF or atrial flutter. In addition, it can be used to control the rate of uncompensated sinus tachycardia when the clinician believes that the tachycardia requires slowing. It has also proved beneficial as adjunctive therapy for the VT of torsades de pointes.31,33,35,42 Esmolol should not be used in patients with second- or third-degree heart block or in frank heart failure. Like all β-blockers, exercise care when using this drug in patients with bronchospastic disease and diabetes.

Dosage. Esmolol has a complicated dose regimen. First, give a loading dose of 0.5 mg/kg over the first 1 minute. Follow this with a maintenance infusion of 50 μg/kg per minute over a 4-minute period. If this is not successful, administer a second bolus dose of 0.5 mg/kg followed by a maintenance infusion of 100 μg/kg over a 4-minute period. The bolus/maintenance dose can be repeated up to a maximum infusion rate of 300 μg/kg per minute for 4 minutes.31,33,42 Similar dosing has been recommended for children: a 100- to 200-μg/kg maintenance rate between 100-μg/kg increases in bolus doses.31

Calcium Channel Blockers

Diltiazem

Diltiazem is a nondihydropyridine calcium channel blocker. It controls the rate of influx of calcium into myocytes during depolarization. This calcium channel blocker slows conduction of impulses through the AV node and prolongs the refractory period of the AV node. As a result, this drug is capable of terminating reentry-based tachycardias that have not converted with the use of adenosine or vagal maneuvers, and it can be used to control the ventricular response rate in a variety of SVTs (AF, atrial flutter). In addition, diltiazem can be used for the treatment of stable, narrow-complex tachycardias that are driven by automaticity (e.g., ectopic, multifocal, or junctional tachycardias). Its effects on AV nodal tissue are selective in
that it reduces AV conduction in tissue responsible for the tachydysrhythmia but spares normal conduction tissue.\textsuperscript{11,35,42,44}

**Indications and Contraindications.** The beneficial effects of diltiazem are: (1) ventricular slowing of rapid AF or atrial flutter without accessory bypass conduction, and (2) rapid conversion of narrow-complex PSVT to sinus rhythm.\textsuperscript{10,11,35,44,45}

Diltiazem is contraindicated in the following settings: (1) sick sinus syndrome, second-degree block, and third-degree block, except in the presence of an internal pacer; (2) severe hypotension or cardiogenic shock; (3) hypersensitivity to diltiazem; (4) use of IV \( \beta \)-blockade within a few hours of needing to use diltiazem; (5) AF or atrial flutter with coexisting accessory bypass tract conduction (WPW, LGL); and (6) VT.

**Dosage.** Give an initial dose of 0.25 mg/kg followed by a repeated dose of 0.35 mg/kg. Start a maintenance infusion of 5 to 15 mg/hr.\textsuperscript{11,35,42,44} Consider pretreating with IV calcium if the patient is hypotensive (see later).

**Verapamil**

Verapamil is also a calcium channel blocker. This medication blocks the slow channel for entry of calcium into myocytes. Verapamil blocks not only the calcium channels in the specialized conduction tissue of the myocardium but also the contracting cells of the heart. As a result, verapamil prolongs the effective refractory period within the AV node and slows conduction.\textsuperscript{17,62} It also has a modest effect on myocardial contractility.\textsuperscript{17}

**Indications and Contraindications.** Verapamil is effective in: (1) converting narrow-complex PSVT to normal sinus rhythm, and (2) controlling the ventricular response in AF or atrial flutter if the AF or atrial flutter is not complicated by the presence of an accessory bypass tract (WPW, LGL). With specific regard to WPW syndrome and rapid AF, caution is advised with the use of verapamil. However, verapamil has been reported to be safe in those with overt or concealed accessory bypass tract conduction (WPW, LGL); and (6) VT.

Verapamil should not be used or be used with caution in the following settings: (1) PSVT with accessory bypass tract conduction, (2) AF or atrial flutter with accessory bypass tract conduction (WPW syndrome), (3) coexistence of sick sinus syndrome or second- or third-degree AV block unless an internal pacer is present, (4) severe left ventricular dysfunction (systolic blood pressure < 90 mm Hg) or cardiogenic shock, and (5) patients with known verapamil hypersensitivity.\textsuperscript{13,35,42,44}

Because of its prolonged activity, verapamil should be used with caution in patients with congestive heart failure.

In the presence of SVT, hypotension may be caused by the negative inotropic and vasodilating effects of verapamil. Administration of calcium before IV verapamil results in a decreased incidence of hypotension without compromising the effectiveness of channel blockers. The most common adverse effect of IV calcium is flushing. Use of digoxin does not contraindicate calcium pretreatment. A pre-verapamil dose of calcium gluconate, 1 g (ionized calcium, 90 mg) administered over a period of 3 minutes, is recommended for preventing or lessening the hypotensive effect of verapamil without affecting its antiarrhythmic effects.\textsuperscript{37}

**Dosage.** Administer 2.5 to 5 mg IV over a 1- to 2-minute period. It may be repeated every 10 minutes to a maximum of 15 to 20 mg total. Pretreatment with calcium (calcium gluconate, 1 g IV over a period of 2 to 3 minutes) is suggested in hypotensive patients.

**Digoxin**

Digoxin was a time-honored drug used for the treatment of AF and atrial flutter. It is the only antidysrhythmic with inotropic characteristics. Digoxin is less useful for the emergency clinician because of its long delay of onset.

Digoxin is a cardiac glycoside found in a number of plants. It is extracted from the leaves of *Digitalis lanata*. Digoxin increases intracellular Na\(^+\) and K\(^+\) by inhibiting sodium-potassium adenosine triphosphatase, the enzyme that regulates the quantity of Na\(^+\) and K\(^+\) inside the cell. An intracellular increase in Na\(^+\) stimulates Na\(^+\)-Ca\(^2+\) exchange, which leads to increased intracellular Ca\(^2+\). Digoxin acts both directly on cardiac muscle and indirectly on the cardiovascular system. The indirect effects are mediated by the autonomic nervous system. The results of these actions are vagomimetic effects on the SA node and the AV node.

The consequences of these actions are: (1) increased force and velocity of myocardial contraction (positive inotropic effect), (2) slowing of HR and AV nodal conduction (vagomimetic effect), and (3) a decrease in symptomatic nervous system effects (neurohormonal-deactivating effect).\textsuperscript{29,30,46–51}

**Indications and Contraindications.** Although its use in controlling the ventricular response rate in chronic AF is well established, digoxin is no longer the mainstay of therapy for narrow-complex tachycardias, as it has been replaced by newer agents. Its inotropic character is still widely used in the setting of heart failure.

Use of digoxin should be avoided in the clinical settings of sinus node disease and AV blockade. It may cause complete heart block or severe bradycardia. Do not use digoxin in patients with accessory bypass tract rhythms (WPW or LGL). It may cause a rapid ventricular response or VF. Patients with idiopathic hypertrophic subaortic stenosis, restrictive cardiomyopathy, constrictive pericarditis, or amyloid heart disease are particularly susceptible to digoxin toxicity.\textsuperscript{12}

**Dosage**

Give an IV loading dose of 10 \( \mu \)g/kg to 15 \( \mu \)g/kg, followed by individual parenteral dosing until the desired rate is achieved.\textsuperscript{29,31,35,42,53–56}

**Electrical Cardioversion**

Because it is more complicated, electrical cardioversion is usually considered after failure of medications, but it is successful in the majority of SVTs. In life-threatening or unstable situations, patients in AF are to be immediately cardioverted because the risk for continued AF outweighs the risk for thromboembolism.\textsuperscript{35,57–65} Cardioversion is specifically indicated when the patient is unstable; that is, a change in mental status occurs, the patient becomes hypotensive, ischemic chest pain develops, heart failure develops, or the ventricular rate exceeds 140 to 150 beats/min.\textsuperscript{42} Urgent restoration of normal rhythm in patients with symptomatic new-onset AF is best achieved by direct cardioversion with either a monophasic or a biphasic defibrillator. Success rates with biphasic defibrillators are reported to be approximately 94% to 95%.\textsuperscript{66–69} The cardioversion procedure is discussed in greater detail in Chapter 12. A brief synopsis, algorithm, and flow chart is included (see Fig. 12.18) to facilitate cardioversion.
Current guidelines for the treatment of symptomatic new-onset AF focus on the length of time that the patient has been in AF or atrial flutter. This is the determining factor for the initiation of anticoagulation when confronted by the need for cardioversion to sinus rhythm. Accordingly, onset within 48 hours or less has been determined to be the time limit that a patient with new-onset AF can undergo cardioversion without the need for anticoagulation. Studies have shown that staying under the 48-hour limit allows cardioversion to occur with the lowest risk for thromboembolism.1,58,61,72 Patients who have been in AF for longer than 48 hours and are not in need of urgent care need to undergo anticoagulation to an international normalized ratio (INR) of 2.0 to 3.0 for a 3-week duration before cardioversion.43 If this approach is not clinically acceptable, transesophageal echocardiography (TEE) should be performed in addition to heparinization.

If no clot in the left atrial appendage is visualized on TEE, the heparinized patient should immediately undergo cardioversion and take anticoagulants for the next 4 weeks. If a clot is visualized in the left atrial appendage, the patient should first be anticoagulated to an INR of 2.0 to 3.0 for 3 weeks before cardioversion3,4,7,73–77 (Boxes 11.5 and 11.6).

An alternative treatment strategy with a reported success rate of 50% to 70% is ibutilide in a bolus IV infusion. Be cautious when using ibutilide in patients with prolonged QT syndrome.
remain an important diagnostic tool and are routinely used prior to medications. Vagal maneuvers are especially important in unmasking the underlying rhythms of narrow-complex tachydysrhythmias and in determining the presence of CSS in patients with syncope.

The advent and availability of medications that quickly and safely control the rate of tachydysrhythmias has given the emergency clinician a more varied and powerful armamentarium to be used in cardioverting these life-threatening dysrhythmias to normal sinus rhythm.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


INTRODUCTION

Defibrillation is an emergency procedure performed to terminate ventricular fibrillation (VF) (Fig. 12.1A). VF is a potentially lethal, but survivable, rhythm commonly found in victims of sudden cardiac arrest (SCA).1,2 VF can be caused by myocardial infarction, myocardial ischemia, undiagnosed coronary artery disease, and electrical injuries. Medications such as tricyclic antidepressants, digitalis, quinidine, and other proarrhythmics can cause QT-segment prolongation and changes in the refractory period of the cardiac cycle that are capable of precipitating VF. Furthermore, chest trauma, hypothermia, cardiomyopathy, electrolyte disturbances, and various toxicidromes can induce conditions favoring the development of VF. Hypoxia is another culprit that frequently precipitates VF in adults and the pediatric population. Congenital malformations of the heart and great vessels have also been associated with an increased incidence of VF in young children. The most effective treatment of VF in its early phase is defibrillation.3 It can also be used to terminate pulseless ventricular tachycardia (VT) (see Fig. 12.1B). Patients with VF or pulseless VT are unresponsive, pulseless, and apneic. These patients sometimes require appropriate integration of cardiopulmonary resuscitation (CPR) with defibrillation to establish the return of spontaneous circulation (ROSC). Other dysrhythmias may also be encountered in patients with SCA, such as pulseless electrical activity (PEA) and even asystole; however, in this chapter the discussion is limited to the treatment of VF and pulseless VT.

Defibrillation entails passing a therapeutic burst of electrical current across the chest wall through the myocardium for the purpose of terminating the chaotic electromechanical activity that is impeding the ventricles from ejecting blood into the circulation (Fig. 12.2). Failure to recognize and terminate VF promptly makes suppression of VF via defibrillation more difficult.4 For every minute that the heart is in VF without treatment, the potential for the initial defibrillation to be successful and for the victim of SCA to survive decreases by 7% to 10%.5 However, the integration of CPR with defibrillation, when appropriate, increases the chance for successful defibrillation and survival from SCA.6

Cardioversion is performed to suppress dysrhythmias that produce a rapid pulse and cause the patient to become unstable; such dysrhythmias include supraventricular tachycardia (SVT), atrial fibrillation (AF), atrial flutter, and unstable monomorphic VT (Fig. 12.3). These patients do have a pulse, albeit weak, but can rapidly decompensate, become hypotensive, experience chest pain, or have a change in mental status that will require rapid intervention (i.e., cardioversion). CPR is obviously not indicated because these patients have a pulse and their peripheral tissues are being perfused. Cardioversion is very similar to defibrillation; however, the shock is administered during the

---

Review Box 12.1 Defibrillation and cardioversion: indications, contraindications, and equipment.

- **Indications**
  - **Defibrillation**
    - Ventricular fibrillation
    - Pulseless ventricular tachycardia
  - **Cardioversion (usually reserved for unstable rhythms)**
    - Ventricular tachycardia with a pulse
    - Supraventricular tachycardia
    - Atrial fibrillation
    - Atrial flutter

- **Contraindications**
  - **Defibrillation**
    - Presence of a pulse
    - Asystole or pulseless electrical activity
    - Obvious signs of death
    - Valid do-not-resuscitate order
  - **Cardioversion**
    - Arrhythmias due to digitalis toxicity
    - Sinus tachycardia

- **Complications**
  - Chest wall burns
  - Shock of a health care worker
  - Myocardial tissue injury

---
are altered. Recommendations from the American Heart Association (AHA) are considered the most reasonable guidelines for the clinician, but many of the principles and caveats are based on minimal data, can be contradictory, and are subject to change; more importantly, any guideline is best applied by considering a specific clinical scenario. Most recently, cardiac resuscitation has been reviewed and new AHA guidelines were released in 2010. On the basis of the strength of the evidence available, the AHA developed recommendations to support the interventions that showed the most promise. The new algorithms reflect alterations in the sequence of actions to be performed and stress high-quality CPR with compressions of adequate rate and depth that allow complete chest recoil after compressions, minimize interruptions in chest compressions, and avoid excessive overventilation. These modifications stress the interposition of effective CPR (Fig. 12.4) with defibrillation and have been organized in such a way that the time until the first shock is minimized and time to initiation of effective chest compressions is not unnecessarily delayed.

PRINCIPLES OF RESUSCITATION

The clinical approach to cardiac resuscitation is an evolving and dynamic endeavor, and guidelines frequently change or

ANATOMY, PHYSIOLOGY, AND PATHOPHYSIOLOGY

The normal human heart rate (HR) is approximately 80 (±20) beats/min. With each beat the heart ejects a stroke volume (SV) of approximately 70 to 80 mL of blood from each ventricle. Multiplying HR by SV produces a value termed cardiac output (CO) (i.e., HR × SV = CO). The product of CO times total peripheral resistance (TPR) produces the value for mean arterial blood pressure (MABP) (i.e., CO × TPR = MABP). When the HR falls to zero or the heart fails to eject an SV (as in VF), MABP drops precipitously. Subsequently, vital organ perfusion is compromised. Hence, blood flow to the brain, the heart, the lungs, and other peripheral organs ceases. Failure to promptly restore blood flow will lead to significant mortality,
morbidity, and SCA. Therefore any interruption in cardiac contraction must be recognized quickly and corrected promptly. Cardiac contraction occurs as a result of a sequence of electromechanical events occurring in myocytes. The human heart has several unique characteristics that enable it to perform its physiologic role. These myocardial characteristics are automaticity, conductivity, excitability, and contractility. Individual cells have a “variable blend” of these characteristics. Some characteristics are more prominent than others, depending on the anatomic location of the cells in the heart. For example, the pacemaker cells have more automaticity, the conduction system has increased conductivity, and ventricular free-wall myocytes have more contractility. The electrical properties of these cells can be assessed by performing regional recordings of the changes in voltage in the tissue with respect to time (i.e., action potentials) (Fig. 12.5). The electrical impulse for myocardial contraction originates spontaneously in the sinoatrial (SA) node and spreads through the atria, which causes it to contract. As the impulse arrives at the atrioventricular (AV) node, it undergoes decremental conduction in which the electrical impulse is slowed down as the atria contract and “preload” the ventricles. Subsequently, the impulse activates the bundle of His and Purkinje fibers, which then cause ventricular contraction via excitation-contraction coupling. The electrical events precede the mechanical events. These events are graphically represented in Fig. 12.6, which depicts the change in membrane voltage with respect to time as a result of temporal changes in ion permeability across the myocyte membranes. These sequential changes in ion permeability occur as the membrane potential varies, thereby producing the characteristic cardiac action potential.

As the original impulse from the SA node travels through the atria and into the ventricles, various action potentials are generated regionally. The summation of all these action potentials Figure 12.4 High-quality cardiopulmonary resuscitation is essential in the resuscitation of victims of sudden cardiac arrest. Push hard to a depth of 2+ inches and +fast at a rate of 100 compressions per minute. Minimize interruptions and avoid overventilating the patient. Allow full recoil of the chest between compressions.

Relation of Action Potential from the Various Cardiac Regions to the Body Surface ECG

![Image of cardiac anatomy with action potential waveforms correlating to the body surface ECG.](https://www.netterimages.com)
Mechanisms of Cardiac Dysrhythmias

As is evident from the preceding discussion, normal cardiac activity is a compendium of complex, sequential electrochemical, physiologic, and mechanical events. It includes three mechanisms: enhanced automaticity, triggered activity, and reentry. If alterations in the action potential phases or a modification of the refractory periods occurs and another impulse stimulates the myocyte at a time that it is out of sync with the normal depolarization-repolarization process, the coordinated normal excitation-contraction coupling becomes asynchronous. If conditions favor the development of ectopic foci, individual loci in the ventricular free walls and septum become “pacemakers” and the myocardium begins to contract uncontrollably (see Fig. 12.2) and produce an irregular ECG tracing (see Fig. 12.1A). CO falls to zero, and SCA ensues. Another proposed mechanism that can precipitate the development of a dysrhythmia is a malfunction in propagation secondary to errors in conductivity and excitability and reentry of already propagated impulses (Fig. 12.8).

Defibrillation and Sudden Cardiac Arrest

When the heart is in VF or pulseless VT, applying a sufficient “burst” of therapeutic current across the myocardium will cause all the membrane channels that are involved in excitation-contraction coupling to be activated and mobilized into the absolute refractory period. As the myocardium enters the relative refractory period and returns to the resting state, the SA node will resume the role of pacemaker of the heart and normal AV contraction will resume. HR will increase, the ventricles will resume ejection of a normal SV, and CO will be adjusted to meet tissue needs. Subsequently, ROSC should ensue.

Cardiopulmonary Resuscitation: Ventricular Fibrillation and Pulseless Ventricular Tachycardia

When SCA occurs and the heart is in VF or pulseless VT, ventricular contraction is absent and circulation of blood comes to a standstill. To initiate CPR, mechanically compress the heart between the sternum and vertebral column. This causes pulsatile ejection of blood into the circulation, including the coronary circulation. For these compressions to be effective, perform them quickly and with sufficient displacement of the sternum (i.e., at least 2 inches) to produce adequate flow. Furthermore, keep interruptions in CPR to a minimum so that adequate perfusion pressure is maintained in the vasculature. Although the flow is not at physiologic levels, enough circulation occurs in the tissues, especially the myocardium, that the by-products of VF are “washed out” and the myocardium becomes less refractory to defibrillation. During VF, the myocytes are actually consuming oxygen and adenosine triphosphate at a rate believed to be the same or higher than during normal contraction. Several other concerns must be reinforced. During chest compressions, make sure that the chest recoils completely to the resting state so that blood can...
INDICATIONS FOR AND CONTRAINDICATIONS TO DEFRIBRILLATION

Prompt electrical defibrillation is the most effective treatment of acute SCA and VF. Prompt initiation of CPR in patients with SCA or VF is also critical for successful resuscitation and ROSC. Starting with the onset of collapse, the survival rate for patients with SCA or VF drops 7% to 10% for every minute of downtime without defibrillation. If CPR is initiated, the survival rate declines less rapidly (i.e., 3% to 4% per minute of downtime). If an SCA is witnessed and immediate CPR is provided, coupled with immediate defibrillation, survival from such events has been reported to increase up to fourfold. Therefore immediate defibrillation is indicated as soon as VF or pulseless VT is diagnosed. Few absolute specific contraindications to early defibrillation exist other than the presence of a pulse, absence of SCA, medical futility for the procedure, or a valid do-not-resuscitate order.

If a patient is found unresponsive, pulseless, and apneic and the downtime is unknown, immediately perform good-quality, effective CPR while preparing for defibrillation. Previous recommendations called for immediate defibrillation in lieu of a short period of CPR. The newest development in the 2010 AHA guidelines for CPR is also a change in the basic life support sequence of steps from the “ABCs” (airway, breathing, chest compressions) to “CAB” (chest compressions, airway, breathing) for adults and pediatric patients (children and infants, excluding newborns). As CPR is performed, prepare for rhythm analysis and initiate defibrillation if indicated. After performing CPR for 2 minutes (5 cycles at a rate of 30 compressions to 2 ventilations), perform rhythm analysis. If VF or pulseless VT is diagnosed, promptly perform defibrillation. When the time until the first shock is delayed during prehospital resuscitation (because of prolonged response times), data have demonstrated that the rate of successful defibrillation increases if patients receive bystander CPR before defibrillation. A scientific evaluation of this information proposed that CPR enhances the defibrillation threshold by restoring substrates to myocytes for the facilitation or resumption of normal excitation-contraction coupling. Furthermore, CPR may wash out myocardial depressants that have built up during prolonged VF. Therefore administration of CPR before defibrillation in patients with suspected, prolonged VF is recommended in the prehospital setting. Data to substantiate this sequence for in-hospital resuscitation have not been presented. Thus, the issue of unknown downtime, though not a definitive contraindication to immediate defibrillation, may be a factor in the clinician’s decision-making process regarding the resuscitation sequence.

Victims of SCA as a result of traumatic injuries do not usually survive. The heart, aorta, and pulmonary arteries may have sustained injury that will prevent resumption of normal cardiovascular function. There is a high probability that the underlying hypovolemia and organ damage may preclude successful resuscitation. However, the cause of the trauma may have been SCA with subsequent loss of consciousness. In such cases, if SCA or VF is present in a trauma patient, attempt treatment with CPR and defibrillation; if unsuccessful, search...
for and treat the underlying cause of the trauma and pursue the SCA. Therefore trauma is not a contraindication to defibrillation, although the resuscitative effort may be futile.

If the victim of VF or pulseless VT is a pregnant female, treatment of the mother is critical. Therefore prompt defibrillation is indicated for the same guidelines and sequencing as for nonpregnant patients. No harm to the fetus has been reported as a result of defibrillation, and thus pregnancy is not a contraindication to defibrillation.

Previous recommendations suggested delivering a “stacked” sequence of up to three shocks without interposed chest compressions if the first shock was unsuccessful in terminating VF. This was done to decrease transthoracic impedance with the monophasic damped sinusoidal (MDS) defibrillators in use and to deliver more current to the myocardium. However, this recommendation has been rescinded because of lack of supporting evidence. Now, with the higher first-shock efficacy (90%) in successfully terminating VF (termination of VF for 5 seconds) through the use of biphasic defibrillators, the recommendation to repeat a shock if the first treatment was unsuccessful is harder to justify. Hence, the AHA now recommends a one-shock protocol for VF. Evidence has accumulated that even short interruptions in CPR are harmful. Thus, rescuers should minimize the interval between stopping compressions and delivering shocks and should resume CPR immediately after delivery of a shock.

Defibrillation is also an effective treatment modality for terminating pulseless VT. If the patient has a pulse, is stable, and has a perfusing rhythm while in VT, defibrillation is contraindicated. However, if the patient in VT becomes unstable and shows signs of poor perfusion, has a change in mental status or if persistent chest pain with pulmonary edema, hypotension, and subsequent shock develop, synchronized cardioversion is recommended. This procedure is addressed later. If the patient becomes unstable as a result of polymorphic VT or becomes pulseless during the episode of VT, an unsynchronized shock (i.e., defibrillation) is indicated.

Patients “found down” or who have just become unresponsive can have other “rhythms present” besides VF or pulseless VT (e.g., PEA or asystole). Defibrillation is contraindicated in individuals with PEA. True asystole is not a shockable rhythm, and current evidence suggests that defibrillating patients with “occult” or false asystole is not beneficial and may actually be harmful. Therefore defibrillation is contraindicated in patients in asystole as long as fine VF has been ruled out (see discussion later).

Some patients who succumb to SCA may have various medication-releasing patches (e.g., nitroglycerin, contraceptive hormones, antihypertensive agents, smoking cessation adjuncts) present on their chest. Their presence is not a contraindication to defibrillation. However, modify the placement of the electrodes or paddles used for defibrillation to avoid contact with these patches. If necessary, remove these items before defibrillation to avoid diversion of current from the myocardium, current arcing, sparks, and other problems.

Developments in defibrillation and computer electronics have led to the availability and use of implantable defibrillators (automatic implantable cardiac defibrillators [AICDs], pacemakers) in the chest of patients who have known coronary artery disease. These patients are prone to dysrhythmias and may have episodes of VT and VF that are automatically detected and defibrillated or cardioverted. However, these devices can malfunction, so if these patients have SCA or VF, perform defibrillation as indicated. The presence of an AICD or pacemaker is not a contraindication to defibrillation.

Current trends in fashion sometimes include piercing of the body in various locations. In addition, certain items of clothing and jewelry may require modification of electrode or paddle placement. The presence of metal in locations proximal to the heart or in locations on the chest should be avoided to minimize the potential for diverting the defibrillating current from the myocardium. In addition, if the metal object provides a potential short circuit from the patient or leads to “ground,” this object should be removed, if feasible, to avoid diversion of current from the myocardium or arcing and burns across the chest. However, the presence of these materials, such as jewelry or body piercings, is not a contraindication to defibrillation.

In this part of the chapter on defibrillation the recommendations are intended for application to an adult patient (defined as older than 8 years or weighing more than 25 kg [55 lb]) with SCA or VF. If the patient is a child (e.g., 1 to 8 years of age or weighing less than 25 kg [55 lb]), modifications in the sequence, defibrillation energy, energy attenuation equipment, and size of the defibrillation paddles are necessary. Pediatric defibrillation details are discussed later in this chapter. If a defibrillator or automated external defibrillator (AED) and equipment suitable for use in children are not available, the health care provider can resort to using a standard AED or defibrillator. Use of AEDs or defibrillators in infants younger than 1 year has not been studied.

Defibrillation can be an ignition source for explosion if arcing occurs or if there are any stray or aberrant electrical discharges that occur as a result of paddle or electrode discharge. Therefore in an environment in which volatile explosive material is present, such as the operating room or other areas of critical care, be careful during defibrillation to avoid electrical arcing and to ensure that electrical conductivity through the patient’s chest is optimal. Avoid using anesthetic agents and oxygen. A potentially explosive environment is a relative contraindication to defibrillation.

When performing defibrillation, take care to avoid excessive moisture on the chest or around the patient. Although it is unlikely that there will be any significant or dangerous current leaks from the patient onto a wet floor, take care to avoid creating an electrical hazard. Try to ensure that the area is not wet; however, a wet surface is not an absolute contraindication to defibrillation. Defibrillation can be performed on ice and wet pavement.

Finally, defibrillation of an occult or false asystole or a very fine VF not detectable because of paddle or electrode position may be considered but is not recommended. Fine VF can occasionally masquerade as ventricular standstill or asystole. This may be a function of perpendicular electrode orientation with respect to the wavefront of depolarization. When evaluating the rhythm of a patient, if there is any doubt or confusion regarding the type of rhythm present, make sure that several leads are checked and rotate the paddles 90 degrees from their
original position to ensure that asystole is indeed present before abandoning the possibility of defibrillation. If fine VF is unmasked, consider providing aggressive CPR before defibrillation. In addition, place the controls on the ECG monitor on maximal gain to ensure adequate amplification of weak signals.

CONDUCTIVE MATERIAL

Use of conductive material is important to lower the impedance or resistance to flow of current at the electrode–chest wall interface. Multiple factors affect the range of impedance (e.g., body weight, chest size, chest hair, moisture on the skin surface of the patient, paddle size [diameter], paddle contact pressure, phase of respiration, and type of conductive material used). High impedance or resistance to flow of current can compromise the amount of current actually delivered to the myocardium and lead to a failed first shock. Inappropriate use of conductive material can result in current bridging or a short circuit and arcing of electrical current secondary to streaking of the material across the chest. This can produce sparks and unnecessary burns on the patient's skin. In addition, arcing of electricity can become a possible explosion hazard, depending on the circumstances. Conductive material needs to be used with the handheld electrodes. Various electrode gels are available on the market and should be kept in the proximity of the defibrillator, on the prearranged cart ready to use (see Fig. 12.11, step 2).

Self-adhesive pad electrodes now have a resistance-reducing, conductive material incorporated into the adhesive, thus rendering the use of a gel or other conductive material unnecessary. Firmly applying the self-adhesive electrode pads to the skin will usually be sufficient to minimize impedance, allow adequate ECG acquisition, and if indicated, defibrillate (see Fig. 12.9B).

PROCEDURE

Witnessed Sudden Cardiac Arrest (Figs. 12.10 and 12.11)

When confronted with a patient who has just become unresponsive, prepare for immediate defibrillation (Fig. 12.12). As soon as the defibrillator is available and the patient is connected to the monitor, assess the rhythm. In the interim, turn on the defibrillation equipment, place the paddles or electrodes on the chest, begin assessment of the patient, and initiate the steps in CPR by applying the CAB principle.

Cardiopulmonary Resuscitation

Perform a pulse check (<10 seconds; see Fig. 12.11, step 1). If a pulse is definitely present, provide 1 breath for 1 second every 5 to 6 seconds or 8 to 10 breaths/min. The breaths can be delivered with either a bag-valve-mask (BVM) or some type of barrier device. Observe the patient for visible chest wall rise and fall so that the thorax does not become overinflated. Hyperinflation of the chest can lead to inadvertent pressurization of the esophagus, which can cause lower esophageal sphincter pressure to be exceeded. This can lead to retrograde flow of gastric contents into the esophagus with the potential for subsequent aspiration of acid and debris into the trachea if the airway is not adequately protected. Overventilation of the thorax can also lead to an increase in intrathoracic pressure and impedance of blood flow to and from the heart, which should be avoided. Reassess the patient’s pulse every 2 minutes.

If no pulse is present, begin a sequence of 30 chest compressions followed by 2 ventilations/breaths (see Fig. 12.11, step 2). Keep your hands on the lower half of the sternum and compress it at least 2 inches (5 cm) at a rate of at least 100 compressions/min. The time allotted for compression should be 50%/50% for compression and relaxation of the chest. Watch for full chest recoil to allow adequate ventricular filling and do not lean on the chest, before the next compression. When the defibrillator or AED arrives, continue the 30:2 ratio of compressions to ventilations during CPR, attach the patient to the defibrillator via electrodes, pads, and paddles applied to the patient’s chest, and initiate the rhythm check (see Fig. 12.11, step 3). Every attempt should be made to minimize interruption of compressions.

Rhythm Assessment

Once the defibrillator is at the bedside, turn on the defibrillator/monitor and place electrodes on the patient’s chest in the form...
**Figure 12.10** Adult cardiac arrest algorithm. Note: The use of epinephrine and its safety and beneficial effects for an improved cardiac arrest outcome has recently been questioned. A worsened final outcome, possibly from prolonged cerebral vasoconstriction, has been associated with the use of epinephrine during resuscitation. Only two medications, epinephrine and amiodarone, are currently recommended for cardiac arrest; however, no ACLS drug has been proved to improve long-term survival. CPR, cardiopulmonary resuscitation; ET, endotracheal; IO, intraseous; IV, intravenous; PEA, pulseless electrical activity; PETCO₂, end-tidal CO₂; pVT, pulseless ventricular tachycardia; VT, ventricular fibrillation. (©2015 American Heart Association.)
DEFIBRILLATION

Assess patient responsiveness, breathing, and circulation. Check for a pulse for <10 seconds. Call for help and have the code cart delivered to the bedside.

Apply the electrodes to the patient’s chest. Place the sternal electrode below the clavicle, to the right of the sternum. Place the apical electrode in the midaxillary line at the fifth intercostal space.

Select the appropriate energy for the initial shock. For biphasic defibrillators, a default energy of 200 J is appropriate (see text for details). For monophasic machines, use 360 J. Make sure that “SYNC” is turned off.

As the “CHARGE” button is depressed, loudly announce “I’m clear, you’re clear, everybody’s clear!” and make sure that no caregivers are in contact with the patient.

Resume CPR immediately and continue for 5 cycles/2 minutes. Additional interventions such as IV/IO access and airway management may be pursued but should not interfere with continuous CPR.

If there is no pulse, begin CPR at a rate of >100 compressions per minute with a ratio of 30 compressions to 2 ventilations. Avoid interruptions in CPR, PUSH HARD, and PUSH FAST.

Check the rhythm on the monitor. If there is a shockable rhythm (i.e., VF or pulseless VT), prepare for immediate defibrillation.

Once the energy has been selected and the decision to defibrillate confirmed, press the “CHARGE” button on the defibrillator.

Check once again to make sure that everyone is clear, and then depress the “SHOCK” button to defibrillate the patient.

After 2 minutes of CPR, reassess the patient and the rhythm. Refer to the text and algorithms in this chapter for additional information.

Figure 12.11 Defibrillation. CPR, Cardiopulmonary resuscitation; IO, intraosseous; IV, intravenous; VT, ventricular fibrillation; VP, ventricular tachycardia.
Defibrillation and Cardioversion

CHAPTER 12

247

Figures 12.11 and 12.12: All resuscitation equipment should be kept on a “code cart” equipped with the monitor/defibrillator, airway supplies, emergency medications, and other essentials.

If a shockable rhythm such as VF or pulseless VT is present, defibrillation is indicated. Proceed to select the appropriate energy level for the anticipated defibrillation.

Energy Selection

As noted previously, two major types of defibrillators are available: biphasic and monophasic (Fig. 12.14). Currently, the biphasic defibrillator, which is more likely to be found in the clinical setting, produces either a biphasic rectilinear waveform or a biphasic truncated exponential (BTE) waveform. However, there are still monophasic defibrillators present that usually produce an MDS waveform. Current data do not support one waveform over another, but biphasic defibrillators appear to be more efficient in achieving defibrillation with the first shock.

Therefore the following recommendations are made: In general, a defibrillator using the biphasic rectilinear waveform should be set to an energy level of 120 J. If a BTE defibrillator waveform is being used, energy levels of 150 to 200 J are suggested for the first shock. If the type of waveform of the biphasic defibrillator is unknown or unavailable, a consensus default energy level of 200 J is suggested.

If the defibrillator is an older monophasic model using the MDS waveform, use 360 J for the first shock.
**Mode Selection**

Before defibrillation, check to make sure that the defibrillator is set to the unsynchronized mode. Most defibrillators default into the unsynchronized mode between shocks. Nonetheless, this control should be checked to make sure that it is in the unsynchronized mode; otherwise, the defibrillator may not discharge when the shock buttons are depressed because it is looking for the QRS complex, which is not present in VF. This is discussed in more detail in the section on Cardioversion later.

**Defibrillate**

Continue CPR until the defibrillator is charged and ready to defibrillate. Once the energy level has been selected and the decision made to defibrillate, clear the patient for defibrillation by loudly stating “I’m clear, you’re clear, everybody’s clear,” and then activate the button to charge the defibrillator (see Fig. 12.11, steps 6 and 7). Once the defibrillator has been charged and everyone is clear, apply firm pressure to the defibrillation paddles (25 lb) to increase contact and deflate the lungs to the end-expiration state. This will decrease impedance at the paddle–chest wall interface. Subsequently, depress the defibrillation controls and deliver the shock (see Fig. 12.11, step 8). This will usually be followed by a perceptible whole-body muscle twitch in the patient. If no obvious response or twitch of the patient is seen, check the defibrillator controls to make sure that it is in the unsynchronized mode and that the paddles are activated. If using the multifunctional pads, no pressure is needed.

**Resume Cardiopulmonary Resuscitation**

Once the shock has been delivered, resume resuscitation with immediate chest compressions (see Fig. 12.11, step 9). Continue chest compressions for approximately 5 cycles of 30 compressions to 2 ventilations, or approximately 2 minutes of CPR. This facilitates the transition from SCA to ROSC after the heart has been stunned by the defibrillation and may not be functioning at optimal contractility for a few minutes after the shock. If additional monitoring devices are in place, such as arterial lines or Swan catheters, modify this step accordingly as dictated by the resuscitation team leader.

Continue CPR for approximately 2 minutes. If the rescuers become fatigued, rotate the compressor and ventilator.

**Reassess the Patient: Management of the Airway and Intravenous Access**

After 2 minutes of CPR (5 cycles at a 30:2 ratio of compressions to ventilations), check the patient’s perfusion status or carotid pulses. If there is no palpable pulse, resume compressions immediately and prepare for delivery of a second defibrillatory shock.

As preparation for the second shock begins, other members of the resuscitation team can work on securing the airway via endotracheal intubation, a laryngeal mask airway, or another appropriate device. Proceed with blood drawing and intravenous (IV) line placement, or intraosseous (IO) if applicable, but do not interfere with chest compressions. The goal is to maintain uninterrupted chest compressions and to avoid any unnecessary interruptions.

**Changes in Cardiopulmonary Resuscitation**

Once an advanced airway has been secured, the compression and ventilation cycles are no longer delivered as described. Now, the compressor will continue to deliver compressions at a rate higher than 100 compressions/min continuously, without pausing for interposition of ventilation. Ask an assistant to deliver the ventilations at a rate of 8 to 10 breaths/min. Be careful to not overinflate the chest or to use too much force during ventilation because this can overpressurize the airways and esophagus, potentiate reflux, and impede venous return to the heart.

**Energy and Mode Selection for the Second Shock**

The energy for the second shock can be the same as that used before, but a higher energy level can be chosen at the discretion of the resuscitation leader. Check the mode selector again to be certain that it is in the unsynchronized position.

**Second Defibrillation**

Once the energy level has been selected, charge the defibrillator. When the defibrillator is ready to shock, halt CPR, clear the patient as discussed earlier, and deliver the second shock.

Resume CPR immediately after delivering the second shock. This step can be modified at the discretion of the resuscitation team leader, if there is clinical evidence of ROSC, or if devices are being used to monitor circulatory status (e.g., central venous pressure monitor, Swan-Ganz catheter, or direct arterial line).
If the patient does not have a shockable rhythm, proceed to the appropriate algorithm for VT, PEA, or asystole. Regarding airway management, consider using a supraglottic airway or endotracheal intubation without causing any significant interruption in chest compressions. In addition, initiate end-tidal carbon dioxide measurements (capnography) to determine the adequacy of CPR and ROSC when applicable.7

It is now believed that when SCA occurs in a presumably nonhypoxic heart, there is enough oxygen in the functional residual capacity (FRC) of the lungs (FRC = ERV [expiratory reserve volume] + RV [residual volume]) that with compressions only, blood will be oxygenated in the lungs for a short period. Therefore airway management is not as urgent as when restoring circulation that has totally ceased. Additionally, there is no need to overventilate the patient because hyperinflation of the lungs will cause an increase in intrathoracic pressure and compromise venous return to the right side of the heart.

**Unwitnessed Arrest**

When encountering a patient who is unresponsive and has been down for an unknown amount of time, assess the patient, summon help, and initiate CPR immediately, if indicated. Perform CPR until the defibrillator or AED is brought to the patient's side. As preparations are being made for defibrillation, consider performing 5 cycles of 30:2 compressions to ventilations before performing defibrillation.

**Automated External Defibrillator Application**

The availability of AEDs or semi-automated defibrillators in hospitals has increased, especially in non–critical care areas (Fig. 12.15). Although AEDs are designed for lay public use, application of these devices may also occur in the clinical setting. As in the algorithm, assess the patient, summon help, and apply the AED. Operation of the AED is guided by voice and visual prompts. Turn the device on, apply the patient electrodes in the appropriate positions, analyze the rhythm, and deliver a shock if a shockable rhythm is present. The AED will determine the rhythm and choose the energy level. Integrate CPR with the shocks to enhance the potential outcome of SCA resuscitation.

**Medication**

As per the 2010 AHA guidelines, there are insufficient data to demonstrate that any drugs or mechanical CPR devices improve long-term outcome after cardiac arrest. There is now some concern that epinephrine, a long-time universally recommended adjunct to CPR, may actually worsen outcomes in patients with SCA. The routine use of medications has been deemphasized, but not abandoned, in the current recommendations for resuscitation of SCA, VF, and pulseless VT (Box 12.1). Whether increased long-term survival from cardiac arrest can be expected with the use of any medications during CPR remains uncertain.

**Complications**

Complications of defibrillation include soft tissue injury, myocardial injury, and cardiac dysrhythmias. The availability of multifunctional electrode pads and better applicators for electrode gel has decreased the potential for soft tissue injuries such as chest burns.19 In fact, many clinicians now prefer to use the multifunctional electrode pads for ECG acquisition and for defibrillation.

The development of new, energy-efficient biphasic defibrillation waveforms, such as the BTE and the rectilinear biphasic waveform, has increased first-shock success and decreased the incidence of dysrhythmias after defibrillation.8 As a result, fewer shocks are needed to defibrillate the myocardium and less current is applied to the myocardium, which results in less electrical damage to myocytes.

Use of AEDs in public access defibrillation programs has not been reported to have produced any significant mishaps or adverse outcomes.20

Some older recommendations, such as use of the precordial thump, have been retracted. This procedure has been reported...
to have caused asystole or complete heart block (or both) when applied.\textsuperscript{21}

In addition, the use of procainamide, though not a complication, has fallen out of favor because of long infusion times and mixed results regarding the efficacy of its effects during the acute phase of VF and pulseless VT resuscitation.\textsuperscript{22}

### PEDIATRIC DEFIBRILLATION

Cardiac arrest in infants and children should initially be considered to be secondary to respiratory arrest. SCA, VF, and pulseless VT are much less likely to occur in children than in adults. However, 5% to 15% of pediatric and adolescent SCA events demonstrate VF in the prehospital setting. In in-hospital arrests, a 20% occurrence of VF at some point during the resuscitation is reported. Nonetheless, rapid intervention and defibrillation improve outcomes from SCA. Causes of SCA, VF, and pulseless VT are more diverse in pediatric patients.\textsuperscript{22} Cardiac arrest does not usually occur as a result of a primary cardiac cause. Therefore the approach to resuscitation of a pediatric patient in VF or pulseless VT may differ depending on the cause of the arrest.

### Ventricular Fibrillation in Children

VF is much less common in children than in adults. The etiology of VF and SCA in children is most likely to be sudden infant death syndrome, respiratory compromise, sepsis, neurologic disease, or injuries from motor vehicle crashes, burns, accidental firearm discharge, and drowning, many of which are preventable.\textsuperscript{21,24} The most common terminal rhythms reported in children younger than 17 years are PEA, bradycardia, and asystole.\textsuperscript{23} The etiology of these pediatric arrhythmias is most often hypoxemia, hypotension, hypoglycemia, and acidemia. In addition, focal electrical ectopy is less likely to initiate VF in a young heart. A significant myocardial mass must be unstable and fibrillating before VF becomes established. In children (from birth to 8 years old) with nontraumatic arrest, only 3% of the dysrhythmias are reported to be VF. In victims aged 8 to 30 years, the number of patients with VF increases by almost sixfold (17%).\textsuperscript{23} Several subpopulations of pediatric patients at various ages with cardiomyopathy or myocarditis or who have undergone heart surgery are at increased risk for a primary dysrhythmia.

As noted previously, the incidence of VF in cardiac arrest rhythms of pediatric patients is reported to range from 7% to 20%.\textsuperscript{26} Patients with rhythms who have been defibrillated from VF have been reported to have a higher survival-to-discharge rate than do children who sustained asystole or PEA.\textsuperscript{27} Therefore there is a definite indication for early defibrillation in the pediatric population.

#### Procedure and Technique

The procedure for pediatric defibrillation is similar to the algorithm for adult defibrillation (Fig. 12.16). However, a few differences must be addressed. These guidelines do not apply to children younger than 1 year.

### Pediatric Sudden Cardiac Arrest

When cardiac arrest occurs in a child, it is usually a terminal event associated with respiratory compromise or shock. The probability of SCA resulting from a primary cardiac cause is extremely low.\textsuperscript{21} Nonetheless, it can and does occur. If resuscitation is prompt, the potential for a positive outcome, including preservation of the patient’s neurologic integrity, is quite high. To enhance the outcome of SCA resuscitation, defibrillation and CPR must be effectively integrated. The pediatric resuscitation guidelines incorporated findings from a comprehensive review of the data.\textsuperscript{20} Revised steps for the recommended resuscitation sequence are described in the following sections.

#### Equipment

To perform pediatric defibrillation, a defibrillator monitor capable of adjustments in energy appropriate for children is needed. If an AED is to be used, it should have an energy attenuator for adjusting the energy to the appropriate level for a child (Fig. 12.17A). In addition, the quick-look electrode paddles (see Fig. 12.17B) should have adapters attached to the adult paddles to ensure appropriate contact with the chest wall in a child without causing the electrodes to overlap. If adhesive pads are used, choose the appropriate size that will not overlap (see Fig. 12.17C). Use gels as in adults while being careful to prevent bridging across the chest wall from streaks of conductive material that may have been carelessly applied to the chest.

#### Paddle and Pad Application and Use of Conductive Material

To acquire the electrical rhythm and subsequently administer an effective defibrillatory shock, place the appropriate-sized pads and paddles correctly on the chest. Use of the appropriate size and placement of paddles or pads will ensure that the appropriate current density is delivered across the myocardium to effectively defibrillate the myocytes. Furthermore, appropriate pad or paddle size—the largest surface area possible without direct electrode-to-electrode contact—will decrease transthoracic impedance and enhance defibrillation.\textsuperscript{29} To accomplish this, use infant paddles for children weighing less than 10 kg. However, use larger paddles if they do not contact each other. If contact is made between the paddles, an electrical arc or short circuit could occur.\textsuperscript{10} In children who weigh more than 10 kg (mean age, 1 year), use adult pads or paddles (8 to 10 cm in diameter).\textsuperscript{29} Use a conductive agent to enhance skin contact and decrease transthoracic impedance. Never use dry paddles because the resistance to flow of current will be very large. However, refrain from using saline-soaked pads in children because they may cause arcing as a result of the proximity of the pads on the chest. Remember that electricity will take the path of least resistance and that the current from defibrillation will travel across the chest if there is a saline bridge between the electrodes. In addition, the use of ultrasound gel and alcohol pads is discouraged because of poor electrical conductivity and potentially high impedance.\textsuperscript{10}

Apply the paddles or pads firmly to the chest, one to the right of the sternum, just below the clavicle, and the other to the left of the left nipple, over the ribs and the apex of the heart (see Fig. 12.13A). An option when using self-adhesive pads is to place one pad just to the left of the sternum and the other over the back so that they approximate the position of the heart (see Fig. 12.13B).

#### Procedure in an Unresponsive Child

When confronted with an unresponsive child, immediately summon assistance and start the CABs of CPR. Bring equipment
Defibrillation and Cardioversion

Figure 12.16 Pediatric defibrillation algorithm. Note: The use of epinephrine and its safety and beneficial effects for an improved cardiac arrest outcome have recently been questioned. No ACLS drug has been proved to improve long-term survival. CPR, Cardiopulmonary resuscitation; ET, endotracheal; IO, intravenous; IV, intravenous; PEA, pulseless electrical activity; pVT, pulseless ventricular tachycardia; VF, ventricular fibrillation. (©2015 American Heart Association.)
Rhythm Assessment. Once the defibrillator or monitor is at the patient’s side, turn it on and place the electrodes on the patient’s chest. The positions of the electrodes on a child correspond to the positions used in an adult (see Fig. 12.13). If quick-look paddles are used, carefully apply the conductive gel to the electrodes’ surface. If self-adhesive multifunctional electrode pads are used, there is no need to use conductive gel. Make sure that the input selector switch is reading from the appropriate source (i.e., paddles or pads). Adjust or increase the gain or sensitivity of the monitor so that fine VF is not missed because of low amplitude. As the ECG rhythm appears on the monitor, assess and diagnose the rhythm. If VF or pulseless VT is present, proceed to select the appropriate energy level for the anticipated defibrillation.

Energy Selection. As mentioned earlier in the adult section of the chapter, two types of defibrillators are available: biphasic and monophasic. As of this writing, there is no specific, detailed differentiation between energy levels to be used by either type of defibrillator. However, the caveat that biphasic shocks are at least as effective as monophasic shocks and that they are less damaging to the myocardium still applies.

Based on a review of adult and pediatric animal data, when a manual defibrillator is used for the first shock attempt, an energy level of 2 J/kg should be used with either a biphasic or a monophasic defibrillator. If a second or subsequent defibrillation is indicated, 4 J/kg should be used with either type. As mentioned earlier in the adult section of the chapter, two types of defibrillators are available: biphasic and monophasic. As of this writing, there is no specific, detailed differentiation between energy levels to be used by either type of defibrillator. However, the caveat that biphasic shocks are at least as effective as monophasic shocks and that they are less damaging to the myocardium still applies.

Mode Selection. Before defibrillation, check to make sure that the defibrillator is set to the unsynchronized mode. Most defibrillators default into the unsynchronized mode between shocks. Nonetheless, this control should be checked to make sure that it is in the unsynchronized mode; otherwise, the defibrillator may not discharge when the shock buttons are depressed because it is looking for the QRS complex, which is not present in VF (this is discussed in more detail in the later section on Cardioversion).

Defibrillate. Once the energy level has been selected and the decision made to defibrillate, simultaneously clear the patient for defibrillation by loudly stating “I’m clear, you’re clear, everybody’s clear,” while the button is activated to charge the capacitor. Continue CPR until ready to shock. Once the defibrillator has been charged and the patient cleared, apply firm pressure to the defibrillation paddles (25 lb) to increase contact and deflate the lungs to the end-expiration state. This will decrease impedance at the paddle–chest wall interface. Subsequently, depress the defibrillation controls and deliver the shock. This will usually be followed by a perceptible whole-body muscle twitch by the patient. If no obvious response or twitch of the patient is seen, check the defibrillator controls to make sure that it is in the unsynchronized mode and that the paddles are activated. No pressure is needed if adhesive multifunctional pads are used.
**Resume Cardiopulmonary Resuscitation.** Once the shock has been delivered, resume resuscitation with immediate chest compressions. Continue the compressions for approximately 5 cycles of 30 compressions to 2 ventilations or approximately 2 minutes of CPR. If two rescuers are available, use a 15:2 ratio and switch compressors when the first compressor fatigues. This is done to facilitate the transition from SCA to ROSC after the heart has been stunned by the defibrillation and may not be functioning at optimal contractility for a few minutes after the shock. If additional monitoring devices are in place in the hospital setting, modify this step accordingly as decided by the resuscitation team leader. Continue CPR for approximately 2 minutes.

**Reassess the Patient, Manage the Airway, and Gain Intravenous Access.** After 2 minutes of CPR, 5 cycles of 30:2, check the patient’s perfusion status or carotid pulses. If no pulse is palpable, resume compressions immediately and prepare to deliver a second defibrillatory shock.

While the operator is preparing for the second shock, other members of the resuscitation team can work on securing the airway via endotracheal intubation, a laryngeal mask airway, or another appropriate device. Blood draws and IV-line placement (or IO if applicable) should proceed but not interfere with chest compressions. The goal is to maintain chest compressions and avoid any unnecessary interruptions.

**Change in Cardiopulmonary Resuscitation.** Once an advanced airway has been secured, compression and ventilation cycles are no longer delivered. Now, the compressor will continue to deliver compressions at a rate of 100/min continuously without pausing for interposition of ventilations. When delivering the ventilations, provide 8 to 10 breaths/min, but be careful to not overinflate the chest or use too much force during ventilation to avoid overpressurizing the airways and esophagus and potentiating reflux. Lesser force also decreases the possibility of compromising CO as a result of elevated intrathoracic pressure.

**Second Shock: Energy Selection and Mode.** The second shock, if indicated, should occur after 2 minutes of CPR. The energy level for the second shock should be 4 J/kg. Be sure to check that the defibrillator or monitor is in the unsynchronized mode.

Immediately after the second shock, resume CPR and continue for approximately 2 minutes. If assessment of the rhythm and circulatory status shows continued VF or pulseless VT and no ROSC, continue CPR and consider medications such as a vasopressor (e.g., epinephrine).

**Medications.** The same caveats concerning the lack of proven benefit of any medications to improve long-term survival in adults also applies to children.

**Automated External Defibrillators in Children**

As mentioned previously, the incidence of VF and pulseless VT in children is low. Nonetheless, the presence of VF or pulseless VT is an indication for using an AED or defibrillator. The age range for use of an AED or defibrillator is 1 to 8 years. No recommendations for the use of a defibrillator or AED in children younger than 1 year have been provided as of this writing.17 A pediatric energy dose attenuator (see Fig. 12.17-4) should be used to prevent the delivery of too much current to the myocardium. If a pediatric dose attenuator is not immediately available, a standard defibrillator should be used at the lowest appropriate setting.

Use of the AED entails bringing the AED to the patient’s side, turning the device on, following the voice or visual prompts, and connecting the electrodes to the patient. Once the pads are applied to the patient, the AED will initiate the rhythm analysis automatically or the rescuer will be prompted to press a button to activate the “analyze mode” of the AED. Subsequently, the AED will diagnose the rhythm and advise a shock if indicated. The energy level and mode are all preprogrammed into the AED electronics.

**CARDIOVERSION**

**Introduction and Physiology**

Cardioversion is the application of a direct current (DC) “shock” across the chest or directly across the ventricle to normalize the conduction pattern of a rapidly beating heart. This shock is delivered during the absolute refractory period of the ECG QRS; it is synchronized to the peak of the R wave.

A patient with significant tachycardia may be asymptomatic or may complain of chest pain or discomfort, light-headedness, or shortness of breath. These symptoms are the result of altered cardiovascular physiology. Rapid cardiac rhythms allow less time for ventricular filling and thereby result in reduced preload and hypotension. The reduced preload, as well as the increased ventricular work caused by the rapid HR, may also result in ventricular ischemia. Pulmonary capillary wedge pressure may also rise despite the shortened filling time because of reduced ventricular compliance secondary to ventricular ischemia. Elevated pulmonary capillary wedge pressure can then lead to pulmonary edema.

Termination of rapid rhythms to alleviate or prevent these symptoms must occur quickly to prevent further deterioration. Persistently poor CO because of a rapid HR results in the development of lactacidosis, which further compromises cardiac function and makes cessation of the dysrhythmia even more difficult. Unchecked myocardial ischemia may lead to infarction with its attendant sequelae. Carotid sinus message drug therapy, rapid cardiac pacing, and cardioversion are the methods available to terminate tachydysrhythmias.

In many cases, DC cardioversion has specific advantages over drug therapy. Although it may require more preparation than drug therapy, the effectiveness of electrical cardioversion enhances its usefulness in the ED setting. Cardioversion is effective almost immediately, has few side effects, and is often more successful than drug therapy in terminating dysrhythmias. In addition, the effective dose of many anti- dysrhythmic medications is variable, and there is often a small margin between therapeutic and toxic dosages. Although they can often suppress an undesirable rhythm, drugs may also suppress a normal sinus mechanism or may create toxic manifestations that are more severe than the dysrhythmia being treated.

In the clinical setting of hypotension or acute cardiopulmonary collapse, cardioversion may be lifesaving, and is preferred over drug therapy. Key concepts in the use of this procedure include understanding the indications for its use,
the equipment involved, the importance of adequate sedation, and the concerns for health worker safety.

Indications and Contraindications

All decisions regarding the need for cardioversion are best made at the bedside by the clinician assessing the given scenario. There are no firm guidelines on exactly what defines an unstable clinical situation, and definitions are relative terms that lend themselves to real-time clinical decision making and clinician interpretation. Cardioversion is often indicated whenever a reentrant tachycardia is causing chest pain, pulmonary edema, light-headedness, or hypotension. This excludes tachydysrhythmias that are known to be caused by digitalis toxicity, as well as a known sinus tachycardia. It is also indicated in less urgent circumstances when medical therapy has failed. In elderly patients, in whom a prolonged rapid heartbeat can be anticipated to cause complications (e.g., clots, thrombi) related to cardiac ischemia or dysfunction, early intervention with cardioversion may also be beneficial.

A reentrant tachydysrhythmia should be suspected when a sudden change in HR occurs within a few beats. Unless the dysrhythmia is noted whilst the patient is being monitored, it can be inferred only from the patient’s history of a sudden onset of symptoms. In the unusual case of sinus node reentrant tachycardia, rapid onset and offset may be the only clues. Other clues to the presence of a reentrant dysrhythmia are a history of Wolff-Parkinson-White (WPW) syndrome or another known accessory pathway syndrome. Ventricular rates in excess of those predicted for age strongly suggest an accessory pathway.

Dysrhythmias caused by enhanced automaticity will not be terminated by uniformly depolarizing myocardial tissue because a homogeneous depolarization state already exists. Enhanced automaticity is the cause of most cases of digitalis toxicity–induced dysrhythmia, sinus tachycardia, and multifocal atrial tachycardia. Although cardioversion will not work in these cases, medications that suppress automaticity, including potassium and magnesium, may be useful.

In digoxin toxicity, not only is cardioversion ineffective, but it is also associated with a higher incidence of post-shock VT and VF. However, in a patient with a therapeutic digoxin level, the risk associated with cardioversion is now thought to be no different from that of other patients. Digoxin is still generally withheld for 24 hours before cardioversion as a precaution against inadvertently elevated levels. Pregnancy at any stage is not a contraindication to cardioversion.

Treatment

Therapy is dictated by the specific wide-complex tachycardia and the patient’s clinical findings (Fig. 12.18). The initial approach must always be led, and modified if necessary, by the patient’s signs and symptoms and subsequent changes. Synchronized monophasic or biphasic cardioversion is the appropriate first choice of treatment for unstable patients. In patients deemed to be stable, the therapeutic options are more diverse. Stable, wide-complex tachycardia can always be considered VT and treated according to current VT algorithms. A reasonable treatment protocol for stable patients may be the use of adenosine, procainamide, lidocaine, and finally, cardioversion. Amiodarone is effective for most SVTs, and its use for stable unknown wide-complex SVT is both appropriate and safe.

Equipment and Setup

The critical components of preparation for cardioversion are IV access, airway management equipment, drugs for sedation, monitoring, and DC delivery equipment (cardioverter) (Fig. 12.19, step 1).

Secure IV access is essential for delivery of sedatives, antidysrhythmics, fluids, and possibly paralytic agents. Although many of these drugs are not used routinely, if they are needed, timing is likely to be critical. A large-bore IV catheter should be inserted and firmly taped to the patient’s skin.

A significant and preventable complication of procedures involving sedation is hypventilation leading to hypoxia. Airway management equipment includes the secure IV catheter discussed previously, working suction with a tonsil-tipped device attached, BVM apparatus, oxygen, and appropriately sized laryngoscope and endotracheal tube. A pulse oximeter is generally recommended for patients undergoing conscious sedation. Another adjunct is continuous monitoring of carbon dioxide pressure (PCO₂). A rising PCO₂ level will be an earlier clue to hypventilation secondary to sedation because oxygen saturation may remain normal for several minutes, especially if the patient has been preoxygenated.
Sedative medications should be ready for use in labeled syringes, along with a prefilled saline syringe for flushing the catheter. Antidysrhythmic medications for ventricular dysrhythmias (e.g., amiodarone, lidocaine) and for unexpected brady-cardia (e.g., atropine) should be readily accessible.

**Technique**

If time permits, metabolic abnormalities such as hypokalemia and hypomagnesemia should be corrected before attempting cardioversion. At a minimum, hypoxia should be corrected with supplemental oxygen. If a patient has metabolic acidosis, compensatory hyperventilation after endotracheal intubation may be indicated before cardioversion. Respiratory acidosis should always be treated before the use of sedative drugs.

**Sedation**

Cardioversion may be extremely painful or terrifying, and patients must be adequately sedated before its use (see Fig. 12.19, step 2). Patients who are not adequately sedated may experience extreme anxiety and fear. Several IV medications are available for sedation of patients before cardioversion,
TABLE 12.1 Commonly Available Intravenous Medications Used for Sedation in Cardioversion

<table>
<thead>
<tr>
<th>DRUG</th>
<th>DOSE</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Midazolam</td>
<td>0.15 mg/kg</td>
<td>Most commonly used. Induction occurs in approximately 2 min. Small drop in blood pressure. Flumazenil antagonist available.</td>
</tr>
<tr>
<td>Methohexital</td>
<td>1 mg/kg</td>
<td>Quicker onset than midazolam. Shorter duration than midazolam. Small drop in blood pressure. Rare complication of laryngospasm.</td>
</tr>
<tr>
<td>Propofol</td>
<td>1.5 mg/kg</td>
<td>Small drop in blood pressure. Painful IV infusion.</td>
</tr>
<tr>
<td>Thiopental</td>
<td>3 mg/kg</td>
<td>Painful IV infusion.</td>
</tr>
<tr>
<td>Fentanyl</td>
<td>1.5 µg/kg</td>
<td>An opiate. Added for more sedation. Can cause respiratory depression.</td>
</tr>
<tr>
<td>Ketamine</td>
<td>1.5 mg/kg</td>
<td>Will not cause hypotension. May be combined with reduced-dose propofol (0.5 mg/kg) or a full-dose benzodiazepine.</td>
</tr>
</tbody>
</table>

including etomidate (0.15 mg/kg), midazolam (0.15 mg/kg), methohexital (1 mg/kg), propofol (0.5–1.0 mg/kg), and thiopental (3 mg/kg). In addition, IV ketamine (1.5 mg/kg), with or without a benzodiazepine or slightly reduced-dose propofol, and IV fentanyl (1.5 µg/kg), a synthetic opioid analgesic, may be administered 3 minutes before induction (Table 12.1).

Midazolam is probably the most commonly used agent, with induction occurring approximately 2 minutes after a dose of approximately 0.15 mg/kg, or at least 5 mg for an average-sized adult. Although induction with midazolam takes slightly longer than with the other medications, it has the advantage that a commercial antagonist, flumazenil, is available for reversal if necessary. Small additional doses of fentanyl (1 to 1.5 µg/kg) may be added for more profound sedation. Fentanyl can cause respiratory depression, but its action can be reversed with naloxone. Methohexital has the advantage of quick onset and a somewhat shorter duration of action than midazolam does, but it has a rare association with laryngospasm. All the drugs except etomidate and ketamine may cause a small drop in blood pressure, and infusion of propofol and etomidate is painful. Ketamine is a reasonable choice in patients with borderline hypotension.

In elderly patients, the pharmacodynamics and kinetics are altered by coexisting illness and polypharmacy rather than by any intrinsic effect of old age. Drug doses should be reduced in these patients.

Administer the anesthetic agent or agents intravenously over a period of approximately 30 seconds and wait until the patient is unable to follow simple commands and loss of the eyelash reflex is noted. Administering the agent too quickly may result in hypotension; administering it too slowly may not allow blood levels to reach a therapeutic range if the agent has a rapid rate of metabolism.

**Cardioverter Use**

Selection of the synchronized or nonsynchronized mode is the next critical step (see Fig. 12.19, step 3). In the synchronized mode, the cardioverter searches for a large positive or negative deflection, which it interprets as the R or S wave. It then automatically discharges an electric current that lasts less than 4 msec, thereby avoiding the vulnerable period during repolarization when VF can easily be induced. Once the cardioverter is set to synchronize, a brief delay will occur after the buttons are pushed for discharge as the machine searches for an R wave. This delay may be disconcerting to an unaware operator.

If concern exists about whether the R wave is large enough to trigger the electrical discharge, the clinician can place the lubricated paddles together and press the discharge button. Firing should occur after a brief delay. When the R- or S-wave deflection is too small to trigger firing, change the lead that the monitor is reading or move the arm leads closer to the chest.

If there is no R or S wave to sense, as in VF, the cardioverter will not fire. Always turn off “synchronization” if VF is noted.

**Electrode Position: Same as for Defibrillation**

Electrode paddles may be positioned just as they are for defibrillation. Safety is a key concern in the performance of cardioversion. Any staff member acting as a ground for the electrical discharge can be seriously injured. The operator must announce “all clear” and give staff a chance to move away from the bed before discharging the paddles. Care must be taken to clean up spills of saline or water because they may create a conductive path to a staff person at the bedside.

**Energy Requirements**

The amount of energy required for cardioversion varies with the type of dysrhythmia, the degree of metabolic derangement, and the configuration and thickness of the chest wall (see Fig. 12.19, step 4). Obese patients may require a higher energy level for cardioversion, and the anteroposterior paddle position is sometimes more effective in these patients. If patients are shocked while in the expiratory phase of their respiratory cycle, energy requirements may also be lower.

VT in a hemodynamically stable patient should be treated with amiodarone, 150 mg intravenously, and this can be repeated as needed up to a dose of 2.2 g/24 hr. If unsuccessful, cardioversion is then performed. Cardioversion with 10 to 20 J is successful in converting VT in more than 80% of cases. Cardioversion will be accomplished with 50 J in 90% of cases, and conversion should initially be attempted at this energy level. Cardioversion should be synchronized unless the T wave is large and could be misread as the R wave by the cardioverter. If the initial attempts at electrical cardioversion
are unsuccessful, the energy level should be doubled, and doubled again if necessary, until a perfusing rhythm is restored. Immediately after conversion of VT, antidysrhythmic medications should be given to prevent recurrence.

Patients with pulseless VT should be initially shocked with 200 J, followed by 300 J if the first shock is not successful. Reentrant SVTs generally respond to low energy levels. Atrial flutter, for example, usually requires less than 50 J for conversion. Cardioversion of atrial flutter in the emergency department (ED) is indicated when the ventricular rate is not slowing in response to pharmacologically enhanced AV node blockade or if the patient is unable to tolerate the aberrant rhythm.

The majority of patients with paroxysmal atrial tachycardia respond to adenosine. If they do not, or if urgent conversion is needed because of a high ventricular rate, an electrical countershock should be administered in the synchronized mode at 50 J and doubled if necessary.

In patients with AF, the response to cardioversion is dependent on the duration of the AF and its underlying cause. Most patients with AF do not require cardioversion in the ED unless their ventricular response is high because of a bypass tract, as in WPW syndrome. They may also require cardioversion when sequelae of rapid ventricular contraction are present or anticipated and the ventricular rate is not responding to drug therapy aimed at slowing AV node conduction. Conversion of AF generally requires more energy than reentrant SVTs do (=100 J in most cases).

Complications

Complications of cardioversion may affect the patient, particularly those with a cardiac pacemaker, as well as health care personnel at the bedside. Injuries to health care personnel during cardioversion or defibrillation include mild shock and burns.

Patient complications are dose related and may involve the airway, heart, or chest wall, or they may be psychological. Hypoxia may develop in patients if sedation is excessive or the airway becomes compromised. With proper preparation and precautions, airway complications can be minimized. Respirations may also be depressed by any of the anesthetic agents, and the adequacy of tidal volume must be continually assessed by either direct observation or end-tidal CO2 monitoring. If another clinician is available, that clinician should be placed in charge of monitoring the patient’s airway. Routine supplemental oxygen is suggested for all patients undergoing sedation.

Chest wall burns resulting from electrical arcing are generally superficial partial-thickness burns, although deep partial-thickness burns have occurred. They are preventable by adequate application of conductive gel and firm pressure on the paddles. Paddles should not be placed over medication patches or ointments, especially those containing nitroglycerin, because electrical discharge may cause ignition and result in chest burns.

Cardiac complications after cardioversion are proportionate to the energy dose delivered. In the moderate energy levels used most commonly, the hemodynamic effects are small. At higher energy levels, however, complications include dysrhythmias, hypotension, and rarely, pulmonary edema, which may occur several hours after the countershock.

The dysrhythmias occurring after high-dose (≈200 J) DC shocks include VT and VF, bradycardia, and AV block, in addition to transient and sustained asystole. Sustained VT or VF was reported following 7 of 99 shocks in a study of patients undergoing electrophysiologic study and requiring cardioversion for VT, VF, or AF. These episodes occurred only in patients with prior VT or VF. Patients with ischemia or known coronary artery disease appear to be at much higher risk for significant post-shock bradycardia, with rate support pacing being required after 13 of 99 shocks in the aforementioned study. Asystole requiring pacing occurred only once in 99 countershocks. Therefore the proclivity for dysrhythmias is greater with high-dose cardioversion of an ischemic heart.

Conclusion

Cardioversion is a safe and effective method of quickly terminating reentrant tachycardia. Complications related to psychological trauma, respiratory depression, and unintentional shock in health care workers can be avoided with proper precautions. Adequate sedation is essential. Synchronized shock should be administered after close scrutiny of the lead used for sensing to be sure that the R or S wave is significantly larger than the T wave. Be prepared for post-shock VT or VF, and if VF occurs, switch the cardioverter to “nonsynchronized” and defibrillate. Atropine and temporary pacing equipment should be available to treat post-shock bradycardia, especially in patients with myocardial ischemia or infarction.

Pediatric Cardioversion

Pediatric cardioversion is similar to adult cardioversion. As described previously, the purpose of the procedure is to depolarize the myocytes completely at the most opportune time, during the peak of the R wave so that VF is not precipitated, and allow a slower perfusing rhythm to resume. However, the energy levels for pediatric cardioversion are different from those for adults. In the pediatric procedure, the initial recommended energy dose is 0.5 to 1 J/kg with the defibrillator in the synchronized mode. If needed, repeated cardioversion may be attempted at 2 J/kg, again while the defibrillator is in the synchronized mode. Remember to resynchronize the defibrillator after each cardioversion attempt and look for the appropriate markers on the monitor to ensure that the current is delivered at the appropriate phase of the cardiac cycle. If medication is needed, amiodarone at a dose of 5 mg/kg intravenously over a 60-minute period or procainamide at a dose of 15 mg/kg over a 60-minute period can be used (do not give these drugs together).

ACKNOWLEDGMENT

The editors and author would like to acknowledge the significant contributions to this chapter in previous editions by Steven Gazak, MD, William Burdick, MD, Jerris R. Hedges, MD, Michael Greenberg, MD, and John Krimm, DO.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Patients with implanted pacemakers or automatic implantable cardioverter defibrillators (AICDs) are commonly seen in the emergency department (ED). Fortunately, the increased reliability of these devices has precluded a marked increase in patients with true emergencies related to device malfunction, but such patients clearly have serious underlying medical problems that must be considered. Pacemaker complications are not uncommon, with rates ranging from 2.7% to 5%. Many pacemakers fail within the first year. AICD complication rates, including inadvertent shocks, occur in up to 34% of patients with the device. The basic evaluation and treatment of patients with cardiac complaints are not substantially different in patients with pacemakers and AICDs than those without. However, a general knowledge of the range of problems, complications, and techniques for evaluating or inactivating pacemakers or AICDs is important for emergency clinicians. These devices are complicated and emergency physicians are not expected to be experts in all complications; therefore appropriate consultation is often necessary depending on the clinical situation.

**PACEMAKER CHARACTERISTICS**

In essence a pacemaker consists of an electrical pulse-generating device and a lead system that senses intrinsic cardiac signals and then delivers a pulse. The pulse generator is hermetically sealed with a lithium-based battery device that weighs approximately 30 g and has an anticipated lifetime of 7 to 12 years. A semiconductor chip serves as the device’s central processing unit. The generator is connected to sensing and pacing electrodes that are inserted into various locations in the heart, depending on the configuration of the pacemaker. Newer models are programmable for rate, output, sensitivity, refractory period, and modes of response. They can be reprogrammed radiotelemetrically after implantation.

Pacemakers are classified according to a standard five-letter code developed by the North American Society of Pacing and Electrophysiology/British Pacing and Electrophysiology Group (Table 13.1). Known as the NBG code, it consists of five positions or digits. The first letter designates the chamber that receives the pacing current; the second, the sensing chamber; and the third, the pacemaker’s response to sensing. The fourth letter refers to the pacemaker’s rate modulation and programmability, and the fifth describes the pacemaker’s ability to provide an antitachycardia function. Whereas standard pacemakers generally do not have an antitachycardia function, AICDs do have this capability and overdrive pacing is the device’s first response to tachycardia. In normal practice, only the first three letters are used to describe the pacemaker (e.g., VVI or DDD).

Pacemaker wires are embedded in plastic catheters. The terminal electrodes, which may be unipolar or bipolar, travel from the generator unit to the heart via the venous system. In a unipolar system, the lead electrode functions as the negatively charged cathode, and the pulse generator case acts as the positively charged anode into which electrons flow to complete the circuit. The pulse generator casing must remain in contact with tissue and be uninsulated for pacing to occur. In the case of bipolar systems, both electrodes are located within the heart. The cathode is at the tip of the lead, and the anode is a ring electrode roughly 2 cm proximal to the tip. Bipolar leads are thicker, draw more current than unipolar leads, and are commonly preferred because of several advantages, including a decreased likelihood of pacer inhibition as a result of extraneous signals, and decreased susceptibility to interference by electromagnetic fields.

The typical entry point for inserting the leads is the central venous system, which is generally accessed via the subclavian or cephalic vein. The terminal electrodes are placed either in the right ventricle or in both the right ventricle and the atrium under fluoroscopic guidance. Proper lead placement, sensing, and pacing thresholds are assessed with electrocardiograms (ECGs). The typical radiographic appearances of implanted pacemakers and implantable cardioverter-defibrillators to include an example of lead fracture are depicted in Fig. 13.1A–E.

The pacemaker is typically programmed to pace at a rate of 60 to 80 beats/min. A significantly different rate usually indicates malfunction. When the battery is low, the rate generally begins to drop and gets slower as the battery fades. Senses of intracardiac electrical activity is a combination of recognizing the characteristic waveforms of P waves or QRS complexes while discriminating them from T waves or external interfering signals, such as muscle activity or movement. The pacing electrical stimulus is a triphasic wave consisting of an intrinsic deflection, far-field potential, and an injury current, which typically delivers a current of 0.1 to 20.0 mA for 2 msec at 15 V.

Pacemakers have a reed switch that may be closed by placing a magnet over the generator externally on the chest wall; this inactivates the sensing mechanism of the pacemaker, which then reverts to an asynchronous rate termed the magnet rate. Essentially, the magnet turns the demand pacemaker into a fixed-rate pacemaker. The magnet rate is usually, but not always the same as the programmed rate.

Several innovations in rate regulation have been incorporated into some pacemakers. When present, the hysteresis feature causes pacing to be triggered at a rate greater than the intrinsic heart rate. When the hysteresis feature is used in a single-chamber ventricular pacemaker, it is designed to maintain atrioventricular (AV) synchrony at rates that are lower than what would be normal for a ventricular-paced rhythm alone. To illustrate, were the hysteresis feature of the pacemaker set at 50 beats/min, an intrinsic rate lower than 50 beats/min would trigger ventricular pacing. Unlike a standard ventricular pacemaker, the hysteresis feature might be set to offer a ventricular pacing rate at 70 beats/min or greater once the pacer is triggered.

Rate modulation by sensor-mediated methods is an additional feature triggered and mediated by a sensed response to various physiologic stimuli. The primary application for this rate modulation feature is in patients with pacemakers who continue to engage in vigorous physical activity. When present, the rate regulation feature is engaged and modulated through motion sensors installed within a pulse generator device, with
A corresponding increase or decrease in the pacing rate depending on the degree of motion sensed by the pacemaker device. Other physiologic sensors that may be installed as part of the pacemaker system include those designed to sense minute ventilation, the QT interval, temperature, venous oxygen saturation, and right ventricular contractions. The latter sensors generally require that additional leads be placed.

One of the newest innovations in cardiac pacing is cardiac resynchronization therapy (CRT), also known as biventricular pacing. In addition to a conventional right ventricular endocardial lead, CRT also employs a coronary sinus lead for left ventricular pacing. A right atrial lead may also be included. The primary objective of CRT is to restore left ventricular synchrony, primarily in patients with dilated cardiomyopathy and widened QRS complex. This condition results primarily from the presence of a left bundle branch block. In such situations CRT may improve left ventricular function, and thus cardiac output.

### Characteristics of AICDs

The basic components of an AICD, including sensing electrodes, defibrillation electrodes, and a pulse generator (Fig. 13.2), can be seen on a chest radiograph. Transvenous electrodes have obviated the previous need for surgical placement. They are inserted into the pectoralis muscle. Many transvenous systems consist of a single lead containing a distal sensing electrode and one or more defibrillation electrodes in the right atrium and ventricle. Leads are inserted through the subclavian, axillary, or cephalic vein into the right ventricular apex. The left side is preferred because of a smoother venous route to the heart and a more favorable shocking vector. In an effort to improve the efficiency of defibrillation, an additional defibrillation coil may be used. Various placements of AICDs are demonstrated in Fig. 13.3.

The pulse generator is a sealed titanium casing that encloses a lithium–silver–vanadium oxide battery. It has voltage converters and resistors, capacitors to store charge, microprocessors and integrated circuits to control analysis of the rhythm and delivery of therapy; memory chips to store electrographic data, and a telemetry module. Although a pacemaker can draw the voltage required for function from its component battery, the energy needed for defibrillation requires a battery that is prohibitively large. To circumvent this problem, an AICD contains a capacitor that maximizes the voltage required by transferring energy from the battery before discharge. To achieve the energy required, AICDs use capacitors that are charged over a period of 3 to 10 seconds by the battery and then release this energy rapidly for defibrillation. The maximal output is 30 J in most units and 45 J in higher-energy units. This energy is high enough that a discharge is very obvious and often distressing to the patient.

Most AICDs use a system in which the pulse generator is part of the shocking circuit, often described as a “can” technology, and most of them have a dual-coil lead with a proximal coil in the superior vena cava and a distal coil in the right ventricle. Current flows in a three-dimensional configuration from the distal coil to both the proximal coil and the generator. This dispersion of the electrical field increases the likelihood of depolarizing the entire myocardium at once, thereby leading to successful defibrillation.

AICDs may have the same programming capabilities as pacemakers and can be single chambered, dual chambered, or used with the aforementioned CRT. Single-chamber devices have only a right ventricular lead. They often have difficulty identifying atrial arrhythmias, which can result in inappropriate defibrillation of atrial tachycardias. Dual-chamber AICDs have right atrial and right ventricular leads and improved ability to discriminate rhythms. In most studies, dual devices have been found to offer improved discrimination between ventricular and supraventricular arrhythmias, thus decreasing inappropriate shocks as a result of rapid supraventricular rhythms or physiologic sinus tachycardia. Of AICDs implanted annually 27% are single chamber, 32% dual chamber, and 41% are CRT systems. In patients requiring both AICD and pacemaker functions, both these devices are placed together. The advent of technology has allowed placement of a single device that can perform both pacemaker and defibrillator functions.

AICDs use a combination of antitachycardia pacing, low-energy cardioversion, defibrillation, and bradycardiac pacing in a combination also known as tiered therapy. They are programmed with specific algorithms that identify and treat specific rhythms. Ventricular arrhythmias may initially be converted (or undergo attempts at conversion) with antitachycardiac pacing as opposed to immediate defibrillation. This overdrive pacing may terminate the rhythm without the need for electrical defibrillation in up to 90% of events. It is most successful for terminating monomorphic ventricular tachycardia with a rate of less than 200 beats/min. Overdrive pacing is better tolerated by patients than cardioversion and reduces the risk for inducing atrial fibrillation. These events may be silent, not felt by the patient, and discovered only by interrogating the device.

If unsuccessful, the next intervention may be low-energy cardioversion (<5 J). The device may be programmed to very

### Table 13.1 North American Society of Pacing and Electrophysiology/British Pacing and Electrophysiology Group Generic Pacemaker Code (NBG Code)

<table>
<thead>
<tr>
<th>I Chamber Paced</th>
<th>II Chamber Sensed</th>
<th>III Response to Sensing</th>
<th>IV Rate Modulation and Programmability</th>
<th>V Antitachycardia Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>0—None</td>
<td>0—None</td>
<td>0—None</td>
<td>0—None</td>
<td>0—None</td>
</tr>
<tr>
<td>A—Atrium</td>
<td>A—Atrium</td>
<td>I—Inhibited</td>
<td>I—Inhibited</td>
<td>P—Antitachycardiac pacing</td>
</tr>
<tr>
<td>V—Ventricle</td>
<td>V—Ventricle</td>
<td>T—Triggered</td>
<td>M—Multiple</td>
<td>S—Shock</td>
</tr>
<tr>
<td>D—Dual</td>
<td>D—Dual</td>
<td>D—Dual</td>
<td>C—Communicating</td>
<td>D—Dual</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>R—Rate modulation</td>
<td></td>
</tr>
</tbody>
</table>

low levels of electricity that are better tolerated by the patient. This works best for ventricular rates higher than 150 and lower than 240 beats/min. This may be followed by a high-energy defibrillation. Traditionally, the energy level of the first shock is set at least 10 J above the threshold of the last defibrillation measured. If the first shock fails a backup shock may be required, but this may induce or aggravate ventricular arrhythmias (see the later section on Pacemaker-Mediated Tachycardia). Unlike the proarrhythmic effects of medication, these arrhythmias are almost never fatal, although they may be associated with increased morbidity. Currently used biphasic waveforms have improved defibrillation thresholds.

This tiered approach obviates the need for unnecessary energy requirements. The devices also have antibradycardic

![Figure 13.1 A](image1) Various radiographs of an implanted pacemaker and automatic implantable cardioverter-defibrillator showing battery and lead wires. Posteroanterior (PA; A1) and lateral (A2) chest radiographs demonstrate a biventricular pacing system. There are three leads—the first is positioned in the right atrium, the second is in the right ventricular apex, and the third courses posteriorly in the coronary sinus and into the posterolateral cardiac vein. B, PA chest radiographs of a dual-chamber pacemaker. B1, The ventricular lead is passing through an atrial septal defect into the left ventricle. B2, The lead is repositioned in the right ventricular apex. C, A dual-chamber implantable cardioverter-defibrillator with active fixation leads has been implanted via a transvenous approach to place the atrial lead in the systemic venous atrium and the ventricular lead across the baffle into the morphologic left ventricle.
pacing that allows these patients to have one device instead of separate units. Additional complications associated with AICDs that have antibraycardiac pacing algorithms include a tendency toward oversensing, increased current drain, potential detection problems, and an increased incidence of hardware and software design problems. At the time of insertion, the amount of energy required for various AICD functions, such as the defibrillation threshold, is determined for any given patient, and output and sensing functions can be adjusted by reprogramming as needed.

In 2012 the Food and Drug Administration approved the use of a subcutaneous defibrillator. This consists of a subcutaneous lead with a shock coil flanked by two sensing electrodes; this coil is tunneled along the parasternal margin from the pulse generator that is implanted in a subcutaneous pocket. It
CARDIAC PROCEDURES

Left Ventricular Assist Devices

As the management of heart failure continues to evolve, emergency physicians may encounter patients who have surgically implanted ventricular assist devices (VADs). VADs are implanted pumps used to assist a failing ventricle. Although they have been used in the right ventricle and for biventricular support, they are primarily used as a left ventricular assist device or LVAD. The first-generation of these devices utilized a pulsatile flow pump. The second and third devices provide continuous flow through the use of axial flow or centrifugal pumps. These devices are smaller and more durable, with higher energy efficiency than the first-generation models, which are no longer in use.

Initially used as a bridge to heart transplantation, they are now also being used as a bridge to recovery of heart function (such as in patients with myocarditis) and for permanent (destination) therapy. The LVAD consists of an inflow cannula that is placed in the apical part of the ventricle and allows blood to flow to a pump. The pump is attached to the outflow cannula, which is placed in the ascending aorta. The device is connected through a percutaneous lead or driveline to batteries located outside the body. Figs. 13.5 and 13.6 demonstrate the components of a left ventricular device.

INDICATIONS FOR PLACEMENT OF IMPLANTABLE PACEMAKERS AND AICDS

The most common indication for placement of a cardiac pacemaker is for the treatment of symptomatic bradyarrhythmias.

is placed by anatomic landmarks and as such no fluoroscopy is required. It delivers an 80-J biphasic shock across the precordium. Fig. 13.4 is a radiograph of a patient with a subcutaneous defibrillator.

The subcutaneous implantable device may be an alternative to patients who have difficult vascular access or who are at risk for transvenous complications. These could include the pediatric populations, patients with congenital disease, complicated vascular anatomy, patients at high risk for infection, or patients on dialysis. The notable limitations of the subcutaneous internal defibrillator when compared with conventional devices is the inability to provide long-term bradycardia pacing or antitachycardic pacing. Patients that need resynchronization therapy are also unsuitable candidates.

**Figure 13.3** Diagrammatic demonstration of various automatic implantable cardioverter defibrillator (AICD) configurations. **A,** Single-chamber AICD. **B,** Dual-chamber AICD. **C,** Biventricular AICD.

**Figure 13.4** Chest radiograph demonstrating a subcutaneous defibrillator.
manifest low-grade AV blocks and left bundle branch block. The resultant delay in left ventricular conduction often results in corresponding biomechanical delays in ventricular contraction, which in turn cause a further decrement in cardiac output and worsening congestive heart failure. Such prolongation may occur in as many as 33% of patients with advanced heart failure. This electromechanical “dyssynchrony” has been associated with increased risk for sudden cardiac death.

CRT comprises atrial-synchronized, biventricular pacing, which overcomes the atrial and ventricular blocks while optimizing both preload and LV EF. Clinical trials and systematic reviews have confirmed the efficacy of CRT, with decrements in mortality of 22% to 30%, in addition to improved LV EF and quality of life. It is therefore likely that emergency physicians will see the CRT configuration with increasing frequency in patients with implanted pacemakers and AICDs.

AICD technology is used principally for both primary and secondary prevention in patients at risk for sudden death. Primary prevention is an attempt to avoid a potentially malignant ventricular arrhythmia in patients identified as being at high risk. Secondary prevention is for patients who have already had a ventricular arrhythmia and are at risk for further events. In addition, AICDs are implanted for a number of other congenital or familial cardiac conditions.

**PACEMAKER AND AICD RESPONSE TO MAGNET PLACEMENT**

In the clinical setting, placement of a magnet over the pulse generator of a pacemaker is a technique that can be used either diagnostically or therapeutically by the emergency clinician. Application of a magnet is used to suspend antitachyarrhythmia detection and therapy of an AICD without changing pacing mode, or to produce asynchronous (fixed rate) pacing of a pacemaker. Hence, a magnet may change a demand pacemaker (synchronous) to a fixed rate (asynchronous) pacemaker. However, it is important to note that each pacemaker is programmed to respond in a specific fashion to a magnet as determined by the manufacturer. The response to magnet placement may not only vary by manufacturer, but also by model and by the particular mode in which the pacemaker is currently operating. Some devices (Boston Scientific, St. Jude, Biotronic devices) may be programmed to disable the magnet function and hence not respond to magnet application. Some leadless pacemakers do not initiate asynchronous pacing in response to a magnet application. Some manufacturers set the asynchronous (fixed rate) baseline pacing rate in a range approximating 70 beats/min. An indicator of aging of the pacemaker and weakening of the battery is that this asynchronous baseline pacing rate will decrease over time as the battery approaches the point at which replacement is required.

Keeping these provisions in mind, there are standard responses that the provider might expect to see in most circumstances. In the case of single-chamber ventricular pacemakers, the response will most likely be asynchronous pacing (V00). In the case of dual-chamber pacemakers, placement of a magnet usually results in dual-chamber asynchronous pacing (D00). In either case, it is important for the clinician to note that placement of a magnet over the pacemaker pulse generator will not turn the pacemaker off.

Roughly 50% of pacemakers are placed in such patients for the treatment of sinus node dysfunction (sick sinus syndrome). Other diagnoses include symptomatic sinus bradycardia, atrial fibrillation with a slow ventricular response, high-grade AV block (including Mobitz type II and third-degree AV block), tachycardia-bradycardia syndrome, chronotropic incompetence, and selected prolonged QT syndromes. Though not classified as absolute indications, pacemakers are sometimes placed for the treatment of severe refractory neurocardiogenic syncope, paroxysmal atrial fibrillation, and hypertrophic or dilated cardiomyopathy.

In recent years, CRT has emerged as a primary approach for patients with severe diastolic dysfunction and a low left ventricular ejection fraction (LVEF). Commonly such patients manifest low-grade AV blocks and left bundle branch block. The resultant delay in left ventricular conduction often results in corresponding biomechanical delays in ventricular contraction, which in turn cause a further decrement in cardiac output and worsening congestive heart failure. Such prolongation may occur in as many as 33% of patients with advanced heart failure. This electromechanical “dyssynchrony” has been associated with increased risk for sudden cardiac death.

CRT comprises atrial-synchronized, biventricular pacemaking, which overcomes the atrial and ventricular blocks while optimizing both preload and L VEF. Clinical trials and systematic reviews have confirmed the efficacy of CRT, with decrements in mortality of 22% to 30%, in addition to improved L VEF and quality of life. It is therefore likely that emergency physicians will see the CRT configuration with increasing frequency in patients with implanted pacemakers and AICDs.

AICD technology is used principally for both primary and secondary prevention in patients at risk for sudden death. Primary prevention is an attempt to avoid a potentially malignant ventricular arrhythmia in patients identified as being at high risk. Secondary prevention is for patients who have already had a ventricular arrhythmia and are at risk for further events. In addition, AICDs are implanted for a number of other congenital or familial cardiac conditions.
standard advanced cardiac life support (ACLS) protocols should be used to stabilize the patient.

Of further note, in some obese patients or those with heavily developed chest wall musculature, the magnetic field emitted by a single magnet device may not be strong enough to elicit the desired effect on the implanted device. In such cases the clinician may find greater efficacy by using two magnets, one on top of the other.

**CLINICAL EVALUATION OF PATIENTS WITH IMPLANTED PACEMAKERS AND AICDS**

**History**

Although patients who go to the ED because of implantable pacemaker–related issues may have one or more of several complaints, those with AICD-related issues are generally seen because their device has discharged. They will often describe a sensation of being kicked or punched in the chest, and the sensation is not subtle. In fact, some patients live in fear of the shock after having experienced it previously, and this is one reason for removal of the device. Ask the patient about the number of discharges and associated symptoms, including
doubt, an arterial line would be the most accurate method to identify MAP. MAP is usually kept between 70 to 80. Pulse oximetry is also unreliable and a true measurement should be obtained with an arterial blood gas or arterial line.

**Radiography**

The imaging modality of choice, at least initially, in a patient with complaints related to an implanted pacemaker or AICD is the plain chest radiograph. If the patient’s stability is in question, obtain a portable anteroposterior film. In addition to the standard cardiac, pulmonary, vascular, and skeletal evaluation, this study will probably confirm the location of the pulse generator case, as well as the current location of any leads. Survey the radiograph for evidence of lead fracture or displacement (see Fig. 13.1)

**Physical Examination**

First, assess for airway patency, adequate ventilation, and cardiovascular status. The patient’s mental status may also be an important clue to the severity of the symptoms. Perform an appropriate physical examination with emphasis on examining the heart and lung to seek murmurs, pericardial friction rubs, and evidence of pulmonary effusion or other abnormalities. Inspect the pacemaker or AICD site for erythema or edema. Palpate the skin for evidence of obvious lead abnormalities. Examine the extremities for evidence of edema or erythema.

There are several unique characteristics to consider in patients with ventricular assist devices. On physical exam, the provider should listen to the chest to determine if an audible “hum” is present. Patients lack a palpable pulse given the device’s continuous flow and because their cardiac function at baseline is poor. The blood pressure should be obtained by using a Doppler on either the radial or brachial artery. Inflate the cuff until you can’t hear flow. The cuff pressure should then be released, listening for the presence of flow.9 There is some degree of controversy about whether this is mean arterial pressure (MAP) or systolic pressure, however.10,11 When in doubt, an arterial line would be the most accurate method to identify MAP. MAP is usually kept between 70 to 80. Pulse oximetry is also unreliable and a true measurement should be obtained with an arterial blood gas or arterial line.

**Electrocardiography**

Perform an ECG to seek evidence of pacing, ectopy, and ST-segment or T-wave abnormalities. A comparison ECG may be helpful. If an AICD shock has been delivered, the shock itself can cause transient electrocardiographic changes, and waiting for several minutes to repeat the ECG may identify whether the changes are caused by the discharge or an ongoing disease process.

**Cardiopulmonary Resuscitation, ACLS Interventions, and External Cardiac Defibrillation in Patients With Implanted Pacemakers or AICDs or LVADS**

In general, ACLS interventions may be performed safely and effectively in patients with pacemakers and AICDs when indicated. Cardiopulmonary resuscitation (CPR) can usually be performed in standard fashion. If an AICD is present,
rescuers may notice mild electrical shocks while performing CPR; these shocks are harmless to the rescuer. If the AICD shocks are impeding rescuer performance of CPR, or if supraventricular tachycardias are noted during resuscitation, disable the AICD by applying a magnet over the corner of the device from which the leads emerge. This location is generally found easily by palpation but may be located blindly by slowly relocating the magnet until AICD activity ceases.

External cardiac defibrillation may be performed safely in patients with pacemakers and AICDs with the standard expected efficacy; however, it is recommended that external paddles or defibrillator pads be placed at a location approximately 10 cm distant from the pulse generator if possible. A transcutaneous cardiac pacemaker may also be used in similar fashion with a recommendation that the pacing pads be placed in anatomically appropriate locations but preferably at a distance of 10 cm from the pulse generator.

Placing the external defibrillation or transcutaneous pacemaker pads in an anteroposterior configuration is advised because this configuration may circumvent energy shunting and shielding. Every attempt should be made to avoid application of the defibrillators directly over the device. Use of the lowest possible energy setting for cardioversion or defibrillation is recommended. If available, biphasic cardioverter defibrillators are further suggested. In the event of successful resuscitation and return of spontaneous circulation, the pacemaker or AICD should be interrogated expeditiously by a cardiologist or electrophysiologist to ensure that no damage was sustained as a result of the resuscitation effort.

Regarding pharmacologic adjuncts, amiodarone has been reported to be more effective for the treatment of potentially lethal arrhythmias in the setting of implanted devices. Antiarrhythmic medications may be required for a resistant malignant rhythm when the AICD is functioning properly but the arrhythmia persists (Fig. 13.10G).

Several additional considerations are unique to the setting of ACLS in patients with a pacemaker or AICD. In cases of acute myocardial infarction involving areas of the myocardium in contact with the pacemaker leads, the implanted pacemaker may experience operative failure. Therefore, maintain a high level of suspicion for the potential requirement for supplemental transcutaneous or transvenous cardiac pacing.

LVAD patients that go into cardiac arrest offer some unique challenges. Ensure that the machine is connected, the battery charged, and the machine has an audible hum. The lack of peripheral pulses also make the use of ACLS problematic. In the case of arrest, follow the standard Advanced Cardiac Life Support algorithms. There is concern that standard CPR may

**Figure 13.9** Heart-Mate II model of left ventricular assist device (LVAD) appearance in A, posterior–anterior and B, lateral chest X-ray, C, computerized tomography, and D, ultrasound examination. ICD, Implantable cardioverter defibrillator; LV, left ventricle; LA, left atrium.
Drug Therapy for AICD-Resistant Ventricular Tachycardia

- Procainamide, 30 mg/min or 1 g over 1 hr (max, 17 mg/kg), and/or:
- Amiodarone, 150 mg over 1–3 min, repeat prn, followed by an infusion at 1 mg/min, and/or:
- Lidocaine, 1–1.5 mg/kg, repeat, max of 3 mg/kg, and/or:
- Metoprolol, 3–5 mg IV every 5–10 min
- If torsades de pointes: magnesium, 2–4 g IV

Figure 13.10 A, Patient with an automatic implantable cardioverter defibrillator (AICD) who experienced multiple discharges from an AICD because of recurrent ventricular tachycardia (VT). B, Electrocardiogram (ECG) of VT in the presence of an underlying tachycardia as the possible cause. C, A rhythm strip demonstrates persistent polymorphic (torsades de pointes) VT. D, After attempts at overdrive pacing failed, the AICD appropriately discharges with temporary termination of VT. E, Interrogation of the device uncovers a history of more than 20 episodes of recurrent VT within 30 minutes. This patient was terrified of a subsequent shock because he could sense the VT and knew of the impending shock. In addition to antiarrhythmic medication, a quiet dark room and aggressive sedation are suggested to reduce catecholamine levels. F, ECG after medical therapy with multiple medications, including magnesium. Metoprolol seemed to be the deciding factor in terminating the VT in this case. G, Current suggested medical therapy for VT in the presence of an appropriately functioning AICD.
dislodge the cannula or damage the unit’s outflow conduit, but there have also been reports that this is not as harmful as previously thought. There was also a report of an emergency physician who replaced the driveline connections in a patient who had cut the wires resulting in subsequent resumption of the pump function.

In the postresuscitation setting involving a patient with an implanted pacemaker or AICD, it is important to maintain a higher index of suspicion for device lead fracture or disruption resulting from CPR. Finally, in the postresuscitation phase the clinician should closely watch for the development of pneumothorax, hemotorax, pericardial effusion, or other aforementioned pathophysiologic processes that could adversely affect the function of the implanted device.

### COMPLICATIONS AND MALFUNCTION OF IMPLANTED PACEMAKERS

Complications associated with pacemakers are listed in Box 13.1. In addition, patients with previously implanted and otherwise stable pacemakers may experience complications related to direct or indirect trauma affecting the pulse generator or leads. Major complications of pacemaker placement or those caused by subsequent injuries that the emergency clinician might encounter include: local or systemic infections resulting from pacemaker placement, thromboembolic event, pneumothorax or hemothorax, pericarditis, air embolism, localized hematoma interfering with pacemaker operation or sensing, lead dislodgement, cardiac perforation, hemopericardium with possible progression to cardiac tamponade, and development of the phenomenon known as pacemaker syndrome. This condition is often seen in patients with single-chamber ventricular pacemakers who have an underlying component of congestive heart failure. It is believed to be a consequence of the loss of AV synchrony resulting from the ventricular pacing, and may be manifested as vertigo, syncope, hypotension, and signs specific to the exacerbation of congestive heart failure.

In addition to the complications associated with initial pacemaker placement, malfunctions of these devices may occur in the short-, intermediate-, and long-term phases of their functional life spans. Most malfunctions result from one or a combination of three primary problems: failure of the pace generator to provide output, failure to capture, or failure to sense the intrinsic cardiac rhythm.

### Pacemaker Output Failure

Pacemaker generator output failure is present when no pacing “spike” is noted on the ECG despite an indication for pacing. This condition may result from battery failure, fracture or loss of insulation in the pacer leads, oversensing of extraneous signals resulting in pacer inhibition, disconnection of the leads from the pacer generator, or in the case of a dual-chamber pacer, erroneous sensing of the pacemaker’s atrial signal by a ventricular sensor. The latter phenomenon is commonly referred to as crosstalk.

Given that the reason for pacemaker implantation in most cases was for the treatment of an underlying bradycardia condition, initial clinical management of patients with some degree of pacer output failure will usually focus on pharma-

**Box 13.1 Complications of Permanent Pacemakers**

- **FAILURE TO PACE (NO PACEMAKER ACTIVITY PRESENT)**
  - Lead fracture
  - Lead disconnection
  - Battery depletion
  - Component failure
  - Oversensing
  - External interference

- **FAILURE TO SENSE (CONSTANT PACEMAKER SPIKES DESPITE ONGOING INTRINSIC CARDIAC ELECTRICAL ACTIVITY)**
  - Lead dislodgement
  - Lead fracture
  - Fibrosis around the tip of the lead
  - Battery depletion
  - Pacer in asynchronous mode
  - External interference
  - Low-amplitude intracardiac signal

- **FAILURE TO CAPTURE (PACEMAKER SPIKES BUT NO SUBSEQUENT CARDIAC ACTIVITY)**
  - Lead dislodgement, including perforation
  - Lead fracture
  - Lead disconnection
  - Poor lead position
  - Fibrosis around the tip of the lead
  - Battery depletion
  - Metabolic abnormalities
  - Medications

- **INAPPROPRIATE PACEMAKER RATE (RUNAWAY PACEMAKER)**
  - Pacemaker reentrant tachycardia
  - Resetting from external interference
  - Battery depletion

- **OTHER**
  - Infections: pocket, wires
  - Lead displacement: cardiac perforation, tamponade, pericarditis, vascular perforation
  - Vascular complications: thrombosis, superior vena cava syndrome
  - Psychiatric: anxiety, panic attacks

Cardiac procedures

Pacemaker generator output failure is present when no pacing “spike” is noted on the ECG despite an indication for pacing. This condition may result from battery failure, fracture or loss of insulation in the pacer leads, oversensing of extraneous signals resulting in pacer inhibition, disconnection of the leads from the pacer generator, or in the case of a dual-chamber pacer, erroneous sensing of the pacemaker’s atrial signal by a ventricular sensor. The latter phenomenon is commonly referred to as crosstalk.

Given that the reason for pacemaker implantation in most cases was for the treatment of an underlying bradycardia condition, initial clinical management of patients with some degree of pacer output failure will usually focus on pharmaco-
Failure to Capture

In the case of failure to capture, pacemaker spikes are present on the ECG. However, some or all the spikes are not followed by atrial or ventricular complexes as appropriate for the pacemaker model in question. Failure to capture may result from: deterioration of lead insulation; fracture or dislodgement of the leads; electrolyte disturbances, including hyperkalemia or hypocalcemia; a new condition requiring an elevated pacing threshold; acid-base disturbance; direct damage to the myocardium, which is in contact with the pacer lead’s tip (such as myocardial infarction or direct trauma); or dysfunction of the microcircuitry of the pulse generator. In addition, flecainide, a class IC antiarrhythmic medication, has been identified as an acute cause of the rise in ventricular capture thresholds in patients with implanted pacemakers. Likewise, all class I antiarrhythmic agents (sodium channel antagonists) may affect pacer capture thresholds and should therefore be identified as potential etiologic agents in patients suffering failure to capture.

Failure to Sense

Failure of an implanted cardiac pacemaker to sense the patient’s intrinsic cardiac rhythm may be subdivided into conditions related to oversensing or undersensing. Oversensing is present when the pacemaker erroneously identifies extrinsic electrical signals, such as those from skeletal muscle potentials or electromagnetic interference (EMI), and is inhibited from delivering an appropriate pacemaker pulse. For additional information on extrinsic EMI, refer to the section on Electromagnetic Interference and Implantable Devices later in this chapter.

Management of pacemaker failure to sense will be driven largely by the patient’s clinical condition. A prudent clinician will order cardiovascular monitoring, intravenous access, and a portable chest radiograph, with additional measures as dictated by the patient’s condition. In the event of symptomatic bradycardia, placement of a magnet over the pacemaker pulse generator may be indicated because this maneuver will usually place the pacemaker in an asynchronous ventricular pacing mode and thereby restore a stable and regular paced ventricular rhythm while a consulting cardiologist or electrophysiologist is summoned.

Undersensing is said to occur when the pacemaker fails to identify intrinsic cardiac depolarization and delivers a pacing signal. This condition may result from damage or dislodgement of the pacemaker leads, myocardial infarction, direct cardiac trauma, failure of the pacemaker’s power source, and even the application of a magnet. Initial ED management of this condition is similar to that performed in the case of oversensing. Magnet placement may also be appropriate in this setting because it will restore a stable, regular, and normal cardiac rhythm until the pacemaker and its leads may be more thoroughly examined.

Runaway Pacemaker Syndrome

This condition is seen almost exclusively in older pacemaker models particularly as they approach the end of their battery life, or when the pulse generator is damaged by exposure to radiation or direct impact. The hallmark of runaway pacemaker syndrome is uncontrolled tachycardia resulting in ventricular rates approaching 300 to 400 beats/min. In addition to initial attempts at stabilization, magnet placement may be attempted. However, this is often ineffective. If the patient is hemodynamically unstable in the setting of runaway pacemaker syndrome, it may be necessary to disconnect the pulse generator. To do this, identify the location of the pacer leads by physical examination or a portable chest radiograph. Dissect through the skin and subcutaneous tissue and then sever the leads with a wire cutter or similar tool.

Pacemaker-Mediated Tachycardia

In some circumstances patients with implanted pacemakers or AICDs may be seen in the ED because of symptomatic tachycardias resulting specifically as a complication of their pacemaker devices. This condition, referred to as pacemaker-mediated tachycardia, and alternatively as pacemaker-induced tachycardia, results most often from one of three clinical scenarios. In patients with dual-chamber pacemakers, one of the pacemaker leads may function as a pathway for either anterograde or retrograde conduction and result in what is referred to as endless loop syndrome and thus a tachycardiac arrhythmia, which may often become hemodynamically unstable. In general, patients suffering from endless loop syndrome will not have a ventricular rate greater than the maximum tracking rate of the pacemaker device. Consequently this condition will rarely be manifested as hemodynamic instability. One caveat, however, is that patients with underlying coronary artery disease and endless loop syndrome may experience coronary ischemia. In such a case or in a patient who is hemodynamically unstable because of the increased ventricular rate, application of a magnet will terminate the syndrome in most cases. Once stabilized, this condition may be prevented or at least mitigated by reprogramming of the pacemaker’s atrial sensor lead by an electrophysiologist.

A second scenario in patients with dual-chamber pacemakers occurs under the circumstance in which the patient experiences an intrinsic atrial tachycardia, at which point the implanted pacemaker begins to continuously discharge at its maximum preprogrammed ventricular rate. This condition may continue until the underlying atrial tachycardia is terminated by intervention.

A third instance of pacemaker-mediated tachycardia occurs in patients with AICD units that have backup antibradyarrhythmia pacing capability. It appears that if this pacemaker feature is switched on in such patients and an ectopic ventricular stimulus is delivered after a sudden pause in the intrinsic ventricular depolarization cycle, a ventricular tachyarrhythmia may be triggered.

DIAGNOSIS OF ACUTE MYOCARDIAL INFARCTION IN THE PRESENCE OF A PACED CARDIAC RHYTHM

Patients undergoing active ventricular pacing from an implanted pacemaker device will normally have ECGs that resemble a left bundle branch block pattern. As a result, electrocardiographic diagnosis of acute ischemic changes is equally challenging in both populations. Sgarbossa and colleagues published a series of criteria in 1996 that offer some utility in the interpretation of ECGs in patients with active ventricular pacing in whom acute coronary syndromes are suspected. These criteria are depicted in Table 13.2.
**TABLE 13.2** Criteria for Electrocardiographic Diagnosis of Acute Myocardial Infarction in the Setting of a Ventricular Paced Rhythm

<table>
<thead>
<tr>
<th>Electrocardiographic Criterion</th>
<th>Sensitivity (%)</th>
<th>Sensitivity (%)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Discordant ST-segment elevation &gt;5 mm</td>
<td>53</td>
<td>88</td>
<td>0.025</td>
</tr>
<tr>
<td>Concordant ST-segment elevation &gt;1 mm</td>
<td>18</td>
<td>94</td>
<td>NS</td>
</tr>
<tr>
<td>ST-segment depression &gt;1 mm in precordial leads V1–V3</td>
<td>2</td>
<td>82</td>
<td>NS</td>
</tr>
</tbody>
</table>

NS, Not significant.

**AUTOMATIC IMPLANTABLE CARDIOVERTER DEFIBRILLATORS—UNIQUE MALFUNCTIONS**

Issues with sensing problems, lead migration, and battery failure are similar to pacemaker complications, and most occur within 3 months after implantation. There is some data that a dual chamber device is associated with a higher risk of device-related complications but a similar 1-year mortality compared to a single-chamber device. A large metaanalysis put the incidence of in-hospital adverse events between 2.8% to 3.6%. Complications related to the insertion include cardiac injury, lead dislodgement, infection, hemorrhage, and pneumothorax. A potential malfunction unique to AICDs is an inappropriate defibrillation or lack of defibrillation of the device.

The AICD may not terminate ventricular arrhythmias, which may or may not be the result of malfunction of the device. AICD malfunction may be a result of battery depletion, component failure, failure-to-sense, or lead malfunction. Failure to cardiovert or defibrillate occurring in the setting of a functioning AICD system may be caused by inappropriate cutoff rates, failure to satisfy multiple detection criteria, completion and exhaustion of therapies, and cross-inhibition by a separate pacemaker. The advent of AV or dual-chamber AICD devices has improved the sensitivity of detecting arrhythmias and thus prevents the delivery of inappropriate shocks.

Inappropriate AICD-delivered shocks occur in 20% to 25% of patients and are the most common adverse events observed in AICD patients. The main causes are atrial arrhythmias, sinus tachycardia, nonsustained ventricular tachycardia, lead fracture or EMI, or electrical storm. By definition this phenomenon occurs when three or more shocks are delivered in a 24-hour period, occurs in 10% to 20% of AICD patients, and constitutes a medical emergency. AICD patients who experience this phenomenon may have end-stage cardiac failure and increased long-term mortality. Specific causes of electrical storm are unclear, but it is associated with ventricular tachycardia in the setting of LVEF lower than 30%, and occurs more frequently in patients with demonstrated coronary artery disease, who have not as yet undergone revascularization procedures. Suggested initial treatment includes the administration of amiodarone and β-adrenergic antagonists to pharmacologically suppress the arrhythmias, and urgent cardiology consultation for possible pacemaker interrogation, overdrive pacing, or even catheter ablation.

The AICD may discharge inappropriately in response to rapid supraventricular rhythms such as atrial fibrillation, supraventricular tachycardia, or even sinus tachycardia. Multiple shocks may be a manifestation of inefficient termination of tachycardia, such as inappropriately low-energy delivery at the first shock, increased defibrillation thresholds, and migration or dislodgement of the defibrillation lead system, or failure of the defibrillator system. Shocks that occur every few minutes may suggest that recurring ventricular tachyarrhythmias are being terminated appropriately (see Fig. 13.10).

AICD discharges in the setting of chest pain may be a result of myocardial ischemia–induced tachyarrhythmias. As noted earlier, any electrocardiographic abnormalities noted immediately after shocks should be interpreted with caution because ST elevation or depression can occur immediately after a shock. If the patient receives shocks in association with chest pain, ischemia is suggested, but other causes, including hypokalemia, hypomagnesemia, drug-induced proarrhythmia, or drugs that can prolong the QT interval (such as phenothiazines), should also be considered as underlying causes.

In some settings the AICD may fail to sense sustained ventricular tachycardia or fibrillation. Such failure may be caused by an intrinsic arrhythmia rate below the programmed detection rate, usually as a result of concurrent pharmacologic therapy. If the patient is hemodynamically stable, it may be advantageous for the cardiologist to interrogate the pacer before initiating further antiarrhythmic therapy. If unsuccessful or if the patient is experiencing a nonperfusing ventricular arrhythmia, other pharmacologic interventions include procainamide or amiodarone. Box 13.2 depicts the outcome of AICD placement in a general population.

Subcutaneous defibrillators have some of the same complications as the venous AICDs including hematoma, infection, lead displacement, and inappropriate shocks.

**LVAD Complications**

In patients with LVAD, infection is the second most common cause of death after cardiac failure. Ventricular assist device infections can be divided into three categories: VAD-specific infections including the pump, pocket, and driveline; VAD-related
infections such as bloodstream infections, endocarditis, and mediastinitis; and non-VAD infections such as pneumonia, urinary tract infection, or cholecystitis. Infections can involve any portion of the LVAD including the surgical site, pocket, pump, and most commonly the driveline. The patient may have nonspecific symptoms, or have fever, localized erythema, or wound drainage. Infections occur most commonly within the first three months of device implantation with the risk of severe infection and sepsis peak at 1 month. Most infections are caused by gram-positive organisms such as *Staphylococcus aureus*. *Enterococcus* is also common, as are gram-negative organisms, most commonly *Pseudomonas*. Treatment includes broad spectrum antibiotics, though localized infection may be treated for gram-positives alone. LVAD-associated endocarditis has a high mortality rate, and can range from minimal symptoms to fever and systemic emboli in a presentation similar to prosthesis valve endocarditis.

Given that most patients are on both antiplatelet therapy and anticoagulant therapy, bleeding is another frequent adverse event in LVAD patients. The increase in bleeding susceptibility is thought to be related to the development of acquired von Willebrand syndrome secondary to polymer deformation by the second- and third-generation LVADs resulting in deficiency of von Willebrand factor. Second- and third-generation LVADs have also led to nonsurgical gastrointestinal bleeding secondary to arterial venous malformations. It is thought that reduced pulse pressure from continuous infusion results in hypoperfusion of the gastrointestinal mucosa and neovascularization that, with friable vessels, is prone to bleeding. These patients need endoscopy and possibly agent reversal with the use of fresh frozen plasma or prothrombin complex concentrates.

LVAD thrombosis is a potentially devastating complication occurring in 2% to 13% of continuous flow LVAD patients. Thrombosis can occur throughout the device resulting in thrombotic stroke, thromboembolism, LVAD malfunction, and failure, resulting in life-threatening hemodynamic impairment, cardiogenic shock, and death. The patient may have very mild and nonspecific symptoms such as tea-colored urine to decompensated heart failure. Useful laboratory tests include coagulation studies to check for patient compliance, and tests for hemolysis to include a urinalysis and lactate dehydrogenase (a level of 3 times normal indicates hemolysis). Chest x-ray may reveal malposition of the outflow cannula and chest CT angiography may diagnose inflow malposition or outflow obstruction. These patients need to be urgently transferred to a center with mechanical support because the definitive therapy is removal of the device and cardiac transplantation.

Dysrhythmias are a common problem in patients with advanced stages of cardiomyopathy, but they are also at risk from placement of the device. Atrial arrhythmias should be treated in the same manner as the heart failure population by initially controlling the rate with β-blockers. LVAD patients are also at risk for ventricular arrhythmias and most of them will have an AICD in place before their placement. Patients with persistent ventricular arrhythmias, in spite of the AICD discharges, may require IV β-blockers in addition to reprogramming the device.

Resuscitation of these patients can be challenging given their sensitivity to both preload and afterload. A decrease in systemic volume may result in collapse of the left ventricular wall, which may crowd the inflow cannula leading to obstruction of blood flow (also known as a suction effect), and subsequent clinical deterioration. Uncontrolled increase in MAP higher than 90 mm may reduce LVAD outflow, cardiac output, and distal perfusion. These patients are at risk for cerebral vascular events and should be treated promptly with afterload-reducing medications.

### BOX 13.3 Method for Inactivation of an Automatic Implantable Cardioverter Defibrillator

1. Determine the orientation of the device in the abdominal pocket radiographically or by palpation.
2. Place a ring magnet over the upper right-hand corner of the device.
3. A beeping tone will sound that corresponds to the sensing of QRS complexes.
4. Leave the magnet in place for at least 30 seconds.
5. When the beeping changes to a continuous tone, the device is inactivated.
6. Remove the magnet.

#### Use of a Magnet for AICD

A patient who is experiencing inappropriate AICD discharges in the ED can be treated by inactivation of the device with a magnet, similar to the approach described earlier for pacemaker patients. If the patient is experiencing recurrent rhythms that require activation of the AICD, do not inactivate the device because it is functioning as required.

#### Technique

The method for inactivating an AICD device is outlined in Box 13.3. The orientation of the device in the abdominal pocket should be determined, with the lead connections normally being cephalad. A ring magnet is then placed over the corner adjacent to the lead connections (usually the upper right-hand corner of the device). A series of beeping tones will sound that correspond to the sensed QRS complexes. In the absence of organized QRS activity, random beeps will sound. When the magnet is left in place for 30 seconds, a continuous beep is heard. This indicates that the AICD is inactivated. The magnet should then be removed, and the AICD will remain inactivated. The AICD may be reactivated by applying the magnet for 30 seconds and removing it when the steady beep changes to intermittent beeping. Note that unlike a pacemaker, in which a magnet will turn a demand pacemaker to a fixed-rate pacemaker, a magnet will not affect the pacing function of an AICD.

### “Twiddler’s Syndrome”

In some cases, patients with implantable pacemakers choose to “twiddle” with the device: they manipulate the pulse generator case within its physiologic pocket under the skin in the chest. Note that a generator may also be placed in the abdominal wall (Fig. 13.11). This practice of twiddling may result in coiling, dislodgement, or disconnection of the pacemaker leads (Fig. 13.12). It may even lead to actual displacement of the pulse generator case. At a minimum, it may result in physical discomfort and may, in fact, precipitate cardiac arrhythmias or other complications local to the site of pacemaker placement.
After initial stabilization of the complaint, these patients may require readjustment or replacement of their pacemaker devices. As part of their care, pacemaker patients should be educated to avoid manipulating their pacemakers.

**Mental Health Issues Related to Implanted Pacemakers and AICDs**

Patients with these devices may manifest a number of anxiety-related complications including adjustment disorder, panic attacks, depression, imaginary shock; and defibrillator dependence, abuse, or withdrawal. These patients may benefit from psychiatric referral either as an outpatient or as part of the admission evaluation if applicable. The conditions may be severe enough that the device is removed by patient request.

**Implantable Pacemaker and AICD Recalls**

Since 1990, more than 60 device or lead advisories have been announced, affecting more than one million device patients. These advisories were issued as a result of unanticipated failure of devices identified after release of the product and widespread clinical use. The decision to remove a device is complex, and there has been difficulty reaching consensus on the optimal management of patients with these recalled devices. The decision to replace them should be multifactorial and take into consideration the estimated malfunction rate of the device, anticipated consequences of failure of the device, the individual center’s procedural risk for complications resulting from generator replacement, and patient preferences and desired level of risk tolerance.

**Electromagnetic Interference and Implantable Devices**

Given the plethora of new technologies there is always concern about the interaction of EMI with pacemakers and AICDs. The sources of EMI comprise a significant spectrum and may involve radiated and conducted sources.

The most common response of implanted devices to EMI is inappropriate inhibition or triggering of pacemaker stimuli, reversion to asynchronous pacing, and spurious detection of tachyarrhythmias by the AICD. Reprogramming of operating parameters and permanent damage to the circuitry of the device or the electrode-tissue interface can also occur but is much less frequent. Additional adverse effects that may occur include inhibition of bradycardia pacing, inadvertent delivery of a shock, or antitachycardia pacing.

The use of hermetic shielding in metal cases, filtering, interference rejection circuits, and bipolar sensing have helped mitigate most of this interference. Nonetheless, the clinician should be familiar with common sources of EMI that may affect pacemakers and AICDs. Several specialty societies have published position statements emphasizing patient specific selection and the need to have the study at an experienced center with close coordination between cardiology and radiology services.

Several caveats will help to avoid the deleterious effects of EMI on implantable devices. Cell phones should not be kept in a pocket over the device. When in use they should be held at least 6 inches away from the device. When hands-free headphones are used, these devices should be placed in the ear opposite the implanted pacemaker or AICD. In addition, patients should be advised to not linger in theft detection areas or airport metal detectors. Box 13.4 identifies different devices and the corresponding potential EMI that can occur with pacemakers and AICDs.

**Out-of-Hospital Discharge of AICD**

Patients are told to adhere to standard advice defining an appropriate response to out-of-hospital discharges of the AICD (Box 13.5). However, many come to the ED for evaluation after every shock. There is no standard ED intervention mandated by historical information, and clinical decisions are made on an individual basis corresponding to the current scenario. Options include prolonged ED observation, consultation, cardiac monitoring, laboratory testing (such as electrolytes and cardiac enzymes), or interrogation.
BOX 13.4 Sources of Electromagnetic Interference and Their Potential Effects on Implanted Pacemakers and Automatic Implantable Cardioverter Defibrillators

**GENERALLY THOUGHT TO BE SAFE WITH EMI**
- Copy machines
- Electric blankets
- Household appliances (microwaves, washer/dryer)
- DVD/CD players, TVs
- Personal computers
- Remote controls
- Heating pads

**SOME EVIDENCE OF INTERACTION**
- Cell phones
- Induction ovens
- Power toothbrushes
- Battery-powered, cordless power tools
- Arc welding equipment
- Chain saws
- Drills
- Hedge trimmers
- Lawn mowers
- Leaf blowers
- Snow blowers
- High-voltage lines
- Theft detection systems
- Airport scanners/metal detectors

**AVOID BECAUSE OF INTERACTION**
- Electrolysis
- MRI
- Jackhammers

**MEDICAL DEVICES REQUIRING CAUTION**
- Electrocautery (especially unipolar) equipment
- High-energy radiation sources
- TENS units
- MRI scanners
- Body fat–measuring scales
- Diathermy equipment
- Electrolysis equipment
- Spinal cord stimulators
- Direct current external cardioversion/defibrillation equipment
- Radiofrequency catheter ablation equipment
- Lithotripsy equipment

**SAFE MEDICAL DEVICES**
- CT scanners
- Dental drills
- Diagnostic x-ray machines
- Electrocardiography equipment
- Ultrasound equipment

CT, Computed tomography; EMI, electromagnetic interference; MRI, magnetic resonance imaging; TENS, transcutaneous electrical nerve stimulation.

BOX 13.5 Follow-Up of Patients With an AICD

Follow-up assessments of patients with an AICD are made on a routine basis, every 3 to 6 months, and when discharge of the device occurs. Analysis of any previous clinical event and testing of defibrillation function are readily accomplished. Internet-based remote follow-up systems may replace some office follow-up. Follow-up can occur remotely with vendor-specific equipment to interrogate and upload data. Remote follow-up, however, permits only device interrogation and retrieval of diagnostic data, not threshold testing or reprogramming. Device interrogation includes the following:

- Determination of pacing and sensing thresholds.
- Analysis of recorded episodes of arrhythmia detection and AICD activation, including pacing and delivered shocks. Data include the date and time of each episode and a stored electrocardiogram from the event.
- Battery status.

**DEFIBRILLATOR DISCHARGE**
An appropriate shock is delivered in approximately 50% of patients by 2 years after implantation. Patients may not sense antitachycardiac (overdrive) pacing to terminate arrhythmias. Not all episodes of defibrillator discharge require immediate medical evaluation, although many patients go to the ED immediately. Patients with a first shock may be seen on an urgent or elective basis to ascertain the specifics of the event and to determine whether the device is functioning properly. Discharges that are accompanied by changes in cognition (syncope, seizure, or loss of consciousness) require ED evaluation.

Per guidelines, patients who have had a single AICD discharge with immediate return to baseline clinical status and no associated symptoms (e.g., chest pain, shortness of breath, light-headedness) may have the device interrogated within 1 to 2 days. Delivery of frequent shocks or clusters of shocks is either appropriate (because of recurrent VT) or inappropriate (because of atrial fibrillation, supraventricular tachycardia, or device malfunction). Such patients generally require emergency evaluation and hospital admission to determine the cause. Additional therapy (such as an antiarrhythmic drug or catheter ablation) may be required.

AICD, Automatic implantable cardioverter defibrillator; ED, emergency department; VT, ventricular tachycardia.
Disposition Criteria

In most cases, patients seen in the ED with pacemaker complications or malfunctions will be symptomatic. Accordingly, and regardless of the clinical requirement for admission, they will probably require device interrogation and possible recalibration or replacement by a cardiologist. In most cases this will be accomplished during hospital admission.

With regard to AICD malfunctions and disposition, patients who have had a single shock and no other specific complaints or comorbidity can be discharged with follow-up in 24 to 48 hours. Patients with symptoms concerning for ischemia, potentially lethal arrhythmias, or symptomatic illness should be admitted and specialty consultation obtained expeditiously. Patients who have had multiple shocks will need admission for observation and interrogation of their AICD device. Interrogation reveals significant information about the device such as why it fired, the rhythm history, and an accurate assessment of the underlying problem.
REFERENCES


**INTRODUCTION**

The electrocardiogram (ECG) is a graphic recording of the electrical activity of the heart. The standard ECG is obtained by applying electrodes over the chest and limbs to record the electrical activity of the cardiac cycle. Developed over 100 years ago, the ECG remains the most important initial diagnostic tool for the evaluation of patients presenting with chest pain, dyspnea, syncope, cardiac arrest, and toxicologic exposures, among many other clinical symptoms.

Electrocardiography is performed widely throughout the health care field, including ambulances, ambulatory clinics, emergency departments (EDs), and in-patient hospital units. Standard electrocardiography machines are small, self-contained, and portable, thus allowing them to be used in virtually any setting.

**BACKGROUND**

In 1903, Dutch physiologist Willem Einthoven first published his recordings of the cardiac cycle with a new device, the string galvanometer, an early version of today's electrocardiogram. Although others had previously recorded cardiac electrical activity, Einthoven's instrument laid the basis for modern clinical electrocardiography. He used bipolar leads, established standards for recording rate and amplitude, and initially described the deflections using the now familiar P, Q, R, S, and T, descriptors.

Thomas Lewis visited Einthoven's laboratory and recognized the potential clinical utility of the electrocardiography machine. Lewis became the leading authority on electrocardiography in the early 1900s and was instrumental in the development and clinical application of this new technology. His electrocardiographic description of atrial fibrillation demonstrated the clinical potential of this technology. In the early 1930s, Francis Wood and Charles Wolferth first reported the use of ECGs to differentiate cardiac and noncardiac chest pain. Along with Frank Wilson, their work also led to development of the unipolar “exploring” electrode, which measured electrical activity anywhere in the body with a zero-potential central terminal as a reference. These electrodes could be placed directly over the chest and formed the basis for the standard precordial leads.

In 1938, the American Heart Association in conjunction with the Cardiac Society of Great Britain established the standard six precordial chest lead positions (V1 to V6). These precordial leads, along with Einthoven’s original bipolar limb lead system (I, II, III) and the augmented unipolar limb leads developed by Emmanuel Goldberger (aVR, aVL, and aVF) in 1942, make up the standard 12-lead ECG used today.

**INDICATIONS**

The most frequent indication for electrocardiography in the ED is the presence of chest pain; other common complaint-based indications include dyspnea, syncope, and palpitations. Beyond complaint-based reasons for ECG performance, one can consider diagnostic investigations of suspected illness (cardiac arrest, acute coronary syndrome [ACS], pulmonary embolism, and toxic ingestion) and system considerations (“rule out myocardial infarction [MI]” protocol, sedations monitoring, admission purposes, and operative clearance). The ECG is used to help establish a diagnosis, select appropriate therapy, determine the response to treatment, assist in correct disposition of the patient, and help predict risk for both cardiovascular complications and death.

The initial 12-lead ECG obtained in the ED can be an important tool for determination of cardiovascular risk and accordingly, the choice of in-hospital admission location. Brush and coworkers classified the initial ECG into high- and low-risk groups. The low-risk electrocardiographic group had normal ECGs, nonspecific ST-T-wave changes, or no change when compared with a previous ECG. High-risk ECGs had significant abnormalities or confounding patterns, such as pathologic Q waves, ischemic ST-segment or T-wave changes, left ventricular hypertrophy, left bundle branch block, or ventricular paced rhythms. Patients with initial ECGs classified as low risk had a 14% incidence of acute myocardial infarction (AMI), a 0.6% incidence of life-threatening complications, and a 0% mortality rate. Patients with initial ECGs classified as high risk had a 42% incidence of AMI, a 14% incidence of life-threatening complications, and a 10% mortality rate. Another approach to risk prediction involves simple calculation of the number of electrocardiographic leads with ST-segment deviation (elevation or depression), with an increasing number of leads being associated with higher risk. Along similar lines, the clinician is also able to predict risk with a summation of the total millivolts of ST-segment deviation; once again, higher totals are associated with greater risk.

The limitations of the ECG must be recognized. The ECG is widely reported to have a sensitivity for AMI of only approximately 55%; in one study of 1000 patients with ischemic symptoms, sensitivity improved to 68% with serial ECGs and monitoring of ST-segment trends. In another series, the sensitivity of the ECG for AMI ranged from 43% to 65% over a 12-hour period after the onset of ischemic symptoms, yet the negative predictive value of a normal ECG (defined as normal or with nonspecific changes or isolated fascicular blocks) for AMI did not improve above 93% during this period. In a large series of more than 10,000 patients evaluated for ACS, 19 (2%) patients—ultimately found to have an AMI—were inappropriately discharged from the ED. A nonischemic ECG emerged as one of five risk factors for that inappropriate disposition decision.

**BASIC EQUIPMENT**

**The 12-Lead ECG**

Although there is variability depending on the workplace, most ECGs in use today are three-channel recorders with computer memory. Such multichannel systems, which record electrical events in several leads concurrently, offer advantages over the...
The standard ECG includes 12 leads derived from 10 electrodes placed on the patient; each is color-coded and represented by a two-character abbreviation (Table 14.1; see also Fig. 14.1B). Distinction is made between the terms lead and electrode with the former referring to the electrical view or perspective and the latter the actual wire attached to the patient. Placement of limb leads includes on the left and right arms (LA and RA, respectively) and on the left and right legs (LL and RL, respectively). Caution is advised as to the correct anatomic placement of the leads; incorrect lead placement produces electrocardiographic findings which can mimic illness with resultant unnecessary clinical interventions and further diagnostic testing; please refer to the section on Electrode Misplacement and Misconnection later in this chapter for a more complete discussion of this important issue.

**Standard 12 Leads**

The standard 12-lead ECG depicts cardiac electrical activity from 12 points of view, or leads, that can be grouped according to planar orientation. Six leads (I, II, III, aVR, aVL, and aVF) are oriented in the frontal, or coronal, plane, and are derived from the four limb electrodes. The six precordial leads (V₁, V₂, V₃, V₄, V₅, and V₆) are oriented in the horizontal, or transverse, plane, with each representing cardiac electrical activity from that perspective. Leads I, II, and III are termed limb leads and are bipolar in that they record the potential difference between two electrodes (Fig. 14.2). The fourth electrode located on the right leg serves as an electrical ground. The positive poles of these bipolar leads lie to the left and inferior, a position approximating the major vector forces of the normal heart. This early convention was established so that the tracing would feature primarily upright complexes. In contrast, augmented leads aVR, aVL, and aVF are unipolar leads with the positive electrode located at the respective extremities. These augmented leads serve to fill the “electrical gaps” between leads I, II, and III. Lead aVR stands alone with a polarity and resultant orientation opposite that of the other limb and augmented leads because of the fact that its positive electrode is located in the opposite direction (superior and to the right) of the major vector force of the normal heart (inferior

<table>
<thead>
<tr>
<th>LOCATION</th>
<th>NOTATION</th>
<th>COLOR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Right arm</td>
<td>RA</td>
<td>White</td>
</tr>
<tr>
<td>Left arm</td>
<td>LA</td>
<td>Black</td>
</tr>
<tr>
<td>Left leg</td>
<td>LL</td>
<td>Red</td>
</tr>
<tr>
<td>Right leg</td>
<td>RL</td>
<td>Green</td>
</tr>
<tr>
<td>Precordial leads</td>
<td>V₁</td>
<td>Brown/red</td>
</tr>
<tr>
<td></td>
<td>V₂</td>
<td>Brown/yellow</td>
</tr>
<tr>
<td></td>
<td>V₃</td>
<td>Brown/blue</td>
</tr>
<tr>
<td></td>
<td>V₄</td>
<td>Brown/green</td>
</tr>
<tr>
<td></td>
<td>V₅</td>
<td>Brown/orange</td>
</tr>
<tr>
<td></td>
<td>V₆</td>
<td>Brown/violet</td>
</tr>
</tbody>
</table>
and to the left); thus its complexes usually appear “opposite” those in most or all of the other leads.

Merging of the vector axes of the limb and augmented leads around a central axis yields a hexaxial system representing cardiac electrical activity in the frontal plane (Fig. 14.3). The six precordial leads, oriented in the horizontal plane, represent six unipolar electrodes with vector positivity oriented toward the chest surface and the central terminal of the hexaxial system serving as a negative pole. In contrast to the frontal-plane leads, the angles between each of the precordial leads in the horizontal plane are not equal. They can vary depending on electrode placement and body habitus.

**Electrode Placement**

The four limb electrodes are conventionally placed on the extremities as follows: RA on the right wrist, LA on the left wrist, RL on the right ankle, and LL on the left ankle. Electrodes may be affixed more proximally on the limbs if necessary (e.g., amputation, severe injuries), ideally with a notation made on the ECG. Others note that the electrodes may be placed on any part of the arms or legs provided that they are distal to the shoulders or inguinal/gluteal folds, respectively.

Mason-Likar electrode placement is commonly used by hospital staff and paramedics; this approach does not alter precordial electrode placement but instead moves the limb electrodes to the torso. Originally described in 1966, the Mason-Likar configuration differs from standard electrode placement in that the arm electrodes are relocated to the infraclavicular fossae (medial to the borders of the deltoid muscles and 2 cm below the clavicles), and the leg electrodes are positioned along the anterior axillary lines (halfway between the costal margins and the iliac crests). Actual torso positioning may differ in practice because of individual variation or an attempt to simulate limb electrode placement. A rightward frontal-plane access shift has been described when torso electrode placement is used for the limb electrodes instead of standard positioning on the extremities. Mason-Likar positioning has also been associated with diminution of inferior Q waves, thus making detection of inferior MI more difficult.

An alternative electrode configuration is the Lund system, in which the arm electrodes are placed laterally on the left and right arms at the level of the axillary folds, and the leg electrodes are positioned laterally on the greater femoral trochanters. The Lund system has been found to more directly approximate the electrocardiographic recordings obtained with conventional positioning than the Mason-Likar configuration does.

The precordial electrodes should be placed as follows: V1—right sternal border, fourth intercostal space; V2—left sternal border, fourth intercostal space; V3—midway between V2 and V4; V4—left midclavicular line, fifth intercostal space; V5—left anterior axillary line, same horizontal level as V6; and V6—left midaxillary line, same horizontal level as V4 and V5. Note that V4 to V6 are placed at the same horizontal level, not all in the fifth intercostal space. If V4 and V6 are situated so that they follow the contour of the intercostal space rather than being on the same horizontal level, they will be superiorly displaced as the ribs curve around the side of
the thorax. Minor changes in the position of the precordial leads will alter the ECG tracing, so it is important to keep the adhesive leads in place throughout the ED stay so that lead placement is identical during serial ECG comparisons.

Intercostal space number can be determined by first palpating the sternal angle (angle of Louis), which is the junction of the manubrium and body of the sternum (see Fig. 14.4). This transverse bony ridge is located about 5 cm caudal from the sternal notch in adults. Immediately lateral and inferior to it is the second intercostal space; two spaces farther down lies the fourth intercostal space, where V1 and V2 should be placed. Alternatively, one can count down from the medial aspect of the clavicle; beneath the clavicle lies the first rib, below which is the first intercostal space. The precordial electrodes should not be simply “eyeballed” by the technician because as little as 1 to 2 cm of electrode displacement can result in significant morphologic alteration in the precordial QRS complexes.

If the patient’s body habitus or pathologic process precludes placement of a precordial electrode as just described, it is permissible to attach it within the radius of the width of one interspace of the recommended position, with appropriate notation on the tracing. If the situation demands further displacement, it is recommended that the lead be omitted, with appropriate documentation on the tracing.

Preformatted lead placement is used by some clinicians. For instance, the use of disposable, prewired electrodes in a manner similar to Mason-Likar positioning allows for more rapid placement, 20% faster compared to standard lead placement. The quality of the resultant ECG tracing is excellent, with lower rates of artifact and minimal impact on standard electrocardiographic structures. These preformatted lead placement systems are more expensive, however, as compared to standard lead placement wiring.

Pediatric Electrode Placement

In addition to the standard 12-lead tracing, leads V4R and V3R should also be recorded; these are mirror images of their left-sided counterparts (see section on Additional Leads later in this chapter). The chest of a tiny infant may not accommodate all the precordial electrodes; in such cases the following array is recommended: V1R or V4R, V5, V6. Limb electrode placement is the same as in adults.

FEATURES OF THE ECG

Information Provided by the Computer

In addition to the patient demographic data entered by the operator, the tracing will often feature computations regarding rate, intervals, and axes along the top of the paper. On some tracings, a computer-generated interpretation, or “reading,” will also be displayed at the top of the tracing. These interpretations are not infallible. A sample of nine of these programs was compared with the readings of eight cardiologists; the gold standard in this study was clinical diagnosis made independently of the interpretations of these tracings based on other objective data (e.g., echocardiography, cardiac catheterization). The performance of the programs was good, with correct interpretations in a median of 91% of cases, but the cardiologists were significantly better (median of 96% correct).

The computer programs demonstrated a median sensitivity for anterior and inferior MI of only 77% and 59%, respectively. Of note, this study did not evaluate interpretation of acute ischemia and cardiac rhythm disturbance, other critical issues in electrocardiographic interpretation. Others have found both the computer programs and clinicians to be lacking in their ability to exclude cardiac disease with the ECG, with a negative predictive value for each of between 80% and 85%. When diagnosing atrial fibrillation, both general practitioners (sensitivity, 80%; specificity, 83%) and computer software (sensitivity, 83%; specificity, 99%) are flawed; when combined, diagnostic accuracy improves but is still imperfect (sensitivity, 92%; specificity, 91%).

It is worthwhile to read and consider the computer reading of the ECG, but the emergency clinician should not be beholden to it. The properly trained and experienced clinician provides more accurate electrocardiographic interpretation, as compared with the machine’s computer algorithm.

Adjustable Features

A notation of electrocardiographic paper speed (in millimeters per second), calibration (in millimeters per millivolt), and the frequency response (in hertz) is displayed, usually in the left lower corner of the recording. Paper speed is usually set at a default of 25 mm/sec. It may be manipulated for purposes of deciphering a dysrhythmia, as described later (see section on Alteration in Amplitude and Paper Speed).

Calibration, or standardization, refers to the amplitude of the waveforms on the tracing. It is usually set at a default value of 10 mm/mV and is graphically depicted by a plateau-shaped waveform that appears at the extreme left side of the tracing, in front of the first complex (Fig. 14.5A). This calibration can be modified by the operator or by the computer itself, as was the case in Fig. 14.5B, in which the patient appeared to have acquired voltage criteria for left ventricular hypertrophy when in reality the tracing was unchanged from his baseline (see Fig. 14.5A). Increasing the calibration to 20 mm/mV is helpful when trying to decipher P-wave morphology. Decreasing the calibration to 5 mm/mV is helpful in cases in which the amplitude of the QRS complex (usually in the precordial leads) is so large that it encroaches on those of adjacent leads. Standardization may not be uniform throughout a given tracing. At times calibration will be automatically adjusted by the computer based on the waveform amplitudes that it perceives.

For example, it is possible to have normal calibration (10 mm/mV) in the limb and augmented leads, with half-standard calibration in the precordial leads (5 mm/mV). This may occur in instances of marked left ventricular hypertrophy. In this case the calibration pulse at the left-hand side of the paper will have a downward stairstep appearance.

It is important that the clinician examine all electrocardiographic tracings for standardization and paper speed parameters before rendering an interpretation.

ADDITIONAL ELECTROCARDIOGRAPHIC LEADS

Though not performed routinely and not an ED standard of care, additional electrocardiographic leads may be used in the evaluation of a patient with possible ACS. These leads can be considered in a patient with a presentation consistent with
Potential indications for 15-lead ECGs in patients with suspected ACS include the following: (1) ST-segment depression in leads V1 through V3, (2) STEMI involving the inferior or lateral regions, (3) isolated ST-segment elevation in lead V1, or (4) high clinical suspicion for AMI without electrocardiographic evidence of STEMI on a 12-lead ECG (i.e., to detect “occult” left circumflex STEMI).

The posterior electrodes V8 and V9 are placed on the patient’s back, V8 at the tip of the left scapula and V9 in an intermediate position between lead V8 and the left paraspinal muscles. An additional electrode, V7, may also be used and is placed on the posterior axillary line equidistant from electrode V8 (Fig. 14.6). The degree of ST-segment elevation in the posterior leads is often less pronounced than the ST-segment elevation in ACS yet demonstrating an unrevealing or nondiagnostic 12-lead ECG. These additional, or nontraditional, leads include posterior leads (V7, V8, and V9), right ventricular leads (especially V4R), and procedural leads (transvenous pacemaker wire placement and pericardiocentesis). Acute posterior and right ventricular MIs are likely to be underdiagnosed because the standard 12-lead ECG does not assess these areas directly. The standard ECG coupled with these additional leads constitutes the 15-lead ECG, the most frequently used extra lead ECG in clinical practice.

### 15-Lead ECG—Posterior and Right Ventricular Leads

Several different investigations have explored the use of the 15-lead ECG. In a study of all ED patients with chest pain, Brady and associates reported that the 15-lead ECG provided a more accurate description of myocardial injury in patients with AMI, yet failed to alter rates of diagnosis or use of reperfusion therapies or to change disposition locations. Looking at a more select population of ED patients, Zalenski and colleagues investigated use of the 15-lead ECG in patients with chest pain and a moderate to high pretest probability of AMI, who were already identified as candidates for hospital admission. In this 15-lead ECG study, the authors reported a 12% increase in sensitivity with no loss of specificity (i.e., no increase in false-positive findings) for the diagnosis of ST-segment elevation AMI (STEMI). They further suggested that in the diagnosis of posterior AMI, leads V7 and V9 are superior to reliance on detecting the reciprocal ST-segment depression seen in leads V1 to V3. Aqel and colleagues used balloon inflation during coronary angiography for ostial or proximal left circumflex disease to simulate proximal left circumflex STEMI; they found that the posterior leads were significantly more accurate at detecting simulated STEMI when this vessel was occluded. Interestingly, 11% of their 53 patients had 1-mm or greater ST-segment elevation in a posterior lead with no ST-segment elevation or depression in any other lead.

Potential indications for 15-lead ECGs in patients with suspected ACS include the following: (1) ST-segment depression in leads V1 through V3, (2) STEMI involving the inferior or lateral regions, (3) isolated ST-segment elevation in lead V1, or (4) high clinical suspicion for AMI without electrocardiographic evidence of STEMI on a 12-lead ECG (i.e., to detect “occult” left circumflex STEMI).

The posterior electrodes V7 and V9 are placed on the patient’s back, V7 at the tip of the left scapula and V9 in an intermediate position between lead V8 and the left paraspinal muscles. An additional electrode, V7, may also be used and is placed on the posterior axillary line equidistant from electrode V8 (Fig. 14.6). The degree of ST-segment elevation in the posterior leads is often less pronounced than the ST-segment elevation.
Invasive Procedural Leads

Patients with severely compromising bradycardia may require a transvenous pacemaker. In such instances, it may be necessary to place the pacing wire without the benefit of fluoroscopy. In such cases the recommendation is to advance the wire under electrocardiographic guidance with the patient connected to the limb leads of a grounded electrocardiographic machine and the pacing wire connected to the V lead (see Chapter 15). As the electrode enters the superior vena cava and high right atrium the P wave and QRS complex will be negative. While traversing the atrium, the P wave and QRS complex will become positive, and the latter will become larger as the ventricle is approached. If a balloon-tipped flotation catheter is used, deflate the balloon once it is in the right ventricle. Next, advance it until contact is made with the endocardium and the ventricle is captured. Ventricular wall contact will be indicated by marked ST-segment elevation. (See Chapter 15 for additional information on this procedure)

For patients with suspected pericardial effusion who are undergoing urgent pericardiocentesis, an electrocardiographic lead may be placed on the syringe needle. This form of monitoring will assist in correct positioning of the catheter in the pericardial space. Monitor the ST segments while advancing the needle. A sudden appearance of ST-segment elevation indicates that the needle has moved too far internally (i.e., beyond the pericardial space) and has made contact with the epicardium. (See Chapter 16 for additional information on this procedure.)

Body Surface Mapping

An emerging electrocardiographic tool, body surface mapping uses numerous leads to provide a more detailed electrical description of the heart than possible with the 12-lead ECG. The body map ECG that is most commonly used is based on an 80-lead ECG with 64 anterior and 16 posterior leads. This more detailed imaging of the myocardium allows potentially greater diagnostic accuracy in the early detection of STEMI, in addition to detection of infarction in more traditionally electrocardiographic “silent” areas of the heart. The body map ECG should not replace the typical 12-lead ECG in patients with chest pain. It should be used only as a second-tier tool in the evaluation of patients with intermediate to high clinical suspicion for ACS and an unrevealing initial 12-lead ECG. In this instance the clinician is in search of STEMI in electrocardiographically silent areas, namely, the far inferior and lateral walls, the posterior wall, and the right ventricle. The use of body mapping does increase the rate of ACS diagnosis as noted in the Optimal Cardiographic Diagnostic Evaluation Enabling Faster Treatment of Myocardial Infarction (OCCULT MI) trial; the OCCULT MI study group obtained simultaneous 12-lead and 80-lead ECGs in patients suspected of having ACS. The 80-lead ECG improved the rate of diagnosis

Figure 14.7 Right-sided lead placement. Place right-sided leads V1R to V6R on the chest as a mirror image of the standard precordial leads. This alteration in lead placement is used to investigate the possibility of a right-sided acute myocardial infarction that may not be appreciated with normal lead placement.

Figure 14.8 Right-sided precordial leads. This alteration in lead placement is used to investigate the possibility of a right-sided acute myocardial infarction (MI) that may not be appreciated with normal lead placement. This tracing displays the right-sided precordial leads in an elderly man with chest pain, and they are consistent with acute coronary syndrome. There is ST-segment elevation that is somewhat subtle in the inferior leads (II, III, and aVF), which together with the reciprocal ST-segment depression seen in lead aVL is consistent with a diagnosis of acute inferior MI. Leads V1 to V6 are in actuality leads V5R to V6R—right-sided precordial leads. The convex upward ST-segment elevation seen in leads V5R to V6R is indicative of concomitant right ventricular infarction. This patient was found to have a subtotal proximal occlusive lesion of his right coronary artery at cardiac catheterization.
PREHOSPITAL ECG

Early treatment, primarily reperfusion therapy, is an important goal in the management of ACS, particularly STEMI. Early treatment is made possible by early STEMI diagnosis, which is achieved by earlier application of the diagnostic tools, namely the 12-lead ECG. Emergency medical services (EMS) systems and strategies that optimize STEMI care include early application of the ECG, which reduces the time to initiation of reperfusion therapy.\(^{37-40}\)

Many studies have demonstrated benefits of prehospital ECGs in decreasing door-to-fibrinolytic treatment and door-to-vascular reperfusion therapy.\(^{35-40}\) Other studies have shown even further reductions in time to revascularization when cardiac catheterization laboratories are activated prior to patient arrival at the hospital.\(^{17-40}\) The early acquisition, interpretation, and transmission of a prehospital 12-lead ECG can dramatically decrease the time to treatment for a patient with ACS, particularly STEMI. Importantly, this reduction in time to management is seen in urban, suburban, and rural settings, regardless of the transport time or type of reperfusion strategy selected for the particular patient.

Beyond establishing the diagnosis of STEMI prior to ED arrival, the prehospital 12-lead ECG can impact EMS management strategies and hospital-based diagnosis in several ways, including the following: more aggressive and focused EMS-based therapies; change of destination hospital (i.e., a PCI-capable facility); rapid transport to hospital (i.e., helicopter) if transport time is significantly prolonged; and the detection of ACS-related changes that resolve prior to the patient’s arrival at hospital.

It must also be pointed out that minimal prolongation in EMS scene time is acceptable when one considers the vital information provided by the ECG and its impact on overall time-to-therapy considerations. The prehospital ECG can be obtained within as little as 2 to 3 minutes; furthermore, it can be obtained while en route to the hospital. The significant reduction in time to definitive therapy for STEMI more than compensates for any small transport delay.

ALTERNATIVE TECHNIQUES FOR ASSESSMENT OF RHYTHM

Electrocardiographic rhythm assessment depends on a clear signal of both atrial and ventricular electrical activity over a period. Although continuous 12-lead electrocardiographic rhythm monitoring has the advantage of recording cardiac activity over multiple leads (thus maximizing atrial and ventricular monitoring), it is often impractical. Moreover, correct identification of the cardiac rhythm on an ECG can be difficult, depending on the clinical setting. Rapid atrial or ventricular rates, especially those above 150 beats/min, often lead to simultaneous or nearly simultaneous deflections that can alter the usual waveform patterns or cause smaller deflections to be buried within larger ones (such as P waves buried within the QRS complex). In addition, rapid rates result in smaller, narrower waveforms, which makes visual recognition on the ECG challenging. Finally, assessment of atrial activity in general is more difficult because of the smaller electrical impulse and resulting electrocardiographic waveform generated by the atria.

Lead V₁ is generally considered the most appropriate lead for detecting the P wave, followed by lead II. In a study of 62 measurements in 28 patients, lead V₁ demonstrated the tallest P wave 53% of the time, followed by lead II (29%), lead I (7%), and lead III (3%).\(^{41}\) A number of alternative techniques have been developed to improve assessment of rhythm, including alterations in the standard 12-lead ECG, as well as the addition of nonstandard leads to monitor cardiac and, in particular, atrial rhythm activity.

Alteration in Amplitude and Paper Speed

Most 12-lead electrocardiography machines today allow alteration of both amplitude and paper speed from the basic 10-mm/mV and 25-mm/sec standards, respectively. Increasing the amplitude to double the standard or 20 mm/mV, can increase the prominence of smaller deflections such as the P wave, and can improve recognition of the atrial rhythm (Fig. 14.9). In addition, clinicians have also used photocopy enlargements of the standard ECG to visually enhance smaller deflections.\(^{42}\)

Increasing the paper speed to double the standard or 50 mm/sec, has the effect of artificially slowing the rhythm. This technique is most advantageous when assessing patients with marked atrial or ventricular tachycardia. Increasing the paper speed exaggerates any existing irregularity (such as in atrial fibrillation) and can improve recognition of smaller deflections, such as P waves, in the presence of a significant tachycardia. Faster paper speeds also make it possible to measure short electrocardiographic intervals (such as PR or R-R) more accurately (Fig. 14.10). Accardi and coworkers\(^{43}\) found that overall diagnostic accuracy improved when clinicians were provided ECGs recorded at the faster 50-mm/sec paper speed, as opposed to a standard 12-lead ECG, in patients with narrow-complex tachycardia.

Alternative Leads

Assessment of rhythm often requires electrocardiographic monitoring over a longer and continuous period; such a need is not met by the 12-lead ECG. A number of alternative lead systems requiring fewer electrodes have been described. Many of these systems use the limb bipolar leads (RA, LA, LL) in alternative positions over the chest. Leads I, II, or III are then recorded depending on the positions of the positive and negative electrodes. Although various alternative leads provide additional information to the clinician, there are no standard guidelines mandating the use of any alternative leads in the ED.

Lewis Leads

In 1910, Thomas Lewis first described alternative positions for the RA and LL leads to enhance detection of atrial fibrillation. The RA lead was placed over the right second costochondral junction, whereas the LL lead was placed in the right fourth intercostal space 1 inch to the right of the sternum, with the LA and RL leads left in their usual positions. Madias compared standard ECG leads to the “Lewis lead,”\(^{44}\) noting enhancement of atrial activity when the RA served as the negative electrode and LL as the positive electrode (lead II) in this new configuration. Other alternative lead placements to enhance detection of atrial activity have also been described (Table 14.2 and Fig. 14.11).
Figure 14.9  
A, Baseline electrocardiogram (ECG) of a patient before the development of an abnormal rhythm (10 mm/mV). Note the P-wave morphologies, especially in leads I, II, and V1.  
B, ECG during ectopic atrial tachycardia (10 mm/mV). Note the change in P-wave morphology, especially in lead V1.  
C, ECG during ectopic atrial tachycardia (20 mm/mV). The P waves are now easier to see in all leads.  
D, ECG after reversion to a normal atrial focus (20 mm/mV). Contrast these accentuated P waves with those in C.

Figure 14.10  
A, Electrocardiogram (ECG) with tachycardia at normal paper speed (25 mm/sec). Because of the rapid rate, the actual P waves are difficult to discern, thus making determination of the rhythm difficult. The computerized interpretation is sinus tachycardia with a first-degree atrioventricular (AV) block.  
B, ECG with tachycardia at double paper speed (50 mm/sec). With increased paper speed, atrial P-wave activity is accentuated, and atrial flutter with a 2:1 AV block is demonstrated. Carotid sinus massage may have slowed the rhythm and also revealed P waves.
**TABLE 14.2 Alternative Leads for Assessment of Rhythm**

<table>
<thead>
<tr>
<th>Lead</th>
<th>Negative Electrode Position</th>
<th>Positive Electrode Position</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lead I</td>
<td>Right second costochondral junction</td>
<td>Right fourth intercostal space, 1 inch right of the sternum</td>
</tr>
<tr>
<td>Lead II</td>
<td>Second right costochondral junction, Center of the sternum</td>
<td>Seventh right costal cartilage, Inferior angle of the scapula 2 inches right of the spine</td>
</tr>
<tr>
<td>Lead III</td>
<td>Third intercostal space along the right sternal border, Left leg, Right arm</td>
<td></td>
</tr>
<tr>
<td>Alternative Lead</td>
<td>Negative Electrode Position</td>
<td>Positive Electrode Position</td>
</tr>
<tr>
<td>Lewis</td>
<td>Right second costochondral junction</td>
<td>Right fourth intercostal space, 1 inch right of the sternum</td>
</tr>
<tr>
<td>Drury</td>
<td>Second right costochondral junction, Center of the sternum</td>
<td>Seventh right costal cartilage, Inferior angle of the scapula 2 inches right of the spine</td>
</tr>
<tr>
<td>Schoenwald</td>
<td>Third intercostal space along the right sternal border, Left leg, Right arm</td>
<td></td>
</tr>
<tr>
<td>Lu</td>
<td>First intercostal space directly above V1</td>
<td>Approximately 3 inches directly below V4</td>
</tr>
<tr>
<td>Vertical sternal (&quot;Barker leads&quot;)</td>
<td>Below the suprasternal notch at the manubrium</td>
<td>Xiphoid process</td>
</tr>
<tr>
<td>MCL1</td>
<td>Left shoulder (1 cm inferior to the left midclavicle)</td>
<td>V1 (fourth intercostal space, right sternal border)</td>
</tr>
<tr>
<td>MCL6</td>
<td>Left shoulder (1 cm inferior to the left midclavicle)</td>
<td>V6 (sixth rib, midaxillary line)</td>
</tr>
</tbody>
</table>

1. First, set the electrocardiographic machine to record the rhythm strip with this lead. If the recording rhythm strip is lead I, the RA wire becomes the negative electrode, which is placed as noted in the table, and the LA wire becomes the positive electrode, which is placed as noted in the table. If lead II or lead III is the lead that is set to record the rhythm strip, the positive and negative electrodes will vary.

2. Example: One way to record the Lewis lead is to set the electrocardiographic machine to record lead I, use the RA wire as the negative electrode, and place it in the right second costochondral junction. Use the LA wire as the positive electrode and place it in the right fourth intercostal space, 1 inch to the right of the sternum. The Lewis lead may also be recorded on lead II and lead III, but the wires that serve as the positive and negative electrodes will vary.

P wave than other systems do, including the Lewis leads. In addition, the vertical sternal leads are placed over bone, which may reduce artifact caused by muscle activity on recordings (see Fig. 14.11).

**Limb-Precordium Leads**
A sequential pattern of bipolar leads on the chest, termed limb-precordium leads, has been proposed in combination with the original Einthoven limb leads. In this system standard limb leads are placed on the patient. The RA electrode is then repositioned sequentially at the fourth intercostal space just to the right of the sternum, the fourth intercostal space just to the left of the sternum (low parasternal), the first intercostal space just to the left of the sternum, and the first intercostal space just to the right of the sternum (high parasternal). During this sequential mapping, tracings are recorded for leads I and II until atrial activity is identified. Brenes-Pereira reported that this mapping system allowed the identification of P waves in a majority of patients when none were detected initially on the standard 12-lead ECG.

**Modified Bipolar Chest Leads**
Modified bipolar chest leads (MCLs) are the most commonly used leads for monitoring cardiac rhythm. The positive electrode is placed on the chest at a precordial position (V) concordant with the MCL desired (e.g., the V1 position for MCL1). The negative electrode is placed on the left shoulder. On standard electrocardiographic machines, the LA electrode is placed at V1, RA at the left shoulder, LL at V6, and RL at a remote location on the chest to serve as ground. Lead I would then reflect MCL1 and lead II, MCL6. MCL1 may be useful in distinguishing atrial activity; MCL4 and MCL6 more commonly
in monitoring of the ST-T wave; and both MCL₁ and MCL₆ in evaluating wide-complex tachycardias

Esophageal Leads
The esophageal lead (E) was first described by Brown in the 1930s. Since that time both unipolar and bipolar esophageal leads have been developed. Because of its posterior location, this lead is often superior in detecting atrial deflections and recording the activity of the posterior surface of the left ventricle. The electrode, which is connected to the ECG by thin wires, is either swallowed or passed through the nares into the esophagus. Once in the esophagus, the location of the electrode is determined either by fluoroscopy or by making a series of low to high esophageal recordings. The position of the electrode in the esophagus is adjusted by slowly pulling the electrode wire out the nares or mouth. In normal adults, leads E₁₅–₂₅ (the electrode is located in the esophagus 15 to 25 cm from the nares) generally records atrial activity; E₂₅–₃₅ activity of the atroventricular groove; and E₃₀–₅₀ activity of the left ventricular posterior surface. The E lead should be recorded through lead channel I simultaneously with lead channel II and the other surface channels.

Central Venous Catheter Intracardiac Leads
For patients in whom a central venous catheter was placed for vascular access (or for other reasons such as cardiac pacing, hemodialysis, or Swan-Ganz monitoring) that catheter, when filled with saline, can be used as a modified intracardiac electrode for recording of atrial activity. Once filled with saline, a needle was then left in a side access port of the catheter and attached via an alligator clip to lead V₁. With this method, the distal port of the saline-filled central venous catheter demonstrated significantly larger P waves than the standard 12-lead ECG and the Lewis lead did.

ELECTRODE MISPLACEMENT AND MISCONNECTION

Limb Electrode Reversal
Although the limb electrodes are not often misplaced, the cables that link them to the ECG are at times improperly connected, resulting in “electrocardiographic changes” that are, in actuality, artifact. A multitude of possibilities for misconnection of the limb electrodes exists; some of the most likely are summarized here. It is helpful to categorize these possibilities into those that are easily recognizable without comparison to an old ECG versus those that are not. Failure to recognize limb electrode reversal may lead to misattribution of ECG “changes” to a disease that is actually caused by technical misadventure. Even though the incidence of electrode reversal in the ED has not been quantified, it has been observed to occur in an intensive care unit setting in 4% to 5% of tracings.

Easily Recognizable without an Old ECG
The most common of all misconnections is reversal of the LA and RA electrodes (Fig. 14.12). The hallmark is a negative P wave and a primarily negative QRS complex in lead I, which creates a right or extreme axis deviation (depending on the principal vector of the QRS complex in lead aVF). Dextrocardia

Figure 14.12 Arm electrode reversal (LA ↔ RA). The most common of limb electrode reversals, the clues lie in leads I and aVR. Lead I features a negative P wave, as well as a principally negative QRS complex and T wave. This could suggest dextrocardia, but the precordial leads demonstrate a normal transition, which is not consistent with dextrocardia. Note also the unusual appearance of aVR in this tracing. should also be considered with such findings; the pattern of precordial lead transition will differentiate between dextrocardia

Not Easily Recognizable Without an Old ECG
One limb electrode reversal that is not readily recognizable without comparison to a previous tracing is transposition of the LA and LL electrodes. This causes transposition of lead I with lead II on the tracing in addition to lead aVL with aVF. In effect, two inferior leads (II and aVF) have become the lateral leads (I and aVL) and vice versa, thus making this misconnection difficult to detect at times without a baseline ECG for comparison. Furthermore, lead III will be upside
Changes are not usually grossly abnormal and therefore can be difficult to detect. Variation often becomes evident when comparing the current tracing with an old ECG. In such cases, it is useful to go to the bedside and examine where the electrodes were positioned relative to the recommended placement (see section on Electrode Placement earlier in this chapter). One cannot ensure, however, that the baseline ECG was done with proper electrode placement. When comparing the precordial leads on the current ECG with a baseline tracing, ST-segment and T-wave changes should be viewed in the context of the relative morphologies of the associated QRS complexes. If a marked difference is noted between the two tracings in the amplitude or polarity of the QRS complex in a given precordial lead—the R/S ratio for that QRS complex—the corresponding ST-T-wave changes may be the result of variability in electrode placement, although cardiac ischemia cannot be completely excluded as the cause.

**Figure 14.13** Right-sided electrode reversal (RA ↔ RL). A, In this tracing, the nearly flat line in lead II (arrow) suggests that the right leg electrode has been switched with the right arm electrode. The trivial voltage in lead II reflects the expected lack of potential difference between the two legs (the right leg electrode is in the right arm position; lead II normally runs from the right arm to the left leg, but now it effectively runs from the right leg to the left leg). B, The second tracing was acquired with correct electrode placement.

**Figure 14.14** A, Left-sided limb electrode reversal (LA ↔ LL). A patient with a history consistent with acute coronary syndrome was brought to the emergency department after this electrocardiogram was recorded in a clinic. Leads I and aVL suggest an acute high lateral infarction, but surprisingly there are no corresponding changes in leads V3 and V6. The deep T-wave inversions in III and aVF were at first thought to be inferior ischemia or reciprocal changes (see also B). B, Correction of electrode reversal (LA ↔ LL). After the electrodes were reconnected, this tracing reveals an acute inferior wall myocardial infarction (MI) and deep T-wave inversion in aVL, a harbinger of acute inferior MI. In comparing this tracing with that in A, note the following: lead I ↔ lead II, lead aVL ↔ aVF, and lead III is inverted. Thus, inferior changes become lateral, and lateral changes become inferior, the hallmark of LA ↔ LL reversal.

**Precordial Electrode Misplacement and Misconnection**

Unlike the limb electrodes, the precordial electrodes are more prone to misplacement, especially when variations in body habitus (e.g., obesity, breast tissue, pectus excavatum, chronic lung disease) make proper electrode placement more difficult. This may cause some variability in the amplitude and morphology of the complexes in the precordial leads. However, these changes are not usually grossly abnormal and therefore can be difficult to detect. Variation often becomes evident when comparing the current tracing with an old ECG. In such cases, it is useful to go to the bedside and examine where the electrodes were positioned relative to the recommended placement (see section on Electrode Placement earlier in this chapter). One cannot ensure, however, that the baseline ECG was done with proper electrode placement. When comparing the precordial leads on the current ECG with a baseline tracing, ST-segment and T-wave changes should be viewed in the context of the relative morphologies of the associated QRS complexes. If a marked difference is noted between the two tracings in the amplitude or polarity of the QRS complex in a given precordial lead—the R/S ratio for that QRS complex—the corresponding ST-T-wave changes may be the result of variability in electrode placement, although cardiac ischemia cannot be completely excluded as the cause.
Electrocardiographic artifact is commonly encountered yet not always easy to recognize. It can be attributed to either physiologic (internal) or nonphysiologic (external) sources; the former includes muscle activity, patient motion, and poor electrode contact with the skin. Tremors, hiccups, and shivering may produce frequent, narrow spikes on the tracing and simulate atrial and ventricular dysrhythmias (Fig. 14.16). A wandering baseline featuring wide undulations in addition to other “noise” on the ECG can often be traced to patient movement and high skin impedance, which leads to inadequate contact of the electrode with the skin. Minimizing skin impedance and artifact may be achieved by: (1) avoiding electrode placement over bony prominences, major muscles, or pulsating arteries; (2) clipping rather than shaving thick hair at electrode sites; and (3) cleaning, and most importantly, drying the skin surface before reapplying the electrode if the tracing features substantial artifact. Nonphysiologic artifact is most often caused by 60-Hz electrical interference, which is ascribable to various other sources of alternating current near the patient. This will be manifested as a wide, indistinct isoelectric baseline.

Some studies have reported that placement of chest electrodes by more than 20 to 25 mm from the standard positions can be associated with clinically significant changes on the ECG. It has been observed that leads V1 and V2 are typically placed too high and that the lateral leads are placed too laterally and too low. McCann and colleagues demonstrated a high degree of variability between experienced clinicians in identifying anatomic landmarks for precordial electrocardiographic electrode placement. There was frequently a large difference in the measured distance from the actual to the “standardized” electrode position that ranged between 0 and 105 mm in the vertical direction (mean, 14 mm; median, 10 mm), and between 0 and 120 mm in the horizontal plane (mean, 17 mm; median, 10 mm). Overall, 20.8% of the paired measurements in the vertical direction and 26.6% of those in the horizontal plane differed by more than 25 mm.

Misconnection of the precordial cables is usually easy to detect. The expected progression of P-, QRS-, and T-wave morphologies across the precordium will be disrupted (Fig. 14.15). An abrupt change in wave morphology evolution—followed by a seeming return to normalcy in the next lead—is a good clue to misconnection of the precordial electrodes.

**TABLE 14.3** Clues to Improper Limb Electrode Connections

<table>
<thead>
<tr>
<th>REVERSED LEADS</th>
<th>NEED OLD ECG FOR DETECTION?</th>
<th>KEY FINDINGS</th>
</tr>
</thead>
<tbody>
<tr>
<td>LA ↔ RA</td>
<td>No</td>
<td>P-QRS-T waves upside down in lead I Precordial leads normal (not dextrocardia)</td>
</tr>
<tr>
<td>LA ↔ LL</td>
<td>Yes</td>
<td>III upside down from baseline I ↔ II, aVL ↔ aVF, no change in aVR</td>
</tr>
<tr>
<td>LA ↔ RL</td>
<td>No</td>
<td>III is a straight line</td>
</tr>
<tr>
<td>RA ↔ LL</td>
<td>No</td>
<td>P-QRS-T waves upside down in all leads except aVL</td>
</tr>
<tr>
<td>RA ↔ RL</td>
<td>No</td>
<td>II is a straight line</td>
</tr>
<tr>
<td>LL ↔ RL</td>
<td>Cannot detect change</td>
<td>Looks like normal electrode placement</td>
</tr>
<tr>
<td>LA ↔ LL + RA ↔ RL</td>
<td>No</td>
<td>I is a straight line, aVL and aVR are the same polarity and amplitude, and II is upside down III</td>
</tr>
</tbody>
</table>

LA, Left arm; LL, left leg; RA, right arm; RL, right leg.


**Figure 14.15** A and B, Precordial electrode reversal (V2 ↔ V3). Note that the usual precordial progression of R-wave growth in leads V1 and V3 is disrupted in the tracing displayed in A. B shows a return to a normal V3 transition zone.

**ARTIFACT**

Electrocardiographic artifact is commonly encountered yet not always easy to recognize. It can be attributed to either physiologic (internal) or nonphysiologic (external) sources; the former includes muscle activity, patient motion, and poor electrode contact with the skin. Tremors, hiccups, and shivering may produce frequent, narrow spikes on the tracing and simulate atrial and ventricular dysrhythmias (Fig. 14.16). A wandering baseline featuring wide undulations in addition to other “noise” on the ECG can often be traced to patient movement and high skin impedance, which leads to inadequate contact of the electrode with the skin. Minimizing skin impedance and artifact may be achieved by: (1) avoiding electrode placement over bony prominences, major muscles, or pulsating arteries; (2) clipping rather than shaving thick hair at electrode sites; and (3) cleaning, and most importantly, drying the skin surface before reapplying the electrode if the tracing features substantial artifact. Nonphysiologic artifact is most often caused by 60-Hz electrical interference, which is ascribable to various other sources of alternating current near the patient. This will be manifested as a wide, indistinct isoelectric baseline.
Electrocardiographic artifact should be considered when the clinical picture indicates stability, and status quo and coincident procedures are in progress (e.g., hemodialysis, blood warmer, bronchoscopy) or devices are in use (e.g., nerve stimulators); the list of causative equipment is long and varied. Other sources of nonphysiologic artifact include those attributable to the monitoring equipment: loose connections, broken monitor cables, and mechanical issues with the machine (e.g., broken stylus, uneven paper transport). The 60-Hz artifact caused by electrical current interference can be minimized by shutting off nonessential sources of current in the vicinity, and also by straightening the lead wires so that they are parallel to the patient’s body in the long axis.

Differentiation of artifact from true electrocardiographic abnormality is intuitively important; moreover, clinical consequences have been reported that are directly attributable to confusion of artifact with disease. Unnecessary treatment and procedures—including cardiac catheterization, electrophysiologic testing, and even implantation of a pacemaker and an automatic defibrillator—have been reported. Characteristics that may aid in differentiating artifact from dysrhythmia include the absence of hemodynamic instability during the event (or even absence of any symptoms), normal QRS complexes occurring during the “dysrhythmia,” instability of the baseline on the tracing during and immediately after the “dysrhythmic” event, association with body movement, and observance of “notches” amid the complexes of the pseudo-dysrhythmia that “march out” with the normal QRS complexes that precede and follow the disturbance (Fig. 14.17).

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Emergency Cardiac Pacing

Edward S. Bessman

The purpose of cardiac pacing is to restore or ensure effective cardiac depolarization. Emergency cardiac pacing may be instituted either prophylactically or therapeutically. Prophylactic indications include patients with a high risk for atrioventricular (AV) block. Therapeutic indications include symptomatic bradycardias and overdrive pacing. Pacing for asystole has very minimal success but it has been used for this condition. Several approaches to pacing can be taken, including transcutaneous, transvenous, transthoracic, epicardial, endocardial, and esophageal. Transcutaneous and transvenous pacemakers are the two techniques most commonly used in the emergency department (ED). Because it can be instituted quickly and noninvasively, transcutaneous pacing is the technique of choice in the ED when time is of the essence. Transvenous pacing should be reserved for patients who require prolonged pacing or have a very high (>30%) risk for heart block. Transcutaneous pacing is generally a temporizing measure that may precede transvenous cardiac pacing. Although it is not an expectation that all emergency clinicians will be adept at placing temporary transvenous cardiac pacemakers, many have mastered the techniques and are often the only clinicians available to perform this lifesaving procedure.

EMERGENCY TRANSVENOUS CARDIAC PACING

The transvenous method of endocardial pacing is commonly used and is both safe and effective. In skilled hands, the semifloating transvenous catheter is successfully placed under electrocardiographic (ECG) guidance in 80% of patients. The technique can be performed in less than 20 minutes in 72% of patients and in less than 5 minutes in 30% (Videos 15.1–15.3). However, in some instances, anatomic, logistic, and hemodynamic impediments can prohibit successful pacing by even the most skilled clinician. As with other medical procedures, it should not be performed without a thorough understanding of its indications, contraindications, and complications.

Because this procedure is essentially performed in a blind manner, sometimes it will not be successful. This may be because the condition is not amenable to pacing (e.g., asystole, drug overdose) or because of technical difficulties inherent with the procedure.

Transcutaneous Cardiac Pacing

**Indications**

**Bradycardias**
- Symptomatic sinus node dysfunction
- Second- and third-degree heart block
- Atrial fibrillation with a slow ventricular response
- With myocardial infarction: new left bundle branch block, bifascicular block, alternating bundle branch block
- Malfunction of an implanted pacemaker

**Tachycardias**
- Supraventricular dysrhythmias
- Ventricular dysrhythmias

**Contraindications**
- Prosthetic tricuspid valve
- Severe hypothermia

**Complications**
- Inadvertent arterial puncture
- Venous thrombosis/thrombophlebitis
- Pneumothorax/other anatomic injury
- Ventricular arrhythmia
- Misplacement of the pacing catheter
- Myocardial/pericardial perforation
- Entanglement of the pacing catheter

**Equipment**

- Sheath introducer
- Transvenous pacing catheter
- 3-mL syringe
- Sterile sleeve
- Connecting cable
- Adapter pins
- Alligator clamp
- Pacing generator

The contents of a typical transvenous pacemaker kit are shown here. Additional equipment required for insertion of the sheath introducer is reviewed in Chapter 22. Individual kits may vary by manufacturer; be familiar with the equipment available at your institution before performing the procedure.

Review Box 15.1 Transcutaneous cardiac pacing: indications, contraindications, complications, and equipment.
Background

The ability of muscle to be artificially depolarized was recognized as early as the 18th century. Initial efforts focused on the transcutaneous approach (see later in this section). Over the succeeding years several scattered experiments were reported, and in 1951 Callaghan and Bigelow first used the transvenous approach to stimulate asystolic hearts in hypothermic dogs.1

Furman and Robinson demonstrated the transvenous endocardial approach in humans in 1958.4 They treated two patients with complete heart block and Stokes-Adams seizures, thus reconfirming that low-voltage pacing could completely control myocardial depolarization. The catheter remained in the second patient for 96 days without complication. Other early clinical studies also demonstrated the utility of transvenous pacing.5 Fluoroscopic guidance was used for placement of the pacing catheter in all these studies.

In 1964 Vogel and coworkers demonstrated the use of a flexible catheter passed without fluoroscopic guidance for intracardiac electrocardiography.6 One year later, Kimball and Killip used this technique to insert endocardial pacemakers at the bedside.7 They noted technical problems in 20% of their patients, including intermittent capture, difficulty passing the catheter, and catheter knotting. During the same year, Harris and colleagues confirmed the ease and speed with which this procedure could be accomplished.8

Before 1965 all intracardiac pacing was done asynchronously, which meant that the pacing catheter could cause electrical stimulation during any phase of the cardiac cycle. Asynchronous pacing frequently resulted in the pacemaker firing during the vulnerable period of an intrinsic depolarization; this occasionally caused ventricular tachycardia or fibrillation. In 1967 a demand pacemaker generator that sensed intrinsic depolarizations and inhibited the pacemaker for a predetermined period was used successfully by Zuckerman and associates in six patients.7 Since then there has been steady progress in the design and functionality of pacemakers. Table 15.1 summarizes the four-letter code that is used to describe modern pacemakers (there is a fifth letter for combined pacemaker-cardioverter/defibrillators). The most commonly used emergency transvenous pacemaker is represented by the code VVI: the ventricle is paced, the ventricle is sensed, and when a native impulse is sensed, the pacemaker is inhibited. Dual-chamber pacing (DDD or DDDR) is the preferred methodology for permanent pacing but is rarely used on an emergency basis because of the increased complexity of the procedure.

Rosenberg and coworkers introduced an improved pacing catheter known as the Elecath semifloating pacing wire.1 The Elecath was stiffer than the Flexon steel wire electrode that was in prevailing use. Rosenberg and coworkers achieved pacing in 72% of their patients with an average procedure time of 18 minutes. They also noted that 30% of their patients were paced in 5 minutes or less. In 1970, Swan and Ganz introduced the technique of heart catheterization with a flow-directed balloon-tipped catheter.9 Schnitzler and colleagues successfully used this method for placement of a right ventricular pacemaker in 15 of 17 patients.10

In 1981 Lang and associates compared bedside use of the flow-directed balloon-tipped catheter with insertion of a semirigid electrode catheter in 111 perfusing patients.11 These researchers found a significantly shorter insertion time (6 minutes 45 seconds versus 13 minutes 30 seconds), a lower incidence of serious arrhythmias (1.5% versus 20.4%), and a lower incidence of catheter displacement (13.4% versus 32%) with the balloon-tipped catheter. They concluded that the balloon-tipped catheter was the method of choice for temporary transvenous pacing.

Although placing temporary transvenous pacemakers has long been considered a core skill in the ED, it has not been well studied. Birkhahn and coworkers retrospectively compared the experience of emergency physicians with that of cardiologists in placing transvenous pacemakers under ECG guidance.12 They reported a 13% risk for major complications in both groups of specialists. They concluded that pacemaker placement by emergency physicians under ECG guidance without fluoroscopy had success and complication rates that were comparable to those of their cardiology colleagues. In a retrospective review of 43 ED patients in whom emergency transvenous pacemaker placement was attempted, there was a 95% success rate and no immediate or delayed complications.13

Indications

The purpose of cardiac pacing is to stimulate effective cardiac depolarization. In most cases the specific indications for cardiac pacing are clear; however, some areas are still controversial. The decision to pace on an emergency basis requires knowledge of the presence or absence of hemodynamic compromise, the cause of the rhythm disturbance, the status of the AV conduction system, and the type of dysrhythmia. The clinician caring for the patient is in the best position to decide on the value or nonvalue of pacing, based on nuances of the clinical scenario that are not possible to unravel by any theoretical

### Table 15.1 Four-Letter Pacemaker Code

<table>
<thead>
<tr>
<th>FIRST LETTER</th>
<th>SECOND LETTER</th>
<th>THIRD LETTER</th>
<th>FOURTH LETTER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chamber Paced</td>
<td>Chamber Sensed</td>
<td>Sensing Response</td>
<td>Programmability</td>
</tr>
<tr>
<td>A = Atrium</td>
<td>A = Atrium</td>
<td>T = Triggered</td>
<td>P = Simple</td>
</tr>
<tr>
<td>V = Ventricle</td>
<td>V = Ventricle</td>
<td>I = Inhibited</td>
<td>M = Multiprogrammable</td>
</tr>
<tr>
<td>D = Dual</td>
<td>D = Dual</td>
<td>D = Dual (A triggered and V inhibited)</td>
<td>R = Rate adaptive</td>
</tr>
<tr>
<td>O = None</td>
<td>O = None</td>
<td>O = None</td>
<td>C = Communicating</td>
</tr>
<tr>
<td>O = None</td>
<td>O = None</td>
<td>O = None</td>
<td>O = None</td>
</tr>
</tbody>
</table>

289
discussion. Controversy exists throughout the literature, and this discussion is not meant to set a standard of care for individual circumstances.

In general, the indications can be grouped into those that cause either tachycardias or bradycardias (see Review Box 15.1). Transcutaneous cardiac pacing (TCP) has become the mainstay of emergency cardiac pacing and is often used pending placement of a transvenous catheter or to determine whether potentially terminal bradyasystolic rhythms will respond to pacing.

**Bradycardias**

**Sinus Node Dysfunction**

Sinus node dysfunction may be manifested as sinus arrest, tachybrady (sick sinus) syndrome, or sinus bradycardia. Although symptomatic sinus node dysfunction is a common indication for elective permanent pacing, it is seldom cause for emergency pacemaker insertion.

Seventeen percent of patients with acute myocardial infarction (AMI) will experience sinus bradycardia.11 It occurs more frequently with inferior than with anterior infarction and has a relatively good prognosis when accompanied by a hemodynamically tolerable escape rhythm. However, sinus bradycardia is not a benign rhythm in this situation; it has a mortality rate of 2% with inferior infarction and 9% with anterior infarction.16 Sinus node dysfunction frequently responds to medical therapy but requires prompt pacing if such therapy fails.

**Asystolic Arrest**

Transvenous pacing in an asystolic or bradyasystolic patient has little value and is not recommended.15 In a study of 13 patients who had suffered cardiac arrest, capture of the myocardium was not present in 4 patients, but there were no survivors.13 Transvenous pacing alone may also not be effective for post-countershock pulseless bradyarrhythmias.19 This failure of pacing has likewise been demonstrated with transcutaneous pacemakers, thus suggesting that failure of effective pacing is primarily related to the state of the myocardial tissue.24 Cardiac pacing may be used as a “last-ditch” effort in bradyasystolic patients but is rarely successful and is not considered standard practice. Early pacing is essential when done for this purpose if success is to be achieved20 (see later in this section). Most importantly, given the continued emphasis on the importance of maximizing chest compressions during cardiopulmonary resuscitation (CPR), interrupting CPR to institute emergency pacing is not recommended.21

**AV Block**

AV block is the classic indication for pacemaker therapy. In symptomatic patients without myocardial infarction (MI) and in asymptomatic patients with a ventricular rate lower than 40 beats/min, pacemaker therapy is indicated.22

In patients with AMI, 15% to 19% progress to heart block: first-degree block develops in approximately 8%, second-degree block in 5%, and third-degree block in 6%.23 First-degree block progresses to second- or third-degree block 33% of the time, and second-degree block progresses to third-degree block approximately one-third of the time.21

AV block occurring during anterior infarction is believed to result from diffuse ischemia in the septum and infranodal conduction tissue. Because these patients tend to progress to high-degree block without warning, a pacemaker is often placed prophylactically. Some patients are prophylactically paced on a temporary basis, even in the absence of hemodynamic compromise.

During inferior infarction, early septal ischemia is the exception and typically block develops sequentially from first-degree to Mobitz type I second-degree and then to third-degree AV block. These conduction abnormalities frequently result in hemodynamically tolerable escape rhythms because of sparing of the bundle branches. A hemodynamically unstable patient who is unresponsive to medical therapy should be paced promptly. Whether and when stable patients should be paced is unclear, but placing a transcutaneous pacemaker is one option that can be attempted before placing a transvenous catheter.

**Trauma**

Pacing is not a standard intervention in traumatic cardiac arrest, but in selected cases it may be considered. Several rhythm and conduction disturbances have been documented in patients with nonpenetrating chest trauma. In these patients, traumatic injury to the specialized conduction system may predispose to life-threatening dysrhythmias and blocks that can be treated by cardiac pacing.23

Hypovolemia and hypotension can cause ischemia of conduction tissue and cardiac dysfunction.26 Marked bradycardias that persist even after vigorous volume replacement may rarely respond to cardiac pacing in patients with such trauma.27

**Bundle Branch Block and Ischemia**

Bundle branch block occurring in AMI is associated with a higher mortality rate and a greater incidence of third-degree heart block than is an uncomplicated infarction. Atkins and colleagues noted that 18% of patients with MI had a bundle branch block.24 Of these patients, complete heart block developed in 43% who had right bundle branch block (RBBB) and left axis deviation, in 17% who had left bundle branch block (LBBB), in 19% who had left anterior hemiblock, and in 6% who had no conduction block. The investigators concluded that RBBB with left axis deviation should be paced prophylactically.

A study by Hindman and associates confirmed the natural history of bundle branch block during MI.25 In their study, the presence or absence of first-degree AV block, the type of bundle branch block, and the age of the block (new versus old) were used to determine the relative risk for progression to type II second-degree or third-degree block (Table 15.2). Because of the increased risk, consider pacing for the following conduction blocks: new-onset LBBB, RBBB with left axis deviation or other bifascicular block, and alternating bundle branch block.26 Though controversial, one authority recommends prophylactic pacing for all new bundle branch blocks when MI is evident.25

Whether to place a transvenous pacemaker prophylactically in patients with LBBB before insertion of a flow-directed pulmonary artery catheter (PAC) remains controversial. Some researchers strongly advocate this procedure because of the risk for transient RBBB and life-threatening complete heart block associated with PAC placement.31 One study noted that this risk is low in patients with previous LBBB but continued to recommend temporary catheter placement for all cases of new LBBB.23 One solution to this problem is to place a transcutaneous pacemaker before catheterization as an emergency measure should heart block develop. In these cases a temporary transvenous pacemaker can be placed in a
TABLE 15.2 Influence of Different Variables on the Risk for High-Degree Atrioventricular Block in Patients With Bundle Branch Block During Myocardial Infarction

<table>
<thead>
<tr>
<th>Patients</th>
<th>Progressing to High-Grade AVB (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infarct location&lt;br&gt;Anterior</td>
<td>25</td>
</tr>
<tr>
<td>Indeterminate&lt;br&gt;</td>
<td>12</td>
</tr>
<tr>
<td>Inferior or posterior&lt;br&gt;</td>
<td>20</td>
</tr>
<tr>
<td>PR interval&lt;br&gt; &gt;0.20 sec&lt;br&gt;</td>
<td>25</td>
</tr>
<tr>
<td>≤0.20 sec&lt;br&gt;</td>
<td>19</td>
</tr>
<tr>
<td>Type of BBB&lt;br&gt;LBBB&lt;br&gt;</td>
<td>13</td>
</tr>
<tr>
<td>RBBB&lt;br&gt;</td>
<td>14</td>
</tr>
<tr>
<td>RBBB + LAFB&lt;br&gt;</td>
<td>27</td>
</tr>
<tr>
<td>RBBB + LPFB&lt;br&gt;</td>
<td>29</td>
</tr>
<tr>
<td>ABBB&lt;br&gt;</td>
<td>44</td>
</tr>
<tr>
<td>Onset of BBB&lt;br&gt; Definitely old&lt;br&gt;</td>
<td>13</td>
</tr>
<tr>
<td>Possibly new&lt;br&gt;</td>
<td>25</td>
</tr>
<tr>
<td>Probably new&lt;br&gt;</td>
<td>26</td>
</tr>
<tr>
<td>Definitely new&lt;br&gt;</td>
<td>23</td>
</tr>
</tbody>
</table>

ABB, Alternating bundle branch block; AVB, atrioventricular block; BBB, bundle branch block; LAFB, left anterior fascicular hemiblock; LBBB, left bundle branch block; LPFB, left posterior fascicular hemiblock; RBBB, right bundle branch block.


Tachycardias

Hemodynamically compromising tachycardias are usually treated with medications or electrical cardioversion. Since 1980 there has been increasing interest in pacing therapy for symptomatic tachycardias. Supraventricular dysrhythmias, with the exception of atrial fibrillation, respond well to atrial pacing. By “overdrive” pacing the atria at rates 10 to 20 beats/min faster than the underlying rhythm, the atria become entrained, and when the rate is slowed the rhythm frequently returns to normal sinus. A similar procedure is done for ventricular dysrhythmias. Overdrive pacing is especially useful for arrhythmias with recurrent prolonged QT intervals, such as those seen with quinidine toxicity or torsades de pointes. Though an attractive thought, there is no reported experience with these techniques in the ED. Transvenous pacing is also useful in patients with digitalis-induced dysrhythmias, in whom direct current cardioversion may be dangerous, or in patients in whom there is further concern about myocardial depression with drugs.

Cardiac Pacing for Drug-Induced Dysrhythmias

Significant dysrhythmias can be caused by excessive therapeutic medication (often in combination therapy) and overdose of cardioactive medications. Because these drugs have direct effects on cells of the myocardial pacemaker and conduction system, cardiac pacing is usually of little therapeutic value. Both bradycardias and tachycardias may result. Tachycardic rhythms from amphetamines, cocaine, anticholinergics, cyclic antidepressants, theophylline, and other drugs do not benefit from cardiac pacing. Drug-induced torsades de pointes may theoretically be overdriven by pacing, but data on this technique are lacking. Any drug that affects the central nervous system (e.g., opiates, sedative-hypnotics, clonidine) may produce bradycardia. Uncommon causes of toxin-induced bradycardia include organophosphate poisoning, various cholinergic drugs, ciguatera poisoning, and rarely, plant toxins. Cardiac pacing is not used for bradycardias from these sources; rather, the underlying central nervous system depression is addressed.

Severe bradycardia and heart block often accompany overdose of digitalis preparations, β-adrenergic blockers, and calcium channel blockers. Although intuitively attractive, cardiac pacing is not generally effective for serious toxin-induced bradycardias, even though there have been rare case reports of success. In β-blocker overdose, pacing may increase the heart rate but rarely benefits blood pressure or cardiac output. Worsening of blood pressure may occur as a result of loss of atrial contractions with ventricular pacing. Likewise, calcium channel blocker overdose and digitalis-induced bradycardia and heart block rarely benefit from cardiac pacing. Pharmacologic interventions, such as digoxin-specific Fab, glucagon, calcium, inotropic medications, and vasopressors, remain the mainstay in the treatment of drug-induced dysrhythmias. Given the lack of success of pacing, possible downsides, and the greater effectiveness of specific antidotes, it is not standard to routinely attempt transvenous cardiac pacing in the setting of drug overdose. However, as a last resort, cardiac pacing can be supported.

Contraindications

The presence of a prosthetic tricuspid valve is generally considered to be an absolute contraindication to trans venous cardiac pacing. In addition, severe hypothermia will occasionally result in ventricular fibrillation when pacing is attempted. Because ventricular fibrillation under these conditions is difficult to convert, caution is advised when considering pacing severely hypothermic and bradycardic patients. Rapid and careful rewarming is often recommended first, followed by pacing if the patient’s condition does not improve.

Equipment

Several items are required to insert a transvenous pacemaker adequately. Like most special procedures, a prearranged tray is convenient. The usual components required to insert a transvenous cardiac pacemaker are depicted in Review Box 15.1.
Pacing Generator

Many different pacing generators are available, but in general they all have the same basic features. The controls will frequently have a locking feature or cover to prevent the generator from being switched off or reprogrammed inadvertently. An amperage control allows the operator to vary the amount of electrical current delivered to the myocardium, usually 0.1 to 20 mA. Increasing the setting increases the output and improves the likelihood of capture. The pacing control mode is determined by adjusting the gain setting for the sensing function of the generator. By increasing the sensitivity, one can convert the unit from a fixed-rate (asynchronous mode) to a demand (synchronous mode) pacemaker. The typical pacing generator has a sensitivity setting that ranges from approximately 0.5 to 20 mV. The voltage setting represents the minimum strength of the electrical signal that the pacer is able to detect. Decreasing the setting increases the sensitivity and improves the likelihood of sensing myocardial depolarization. In the fixed-rate mode, the unit fires despite the underlying intrinsic rhythm; that is, the unit does not sense any intrinsic electrical activity. In the full-demand mode, however, the pacemaker senses the underlying ventricular depolarizations, and the unit does not fire as long as the patient’s ventricular rate is equal to or faster than the set rate of the pacing generator. A sensing indicator meter and a rate control knob are also present.

Temporary pacing generators are battery operated, and thus it is always good practice to install a fresh battery whenever pacing is anticipated. An example of a pacing generator is shown in Fig. 15.1.

Pacing Catheters and Electrodes

Several sizes and brands of pacing catheters are available. In general, most range from 3 to 5 Fr in size and are approximately 100 cm in length. Lines are marked along the catheter surface at approximately 10-cm intervals and can be used to estimate catheter position during insertion. Pacing catheters differ with respect to their stiffness, electrode configurations, floating characteristics, and other qualities. For emergency pacing, the semifloating bipolar electrode catheter with a balloon tip is used most frequently (Fig. 15.2). The balloon holds approximately 1.5 mL of air, and the air injection port has a locking lever to secure balloon expansion. Before insertion, check the balloon for leakage of air by inflating and immersing it in sterile water. The presence of an air leak is noted by a stream of bubbles rising to the surface of the water. An inflated balloon helps the catheter “float” into the heart, even in low-flow states, but is not advantageous in the cardiac arrest situation.

For all practical purposes, temporary transvenous pacing is accomplished with a bipolar pacing catheter. The terms unipolar and bipolar refer to the number of electrodes in contact with the portion of the heart that is to be stimulated. All pacemaker systems must have both a positive (anode) and a negative (cathode) electrode; hence, all stimulation is bipolar. In the typical bipolar catheter used for temporary transvenous pacing, the cathode (stimulating electrode) is at the(919,522),(983,625) of the pacing catheter. The anode is located 1 to 2 cm proximal to the tip, and a balloon or an insulated wire separates the two electrodes. The distinction between unipolar and bipolar pacing catheters is that a bipolar catheter has both electrodes in relatively close proximity on the catheter and both may contact the endocardium. In a bipolar catheter, the electrodes are usually stainless steel or platinum rings that encircle the pacing catheter. When properly positioned, both electrodes will be within the right ventricle so that a field of electrical excitation is set up between the electrodes. With a bipolar catheter, the cathode does not need to be in direct contact with the endocardium for pacing to occur, although it is preferable to have direct contact.

A unipolar system is also effective but is used infrequently for temporary transvenous pacing. In a unipolar system, the cathode is at the tip of the pacing catheter and the anode is located in one of three places: in the pacing generator itself, more proximally on the catheter (outside the ventricle), or on the patient’s chest. A bipolar system may be converted to a unipolar system by simply disconnecting the positive proximal connection of the bipolar catheter from the pacing generator and running a new wire from the positive (pacing generator) terminal to the patient’s chest wall. Such a conversion may be required in the unlikely event of failure of one lead of the bipolar system.
**ECG Machine**

An ECG machine can be used to record the heart’s inherent electrical activity during insertion of the pacer and to aid in localization of the tip of the catheter without fluoroscopy. The ECG machine must be well grounded to prevent leakage of alternating current, which can cause ventricular fibrillation. Such leakage should be suspected if interference of 50 to 60 cycles per second (Hz) is noted on the ECG tracing.

The ECG machine should be placed in a manner that allows easy visibility of the rhythm during insertion. One method is to place the machine near the level of the patient’s midthorax, facing the operator, on either side of the patient as logistics and operator preference allow (Fig. 15.3). Note that the operator stands at the head of the patient during passage of the catheter through the internal jugular or subclavian vein and at the midabdomen for insertion through the femoral or brachiocephalic vein. Newer patient monitors may be equipped with suitable ECG connections to allow their use in place of a stand-alone ECG machine. Because these patients will already be attached to a monitor, it may prove convenient to use the same piece of equipment to assist in insertion of the pacemaker.

**Introducer Sheath**

An introducer set or sheath is required for venous access (see Chapter 22). Some pacing catheters are prepackaged with the appropriate equipment, whereas others require a separate set. The introducer set is used to enhance passage of the pacing catheter through the skin, subcutaneous tissue, and vessel wall. The sheath must be larger than the pacing catheter to allow it to pass. The size of the pacing catheter refers to its outside diameter, whereas the size of the introducer refers to its inside diameter. Thus, a 5-Fr pacing catheter will fit through a 5-Fr introducer. Introducer sheaths are available with a perforated elastic seal covering the opening through which the pacing catheter is passed (pacer port). The seal allows the catheter to be manipulated while preventing blood from escaping or air from entering the vein. A side port allows the sheath to be used for central venous access. A makeshift sheath can be fashioned with an appropriately sized intravenous (IV) catheter. A 3-Fr balloon-tipped catheter will fit through a 14-gauge IV catheter is suitable. A 4-Fr balloon-tipped catheter will also fit through a 14-gauge catheter or needle. However, without a seal over the hub, blood will leak from the end of the IV catheter.

Overall, the key to success with this procedure is preparation. In a typical ED there are often a variety of vascular access kits and devices, not all of which will work well, if at all, for passing a pacing catheter. It is imperative that one examine all the components of the tray before starting the procedure to ensure that all wires, sheaths, dilators, and syringes fit as expected. Ideally, all the equipment and accessories needed for emergency pacemaker insertion should be kept together in a designated location.

**Procedure**

A checklist for the preparation and initial setup of a pacing generator is shown in Box 15.1. It may be useful to have a copy of this checklist or a similar list stored with the pacemaker, to have on hand in emergency situations.

**Patient Preparation**

Patient instruction is an extremely important aspect of any procedure. Frequently, there is not enough time to give patients a detailed explanation or to obtain written informed consent. Nonetheless, sufficient information should be provided so that the patient feels at ease. It is always prudent to obtain and document informed consent from the patient, if possible, before any invasive procedure or to document that the circumstances

---

**Figure 15.3** Position of operator and electrocardiographic device during insertion of a pacemaker catheter through the left subclavian vein (a preferred site).
The left subclavian vein is preferred because of the less acute angle traversed than with the right-sided approach, but either side may be used.\textsuperscript{46,47} The internal jugular approach may also be used. In this case, the right internal jugular vein is preferred because of the direct line to the superior vena cava. Problems with this approach include dislodgment of the pacemaker with movement of the head, puncture of the carotid artery, and thrombophlebitis.

During CPR, use of the right internal jugular vein and the left subclavian vein for pacemaker insertion has been demonstrated to result in the highest rates of proper placement.
in the right ventricle. The right internal jugular vein is the more direct route of the two and may be the most appropriate site.

Femoral veins, like neck veins, are compressible and easily catheterized. Problems include easy dislodgement, infection, and increased risk for thrombophlebitis.

Brachial vein catheterization is easy to perform but results in a high incidence of infection and vessel thrombosis. In addition, the catheter is easily dislodged with arm motion. This approach is seldom used in the emergency setting.

Although the left subclavian and right internal jugular veins are the preferred routes for access, in emergency situations clinicians should use the approach with which they are most experienced to minimize the time spent cannulating the vein and to reduce the potential for complications from the venipuncture.

**Skin Preparation and Venous Access**

Clean the skin over the venipuncture site twice with an antiseptic solution such as chlorhexidine or povidone-iodine. Prepare a wide area because of the tendency for guidewires and catheters to spring from the hands of an unsuspecting operator. Similarly, drape widely in the standard manner to maintain a sterile field and to allow clear visibility of the venipuncture site.

The infraclavicular approach is used in this chapter to illustrate venous access, although the mechanics are generally the same as for other vascular approaches.

Occasionally, a patient who already has a central venous line in place requires emergency placement of a pacing catheter. An existing central venous pressure (CVP) line can be used to place the pacing catheter if the lumen of the catheter is large enough to accept a guidewire. Withdraw the CVP line 3 to 5 cm to expose an area of sterile tubing. Transsect the tubing through a sterile area while holding it firmly at skin level. Pass a guidewire through the tubing, and then withdraw the tubing so that only the wire is left in the vein. Never release the guidewire and the tubing because embolization may result.

With the guidewire in place, pass a dilator and introducer sheath together over the guidewire, as is done in the Seldinger technique. Remove the dilator and guidewire and pass the pacing catheter through the introducer sheath (Fig. 15.5, step 1). One key additional step to help preserve sterility while manipulating the pacing catheter is to attach an extensible sleeve on the end of the introducer before inserting the pacing catheter (see Fig. 15.5, steps 1 and 6). In this way the pacing catheter can be advanced and withdrawn multiple times without fear of contamination.

Bedside ultrasound (US) can be useful as an aid in securing central venous access, and its use in the setting of emergency transvenous pacing has been reported.

**Pacemaker Placement**

**Emergency Blind Passage**

A transvenous pacemaker wire can be advanced blindly, or under ECG guidance. In an emergency, blind advancement is preferred. The right internal jugular route is preferred. Attach the pacing wire to the pacemaker energy source and turn it on at a rate of 80 beats/min, output at 20 mA of current, and set the asynchrony (fixed rate) mode. After the wire is blindly advanced to the 20-cm depth (two hash marks on the wire) inflate the pacing-wire balloon and advance the pacing wire while viewing the patient’s cardiac monitor. When capture is seen on the ECG monitor by observing a pacer spike followed by QRS ST segment elevation, the pacing wire is in the proper position. Deflate the balloon and secure the pacing wire at the skin. When time permits, or if the blind passage procedure is not successful, attach the pacing wire to the ECG V1 lead with an alligator clamp. Turn on the ECG machine to monitor the V1 lead and hence the position of the catheter tip. The following is a detailed summary of that ECG-guided procedure.

**ECG Guidance**

Connect the patient to the limb leads of an ECG machine, and turn the indicator to record the chest (V) lead.

With newer ECG machines, the pacemaker may be attached to any of the V leads (usually V1 or V5) that are displayed during rhythm monitoring. The distal tip of the pacing catheter (the cathode or lead marked “negative”, “−”, or “distal”) must be connected to the V lead of the ECG machine by a male-to-male connector or by an insulated wire with an alligator clip on each end. Some prepackaged kits contain an alligator clamp that can be connected to the lead with an adapter pin (see Fig. 15.5, step 2). The pacing catheter is thus an exploring electrode that creates a unipolar electrode for intracardiac ECG recording. The ECG tracing recorded from the electrode tip localizes the position of the tip of the pacing electrode. Because the tracing on the ECG machine may be slightly delayed, advancement of the catheter after initial insertion must be evaluated carefully. If a balloon-tipped catheter is used, inflate the balloon with air after the catheter enters the superior vena cava (= 10 to 12 cm for a subclavian or internal jugular insertion) (see Fig. 15.5, step 3). The inflation port should be locked and the syringe left attached.

Advance the pacing catheter both quickly and smoothly. Monitor the ECG V lead, and observe the P wave and QRS complex to ascertain the location of the tip of the pacing catheter (see Fig. 15.5, step 4). Use of electrocardiography to guide placement of a pacing catheter is based on two concepts. First, the complex will vary in size depending on which chamber is entered. For example, when the tip of the pacing catheter is in the atrium, one will see large P waves, often larger than the corresponding QRS complex. Second, the sum of the electrical forces will be negative if the depolarization is moving away from the catheter tip and positive if the depolarization is moving toward the catheter tip. Therefore if the tip of the catheter is above the atrium, both the P wave and QRS complex will be negative (i.e., the electrical forces of a normally beating heart will be moving away from the catheter tip). As the tip progresses inferiorly in the atrium, the P wave will become isoelectric (biphasic) and will eventually become positive as the wave of atrial depolarization advances toward the tip of the catheter. The ECG tracing resembles an aVR lead initially when in the left subclavian vein (Fig. 15.6D) or the midportion of the superior vena cava (see Fig. 15.6B). At the high right atrial level, both the P wave and QRS complex are negative. The P wave is larger than the QRS complex and is deeply inverted (see Fig. 15.6C and D). As the center of the atrium is approached, the P wave becomes larger and biphasic (see Fig. 15.6E). As the catheter approaches the lower atrium (see Fig. 15.6F), the P wave becomes smaller and upright. The QRS complex is fairly normal. When striking the right atrial wall, an injury pattern with a P-Ta segment is seen (see Fig. 15.6G). As the electrode passes through the tricuspid valve, the P wave becomes smaller and the QRS complex becomes larger (see Fig. 15.6H). Placement in the inferior vena cava may be recognized by a change in the morphology of the P
Emergency Transvenous Cardiac Pacing

1. Establish a sheath introducer in the selected vein. Attach the still-compressed sterile sleeve to the introducer hub. Check the balloon for integrity and then advance the catheter into the sleeve. (Full sterile drapes should be used but are not depicted here.)

2. Instruct an assistant to make the following nonsterile connections. Attach the proximal (+) lead to the positive terminal of the connecting cable and the distal (−) lead to the V1 lead of an ECG machine with an alligator clamp. (Alternatively, an insulated wire with an alligator clip on each end may be used.)

3. Advance the catheter approximately 10 to 12 cm so that the tip lies within the superior vena cava. Then inflate the balloon with 1.5 mL of air. Close the stopcock valve to keep the balloon inflated.

4. Advance the pacing catheter quickly and smoothly. Monitor the V1 lead on the electrocardiogram to ascertain the location of the tip of the pacing catheter. The P wave and QRS complex will vary in size depending on which chamber the tip is in, and the sum of the electrical forces will be negative if depolarization is moving away from the catheter tip and positive if depolarization is moving toward it. ST-segment elevation will occur when the tip contacts the endocardium.

5. When the pacing catheter is in the desired position, deflate the balloon by unlocking the stopcock and allow the syringe to spontaneously refill with air.

6. Extend the sterile sleeve so that it fully covers the pacing catheter. If your sheath and sleeve have valves, close them by turning clockwise to keep the wire and sleeve in place.

Figure 15.5 Emergency transvenous cardiac pacing. Note that by attaching the pacing catheter to the electrocardiographic (ECG) machine (step 2) and observing the V1 ECG tracing, the negative lead now becomes an exploring lead that tells the operator the position of the tip of the pacing catheter in the body (see Fig. 15.6).
Emergency Transvenous Cardiac Pacing, CONTINUED

Disconnect the catheter's distal (negative) lead from the ECG machine and attach it to the energy source via the connector cable. Set the rate to 80 beats/min, the output to 5 mA, and the sensitivity to full-demand mode (most sensitivity), and turn the pacer on.

Assess for electrical capture by looking for a QRS complex to follow each pacer spike on the ECG monitor. Assess for mechanical capture by checking for a palpable pulse that equals the pacemaker rate. Reposition the catheter if needed.

Testing Threshold
a. Set rate to 80 beats/min.
b. Set output to 5 mA.
c. Set sensitivity to maximum.
d. Reduce output slowly until capture is lost. This output is the threshold.
e. Increase output to 2.5 times the threshold to ensure consistency of capture (usually 2 to 3 mA).

Testing Sensing
a. Set the rate at 10 beats/min greater than intrinsic rhythm.
b. Place the pacemaker in asynchronous mode (minimum sensitivity) and ensure that there is complete capture.
c. Adjust the sensitivity to midposition (approx. 3 mV).
d. Decrease the rate until pacing is suppressed by the intrinsic rhythm.
e. Check that the sensing indicator signals each time that a native beat is sensed.
f. If the pacer fails to sense an intrinsic rhythm, increase the sensitivity.
g. If the pacer oversenses (e.g., triggered by P or T waves or artifact), decrease the sensitivity.
h. Once the sensitivity threshold is determined, set the millivoltage to half that value.

Figure 15.5, cont’d

wave and a decrease in the amplitude of both the P wave and the QRS complex (see Fig. 15.6).

Once the pacing catheter is in the desired position, deflate the balloon by unlocking the port, observe that the syringe refills with air spontaneously, and then remove the syringe (see Fig. 15.5, step 5). Avoid drawing back on the syringe because this may rupture the balloon. If the syringe does not refill spontaneously, the balloon might be ruptured. Do not inflate the balloon. Withdraw the pacing catheter, and check the balloon for leaks. If a leak is found, use a new pacing catheter.

After successful passage of the catheter into the right ventricle, advance the tip until contact is made with the endocardial wall. When this occurs, the QRS segment will show ST-segment elevation (see Fig. 15.6). Ideally, the tip of the catheter should be lodged in the trabeculae at the apex of the right ventricle; however, pacing may also be successful if the catheter is in various other positions within the ventricle or outflow tract.

If the pacer enters the pulmonary artery outflow tract, the P wave again becomes negative, and the QRS amplitude diminishes (see Fig. 15.6K). If the catheter is in the pulmonary artery, withdraw the pacing catheter into the right ventricle and readvance it. Sometimes a clockwise or counterclockwise twist of the catheter will redirect its path in a more favorable direction. If catheter-induced ectopy develops, withdraw the catheter slightly until the ectopy stops, and then readvance.
it. Occasionally, an antidysrhythmic drug such as lidocaine may be needed to desensitize the myocardium. Once ventricular endocardial contact is made, disconnect the catheter from the ECG machine and connect the distal lead to the negative terminal on the pacing generator (see Fig. 15.5, step 7). Set the pacing generator at a rate of 80 beats/min or 10 beats/min faster than the underlying ventricular rhythm, whichever is higher. Select the full-demand mode with an output of approximately 5 mA. Then turn on the pacing generator.
Assess the patient for electrical and mechanical capture. Electrical capture will be manifested on the ECG monitor as a pacer spike followed by a QRS complex (see Fig. 15.5, step 8). If a pacer spike is seen but no QRS follows, capture is not occurring. Mechanical capture means that a pacer spike with its corresponding QRS triggers a myocardial contraction. This can be assessed by checking that a palpable pulse is present and equal to the rate set on the pacemaker. If complete capture does not occur or if it is intermittent, the pacer will need to be repositioned. When proper capture occurs, assess the pacer for optimal positioning. This is done by testing the thresholds for pacing and sensing, and by physical examination, electrocardiography, and chest radiography.

Catheter Placement in Low-Flow States
If cardiac output is too low to float a pacing catheter or if the patient is in extremis, there may not be enough time to advance a pacing catheter via the previously described techniques. Such a situation would be asystole or complete heart block with malignant ventricular escape rhythms (although one can make a case for TCP in such conditions). In such emergency situations, connect the pacing catheter to the energy source, turn the output to the maximum amperage, and select the asynchronous mode (see earlier section on Emergency Blind Passage). Advance the catheter blindly in the hope that it will enter the right ventricle and pacing will be accomplished. Rotate, advance, withdraw, or otherwise manipulate the pacing catheter according to clinical response. The right internal jugular approach is the most practical access route in this situation.

US Guidance
As bedside US has become more widely available in the ED, new uses have been discovered. One promising technique involves using US to assist in the placement of emergency transvenous pacing catheters. US may also help demonstrate whether mechanical capture has been achieved. The advantages of US over fluoroscopy are: no exposure to ionizing radiation, lower rates of complications, and faster times to pacemaker insertion. Further experience will be necessary to confirm its utility (Ultrasound Box 15.1).

Testing Threshold
The threshold is the minimum current necessary to obtain capture. Ideally, it is less than 1.0 mA, and usually it is between 0.3 and 0.7 mA. If the threshold is in this ideal range, good contact with the endocardium can be presumed.

To determine the threshold, set the pacing generator to maximum sensitivity (full-demand mode) at 5-mA output and a rate of approximately 80 beats/min (or at least 10 beats/min greater than the patient’s intrinsic rate) (see Fig. 15.5, step 9). Reduce the current (output) slowly until capture is lost. This current is the threshold. Carry out this maneuver two or three times to ensure that this value is consistent. Increase the amperage to 2.5 times the threshold to ensure consistency of capture (usually between 2 and 3 mA).

Testing Sensing
Test the sensing function in patients who have underlying rhythms. Set the rate at approximately 10 beats/min greater than the endogenous rhythm, place the pacemaker in asynchronous mode (minimum sensitivity, which is the maximum setting on the sensitivity voltage control), and ensure that there is complete capture (see Fig. 15.5, step 10). Then adjust the sensitivity control to its midposition or approximately 3 mV, and gradually decrease the rate until pacing is suppressed by the patient’s intrinsic rhythm. The sensing indicator on the pacing generator should signal each time a native beat is sensed and should be in synchrony with each QRS complex on the ECG monitor. If the pacer fails to sense the intrinsic rhythm, increase the sensitivity (decrease the millivolts) until the pacer is suppressed. Conversely, if the sensing indicator is triggered by P or T waves or by artifact, decrease the sensitivity until only the QRS complex is sensed. Once the sensitivity threshold is determined, set the millivoltage to approximately half that value.

Securing and Final Assessment
After the pacemaker's position has been tested for electrical accuracy, it must be secured in place. If a sealed introducer sheath was used, the hub should be fixed firmly to the skin with suture (e.g., 4-0 nylon or silk). A fastening suture should be sewn to the skin and the hub tied securely in place. If a plain introducer was used, withdraw it to prevent leakage and suture the catheter in place. In either case, coil the excess pacing catheter and secure it in a sterile manner underneath a large sterile dressing. Assess pacemaker function again, and take a chest film to ensure proper positioning. Ideal positioning of the pacing catheter is at the apex of the right ventricle (Fig. 15.7).

A 12-lead ECG tracing should be obtained after placement of the transvenous pacemaker. If the catheter is within the right ventricle, an LBBB pattern with left axis deviation should be evident in paced beats (Fig. 15.8). If an RBBB pattern is noted, coronary sinus placement or left ventricular pacing secondary to septal penetration should be suspected (Fig. 15.9).

With a properly functioning ventricular pacemaker, large cannon waves may be noted on inspection of the venous pulsations at the neck. When the pacemaker achieves ventricular capture, there may be times when the atria contract against a closed tricuspid valve and a cannon wave results. On auscultation of the heart a slight murmur may be evident, secondary to tricuspid insufficiency as a result of the catheter interfering with the tricuspid valve apparatus. Following each pacemaker impulse a clicking sound may be heard during expiration; this is believed to represent either intercostal or diaphragmatic muscular contractions caused by the pacemaker. Note that this can also be a sign of cardiac perforation. On auscultation of the second heart sound, paradoxical splitting may be noted. This represents a delay in closure of the aortic valve because of delayed left ventricular depolarization.

As in any procedure, assess the patient for improvement in clinical status. Evaluate vital signs, mentation, congestive symptoms, and urinary output. In addition, look for complications secondary to the procedure and treat as needed.

Complications
The complications associated with emergency transvenous cardiac pacing are numerous and represent a compendium of those related to central venous catheterization, those related to right-sided heart catheterization, and those unique to the pacing catheter itself. Overall rates of adverse events using modern flexible, balloon-tipped catheters are in the range of 20%. However, the majority of complications are a result of obtaining venous access, and when US guidance is used the rate is markedly decreased.
Using ultrasound to guide placement of a transvenous pacemaker (TVPM) gives the physician the advantage of direct visualization of the tip of the pacemaker and the ability to directly maneuver it into the correct position to obtain capture. Although placement of a TVPM is a relatively rare procedure in the emergency department, the use of ultrasound has been described to facilitate the procedure. Ultrasound can also be used to evaluate placement in cases in which blind attempts at transvenous pacing are unsuccessful. Additionally, ultrasound is useful in achieving the central venous access necessary to introduce the pacemaker.

**Equipment**

A low-frequency (2 to 5 mHz) transducer offers the optimum depth to evaluate the heart. A phased-array or microconvex transducer is ideal for this purpose because its small footprint enables the sonographer to "see" between the patient’s ribs. In the absence of a phased-array or microconvex transducer, a curvilinear transducer may be adequate. Although insertion of a TVPM is a sterile procedure, the transducer does not need to be sterile if it is used away from the field. If the transducer will be used near or on the field, it can be placed inside a sterile cover or a sterile glove.

**Image Interpretation**

The subxiphoid view of the heart is the optimum view for this procedure (Fig. 15.US1). It enables the operator to view all four chambers of the heart at once. The subxiphoid view is obtained by placing the transducer just inferior to the xiphoid process with the indicator pointing toward the patient’s right side (Fig. 15.US2). A hand should be placed on top of the transducer to enable the sonographer to push downward into the epigastric area. The transducer can then be aimed toward the left side of the chest until the four-chambered view of the heart is seen. In this view the left lobe of the liver is seen at the top of the image. Deep to this is the heart, with the right atrium and ventricle abutting the liver.

**Figure 15.US1** Ultrasound subxiphoid image of the heart. In this ultrasound the liver is seen at the top of the image. The heart is seen below, with the right ventricle (RV), right atrium (RA), and left atrium (LA) abutting the left lobe of the liver. The left ventricle (LV) and atria are deep to the right side at the bottom of the image.

**Figure 15.US2** Placement of the ultrasound transducer to obtain a subxiphoid image of the heart. (Courtesy Christy Butts, MD.)

**Figure 15.US3** Pacing wire within the right ventricle (arrow). A, Sonographic image. B, Schematic representation. The right ventricle is seen through the subxiphoid window, which provides excellent views of the heart without interfering with placement of the pacemaker line. The pacemaker wire is seen as a brightly echogenic structure within the right ventricle.
Problems Related to Central Venous Catheterization

Inadvertent arterial puncture is a well-known complication of the percutaneous approach to the venous system. This problem is usually recognized quickly because of the rapid return of arterial blood. Firm compression over the puncture site will almost always result in hemostasis in 5 minutes or less.

Venous thrombosis and thrombophlebitis are also potential problems with central venous catheterization. Thrombophlebitis, which occurs early after insertion, is an uncommon complication. Thrombosis of the innominate vein is also a rare problem, with pulmonary embolism being an even more uncommon event. Femoral vein thrombosis, however, appears to be a much more common event associated with femoral vein catheterization. Studies using noninvasive techniques have shown a 37% incidence of femoral vein thrombosis, with 55% of these patients having evidence of pulmonary embolism on ventilation-perfusion scans. At least one study suggests that the use of low-dose enoxaparin is safe and effective in reducing pacemaker-related femoral vein thrombosis.

Pneumothorax is consistently a problem with the various approaches to the veins at the base of the neck. The decision to place a chest tube in patients with this complication depends on the extent of the air leak and the clinical status of the patient. In addition, laceration of the subclavian vein with hemothorax, laceration of the thoracic duct with chylothorax, air embolism, wound infections, pneumomediastinum, hydromediastinum, hemomediastinum, phrenic nerve injury, and fracture of the guidewire with embolization are all potential complications.

Complications of Right-Sided Heart Catheterization

A frequent complication of the pacing catheter is dysrhythmia, with premature ventricular contractions being a common occurrence. One study noted a 1.5% incidence of serious dysrhythmias with a balloon-tipped catheter inserted under ECG guidance, as opposed to a 32% incidence with insertion of a semirigid catheter under fluoroscopic guidance, thus suggesting that the balloon catheter was the preferred type of catheter. Another study noted a 6% incidence of ventricular tachycardia during insertion. An ischemic heart is more prone to dysrhythmias than a nonischemic heart. Therapy for catheter-induced ectopy during insertion involves repositioning the catheter in the ventricle. This usually stops the ectopy; however, if after repeated attempts it is found that the catheter cannot be passed without ectopy, myocardial suppressant therapy may be used to desensitize the myocardium.

Misplacement of the pacing catheter has been well studied. Passage of the catheter into the pulmonary artery can be
Figure 15.8 Electrocardiographic pattern of a right ventricular pacemaker.

Figure 15.9 Coronary sinus pacing. Note the paced right bundle branch block pattern.
diagnosed electrocardiographically by observing the return of an inverted P wave and a decrease in the voltage of the QRS complex. Misplacement in the coronary sinus may occur and should be suspected in patients in whom a paced RBBB pattern on the electrocardiogram is seen with right ventricular pacing (see Fig. 15.9). Rarely, an RBBB pattern can be seen with a normal right ventricular position; therefore all RBBB patterns do not represent coronary sinus pacing. Further evidence of coronary sinus location can be obtained by viewing the lateral chest film. Normally the tip of the catheter should point anteriorly toward the apex of the heart; however, with placement in the coronary sinus, the tip of the catheter is displaced posteriorly and several centimeters away from the sternum (Fig. 15.10). Other potential forms of misplacement include left ventricular pacing through an atrial or ventricular septal defect, septal puncture, extraluminal insertion, and arterial insertion.

Perforation of the ventricle is a well-described complication that can result in loss of capture, hemopericardium, and tamponade. Reported symptoms and signs of this problem include chest pain, pericardial friction rub, and diaphragmatic or chest wall muscular pacing. At least one case of a postpericardiotomy-like syndrome and two cases of endocardial friction rub have been reported without perforation.

Pericardial perforation is suggested radiographically when the pacing catheter is outside or abuts the cardiac silhouette and is not in proper position within the right ventricular cavity (Fig. 15.11). ECG clues include a change in the QRS complex and T-wave axis or failure to properly sense. In suspected cases a two-dimensional echocardiogram usually demonstrates the catheter’s extracardiac position. Simply pulling the catheter back and repositioning it in the right ventricle can usually treat uncomplicated perforation.

During insertion of a temporary pacing catheter when a nonfunctioning permanent catheter is in place, there is a small risk of entanglement or knotting. This potential also exists with other central lines and PACs. Even without the presence of other lines, the pacing catheter can become knotted. Frequently, these lines can be untangled under fluoroscopy with the use of specialized catheters.

Local and systemic infection, balloon rupture, pulmonary infarction, phrenic nerve pacing, and rupture of the chordae tendineae are also potential complications.

Complications of the Pacing Electrode

Complications related to the pacing electrode can be separated into three groups: mechanical, organic, and electrical.

Mechanical failures include displacement, fracture of the catheter, and loose leads. Displacement can result in intermittent or complete loss of capture or improper sensing, malignant dysrhythmias, diaphragmatic pacing, or perforation. Displacement should be suspected with changes in amplitude, with changes of greater than 90 degrees in vector, or with a change in threshold. Frequent catheter fractures may be detected by careful review of the chest film or may be suspected because of a change in the sensing threshold. As with displacement, catheter fractures may result in intermittent or complete loss of capture.

Organic causes of pacemaker failure result in changes in the threshold or sensing function. Progressive inflammation, fibrosis, and thrombosis may result in more than doubling of the original threshold. However, this process takes several weeks and is of no concern in the setting of pacemaker placement in the ED.

Electrical problems with pacing in the past have included failure of the pacemaker generator, dysrhythmias, and outside interference. Modern devices are extremely reliable and resistant to outside interference. Although ventricular tachycardia and ventricular fibrillation have been reported to result from pacemakers, these dysrhythmias are rare. Therefore patients with such dysrhythmias should be evaluated for a non–pacemaker-induced cause. Defibrillation and cardioversion are safe in patients who have temporary pacemakers.

EMERGENCY TCP

TCP is a rapid, minimally invasive method of emergency cardiac pacing that may temporarily substitute for transvenous pacing. Electrodes are applied to the skin of the anterior and posterior chest walls, and pacing is initiated with a portable pulse.
Transcutaneous Cardiac Pacing

**Indications**
General indications are identical to those for transvenous pacing (see Review Box 15.1)
Initial stabilization of patients requiring pacing while arrangements for transvenous pacing are being made
Pacing in various environments (prehospital, hospital ward, etc.)
Pacing in patients treated with thrombolytics or other anticoagulants

**Contraindications**
No absolute contraindications

**Complications**
Failure to recognize the presence of underlying treatable ventricular fibrillation
Induction of ventricular fibrillation (rare)
Soft tissue discomfort

---

**Equipment**

---

Generator. In an emergency setting, this pacing technique is faster and easier to initiate than transvenous pacing. Pulse generators are sufficiently portable to be used in EDs, hospital wards, intensive care units, and mobile paramedic vehicles.

In 1872, Duchenne de Boulogne reported successful resuscitation of a child by attaching one electrode to a limb while a second electrode was rhythmically touched to the precordium of the thorax. Successful overdrive pacing of the human heart with a precordial electrode was reported by VonZiemssen in 1882.

In 1952, Zoll introduced the first practical means of TCP. Using a ground electrode attached to the skin and a subcutaneous needle electrode over the precordium, he reported successful resuscitation of two patients in ventricular standstill. One patient was paced for 5 days and subsequently discharged from the hospital. Zoll later introduced a machine that delivered impulses lasting 2 msec through 3-cm diameter metal paddles pressed firmly against the anterior chest wall. This device was the first commercial transcutaneous cardiac pacemaker. During the 1950s, both Zoll and coworkers and Leatham and colleagues demonstrated the effectiveness of TCP in patients with bradycardia and asystole. Leatham and colleagues used larger electrodes (4 x 6 cm) and a longer pulse duration (20 msec) to successfully pace two patients with bradydysrhythmias.

Until the late 1950s, TCP was the only clinically accepted method of cardiac pacing. The original technique involving bare metal electrodes had adverse effects, including local tissue burns, muscle contraction, and severe pain. With the development of the first implantable pacemakers from 1958 to 1960, and the improvement in transvenous electrodes during the early 1960s, TCP was rapidly discarded.

Refinements in electrode size and pulse characteristics led to the reintroduction of TCP into clinical practice. Increasing the pulse duration from 2 to 20 msec or longer was found to decrease the current output required for cardiac capture. Longer impulse durations also make induction of ventricular fibrillation less likely. Electrodes with a larger surface area (80 to 100 cm²) decrease the current density at the underlying skin and therefore decrease pain and the possibility of tissue burns.

**Indications and Contraindications**

General indications for cardiac pacing were discussed earlier. TCP is the fastest and easiest method of emergency pacing. This technique is useful for initial stabilization of patients in the ED who require emergency pacing while arrangements or decisions about transvenous pacemaker insertion are being made. The equipment is readily mastered, the procedure is fast, and it is minimally invasive. TCP is also widely used in the prehospital environment, as well as in the hospital in the cardiac catheterization laboratory, operating room, intensive care units, and on general medical floors. The technique may be preferable to transvenous pacing in patients who have received thrombolytic agents or other anticoagulants. No central venous puncture, with the attendant risk of hemorrhage, is required. Limited experience suggests that TCP may also be useful in the treatment of refractory tachydysrhythmias by overdrive pacing. Although small pediatric electrodes for TCP have been developed, experience with pediatric TCP has been limited.

TCP is indicated for the treatment of hemodynamically significant bradydysrhythmias that have not responded to medical therapy. Hemodynamically significant implies hypotension, anginal chest pain, pulmonary edema, or evidence of decreased cerebral perfusion. This technique is temporary and is indicated for short intervals as a bridge until transvenous pacing can be initiated or the underlying cause of the bradydysrhythmia (e.g., hyperkalemia, drug overdose) can be reversed. Though generally unsuccessful, TCP may be attempted for the treatment of asystolic cardiac arrest. In this setting the technique is efficacious only if used early after the onset of arrest (usually within 10 minutes). TCP is not
indicated for the treatment of prolonged arrest victims with a final morbid rhythm of asystole.\textsuperscript{111,118-120}

Delay from the onset of arrest to the initiation of pacing is a major problem that limits the usefulness of TCP in prehospital care. Hedges and associates reported that the everyday availability of pacing increased the number of patients who underwent pacing within 10 minutes of hemodynamic decompensation and also increased long-term patient survival.\textsuperscript{117} Prehospital pacing may be most useful in the treatment of patients with a hemodynamically significant bradycardia who have not yet progressed to cardiac arrest (e.g., heart block in the setting of AMI) or in patients who arrest after the arrival of prehospital providers.\textsuperscript{116,117}

In conscious patients with hemodynamically stable bradycardia, TCP may not be necessary. It is reasonable to attach electrodes to such patients and to leave the pacemaker in standby mode against the possibility of hemodynamic deterioration while further efforts at treatment of the patient’s underlying disorder are being made. This approach has been used successfully in patients with new heart block in the setting of cardiac ischemia.\textsuperscript{121} Generally, when a transvenous pacemaker becomes available it is preferred because of better patient tolerance.

**Sedation/Analgesia**

Most patients cannot tolerate TCP for prolonged periods because they will sense each depolarization spike. Hence, TCP is often rapidly replaced by transvenous pacing. Standard sedatives (such as benzodiazepines) and opioids (such as morphine) should be administered as needed to control anxiety and discomfort.

**Equipment**

Since their reintroduction, transcutaneous pacemakers have undergone rapid evolution and are now standard equipment in most EDs, as well as in other hospital settings and the prehospital environment. The pacemakers introduced in the early 1980s tended to be asynchronous devices with a limited selection of rate and output parameters. Units introduced more recently have demand mode pacing and more output options and are often combined with a defibrillator in a single unit. Combined defibrillator-pacemakers offer advantages in cost, ease, and rapidity of use when compared with stand-alone devices. An example of a combined unit is shown in Review Box 15.2.

All transcutaneous pacemakers have similar basic features. Most allow operation in either a fixed rate (asynchronous) or a demand mode (VVI). Most allow rate selection in a range from 30 to 200 beats/min. Current output is usually adjustable from 0 to 200 mA. If an ECG monitor is not an integral part of the unit, an output adapter to a separate monitor is required to “blank” the large electrical spike from the pacemaker impulse and allow interpretation of the much smaller ECG complex. Without blanking protection, the standard ECG machine is swamped by the pacemaker spike and is uninterpretable. This could be disastrous because the large pacing artifacts can mask treatable ventricular fibrillation (Fig. 15.12). Pulse durations on available units vary from 20 to 40 msec and are not adjustable by the operator.

Two sets of patient electrodes are usually required for operation of the device. One set of standard ECG electrodes is used for monitoring. The much larger pacing electrodes deliver electrical impulses for pacing. Newer combined defibrillator-pacemakers can use a single set of electrodes for ECG monitoring, pacing, and defibrillation. This approach makes the device simpler to use, although the ECG waveform and analysis may be suboptimal. Provisions are generally made for separate ECG monitoring electrodes for use as desired by the operator.

Along with the widespread use of TCP come problems arising from lack of standardization of equipment. Pacing electrodes placed on a patient before arrival at the hospital may be incompatible with the transcutaneous pacemaker used in the ED, and likewise, the equipment in the ED may differ from inpatient units. Efforts should be made to establish a single standard for pacing electrode connectors within an institution and out to the prehospital environment if possible.

To facilitate rapid setup, the pacing electrodes should be connected to the pacemaker at all times. With conventional packaging the leads are inside the packet with the pads, which means that the packet must be opened to allow connection to the pacing unit. However, exposure to air causes the electrodes to dry out and lose their conductivity, thus requiring continual replacement of the unused electrodes. Newer packaging leaves the connectors outside the packet, thus allowing connection to the pacemaker while preserving the shelf life of the pacing electrodes.

**Technique**

**Pad Placement**

The pacing electrodes are self-adhesive. Position them as shown in Fig. 15.13. Take care to avoid placing the electrodes over an implanted pacemaker or defibrillator. Remove any transdermal drug delivery patches if they are in the way. Remove excessive hair if time permits. Place the anterior electrode (cathode or negative electrode) as close as possible to the point of maximal impulse on the left anterior chest wall. Place the second electrode directly posterior to the anterior electrode (see Fig. 15.13A and B). The posterior electrode serves as the ground. An alternative arrangement for the pacing electrodes is shown in Fig. 15.13C. On females, place the electrode beneath the breast and against the chest wall. Because data regarding optimum electrode placement are scarce, selection can be based on the clinician's preference and the patient's habitus.\textsuperscript{122,123} Nonetheless, suboptimal capture as a result of poor electrode placement may be rectified with a small change in electrode position. Although the polarity of the electrodes does not appear to be important for defibrillation, at least one study has indicated that it may be important for pacing.\textsuperscript{124} The electrodes are labeled by the manufacturer to indicate which one should be placed over the precordium, and it is prudent to observe this recommendation. ECG electrodes (if used) are placed on the chest wall or limbs, or both, as required and connected to the instrument cable. Some clinicians prophylactically apply pacing electrodes to all critically ill patients with bradycardia to facilitate immediate TCP should decompensation occur.

There is little risk for electrical injury to health care providers during TCP. The power delivered during each impulse is less than 1/1000 of that delivered during defibrillation.\textsuperscript{125} Chest compressions (CPR) can be administered directly over the insulated electrodes while pacing.\textsuperscript{126} Inadvertent contact with the active pacing surface results in only a mild shock.
Generally, a heart rate of 60 to 70 beats/min will maintain adequate blood pressure (by blood pressure cuff or arterial catheter) and cerebral perfusion.

Assess electrical capture by monitoring the ECG tracing on the filtered monitor of the pacing unit (see Fig. 15.14, steps 4). Rate and current (output) selections are adjustable.

**Figure 15.12** A-C, The top three rhythm strips are taken from a standard wall-mounted electrocardiographic monitor. They all demonstrate large pacer spikes without capture (arrows). The underlying rhythm cannot be determined and could be treatable ventricular fibrillation. D, The bottom rhythm strip demonstrates a tracing on the same patient with the external pacer monitor (special dampening). Note that the pacing spikes are much smaller, and it is easy to see that the underlying rhythm is asystole, without pacer capture. The presence of a T wave after the QRS complex is a good indicator of ventricular capture.

**Figure 15.13** Correct placement of transcutaneous pacemaker electrodes. A and B, Anteroposterior positions. C, Anterolateral positions (see text).

**Pacing Bradycardic Rhythms**

To initiate TCP, apply the pacing electrodes and activate the device (Fig. 15.14, steps 1 and 2). Slowly increase the output from minimal settings until capture is achieved (see Fig. 15.14, step 4). Rate and current (output) selections are adjustable. Generally, a heart rate of 60 to 70 beats/min will maintain adequate blood pressure (by blood pressure cuff or arterial catheter) and cerebral perfusion.

Assess electrical capture by monitoring the ECG tracing on the filtered monitor of the pacing unit (see Fig. 15.14, steps 4).
Emergency Transcutaneous Cardiac Pacing

Apply the pacing pads in either the anteroposterior or the anterolateral positions.
Excessive hair may be removed if time permits.

Select the pacing rate by using the Rate button.
Generally, a rate of 60 to 70 beats/min will maintain adequate blood pressure and cerebral perfusion.

Note the presence of pacer spikes (arrows) and the absence of subsequent QRS complexes.
Electrical capture has not been achieved and the current needs to be increased.

Turn the pacemaker on (long arrow), and check to make sure that the pacemaker is sensing the intrinsic rhythm (short arrows).
Slowly increase the current output from minimal settings until capture is achieved.
With higher current electrical capture has been achieved (QRS after every pacer spike, arrows).
Check for mechanical capture by palpating for a pulse.

Figure 15.14 Emergency transcutaneous cardiac pacing.

5 and 6). Assess mechanical capture by palpating the pulse as for transvenous pacing. Because of muscular contractions triggered by the pacer, carotid pulses may be difficult to assess, so palpating the femoral pulse may be easier. Additionally, bedside US may prove useful in determining ventricular capture.127,128 Ideally, continue pacing at an output level just above the threshold of initial electrical capture to minimize discomfort. One study involving 16 normal male volunteers who were paced without sedation noted cardiac capture at a mean current of 54 mA (range, 42 to 60 mA).129 Most volunteers could tolerate pacing at their capture threshold; only one participant required discontinuation of pacing at 60 mA because of intolerable pain. Heller and coworkers compared subjective pain perception and capture thresholds in 10 volunteers paced with five different transcutaneous pacers.130 Capture rates (40% to 80%), thresholds (66.5 to 104 mA), and subjective discomfort varied from pacemaker to pacemaker. Failure to capture with TCP may be related to electrode placement or patient size. Patients with barrel-shaped chests and large amounts of intrathoracic air conduct electricity poorly and may prove refractory to capture. In one study, the scarring associated with thoracotomy was found to nearly double the pacing threshold.131 A large pericardial effusion or tamponade will also increase the output required for capture.132 Failure to electrically capture with a transcutaneous device in these settings is an indication to consider immediate placement of a transvenous pacer.

Patients who are conscious or who regain consciousness during TCP will experience discomfort because of muscle contraction.121,129,130 Analgesia with incremental doses of an opioid agent, sedation with a benzodiazepine compound, or both, will make this discomfort tolerable until transvenous pacing can be instituted.

Overdrive Pacing
Overdrive pacing of ventricular tachycardia or paroxysmal supraventricular tachycardia is performed in patients who are stable enough to tolerate the brief delay associated with the preparation needed for this technique.106–111 Few data exist on the efficacy or use of this procedure in the ED. Sedate the patient as explained earlier, place pacing and monitoring electrode pads in the standard positions as detailed earlier, and initiate brief trains (6 to 10 beats) of asynchronous pacing. Set the pacer rate at approximately 20 to 60 pulses/min greater...
than the dysrhythmia rate. Generally, an impulse rate of 200 pulses/min is used for ventricular tachycardias (the rate is usually 150 to 180 beats/min), and a rate of 240 to 280 pulses/min is used for paroxysmal supraventricular tachycardias (the rate is commonly 200 to 250 beats/min).

Because rhythm acceleration is possible during overdrive pacing, it is essential to keep resuscitation equipment, including a defibrillator, at the bedside.

**Complications**

The major potential complication of TCP is failure to recognize the presence of underlying treatable ventricular fibrillation. This complication is primarily due to the size of the pacing artifact on the ECG screen, a technical problem inherent in systems without appropriate dampening circuitry.

A rare complication of TCP is induction of ventricular fibrillation. Studies of fibrillation thresholds using large precordial electrodes have shown that the longer impulse durations used in modern devices seem to decrease the chance of inducing ventricular fibrillation with TCP. Nonetheless, asynchronous TCP for tachydysrhythmias has been associated with acceleration in rhythm and the development of ventricular fibrillation.

If the patient has an implanted cardioverter defibrillator that is not delivering effective pacing, disable the defibrillator by placing a magnet over it prior to initiating TCP. Failure to do so may trigger the device to administer an unnecessary shock to the patient.

Studies looking at prolonged TCP in humans have not been extensive. Zoll and colleagues reported 25 humans paced for up to 108 hours with impulses of 20-msec duration. Pacer-induced dysrhythmias did not occur. Leatham and colleagues paced one patient for 68 hours with impulses 20 msec in duration. The patient died 2 days after pacing was discontinued. Pathologic examination revealed no evidence of pacer-induced myocardial damage. Madsen and colleagues paced 10 healthy volunteers at threshold for 30 minutes and found no enzyme or echocardiographic abnormalities. TCP appears unlikely to produce cardiac injury with short-term use in the ED.

Soft tissue discomfort with potential injury may still occur with current transcutaneous pacemakers. Most patients are able to tolerate the discomfort, especially after sedation and analgesia, which should be routine. Nonetheless, prolonged use may still induce local cutaneous injury, particularly in pediatric patients because of the use of smaller electrodes. Patients who cannot tolerate TCP or who will need long-term pacing are candidates for transvenous pacing.
REFERENCES


77. Duchenne de Boulogne: De l’electrification localise et son application a la pathologie et a la therapeutique, Paris, 1872, Bailliere.


DEFINITION

Pericardial effusion, the presence of fluid within the pericardial space, has a number of causes. As fluid accumulates, a critical point is reached at which pericardial pressure negatively affects cardiac filling and causes circulatory insufficiency. This is called pericardial tamponade. Once compensatory mechanisms begin to fail, obstructive shock ensues and a failure to restore hemodynamics eventually leads to cardiac arrest. Only removal of fluid can stabilize hemodynamics at this point.

ANATOMY AND PHYSIOLOGY

Pericardium and Pericardial Space

The pericardium is a two-layered fibroelastic sac surrounding the heart. The pericardium is avascular but well innervated, so inflammation induces pain. The visceral pericardium is a single-cell layer that adheres to the epicardium. The outer parietal pericardium consists mostly of collagen with some elastin. These two layers create the pericardial space, which normally contains 15 to 50 mL of serous fluid and is under negative pressure to promote filling of the right ventricle (RV) during diastole. Pericardial fluid provides lubrication for cardiac contractility and acts as a “shock absorber” for deceleration forces.

The pericardium is a tense structure, but it also has some elasticity. These properties limit the amount of cardiac dilation that is possible during diastole and enhance mechanical interactions between the atria and ventricles during systole. This semi-elastic property can also tolerate an acute (i.e., over a period of hours to days) accumulation of pericardial fluid (80 to 120 mL) without significantly increasing intrapericardial pressure, which is the flat portion of the pressure-volume curve for pericardial pressure (Fig. 16.1). Once a critical volume is reached, adding as little as 20 to 40 mL can double intrapericardial pressure (the steep portion of the pressure-volume curve [see Fig. 16.1]) and cause clinical decompensation from cardiac tamponade. Cardiac tamponade typically occurs with an intrapericardial pressure of 15 to 20 mm Hg.

Slow and chronic accumulation of pericardial fluid (over a period of weeks to months) causes the pericardium to expand circumferentially, and it can accommodate several liters of fluid with minimal alteration in intrapericardial pressure. Patients with this condition may be asymptomatic despite large effusions. No specific pericardial volume predicts the hemodynamic consequences of an effusion; such consequences depend on the acuity of the accumulation of the fluid.

Pericardiocentesis

**Indications**

*Diagnostic*
- Determining the cause of pericardial effusion

*Therapeutic*
- Shock/hemodynamic instability
- Cardiac arrest, pulseless electrical activity

**Contraindications**

*Absolute*
- None (if hypotension or hypoperfusion is evident or if the patient is in cardiac arrest)

*Relative*
- Coagulopathy
- Prosthetic heart valve
- Pacemakers and cardiac devices
- Lack of direct visualization (e.g., ultrasound) during the procedure
- Traumatic pericardium (thoracotomy preferred)

**Complications**

- Dysrhythmias
- Intra-abdominal injury (e.g., liver injury)
- Air embolism
- Fluid reaccumulation
- Hemothorax
- Pneumothorax
- Pneumopericardium
- Costochondritis
- Suppurative pericarditis
- Coronary artery injury (puncture or laceration)
- Internal mammary artery injury
- Intercostal vessel or nerve injury
- Atrial or ventricular puncture or laceration

**Equipment**

- Chlorhexidine
- Sterile drape
- 10-mL syringe/25-gauge needle
- Ultrasound
- J-tipped guidewire
- ECG wire with alligator clips
- 60-mL syringe
- 18-gauge spinal needle
- Dilator
- 8-Fr pigtail catheter
- 3-way stopcock

Review Box 16.1 Pericardiocentesis: indications, contraindications, complications, and equipment.
Pulmonary capillary wedge pressure are virtually identical. This diastolic pressure, pulmonary artery diastolic pressure, and a "pressure plateau" occurs in which right atrial pressure, RV to rising intrapericardial pressure (Fig. 16.2). Late in tamponade pressure than systemic arterial pressure and are also vulnerable of the RV.

Walled structures then become compressed and reduce filling to the inferior vena cava (IVC) and right atrium. These thin-walled structures then become compressed and reduce filling. In this situation, the free wall of the RV cannot expand against the pericardial fluid during inspiration. To accommodate RV filling, the interventricular septum bows abnormally into the LV, which reduces its volume; this is also known as ventricular interdependence. LV filling, stroke volume, and ultimately distal tissue perfusion is reduced. This phenomenon is responsible for pulsus paradoxus (PP; described later in this chapter), which is sometimes observed with tamponade. LV filling is also reduced by the collapse of right-sided structures. After a critical volume is reached on the pressure-volume curve (see Fig. 16.1), the intrapericardial pressure is transmitted to the inferior vena cava (IVC) and right atrium. These thin-walled structures then become compressed and reduce filling of the RV.

The atria and pulmonary circulation are at much lower pressure than systemic arterial pressure and are also vulnerable to rising intrapericardial pressure (Fig. 16.2). Late in tamponade a “pressure plateau” occurs in which right atrial pressure, RV diastolic pressure, pulmonary artery diastolic pressure, and pulmonary capillary wedge pressure are virtually identical. This equalization of chamber pressure leads to a reduction in venous return and the echocardiographic hallmark of tamponade: diastolic collapse of the RV (discussed later in this chapter). At this point, hemodynamic collapse is imminent, with severe hypotension, and potentially pulseless electrical activity (PEA) developing. Unless intrapericardial pressure is decreased immediately, cardiac arrest will ensue.

**Compensatory Mechanisms and Pericardiocentesis**

To maintain a physiologic cardiac output early in tamponade, the sympathetic nervous system increases the heart rate, arterial vasoconstriction (to maintain mean arterial blood pressure), and venoconstriction (to maintain normal venous-atrial and atrioventricular filling gradients). Early in tamponade, these compensatory mechanisms are usually effective in maintaining adequate cardiac output.

Compensatory mechanisms also preserve normal cardiac contractility and myocardial perfusion. However, when the pericardial pressure overwhelms the compensatory mechanisms, coronary perfusion pressure is reduced, which leads to myocardial ischemia, systolic dysfunction, and ultimately a reduction in cardiac output. Experimental induction of severe tamponade
demonstrated microscopic ischemic cardiac injury. Lactic acidosis (resulting from reduced cardiac output and systemic hypoperfusion) also leads to further reduction in cardiac contractility and, ultimately, cardiac output. Removal of pericardial fluid (i.e., pericardiocentesis) reverses the pathophysiologic processes just described by improving coronary filling, cardiac filling, and hence cardiac output. Interestingly, the pressure-volume relationship of the pericardial space demonstrates hysteresis; that is, withdrawing a certain quantity of fluid reduces intrapericardial pressure more than addition of the same amount of fluid increases intrapericardial pressure. This effect, however, is not universal and may vary among patients and in various disease states (Fig. 16.3).

Special Considerations in Patients With Pericardial Effusion and Tamponade

Under normal circumstances, positive pressure ventilation (e.g., mechanical ventilation) reduces venous return to the right side of the heart by increasing intrathoracic pressure. This could be detrimental for patients with tamponade because right-sided filling is already compromised and further reductions can lead to severe hemodynamic instability. Therefore positive pressure ventilation (i.e., noninvasive or invasive) should be avoided in patients with known or suspected tamponade unless it is absolutely necessary (e.g., respiratory failure).

Low-pressure pericardial tamponade is defined as a hemodynamically significant effusion with an intrapericardial pressure that is lower than expected. This category of tamponade occurs in patients with subacute or chronic effusions, but present with a superimposed hypovolemia (e.g., associated with long-term diuretic use, dehydration, or excessive dialysis). The diagnosis may be challenging because the classic symptoms and findings on physical examination (e.g., distended neck veins) may be absent. Ultrasound is most helpful in making the diagnosis in these patients. Fluid boluses may temporize the hemodynamic compromise while pericardial decompression is being arranged.

EPIDEMIOLOGY

The major categories of pericardial effusion include infection, inflammation, malignancy, trauma, and metabolic abnormalities. Effusion may also be associated with aortic disease, connective tissue disease, or idiopathic causes. It is often difficult to report the exact incidence of each type of pericardial effusion because of variations in patient populations, local epidemiology, and the diagnostic protocols used during evaluation. The prevalence of a chronic effusion is also difficult to ascertain because it is often asymptomatic and underreported. General autopsy studies demonstrate an overall prevalence of 3.4%.

CAUSES OF PERICARDIAL EFFUSION (BOX 16.1)

Acute Hemopericardium

Acute hemopericardium, or rapid accumulation of blood in the pericardial space, can have a traumatic or nontraumatic etiology. It is one of the most feared causes of tamponade because the semi-elastic pericardium cannot accommodate acute increases in pericardial fluid and clinical deterioration can be rapid. This diagnosis can be challenging to make because there may be little or no evidence during the initial evaluation (e.g., pericardial size might be normal on a chest radiograph). Common causes include trauma and aortic dissection retrograde into the pericardial sac.

Traumatic Hemopericardium

Penetrating Trauma

Penetrating cardiac trauma can cause acute hemopericardium by either external forces (e.g., a stab wound to the heart) or internal forces (e.g., iatrogenic injury during placement of a pacemaker). Cardiac perforation can lead to rapid clinical deterioration and PEA.

External cardiac puncture is associated with stab wounds or projectile injuries (e.g., gunshot wounds). Tamponade develops in 80% to 90% of patients with cardiac stab wounds as opposed to 20% of those with gunshot wounds. Stab wounds cause tamponade more frequently because if the pericardial injury is small, it can reseal and trap blood within the pericardial space. On the other hand, a gunshot typically produces both large myocardial and pericardial wounds that allow continuous drainage into the mediastinal and pleural space. Clinical deterioration is usually secondary to hypovolemia. Any penetrating injury to the chest, back, or upper part of the abdomen may injure the pericardium and cause tamponade.

Internal penetrating trauma is typically caused by invasive diagnostic or therapeutic procedures. The procedures most often associated with this injury are cardiac catheterization (angioplasty or valvuloplasty) and pacemaker insertion. Hemopericardium results from puncturing a cardiac chamber, a coronary artery, or a great vessel (e.g., the superior vena cava). Ironically, pericardiocentesis itself (treatment of a pericardial effusion) can cause hemopericardium if a coronary vessel or the myocardium is injured during the procedure.
Internal jugular and subclavian venous catheters (e.g., central venous or hemodialysis catheters) are commonly inserted in the emergency department (ED). During such procedures, hemopericardium results from perforation of the superior vena cava, right atrium, or RV. Hemopericardium can occur immediately or can be delayed for days subsequent to erosion of the catheter through myocardial or vascular tissue.\textsuperscript{27,28} Although this complication seldom occurs, it should always be considered when a patient experiences sudden hemodynamic deterioration following an invasive procedure.

**Blunt Trauma**

Major blunt chest trauma can cause hemopericardium with or without obvious signs of injury, from rupture of a cardiac chamber or, less commonly, damage to a coronary artery.\textsuperscript{29} Myocardial rupture can be uncontained or contained.\textsuperscript{16,31} Patients with uncontained rupture do not typically survive long enough to reach the hospital.\textsuperscript{32} Contained rupture may be found soon after injury or may be a late finding.\textsuperscript{17,18} Tamponade can also be caused by a deceleration mechanism of injury that induces either aortic or vena caval disruption.\textsuperscript{34} In one case series the incidence of tamponade following deceleration injury was found to be 2.3% (1 in 43 patients).\textsuperscript{35}

**Miscellaneous Trauma**

Chest compressions during cardiopulmonary resuscitation (CPR) can also cause hemopericardium from broken ribs or bleeding intercostal vessels.\textsuperscript{36} Hemopericardium following CPR has been described in case reports\textsuperscript{17,18} but is unlikely to be significant, much less to cause tamponade.

**Atraumatic Hemopericardium**

Atraumatic hemopericardium is difficult to diagnose. Diagnosis is often delayed because it occurs spontaneously and the clinical findings can be less obvious than those of hemopericardium from traumatic causes. Maintain a high index of suspicion for this condition in patients with risk factors, such as certain malignancies (e.g., lung cancer) or tuberculosis within endemic areas. Common causes of atraumatic hemopericardium are discussed later.
Bleeding diathesis is an important cause of spontaneous hemopericardium and may be associated with the use of anticoagulants (reported incidence of 2.5% to 11%) or thrombolytic therapy (incidence <1%). Patients who have undergone cardiac surgery are at increased risk because of the anticoagulative effects of the cardiopulmonary bypass machine and medications started postoperatively (e.g., clopidogrel, warfarin). Fortunately, tamponade has a low incidence and is generally detected in the postoperative period before discharge. This complication is usually prevented by the intraoperative placement of mediastinal or pericardial drains.

Hemopericardium can develop following myocardial infarction (MI). Early after a transmural MI (1 to 3 days), the necrotic myocardium causes inflammation of the overlying pericardium and then effusions can form. Late-developing effusions (weeks after an MI) are caused by an autoimmune pericarditis called Dressler’s syndrome. Improved reperfusion techniques have drastically reduced the incidence of post-MI pericarditis and effusion.

Ascending aortic dissection causes rapid and usually fatal hemopericardium. The dissection may expand in a retrograde fashion by extending to the base of the aorta and into the pericardial sac. This is a very difficult diagnosis, and best visualized and confirmed by bedside ultrasound. Risk factors for aortic dissection include hypertension, atherosclerosis, vasculitis (e.g., giant cell arteritis, syphilis), collagen vascular disease (e.g., Marfan’s syndrome), and the use of sympathomimetics (e.g., cocaine).

Ventricular free-wall rupture is a rapidly fatal cause of acute hemopericardium that can occur after MI leading to cardiac failure and shock. This complication is less common today than in the past (<1%) secondary to improved revascularization techniques, better therapeutic medications, and faster intervention times (shorter door-to-balloon times) for coronary ischemia. Despite a reduction in its overall incidence, 7% of all deaths related to MI are caused by this complication. Survival is theoretically possible with prompt recognition and treatment, but the prognosis is grim once tamponade occurs.

### Nonhemorrhagic Effusions

Nonhemorrhagic pericardial effusions usually accumulate slower than acute hemopericardium does (over a period of weeks to months). Chronic fluid accumulation allows the pericardium to stretch circumferentially and accommodate up to 2000 mL of fluid without any hemodynamic compromise. Effusions that grow slowly allow the circulatory system to adapt to increased hemodynamic stability. Thus, asymptomatic patients with moderate to large effusions may not need emergency pericardiocentesis, in contrast to patients with acute hemopericardium.

Nonhemorrhagic effusions have several causes (see Box 16.1), and the exact one may not be obvious during the initial evaluation without diagnostic pericardiocentesis. Common causes of nonhemorrhagic effusions are discussed in the following sections.

### Idiopathic Effusions

Most idiopathic effusions are believed to be viral in origin and most commonly caused by infection with coxsackievirus, echovirus, cytomegalovirus, or human immunodeficiency virus (HIV) (discussed later in the chapter). Idiopathic pericardial effusions may be asymptomatic or have an associated component of pericarditis (e.g., positional pain or diffuse ST-segment changes on the electrocardiogram [ECG]). These effusions are often labeled idiopathic because the diagnosis cannot be made noninvasively (i.e., based on the history, physical examination, or serum testing) and the risk associated with diagnostic pericardiocentesis often outweighs the risk of observation in asymptomatic adults who appear to be well. Diagnostic pericardiocentesis may be recommended for idiopathic effusions that are persistent or symptomatic without a known cause.

### Neoplastic Effusion

Tumors of the pericardium or myocardium may cause nonhemorrhagic effusions. Primary cardiac tumors are less common (0.001% to 0.003%) than metastases from another site (2% to 18%), but either may cause a malignant effusion. Although no malignancy preferentially metastasizes to the heart, certain tumors commonly involve the heart when they metastasize; frequently implicated are lung cancer, breast cancer, mediastinal tumors, malignant melanoma, leukemia, and lymphoma.

Cardiac metastasis is usually a late finding in cancer, as other foci are generally evident first. The classic signs and symptoms of tamponade (e.g., chest pain and dyspnea) may not be obvious with malignant tamponade. When present, they may be mistakenly attributed to the underlying malignancy. Thus, in the relevant clinical scenario, consider screening patients with malignancy for pericardial effusion (e.g., ultrasound) before the clinical findings of tamponade appear.

### Radiation

Pericardial effusions (secondary to radiation-induced pericarditis) can develop acutely during radiation therapy or may be delayed for years. Risk factors include the radiation dose, duration of exposure, and age of the patient. Patients treated with radiation for Hodgkin’s disease have the highest association of radiation-induced pericarditis and subsequent effusions. These effusions can be serous, hemorrhagic, or fibrinous.

### Congestive Heart Failure

Congestive heart failure (CHF) is a cause of pericardial effusion. Diagnosis may be difficult because of overlapping signs and symptoms with exacerbations of CHF (e.g., chest pain or dyspnea). Adding to the diagnostic complexity is that 12% to 20% of patients with CHF have a coexisting pericardial effusion. Fortunately, treatment of CHF-associated pericardial effusion does not differ from that for an effusion from other causes: treat the underlying cause unless the patient has evidence of hemodynamic compromise.

### HIV-Associated Effusions

HIV can cause nonhemorrhagic pericardial effusion and tamponade. The incidence has been reported to be approximately 11% in patients with HIV infection or acquired immunodeficiency syndrome, and 13% of cases are classified as moderate to severe. It is unclear whether antiretroviral therapy has affected these data. HIV-related effusions have been attributed to bacterial (e.g., Staphylococcus aureus), viral (e.g., cytomegalovirus), fungal (Cryptococcus neoformans), and mycobacterial causes (e.g., tuberculosis, which is the most common cause of HIV-related effusions worldwide). Kaposis’s sarcoma and lymphoma can cause noninfectious pericardial effusions in HIV patients.
Renal Failure and Uremia

Pericardial effusion develops in approximately 15% to 20% of dialysis patients, and tamponade may eventually occur in as many as 35% of those people.66,67 Up to 7% of chronic dialysis patients have effusions with volumes of 1000 mL or greater.68 In many cases, effusions secondary to renal failure can be managed solely with aggressive dialysis without pericardiocentesis. Any sign of hemodynamic compromise, however, warrants strong consideration of pericardiocentesis.

Hypothyroidism

Hypothyroid patients are at risk for pericardial effusions (up to 30%), but the fluid accumulates gradually, so tamponade develops in only a few patients.69 If a pericardial effusion is present, other areas of the body usually demonstrate serositis (e.g., pleural effusions). Medically managing the underlying hypothyroidism often reverses the effusion without the need for pericardiocentesis.

Special Considerations in Pericardial Disease

Pericardial tamponade is classically described as being secondary to circumferential effusion, which causes a generalized increase in pericardial pressure and compression of multiple cardiac chambers. Loculated effusions (caused by a local hematoma or an infectious process) or pericardial adhesions (from previous inflammation) can lead to tamponade by compressing one or more cardiac chambers and thus reducing both cardiac filling and cardiac output.69,70

Constrictive pericarditis occurs following chronic pericardial inflammation, infection, or mediastinal irradiation. These processes cause scarring, fibrosis, or calcification, and the pericardium eventually becomes a nonelastic and “constrictive” sac around the heart. Myocardial relaxation and cardiac filling are impaired, and diastolic dysfunction ensues. Without echocardiography, constrictive pericarditis can be difficult to distinguish from pericardial tamponade.71

Effusive-constrictive pericarditis is defined by the presence of both pericardial effusion and pericardial constriction. It may be quite difficult to differentiate between effusive-constrictive pericarditis and pericardial tamponade in stable patients because both are associated with effusions.72 Fortunately, distinguishing between these diagnoses is less important in hemodynamically unstable patients because they are treated identically (i.e., with pericardiocentesis).72

Pneumopericardium is an interesting, though rare cause of cardiac tamponade. It is most commonly associated with pneumothorax caused by barotrauma (e.g., mechanical ventilation).73 It also occurs spontaneously during acute asthma exacerbations,74 and it can follow blunt chest injury.75,76 Although typically benign, tension pneumopericardium has been reported as a cause of life-threatening tamponade after blunt77,78 and penetrating chest trauma.79,80

DIAGNOSING CARDIAC TAMPOONADE

Diagnosis of pericardial effusions requires careful integration of the patient’s history, findings on physical examination, and diagnostic testing. Unfortunately, even experienced clinicians may not initially consider pericardial effusion because the clinical findings are often vague and nonspecific. Nonspecific symptoms, such as chest pain and dyspnea, can be ascribed to more common conditions (e.g., CHF or pulmonary pathology), so the diagnosis might be delayed until diagnostic testing is performed (e.g., computed tomography [CT] of the chest for pulmonary embolism),81 or until hypotension develops and bedside ultrasound is performed.82

Acute pericardial tamponade (e.g., secondary to blunt chest wall trauma) is usually challenging to diagnose because the findings on physical examination may resemble those of other life-threatening conditions (e.g., tension pneumothorax, hemothorax, hypovolemia, pulmonary edema, severe contusion of the RV, aortic dissection, or pulmonary embolism).67 In hemodynamically unstable patients, diagnostic (e.g., bedside ultrasound) and therapeutic (e.g., pericardiocentesis) interventions must be performed even with a paucity of findings on physical examination because rapid clinical deterioration and cardiac arrest can occur before a definitive diagnosis can be made.

Once a pericardial effusion is suspected (or diagnosed), the next step is to determine its size and hemodynamic significance and presence of underlying or associated diseases.83 Specific therapies will hinge on this information and is discussed in the following sections.

History: Patient Profile and Symptoms

The historical features of pericardial effusions are nonspecific and the diagnosis may easily be overlooked. An astute clinician might be suspicious based on comorbid conditions (e.g., warfarin therapy or a history of myxedema) and the time course of the symptoms (e.g., free-wall rupture several days after MI, dyspnea in a patient with uremic pericarditis). Patients are likely to present with symptoms relating to the underlying disease rather than the pericardial effusion itself. Box 16.2 lists important details to be ascertained from the history when pericardial effusion is suspected.

Physical Examination

Physical examination of patients with pericardial effusion (e.g., displaced point of maximal impulse, muffled heart sounds) lacks sensitivity and specificity. If the history suggests pericardial effusion, the physical examination should focus on determining the underlying cause (e.g., stigmata of hypothyroidism) to guide definitive diagnostic testing (e.g., echocardiography). Ironically, many pericardial effusions are not diagnosed from the history or findings on physical examination but are found incidentally during the evaluation for other diseases.

In 1935, Beck characterized the physical manifestations of tamponade with two triads, one for chronic and one for acute tamponade.85 Beck’s chronic triad consists of increased central venous pressure (CVP) (i.e., distended neck veins), ascites, and a small, quiet heart. Beck’s triad is a classic description of acute cardiac compression, which includes increased CVP, decreased arterial pressure, and muffled heart sounds. Almost 90% of patients have one or more of these “acute” signs,86 but only approximately 33% demonstrate the complete triad.87 The utility of this triad is further limited because all three signs are usually observed shortly before cardiac arrest.

It would be clinically desirable to identify patients in early tamponade, before hemodynamic collapse. Unfortunately, findings on physical examination in early tamponade are nonspecific and may be indistinguishable from those of other critical diseases (e.g., septic shock, right heart failure).87 Patients
initially seen in late tamponade also have nonspecific findings. They may be agitated, panic-stricken, confused, uncooperative, restless, cyanotic, diaphoretic, acutely dyspneic, or hemodynamically unstable. Such patients should undergo a brief and focused physical examination followed by a rapid hemodynamic assessment with bedside ultrasound because the time between initial evaluation and full arrest may be brief.

Some of the findings on physical examination associated with tamponade are described later. A more comprehensive list is presented in Box 16.3.

**Vital Sign Abnormalities**

There are three sequential stages that are typically described to reflect the natural history of acute tamponade (Table 16.1). The time course within each stage varies from patient to patient. Some patients are stable within a given stage for hours, whereas others proceed through all three stages and develop cardiac arrest within minutes. Grade I tamponade is characterized by normal blood pressure and cardiac output with an increase in the heart rate and CVP (measured invasively with a central venous catheter). Grade II tamponade is defined by normal or slightly reduced blood pressure; CVP and the heart rate remain increased. Grade III tamponade is identified on the basis of Beck's triad: hypotension, tachycardia, and elevated CVP.

Nearly all patients with tamponade present with sinus tachycardia, although its specificity is low. The physiologic purpose of tachycardia is to maintain normal cardiac output.

**ONSET AND DURATION OF SYMPTOMS**

**Acute**
- Trauma
- Recent cardiac surgery
- Recent myocardial infarction
- Acute aortic dissection
- Recent diagnostic or therapeutic intervention (e.g., catheterization)
- Recent upper thoracic vascular procedure (e.g., hemodialysis catheter, central line, peripherally inserted central catheter line, Mediport)
- Recent placement of a pacemaker or automatic implantable cardioverter-defibrillator

**Subacute/Chronic**
- Metabolic (e.g., uremia)
- Endocrine (e.g., hypothyroidism)
- Infectious
  - Viral (e.g., human immunodeficiency virus)
  - Bacterial (e.g., *Staphylococcus*)
  - Fungal (e.g., *Aspergillus*)
- Neoplastic
  - Primary cardiac
  - Metastatic
- Autoimmune disorders (e.g., lupus)
- Inflammatory
  - Vasculitis

**MEDICAL/SURGICAL HISTORY**

- Autoimmune disorders
  - Lupus
  - Mixed connective tissue disease
  - Vasculitis
- Endocrine disease
  - Hypothyroidism
  - Ovarian hyperstimulation syndrome
- Metabolic diseases
  - End-stage renal disease and uremia
  - Thyroid disorders
  - Coagulopathies
- Artificial cardiac valves
  - Anticoagulation medications
  - Risk for myocarditis
- Cardiac disease
  - Anticoagulation medications
  - Pericarditis
  - Aneurysm
- Recent placement of a vascular catheter
- Recent cardiac surgery
- Recent cardiac intervention
- Recent thoracic radiation

**MEDICATIONS**

- Anticoagulants
  - Aspirin
  - Warfarin
  - Clopidogrel
- Antiarrhythmic
  - Procainamide (drug-induced lupus)
- Tuberculosis therapy
  - Isoniazid (drug-induced lupus)

**COMMON SYMPTOMS**

- Altered mental status, confusion
- Fatigue
- Dizziness, light-headedness
- Orthostatic changes
- Exercise intolerance
- Hoarseness
- Hiccups
- Fever
- Chills
- Chest pain
  - Substernal
  - Pleuritic
  - Positional
  - Scapula (phrenic nerve irritation)
  - Palpitations
- Cough
- Dyspnea
- Myalgia
- Arthralgia

**BOX 16.2** Information to Obtain from Patients When Pericardial Effusion Is Suspected

**ONSET AND DURATION OF SYMPTOMS**

**Acute**
- Trauma
- Recent cardiac surgery
- Recent myocardial infarction
- Acute aortic dissection
- Recent diagnostic or therapeutic intervention (e.g., catheterization)
- Recent upper thoracic vascular procedure (e.g., hemodialysis catheter, central line, peripherally inserted central catheter line, Mediport)
- Recent placement of a pacemaker or automatic implantable cardioverter-defibrillator

**Subacute/Chronic**
- Metabolic (e.g., uremia)
- Endocrine (e.g., hypothyroidism)
- Infectious
  - Viral (e.g., human immunodeficiency virus)
  - Bacterial (e.g., *Staphylococcus*)
  - Fungal (e.g., *Aspergillus*)
- Neoplastic
  - Primary cardiac
  - Metastatic
- Autoimmune disorders (e.g., lupus)
- Inflammatory
  - Vasculitis

**MEDICAL/SURGICAL HISTORY**

- Autoimmune disorders
  - Lupus
  - Mixed connective tissue disease
  - Vasculitis
- Endocrine disease
  - Hypothyroidism
  - Ovarian hyperstimulation syndrome
- Metabolic diseases
  - End-stage renal disease and uremia
  - Thyroid disorders
  - Coagulopathies
- Artificial cardiac valves
  - Anticoagulation medications
  - Risk for myocarditis

**COMMON SYMPTOMS**

- Altered mental status, confusion
- Fatigue
- Dizziness, light-headedness
- Orthostatic changes
- Exercise intolerance
- Hoarseness
- Hiccups
- Fever
- Chills
- Chest pain
  - Substernal
  - Pleuritic
  - Positional
  - Scapula (phrenic nerve irritation)
  - Palpitations
- Cough
- Dyspnea
- Myalgia
- Arthralgia
316  SECTION III  CARDIAC PROCEDURES

BOX 16.3  Physical Examination Findings Suggestive of Pericardial Effusion or Tamponade

VITAL SIGNS
Normal vital signs
Tachycardia (tamponade)
Hypotension (tamponade)
Narrow pulse pressure
Pulsus paradoxus
Fever (if the cause is infectious or neoplastic)

APPEARANCE
Normal appearance
Anxiety
Sense of impending doom
Diaphoretic
Cold and clammy
Pallor
Altered mental status, confusion

NECK
Supraclavicular retractions
Distended neck veins (may be flat with hypotension)

CARDIOVASCULAR
Normal findings on examination
Tachycardia
Increased pain with a supine position, relieved by leaning forward
Muted, distant heart sounds
Pericardial friction rub (inflammatory pericarditis)
Displaced point of maximal impulse
Increased cardiac borders with percussion
Distended neck veins
Auscultatory dullness along the left scapular area

PULMONARY
Telegraphic speech
Respiratory distress
Supraventricular retractions, abdominal breathing
Cough
Hoarseness
Clear breath sounds (may distinguish tamponade from congestive heart failure)

ABDOMINAL
Hepatomegaly
Splenomegaly

SKIN
Cool extremities
Clammy extremities
Dilated head and scalp veins
Peripheral edema
Anasarca


TABLE 16.1  Shoemaker System of Grading Cardiac Tamponade

<table>
<thead>
<tr>
<th>GRADE</th>
<th>PERICARDIAL VOLUME (mL)</th>
<th>CARDIAC INDEX</th>
<th>STROKE INDEX</th>
<th>MEAN ARTERIAL PRESSURE</th>
<th>CENTRAL VENOUS PRESSURE</th>
<th>HEART RATE</th>
<th>BECK’S TRIAD</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>&lt;200</td>
<td>Normal or ↑</td>
<td>Normal or ↓</td>
<td>Normal</td>
<td>↑</td>
<td>↑</td>
<td>Venous distention, hypotension, muffled heart sounds usually not present</td>
</tr>
<tr>
<td>II</td>
<td>≥200</td>
<td>↓</td>
<td>↓</td>
<td>Normal or ↓</td>
<td>↑ (≤12 cm H₂O)</td>
<td>↑</td>
<td>May or may not be present</td>
</tr>
<tr>
<td>III</td>
<td>&gt;200</td>
<td>↓↓</td>
<td>↓↓</td>
<td>↓↓</td>
<td>↑↑ (30–40 cm H₂O)</td>
<td>↑</td>
<td>Usually present</td>
</tr>
</tbody>
</table>

Despite reductions in stroke volume from worsening tamponade. Exceptions to the pairing of tachycardia with tamponade usually relate to the underlying cause of the effusion (e.g., myxedema) or the concomitant use of certain medications (e.g., β-blockers).

Adding to the diagnostic complexity, not all patients in tamponade have a reduction in blood pressure. In fact, Brown and co-workers described several tamponade patients with elevated blood pressure. These patients were previously hypertensive and paradoxically had reduced systolic blood pressure following pericardiocentesis.

Pulsus Paradoxus
PP is an exaggerated decrease in systolic blood pressure (>12 mm Hg) during inspiration secondary to reduced stroke volume (Fig. 16.4A) . Patients with moderate to severe tamponade typically demonstrate PP greater than 20 mm Hg. Unfortunately, PP is not pathognomonic for tamponade. It is observed in other conditions, such as hypotension associated with labored breathing (secondary to extreme reductions in intrathoracic pressure) (see Fig. 16.4B), severe emphysema, severe asthma, obesity, cardiac failure, constrictive pericarditis, pulmonary embolism, and cardiogenic shock.

The absence of PP does not rule out tamponade because it can occur with several conditions: atrial septal defects, aortic insufficiency, positive pressure ventilation, loculated pericardial effusions, and elevated left ventricular diastolic pressure (e.g., poor left ventricular compliance secondary to chronic hypertension). Finally, PP should be interpreted with caution in patients
Neck Vein Distention and Elevated CVP

Neck vein distention (a surrogate for measuring CVP) occurs late in tamponade, when right-sided chambers (e.g., the RV) collapse. Neck vein distention may be obvious on examination (Fig. 16.5A), but visualization of such distention is less accurate than measuring CVP by central venous catheter or evaluation of the IVC with ultrasound (Table 16.2). Patients with significant tamponade typically have a CVP of 12 cm H₂O or higher. Finally, although initial CVP readings are useful and diagnostic when grossly elevated (e.g., 20 to 30 cm H₂O), upward trends in CVP can be a more sensitive diagnostic tool. Overreliance on increased CVP and venous distention should be avoided because they do not always indicate tamponade. For example, increased intrathoracic pressure (as induced by positive pressure ventilation or Valsalva maneuvers) increases CVP and causes neck vein distention even without pericardial effusion. Conversely, hypovolemic patients may have reduced CVP and no neck vein distention despite having clinical tamponade. The absence of distended neck veins may also result from severe venoconstriction secondary to intrinsic sympathetic discharge, vasopressor use, or severe hypovolemia.

Diagnostic Testing

Diagnostic testing should be initiated when pericardial effusion or tamponade is suspected. Definitive diagnosis requires imaging, which may be done by CT but preferably is done by cardiac ultrasound (see Fig. 16.5B). Bedside ultrasound is the fastest and most reliable diagnostic tool because it is noninvasive, does not emit radiation, and can be performed at the bedside without transporting unstable patients outside the ED.

As discussed previously, pericardial effusions are occasionally discovered incidentally during evaluation for other disorders. For example, a chest x-ray and ECG performed on a patient being assessed for acute coronary syndrome (ACS) may suggest tamponade. If the clinical context supports pericardial effusion as the primary diagnosis rather than ACS, further workup for the effusion (e.g., bedside echocardiogram) should be ordered. If neither diagnosis is more likely than the other, dual workups may be necessary.

Chest Radiography

Chest radiographs are not diagnostically useful in patients with acute traumatic tamponade because the pericardium does not have sufficient time to change size or shape (see section on Pathophysiology of Pericardial Tamponade). Radiographs, however, may reveal other associated findings such as hemotorax, bullets in the thorax, or even pneumopericardium. Chest radiography may also be helpful when other diagnoses (e.g., CHF) have clinical findings similar to those of pericardial effusion. For example, a dyspneic patient with a clear chest film is less likely to have decompensated CHF than tamponade.

In patients with chronic pericardial effusions, chest films often demonstrate an enlarged, saclike, “water-bottle” cardiac shadow or a pleural effusion (see Fig. 16.5C). Unfortunately, it is difficult to differentiate a large effusion from myocardial enlargement (e.g., dilated cardiomyopathy) because radiographs demonstrate only the cardiac silhouette and do not reveal the physiologic differences between these two diagnoses.

Electrocardiography

Pericardial effusion secondary to acute pericarditis is suspected on the basis of typical changes on the ECG. Pericarditis has
QRS complexes can be identified by a sum of 10 mm or less for all precordial lead QRS amplitudes. Electrical alternans is a beat-to-beat alternation in QRS axis caused by the pendulum motion of the heart within the fluid-filled pericardial sac. Electrical alternans has been observed in 22% of medical patients with tamponade and in 5% of patients with tamponade secondary to cancer. Electrical alternans of P waves and QRS complexes (i.e., total electrical alternans) is a rare finding but, when seen, is pathognomonic of tamponade (see Fig. 16.6).

Echocardiography is the best tool for diagnosing pericardial effusion or tamponade. It not only demonstrates the presence of a pericardial effusion but is usually abnormal when significant chronic effusions are present. Overall, the echocardiogram has low sensitivity for pericardial effusion and tamponade. Notice the beat-to-beat alternation in the voltage in the P-QRS-T axis; this is caused by the periodic swinging motion of the heart in a large pericardial effusion. Relatively low QRS voltage and sinus tachycardia are also present. The four most commonly described electrocardiographic findings in pericardial effusion are sinus tachycardia, PR depression, low-voltage QRS complexes, and electrical alternans. This alternans is defined as a sum of 10 mm or less for all precordial lead QRS amplitudes. Electrical alternans is a beat-to-beat alternation in QRS axis caused by the pendulum motion of the heart within the fluid-filled pericardial sac. Alternans has been observed in 22% of medical patients with tamponade and in 5% of patients with tamponade secondary to cancer. Electrical alternans of P waves and QRS complexes (i.e., total electrical alternans) is a rare finding but, when seen, is pathognomonic of tamponade (see Fig. 16.6).

**Table 16.2** Noninvasive Estimation of Right Atrial Pressure With Ultrasound

<table>
<thead>
<tr>
<th>DIAMETER (cm) OF INFERIOR VENA CAVA</th>
<th>CHANGE IN DIAMETER WITH RESPIRATION</th>
<th>ESTIMATED RIGHT ATRIAL PRESSURE (mm Hg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal (&lt;2.1)</td>
<td>Decrease &gt;50%</td>
<td>~3 (normal, 0–5)</td>
</tr>
<tr>
<td>Dilated (2.1)</td>
<td>Decrease &lt;50%</td>
<td>~8 (normal, 5–10)</td>
</tr>
<tr>
<td>Dilated (&gt;2.1)</td>
<td>Decrease &gt;50%</td>
<td>~8 (normal, 5–10)</td>
</tr>
<tr>
<td>Dilated (&gt;2.1)</td>
<td>Decrease &lt;50%</td>
<td>&gt;15 (10–20)</td>
</tr>
</tbody>
</table>

Pericardiocentesis

in thickness between the visceral and parietal pericardium. Moderate effusions typically contain 100 to 500 mL with 10 to 20 mm of echo-free space. Large effusions contain more than 500 mL and have an echo-free space greater than 20 mm. Large effusions tend to have an echo-free space seen circumferentially around the heart. A loculated pericardial effusion can present in an atypical location and may be more difficult to appreciate compared to the non-loculated types described previously.

Once a pericardial effusion is discovered, the next step is to evaluate the patient for evidence of hemodynamic compromise. Echocardiographic signs include (1) diastolic collapse of the right atrium (highly specific and sensitive for tamponade, especially when the collapse occurs for more than a third of the cardiac cycle); (2) early diastolic collapse of the right ventricular free wall (less sensitive than right atrial collapse but specific for tamponade); (3) left atrial collapse (a very specific sign of tamponade); (4) a small, slit-like, hyperkinetic left ventricle; (5) dilation of the hepatic veins and IVC; (6) respiratory variations in the velocity of blood flow through the tricuspid and mitral valves; and (7) visualizing the heart swinging to and fro within the pericardial sac. The presence of any of these signs should alert the clinician to the possibility of hemodynamic instability. Obtain expert consultation if there is any uncertainty about the ultrasound findings.

Limitations of Ultrasound

Ultrasound is the best diagnostic tool for pericardial effusion and tamponade, but great care must be taken when it is used as the only diagnostic modality. Echocardiographic signs include (1) diastolic collapse of the right atrium (highly specific and sensitive for tamponade, especially when the collapse occurs for more than a third of the cardiac cycle); (2) early diastolic collapse of the right ventricular free wall (less sensitive than right atrial collapse but specific for tamponade); (3) left atrial collapse (a very specific sign of tamponade); (4) a small, slit-like, hyperkinetic left ventricle; (5) dilation of the hepatic veins and IVC; (6) respiratory variations in the velocity of blood flow through the tricuspid and mitral valves; and (7) visualizing the heart swinging to and fro within the pericardial sac. The presence of any of these signs should alert the clinician to the possibility of hemodynamic instability. Obtain expert consultation if there is any uncertainty about the ultrasound findings.

Diagnosing Pericardial Effusions and Tamponade

Fluid found within the pericardial space is not always pathologic as it normally holds 15 to 50 mL of fluid. Small effusions may be clinically insignificant. The sonographer/clinician should exercise caution to not over-read an effusion, particularly when the patient is hemodynamically stable. Small effusions typically contain 50 to 100 mL with an echo-free space less than 10 mm in thickness between the visceral and parietal pericardium. Moderate effusions typically contain 100 to 500 mL with 10 to 20 mm of echo-free space. Large effusions contain more than 500 mL and have an echo-free space greater than 20 mm. Large effusions tend to have an echo-free space seen circumferentially around the heart. A loculated pericardial effusion can present in an atypical location and may be more difficult to appreciate compared to the non-loculated types described previously.

Once a pericardial effusion is discovered, the next step is to evaluate the patient for evidence of hemodynamic compromise. Echocardiographic signs include (1) diastolic collapse of the right atrium (highly specific and sensitive for tamponade, especially when the collapse occurs for more than a third of the cardiac cycle); (2) early diastolic collapse of the right ventricular free wall (less sensitive than right atrial collapse but specific for tamponade); (3) left atrial collapse (a very specific sign of tamponade); (4) a small, slit-like, hyperkinetic left ventricle; (5) dilation of the hepatic veins and IVC; (6) respiratory variations in the velocity of blood flow through the tricuspid and mitral valves; and (7) visualizing the heart swinging to and fro within the pericardial sac. The presence of any of these signs should alert the clinician to the possibility of hemodynamic instability. Obtain expert consultation if there is any uncertainty about the ultrasound findings.
Pericardiocentesis has traditionally been performed blindly. This approach was associated with a low success rate and a high rate of complications, such as inadvertent puncture of the lung, ventricle, or epicardial vessels.\(^1\) Using ultrasound to both diagnose and guide pericardiocentesis has resulted in increased success rates, as well as a lower rate of complications.\(^2,3\)

Bedside ultrasound may additionally allow the emergency physician to make a rapid diagnosis of pericardial effusion. Evidence of impending cardiac tamponade, including right ventricular collapse and distention of the inferior vena cava, can also be identified.

**Equipment**

The pericardium should be imaged with a low-frequency (2 to 4 mHz) transducer to achieve adequate depth. A phased-array or microconvex transducer is preferred for its smaller footprint, which enables the sonographer to image between the ribs. However, a curvilinear transducer may be used if that is what is available.

**Image Interpretation**

The initial step in the procedure is to evaluate the pericardium in multiple windows. This will allow identification and characterization of the effusion, as well as planning of the best approach for drainage. This chapter focuses primarily on the views most commonly used in this procedure, the subxiphoid and parasternal.

The subxiphoid view is best known to most emergency physicians as part of the focused abdominal sonography in trauma (FAST) examination. This view provides a four-chamber view of the heart and uses the left lobe of the liver as an acoustic window. To obtain this view, the transducer is placed just inferior to the xiphoid process in the midline. The indicator faces the patient’s right side (Fig. 16.US1). To obtain the best image possible it is best to place the hand over the transducer and press down into the epigastric area. The transducer can then be aimed toward the left side of the chest until the heart comes into view. The depth may need to be adjusted to view all four chambers of the heart, as well as the pericardium.

In this view the left lobe of the liver can be seen at the top of the image. Deep to the liver, a four-chambered view of the heart should be seen, surrounded by the brightly echogenic (white) border of the pericardium (Fig. 16.US2). The right ventricle will abut the liver, with the left ventricle located deeper into the body. A pericardial effusion can be identified as an anechoic (black) or hypoechoic (dark gray) collection between the heart and pericardium (Fig. 16.US3). Although fluid will typically collect at the most gravity-dependent area, loculated collections may not follow this rule.

The parasternal view is obtained by placing the transducer to the left of the patient’s sternum in the fourth to fifth intercostal space. The indicator should be pointing toward the patient’s right shoulder (Fig. 16.US4). Slight adjustments in angle may be needed to obtain the best image of the heart. If the patient’s hemodynamic status allows, placing the patient in a left lateral decubitus position may improve this view by moving the heart closer to the anterior chest wall and displacing the air-filled lungs.

The parasternal view will demonstrate the left atrium, left ventricle, and a small portion of the right ventricle (Fig. 16.US5). The pericardium can be seen as an echogenic (white) border surrounding the heart. As in the subxiphoid view, an effusion will appear as an anechoic (black) or hypoechoic (dark gray) collection between the heart and pericardium (Fig. 16.US6).

**Procedure and Technique**

Once the views have been evaluated and an effusion has been identified, evaluate the pericardium in multiple views to determine the

---

**Figure 16.US1** Placement of the ultrasound transducer to obtain a subxiphoid image of the heart.

**Figure 16.US2** Normal subxiphoid view of the heart seen with ultrasound. The pericardium (arrow) can be identified as a bright white (hyperechoic) outline, typically best seen at the inferior aspect of the heart. The pericardium should directly abut the heart, as seen in this image.

**Figure 16.US3** Subxiphoid view demonstrating a pericardial effusion. When comparing this image with the normal view, a black (anechoic) fluid collection (arrow) can be seen between the pericardium and the left ventricle.
best area to attempt pericardiocentesis. Tsang and colleagues described the procedure for echocardiographically guided pericardiocentesis in detail in 1998. Ideally, the procedure should be attempted at the site at which the largest fluid collection is closest to the skin surface. Typically, the anterior chest wall is preferred because of its proximity to the pericardium and the absence of vital interfering structures such as the liver. The air-filled lung creates a scatter artifact that does not allow the ultrasound beam to pass. Therefore if the heart and pericardium can be viewed clearly, avoidance of the lung can be ensured. The sonographer should also attempt to approximate and avoid the location of the internal mammary artery, which lies 3 to 5 cm lateral to the sternum. Once the optimum site is identified, the practitioner should note the trajectory of the ultrasound beam. This is the trajectory that should be followed by the needle.

Sterilize the field. Anesthetize the area to be traversed with local anesthetic. Place the ultrasound transducer in a sterile covering. Take care to not reposition the patient after initial assessment because this will alter the position and trajectory. Use a 16-gauge catheter with a retractable needle to minimize potential injury to the underlying structures. After the field is prepared, attach the catheter to an attached syringe. Advance the needle in the predetermined location along the predetermined trajectory. This can be done blindly or can be guided directly by the transducer. Apply gentle continuous pressure to the syringe until the pericardial space is entered and fluid is obtained.

If there is any question of whether the needle tip is in the pericardial space, inject agitated saline through the catheter under direct ultrasound guidance. This will allow analysis of the location of the needle tip. Prepare saline echocardiographic contrast medium by using two 5-mL syringes, one with saline and the other air, connected via a three-way stopcock to the needle catheter sheath. Saline in one syringe is rapidly injected between the syringes and then injected into the sheath once it is agitated. Entrance of the agitated saline into the pericardial space will appear sonographically as a brightly echogenic area.

Complications

Complications may occur when a pericardial effusion is misdiagnosed. The most common factor causing misdiagnosis is the presence of a fat pad anterior to the heart. Unless the effusion is loculated, it should lie within the most dependent portion of the heart and should be circumferential, depending on its size. Considering these factors and evaluating the pericardium in multiple views will aid in decreasing this misdiagnosis. Additionally, take care when performing pericardiocentesis in stable patients, particularly when the effusion is small. Smaller effusions may be more difficult to access and therefore lead to increased complications.

References

pericardial effusions, although several details help distinguish between the two entities. First, epicardial fat pads tend to occur anteriorly, unlike circumferential effusions, which occur posteriorly. If a fat pad is suspected, multiple ultrasound views should be obtained to rule out a posterior effusion. Second, the echocardiographic appearance of an epicardial fat pad is isoecholic and homogeneous. This differs from blood in the pericardial space, which may look like fronds of clot waving within an anechoic (black) pericardial space. Third, an epicardial fat pad does not alter hemodynamics like tamponade does; it should not cause diastolic collapse of the right ventricle free wall, dilatation of the IVC, or any signs indicating hemodynamic compromise. After careful examination, if doubt still exists regarding the presence of an effusion, hemodynamically stable patients should have a formal echocardiogram; consider CT scan if an echocardiogram is not available.

**CT Scan**
CT may be the only diagnostic option at institutions where ultrasound or formal echocardiography is not available in the ED. It is also helpful when concomitant diseases of the lung or mediastinum are considered. CT can demonstrate dilated hepatic veins, a plethoric IVC, and interventricular septal bowing.\(^{117}\) This modality is less desirable than bedside ultrasound because it requires patients to be transported to the radiology suite. Thus, patient safety and cardiopulmonary stability must be considered before the decision is made to transport the patient. Unstable patients should never be transported until they are fully stabilized.

For stable patients, CT is effective in defining the presence, severity, and extent of pericardial effusions (i.e., circumferential versus loculated). In certain circumstances, it even provides a more definitive diagnosis than echocardiography does because it may reveal the type of pericardial fluid (by differences in tissue density) and pericardial disease (e.g., constrictive pericarditis).\(^{118}\) In one series, eight equivocal echocardiograms were evaluated by follow-up CT.\(^{119}\) Two patients thought to have pericardial effusion by ultrasonography were found to have pleural effusions. Another patient in whom a pericardial effusion was diagnosed on ultrasonography was found by CT to have an epicardial lipoma. CT defined three loculated pericardial effusions not identified by ultrasonography. Finally, two patients had hemopericardium visualized by CT but not by ultrasonography.

**Treating Pericardial Effusions and Tamponade**
Treatment of pericardial effusions in the ED depends on the degree of hemodynamic compromise. Patients with stable effusions should be treated supportively while the underlying cause (known or suspected) is addressed. For example, stable patients with pericardial effusions secondary to uremia may best be treated with dialysis, observation, and serial echocardiograms.\(^{120}\) Even when the diagnosis is unknown there may be no need to perform emergency pericardiocentesis if the effusion is small to moderate.\(^{121}\) Deferring diagnostic pericardiocentesis to the inpatient setting may be preferred because the cardiac catheterization laboratory or operating room is a more sterile and controlled environment than the ED.\(^{17}\)

Patients with evidence of tamponade need urgent pericardiocentesis (discussed in the next section). Even those with early tamponade who are stable can decompensate quickly with little warning. Fluid boluses may improve hemodynamics temporarily, especially in patients with concomitant hypovolemia.\(^{10}\) Administration of a vasopressor (e.g., norepinephrine) is a temporizing measure and should be initiated while preparing for emergency pericardiocentesis. Finally, positive pressure ventilation (e.g., mechanical ventilation) should be avoided if possible because, as discussed in the section on physiology, it can lead to hemodynamic collapse secondary to changes in intrathoracic pressure.

**INDICATIONS FOR PERICARDIOCENTESIS**
There are two indications for pericardiocentesis: (1) to diagnose the presence and cause of a pericardial effusion (diagnostic pericardiocentesis) and (2) to relieve tamponade (therapeutic pericardiocentesis). Diagnostic pericardiocentesis is an elective procedure. It is ideally performed under visual guidance (e.g., ultrasound or fluoroscopy). Therapeutic pericardiocentesis may be an urgent or emergency procedure based on patient hemodynamics. In the semi-stable patient, ultrasound guidance should be utilized for pericardial fluid aspiration; however, in the case of cardiovascular collapse or cardiac arrest, a blind approach may be warranted.

**Diagnostic Pericardiocentesis**
The use of pericardiocentesis to determine the cause of nonhemorrhagic effusions is common practice despite varying opinions on its utility.\(^{52,122,123}\) Recovery of pericardial fluid to assess for tumor, autoimmune, and biochemical markers in addition to bacterial and viral cultures with Gram stain can be valuable in making the diagnosis. Measurement of pericardial fluid pH can also be helpful because inflammatory fluid is significantly more acidic than noninflammatory fluid.\(^{124}\) When a specific cause is suspected, additional diagnostic testing may be useful (e.g., adenosine deaminase in patients with tuberculosis and carcinoembryonic antigen in those with suspected malignancy).\(^{125}\)

The diagnostic accuracy of pericardiocentesis is variable, and certain diagnoses are unlikely to be made from pericardial fluid. In one large series, fluid samples were obtained in 90% of aspirations, but the specific cause was determined from only 24% of those specimens.\(^{126}\) Another series demonstrated false-negative cytologic results in certain cases of lymphoma and mesothelioma.\(^{126}\) In some HIV patients, effusions secondary to Kaposi’s sarcoma and cytomegalovirus infection have been diagnosed by pericardial biopsy following nondiagnostic fluid analysis.\(^{127,128}\) Nevertheless, pericardial biopsy is not routinely indicated and is considered only in cases of recurrent effusions, such as a neoplasm or granulomatous disease.\(^{129}\) When a biopsy is obtained it is often during implementation of a subxiphoid pericardiotomy (pericardial window).

A pericardial window is typically employed to obtain both pericardial fluid and a pericardial biopsy specimen, because once a tissue sample is obtained, a definitive diagnosis is much more likely.\(^ {110}\) It is reserved for those with recurring episodes of tamponade.\(^{130}\) This procedure can be performed safely in the operating suite without general anesthesia.\(^{151}\) In a prospective series of 57 patients who underwent subxiphoid pericardiotomy, a definitive diagnosis was obtained in 36%, a probable diagnosis in 40%, a possible diagnosis in 16%, and no diagnosis in 7%. It is uncertain whether this technique is safer than
ultrasound-guided pericardiocentesis, but published reports show low rates of complications in experienced hands.138

The most valuable clinical predictors in diagnosis are size of the effusion (larger effusions and tamponade are more likely to yield a diagnosis) and signs of inflammation (e.g., fever, pericardial friction rub, ST-segment elevation).138 Unfortunately, the only methods of making definitive diagnosis are pericardial fluid aspiration with analysis and pericardial biopsy.

Diagnostic pericardiocentesis has limited utility for hemopericardium secondary to traumatic causes. When used diagnostically after trauma to assess for the presence of pericardial bleeding, the procedure has a false-negative rate (i.e., no blood aspirated) of between 20% and 40%.95,132–134 The high false-negative rate is due to the fact that posttraumatic blood tends to clot within the pericardial space and therefore cannot be aspirated.17 Furthermore, pericardiocentesis should not delay emergency thoracotomy if cardiac tamponade is suspected.135

If there is uncertainty about the presence of tamponade, the focused abdominal sonography in trauma (FAST) examination rapidly and noninvasively identifies pericardial fluid.136–138

**Therapeutic Pericardiocentesis**

The ultrasonographic finding of a large pericardial effusion (>20 mm) in a stable patient should lead to early cardiology or cardiothoracic surgery consultation for percutaneous drainage or placement of a pericardial window. Patients with a pericardial effusion who remain hypotensive despite fluid resuscitation require urgent therapeutic drainage. The decision to wait for consultants is best made by the emergency physician at the bedside and should be based on clinical judgment. Until recently, few algorithms existed to assist in the risk stratification of patients with pericardial effusions and who should undergo emergency versus delayed pericardiocentesis. In 2012, Halpern and colleagues introduced a scoring system based on effusion size, echocardiography derangements, and clinical criteria.139

Similarly, Ristic and colleagues in 2014 created a system incorporating etiology, clinical factors, and diagnostic imaging.132 Such tools can aid the emergency physician in identifying who would benefit most from ED pericardiocentesis and who could await consultant recommendations. (Fig. 16.10)

For patients in extremis, pericardiocentesis should be performed immediately even if consultation is unavailable. The clinician who performs the procedure should be the one who is most experienced in both sonography and pericardial fluid aspiration.

**Tamponade of Uncertain Cause:** Pulseless Electrical Activity

A major indication for emergency pericardiocentesis is a patient in cardiac arrest with PEA. Always consider cardiac tamponade in the differential diagnosis for PEA, especially if jugular venous pressure is elevated or a pericardial effusion is demonstrated on ultrasound. In a series of 20 patients with PEA, 3 had tamponade and another 5 had some degree of pericardial effusion.141

In this setting, blind (i.e., landmark method), ECG-guided, or ultrasound-guided pericardiocentesis can be lifesaving.

**Tamponade Caused by Nonhemorrhagic Effusions**

Most nonhemorrhagic effusions are liquid. They can be drained by pericardiocentesis with a small needle or by a catheter left in the pericardial space. Removal of even a small amount of fluid can cause immediate and dramatic improvement in blood pressure and cardiac output. Pericardiocentesis relieves tamponade caused by nonhemorrhagic effusions in 60% to 90% of cases.54,126,142

If aspiration with small-needle pericardiocentesis fails, the patient may have a purulent, malignant, or loculated effusion. Placement of a pericardial catheter is more useful in the long-term management of these patients. In Krikorian and Hancock’s series,126 24% of patients were managed successfully with a single pericardiocentesis procedure, but 37% needed multiple aspirations or an indwelling catheter (39% required surgical drainage and 55% of these patients had traumatic hemopericardium). Therefore if time and patient stability permit, consider pericardiocentesis with a catheter to reduce the necessity for repeated aspirations.

Patients with renal failure and tamponade require urgent pericardiocentesis. Without signs of tamponade, however, these patients may be better managed by dialysis. In one series, 63% of renal failure patients were managed successfully with only dialysis.126 There is some evidence that needle pericardiocentesis is a poor choice in patients who need pericardial drainage; in one series, 9 of 10 patients had serious complications following this procedure.126 Consultation with specialists is advised when there is no evidence of hemodynamic collapse but pericardial drainage is still being considered.

An algorithm for the urgent management of nonhemorrhagic cardiac tamponade is presented in Fig. 16.11.

**Pericardiocentesis in Patients With Hemorrhagic Tamponade**

For hemorrhagic tamponade, pericardiocentesis is never the definitive treatment because this strategy has several drawbacks.143,144 Aspiration of a small quantity of fluid may cause a dramatic improvement in hemodynamics, but pericardial clots can prevent adequate drainage, so blood usually reaccumulates.91,136 Patients with traumatic pericardial hemorrhage ultimately require thoracotomy to explore and repair the cardiac injury. Pericardiocentesis simply delays this definitive procedure. A study investigating traumatic cardiac injury found that all patients who underwent surgery within 2 hours of the injury survived but the mortality rate was higher in patients who experienced longer operating room delays.145

Sugg and associates documented a 43% mortality rate when pericardiocentesis was the sole treatment of traumatic tamponade, as compared with 16% in those who also underwent surgical intervention.133 All patients managed by pericardiocentesis had repairable wounds at autopsy, thus suggesting that in this population, operative repair is preferable. The number of deaths from stab wounds has been decreasing over time in response to a shift in trauma philosophy in which early thoracotomy is supported rather than repeated pericardiocentesis.95,134,136,145

In the austere setting where no surgeon is immediately available and hemopericardium is identified on FAST exam, pericardiocentesis can serve as a life-saving bridge until definitive surgical intervention is arranged (Fig. 16.12). A study of penetrating trauma patients with tamponade found that preoperative pericardiocentesis decreased the mortality rate from 25% to 11%.146

**CONTRAINDICATIONS**

There are no absolute contraindications to pericardiocentesis in hemodynamically unstable patients. Relative contraindications
Figure 16.10 Three-step scoring system triage patients requiring urgent percutaneous or surgical drainage of pericardial effusion. Diagnosis of cardiac tamponade is based on clinical symptoms, signs, and echo findings. Total score ≥6 indicates urgent pericardiocentesis. HR, Heart rate; IVC, inferior vena cava; PE, pulmonary embolism; SBP, systolic blood pressure. (From Ristic A, Imazio M, Adler Y, et al: Triage strategy for urgent management of cardiac tamponade: a position statement of the European Society of Cardiology Working Group on Myocardial and Pericardial Diseases, Eur Heart J 35:2279, 2014. Oxford University Press.)
OVERVIEW OF TECHNIQUES AND EQUIPMENT

In an urgent situation (e.g., PEA arrest) with no adjunctive equipment available, pericardiocentesis can be performed with minimal equipment. Use a long 18-gauge spinal needle attached to a 10-mL syringe to withdraw fluid from the pericardial sac. Although the procedure can be done with these two simple devices, access to several others would be beneficial during the procedure. Of all the components, the most essential is an ultrasound machine, which first is used to determine whether a pericardial effusion is present and then assists in accurate needle placement. Ideally, use a probe with a small-footprint and a frequency of 2 to 4 MHz. If this type of probe is not available, use a 2- to 3.5-MHz curvilinear probe in the subxiphoid view, which will also provide excellent images of the heart.

Before the introduction of real-time sonography to guide needle placement, electrocardiographic monitoring was used to indicate appropriate needle placement. This is done by connecting the electrocardiographic machine to one of the precordial leads (e.g., V1). That precordial lead is then attached to the distal end of a spinal needle with an alligator clamp to pericardiocentesis include coagulopathy; previous thoracoabdominal surgery; the presence of prosthetic heart valves, pacemakers, or cardiac devices; inability to visualize the effusion with ultrasound during the procedure; and situations in which better treatment modalities are immediately available (e.g., thoracotomy for trauma patients).

Ideally, pericardiocentesis is performed in the cardiac catheterization laboratory under fluoroscopic or echocardiographic guidance. With the advent of bedside ultrasound and immediate visualization of large pericardial effusions, pericardiocentesis is being performed in the ED more frequently. Ultrasound can accurately identify the area of the heart with the greatest fluid accumulation and clarify its relationship to the body wall. This allows the physician to choose an entry site and angle of penetration with the greatest likelihood of obtaining fluid while avoiding vital structures.
temporizing methods are the mainstay of therapy unless the patient is unstable (e.g., in PEA arrest). The most common therapeutic procedures used as temporizing measures in the setting of tamponade are intravascular volume expansion with crystalloids and administration of vasopressors or inotropes. Although most textbooks and protocols encourage the use of Norepinephrine, isoproterenol, dopamine, and dobutamine, these temporizing methods are backed by only sparse scientific evidence. Studies in animals have shown that any of these agents may be beneficial as a temporizing agent only, but theoretically, dobutamine is preferable because of its greater β-adrenergic activity.

**Preparation**

Before preparing for pericardiocentesis, place all resuscitation equipment at the bedside in anticipation of clinical deterioration. Most patients undergoing pericardiocentesis in the ED have already experienced hemodynamic collapse and are lying supine. If the patient is able to cooperate, elevate the chest 30 to 45 degrees to bring the heart closer to the chest wall. Sedation of stuporous patients is typically forborne because of the risk for further hemodynamic collapse. If the patient is awake and undergoing the procedure without obvious hemodynamic compromise, short-acting medications (e.g., ketamine, midazolam, or fentanyl) are preferred.

Every effort should be made to ensure aseptic technique. Prepare the chest and upper part of the abdomen with a chlorhexidine-based solution. Drape the patient and ensure that all care providers involved in the procedure are wearing a sterile hat, mask, gown, and gloves. If the patient is awake, anesthetize the skin and the proposed route with 1% lidocaine (see Review Box 16.1). Because the pericardium is extremely sensitive, it should be anesthetized.

The approach to pericardiocentesis depends on the clinical status of the patient, the availability of ultrasound, and the distribution of pericardial fluid. Pericardial fluid is not always distributed circumferentially in the pericardial sac, so ultrasound can quickly identify the maximal effusion pocket and demarcate the appropriate site for needle placement. Pericardiocentesis with ultrasound guidance is currently the safest and most reliable method for the diagnosis and treatment of pericardial effusion and tamponade. Studies of echocardiography-directed pericardiocentesis have found that the apical approach is the best site for puncture. Cadaver studies have corroborated this finding and have demonstrated greater safety with an apical approach. However, these studies also revealed that an apical approach is associated with a greater incidence of pneumothorax than a traditional substernal approach is. Before the advent of ultrasound guidance, the subxiphoid approach (discussed later in this chapter) was the preferred method of pericardiocentesis. It is still used frequently during cardiac arrest and when ultrasound is not readily available.

**PROCEDURE**

**Temporizing Measures**

Cardiac tamponade is an emergency that requires urgent therapy. Therapy typically consists of either pericardial drainage by needle aspiration or placement of a pericardial window. These procedures are not classically performed in an ED, so temporizing methods are the mainstay of therapy unless the patient is unstable (e.g., in PEA arrest). The most common therapeutic procedures used as temporizing measures in the setting of tamponade are intravascular volume expansion with crystalloids and administration of vasopressors or inotropes. Although most textbooks and protocols encourage the use of these temporizing methods, they are backed by only sparse scientific evidence. Studies in animals have shown an increase in cardiac output and improvement in blood pressure with expansion of central blood volume; the validity of this in humans with cardiac tamponade is uncertain. Fluid resuscitation in a trauma patient with penetrating cardiac injury might cause deterioration. Animal experiments indicate that the response depends on whether fluid boluses produce recurrent bleeding from the cardiac wound. Despite the lack of evidence, judicious volume expansion with or without an adjunctive vasopressor before definitive therapy may be the only option for patients with tamponade.

Norepinephrine, isoproterenol, dopamine, and dobutamine have all been evaluated as the vasopressor or inotropic of choice in patients with cardiac tamponade. Norepinephrine and isoproterenol increased cardiac output in animal models of tamponade, but failed to increase it in humans. Dopamine and dobutamine increased cardiac output and improved hemodynamics in the setting of tamponade. Any of these...
is probably touching the epicardium. Withdraw it several millimeters to prevent laceration of the myocardium or coronary vessels. After slight withdrawal, the needle should be within the pericardial space. Aspirate any fluid, but watch for changes on the ECG. Electrocardiographic monitoring is not infallible: if the patient has an abnormal myocardium from conditions such as a previous MI or the formation of scar tissue, no current-of-injury pattern will be generated on the rhythm strip.

**Ultrasound-Guided Pericardiocentesis**

Pericardiocentesis has traditionally been performed blindly. This approach was associated with a low success rate and a high rate of complications, such as inadvertent puncture of the lung, ventricle, or epicardial vessels. Using ultrasound to both diagnose and guide pericardiocentesis has resulted in increased success rates, as well as a lower rate of complications. The techniques are described in the Ultrasound Box.

**Subxiphoid/Subcostal Approach**

As mentioned earlier, the traditional blind subxiphoid approach can still be used for pericardiocentesis in the ED (e.g., for patients in cardiac arrest and when ultrasound is not available). The technique is performed as follows: introduce the needle 1 cm inferior to the left xiphocostal angle at a 30-degree angle to the skin (Fig. 16.15). Because the heart is an anterior structure, angles greater than 45 degrees may lacerate the liver or stomach. Aim toward the left shoulder and advance the needle slowly while continuously maintaining negative pressure on the syringe to aspirate any fluid. Aspirate with an “in-and-out” vector only, not “side-to-side,” which may lacerate tissue. If no fluid is aspirated, withdraw the needle completely and redirect it in a deeper posterior trajectory. If no fluid is aspirated after redirecting the needle, withdraw the needle and redirect it, working from the patient’s left to right, until it is aimed at the right shoulder. Recommendations regarding needle trajectory vary widely, including toward the right shoulder, sternal notch, and left shoulder.189,135

**Apical Approach**

The apical approach is occasionally used as an alternative to the subcostal approach to drain a pericardial effusion when ultrasound is available. Use ultrasound to identify the largest area of the apical effusion (Fig. 16.16A) or simply feel for the apex. If the apex cannot be palpated, it typically lies within the area of cardiac dullness, often between the fifth, sixth, or seventh intercostal space, between the midclavicular and midaxillary lines. Introduce the needle 1 cm lateral to and into the intercostal space below the apical heartbeat. Advance the needle over the cephalad border of the rib and aim it toward the right shoulder to avoid the neurovascular bundle located caudal to the rib space. This area is close to the lingula and the left pleural space, thus making pneumothorax a frequent complication. Theoretically, this technique is used because the coronary vessels are small at the apex; therefore if a ventricle is entered, it is the thick-walled LV, which is more likely to seal off after ventricular injury. With echocardiographic guidance, the apical approach is used more commonly.109

**Parasternal Approach**

The parasternal approach is an alternative approach to the previously described techniques. First, identify the largest area of the parasternal effusion on ultrasound if possible. If ultrasound is not available or if the effusion is not clearly identified on the ultrasound image, proceed by introducing the needle 1 cm lateral to the sternal border at the left fifth or sixth intercostal interspace. Advance the needle over the cephalad border of the rib to avoid the neurovascular bundle on the caudal aspect of the rib. Avoid going too far laterally from the sternal border because of potential injury to the internal mammary artery.160 Occasionally, a right parasternal approach may be used when ultrasound predicts superior access to an effusion from this direction.

Tsang and co-workers161 described this technique for ultrasound-guided pericardiocentesis in 1998. The ideal site for skin puncture is where the largest area of fluid accumulation is closest to the skin surface. On ultrasound, this is indicated by a large anechoic (black) area at the top of the screen, usually corresponding to the left anterior chest wall (rather than the subcostal region). This approach also avoids injury to the liver (common with the subcostal approach). Inadvertent puncture of the lung is also prevented with this approach because air in the lung will not conduct sound waves and will prevent visualization of the heart when located immediately beneath the probe. Avoid choosing a site that could puncture the internal mammary artery, which lies 3 to 5 cm from either parasternal border, or the neurovascular bundle, which is located at the inferior border of the rib. Mark the best site with a sterile pen.

**Procedure and Technique**

Confirm the trajectory and depth of the needle before puncturing the skin. Be aware that repositioning the patient alters the position of the heart and pericardial sac within the chest, so reassessment will be necessary. Prepare the skin antisепtically and place a sterile cover over the ultrasound probe. If time permits, anesthetize the selected area with 1% lidocaine, with the superior border of the adjacent rib being used as a landmark. Select an 18-gauge spinal needle. Ideally, the needle should have a sheath that allows it to be withdrawn after the pericardial space is entered. This helps avoid injury to the heart and other vital structures. Attach a saline-filled syringe to the needle, and gently aspirate while slowly advancing the needle. Keep the ultrasound probe on the chest wall, immediately adjacent to the aspiration site.

Once the pericardial space is entered, inject agitated saline to confirm needle placement, particularly if the pericardial fluid is grossly bloody or if there is any question about needle
**PERICARDIOCENTESIS (SUBXIPHOID APPROACH)**

1. Examine the patient and identify the xiphoid process and the costal margin. Prepare the area with antiseptic and administer local anesthetic.

2. Introduce the needle 1 cm inferior to the left xiphocostal angle at a 30-degree angle to the skin. Aim toward the left shoulder.

3. Aspirate during needle advancement and monitor for fluid return. Stop advancing once fluid is returned.

4. Needle withdrawn. ST segment elevation returns to normal.

5. Advance a J-tipped guidewire through the needle and into the pericardium.

6. Remove the needle, while leaving the guidewire in place in the pericardium.

7. Advance a 6- to 8-Fr dilator over the wire and then remove the dilator.

8. Advance a 6- to 8-Fr pigtail catheter over the wire and into the pericardium.

9. Remove the wire and drain the pericardial fluid.

**Figure 16.15** Pericardiocentesis (preferred subxiphoid approach). (From Custalow CB: *Color atlas of emergency department Procedures*, Philadelphia, 2005, Saunders, p 123.)
cardiographic contrast agent (e.g., Definity) as a safe and successful alternative. Contrast agents contain gas microbubbles, which markedly enhance the fluid echo by introducing multiple liquid-gas interfaces. Inject this solution as a bolus. If the contrast material clears immediately after administration (as occurs with agitated saline) or persists temporarily within the cardiac chambers, an intracardiac location is suggested.

**Fluid Aspiration and Evaluation**

Removal of even a small amount of pericardial fluid (e.g., 30 to 50 mL) usually results in either return of spontaneous circulation or hemodynamic improvement. After any approach used for pericardiocentesis, place a temporary drain not only to ensure rapid access into the pericardial sac but also to allow more fluid to be removed quickly if hemodynamic collapse recurs. After needle placement is confirmed, a temporary drain can be placed by the Seldinger technique, described in Chapter 22.

Remove the syringe from the needle, advance a guidewire through the needle, and then remove the needle (see Fig. 16.15). Position a dilator (6 to 8 Fr Cordis) over the wire. If a dilator is not used, particularly with the subxiphoid approach, the pigtail catheter tip may get caught in the subcutaneous tissue and make placement of the catheter difficult. Remove the dilator and slide an introducer sheath dilator (6 to 8 Fr Cordis) over the wire. Remove the wire and the dilator while leaving the introducer sheath in place. Insert the pigtail angiocatheter through the introducer sheath, and aspirate fluid to confirm placement. If drainage of pericardial fluid becomes sluggish, flush the catheter with a heparinized saline solution to ensure patency of the lumen.

Aspiration of blood during pericardiocentesis raises the possibility of cardiac puncture. Blood retrieved from the ventricle usually clots faster than bloody fluid aspirated from the pericardium. In general, hemorrhagic pericardial effusions have local fibrinolytic activity, which prevents clot formation. If the bleeding is brisk enough, however, blood may still clot and does not necessarily point toward ventricular puncture. The hematocrit of pericardial fluid should always be lower than that of a sample from the systemic vascular system, except in patients with aortic dissection or acute myocardial rupture. These circumstances aside, a hematocrit value similar to that for systemic blood should raise concern for an intracardiac needle location. Several other simple laboratory tests can differentiate normal from abnormal pericardial fluid, but they require the availability of a centrifuge system and time.

Under normal conditions, pericardial fluid is less than 50 mL in volume and clear to pale yellow in color with no red or white blood cells, inflammatory markers, bacteria, or cancer cells and with a glucose concentration similar to that of blood. Immediately following the procedure, obtain a chest film to ensure the absence of pneumothorax and free air under the diaphragm. Place the patient on continuous cardiac monitoring for 24 hours and watch for signs of reaccumulating fluid or

![Figure 16.16](image-url) Placement of a pigtail catheter in the pericardial space under ultrasound assistance: subcostal view of a small but hemodynamically significant pericardial effusion during pericardiocentesis. 

- **A**, The effusion. 
- **B**, After the injection of approximately 0.1 mL of agitated saline through the pericardiocentesis needle to confirm its position in the pericardial space. 
- **C**, The shaft of the pigtail catheter (arrowhead, two discrete parallel echogenic lines reflect the catheter walls; the echo-free area represents the catheter lumen) lying in the pericardial space after the majority of fluid has been drained. LI, Liver; PE, pericardial effusion; RV, right ventricle; SC, saline contrast.
iatrogenic complications. Repeating the ultrasound examination in 24 hours is recommended. Diagnostic evaluation of non-hemorrhagic fluid is similar to that for pleural fluid (see Chapter 9).

Suture the pigtail catheter to the skin, but be careful not to occlude the catheter by tying it too tightly. Wrap the catheter in gauze at the skin and cover it with a sterile dressing. Attach the catheter to suction tubing and a drainage system.

COMPLICATIONS

Emergency physicians often perform pericardiocentesis under duress on a patient in PEA arrest. Many also perform the technique blindly because they have little or no time to gather adjunctive assistance or tools. It is critical for the emergency physician to be aware of both the traditional and contemporary methods of performing the procedure and the complications that can be associated with these methods (see Review Box 16.1).

With the advent of ultrasound- and CT-guided pericardiocentesis, the complication rate has been greatly reduced. Complication rates as low as 4% have been reported in large observational studies. Earlier studies of blind pericardiocentesis documented morbidity rates of 20% to 40% and mortality rates as high as 6%. Because pericardiocentesis is performed in moribund patients, the likelihood of cardiac arrest and death is high. However, they are not usually a direct complication of pericardiocentesis but of poor cardiopulmonary reserve. Cardiac arrest and death are rarely associated with echocardiographically guided pericardiocentesis. When blind or electrocardiographically guided pericardiocentesis is performed, the patient is usually already in full arrest and attributing the cause of death to the procedure is nearly impossible. In a series of 52 patients the only death occurred in a patient in cardiogenic shock in whom pericardiocentesis was nonproductive and who was found to have severe arteriosclerotic heart disease, not shock in whom pericardiocentesis was nonproductive and who responded to atropine and fluid loading. Duvernoy and associates reported 1 case of ventricular tachycardia and 1 case of atrial fibrillation in 352 procedures. Maggiolini and co-workers reported transient third-degree heart block in a single patient.169

The traditional subxiphoid approach carries a risk for liver laceration. Fortunately, inadvertent needle passage into the liver has not been reported to cause significant hemorrhage or death.

The parasternal and apical approaches have been documented as causing pneumothorax and pneumopericardium in several case series, but without any clinical consequence (Fig. 16.17). The pneumothoraces were treated with 100% oxygen or thoracostomy. There have also been infrequent reports of pneumopericardium after removal of a pericardiocentesis catheter. The cause of the pneumopericardium is thought to be the formation of a bronchopericardial fistula, but the exact mechanism is unclear. The mortality rate associated with tension pneumopericardium is approximately 50%, so consider pneumopericardium when patients complain of dyspnea and hypotension after removal of their catheter.

Very few studies have reported ventricular or coronary vessel laceration during pericardiocentesis. These complications occur more frequently during blind or electrocardiographically guided procedures. Most cardiac perforations occur in the RV, but punctures in the LV and atria have also been reported. When these perforations occur, they tend to be silent and result in hemopericardium and death. In patients taking anticoagulants, it is important to check coagulation factors and monitor them closely after a seemingly insignificant pericardiocentesis because hemopericardium could develop just from the procedure itself.

In the series compiled by Krikorian and Hancock, hemopericardium developed in 13 of 123 patients as a result of pericardiocentesis, one as a result of a lacerated coronary artery. One patient died of a punctured ventricle. Surgical control was necessary in four patients in whom tamponade developed, whereas it did not develop in eight patients with hemopericardium, and they were managed conservatively. Guberman and colleagues reported one episode of ventricular tachycardia and several “hypotensive vasovagal reactions” that were associated with bradycardia and

Figure 16.17 Air-fluid level (arrow) in the pericardial space immediately after pericardiocentesis. A minor pneumopericardium is inconsequential; a larger collection may cause tamponade.
pericardiocentesis, but none caused any adverse sequela. In
their series of 352 procedures, Duvernoy and associates168
reported 23 penetrations. In two cases both the RV and LV
had been perforated, and in all other cases the RV had been
entered.

Researchers differ in their opinions regarding the adverse
effects of ventricular puncture. Most ventricular punctures
involve the lower aspect of the RV. The wall of the RV is thin
and therefore vulnerable to laceration. However, pressure in
the RV is low,2 so a puncture should cause little bleeding. In
a series of patients who underwent ultrasound-directed peri-
cardiocentesis, ventricular puncture occurred in 1.5% but was
without consequence.157 In another study, laceration of the RV
occurred in 1 patient despite the use of echocardiography; it
resulted in tamponade and necessitated emergency surgery.123
Of the 23 perforations in the series by Duvernoy and associates,
3 were considered major complications (2 patients required
thoracotomy). Left ventricular pseudoaneurysm typically occurs
as a complication of MI. It is rarely seen after surgery, trauma,
or infection. Rare cases of severe left ventricular pseudoaneu-
rysm after pericardiocentesis have been reported.174,175

Even when pericardiocentesis has induced no physical injury,
adverse events have been documented. Most have to do with
the fact that during pericardiocentesis the stroke volume of
the previously collapsed RV increases 75% after the first 200 mL
of fluid is removed.7 In general, this increase in stroke volume
is greater initially than that demonstrated by the LV. This
imbalance can cause significant consequences for both right
and left ventricular function. Three of six patients in whom
large effusions were removed by pericardiocentesis experienced
right ventricular dilation and overload, abnormal septal motion,
and either no increase or a decrease in the right ventricular
ejection fraction.176 These patients subsequently and slowly
returned to normal hemodynamic status.

Pulmonary edema following pericardiocentesis has also been
reported, presumably caused by a sudden increase in venous
return to the LV when peripheral vascular resistance is still
high from compensatory catecholamine secretion.177–181 Sup-
porting evidence for this explanation is that right ventricular
stroke volume increases more than left ventricular stroke volume
after relief of tamponade.16 Circulatory collapse with persistently
low arterial blood pressure has been reported in a patient from
whom 700 mL of clear fluid was drained at a rate of 100 mL/
min.182 Thus, many authors recommend that the pericardial
drainage rate not exceed 50 mL/min.

Acknowledgments
The editors and authors acknowledge the contributions of
Richard J. Harper to this chapter in previous editions. Linda
J. Kesselring, MS, ELS, from the Department of Emergency
Medicine at the University of Maryland School of Medicine,
copyedited the manuscript for the previous edition.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES

26. Solod S, Thomas HM, Jr, Evans R: Myocardial laceration not demonstrated by continuous electrocardiographic monitoring occurring during peri-
CONVENTIONAL CPR

Although CPR is widely taught to health care personnel and reassessed periodically, the importance of high-quality CPR cannot be stressed enough. High-quality CPR immediately before defibrillation increases the chance of successful restoration of circulation.1,2 Consensus opinion, based on a growing foundation of clinical data, is that early CPR and defibrillation (when appropriate) have a significant impact on patient survival and recovery.2-11 Quality chest compression also increases the efficacy of drugs administered during resuscitation, whereas inadequate circulation leads to minimal effects from peripherally delivered drugs.2 Hyperventilation is also widely prevalent and dramatically compromises hemodynamics. In animal studies, hyperventilation leads to reduced survival from arrest. In this section we review the key procedural aspects of manual CPR.

Compressions

The 2015 resuscitation guidelines emphasize the importance of quality chest compression4 by recommending that clinicians focus on maintaining proper chest compression depth and rate. Compress the sternum to a depth of between 2.0 and 2.4 inches with a rate of between 100 and 120 compressions/min. Box 17.1 provides a summary of procedural recommendations for CPR. If possible, place a backboard under the victim to ensure appropriate thoracic compression. In addition, adjust the height of the bed or have the rescuer stand on top of a stepstool so that the entire weight of the rescuer above the waist is directed onto the patient’s sternum (Fig. 17.1A). This enhances the depth of compressions and helps prevent leaning on the patient’s chest between compressions, which is another key deficiency that has been widely observed. Extend the arms fully and place them perpendicular to the patient’s chest while making sure to pull away from the chest sufficiently between compressions to allow full chest recoil. Rotate rescuers aggressively (approximately every 2 to 3 minutes) to avoid deteriorating quality of compressions because of exhaustion. Properly delivered compressions are highly fatiguing, and rescuer bravado often interferes with the realization of declining CPR quality over time.

Minimize pauses in chest compressions because even short pauses have profound effects on coronary perfusion pressure and outcomes.13 As stated earlier, long pauses in chest compressions before delivery of a shock are associated with failure of defibrillation.3 Do not stop CPR to deliver medications because the drugs can be administered at the same time as the compressions. Keep pauses in chest compressions to a minimum (e.g., for procedures such as intubation or pulse checks).

Ventilations

Deliver ventilations at a rate of 8 to 10 breaths/min (see Fig. 17.1B). Hyperventilation (e.g., ventilation rates greater than 30/min) is common during resuscitation. To prevent unwittingly hyperventilating the patient, one practical technique is to ask the rescuer who is providing ventilations to remove his or her hand completely off the bag-valve-mask apparatus between ventilations. The team leader should be vigilant in the observation of delivery of ventilations and should be ready to verbally prompt rescuers to ventilate the patient at the appropriate rate if hyperventilation is performed.

Pulse Checks

Pulse checks are generally performed too frequently during resuscitation efforts and take too much time. If a pulse cannot be readily felt within seconds, return to chest compressions as soon as possible. No studies have suggested that CPR is harmful to a patient with a very weak pulse, so use of a Doppler ultrasound device to detect the pulse is discouraged, as it is unnecessary and leads to increased pause times. If rescuers need ultrasound to find a pulse, the patient is at the very least markedly hypotensive and should probably be receiving CPR. Attempt pulse detection at the location of the carotid or femoral artery because peripheral pulse checks during profound shock or cardiac arrest states are notoriously unreliable. Frequently, a “pulse” can be detected during CPR itself; this phenomenon is often due to venous back pressure during compressions and does not indicate that compressions should be stopped, nor
CHAPTER 17 Cardiopulmonary Resuscitation and Artificial Perfusion During Cardiac Arrest

333 of low blood flow. If the patient achieves return of spontaneous circulation (ROSC), a sharp increase in the PETCO₂ value (usually greater than 25 to 30 mm Hg) is consistent with return of adequate perfusion.¹⁴

Leadership and Teamwork

Cardiac arrest resuscitations are often crowded, chaotic events filled with stress and anxiety. To maximize calm and efficiency and to ensure quality of care, establish a team protocol. Designate someone to be the leader of the resuscitation, and make sure that all participants are clearly aware of this designation. The designated team leader should be responsible for monitoring the rhythm, for giving orders to initiate and terminate chest compressions, and for delivery of drugs and other therapies. The team leader should be situated either at the head of the bed or at a place where they can direct the resuscitation. It is important that the team leader does not actually perform compressions, ventilations, or other specific procedures unless absolutely necessary because they may quickly lose control of the resuscitation. As most rescuers are unable to detect when their own quality of compressions is diminishing, the team leader must observe CPR closely and order rescuer rotations throughout the duration of the resuscitation.¹⁵

Important Option for Layperson CPR: Chest Compression-CPR

Chest compression-only CPR (CC-CPR) has been shown in a number of investigations to be as effective as standard CPR in resuscitation efforts initiated by members of the lay public.¹⁶,¹⁷ Give compressions at a rate between 100 and 120 per minute, as with conventional CPR with interposed ventilations. Because of its simplicity, CC-CPR minimizes pauses in chest compressions while maintaining proper rate and depth. Lay rescuers in the community may be less experienced with standard CPR and uncomfortable with the performance of mouth-to-mouth resuscitation. The simplicity of CC-CPR makes it relatively easy for first responders to initiate resuscitation efforts and for emergency medical dispatchers to guide lay rescuers remotely. Starting in 2010, and affirmed in 2015, the American Heart Association guidelines have shifted emphasis from “ABC” (“airway, breathing, compressions”) to “CAB” (“compressions, airway, breathing”) for lay rescuers. Their endorsement of “hands-only CPR” (a synonym for CC-CPR) educational programs reflects additional evidence that a focus on chest compressions during CPR may lead to an increase in bystander CPR, as well as improvements in patient outcomes.¹⁸ Recent investigations have shown that CC-CPR is associated with improved survival of patients with out-of-hospital cardiac arrest when performed by lay bystanders. A period of CC-CPR before intubation and rhythm evaluation also improves outcomes when used by emergency medical service (EMS) personnel.¹⁹,²⁰ The EMS community is likely to see more widespread adoption and use of CC-CPR by lay public educational programs in the upcoming years.

ADJUNCTS TO IMPROVE THE QUALITY OF CPR

Numerous techniques and adjunctive devices have been investigated in attempts to improve long-term survival rates

---

**BOX 17.1** Key Procedural Elements of Manual CPR

**COMPRESSIONS**
- Between 100 and 120 compressions/min
- Depth of between 2.0 and 2.4 inches/compression
- Allow full chest recoil between compressions
- Minimize pauses in compressions

**VENTILATIONS**
- 8–10 ventilations/min (avoid hyperventilation)
- Minimize pauses in chest compression for intubation
- Use of continuous capnography recommended for intubated patients

**Figure 17.1** Conventional cardiopulmonary resuscitation (CPR). Note: no alternative technique or device in routine use has consistently been shown to be superior to conventional CPR. **A**, Compress the sternum to a depth between 2.0 and 2.4 inches at a rate between 100 and 120 compressions/min. Better CPR can be achieved by having the rescuer stand on a stepstool during compressions, rotating rescuers every 2 to 3 minutes, and minimizing pauses. **B**, Deliver ventilations at a rate of 8 to 10 breaths/min. Avoid hyperventilation during resuscitation.

**CPR**, Cardiopulmonary resuscitation.
with CPR. Data are conflicting and contrary, and as of this writing, no alternative technique or device in routine use has consistently been shown to be superior to conventional CPR. Unless breakthrough technology or new information on the parameters affecting the outcome of CPR emerge, this admonition will probably endure. Nonetheless, a variety of technologies have been developed to improve the quality of CPR. Some of these tools directly improve chest compressions, whereas others are less direct and aim to improve human performance or enhance hemodynamics during the delivery of chest compressions. This section describes some of these promising, intuitively useful, yet still unproven techniques.

**Active-Compression Decompression CPR**

Active compression-decompression CPR (ACD-CPR) is a variant of CPR in which the passive relaxation phase of CPR is converted into an active phase by means of a handheld or mechanical suction device, which can theoretically improve both myocardial and cerebral circulation when compared with traditional CPR. However, data on these devices are mixed; there have been studies on out-of-hospital cardiac arrest using this technique that did not find any improvements in either initial outcome or survival to discharge, and as with many devices, there are instances when its application is impractical.

**Impedance Threshold Device**

The impedance threshold device (ITD) optimizes chest compression hemodynamics via manipulation of intrathoracic pressure. From a practical standpoint, the ITD is a relatively simple device that is placed between the endotracheal tube and the bag-valve apparatus, much like a colorimetric PETCO₂ detector, which is familiar to most emergency department (ED) clinicians (Fig. 17.2). The ITD contains a valve that prevents air that is less than 10 cm H₂O in pressure from flowing through the device. During resuscitation, the ITD prevents air from entering the thorax during recoil of the chest wall after each compression by generating a small but hemodynamically significant negative pressure within the chest. In laboratory studies, this negative pressure enhances venous return to the heart and results in increased cardiac output with each subsequent chest compression.

The ITD can be used during resuscitation either with mask ventilation or via an endotracheal tube and is therefore appropriate for both basic life support care in the field and ED resuscitation. Apply the device and administer ventilations at a rate of 8 to 10 breaths/min as per standard resuscitation guidelines. The Res-Q-Pod ITD (Advanced Circulatory Systems, Inc., Roseville, MN) also has a flashing light timed to prompt the appropriate ventilatory rate. When using it with a face mask, it is important to continuously maintain a tight seal between the patient’s face and the mask during CPR to maintain efficacy of the ITD. This is best accomplished with a two-person ventilation technique in which one person holds the face mask and the second person squeezes the bag. If a pulse is restored, remove the ITD from the respiratory circuit.

Current data are conflicting on whether ITDs improve clinical outcomes when used as an adjunct to resuscitation efforts. Numerous studies and clinical trials using one particular model of ITD (Res-Q-Pod, Advanced Circulatory Systems, Inc.) have demonstrated improved hemodynamics during CPR and have suggested that use of an ITD during resuscitation efforts may lead to improved survival and patient outcomes. However, the findings from randomized controlled trials of ITD use in patients suffering out-of-hospital cardiac arrest have offered opposing data, thus suggesting that there is not a significant improvement in patient outcomes when these devices have been used.

**Monitoring and Feedback Devices**

Emphasis on CPR quality and minimizing interruptions has spurred the development of devices to monitor the quality of chest compressions and ventilations and then provide audio or visual prompts to improve performance. These devices aim to improve human delivery of CPR and, unlike ACD-CPR or the ITD, do not enhance hemodynamics or patient physiology directly.

One method of monitoring chest compressions involves placing a relatively small external device on the patient’s sternum and performing chest compressions on top of the device (Fig. 17.3). The device measures the quality of compressions via a force detector or accelerometer (or both) that determines the rate and depth of chest compressions. Different versions of these CPR quality-monitoring and feedback devices are on the market. Some are incorporated into defibrillators (MRx-QCPR, Philips Healthcare, Andover, MA; R series with Real CPR Help, Zoll Medical Corp., Chelmsford, MA), whereas others are stand-alone devices applied to the chest. In trials, use of such a defibrillator with CPR monitoring and feedback improved CPR performance and, in one out-of-hospital and one in-hospital investigation, improved the rate of initial resuscitation. Further research will be required to assess the magnitude of improvement in survival that these devices can offer and what training mechanisms can maximize team responses to feedback messages.
CHAPTER 17  Cardiopulmonary Resuscitation and Artificial Perfusion During Cardiac Arrest

Mechanical CPR Devices

The adjuncts described previously all rely on human performance of CPR. Another general approach to improve CPR quality is to provide compressions via a mechanical device that is independent of human fatigue or vagaries in performance. Such tools have been introduced in previous decades but fell out of favor because of unwieldy design and other practical considerations.

A newer generation of devices has brought the notion of mechanical CPR back to active consideration. One such device uses a “load-distributing compression band” (Autopulse, Zoll Medical Corp.). The Autopulse device works via a wide band that is attached to a backboard and battery-powered motor and is placed across the torso. Through cycles of constriction and relaxation, the band compresses the chest in a circumferential manner at a fixed rate and depth consistent with resuscitation guidelines. In this fashion, pauses are also minimized by eliminating rescuer switching. Such devices have a unique role in out-of-hospital arrest because compressions can be delivered while transporting a patient down stairs or into an ambulance.

Studies to determine the efficacy of the Autopulse have had mixed results. Although initial smaller investigations appeared promising, a large multicenter randomized trial was stopped early because patients in the manual CPR arm had survival equivalent to those receiving care via the Autopulse, with a trend toward worse outcomes in the Autopulse group. A separate nonrandomized trial showed a marked improvement in survival when using the device. A 2007 randomized trial in Europe demonstrated the utility and feasibility of automated compression devices (in this case the Autopulse) in the resuscitation of out-of-hospital cardiac arrest patients. A 2015 pragmatic trial in the United Kingdom, evaluating a mechanical CPR device known as LUCAS (Physio-control, Redmond, WA), demonstrated equivalency between mechanical CPR and traditional manual CPR. The LUCAS device (Fig. 17.4 and Video 17.1), in contrast to the band mechanism of the Autopulse, uses a piston/suction cup to compress the anterior aspect of the chest, much like during manual CPR, with the suction cup providing some degree of active compression-decompression, as described earlier in this chapter.

To highlight an intriguing opportunity available with mechanical CPR devices, there has been much discussion about the potential utility of these tools in clinical situations in which coronary angiography might be performed concurrently with ongoing resuscitation efforts. If clinical evidence suggests a major coronary event as the cause of the arrest, mechanical devices could be used to perform high-quality, continuous chest compressions as percutaneous coronary intervention is being performed. Case studies have demonstrated the feasibility of using the LUCAS device during intrarrest coronary angiography with good patient outcomes, but additional data will be necessary to draw clearer conclusions about clinical practices and patient outcomes in these situations.

Extracorporeal CPR

Extracorporeal CPR (E-CPR) is an emergency technique that has been investigated as a “last resort” for cardiac arrest patients who have failed to achieve ROSC despite ongoing resuscitation efforts (Fig. 17.5). Several clinical studies have demonstrated successful outcomes for patients in whom E-CPR was used (and thus indicate that E-CPR could be a feasible addition to resuscitation efforts). One prospective trial in Japan, which identified patients who failed to respond to other traditional resuscitation efforts, demonstrated a favorable neurologic outcome in patients who were able to undergo both emergency cardiac bypass and therapeutic hypothermia treatment; rapid initiation of E-CPR and attainment of the target temperature were associated with positive neurologic outcomes in this cohort. The specialized training necessary to perform the procedure, as well as significant logistic issues surrounding rapid establishment of extracorporeal membrane oxygenation (Video 17.2) in the ED setting, raises concern about the widespread applicability of this intervention. Other investigations have highlighted the potential complications related to E-CPR in these critically ill patients. Additional research is needed on this topic, and more information will be necessary to clearly identify patients who are likely to benefit from E-CPR.
Arterial blood gas monitoring during cardiac arrest is not a reliable indicator of the severity of tissue hypoxemia, hypercapnia (and therefore the adequacy of ventilation during CPR), or tissue acidosis. Current evidence in patients with ventricular fibrillation neither supports nor refutes the routine use of intravenous fluids. There is no evidence that any antiarrhythmic drug given routinely during human cardiac arrest increases survival to hospital discharge. There is insufficient evidence to recommend for or against the routine use of fibrinolysis for cardiac arrest.

No blood testing is considered routine or standard during the initial stages of cardiopulmonary arrest, although early serum potassium and blood glucose monitoring is prudent if resuscitation is successful.\textsuperscript{14}

### Capnography

Capnography measures respiratory CO\textsubscript{2}, which is delivered to the lungs and expelled during exhalation (Fig. 17.6). The highest CO\textsubscript{2} levels occur at the end of each exhalation, called PETCO\textsubscript{2}. During cardiac arrest, PETCO\textsubscript{2} falls abruptly at the onset of cardiac arrest, increases during the delivery of effective CPR, and returns to physiologic levels after ROSC. PETCO\textsubscript{2}
as a potential method of measuring hemodynamics and perfusion during cardiac arrest, as well as for determining the outcome of resuscitation efforts (specifically, detection of ROSC). The 2015 resuscitation guidelines recommend continuous waveform capnography for all intubated patients during resuscitation efforts.\textsuperscript{14}

### Ultrasound Monitoring

With advances in ultrasound equipment, properly trained users can portably and accurately monitor cardiac function in real time. Preliminary studies have demonstrated that trained physicians can assess cardiac function and obtain adequate images rapidly by using a subcostal approach to standard echocardiography in the cardiac arrest setting.\textsuperscript{44} If you are adequately trained in this technology, use it during resuscitation efforts to clinically diagnose conditions such as pulseless electrical activity (PEA) and to make a global assessment of cardiac motion during CPR and pulse restoration. Use ultrasound during arrest to rapidly diagnose and treat conditions such as cardiac tamponade. Get the ED ultrasound machine ready to use when preparing for an incoming cardiac arrest. Remember, however, that ultrasound is only a secondary diagnostic adjunct and should not interfere with the performance of high-quality CPR. Minimize interruptions to perform ultrasound and use it only during resuscitation for specific purposes (e.g., diagnosis of PEA versus hypotensive sinus rhythm). In most cases of arrest, ultrasound is probably of little value. Finally, there is ongoing research on the use of transcranial Doppler ultrasound to determine the prognosis after cardiac arrest. One preliminary study concluded that patients with severely disabling or fatal outcomes could be identified within the first 24 hours with this method.\textsuperscript{45}

### CONCLUSION

Physicians and other health care workers have been performing CPR for more than 50 years, but only since the 1990s has the full importance of the quality of CPR become apparent through an evidence-based approach. Chest compressions and ventilations appear to be deceptively easy to the newly trained, but in fact they are highly complex skills and are difficult to perform well under stress. New technologies have been developed to assist in delivery of CPR, and use of these tools may improve the ability to save lives from cardiac arrest in the coming years.
REFERENCES


In the United States trauma is the leading cause of death in people aged 1 through 44. Blunt trauma accounts for the majority of trauma mortality overall, but in urban settings penetrating trauma, including firearm-related injuries, accounts for an increased proportion of trauma deaths. In 2013, more than 33,000 firearm-related deaths occurred in the United States, with many victims arriving at the emergency department (ED) in extremis.

Severe traumatic injuries can be associated with a very high mortality rate. However, on rare occasions an aggressive approach involving the use of emergency department thoracotomy (EDT) leads to survival in patients with impending or recent traumatic arrest. EDT is a dramatic, heroic intervention performed outside the operating room and often in the absence of trained cardiothoracic or trauma surgeons. Though supported as a potential lifesaving procedure, EDT is not a mandated standard of care nor a procedure that is expected

**Resuscitative Thoracotomy**

**Indications**
- Trauma victims with severe refractory instability or cardiac arrest
- Nontraumatic hypothermic cardiac arrest

**Contraindications**
- Blunt trauma cardiac arrest patients without signs of life
- Trauma patients with obvious non-survivable injuries
- Initial rhythm of asystole without signs of life
- Signs of prolonged arrest including lividity or rigor

**Equipment**

**Complications**
- Phrenic nerve injury
- Coronary artery injury
- Infection
- Injury/disease transmission to health care worker

---

Review Box 18.1 Resuscitative thoracotomy: indications, contraindications, equipment, and complications.
to be performed in most EDs. The first successful thoracotomy was reported more than 100 years ago, and the first EDT was reported in 1966. Since then, multiple studies have reported outcomes, indications, techniques, and risks associated with the procedure. In 2003, the National Association of EMS Physicians Standards and Clinical Practice Committee and the American College of Surgeons Committee on Trauma (ACSCOT) proposed specific guidelines for EDT. Several other guidelines have been published without consensus guidelines on the ideal candidates for EDT.

Given the circumstances surrounding the procedure and the associated injuries, few patients survive. The poor overall survival rates, however, should not discourage performance of the procedure in the correct setting and when appropriate surgical backup is available for definitive care.

EDT is not a simple procedure (Video 18.1). Identifying specific structures within a chest cavity filled with blood, coupled with a collapsed lung and an injured heart and major vessels, can be formidable. Localizing the injuries that can be reversed quickly and safely is even more difficult.

This chapter focuses on three major objectives: (1) identifying possible indications for and contraindications to EDT, (2) describing the technical aspects of the procedure and adjunctive maneuvers to repair specific injuries, and (3) recognizing the associated risks and complications.

Because of the controversies surrounding EDT, every institution should have guidelines for its appropriate use. An institutional plan for chest wound management and postprocedural care should also be established with the service that will provide backup when members of the surgical team cannot be on site at the time of resuscitation. Debate regarding who should perform EDT is not necessary because everyone who is licensed to perform resuscitative thoracotomy should be trained, competent, and prepared for the technical and initial critical care aspects of patient management. Patient care needs in the event of successful resuscitation should be considered in advance and the surgical and intensive care teams notified so that they can mobilize the appropriate supplies, equipment, and personnel.

INDICATIONS AND CONTRAINDICATIONS

In the ED the vast majority of thoracotomies are performed on penetrating trauma patients in cardiac arrest. Beall and coworkers initially proposed EDT for the treatment of penetrating cardiac injuries in 1966. Since then it has been expanded to include extrathoracic injuries, blunt trauma, and nontraumatic pathology. Studies show wide variation in survival rates and outcomes. Taking 50 years of collective EDT data into account, the survival rate of patients undergoing EDT for blunt trauma is nearly 2%, whereas that for penetrating trauma is nearly 10%, but these survival rates depend on many variables and are not applicable to every situation. There is a paucity of data concerning survival rates in patients with EDT performed for nontraumatic causes, and it is not recommended that this procedure be regularly used in these settings.

Make the decision to perform EDT quickly based on whether the patient is likely to benefit from the procedure, has a reasonable chance of survival, and cannot tolerate a delay in operative intervention. In addition, consider the risks associated with performing the procedure. Trauma researchers have identified several factors that are considered crucial when determining who will benefit from EDT (Box 18.1).

The first assessment is made in the prehospital setting, where determination of the mechanism of injury and the presence or absence of a pulse is critical. Multiple guidelines recommend against EDT in patients without a pulse, blunt traumatic mechanisms, and without signs of life. Such patients do not survive, regardless of the intervention. In one of the largest EDT series to date, Branney and coworkers reviewed 868 consecutive patients over a 23-year period. They found that no blunt trauma patients survived EDT when they had no vital signs in the field, but that 2.5% of blunt trauma patients survived EDT when vital signs were present in the field. In 2000 Rhee and colleagues examined 4620 cases of EDT from 24 studies over a 25-year period. The overall survival rate after blunt trauma was just 1.4%, which led to EDT falling out of favor for this indication. Recent articles have challenged the idea of limiting EDT to those in cardiac arrest from penetrating injury only.

The survival rate of pulseless trauma patients sustaining penetrating injury is significantly higher than that of blunt trauma patients. Several penetrating injury subtypes have been studied as follows: firearm injuries, stab wounds, and penetrating explosive injuries. Thoracic stab wounds consistently show the highest rates of survival after EDT. This is theoretically as a result of the decreased amount of tissue damage related to the weapon and the ability to quickly identify anatomic structures and injuries. Penetrating firearm injuries are more likely to result in death because of increased tissue damage from the missile and concussive surrounding forces. Patients with firearm injuries are more likely to have multiple wounds, and the depth of penetration is increased in comparison to stab wounds. One published cohort of combat casualties from explosive penetrating injuries reported similar survival rates to those after firearm-related penetrating injuries.

The location of the penetrating injury helps determine the futility of EDT. A trend toward increased survival rates in patients with thoracic injuries was found in historical data. Isolated cardiac wounds have the highest survival rate after EDT, with approximately 17% of patients surviving the procedure. Penetrating abdominal injuries have beneficial outcomes when EDT is performed to cross-clamp the aorta, with survival rates in the midteens. Extremity injuries rarely require EDT because the use of a tourniquet can control the hemorrhaging until the patient can be transported to the operating room. When EDT is used for traumatic extremity exsanguination survival rates range from 10% to 25%. Patients in cardiac arrest associated with head injuries, especially those with open cranial wounds, have dismal survival rates and are considered poor candidates for further resuscitative efforts, including EDT.
The type of cardiac electrical activity can be helpful in determining who may benefit from EDT. Battistella and colleagues\(^{26}\) reviewed 604 patients undergoing cardiopulmonary resuscitation (CPR) for traumatic cardiopulmonary arrest and found that of the 212 patients who were in asystole, none survived. Fulton and associates\(^{27}\) found that of patients in traumatic arrest, survival was improved when the patients exhibited ventricular fibrillation, ventricular tachycardia, or pulseless electrical activity rather than asystole or an idioventricular rhythm. In another study of EDT for traumatic arrest, asystole, idioventricular rhythm, or severe bradycardia was indicative of poor outcomes or an unsalvageable patient.\(^{3}\)

The duration of pulselessness prior to EDT has traditionally been used as a decision point. With traumatic injury, survival rates and meaningful neurologic outcomes diminish as the duration of CPR increases. The traditional dogma is that any trauma patient who has undergone CPR for longer than 15 minutes has an exceedingly dismal survival rate and further resuscitation should be considered futile.\(^{3,6,12,13,27-30}\) Guidelines from 2013 have questioned the 15-minute dogma and consensus timing on futility of the procedure remains elusive.\(^{5,31}\) It is recommended that each institution should develop protocols directing futility decision as it pertains to timing of pulseless activity prior to arrival of the patient in the ED.

Perhaps the most critical determinant of the appropriateness of EDT is whether the patient demonstrates “signs of life.” Signs of life are objective physiologic parameters that are present in patients who survive EDT. They include pupillary response, extremity movement, cardiac electrical activity, measurable or palpable blood pressure, spontaneous ventilation, or the presence of a carotid pulse. The presence of one or more of these indicators has been associated with good neurologic outcomes and increased rates of survival.\(^{3-11,32}\)

Although survival remains the ultimate gauge of the effectiveness of EDT, it is also essential to consider quality of life, especially neurologic function of the patient. It is somewhat surprising that survivors of EDT generally have good neurologic outcomes. Rhee and colleagues\(^{33}\) reported 100% sensitivity and specificity in detecting pericardial tamponade.

The progression from compensated cardiac function to uncompensated tamponade can be sudden and profound. Although one may suspect tamponade based on well-described signs, clinical diagnosis of pericardial tamponade in an unstable trauma patient is difficult because of the combined effect of hemorrhagic and cardiogenic shock. The classic signs of Beck’s triad (distended neck veins, hypotension, and muffled heart sounds) described in 1926\(^{46}\) have limited diagnostic value for acute penetrating cardiac trauma.\(^{17}\) The most reliable signs of tamponade are elevated central venous pressure, hypotension, and tachycardia.

The advent of ultrasound and the focused assessment with sonography in trauma (FAST) examination has improved the diagnosis of pericardial effusion and tamponade. Findings indicative of tamponade include the presence of pericardial fluid with right atrial or ventricular collapse during diastole (Fig. 18.1). FAST is a rapid bedside screening examination used to detect hemopericardium and hemoperitoneum and is now important in the evaluation of unstable trauma patients.\(^{38,39}\) From data collected in 1,340 patients, Rozycki and associates\(^{40}\) reported 100% sensitivity and specificity in detecting pericardial and peritoneal fluid in a hypotensive, unstable trauma patient.

**Cardiac Injuries—Penetrating**

Sixty percent to 80% of cardiac stab wounds result in pericardial effusion regardless of the presence of shock.\(^{13}\) Tamponade can occur if the wound is smaller than 1 cm in size, depending on which chamber is involved. Wounds larger than 1 cm usually continue to bleed regardless of which chamber is involved. Low-pressure atrial wounds generally form a thrombus before tamponade develops. The thicker-walled left ventricle may spontaneously seal stab wounds up to 1 cm in length. As little as 60 to 100 mL of blood acutely filling the pericardium will impede diastolic filling, reduce stroke volume, decrease cardiac output, and increase release of catecholamine. Catecholamine release may mask the severity of illness because it maintains blood pressure through an increase in peripheral vascular resistance. In penetrating cardiac injury, the right ventricle is the chamber most likely to be involved because of its anterior location, followed by the left ventricle and the atria.\(^{10,31}\)

![Figure 18.1](image-url)
Ultrasound can have rare false-negative results when pericardial fluid from a cardiac injury decompresses into the thoracic cavity through a wound in the pericardium. A 2015 study by Inaba and colleagues showed a high sensitivity and negative predictive value of FAST exam in determining which patients will not benefit from EDT. In this study the likelihood of survival if pericardial fluid and cardiac motion were both absent was zero.

After EDT for penetrating cardiac wounds, survival is also related to the mechanism of injury. Patients with stab wounds fare better than do patients with gunshot wounds. Rhee and colleagues noted that 16.8% of patients with stab wounds survived to hospital discharge after EDT. Branney and coworkers reported a 29% survival rate in stab wound patients with tamponade, and a 15% survival rate in those without tamponade.

In contrast, gunshot wounds are often large injuries unable to seal themselves; tamponade occurs in only 20%. Patients with penetrating cardiac injuries from gunshot wounds are more likely to initially be seen with profoundly compromised hemodynamics. In addition, the increasing popularity of larger-caliber weapons has made it more difficult to resuscitate patients with gunshot wounds to the chest. Of 112 patients with gunshot wounds to the heart, only 2% survived neurologically intact.

**Cardiac Injuries—Blunt**

Blunt trauma to the heart can range from minor contusion to cardiac rupture. The most common cause of death after nonpenetrating cardiac injuries is myocardial rupture, and in approximately 25% of such patients the ascending aorta is ruptured simultaneously. Branney and coworkers observed a 2% survival rate in blunt trauma patients resuscitated with EDT. Those who survived had vital signs present in the field. The poor outcomes associated with this type of injury are a result of the poor cardiac function caused by myocardial contusion, even if the hemorrhage has been treated.

**Pulmonary Injuries**

Pulmonary injuries can be divided into three types: parenchymal, tracheobronchial, and large vessel. Parenchymal and tracheobronchial injuries rarely require EDT because they are either rapidly fatal or treated initially by tube thoracostomy. Tracheobronchial injury is more common in blunt than in penetrating trauma. Bertelsen and Howitz reviewed 1128 patients at autopsy and found only three to have this injury. The airway is usually maintained, even with complete transection. The stiff tracheobronchial cartilage tends to hold the lumen open, and the paratracheal and parabronchial fasciae preserve the relationship of the proximal to distal bronchi. Ninety percent of tracheobronchial tears occur within 2.5 cm of the carina and most commonly involve the main stem bronchi. Complete division of the trachea is extremely rare. Depending on the size and location of the injury, patients may have massive hemoptysis, airway obstruction, pneumomediastinum, pneumothorax, or tension pneumothorax. Massive subcutaneous emphysema and pneumomediastinum are usually seen, although up to 10% of patients with this injury have no abnormal findings on the initial radiograph. If hemorrhage is profuse or if the site of the injury can be determined, use of a bifid endotracheal tube or unilateral intubation of a main stem bronchus will help secure the airway.

Lacerations of the lung parenchyma that are not accompanied by injury to major vessels generally respond to tube thoracostomy. If the initial chest tube drainage is more than 1500 mL or if there is persistent hypotension or cardiac arrest, consider immediate thoracotomy. For pulmonary injuries, survival after EDT is also related to the mechanism of injury. Branney and coworkers reported a 17% survival rate after pulmonary stab wounds, 3% after gunshot wounds, and 5% after blunt trauma.

**AIR EMBOLISM**

Air embolism is a complication of pulmonary parenchymal injuries that may require immediate thoracotomy if the patient is hemodynamically unstable. The development of air embolism after penetrating injuries of the lung is often insidious, and the diagnosis is usually made at the time of thoracotomy. Preoperative and postmortem diagnosis of air embolism is difficult, and it is likely that most air emboli are not detected. Air embolism is confirmed at thoracotomy by needle aspiration of a foamy air-blood mixture from the left or right ventricle or by visualization of air within the coronary arteries.

Air embolism may appear in either the right or the left side of the circulatory system. Involvement of the right side of the circulation is referred to as venous or pulmonary air embolism. Generally, venous air is well tolerated, but death can occur when the volume of air reaches 5 to 8 mL/kg. The rate at which air moves into the circulation and the body’s position are important determinants of the volume that can be tolerated. If the body’s position allows dispersion of air into the peripheral circulation, more air can be tolerated, although the damage to peripheral structures and end-organs can be extensive. Rapid death usually results from obstruction of the right ventricle or the pulmonary outflow tract. Injuries to the vena cava or the right ventricle can also create portals of entry into the right circulatory system.

Air embolism involving the left side of the circulatory system is referred to as arterial or systemic air embolism. The lethal volume depends on the organs to which it is distributed. As little as 0.5 mL of air in the left anterior descending coronary artery can lead to ventricular fibrillation. Two milliliters of air injected into the cerebral circulation can be fatal. The formation of traumatic bronchovenous fistulas creates potential entry points for air to move into the left side of the circulatory system. The only requirement is the formation of an air-blood gradient conducive to the inward movement of air. Although lowered intravascular pressure from hemorrhage is a risk factor, the most important element in all reports of air embolism has been the use of positive pressure ventilation.

In a review of 447 cases of major thoracic trauma, Yee and coworkers found adequate chart data to suggest the diagnosis of air embolism in 61 patients. Approximately 25% of patients with air embolism have blunt trauma, with associated lung injury secondary to multiple rib fractures or hilar disruption. The overall mortality is higher than 50%.

The diagnosis of air embolism is easily overlooked because the signs and symptoms are similar to those of hypovolemic shock. Two valuable signs that are present in 36% of patients are hemoptysis and the occurrence of cardiac arrest after intubation and ventilation. The development of focal neurologic changes, seizures, or central nervous system dysfunction in the absence of head injury is also suggestive of the diagnosis.

Overall, the diagnosis is subtle and must be considered when
there is no evidence of the more common causes of extremis in a trauma patient.

Blunt and Penetrating Abdominal Injury

In the setting of penetrating abdominal injury, thoracotomy with cross-clamping of the thoracic aorta has been advocated as a means of controlling hemorrhage, redistributing blood flow to the brain and heart, and reducing blood loss below the diaphragm. Unfortunately, aortic cross-clamping can also have detrimental effects. Kraulovich and colleagues studied the hemodynamic consequences of aortic occlusion in a swine model of hemorrhagic arrest. There was no difference between groups in return of spontaneous circulation; however, the occluded aorta group experienced statistically greater impairments in left ventricular function and systemic oxygen utilization in the postresuscitation period. Branney and coworkers found that 8 of 76 patients undergoing EDT for penetrative abdominal injury survived neurologically intact. More recently, Seamon and colleagues achieved a 16% survival rate with good neurologic outcomes (8 of 50 patients) when EDT was used before laparotomy for abdominal exsanguination from trauma. Of note, none of the survivors in this study were in cardiac arrest at the time of EDT, but they did have severe hemorrhagic shock, and six of the eight had unmeasurable blood pressure. The resurgence of resuscitative endovascular balloon occlusion of the aorta (REBOA) has changed many institutional trauma algorithms dealing with exsanguinating abdominal and pelvic hemorrhage. Current recommendations suggest that EDT be performed judiciously in patients with abdominal trauma as an adjunct to definitive repair of the abdominal injury.

Open-Chest Resuscitation for Nontraumatic Arrest

At present, less than 6% of CPR attempts conducted outside hospital special care units result in survival. The first case of a human survivor of open-chest cardiac massage (OCCM) was reported in 1901. In 1960, Kouwenhoven and associates published favorable survival rates with closed-chest CPR as opposed to OCCM. After further refinement by Pearson and Redding, closed-chest CPR gradually became the preferred method of cardiac compression.

The goal of CPR is to restore coronary perfusion pressure (CPP), which is the prime determinant for return of spontaneous circulation as established in animal models. Paradis and associates found that humans need a minimal CPP of 15 mm Hg to achieve return of spontaneous circulation. Although a CPP of 15 mm Hg does not guarantee return of spontaneous circulation, there is 100% failure of resuscitation if CPP is below this level. Despite the limited number of human studies on OCCM, its hemodynamic superiority over closed-chest CPR is compelling. Del Guercio and coworkers measured cardiac output during both closed-chest CPR and OCCM in in-hospital cardiac arrest patients. OCCM produced a mean cardiac index of 1.31 L/min per m² as opposed to 0.6 L/min per m² during closed-chest CPR. Boezaar and colleagues further examined 10 patients who were unresponsive to closed-chest CPR and measured CPP during closed-chest CPR followed by OCCM. Mean CPP in the closed-chest group was 7.3 mm Hg versus 32.6 mm Hg in the open-chest group. All patients achieved a CPP of at least 20 mm Hg at some time during their OCCM phase. This easily surpassed the minimal CPP required for return of spontaneous circulation. Outcomes after OCCM have not been well established. Animal models suggest not only improved hemodynamic parameters, but also a possible increase in 24-hour survival rates. However, neurologic outcomes are unknown, and the American Heart Association guidelines for CPR do not promote the regular use of OCCM in patients with out-of-hospital cardiac arrest.

At present, the precise indications for open-chest resuscitation after nontraumatic arrest are not well defined, and the procedure is not considered the standard of care. Despite demonstrated hemodynamic superiority in both animal and human models of open-chest versus closed-chest CPR, outcome benefit is lacking. There is a paucity of human data evaluating the window of time during which this treatment can be effective.

Nontraumatic Hypothermic Cardiac Arrest

In the setting of cardiac arrest from hypothermia, one can consider the use of EDT and OCCM. Cardiopulmonary bypass (CPB) or extracorporeal membrane oxygenation are the most rapid methods of core rewarming, but these methods are sometimes not available immediately. Open thoracotomy with mediastinal irrigation has been used successfully in cases of severe hypothermia with cardiac arrest. Brunette and McVaney reported 11 patients with hypothermic cardiac arrest, 7 of whom underwent EDT with OCCM and mediastinal rewarming. Five patients survived, and all had positive neurologic outcomes despite cardiac arrest times of between 10 and 90 minutes (although one patient died of gastrointestinal hemorrhage and sepsis following resuscitation; the other four patients survived with full neurologic recovery). The other four patients who did not undergo EDT did not survive despite being taken promptly to the operating room for CPB rewarming. Although the number of cases is limited, this study is evidence that OCCM can provide prolonged hemodynamic support and good neurologic outcomes. It should be noted that similar case reports also exist in which closed-chest CPR was maintained for prolonged periods and resulted in successful hypothermic resuscitation. EDT with mediastinal irrigation can produce core rewarming rates as fast as 8°C/hr, with the heart and lungs preferentially being rewarmed first. Mediastinal irrigation involves heating sterile saline in a microwave oven to 40°C and then pouring it slowly over the heart and into the thorax. Performing a thoracotomy for hypothermic arrest does not preclude the use of CPB inasmuch as it was subsequently used after EDT in three of the survivors from the Brunette and McVaney study.

EQUIPMENT

Carefully select the instruments to be included in the EDT equipment tray. See Review Box 18.1 for a complete list.

PROCEDURE

Preliminary Considerations

All trauma patients arriving at the ED with hypotension must be assumed to be hypovolemic and should be treated accordingly. Rapidly exclude other causes as well, including tension
pneumothorax, cardiac tamponade, air embolism, and neurogenic or cardiogenic shock. A useful algorithmic overview of the approach to chest trauma is presented in Fig. 18.2. The use of autotransfusion, if available, has several benefits, but its use is not widespread or considered standard.\(^6\)

**Airway Control**

Intubate the patient orotracheally, if possible, but be aware that access to the thoracic organs, surgical repairs, or surgical procedures may be hampered by frequent inflations of the left lung. If necessary, selectively intubate the right lung by blindly advancing a standard single-lumen endotracheal tube to a depth of 30 cm (measured from the corner of the mouth) in adult patients.\(^6\) Although the left lung and the right upper lobe are not ventilated with the tracheal tube in this position, both animal studies and data from humans suggest that selective right lung ventilation provides adequate oxygenation and ventilation for at least 60 minutes.\(^6\) With the left lung deflated one can expedite thoracotomy by maximizing space in the left thoracic cavity. Keep in mind that extending the thoracotomy into the right thoracic cavity may necessitate switching to bilateral lung ventilation or left lung ventilation to allow maximum right thoracic exposure.

**Anesthesia and Amnesia**

Comatose patients undergoing resuscitation may regain consciousness during successful EDT, but this awareness may not be apparent if they are still pharmacologically paralyzed. Anticipate and recognize this phenomenon and administer adequate analgesic, amnesic, and muscle-relaxing agents to a ventilated patient who may also be in shock. No specific regimen has been studied, but ketamine appears to be an ideal agent to use in the ED. It is prudent to administer anesthetic agents routinely if a paralyzed patient demonstrates perfusion during resuscitation. This is not only humane but decreases systemic oxygen consumption.

**Anterolateral Thoracotomy Incision**

Manually ventilate the patient during the procedure. Ask an assistant to pass a nasogastric tube, which helps differentiate the esophagus from the aorta, but do not allow this procedure to delay thoracotomy. If CPR is being performed, ask an assistant to continue closed-chest compressions up to the point of making the initial incision. Take universal precautions to avoid blood exposure and use a suction catheter to minimize contact with blood. Sterile gloves (consider double-glove protection), gown, mask with an eye shield, and an operative surgical cap are recommended for the procedure.

Prepare the skin of the left anterior aspect of the chest with antiseptic if readily available. Wedge towels or sheets under the left posterior part of the chest and place the patient's left arm above the head (Fig. 18.3). On the left side of the chest, make an anterolateral incision at the fourth to fifth intercostal space with a No. 20 blade scalpel (Fig. 18.4, step 1). Do not take the time to count ribs; simply estimate the location to be just beneath the nipple in males and at the inframammary fold in females (see Fig. 18.3). It is important to establish wide exposure by beginning the incision on the right side of the sternum and extending the skin incision past the posterior axillary line. Beginning here will save time if the right side of the chest needs to be opened as well. Inadequate exposure, rib fractures, and additional delays occur when the skin incision is too limited. With the first sweep of the scalpel, separate skin, subcutaneous fat, and the superficial portions of the pectoralis and serratus muscles.

Cut the intercostal muscles with scissors to expose the thoracic cavity (see Fig. 18.4, step 2). Use scissors to divide the intercostal muscles because the risk for lung laceration is greater when a scalpel is used. Make the incision just over the top of the rib to avoid the intercostal neurovascular bundle. Just before opening the pleura, stop ventilations momentarily. This will allow the lung to collapse away from the chest wall. Should the internal mammary artery be transected during the procedure, hemorrhage is generally minimal until after perfusion is reestablished, at which time bleeding may be profuse. If actively bleeding, the internal mammary artery should be ligated or clamped. Do not forget to address the internal mammary artery if perfusion is reestablished because this can be a source of significant bleeding.\(^6\)
Resuscitative Thoracotomy General Technique

1. Make an anterolateral incision at the 4th to 5th intercostal space.
   - Begin at the right side of the sternum and extend the incision past the posterior axillary line.

2. Cut the intercostal muscles with scissors.
   - Incise along the top of the rib to avoid the intercostal artery.

3. Use scissors to incise the parietal pleura and gain entry into the thoracic cavity.

4. Use your hands to spread the ribs.

5. Place a rib spreader between the ribs with the handle and ratchet bar facing downward.
   - Carefully spread the ribs open.

6. PERICARDIOTOMY
   - Lift the pericardial sac with forceps, and cut pericardium with scissors.
   - Incise in a caudal-to-cephalad direction; stay anterior and parallel to the phrenic nerve.

7. AORTIC CROSS-CLAMPING
   - Bluntly dissect the surrounding fascia and temporarily apply an aortic clamp.

Additional injury-specific procedures are depicted elsewhere in this chapter.

Figure 18.3 Left anterolateral thoracotomy. A, Place several towels or sandbags under the left scapula and raise the arm above the head. The patient should be intubated. Insert a nasogastric (NG) tube to differentiate the esophagus from the aorta. Make a left anterolateral submammary incision. B, Dashes indicate the incision site in the inframammary fold in women. ET, Endotracheal.

Figure 18.4 Resuscitative thoracotomy, general technique. (From Custalow CB: Color atlas of emergency department procedures, Philadelphia, 2005, Saunders.)
After entering the pleural cavity, gain good exposure. Karmy-Jones and associates found that 20% of their patients undergoing EDT via the anterolateral approach needed to have the initial incision extended. To gain better exposure first use your hands to open the chest cavity. Then place a Finocchietto retractor (rib spreader) between the ribs with the handle and the ratchet bar directed downward toward the axilla (see Fig. 18.4, step 5). If the retractor were to be placed with the handle up, the ratchet bar would prevent extension of the incision into the right side of the chest. Ribs may be broken during spreading, so be careful to not get cut on the sharp bone edges. If massive hemothorax is encountered, remove the clots manually, suction out the blood, and use towels to absorb any blood spilling from the chest. If the site of injury is to the right of the heart and cannot be reached, extend the incision into the right side of the chest with a Gigli saw, Liebsche knife, sternal osteotome, or standard trauma shears. Before performing EDT it would be prudent to familiarize oneself with the instruments included in the EDT tray, but do not take the time for this during an actual case.

**Pericardiotomy**

For a patient in cardiac arrest, if there is no other obvious injury in the chest and the myocardium cannot be visualized, open the pericardium. It may be difficult to definitively rule out pericardial tamponade by visual inspection alone. If in doubt, use forceps to elevate a portion of pericardium and carefully incise it to assess for hemopericardium.

If you are confident that there is no pericardial tamponade, leave the pericardial sac closed while you address other life-threatening injuries. Opening the pericardium increases the risk for complications such as delay in the onset of cardiac compressions, damage to the myocardium or coronary vessels, or cutting of the left phrenic nerve. Previous pericardial disease may also have caused adhesions, and if you attempt to separate these adhesions rapidly, you can tear the atria or right ventricular wall. The incidence of traumatic rupture of the atria or the right ventricle during massage is greater when the pericardium is open. Furthermore, with an intact pericardium pressure is distributed over a larger area, and the pericardial fluid seldom allows compressing fingers to remain in one spot for a prolonged period.

Perform pericardiotomy if tamponade is present or suspected. This procedure is performed anterior and parallel to the left phrenic nerve (see Fig. 18.4, step 6). Begin the incision near the diaphragm to avoid injury to the coronary arteries. Lift the pericardial sac with toothed forceps, and use scissors to make a small hole. Extend the incision with scissors in a cephalad direction along the anterior aspect of the pericardium. Extend the incision so that it reaches from the apex of the heart to the root of the aorta. When the pericardium is under tension, it may be difficult to grasp the pericardium with forceps. In this case, use sharp, straight Mayo scissors to divide the pericardium by layers. If the heart is in arrest, speed is important, so use sharp scissors to “catch” the pericardium and start the pericardiotomy. To do this procedure, hold the point of the scissors almost parallel to the surface of the heart and use enough pressure to create a wrinkle in the pericardium to puncture it as the scissors move forward. Apply moderate pressure to puncture the fibrous pericardium. Be cautious because if the point of the scissors is unnecessarily angled toward the heart, the sudden “give” that occurs when you open the pericardium may result in laceration of the myocardium. Remove clots of blood from the pericardial sac with a sweeping motion using your gloved hand, sterile lap sponges, or gauze pads. If cardiac repair or cardiac compressions are needed, deliver the heart from the pericardial sac. To do this, place your right hand through the pericardial incision and encircle the heart, pull it into the left side of the chest, and place the pericardial sac behind the heart.

**Internal Cardiac Defibrillation**

It is not uncommon to encounter cardiac dysrhythmias during EDT resuscitative efforts. Recommendations for internal defibrillation are the same as those for external defibrillation: ventricular fibrillation and tachycardia without pulses are immediate indications for defibrillation. With an open chest, internal defibrillation is the procedure of choice.

To perform internal defibrillation, place the internal paddles on the anterior and posterior aspects of the heart. The current is delivered through the circular tips of the paddles onto the surface of the heart. There is decreased electrical impedance with direct myocardial contact, and as a result less energy is needed than with standard defibrillation (typically 10 to 50 J). If internal paddles are not available, perform defibrillation in the usual manner with external paddles or electrodes.

**Direct Cardiac Compressions**

Three techniques for cardiac compression have been described: one-handed compression, one-handed sternal compression, and two-handed (bimanual) compression (Fig. 18.5). To perform the one-handed compression method, place your thumb over the left ventricle, the opposing fingers over the right ventricle,
and your palm over the apex of the heart. To perform one-handed sternal compression, hold your fingers flat. Keep the fingers tightly together to form a flat surface over the left ventricle and compress the heart up against the sternum with your fingers. To perform two-handed compression, cup the left hand and place it over the right ventricle. Hold the fingers of the right hand tightly together to form a flat surface supporting the left ventricle. Push the flat surface of your right-hand fingers to compress the heart against the cupped surface of the left hand. The bimanual technique has been shown to be consistently superior.64

A difference of opinion exists regarding the optimal rate at which the heart should be compressed. Some recommend a rate of 50 to 60 compressions/min; however, no physiologic data support such a recommendation. American Heart Association guidelines for closed-chest CPR recommend a rate of at least 100 compressions/min.51 It is conceivable that this would also apply to OCCM.

It is important to remember the following points while performing cardiac compression:

1. Use the entire palmar surface of the fingers. Avoid fingertip pressure.
2. For each method, adjust the force of compression so that it is perpendicular to the plane of the septum. The anterior descending coronary artery is located over the interven- tricular septum, making this a helpful landmark to orient your hand properly.
3. Position your fingers so that the coronary arteries will not be occluded.
4. Venous filling of the heart is especially sensitive to changes in position. Maintain a relatively normal anatomic position of the heart to prevent kinking of the vena cava and pulmonary veins. Do not angle the heart more than 30 degrees into the left side of the chest.
5. It is essential to completely relax the heart between compressions to allow it to refill completely.

**Control of Hemorrhagic Cardiac Wounds**

To partially control active bleeding from a ventricular wound, place one finger over the wound and use the other hand to stabilize the beating heart. This maneuver buys time while you begin to repair the injury and continue with resuscitation. If the heart is not beating, you may perform closure of the injury before resuscitation and defibrillation, although this is controversial. If you elect to repair the injury before attempting to restart the heart, perform intermittent cardiac massage. Surgical staples can be used to close a ventricular wound and are an extremely rapid method for controlling hemorrhage (Fig. 18.6).65 This technique is particularly useful with large or multiple lacerations. Another advantage is that stapling does not expose the operator to the risk associated with needlestick. Macho and coworkers66 reported a 93% success rate in temporarily controlling hemorrhage in 28 patients (33 lacerations) with penetrating injuries to both the atria and ventricles by using a standard skin stapler with wide (6-mm) staples placed at 5-mm intervals (Auto-Suture 35W, US Surgical Corp., Norwalk, CT). The rotating long neck of the Ethicon Proximate Quantum Skin Stapler (Model PQW-35, Ethicon, Inc., Somerville, NJ) is helpful in obtaining proper orientation of the staples during placement. The staples can be left in place and reinforced or replaced on further exploration in the operating room.

Alternatively, the wound may be repaired by suturing: 1. Using appropriately sized suture, obtaining a generous “bite” of the wound edges and reduce bleeding. Use a skin stapler with wide (6-mm) staples and place them 5 mm apart. This technique may be used for atrial and ventricular lacerations. After stabilizing the patient’s condition, revise the wound in the operating room.

If exsanguinating hemorrhage is not controlled by the aforementioned methods, temporarily occlude inflow to the heart. Apply inflow occlusion intermittently for 60 to 90 seconds. During occlusion, the heart shrinks, hemorrhage is controlled, and you can place sutures in a decompressed injury. Two techniques that are useful are vascular clamping of the superior and inferior vena cava for partial inflow occlusion,69 and the Sauerbruch grip (Fig. 18.9).70 The Sauerbruch grip can be performed quickly with the added advantage of cradling and stabilizing the heart while you repair the wounds over either the ventricle or the left atrium. It involves occlusion of the
gently traction (Fig. 18.10). Apply enough traction to slow the bleeding and provide an acceptable level to visualize and repair the wound. Excessive traction can pull the catheter out and enlarge the wound. The balloon will effectively occlude the wound internally. Use a purse-string suture to close the wound. When repairing the wound be careful with the suture needle because it can easily rupture the balloon. Temporarily pushing the balloon into the ventricular lumen during passage of the needle can be done to avoid this complication. Use normal saline when inflating the balloon because using air could result in air embolism if the suture needle ruptures the balloon.

Foley catheters have several advantages over other methods for controlling cardiac wounds. With the digital method, your fingertip will often slip if there is a strong heartbeat, you cannot visualize the wound during repair, and digital pressure significantly interferes with cardiac massage. Intermittent total venous inflow occlusion is an effective method of controlling bleeding and decompressing the heart, but such control will be at the expense of poor cardiac output. Attempting to elevate the heart for control and repair of posterior cardiac wounds often results in cardiac arrest by reducing both venous and arterial flow. With posterior injuries use of a Foley catheter does not require continued viewing after initial placement. If bleeding can be controlled, repairs in this location should await full-volume expansion or cardiopulmonary bypass. Regardless of location the most valuable feature of the Foley catheter is that you can control hemorrhage without interfering with cardiac compression. The catheter can also be used for infusion of fluids (see Fig. 18.10B).

To initially manage wounds of the atria, use partial-occlusion clamps (Fig. 18.11). Because of the thin structure and instability of the atrial wall, you will not be able to effectively stop bleeding with digital pressure. Injuries near the caval-atrial junction are not amendable to clamping; in this location, use a Foley catheter to tamponade the wound. Be careful not to obstruct atrial filling with the inflated balloon. Skin staples may also be used for closure of atrial wounds.

Wounds of the septa, valves, and coronary arteries require definitive repair in the operating suite. Hemorrhage from a coronary artery can generally be controlled with digital pressure. Avoid ligation of a coronary artery whenever possible.
Figure 18.10 A, Serial illustration. Gentle traction on an inflated Foley catheter may control hemorrhage and allow repair. Inflate the balloon with saline while taking care to not rupture the balloon with the suture needle. This technique is particularly useful for injuries to the inferior cavoatrial junction, for posterior wounds, and during cardiac massage. Volume loading can be achieved by infusion of blood or crystalloid solution through the lumen of the catheter. Take care to avoid an air embolus through the lumen of the catheter during placement. B, Foley catheter in an atrial stab wound. Keep gentle traction on an inflated balloon and inject saline directly into the heart via the catheter. Note how difficult it is to identify structures in the chest cavity. The heart is collapsed and not beating, which makes even this organ hard to find. The aorta could not be isolated before stopping resuscitation efforts.
occlusion of the descending thoracic aorta. This maneuver can maintain myocardial and cerebral perfusion (Fig. 18.13; see also Fig. 18.4, step 7), although Kravolich and colleagues suggested that the benefit of significant improvement in CPP may be overstated. When the aorta has been injured by blunt trauma, selective clamping is necessary (Fig. 18.14). Aortic occlusion has a limited role in controlling hemorrhage below the diaphragm. In patients with a tense abdomen and massive hemorrhage from the subclavian artery, use laparotomy pads for compression in the apex of the pleura from below and the supraclavicular fossa from above (Fig. 18.12) to prevent further bleeding as the patient is stabilized and moved to the operating suite.

Aortic Cross-Clamping

For persistent hypotension (systolic blood pressure <70 mm Hg) after thoracotomy and pericardiotomy, perform temporary occlusion of the descending thoracic aorta. This maneuver can maintain myocardial and cerebral perfusion (Fig. 18.13; see also Fig. 18.4, step 7), although Kravolich and colleagues suggested that the benefit of significant improvement in CPP may be overstated. When the aorta has been injured by blunt trauma, selective clamping is necessary (Fig. 18.14). Aortic occlusion has a limited role in controlling hemorrhage below the diaphragm. In patients with a tense abdomen and massive hemorrhage from the subclavian artery, use laparotomy pads for compression in the apex of the pleura from below and the supraclavicular fossa from above (Fig. 18.12) to prevent further bleeding as the patient is stabilized and moved to the operating suite.

Control of Hemorrhagic Great-Vessel Wounds

Wounds of the great vessels can be controlled with digital pressure or partial-occlusion clamps. If desired, close small aortic wounds with 3-0 Prolene suture. To prevent exsanguinating hemorrhage from the left subclavian artery, try to cross-clamp the intrathoracic portion of the artery. Cross-clamping of the right subclavian artery is very difficult. For injuries to this vessel, use laparotomy pads for compression in the apex of the pleura from below and the supraclavicular fossa from above (Fig. 18.12) to prevent further bleeding as the patient is stabilized and moved to the operating suite.

Aortic Cross-Clamping

For persistent hypotension (systolic blood pressure <70 mm Hg) after thoracotomy and pericardiotomy, perform temporary occlusion of the descending thoracic aorta. This maneuver can maintain myocardial and cerebral perfusion (Fig. 18.13; see also Fig. 18.4, step 7), although Kravolich and colleagues suggested that the benefit of significant improvement in CPP may be overstated. When the aorta has been injured by blunt trauma, selective clamping is necessary (Fig. 18.14). Aortic occlusion has a limited role in controlling hemorrhage below the diaphragm. In patients with a tense abdomen and massive hemorrhage from the subclavian artery, use laparotomy pads for compression in the apex of the pleura from below and the supraclavicular fossa from above (Fig. 18.12) to prevent further bleeding as the patient is stabilized and moved to the operating suite.
SECTION III CARDIAC PROCEDURES

Given the need for timely intervention, the simplest and most desirable approach to aortic occlusion is to have an assistant digitally compress it or use the aortic tamponade instrument (Fig. 18.16). The aortic tamponade instrument, however, may be applied blindly to the vertebral column and permits safe, quick, and complete aortic occlusion. This technique may be most prudent when isolation of the aorta is difficult. The instrument's unique shape allows it to remain in place and to provide atraumatic occlusion with little interference in the hemoperitoneum, aortic cross-clamping is beneficial when applied just before laparotomy.24

The aorta can be very difficult to identify in an ED, especially when collapsed as a result of exsanguination. It is situated immediately anterior to the vertebrae and actually lies on the vertebral bodies themselves. The esophagus lies anterior and slightly medial to the aorta. Passing a nasogastric tube from above may help in identifying the esophagus. To expose the descending aorta, ask an assistant to retract the left lung in a superomedial direction. To achieve adequate exposure, it is sometimes necessary to divide the inferior pulmonary ligament (be careful not to injure the inferior pulmonary vein). Identify the aorta by advancing the fingers of the left hand along the thoracic cage toward the vertebral column. Open the mediastinal pleura and bluntly dissect the aorta away from the esophagus anteriorly and the prevertebral fascia posteriorly before clamping. To locate the aorta, use a DeBakey aortic clamp or a curved Kelly clamp for blunt dissection, and spread the pleura open above and below the aorta (Fig. 18.15). Alternatively, if excessive hemorrhage limits direct visualization, bluntly dissect with the thumb and fingertips. Separate the aorta from the esophagus, which lies medially and slightly anteriorly. It may be difficult to separate the esophagus from the aorta by feel in a hypotensive or cardiac arrest situation. When the aorta is completely isolated, use the index finger of the left hand to flex around the vessel and apply a large Satinsky or DeBakey vascular clamp with the right hand. Check brachial blood pressure immediately after the occlusion. If systolic pressure is higher than 120 mm Hg, slowly release the clamp and adjust it to maintain a systolic pressure of less than 120 mm Hg.25

Figure 18.15 A, Identification of the aorta is very difficult during emergency department thoracotomy. If possible, first pass a nasogastric tube to help identify the esophagus. The aorta is in the posterior mediastinum, directly anterior to the vertebral bodies. The esophagus is anterior and slightly medial to the aorta. In the lower part of the thorax, both are covered on the anterolateral surface by the mediastinal pleura, which must be dissected before isolating the aorta for cross-clamping. B, Aortic cross-clamping. Using blunt dissection, spread the pleura above and below the aorta. Fully mobilize the vessel and clearly separate the esophagus before clamping. The aorta is the more posterior structure and is in contact with the vertebral bodies.

Figure 18.16 Use of a Conn aortic compressor is an excellent method for aortic occlusion because it is fast, does not interfere with the operative field, and is associated with minimal risk for injury. Alternatively, direct digital occlusion can be used, but this technique is more awkward.
operative field when compared with digital compression. The degree of occlusion can be varied by the amount of pressure exerted by the operator.

Potential complications of aortic cross-clamping include ischemia of the spinal cord, liver, bowel, and kidneys. In addition, iatrogenic injury to the aorta and the esophagus may occur. Fortunately, these complications are infrequent. The metabolic penalty of aortic cross-clamping becomes exponential when occlusion time exceeds 30 minutes. Whenever possible, unclamp the aorta for 30 to 60 seconds every 10 minutes to increase distal perfusion. Perform final release of the aorta gradually.

Management of Air Embolism

In those at risk for air embolism spontaneous ventilation is preferred. It is essential that the source of the air embolism be controlled rapidly. Place the patient immediately in the Trendelenburg (head-down) position to minimize cerebral involvement and direct the air emboli to less critical organs. If the chest injury is unilateral, consider isolating the injured lung by selectively intubating the contralateral lung. If this is unsuccessful, perform a left anterolateral thoracotomy. Flood the exposed thorax with sterile saline and look for bloody froth created during positive pressure ventilation to identify peripheral bronchovenous fistulas. Carry out a quick search for hiliar injuries. If the source of the air embolism is not readily apparent, perform a contralateral thoracotomy. Once the bronchovenous communication is controlled, use a needle to aspirate the residual air that commonly remains in the left ventricle and the aorta. If the patient is hypotensive, consider cross-clamping the aorta. Be aware, though, that cross-clamping the aorta before controlling bronchovenous fistulas and removing residual air may result in further dissemination of air to the heart and brain.

Air emboli traverse capillary beds if the blood pressure is high enough. After controlling the bronchovenous fistula, produce a brief period of proximal aortic hypertension by cross-clamping the descending aorta. Maintain systemic arterial pressure with adequate fluid resuscitation. Vasopressors such as dopamine, epinephrine, or norepinephrine may be required to increase systemic pressure and facilitate the passage of air bubbles from left to right.

Maintain left atrial pressure at a high level. Keep the ventilator inspiratory pressure as low as possible, and use 100% oxygen to facilitate diffusion of nitrogen from emboli. Consider high-frequency ventilation, which allows the use of small volumes and has been used successfully in individual patients. The most important adjunctive therapy is hyperbaric oxygen. Although it is best to begin treatment within 6 hours of the traumatic insult, there are cases of success and improvement when hyperbaric oxygen has been started even 36 hours after injury.

INTERPRETATION AND HEMODYNAMIC MONITORING

Use systolic blood pressure after the first 30 minutes of resuscitation as a decision point for further treatment. A report of EDT for blunt and penetrating trauma demonstrated a relationship between blood pressure at 30 minutes and the eventual outcome. Of the 146 cases reviewed, 45 patients (31%) were transferred to the operating room after initial resuscitation and aortic cross-clamping when necessary. For patients who survived with full neurologic recovery, the average systolic blood pressure after the first 30 minutes of resuscitation was 110 mm Hg. In those who were long-term survivors but had significant brain damage, the average systolic blood pressure was 85 mm Hg. No survivals were recorded when mean systolic blood pressure was lower than 70 mm Hg. Transfer of these patients to the operating room for definitive repair of these mortal wounds would be futile.

EDT IN CHILDREN

Trauma is the leading cause of death and morbidity in children older than 1 year. Just as with adults, improved transportation of injured children to the hospital has resulted in the survival of more patients who would have been pronounced dead at the scene. Although the role of EDT has been reviewed extensively in the adult population, experience and data in the pediatric population are limited. Overall survival rates in children undergoing EDT are less than that of the adult population with one study showing an overall survival of 3% to 6%. This is likely as a result of a higher proportion of blunt trauma in the pediatric population. Anatomic injury patterns also showed a higher proportion of concomitant head injuries in the pediatric population. Rates of survival below age 15 are very poor. Despite poor survival rates, the general consensus is that EDT is indicated for traumatic pediatric patients with similar indications discussed earlier in this chapter.

COMPLICATIONS

A variety of significant complications occur in patients surviving EDT, but most of them are related to the primary injury rather than the thoracotomy. Techniques to avoid iatrogenic complications include noting the position of the left phrenic nerve and coronary arteries during EDT. Surprisingly, serious infections are uncommon. Patients receiving antibiotics during or immediately after the procedure have a low rate of infectious complications. Standard antibiotic regimens targeting skin flora are recommended, but should not delay the procedure. As a general rule, excessive attention to antiseptic skin preparation should be avoided because it may delay performance of the procedure.

Another potentially serious complication of EDT is injury or transmission of disease to health care workers. In an emotionally charged environment in which many clinicians are attempting to perform a lifesaving procedure under difficult conditions, it is easy for a needle, scalpel, or scissors to cause injury. Sharp ribs can cut clinicians quite easily (Fig. 18.17). The seroprevalence of human immunodeficiency virus (HIV) in US EDs is estimated to range from 2% to 6% or 9% in urban areas. Tardiff and colleagues noted an HIV-positive rate of 7.2% in a study of trauma patients taken to their urban ED. Occupational exposure to both hepatitis B and hepatitis C virus is also of concern to health care workers. Sloan and associates found a 3.1% incidence of hepatitis B in trauma patients brought to their inner-city ED. Hepatitis C is now the most common viral hepatitis seen in health care workers since the advent of the hepatitis B vaccine. Approximately 2200 health care workers per year seroconvert after occupational exposure. The risk for
Seroconversion after occupational exposure to HIV is estimated to be 0.3% for needlestick and as high as 4.5% for deep injury. Hepatitis C seroconversion rates after occupational exposure are estimated to be between 1.8% and 10%. Keep the risk for exposure to staff in mind, and rigorously follow universal precautions. Although EDT must be performed rapidly to be of value to the patient, excessive haste is not warranted if it threatens the health of members of the medical team.

**ACKNOWLEDGMENT**

The editors and authors wish to acknowledge the contributions of Robert L. Bartlett, MD, and Michael E. Boczar, MD to this chapter in previous editions.

**Figure 18.17** Note the sharp edges of this rib (indicated by the circled clamp), which was fractured during unsuccessful emergency department thoracotomy. The incidence of blood-borne diseases (e.g., human immunodeficiency virus, hepatitis) in trauma patients may be high and transmission to a health care worker may occur if sharp bone fragments puncture the gloves and skin.
REFERENCES

Obtaining vascular access and blood samples in an infant or child can challenge and frustrate even the most skilled emergency clinician and can be especially challenging in children who are dehydrated or in shock. Resuscitation of critically ill and injured children should not be delayed for lack of vascular access, and intraosseous (IO) access is the preferred technique if peripheral vascular access cannot be secured rapidly. This chapter reviews the basic principles and techniques of blood sampling and placement of peripheral and central intravenous (IV) and intraarterial catheters in infants and children, including the use of umbilical catheters in neonates. Hydration techniques for dehydrated children are also reviewed. Though very rarely required, emergency cutdown is occasionally useful in obtaining vascular access, and a section of this chapter is devoted to cutdown techniques.

PATIENT PREPARATION AND RESTRAINT

Fear and anticipation of procedural pain can make the hospital experience traumatic for children. Before beginning any painful procedure in a stable child, explain the procedure to the parents, as well as the reasons that it needs to be done. For children capable of understanding, explain the procedure in developmentally appropriate language before starting and before each successive step. Parents should generally be present during emergency department (ED) procedures but should not assist with painful procedures. Parental presence is generally comforting to children, and children with a parent present have been shown to demonstrate less stress during procedures. Nonpharmacologic techniques such as music, toys, or being held by a parent may further reduce children’s distress during attempts at vascular access. Many applications for smart phones, including some that produce “bubbles” for the child to “pop” on the screen are excellent and typically available distractions.

The success of blood sampling or obtaining vascular access depends in part on proper positioning and restraint of the patient. In most cases this requires the assistance of at least one other staff person and restraint of the extremity a joint above and below the intended insertion site. A significant amount of time may be required to perform venipuncture or vessel cannulation in neonates or young infants. Consequently, they may become hypothermic if disrobed and exposed for a prolonged period, especially if perfusion is compromised due to sepsis or hypovolemic shock. Use overhead lights, warm blankets, or other warming modalities to prevent accidental hypothermia in vulnerable patients. Furthermore, there is some evidence that radiant warmth is effective as an analgesic for minor procedures in newborns.

Anesthesia

Many products are available to decrease the pain associated with vascular access. Do not delay IV access in critically ill or injured children to use these medications or devices, but consider using them in stable patients. These medications and products are discussed in more detail in Chapter 29. Options include vapocoolants, topical heat-enhanced anesthetic delivery such as lidocaine-prilocaine, 4% liposomal lidocaine, and injection of lidocaine via a needleless jet injector (J-tip), and vibrating devices. Orally administered sucrose solution has been demonstrated to decrease the pain response in young infants during procedures.

Procedural sedation is commonly used during central venous and arterial cannulation in children (Chapter 33).

BLOOD SAMPLING TECHNIQUES

Capillary Blood Sampling

Indications and Contraindications

Capillary blood sampling is frequently used to obtain blood samples from young infants. In infants, the heel is the most common location for capillary blood sampling, whereas in older children and adults, blood samples are more commonly obtained from the finger tip. This technique is useful when repeated measurements such as blood glucose or serial hemoglobin are needed. It is also an option for obtaining “arterialized” blood for blood gas analysis. If a sufficient volume is obtained, blood from a capillary sample can also be sent for other routine laboratory studies. Heel sticks are more painful than venipuncture but are useful in the event of difficult access or when arterialized samples are needed. Avoid sampling from an area of local inflammation or hematoma. Avoid repetitive sampling from the same site because it may induce inflammation and subsequent scarring. In general, heel stick sampling is not ideal for blood gas analysis when the infant is hypotensive, the heel is markedly bruised, or there is evidence of peripheral vasoconstriction.
Equipment and Setup

Use a 3-mm lancet (Becton-Dickinson, Rutherford, NJ) or an automated disposable incision device (e.g., Tenderfoot, Tenderfoot, Accrivia Diagnostics, San Diego, CA) to perform this procedure. Perform blood collection with either heparinized capillary tubes or 1-mL Microtainer tubes with a collector attachment (Becton-Dickinson).

Technique

The heel stick method of capillary blood sampling will be described, but capillary blood samples can also be obtained from a finger, toe, or earlobe (Fig. 19.1). Traditional teaching is to puncture only the most medial and lateral portions of the plantar surface of the heel to avoid puncture of the calcaneus. However, newer research has found that although soft tissue thickness is greater on the medial and lateral portions of the heel, if necessary any site can be safely punctured in term infants, as long as automated lancets are used.9,10 Prewarm the foot by wrapping it in a hot moist towel (microwaved) for 5 minutes to produce hyperemia and enhance blood flow. Immobilize the foot in a dependent position with one hand. First, cleanse the heel with antiseptic solution and allow it to dry. Next, puncture the skin with the lancet. Allow the alcohol to dry to avoid false elevations in the glucose level. Avoid squeezing the foot because it may inhibit capillary filling and actually decrease the flow of blood. Furthermore, squeezing may lead to hemolysis of the sample. If blood does not flow freely, another puncture may be required.

Wipe away the first small drop of blood with gauze and allow a second drop to form. Place a heparinized capillary tube in the drop of blood and invert the proximal end of the tube to allow it to fill by capillary action. Fill the capillary tube until blood reaches the demarcation line on the tube. Over- or underfilling may result in clotting or erroneous test results. If 1-mL Microtainer tubes are used, hold the tube at an angle of 30 to 45 degrees from the surface of the puncture site. Touch the collector end of the tube to the drop of blood and allow the blood to drain into the tube. Gently tap the tube to facilitate flow to the bottom. Once filled, seal the tube with the accompanying cap. After an adequate specimen is obtained, apply a dry dressing to the puncture site.

When a heel stick is performed for an arterialized blood sample, use the same technique, but take care not to introduce ambient air into the sample. Place the tip of the tube as near the puncture site as possible to minimize exposure of the blood to environmental oxygen. Fill the tube as completely as possible and avoid collecting air in the tube. When the tube is full, occlude the free end with a gloved finger to prevent the entry of air, and cap both ends. Excessive squeezing of the foot may artificially lower the partial pressure of oxygen (Po2).

Complications

When performed properly, heel sticks are associated with a low incidence of complications. Use an automatic incision device to prevent lacerations. Rare complications include infection, scarring, and calcified nodules.

Interpretation

Multiple studies have demonstrated a good correlation between arterial and capillary specimens for determining pH and partial pressure of carbon dioxide (PCO2) in hemodynamically stable patients.11,12 However, determination of Po2 is not reliable when performed on blood obtained by capillary sampling.

Venipuncture

Indications and Contraindications

Venipuncture is used to obtain larger quantities of blood from infants and children and to obtain blood for culture. When collecting blood for culture, prepare the area of venipuncture with an appropriate antiseptic solution and allow the skin to dry. Wash off the cleanser promptly after blood has been collected because antiseptic solutions can irritate infant skin.

Equipment and Setup

A small-gauge butterfly or straight needle with a syringe are generally preferred for specimen collection in infants and young children because the negative pressure generated by standard specimen collection tubes (e.g., Vacutainer, Becton-Dickinson) may collapse small veins. A 3-mL or 5-mL syringe is less likely than a 10-mL syringe to cause vein collapse. Many providers find that it is easier to control needle position with a butterfly needle rather than a straight needle. If other access is not available, the butterfly needle may also serve as an infusion line after an adequate amount of blood is obtained. A 23-gauge butterfly needle will generally suffice for venipuncture, regardless of patient size.

Technique

As in adults, the usual site for venipuncture in infants and children is the antecubital fossa. However, any reasonably accessible or easily visible peripheral vein that will not be needed for IV cannulation may be used. Veins on the hands, feet, or scalp are frequently visible in young children (Fig. 19.2). Imaging devices (e.g., ultrasound, transillumination, or infrared devices) may also be used to locate and identify veins for venipuncture. These devices are discussed later in this chapter (see section on Vascular Line Placement: Venous and Arterial).

Assemble all necessary equipment, especially needles, out of sight of the child and have equipment within easy reach before beginning. Ask an assistant to help immobilize the patient when drawing blood from infants and small children. If an extremity vein is to be used, apply a tourniquet proximal to the selected vein. In small infants a rubber band can be used as a tourniquet. Be sure that the tourniquet is not so tight that it impedes arterial filling.

Cleanse the area surrounding the chosen site of skin penetration with antiseptic solution and allow it to dry. Apply slight distal traction to the skin to immobilize the vein. Insert the needle quickly through the skin and advance it slowly into the vein at an angle of approximately 30 degrees with the bevel facing up (Fig. 19.3). Successful vessel penetration is heralded by a flashback, or flow, of blood into the needle hub or butterfly tubing. Apply gentle suction by slowly withdrawing the plunger of the syringe. If the required amount of blood is greater than the capacity of the attached syringe, pinch off the tubing, remove the filled syringe, attach a new syringe, and apply gentle suction again after releasing the pinched tubing. Remove the tourniquet. After the required amount of blood is withdrawn, remove the needle and apply a sterile dressing and direct pressure to the puncture site.

Although peripheral sites for venous blood sampling are preferable in infants, the external jugular and femoral veins may also be used for venipuncture during resuscitation or when peripheral sites are inadequate. The external jugular vein lies in a line from the angle of the jaw to the middle of the clavicle
Capillary Blood Sampling

1. Obtain the blood sample from the medial or lateral portions of the plantar surface of the heel.
2. Prewarm the foot for 5 minutes to enhance blood flow. Immobilize the foot in a dependent position, cleanse with antiseptic, and allow to dry. Puncture the skin with the lancet.

Capillary Tube Method

3. Wipe away the first drop of blood with gauze and allow a second drop to form. Place a heparinized capillary tube in the drop of blood and allow it to fill by capillary action.
4. Fill the tube until the blood reaches the demarcation line on the tube. Maintain the blood in the capillary tube using your index finger to maintain capillary tension on the end of the tube.
5. Seal both ends of the tube with the caps provided or with wax or clay.

Microtainer Tube Method

6. If using a Microtainer tube, touch the collector end to the drop of blood and allow the blood to flow down the wall of the tube to the bottom.

Figure 19.1 Capillary blood sampling: capillary tube and Microtainer tube methods.
Venous access is unavailable, arterial blood may be used for most laboratory tests, including blood cultures.

When the infant is crying, this vein is more prominent. Ask an assistant to position the patient’s hips in mild abduction and extension while you palpate the artery. Identify its location by placing a mark on the skin just superior to the femoral triangle. If available, use ultrasound to assess the position of the femoral vessels. Prepare the femoral triangle with alcohol or another antiseptic agent. Use a povidone-iodine or chlorhexidine scrub when obtaining blood for culture. Puncture the skin and then direct the needle or catheter toward the umbilicus at a 30- to 45-degree angle to the skin and just medial to the pulsation of the femoral artery (see Fig. 19.5B). Apply slight negative pressure constantly throughout insertion. After the needle enters the femoral vein, withdraw the desired blood samples. Afterward, remove the needle or catheter unless an IV catheter for venous access is desired in this location. Apply pressure over the puncture site in the femoral triangle for a minimum of 5 minutes. Observe closely for recurrent bleeding.

Scalp veins can be very useful for venous sampling in small infants when other options are not possible or readily available. The anatomic considerations and technique are discussed later (see sections on Peripheral Venous Catheterization: Percutaneous and Peripheral Venous Catheterization: Venous Cutdown).

**Complications**

Complications of venipuncture include hematoma formation, arterial puncture, local infection, injury to adjacent structures, and phlebitis. Serious complications are uncommon. Use special care when attempting to puncture the external jugular or femoral vein. Inadvertent deep puncture in the neck can injure the carotid artery, vagus nerve, phrenic nerve, or apex of the lung. The femoral artery or nerve can be injured during puncture of the area around the femoral triangle. Fortunately, these injuries are unlikely when proper technique is used.

**Arterial Blood Sampling**

**Indications and Contraindications**

Arterial blood sampling is most often indicated to obtain an arterial blood gas (ABG) analysis. Consider sending venous blood for blood gas analysis because all parameters except $PO_2$ may be clinically useful. Arterial blood may be drawn for routine laboratory analysis or blood culture if venous blood is difficult to obtain. Avoid puncture of an artery if the overlying skin is infected, burned, or otherwise damaged. In addition, consider the presence of adequate collateral circulation and any potential coagulation disorders.

**Equipment and Setup**

For arterial puncture in infants and children, a small-gauge butterfly needle is preferable to a needle and syringe. A 23-gauge butterfly needle is used most often, although a 21-gauge
CHAPTER 19 Pediatric Vascular Access and Blood Sampling Techniques

357

The radial artery has several advantages that make it the most commonly used artery for blood sampling. First, its location makes it easy to palpate and puncture (Fig. 19.6A). The ulnar artery is more difficult to locate. Second, no vein or nerve is immediately adjacent to the radial artery, minimizing the risk of obtaining venous blood or damaging a nerve. Another advantage of the radial artery is the presence of good collateral circulation from the ulnar artery. The brachial artery has little collateral circulation and should be avoided unless an arterial sample is absolutely necessary and no other options are available. Limit use of the ulnar artery to preserve collateral circulation to the hand. As a general rule, do not use the femoral artery for obtaining routine blood samples.

Because the radial artery is used most frequently to obtain percutaneous arterial blood samples, the technique for radial artery puncture will be described. (See Chapter 20 for a discussion of the Allen test and the effect of heparin on arterial blood sampling.)

Hold the infant’s wrist and hand in your nondominant hand (see Fig. 19.6B). Hold the hand fully supinated with the wrist
Femoral Venipuncture

The femoral vein lies medial (in most patients) to the femoral artery and inferior to the inguinal ligament.

Position the hips in mild abduction and extension. Palpate the artery just superior to the femoral triangle. Puncture the skin and direct the needle toward the umbilicus at a 30° to 45° angle to the skin while remaining just medial to the femoral artery pulsation. Maintain slight constant negative pressure throughout the insertion. Apply pressure for a minimum of 5 minutes after the procedure and monitor for bleeding.

Figure 19.5 Femoral venipuncture. A, Anatomy. B, Technique.

Radial Arterial Blood Sampling

The radial artery is preferred for sampling procedures. Its superficial location is easy to palpate and puncture, and there are no immediately adjacent nerves or veins that may be injured or accidentally punctured.

Hold the wrist in a position of supination and slight extension. Insert the needle at a 30° to 45° angle while applying continuous, gentle suction with the plunger. Advance the needle until the radial artery is punctured and blood is returned or until resistance is met (see Fig. 19.7).

Figure 19.6 Radial arterial blood sampling. A, Anatomy. B, Technique.
Intravascular lines are indicated when ongoing access to the vascular system is necessary. Remember to consider using local anesthetics or procedural sedation if the clinical situation permits.

**Peripheral Venous Catheterization: Percutaneous**

**Indications and Contraindications**
In general, peripheral IV lines are indicated when the patient is unable to attain medical and nutritional goals with enteral therapy. These lines are used to provide fluids for resuscitation and maintenance needs and for the administration of medications.

**Equipment and Setup**
Over-the-needle catheters such as the Angiocath, JELCO I.V. (Smiths Medical, Dublin, OH) are the mainstay of peripheral venous catheterization. These thin-walled, flexible catheters range in size from 14 to 24 gauge. Select the appropriate gauge and length of the catheter based on the size of the child and the clinical situation. Larger-diameter catheters allow more rapid administration of fluids in emergency situations but may decrease the success of cannulation in young children with small veins. In general, use the smallest-gauge catheter that is appropriate for the clinical situation. For infants, a 22- to 24-gauge catheter is generally appropriate.

Attach T-connector extension tubing to the catheter after insertion to facilitate blood collection. This device makes flushing the catheter easier, especially while taping and securing the IV line. It also allows dressing changes without disturbing the IV insertion site.

Use either a homemade or commercially available device to protect the IV site from a child’s attempts to remove the line. An arm or leg board appropriate for the size of the child provides stabilization of the extremity after insertion. Macrodrip tubing and liter bags are inappropriate for use in infants because they can result in the inadvertent infusion of large amounts of fluid. An infusion pump is an ideal way of limiting fluid infusion while keeping the vein open.

**Vein Imaging Devices**
A variety of imaging modalities, including ultrasound, transillumination, and infrared technologies, can be used to help locate peripheral veins for cannulation. In adults, data support the use of ultrasound to facilitate peripheral vein cannulation in those with difficult access. The data is more limited in children. Several small pediatric studies demonstrated a modest benefit of ultrasound, especially in patients with difficult access and other studies have shown no benefit. Transillumination has been shown to facilitate peripheral venous cannulation in infants and small children. These devices work by projecting a high-intensity light into the patient’s subcutaneous tissue, causing the veins to contrast with surrounding tissue, and making them easier to locate. Newer vein imaging technology, available as VeinViewer (Christie Medical Holdings, Memphis, TN), AccuVein (Cold Spring Harbor, NY), Vein-Eye (Newmaw Medical Ltd, Liverpool, United Kingdom), Veinsite (VueTek Scientific, Gray, ME), Vasculuminator (De Koningh Medical Group, Arnhem, the Netherlands) and others, uses near-infrared technology to project an enhanced image of the subcutaneous veins onto the patient’s skin. Theoretically, knowledge of the location of the venous valves and the course of the vessel can assist the clinician in selecting the best area to be cannulated. However, multiple

**VASCULAR LINE PLACEMENT: VENOUS AND ARTERIAL**

Intravascular lines are indicated when ongoing access to the venous or arterial circulation is necessary. Remember to consider...
Technique

Fig. 19.2 shows sites available for placement of a peripheral IV line. The most common sites chosen for IV insertion in infants and children are the superficial veins of the dorsum of the hand, the antecubital fossa, the dorsum of the foot, and the scalp (in newborns and small infants) (Video 19.1). The veins on the dorsum of the hand are relatively straight and are easily stabilized because they lie flat on the metacarpals. If the hand is chosen, take the child's hand preference into consideration and avoid the dominant hand or hand used for thumb sucking whenever possible. Veins in the antecubital fossa (cephalic and basilic veins) are easily accessible; however, the angulation across the fossa makes advancement of the catheter difficult. These veins may be palpable even if they are not easily visible. Tributaries of the dorsal venous arch on the dorsum of the foot, like those on the dorsum of the hand, are relatively straight, and the extremity is easily immobilized after insertion. Because indwelling catheters in this location prevent mobility, consider using this site only in preambulatory patients or after attempts at other sites have been unsuccessful. The scalp veins are easy to cannulate, but their use is primarily limited to small infants. The particular site is a matter of preference, so choose the vein that appears to be the easiest to cannulate. When possible select the most distal vein that is large enough to accommodate the catheter and leave the larger, more proximal veins in case the initial attempts are unsuccessful or prolonged IV therapy is needed. If peripherally inserted central venous catheter (PICC) placement may be required during the hospitalization, do not use PICC insertion sites if possible.

With few exceptions, the same techniques used for IV insertion in adults may be used in infants and children, especially in the veins of the distal ends of the extremities. If an extremity location is chosen, place a tourniquet proximal to the planned site of insertion. Warm the extremity to induce vasoconstriction in the surface veins. Flush the tubing of the T-extension set before venipuncture with a sterile IV solution, such as normal saline (NS), to prevent air embolism. Direct the IV catheter through the skin at a 10- to 20-degree angle and slowly advance it until blood return is noted (Fig. 19.8, step 1). Next, advance the catheter over the needle and into the vein. Retract the needle and connect the IV line to the hub of the catheter by means of a T-extension set (see Fig. 19.8, step 2). After 1 mL of saline has been flushed through the line, inspect the site for swelling, which may indicate infiltration. If an artery is entered during placement of the needle and fluid is infused, blanching will occur in the area. If this happens, remove the IV catheter, maintain slight pressure for 5 minutes, and repeat the procedure at another site.

Secure the catheter to the skin with a 0.5-inch piece of tape passed over the catheter hub and skin. Place a second piece of tape adhesive side up and slip it under the catheter hub. Cross it over the catheter hub in a V shape (see Fig. 19.8, step 2). After securing the catheter with tape, cover the entire area with a transparent sterile dressing such as Tegaderm (3M, St. Paul, MN) or OP Site (Smith and Nephew Medical, Massillon, OH). Loop back the tubing of the T-extension set, place a piece of tape midway over the tubing, and secure it to the skin. This ensures that the IV tubing will not be accidentally dislodged if it is suddenly pulled. Securely tape the hand and forearm to an arm board for immobilization (see Fig. 19.8, step 3). Occasionally, the flow rate of the infusion may depend on the position of the catheter, especially if the catheter spans a joint or the tip abuts a venous valve. Adjust the hand position or catheter with strategically placed sterile gauze or withdraw the catheter slightly to remedy the problem.

Obtain blood specimens just after IV insertion. If the IV line has been flushed, the initial blood draw will be diluted. To prevent dilution, withdraw 5 mL of blood from the catheter before collecting the samples. This 5 mL of “waste” can be either discarded or reinfused into the patient. Next, remove the syringe. Connect the T-extension tubing to the IV infusion tubing, and set the infusion pump at the desired rate. Send blood for culture only if the skin was cleaned with an appropriate antiseptic before insertion of the IV catheter.

If a butterfly infusion set is used, grasp the wings of the butterfly between the thumb and forefinger and introduce the needle beneath the skin approximately 0.5 cm distal to the anticipated site of vein entrance (Fig. 19.9, step 1). Advance the needle slowly toward the vessel until blood appears in the tubing, which indicates that the vessel has been entered. Next, remove the tourniquet if one was used. Flush the needle with 0.5 to 2 mL of IV fluid, such as NS, to ensure that the needle is properly in place within the vein. If infiltration occurs, as noted by a subcutaneous bump, remove the IV line and repeat the process at another site.

After the wings are secured, tape the tubing of the butterfly set in a loop so that it is not pulled inadvertently. Place a wisp of cotton under the wings of the butterfly if the flow rate of the infusion is affected by the position of the catheter. Tape a small medication cup over the wings and the needle to protect the IV line (see Fig. 19.9, step 2). Connect the tubing of the butterfly set to the tubing from the IV system. It is generally preferable to use standard over-the-needle IV catheters whenever possible, whether for extremity or scalp IV lines, because they are less likely to infiltrate and will last longer. However, a small butterfly needle that can be inserted temporarily until achievement of additional access is sometimes the only option short of IO access.

External Jugular Venous Catheterization

The external jugular vein is superficial and easily visible. It can be used when other attempts at peripheral IV access have been unsuccessful but can be difficult to access if airway or other procedures are occurring simultaneously. The procedure should be discussed with parents because laypersons may have fears about a needle being placed in their child's neck.

Technique. The external jugular vein is most often entered with a standard over-the-needle IV catheter. The vein lies in a line from the angle of the jaw to the middle of the clavicle and is usually visible on the surface of the skin. It is more prominent when the child is crying or in Trendelenburg position. Ask an assistant to restrain the patient in a supine position with the head and neck extended and a towel roll placed under the shoulders. Turn the head approximately 40 to 70 degrees from the midline and away from the side to be punctured (see Fig. 19.4). Cleanse the skin surrounding the area to be punctured with alcohol (or another antiseptic solution). If time allows, infiltrate the overlying skin with 1% lidocaine. Place a finger just above the clavicle to distort the jugular vein.

Puncture the skin approximately one-half to two-thirds of the distance from the angle of the jaw to the clavicle. Advance the needle slowly until the jugular vein is entered. Once there is a flash in the needle hub, advance the catheter into the vein.
Peripheral Venous Catheterization: Venous Cutdown (See also Chapter 23)

Indications and Contraindications

Peripheral venous cutdown is rarely performed in the ED because IO access is rapid and safe. Even in experienced hands, a saphenous vein cutdown may take more than 10 minutes and is associated with a higher rate of infection than other routes of vascular access. Nevertheless, if peripheral venous, central venous, or IO access cannot be obtained, venous cutdown may provide an alternative means of emergency venous access. Contraindications to peripheral venous cutdown are major trauma to the extremity, vascular injury of the extremity proximal to the intended site of cutdown, or soft tissue infection of the intended incision site. A coagulation disorder is a relative contraindication.

Complications

Serious complications of peripheral IV lines and IV fluid therapy are rare. Complications include infection, phlebitis, inadvertent arterial puncture, injection of sclerosing agents into the subcutaneous space with resultant necrosis of the skin (especially in small infants), pneumothorax, air embolism, and administration of inappropriate volumes of fluid. Puncture of the apex of the lung leading to pneumothorax is possible with external jugular puncture if the needle is inserted too far. Attention to antiseptic procedures during insertion and maintenance of the IV system decreases the risk of infection.
Pass two 4-0 silk sutures under the vein. Place one distally to stabilize the vein, and place the other proximal to the intended site of venipuncture. The distal suture may be tied, but if left untied it can still be used for stabilization of the vein and to occlude blood flow by applying gentle traction. Removing the untied distal suture after cannulation of the vein may allow subsequent recannulation after eventual catheter removal. Once the vein is isolated, either place a catheter surgically through a venotomy or cannulate the vein directly with a standard IV catheter. If venotomy is chosen, use fine scissors or a scalpel blade to make an oblique or V-shaped incision in the anterior wall of the vein between the sutures (see Fig. 19.10, step 4).

Grasp the silastic catheter (prefilled with saline solution) with forceps and advance it into the vein for a distance of 2 to 3 cm (see Fig. 19.10, steps 5 and 6). This is usually the most difficult and time-consuming portion of the procedure. Use a vein dilator, an L-shaped lifter, or a forceps to hold the venotomy incision open (see Fig. 19.10, steps 7 and 8). Pull downward on the distal tie to provide countertraction and stabilize the vein during advancement of the catheter. Remove the tourniquet and tie the proximal suture around the vein with the catheter inside. Take care to avoid tying the suture so tightly that it occludes the catheter. If the distal suture was tied, tie the free ends of the suture around the catheter to provide additional stability. If the distal suture was not tied, remove it at this point. Tie the proximal suture to secure the catheter, but leave the ends long enough so that the suture can later be removed. This allows recannulation once the catheter is removed.

Alternatively, direct cannulation of the vein may be achieved more rapidly and, unlike venotomy, may preserve the vein for future cannulation. If direct cannulation is chosen, isolate and immobilize the vein, using a standard IV catheter to directly cannulate the vein. Essentially a percutaneous catheter insertion technique is used to catheterize a directly visualized vein (Fig. 19.11).

**Equipment and Setup**

Successful venous catheterization via cutdown in infants and small children requires an assistant, good lighting, and a selection of standard IV catheters or silastic catheters prefilled with NS. Sterile instruments may be available in a “cutdown tray” or can be individually assembled. Assemble regular and mosquito hemostats, fine forceps, a vein dilator, a vein lifter, silk suture, a needle driver, and fine scissors or a scalpel.

**Technique**

For the purpose of illustration, exposure and cannulation of the saphenous vein will be discussed (Fig. 19.10). The same principles apply when cutdown is performed on most peripheral veins. Begin with complete immobilization of the thigh, leg, ankle, and foot by taping them to a padded leg board. Attach this board in turn to the table or bed where the procedure is being performed (see Fig. 19.10, step 1). Prepare the area around the medial malleolus with antiseptic solution and drape it with sterile towels. Infiltrate 1% lidocaine in an area approximately 1 cm proximal and 1 cm anterior to the medial malleolus. No major nerves or tendons accompany the saphenous vein in this location.

Place a tourniquet proximally on the leg and make a transverse skin incision (usually approximately 2 cm in length) in the anesthetized area. Insert a small mosquito hemostat into the wound with the concavity of the clamp upward. Using the tip of the hemostat, “scoop up” all tissue lying against the bone (see Fig. 19.10, step 2). This will invariably lift the vein out of the wound along with the surrounding tissue. Use fine forceps or a mosquito hemostat to separate and remove all nonvenous structures so that only the saphenous vein is left tented over the hemostat (see Fig. 19.10, step 3). To avoid injury to the vein during dissection, spread the ends of the hemostat parallel to the direction of the vein, never transversely.

**Fig 19.9** Scalp vein intravenous catheterization using a butterfly needle.
**Venous Cutdown**

1. Immobilize the ankle and the site of skin incision.

2. Use a curved hemostat to scoop up the vein. Keep the point of the hemostat against the bone.

3. Dissect the vein free with the curved hemostat.

4. Place silk sutures around the distal and proximal end of the exposed vein. Tie only the distal suture. Make a small venotomy with a scalpel or fine scissors.

5. Grasp the catheter (prefilled with saline) with a hemostat.

6. Advance the catheter into the vein.

7. Use a vein dilator or forceps to hold the venotomy incision open.

8. Advance the catheter 2–3 cm into the vein.

9. Close the incision with 4–0 nylon sutures. Place a sterile dressing over the wound and secure the ankle to the foot board.

Figure 19.10 Venous cutdown (saphenous vein). (From Suratt PM, Gibson RS: Manual of medical procedures, St. Louis, 1982, Mosby. Reproduced by permission.)
Continued infusion of saline through the catheter from an attached syringe will ensure patency. Orient the catheter into one corner of the incision and close the incision with interrupted 4-0 nylon suture. Wrap the skin suture nearest the catheter around it and tie it to hold the catheter in place. Control bleeding with direct pressure. Place antibiotic ointment over the wound and apply a sterile occlusive dressing. Connect the IV tubing and tape it securely to the foot board to prevent inadvertent removal of the catheter (see Fig. 19.10, step 9).

Change the dressing carefully every day with sterile technique and reapply antibiotic ointment. When cared for properly catheters can remain in place for as long as 7 to 10 days, but generally it is best to replace the catheter and insert it into another site after 3 to 4 days. At the first sign of infiltration or infection, remove the catheter. See Chapter 23 for more information on venous cutdown techniques.

Complications
In addition to the problems with percutaneous catheter placement that were discussed previously, venous cutdown can result in wound infection and thromboembolism. Adjacent structures may be injured during the incision and subsequent blunt dissection. Extravasation of infusate may result if the vein is not ligated. Light pressure on the closed wound will generally prevent continued extravasation.

Central Venous Catheterization: Percutaneous

Indications and Contraindications
Central venous cannulation is indicated in the ED when peripheral venous access is limited or impossible. In young children central line placement can almost always be delayed until after initial stabilization. Central venous lines have traditionally been used for hemodynamic monitoring but the use of point-of-care ultrasound in the ED may decrease this need. Ultrasound evaluation of the inferior vena cava (IVC) can be used to estimate intravascular volume status.28 Central venous access is preferred for infusion of vasoactive and sclerosing medications. Nevertheless, 2007 pediatric sepsis guidelines state that vasoactive infusions, including epinephrine, can be started via IO or peripheral venous lines, so placement of a central venous line is not required prior to starting vasopressors.29

Contraindications to percutaneous placement of central venous catheters include local infection or burns at the insertion site; vascular insufficiency of an extremity; or obstruction or compression of the access veins by tumor, abnormal vessels, hematoma, or thrombus. Caution should be used in patients with uncorrected coagulopathies or with malformations or deformations distorting vascular anatomy. Attempts at central venous access should not result in a delay in medication and fluid administration during resuscitation efforts, but if a central line is in place, medications should be given centrally rather than peripherally.10

Equipment and Setup
Commercially available kits are convenient because they contain most of the items needed for the procedure. The catheters in these kits are typically made of a silicone elastomer, polyvinyl chloride, or polyethylene; some are available with an antimicrobial coating, which may reduce infection rates. Catheter length is variable and one- to three-lumen catheters are available. Rapid volume replacement, as in the case of severe dehydration or acute blood loss from trauma, is best achieved by inserting a short, large-bore catheter for the initial resuscitation and stabilization. Table 19.1 lists appropriate central line sizes by age and weight.

Other necessary equipment includes antiseptic solution, gauze pads, sterile drapes, gowns, sterile gloves, caps, masks, syringes (3, 5, and 10 mL), sterile transparent skin coverings, Luer-Lok three-way stopcocks, 0.25% to 1.0% lidocaine, flush solution (1 to 2 Units heparin/mL NS), and IV tubing with a T-connector extension. Depending on the vein to be accessed, restraint of the extremity, pelvis, or head may require a padded support, an assistant, or both.

Technique
Young patients should ideally be sedated, or if that is not possible, well restrained, to minimize movement during the procedure. Local anesthesia should be administered prior to central line placement. Equipment should be placed in an easily accessible location to minimize the need to reach or move while inserting the catheter. All catheter lumens should be prefilled and clamped before beginning the procedure.

Strict sterile precautions with cap, gown, mask, and sterile drapes should be used when placing all but the most emergent central lines to decrease the risk of catheter-associated bloodstream infections. The skin should be cleaned with chlorohexidine in preparation for the procedure unless there is a specific contraindication to chlorhexidine.11,12 In this case, betadine may be used.

Percutaneous placement of central venous catheters can be accomplished with either a guidewire (Seldinger) technique or an over-the-needle catheter. It can be difficult to thread a

<table>
<thead>
<tr>
<th>TABLE 19.1 Pediatric Central Line Sizes</th>
</tr>
</thead>
<tbody>
<tr>
<td>AGE (YEARS)</td>
</tr>
<tr>
<td>&lt;1</td>
</tr>
<tr>
<td>1–5</td>
</tr>
<tr>
<td>6–10</td>
</tr>
<tr>
<td>11 +</td>
</tr>
</tbody>
</table>
guidewire in young children; one technique to maximize the chance of placing a usable catheter is to use a modified Seldinger technique. Use the over-the-needle catheter from the central line kit to place a catheter. The catheter can be left in place for initial resuscitation or the guidewire can immediately be advanced through the catheter. If the guidewire can be threaded into the vein, the catheter can then be removed, the skin dilated, and the longer central catheter can be placed over the wire. If the guidewire cannot be advanced sufficiently through the catheter, remove the guidewire and leave the catheter in place for venous access. The catheter-over-the-needle technique can be used in the femoral, subclavian, or internal jugular veins.

Ultrasound guidance for internal jugular, subclavian, and femoral line placement has been shown to improve safety in adult patients and the Agency for Healthcare Research and Quality has identified the use of ultrasound guidance for placement of central venous catheters as an important patient safety practice. Although there is less literature on the use of ultrasound for placement of central lines in children, ultrasound-guided central line placement has been associated with decreased numbers of punctures, complications, and increased procedural success in pediatric patients.

**Femoral Catheterization**

Femoral venous catheterization is the central venous access route most commonly used in infants and children in emergency situations (Fig. 19.12). The anatomy is fairly straightforward and the femoral arterial pulse provides a good landmark for femoral venous catheterization. The femoral site is advantageous because the risk of mechanical complications is low. In the event of inadvertent arterial puncture or venous laceration, hemostasis can be achieved by the application of direct pressure. Femoral vein catheterization does not carry any risk for hemothorax or pneumothorax. Studies in children have found that when controlling for other factors, such as number of days the catheter is left in place, use of the femoral location does not pose a higher risk of catheter-associated bloodstream infection when compared to subclavian or internal jugular vein locations. Another advantage is that placement of a femoral catheter is less likely to interfere with other resuscitative procedures.

![Central Venous Catheterization: Femoral Approach](image)

**Figure 19.12** Central venous catheterization: femoral approach.
procedures than placement of a subclavian or internal jugular catheter.

**Technique.** Restrain the child adequately to permit exposure of the inguinal region. Use an introducer needle with or without a syringe to enter the femoral vein. If using landmarks rather than ultrasound to locate the vein, palpate the femoral artery with one finger, and puncture the skin just medial to the artery. Enter the skin at a 30- to 45-degree angle approximately 1 cm below the inguinal ligament. Direct the needle in a line toward the umbilicus. If using a finder needle with a syringe, apply continuous, gentle suction while inserting the needle. Gently pass the wire through the needle into the proximal end of the vein when blood return is noted. The wire should not meet resistance when introduced gently. Make sure that the proximal end is always visibly protruding from the hub of the needle. If resistance to passage of the wire is encountered, remove it to assess the needle's position within the vessel. If resistance to removal of the wire is encountered, withdraw the needle and the wire together to prevent shearing off the end of the wire.

An alternative method is to use the catheter over the needle that is supplied in the central line kit to enter the vein and advance the catheter into place. Once the catheter is placed, remove the needle and advance the wire through the catheter. If the wire threads completely, leave the wire in place and remove the catheter. If the wire cannot be threaded into the vein, remove the wire, leaving the catheter in place, and secure the catheter to be used for access until alternative access is obtained. Always occlude the hub of the open finder needle or catheter with a sterile gloved finger when the wire is not in place to prevent excessive blood loss or air embolism.

Once the wire is in place and the catheter or needle removed, make a small incision (1 to 2 mm) with a No. 11 scalpel blade in the skin at the wire's entry point to allow passage of the dilator (see Fig. 19.12, step 2) and the catheter itself. When making the incision, point the sharp edge of the blade away from the wire. Advance the dilator gently over the wire and then remove the dilator. Advance the catheter over the wire and into the vein (see Fig. 19.12, step 3). Remove the wire. Withdraw blood from the catheter ports and then flush them with a sterile saline solution. Secure the catheter to the skin with silk or nylon sutures (see Fig. 19.12, step 4). Place a sterile, transparent, and impermeable dressing over the exit site.

Note that during cardiopulmonary resuscitation (CPR), palpable pulsations or Doppler tones in the femoral vein may be detected. Hence, if the vein is not found medial to the pulsations, consider catheterization of the pulsating vessel during CPR as a last resort when other options for vascular access or drug delivery are unavailable. Alternatively, ultrasound can be used to locate the femoral vein.

**Internal Jugular Venous Catheterization**

The internal jugular veins lie within the carotid sheath along with the carotid artery and vagus nerve. The lower part of the vein lies within the triangle formed by the sternal and clavicular heads of the sternocleidomastoid muscle and then becomes more lateral and anterior to the artery as it joins the subclavian vein. The right internal jugular vein is preferred over the left because the internal jugular, the innominate vein, and the superior vena cava form a nearly straight line into the right atrium and complication rates appear to be lower. Placement of internal jugular venous access may not be possible when other critical procedures are occurring and it is not possible if the patient is in a cervical spine immobilization device.

**Technique.** Three approaches to internal jugular catheterization are possible (including the anterior, median or central, and posterior approaches, as discussed in Chapter 22). The median or central approach is described here. Use of ultrasound to guide cannulation of the internal jugular vein appears to improve success rates (see Chapters 22 and 66). Position the child in the same fashion as described for external jugular venous catheterization. The Trendelenburg position will distend the vein. For the medial or central approach, use the apex of the angle formed by the sternal and clavicular heads of the sternocleidomastoid muscle as the puncture site. If a line were drawn from the mastoid process to the sternal notch, the apex of the angle formed by the two muscular heads would fall approximately along the middle third of that line. Introduce a needle attached to a syringe at the apex of the triangle at an angle of 30 degrees downward and toward the ipsilateral nipple (Fig. 19.13). Advance the needle slowly until the jugular vein is entered. Keep the syringe connected to the needle at all times to maintain constant negative pressure and avoid air embolism. After blood flow is obtained, remove the syringe and place a finger over the hub of the needle. Insert a guidewire during a positive pressure breath or exhalation. Remove the needle and introduce a catheter via the Seldinger technique. Pass the catheter far enough to reach the junction of the superior vena cava and right atrium. Check the catheter for blood return, secure the line with sutures, and apply a sterile occlusive dressing. Confirm proper location of the catheter and rule out pneumothorax with a chest radiograph.

**Subclavian Venous Catheterization**

The subclavian vein is used less frequently for central venous access in children than in adults (see Fig. 19.13). The technique is more difficult in pediatric patients because of the smaller size of the vessel, as well as a more cephalad location under the clavicles. Placement of subclavian venous access may not be possible when other critical procedures are occurring or if the patient cannot be properly positioned due to cervical spine immobilization.

**Technique.** The approach to the vein is more lateral in children than in adults. The infraclavicular approach is described hereafter. Turn the patient's head away from the side to be punctured and place a towel roll under the shoulders. The right side is preferred because the dome of the lung is more cephalad on the left side. The needle insertion site is at the distal third of the clavicle in the depression created between the deltoid and the pectoralis major muscles.

Introduce the finder needle bevel up and advance it slowly while applying negative pressure with the attached syringe. Keep the syringe and needle parallel to the frontal plane. Direct the needle medially and slightly cephalad, beneath the clavicle toward the posterior aspect of the sternal end of the clavicle (i.e., toward a fingertip placed in the sternal notch). Advance the needle until blood return is obtained. Turn the syringe so that the bevel of the needle points caudad to direct the guidewire to the superior vena cava. Remove the syringe from the needle and insert the wire during a positive pressure breath or natural exhalation. Remove the needle and introduce the catheter over the wire via the Seldinger technique, as previously described for femoral catheterization. The cardiac monitor may show a
Emergency Vascular Access

Vascular access is a key component of any resuscitation to allow the administration of fluid and medications and to obtain blood samples. However, achieving venous access during pediatric resuscitation can challenge even the most seasoned clinician, not infrequently requiring 10 or more minutes. Children who are successfully resuscitated have vascular access achieved significantly sooner than those not resuscitated. Emergency IV access is most prolonged in children younger than 2 years, the age group in which the majority of cardiopulmonary arrests occur.

For these reasons, clinicians should obtain IO access in critical patients if IV access is not established quickly, rather than delay treatment with attempts to place peripheral or central venous catheters (see Chapter 25). IO lines are easily and rapidly placed, can accept any drug given parenterally in similar doses, and are likely underused in many EDs. If no IV or IO line is available, give certain drugs via the endotracheal tube (see Chapter 26) while attempts at venous access are initiated. However, endotracheal drug administration has been shown to be less efficacious than IV administration in adults, and this route of administration should be used only when no alternatives are available. Placement of an internal jugular, subclavian, or femoral line can prove time-consuming and is thus discouraged in the unstable patient. Umbilical and IO lines are effective alternatives.

Umbilical Vein Catheterization

Indications and Contraindications

The major indication for umbilical vein catheterization is the need to access the vascular system for emergency resuscitation and stabilization of neonates. It may also be used for exchange transfusions and short-term central venous access in newborns. The umbilical vein may remain patent for up to 2 weeks after birth. In neonates who require emergency access in the ED,
a peripheral vein would be preferable, but attempting to
cannulate one of the umbilical vessels could be lifesaving.
Omphalitis, omphalocele, gastrochisis, and peritonitis are all
contraindications to access of the umbilical artery or vein.

**Equipment and Setup**

Place the infant beneath a radiant warmer because keeping
the infant warm during the procedure is critical. Restrain the
extremities. Wear a mask, cap, gown, and sterile gloves. Assess
the length of catheter needed before the procedure. Methods
to estimate the best length for both umbilical vein and artery
lines are discussed hereafter.

**Technique**

Hold the umbilical stump upright and scrub the cord with a
bactericidal solution. Avoid pooling of liquid at the infant’s
side because this may be associated with blistering of the skin
under a radiant warmer. Drape the umbilical area in sterile
fashion with the infant’s head left exposed for observation
(Video 19.2).

Place a loop of umbilical tape or a purse-string suture at
the junction of the skin and the cord to provide hemostasis
and to anchor the line after placement (Fig. 19.14, step 1). Cut
the cord with a scalpel approximately 1 cm from the skin (often
less in infants who are not newly born), and identify the vessels.
There are two arteries and one vein in the umbilicus. The
vein is usually located at 12 o’clock and has a thin wall and
large lumen. It may continue to bleed after cutting. The two
arteries have thicker walls and smaller lumens. Constriction
reduces bleeding after the arterial vessels are cut. Occasionally,
a persistent urachus may be mistaken for the umbilical vein,
but the presence of urine in the urachus may help correctly
identify that structure.

Flush the catheter (3.5 Fr [preterm infants] to 5.0 Fr [term
infants]) with heparinized saline and attach it to a three-way
stopcock. Place the catheter into the lumen of the umbilical
vein and advance it gently (see Fig. 19.14, step 2). An umbilical
vein line can be placed in a fashion that is intended either for
emergency use or for more chronic use. In an emergency case,
advance the catheter only 1 to 2 cm beyond the point at which
good blood return is obtained. This is usually only 4 to 5 cm
in a term-sized infant. Beyond this distance, the umbilical vein
branches. Therefore the catheter may enter a branch of the
portal vein within the liver (as evidenced by resistance at 5 to
10 cm). Infusion of medications into a catheter in the portal
vein can result in liver necrosis. In an emergent case, a line
placed no greater than 5 cm into the patient can be used without
confirmatory radiographs.

If the patient does not require emergent access and a more
enduring line is desired for blood draws, monitoring central
venous pressure or infusion of medications, high concentrations
of glucose (>10%), IV fluids, and hyperalimentation solutions,
then the line is instead inserted 10 to 12 cm in a term-sized
infant to reach the IVC. Because of the potential for placement
into the portal vein and resultant complications, radiographs
are required prior to usage to confirm placement into the IVC.
Note that an umbilical venous catheter will proceed directly
cephalad (without making a downward loop) until it passes
through the ductus venosus (Fig. 19.15).

Some practitioners use standardized graphs to estimate the
length of insertion. Such graphs are based on the shoulder-to-
umbilicus length (Fig. 19.16). Shoulder-to-umbilicus length
is the perpendicular line measured from the tip of the shoulder
to the horizontal level of the umbilicus. If the graph is not
available, the shoulder-to-umbilicus length multiplied by 0.6
gives an approximate insertion length that will result in the

---

**Umbilical Vein Catheterization**

1. Hold the umbilical stump up and scrub it with bactericidal
   solution. Place a loop of umbilical tape or a purse-string
   suture at the junction of the skin and the cord. Cut the
   cord with a scalpel approximately 1 cm from the skin and
   identify the vessels. The single vein is thin walled, has a large
   lumen, and may continue to bleed after cutting. The two
   arteries have thicker walls and smaller lumens and do not usually
   bleed.

2. Advance a preflushed catheter into the vein and gently advance it.

   For emergency use, advance only 1–2 cm beyond the point at
   which good blood return is obtained. This is usually only 4–5 cm
   for a term-sized infant.

   For longer-term use, the catheter may be passed into the inferior
   vena cava. (See text for details.)

---

**Figure 19.14** Umbilical vein catheterization. (From Zaoutis L, Chiang V: *Comprehensive pediatric hospital medicine*, Philadelphia, 2007, Elsevier.)
tip of the catheter being above the diaphragm but below the right atrium in the IVC.\textsuperscript{45} Formulas based on birth weight, such as \((1.5 \times \text{birth weight in kg}) + 5.5\) cm, can further guide placement.\textsuperscript{46}

Air embolism may occur at the time of catheter removal if the infant generates sufficient negative intrathoracic pressure (as during crying) and causes air to be drawn into the patent umbilical vein. Therefore caution must be used during removal of the catheter to ensure that the vein is promptly occluded (by tightening a purse-string suture or applying pressure on or just cephalad to the umbilicus).

An alternative method has been described. In this method, maintain the clamp on the umbilical cord, then make a cut proximally in the amniotic membrane over the umbilical vein and bluntly dissect through Wharton’s jelly down to the vein. At this point, place a catheter into the umbilical vein through this site. The remainder of the procedure is identical to the standard technique. This option may be attractive in situations where transecting the cord proximal to the site of the clamp is not possible or not desired.\textsuperscript{47} Similarly, use of a 14-gauge angiocatheter with the needle removed has been described, and may be an option if an umbilical line or feeding tube is not readily available.\textsuperscript{48}

**Complications**

Complications of umbilical venous catheters include hemorrhage, infection, injection of sclerosing substances into the liver (resulting in hepatic necrosis), air embolism, catheter tip embolism and resultant ischemia (including myocardial infarction), liver hematoma, arrhythmias, effusions, malplacement, and vessel perforation.\textsuperscript{49–51} Although careful technique during insertion and maintenance should be maintained to minimize complications, studies suggest that the overall complication rate of umbilical venous catheters is lower than femoral catheters, predominantly due to a lower thrombosis rate.\textsuperscript{52}

![Figure 19.15](image1.png)

**Figure 19.15** An umbilical venous catheter will proceed directly cephalad (without making a downward loop) until it passes through the ductus venosus.

![Figure 19.16](image2.png)

**Figure 19.16** \(A\), After measuring the shoulder-to-umbilicus length, a standardized graph can be used to determine the appropriate length of the umbilical venous catheter, or \(B\), the umbilical arterial catheter. The venous catheter should be inserted into the inferior vena cava below the level of the right atrium. The appropriate length of the arterial catheter depends on whether a “high” or “low” line is desired (see text for explanation). (\(A\) and \(B\), From The Johns Hopkins Hospital, Nechyba C, Gunn VL: The Harriet Lane Handbook: A Manual for Pediatric Home Officers, ed 16, St. Louis, 2002, Mosby.)
Umbilical Artery Catheterization

Indications and Contraindications
Umbilical artery catheterization is a useful procedure in the care of newborn infants who require frequent monitoring of ABGs and arterial blood pressure, fluid and medication administration, and exchange transfusions. Either one of the two umbilical arteries may be cannulated for resuscitation purposes, but an umbilical vein is generally technically easier to cannulate and may be preferred in an emergency. These arteries can typically be accessed up to 7 days after birth. Contraindications include those described in umbilical venous access, as well as necrotizing enterocolitis.

Equipment and Setup
The equipment required for umbilical artery catheterization is identical to that used for umbilical venous catheterization. Additional equipment needed for continuous arterial pressure monitoring and infusion should be readily available. Estimate the catheter length needed before starting the procedure. Do this by using a standard graph (see Fig. 19.16B) or a birth weight regression formula.

Technique
The technique of umbilical artery catheterization is similar to that described for umbilical vein catheterization in the preceding section. After the umbilical arteries have been located (Fig. 19.17, step 1), grasp the cord with a curved hemostat near the selected artery. Ask an assistant to use two hemostats to grasp each side of the cord and slightly evert the edges to aid in exposure of the arteries. Using curved iris forceps without teeth, gently dilate the artery (see Fig. 19.17, step 2). Sometimes, repeated passes of the forceps are required because the umbilical artery can spasm and make the procedure difficult. Attach a

---

**Figure 19.17** Umbilical artery catheterization.

1. Prepare the umbilical stump by placing a purse-string suture or loop of umbilical tape at the base of the cord. Make a fresh cut across the cord and identify the single umbilical vein and two umbilical arteries.

2. Grasp the cord with a hemostat near the selected artery. Use toothless curved iris forceps to gently dilate the artery.

3. Introduce the preflushed catheter into the artery. Maintain gentle cephalad tension on the cord, and advance the catheter with slow, constant pressure toward the feet.

4. Secure the catheter in place with the previously placed purse-string suture.

5. For additional protection, use a strip of pleated tape to secure the catheter to the abdominal wall.
3.5-Fr to 5-Fr catheter to a three-way stopcock and flush it with a sterile heparinized solution. Introduce the catheter into the dilated artery (see Fig. 19.17, step 3). Use a 3.5-Fr to 4-Fr catheter for infants weighing less than 2 kg and a 5-Fr catheter for those weighing 2 kg or more.

When the catheter is being inserted, place gentle tension on the cord in a cephalad direction, and advance the catheter with slow, constant pressure toward the feet. Resistance is occasionally felt at 1 to 2 cm. Overcome this with gentle, sustained pressure. On the other hand, if the catheter passes 4 to 5 cm and then meets resistance, this generally indicates that a “false passage” through the vessel wall has occurred, with the catheter curving caudad. Occasionally, one may bypass the perforation by reattempting catheterization with a larger catheter.

Acceptable positions of umbilical artery catheters are between T6 and T9 (“high line”) and between L3 and L5, just above the aortic bifurcation (“low line”). High lines are associated with a lower incidence of thrombotic complications.34 Use graphs and formulas to estimate the proper catheter length for insertion at both the high and low positions (see Fig. 19.16B), but confirm placement with a radiograph. These formulas may not be as accurate in babies at the extremes of weight. A new formula, developed by Wright ([4 × birth weight in kg] + 7 cm), accurately predicts high-line location 83% of the time.35 Once sterile technique is broken, do not advance the line. It is therefore preferable to position the catheter too high and to withdraw it as necessary according to postinsertion radiographs. This will show the catheter proceeding from the umbilicus down toward the pelvis, making an acute turn into the internal iliac artery, continuing toward the head into the bifurcation of the aorta, and then moving up the aorta slightly to the left of the vertebral column (Fig. 19.18). After it has been properly positioned, tie the catheter with the previously placed suture and tape it to the abdominal wall (see Fig. 19.17, steps 4 and 5).

Complications
Complications of umbilical artery catheterization include hemorrhage, infection, thromboembolic phenomena (especially involving the kidneys, gastrointestinal tract, and lower extremities), aortic thrombosis, aortic aneurysm, vasospasm, air embolism, vessel perforation, peritoneal perforation, nerve lesions, and hypertension from renal artery obstruction.

If the catheter becomes plugged or fails to function properly, or if there is blanching or discoloration of the buttocks, heels, or toes, remove the catheter at once. Umbilical arteries are most easily cannulated in the first few hours of life but may provide a viable vascular route as late as 5 to 7 days after birth.

IO Catheterization (See also Chapter 25)

Compared to other forms of access, the IO line is rapid, cost-effective, and has a high success rate.36 In the simulation setting, IO lines were placed an average of 46 seconds more quickly than umbilical lines without a significant difference in adverse events for neonatal resuscitations.37 This delivery system makes use of the venous sinuses in the medullary cavity of the long bones to infuse fluids, medications, and blood products into the venous system. Potential sites include the flat portion of the proximal tibia (1 cm medial and 1 fingerbreadth below the tibial tuberosity), distal femur, medial malleolus (2 cm above), humeral head, distal radius, and ulna. The sternum is a potential IO site; however, its use is not recommended with most modern drill sets. In growing children, care must be taken to avoid the growth plate.38

Indications and Contraindications
IO access is recommended when peripheral access cannot be obtained safely at a pace that matches the illness of the child. Because the fluid can extravasate into the surrounding tissue causing a compartment syndrome, IO access should be avoided in a bone that is fractured or in which an IO line has been attempted within the previous 24 hours.39 A proximal vascular injury, bony or overlying infection, and osteogenesis imperfecta are also contraindications. Typically, the duration of use of an IO line is limited to 24 hours.39

Equipment and Setup
IO access can be achieved with the use of manual or mechanical puncture. The mechanical options include spring-loaded devices and reusable drills. In infants and children, both are successful, with more time required for manual insertion.39 If a reusable drill is used, different sized needles are available depending on the weight and habitus of the patient. Needle options for the EZ-IO (Teleflex, Morrisville, NC) include pink 15-mm (3 to 39 kg), blue 25-mm (>40 kg), and yellow 45-mm (>40 kg with excess subcutaneous tissue) needles. Typically, the pink needles are used for pediatric patients receiving tibial IO lines; however, consideration should be given to use of a longer needle if the line penetrates more soft tissue (e.g., femoral IO).

Technique
All IO lines should be performed with sterile technique. The proximal tibia is preferred in children. After sterile preparation, the needle is held at a 90-degree angle to the bone and inserted. If a manual IO needle is used, a rotary motion will ease passage through the bone, whereas the drills are held stationary. A slight “give” or “pop” signifies that the needle is through the first cortex and into the medullary space. It is imperative to stop at this point to prevent penetration out of the bone into the soft tissue. The needle should stand unassisted if it is firmly

![Figure 19.18](image-url)

Figure 19.18 The umbilical artery catheter makes a loop downward before heading cephalad (schematic drawing of a radiograph interpretation).
implanted in the bone. Often, bone marrow can be aspirated to confirm placement, and to send for laboratory analysis (excluding a complete blood count). Catheters are provided with some of the needles. The line should be flushed with 5 to 10 mL of saline while the provider’s hand is on the patient’s calf to confirm that fluid is not extravasating into the soft tissue.

In an awake patient, infusion of preservative-free lidocaine (typically, lidocaine intended for IV use in arrhythmias is available and appropriate for IO use) prior to the initiation of fluid can mitigate the pain associated with distension of the medullary cavity. The maximum dose is 20 mg to 50 mg, and children receive 0.5 mg/kg. Maximum gravity flow rates of 20 mL/min and pressure-driven rates of 40 mL/min are achievable, and standard resuscitation medications, including vasopressors, glucose, and blood products can be infused. Pushing fluid via a syringe or the use of a pressure bag may be necessary to allow an acceptable rate of delivery in a critically ill patient. Securing an IO line is difficult, but a simple solution is to surround the needle and proximal tubing with a small inverted paper cup with the bottom removed and the cup taped to the leg (Fig. 19.19) if a device to secure the line is not included with the device.

Complications
Complications of IO access are rare, and include cellulitis, osteomyelitis, bony fracture, fat or bone marrow embolism, and extravasation of fluid into the extraosseous tissue leading to compartment syndrome. If both cortices of a bone have been penetrated and fluid extravasates, a new bone must be selected. Clearly, use of the sternum may be associated with additional risks, due to the potential for trauma if both cortices of the sternum are penetrated.

Percutaneous Arterial Catheterization

Indications and Contraindications
Despite the growing use of noninvasive devices for monitoring transcutaneous oxygen and carbon dioxide, percutaneous peripheral artery catheterization may be indicated when there is a need for frequent blood gas sampling, continuous monitoring of arterial blood pressure, or both. However, mounting evidence supports minimizing the frequency of ABG sampling; therefore this is not an intervention that should be considered mandatory in the intubated intensive care unit patient. Arteries used for peripheral catheters in infants and children include the radial, ulnar, femoral, dorsalis pedis, and posterior tibial arteries. If preductal blood gas values are required, the right radial artery is used. This procedure is rarely performed in the ED.

Peripheral arterial catheterization is contraindicated when: (1) adequate peripheral artery samples can be obtained by percutaneous puncture, (2) the circulation of the extremity to be catheterized is compromised, (3) occlusion of the vessel to be catheterized compromises perfusion of the extremity, (4) there is an ongoing bleeding diathesis, (5) localized infection or inflammation overlies the artery to be cannulated, (6) intensive monitoring of line function is not available, or (7) vascular grafts are present in the extremity.

Equipment and Setup
The equipment needed for arterial catheterization is essentially the same as that required for percutaneous peripheral venous catheterization. Some centers use commercially available arterial line kits, which come with all the necessary supplies and equipment. Alternatively, standard 22- or 24-gauge over-the-needle IV catheters, a T-piece connector, and a three-way stopcock can be used. Connect the T-piece and the stopcock and then fill them with NS solution. Prepare an infusion pump with heparinized saline.

Technique
Perform the procedure with good lighting and an adequate work area and monitor the infant’s heart rate and respiratory rate closely. Palpate the radial artery proximal to the transverse wrist crease on the palmar surface of the wrist, medial to the styloid process of the radius. Before the procedure, compress the artery and observe the hand and fingers for change in color. If blanching or cyanosis is noted (indicating poor collateral circulation), do not perform catheterization. If locating the artery by palpation is difficult, a transillumination device, Doppler probe, or ultrasound may be helpful. Use of ultrasound improves success rate over Doppler alone.

Secure the infant’s or child’s hand and lower part of the forearm to an arm board, with the wrist dorsiflexed 45 to 60 degrees by placing a roll of gauze underneath it. Take care to leave the fingers exposed to allow assessment of the peripheral circulation. Palpate the radial artery at the point of maximal impulse and mark it by making a gentle indentation with a gloved fingernail. Prepare the area over the radial artery with povidone-iodine or another antiseptic solution and wash it with alcohol. Use a topical or local anesthetic (such as 1% lidocaine without epinephrine), or both, at the planned insertion site. Insert the catheter with the needle through the skin just proximal to the transverse wrist crease at a 10- to 20-degree angle (Fig. 19.20A). With the bevel of the needle facing proximally, advance the catheter slowly until blood appears in the catheter hub, which signifies puncture of the anterior arterial wall. Slowly advance the catheter until blood appears in the needle and then carefully lower the angle of the needle to approximately 10 degrees. Slowly advance the catheter over the needle and into the lumen of the artery. Remove the needle. Attach the stopcock and T-piece connector to the catheter hub. Open the stopcock to the syringe to confirm return of pulsatile blood. Flush it with 0.5 mL of heparinized solution.

Figure 19.19 Intraosseous line protected by a paper cup.

Intraosseous line protected by a paper cup.
monitored. Additionally, concerns regarding the type of flush used in the arterial line and the impact on results of blood samples drawn from that line must not be underestimated. Normoglycemic patients have received insulin therapy based on falsely elevated glucose readings in samples taken from an arterial line flushed with dextrose-containing fluid.67,68 When decannulating the artery, apply pressure over the site for 3 to 5 minutes following removal to minimize bleeding.

REHYDRATION TECHNIQUES IN INFANTS AND CHILDREN

Approach to Dehydration

The literature on clinical assessment of the degree of dehydration in children with acute gastroenteritis is inconsistent. Physician gestalt alone typically performs poorly, with a reported sensitivity of 42%.69 Parental report of symptoms associated with dehydration (emesis, diarrhea, poor fluid intake, decreased urine output, weak cry, sunken fontanelle, sunken eyes, decreased tears, dry mouth, and cool extremities) is sensitive, but not specific. Table 19.2 presents a more thorough list of the signs, symptoms, and definitions of mild, moderate, and severe dehydration. Different dehydration scales exist to improve assessment, with a wide range of accuracy. A 2015 paper suggests a simpler model in which sunken eyes, decreased skin elasticity, weak radial pulse, and general appearance are used, with two factors being predictive of at least 5% dehydration and three or four factors predictive of 10% dehydration.70 When decannulating the artery, apply pressure over the site for 3 to 5 minutes following removal to minimize bleeding.

Radial Artery Catheterization

A B

Introduce the catheter assembly into the radial artery through the skin at a 10° to 20° angle. This is a smaller angle than that used for simple arterial puncture.

One technique of taping the arterial catheter. The arm board should be well padded and secured.

Figure 19.20 Radial artery catheterization. (See Fig. 20.5 for additional details on this technique.)

Complications

Complications, which have been reported with every type of arterial catheter, include hemorrhage, thrombosis, spasm, infection, scars, air embolism, retrograde blood flow, transient elevation in blood pressure with rapid (<1 second) infusion, arteriovenous fistula formation, and nerve or tendon sheath damage. There is the potential for loss of digits, an entire extremity, or large areas of skin, and perfusion must be carefully monitored.
Oral Rehydration

Oral rehydration therapy (ORT) has been shown to be equivalent to IV therapy in children with acute gastroenteritis in terms of rehydration, subsequent diarrheal episodes, and the development of sodium abnormalities. Failure rates are 4.9% as compared with 1.3% for IV therapy, and use of ORT can avoid the need for IV hydration in 24 of every 25 patients. This is the modality of choice for rehydration of a mildly to moderately dehydrated infant or child aged 1 month or older. Oral rehydration solutions (ORSs) come in a number of commercial forms (World Health Organization rehydration salts, Enfalyte [Mead Johnson Nutrition, Chicago, IL], Pedialyte [Abbott, Abbott Park, IL], Rehydralyte [Abbott], CeraLyte [Cera Products, Hilton Head, SC]). The ideal composition is 50 mEq/l of sodium, 25 g/l of dextrose, and 30 mEq/l of bicarbonate, and a solution can be made at home with five cups of water, eight teaspoons of sugar, and one teaspoon of salt. A JAMA paper came out this year showing that rehydration of mildly dehydrated patients with dilute apple juice instead of oral rehydration solution resulted in fewer treatment failures, and that is another option. The target volume is 50 to 100 mL/kg over a period of 2 to 4 hours. Typically, this is started with a spoon, cup, or syringe delivering 5 mL to children younger than 2 years and 10 mL to children older than 2 years every 5 minutes and then increasing volumes slowly as tolerated (Fig. 19.21). Vomiting is not a contraindication to attempting ORT because children can typically tolerate these small volumes. Some practitioners fear accompanying electrolyte abnormalities; however, hypotonic dehydration responds well to ORT.

Oral rehydration can be facilitated with the use of an antiemetic. Ondansetron has been shown to decrease ED vomiting (number needed to treat [NNT] = 5), decrease the need for IV hydration (NNT = 5), and possibly decrease the need for admission (NNT = 14). Ondansetron is administered at a dose of 0.15 mg/kg in the form of syrup or oral dissolving tablets. An alternative, simplified dosing regimen is to give 2 mg for children 8 to 15 kg, 4 mg for children 15 to 30 kg, and 8 mg for children greater than 30 kg. Oral rehydration should be initiated 15 to 30 minutes after administration of ondansetron. Most oral ondansetron studies include children 6 months and older, but several studies using IV ondansetron have included infants as young as 1 month. Concern regarding the possibility of ondansetron delaying the identification of more pathologic diagnoses is unfounded. A metaanalysis has shown no difference in return visit rates in those who do not receive antiemetic therapy, and it has been shown that antiemetic therapy does not mask alternative diagnoses. Outpatient prescriptions for ondansetron do not affect recidivism or readmission rates. Ondansetron has received attention for its ability to prolong the QT interval. In children, it lengthens

### Table 19.2 Estimating Dehydration

<table>
<thead>
<tr>
<th>DEHYDRATION</th>
<th>SYMPTOMS</th>
<th>SIGNS</th>
<th>FLUID LOST IN INFANTS</th>
<th>FLUID LOST IN OLDER CHILDREN</th>
<th>TREATMENT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild</td>
<td>Thirsty</td>
<td>Slightly dry mucous membranes</td>
<td>50 mL/kg</td>
<td>3–5% total fluid</td>
<td>ORS, 50–100 mL/kg over a 2- to 4-hr period</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Consider other methods if not tolerated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>No laboratory studies</td>
</tr>
<tr>
<td>Moderate</td>
<td>Decreased urine output</td>
<td>Absent tears, Tachycardia, Capillary refill &gt;2 sec, Dry mucous membranes, Weak pulse, Abnormal respirations, Sunken fontanelle, Hypotension</td>
<td>100 mL/kg</td>
<td>6–9% total fluid</td>
<td>ORS, 50–100 mL/kg over a 2- to 4-hr period</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Consider other methods if not tolerated</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>No laboratory studies</td>
</tr>
<tr>
<td>Severe</td>
<td>Abnormal mental status, lethargy</td>
<td>Abnormal skin turgor, Sunken fontanelle, Hypotension</td>
<td>150 mL/kg</td>
<td>&gt;9% total fluid</td>
<td>20–60 mL/kg of NS by IV bolus</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Serum chemistries</td>
</tr>
</tbody>
</table>

IV, Intravenous; NS, normal saline; ORS, oral rehydration solution.
the duration of the QT interval by 10 to 17 msec, and is unlikely to induce torsades de points in an otherwise healthy child.\(^7^8\) It may contribute to mild and self-limited diarrhea.\(^7^6\) Other antiemetics and antimotility agents are not recommended by the American Academy of Pediatrics for use in children younger than 5 years, and they have a high incidence of side effects. There is a Food and Drug Administration “boxed warning” against the use of promethazine in children younger than 2 years and a strong recommendation against its use even in older children. For patients with diarrhea, the antisecretory agent rabeprazol is used in several European countries in conjunction with ORS, with a favorable treatment and adverse event profile.\(^7^9\)

**Laboratory Tests**

The American Academy of Pediatrics recommends electrolyte testing in patients with severe dehydration, those who need IV rehydration, or children with moderate dehydration and findings on the history or physical examination inconsistent with straightforward diarrheal episodes. A prospective study of patients 2 months to 9 years of age, in whom parenteral rehydration was deemed required on clinical grounds, demonstrated a 48% rate of laboratory abnormalities, 10.4% of which changed treatment management. More than 9% of their patients had hypoglycemia, so it is recommended to obtain a glucose level urgently, as well as electrolytes when feasible, in patients who require IV rehydration.\(^4^0\) It is difficult to provide recommendations regarding the management of low bicarbonate because the level has been found to correlate poorly with the need for admission and unscheduled revisit rate in some studies,\(^8^1\) whereas it correlates well in others.\(^8^2\) Urine specific gravity tends to lag behind the child’s hydration status and is particularly unreliable in infants younger than 1 year. Blood urea nitrogen levels are also poorly sensitive in young children,\(^8^3\) but significant elevations in these levels and in creatinine may be specific.\(^8^4\) Serum ketones have been shown in one study to correlate with clinical dehydration scores.\(^8^5\)

**Parenteral Rehydration**

Obtaining IV access is covered elsewhere in this section. If parenteral administration of fluids is deemed appropriate, standard therapy involves 20-mL/kg boluses of NS or lactated Ringer’s solution infused over a 60-minute period each until normalization of pulse, perfusion, and mental status occurs. Recent literature has demonstrated comparable efficacy of rapid rehydration protocols involving the administration of 50 to 60 mL/kg over a 60-minute period in otherwise healthy children,\(^8^5\) though it may paradoxically lengthen time to discharge. Adverse events were minimal in the studies; however, they were not entirely powered to ensure safety.\(^8^6\) A handful of small studies have looked at bolusing with dextrose-containing fluids. This practice seemed safe and decreased ketone levels, but did not significantly decrease the admission rate.\(^8^7\)

Children who have received parenteral fluids and subsequently tolerated oral fluids can typically be discharged home from the ED, although an unscheduled revisit rate of 16% has been reported.\(^8^8\) If the child requires admission for continuance of maintenance fluids, isotonic saline is recommended because of a documented 17% to 45% risk for hyponatremia with the use of hypotonic fluid.\(^8^9\) The amount of maintenance fluid is calculated as 100 mL/kg for the first 10 kg of body weight, 50 mL/kg for 10 to 20 kg of body weight, and 20 mL/kg for the remaining body weight in a 24-hour period. For example, a 50-kg child would require (100 mL × 10 kg) + (50 mL × 10 kg) + (20 mL × 30 kg) divided evenly over 24 hours.

**Nasogastric Tube Rehydration**

Nasogastric (NG) rehydration is a potentially effective, safe, and cost-effective alternative to parenteral rehydration in children who refuse to take fluids orally in the quantity required for rehydration. In developing nations, NG hydration has been shown to be effective for severe dehydration as well.\(^9^5\) NG rehydration is accomplished by placement of an NG tube, with a 5-Fr feeding tube typically being adequate for children younger than 3 years. For comfort, the use of topical anesthetic or nebulized lidocaine (5 mg/kg diluted in NS) before insertion is recommended. A bolus of 50 mL/kg of electrolyte solution (e.g., Pedialyte [Abbott]) is administered by continuous infusion via an enteral feeding pump over a 3-hour period. Twenty-four-hour rehydration regimens (50 to 100 mL/kg for the first 10 kg of body weight and then 25 to 50 mL/kg for the remaining body weight) have also been proposed but seem to have equivalent outcomes to more rapid rehydration protocols.\(^9^6\) Unscheduled revisit rates are not significantly different compared with patients who have received IV rehydration. Additionally, NG hydration has been used for infants with bronchiolitis with hospital lengths of stay comparable to those receiving IV hydration, and also with better placement success.\(^9^7\) No study has evaluated the use of frequent, small boluses; however, experience with ORT would indicate that 5- to 10-mL parenterally administered syringe boluses to equal 50 mL/kg would be an option in the event that a pump for continuous administration is unavailable.

**Subcutaneous Rehydration**

Subcutaneous rehydration therapy, also known as hypodermolysis, is an option in children with mild to moderate dehydration in whom IV or IO access is not available or feasible. Its utility as a bridge to facilitating commencement of IV hydration in severely dehydrated children has been postulated, but not studied. Although the potential has been recognized since the early 1900s, the hyaluronan-containing intercellular matrix of subcutaneous tissue prevents easy delivery and absorption of large quantities of fluid, and the previous preparations of hyaluronidase (enzymes that hydrolyze hyaluronan) were animal based and allergenic. More recently, a recombinant formulation of human hyaluronidase, HuPH20 (Hylenex), has become available, thus making this technique more accessible. Its primary utility has been in geriatric and hospice patients requiring hydration, but several studies have substantiated its safety and efficacy in pediatric patients as young as 2 months, with a mean ED volumes deliverable comparable to IV hydration. In assessing recidivism, pediatric patients treated with subcutaneous hydration and discharged from the ED had an unscheduled revisit rate comparable to those receiving IV hydration.\(^9^8\) The cost-effectiveness profile is advantageous when compared with equivalent IV therapy, with children below 3 years of age representing the most difficult cases for IV access and, thus, the greatest cost savings.\(^9^9\)

After sterile preparation, a 22- or 24-gauge angiocatheter or 25-gauge butterfly needle is inserted at a 30- to 45-degree
angle to the hub into subcutaneous tissue. After placement of a small piece of gauze under the hub of the catheter to maintain the angle, it is secured with a sterile occlusive dressing. Although the interscapular area is often used, the thigh, abdomen (left iliac fossa), chest, and upper part of the arm are also options. If possible, an area with a pinchable fat fold of 1 inch should be selected. The catheter should be aspirated for absence of blood return. A 150-unit dose of hyaluronidase is injected through the catheter, and the fluid infusion is started. Typically, isotonic fluids such as 0.9% NS are used. Mean fluid infusion rates are highly variable in adults and range in children from 18.9 mL/kg over the first hour to 38.4 mL/kg for a 4-hour infusion. An event infusion rate of approximately 20 mL/kg per hour should be selected and titrated up. The infusion pump indicates occlusion. Because the hyaluronidase requires approximately 15 minutes to take full effect, some providers prefer to start more slowly and titrate upward. Initially, an erythematous swelling may appear at the infusion site (Fig. 19.22), but it will decrease in size after 5 minutes and disappear 1 to 2 hours after cessation of the fluid infusion. Some pain may occur at the infusion site. Suggestions for troubleshooting subcutaneous infusion problems can be found in Table 19.3. Maximal diffusion of fluid into the IV compartment occurs in 1 hour. An injection of hyaluronidase allows subcutaneous infiltration for 48 hours, and maintenance fluids, including those containing dextrose and potassium, can be infused through the catheter after the bolus if required. No special care is required following removal of the catheter.

Subcutaneous rehydration is not recommended in patients with signs of shock, coagulation deficits, gross edema, significant electrolyte abnormalities (sodium >150 mEq/L or <130 mEq/L), or no intact skin sites (e.g., massive burns). The most common adverse events reported are redness, swelling, or bruising at the infusion site. Allergic reactions have not been reported with the newer formulation of hyaluronidase, and skin infections are rare.

Discharge

Commonly held beliefs include avoidance of dairy products, need for formula change or overdilution of formula with water, gut rest for 24 hours or longer, and the BRAT diet (bananas, rice, applesauce, toast). These beliefs are not supported by the literature in most cases. Once rehydration has been achieved, returning to a normal diet is suggested as soon as possible, with continued replacement of ongoing losses with an ORS or potentially dilute juice.

Most children seen in the ED with acute dehydration will be corrected and discharged home. Admission criteria include inadequate care at home or inability to follow return precautions, inability to take or failure of rehydration with ORT, concern for more serious pathology, severe dehydration, and neonatal dehydration. Probiotics can help reduce the duration of diarrhea in children with gastroenteritis.

Table 19.3 Troubleshooting Subcutaneous Rehydration Complications

<table>
<thead>
<tr>
<th>The catheter becomes dislodged</th>
<th>Restart in the same area without an additional dose of hyaluronidase</th>
</tr>
</thead>
<tbody>
<tr>
<td>The infusion pump indicates occlusion</td>
<td>Slow the infusion rate by 10 mL/hr</td>
</tr>
<tr>
<td>The area becomes firm, indurated, and painful</td>
<td>Administer an additional dose of hyaluronidase</td>
</tr>
<tr>
<td>The child fails outpatient therapy</td>
<td>Continue infusion of fluid through the catheter for 48 hours as maintenance</td>
</tr>
</tbody>
</table>

Table 19.3 Troubleshooting Subcutaneous Rehydration Complications

---

**Table 19.3** Troubleshooting Subcutaneous Rehydration Complications

<table>
<thead>
<tr>
<th>The catheter becomes dislodged</th>
<th>Restart in the same area without an additional dose of hyaluronidase</th>
</tr>
</thead>
<tbody>
<tr>
<td>The infusion pump indicates occlusion</td>
<td>Slow the infusion rate by 10 mL/hr</td>
</tr>
<tr>
<td>The area becomes firm, indurated, and painful</td>
<td>Administer an additional dose of hyaluronidase</td>
</tr>
<tr>
<td>The child fails outpatient therapy</td>
<td>Continue infusion of fluid through the catheter for 48 hours as maintenance</td>
</tr>
</tbody>
</table>

---

**REFERENCES ARE AVAILABLE AT www.expertconsult.com**
REFERENCES


Arterial puncture is the most accurate blood sampling technique for true arterial blood gas (ABG) and acid-base determination. The absence of arterial blood pressure defines cardiac arrest and serves as a definitive end point for resuscitative efforts. Intraarterial cannulation with continuous blood pressure measurement remains an accepted standard in critically ill patients. Intraarterial monitoring of blood pressure better reflects the force of systemic perfusion and is one of the most important determinants of cardiac work. In recent years, noninvasive technologies have achieved an accuracy that is nearly equal to that of invasive monitoring, but these techniques also have limitations. Invasive modalities require specific expertise and support to perform.

**INDICATIONS AND CONTRAINDICATIONS**

Indications for and contraindications to arterial puncture and cannulation are listed in Review Box 20.1. The use of arterial lines for continuous monitoring is generally reserved for the intensive care setting; however, arterial cannulation may be initiated in the emergency department (ED). The indications for placement of an arterial catheter fall into two major categories:

1. Repetitive and direct arterial blood sampling. Catheter access removes the need for multiple arterial punctures and allows either repeated sampling or placement of sensors for continuous monitoring of blood gas and other chemistry values.

2. Continuous real-time monitoring of blood pressure. Catheter access allows superior monitoring and moment-to-moment detection of changes. Intraoperative and intensive care unit (ICU) management is often facilitated by placement of an arterial line. Some patients, such as those with severe burns, dialysis grafts or shunts, or morbid obesity, may need ongoing monitoring of perfusion, which can best be accomplished by arterial catheterization.

---

**Review Box 20.1** Arterial puncture and cannulation: indications, contraindications, complications, and equipment.
Although acute respiratory decompensation and metabolic emergencies are the most common reasons for ABG sampling, all blood tests performed on venous blood are also possible on an arterial sample. Cultures performed on blood obtained from an indwelling arterial line have a sensitivity and specificity similar to that of cultures performed on blood obtained from a venipuncture site. Patients with moderate respiratory decompensation may be managed without arterial puncture by using continuous, noninvasive pulse oximetry, end-tidal or transcutaneous carbon dioxide monitoring, carboxyhemoglobin and methemoglobin monitoring, or any combination thereof. Nonetheless, a role still exists for arterial blood sampling. The initial correlation between noninvasive values and acid–base status via arterial sampling is often important in critical illness to set a baseline or verify a trend. Some authors use ABG sampling in the initial evaluation of critically ill trauma patients. Vasoactive drugs (e.g., nitroprusside and norepinephrine) are best administered with continuous monitoring of arterial pressure to guide titration. The response of trauma and post-cardiac arrest patients to acute resuscitative efforts may also be more easily monitored with the use of arterial catheterization.

Few contraindications to arterial puncture exist; none are absolute. For example, after thrombolysis, arterial cannulation should be performed only if it will provide essential data that cannot be obtained by any other method. If absolutely necessary, a single arterial puncture of the readily compressible radial artery is preferred. Arterial puncture can be performed safely in patients who are anticoagulated or who have other coagulopathies, but it should be undertaken with extreme caution in patients with severe disseminated coagulopathies.

There are reports of patients with bleeding complications who require transfusion. Some patients have suffered compression neuropathies secondary to hematomas at the puncture site. Repeated arterial sampling in such patients should be accomplished by insertion of an indwelling cannula to minimize trauma to the arterial wall.

The presence of severe arteriosclerosis, with or without diminution in flow, is a relative contraindication to arterial puncture. In hemodynamically unstable patients with advanced cardiovascular disease, the benefits of invasive monitoring may nonetheless outweigh its risks. Consider an alternative site if an isolated, decreased palpable pulse or bruit is felt over the site selected. Additionally, consider an alternative site if there is evidence of decreased or absent collateral flow in areas where flow normally exists, such as in Raynaud’s syndrome or an abnormal result on the modified Allen test (discussed later in the section on Techniques). Avoid puncturing a specific arterial site when infection, burn, or other damage to cutaneous defenses exists in the overlying skin or through or distal to a surgical shunt.

Arterial Versus Venous Analysis

Arterial sampling has been the traditional approach to evaluating acid-base abnormalities in critically ill patients, especially those being maintained on a ventilator. In most ED settings, however, venous blood gas analysis will suffice. Studies have demonstrated that analysis of venous blood (especially central venous blood) for pH, bicarbonate, lactate, base excess, and carbon dioxide pressure (PCO₂) are within 95% limits of agreement with arterial sampling and can safely supplant it. On the other hand, arterial blood sampling is still required for accurate analysis of oxygen pressure (PO₂).

EQUIPMENT: ARTERIAL PUNCTURE

Arterial Puncture With a Needle/Syringe

To obtain a single sample of arterial blood by the percutaneous method, attach a 3-mL syringe (preferred and most common) to a needle. Select the needle size based on puncture location and patient size and age. For an adult, use a 20-gauge, 2.5-inch needle for a femoral sample and a 22-gauge, 1.25-inch needle for a radial artery puncture. For pediatric arterial sampling, use a needle with a slightly shorter length in the range of 22- to 24-gauge at the same sites as in adults.

Precoated blood gas plastic syringes (with dry lithium heparin) are commonly used and allow a longer shelf life and ready use (Fig. 20.1). Such devices are designed to minimize sampling error as a result of heparin. If necessary, prepare a regular syringe with 1 or 2 mL of a heparinized saline solution (1000 International Units [IU]/mL) drawn into the syringe to coat the barrel and needle. Fully eject the heparin through the needle immediately before skin puncture to minimize heparin-related errors. Although the syringe may appear devoid of heparin, enough heparin remains in the needle and syringe to provide anticoagulation. Even dry heparin may produce abnormalities in ABG results because of a heparin-induced dilutional effect.

The latest blood gas and chemistry analyzers require only 0.2 mL of whole blood for accuracy, and some point-of-care devices can perform analyses on single drops of blood. However, sample sizes of less than 1.0 mL of blood aspirated into heparin-coated syringes may result in a heparin-related error on ABG values.

Stored heparin solution has higher PO₂ and lower PCO₂ values than blood does. A dilutional effect from heparin would mean that the addition of 0.4 mL of heparin solution to a 2-mL sample of blood (dilution of 20%) will lower PCO₂ by 16%. Proper technique with dry lithium heparin-prefilled syringes or full ejection of excess heparin will prevent such problems if more than 2 mL of blood is collected. A falsely
low \(\text{PCO}_2\) is the most clinically significant change caused by excess heparin.\(^{14,15}\) Neither \(\text{PO}_2\) nor pH levels are significantly altered by the addition of heparin in most instances, although a slight increase in \(\text{PO}_2\) and a minimal decrease in pH may occur if high concentrations of heparin (25,000 IU/mL) are used.\(^{16}\) If 2 to 3 mL of blood is collected, heparin-related effects are likely to be clinically inconsequential.

### Continuous Monitoring via Arterial Catheter

The fluid-filled recording systems used with arterial cannulation have a great influence on the accuracy of pressure measurements. The frequency responses of tubing, transducers, and other components of the monitoring system influence the accuracy of systolic and diastolic pressure measurement. Failure to recognize recording system artifacts will lead to errors in interpretation of the pressure.

Various catheter types have demonstrated similar frequency-response characteristics, but some studies have found different complication rates. Teflon catheters may carry an increased rate of thrombosis.\(^{17,18}\) Another contributing element leading to thrombosis is catheter diameter; the incidence of thrombosis is inversely related to the ratio of vessel lumen to catheter diameter.\(^{19,20}\) Thus the risk for thrombosis increases as the diameter of the catheter decreases. The incidence of thrombosis also increases with increased duration of catheter placement. In contrast, a higher risk for thrombosis was seen in the femoral artery than in the radial artery in a study involving a pediatric population.\(^{21}\) Catheters coated with a combination of chlorhexidine and silver sulfadiazine have produced lower infection rates.\(^{22}\)

### Preparation for Arterial Cannulation

Box 20.1 lists the usual equipment for arterial cannulation, although the majority of prepackaged kits contain the supplies most needed (see also Review Box 20.1). Shorter catheters are ideal for peripheral artery cannulation, whereas use of a longer catheter and the Seldinger technique is preferable for the femoral artery.

For arterial cannulation in adults, use a 16- to 18-gauge catheter for the femoral artery and a 20-gauge catheter for the radial artery (Fig. 20.2). Small children and infants require a 22- to 24-gauge catheter, which may need to be inserted percutaneously via the Seldinger technique or through a femoral cutdown. Based on patient size, older pediatric patients usually require 20- to 22-gauge catheters.

The tubing that connects the catheter to the pressure transducer has a significant effect on the accuracy of the monitoring system. The higher the frequency response of the entire system, the more accurate the determination of systolic and diastolic pressure; however, artifact also becomes more of a problem.\(^{1,23}\) Use stiff, low-capacitance plastic tubing for arterial catheterization and monitoring. Place the electronic pressure transducer connection as close as possible to the patient and zero it appropriately because the frequency response of a tube is inversely related to its length.\(^{24,25}\)

The pressure wave produced with each contraction is transmitted from the artery through the catheter and connecting tubing to a measuring device. The arterial fluid wave is received by an electromechanical transducer that changes the mechanical pressure wave into an electrical signal that can be displayed on the monitor. The most basic technique for obtaining blood pressure values involves the use of a simple manometer.\(^{26}\) This system can be assembled quickly if the material is available.

A continuous method of flushing the pressure tubing is required to maintain patency of the catheter lumen during intraarterial pressure monitoring. A three-way stopcock through which the tubing is intermittently flushed with saline (a minimum of every 15 to 30 minutes) is a simple, effective method. Continuous flush devices push a set amount of fluid.
SITE SELECTION

The radial, brachial, and femoral arteries are the sites usually punctured for blood gas sampling in adults (Video 20.1). (usually 2 to 3 mL/hr) through the line.16 A typical monitoring system that includes this device is shown in Fig. 20.3. The pressure transducer must be mounted at the level of the patient’s heart. Current pressure-monitoring setups include not only built-in stopcocks but also in-line flushing plungers to facilitate clearance of blood after sampling.

Intravascular transducers were initially seen as an improvement over the external electromechanical transducers in use since the mid-1970s. Many of the numerous brands are fragile, temperature sensitive, of variable quality, and much more difficult to place in vessels than catheters are. Despite anecdotal reports of fibrin deposition on these devices, no complications of arterial catheterization is bleeding, the ability to control hemorrhage must also be considered. For these reasons, the radial and femoral arteries are favored because of their good collateral blood flow and ease of compression in case of hemorrhage. Patient comfort and nursing care concerns should also be considered during selection of the site.

Pediatric sites commonly used for arterial puncture include arteries in the foot and the umbilical artery in newborns.

When an artery is cannulated for longer-term use, there is a risk of complete loss of blood flow through a vessel as a result of intraluminal thrombosis. This is important when choosing a site for arterial puncture. Because the most frequent complication of arterial catheterization is bleeding, the ability to control hemorrhage must also be considered. For these reasons, the radial and femoral arteries are favored because of their good collateral blood flow and ease of compression in case of hemorrhage. Patient comfort and nursing care concerns should also be considered during selection of the site.

**TABLE 20.1 Parameters That Affect Interpretation of Arterial Blood Gases**

<table>
<thead>
<tr>
<th>PARAMETER</th>
<th>HEPARIN</th>
<th>AIR BUBBLE</th>
<th>DELAYED ANALYSIS</th>
</tr>
</thead>
<tbody>
<tr>
<td>PO₂</td>
<td>No significant change</td>
<td>Elevated</td>
<td>Variable</td>
</tr>
<tr>
<td>PCO₂</td>
<td>Lowered</td>
<td>No significant changes</td>
<td>Elevated</td>
</tr>
<tr>
<td>pH</td>
<td>Unchanged</td>
<td>No significant changes</td>
<td>Lowered</td>
</tr>
</tbody>
</table>

*aUse only a 1000-IU/mL concentration. Fill the dead space of the needle and syringe only and collect 3 mL of blood.

*bChanges unpredictable at 20 minutes regardless of the storage method.

*cThere are reports of slight increases in PO₂ with excessive heparin.

*dThe falsely lowered PCO₂ that occurs with added heparin is the most clinically significant change noted. pH may be decreased if a large volume of concentrated heparin (25,000 IU/mL) is used.

*eIf stored at 4°C for 20 minutes. Anaerobic storage at room temperature for 20 minutes results in no significant change.

*fMinimal changes up to 2 hours if stored at 4°C.

**FIGURE 20.3** Arterial pressure monitoring systems. A, System for continuous flushing. A 1-L bag of normal saline is pressurized to 250–300 mm Hg with a metered blood pump (not shown). The continuous flush device is set to deliver 3 mL/hr of saline. A mechanical pressure transducer is depicted. The transducer device is a sterile, inexpensive, fully assembled monitor that can be used during patient transfer. Alternatively, the electronic transducer depicted in B may be used. B, System for manual flush. A saline flush solution can be injected manually through a syringe at the proximal or distal port. The transducer dome should be maintained at the level of the patient’s heart. (From Beal JM, editor: Critical care for surgical patients, New York, 1982, Macmillan. Reproduced by permission.)

**TECHNIQUES**

**Arterial Puncture**

Palpate the arterial pulse to ascertain the location of the vessel and prepare the overlying skin with an antiseptic solution (Fig. 20.4, step 1). Anesthetize the patient’s skin with a wheal of local anesthetic (e.g., 1% lidocaine without epinephrine) through a small needle (25- or 27-gauge) (see Fig. 20.4, step 2). If local anesthesia is to be given, take care to use only a small amount of local anesthetic because a large wheal may obscure the pulse. One study found no significant alterations in PCO₂ or pH from the pain or anxiety of an unanesthetized arterial puncture (Table 20.1).33 If the patient is in extremis or unresponsive to pain in the area to be punctured, anesthetic infiltration may be omitted.

Isolate the arterial pulsation with the index and middle fingers of the gloved, nondominant hand and identify the course of the vessel. Puncture the skin through the anesthetic wheal, immediately distal to the palpated pulse under the index finger (see Fig. 20.4, step 3). The older technique of placing the needle between the index and middle finger risks self-puncture and is no longer advised. Hold the syringe like a dart with the bevel up and the syringe kept in view so that blood flow can be seen immediately. Advance the needle slowly toward the
Arterial Puncture (Radial Artery)

1. Position the wrist in slight dorsiflexion, cleanse the skin with antiseptic solution, and palpate the radial pulse.

2. Optionally, place a small wheal of local anesthetic (e.g., 1% lidocaine without epinephrine) over the entry site. Avoid placing too large of a wheal, which may obscure the artery.

3. Hold the syringe in your hand like a dart, with the bevel up. Palpate the artery with the index and middle fingers of your other hand. Puncture the skin distal to your finger, and slowly advance the needle at a 30° angle toward the pulsating vessel.

4. As soon as blood flows, stop advancing the needle. Allow the syringe to fill on its own. If bone is encountered, withdraw slowly because both vessel walls may have been penetrated and the lumen may be entered as the needle is withdrawn.

5. Remove the needle from the artery after the syringe has filled. Apply a bandage and firm pressure to the puncture site for a minimum of 3 to 5 minutes.

6. Remove all air from the syringe by holding it upward, gently tapping it, and depressing the plunger. Attach the end cap to the syringe to maintain anaerobic conditions, and submit the sample to the laboratory.

Figure 20.4 Arterial puncture (radial artery).
pulsating vessel at an approximately 30-degree angle. A larger angle is required to puncture the deeper femoral artery. Once the needle enters the arterial lumen, allow the syringe plunger to rise with the arterial pressure on its own to discriminate between arterial and venous sampling (see Fig. 20.4, step 4). As soon as blood flows, stop advancing the needle and allow the syringe to fill. If no blood flow is obtained or if bone has been hit, withdraw the needle slowly because both walls of the vessel may have been punctured and the lumen may be entered as the needle is withdrawn. Redirect the needle only when the needle has been retracted to a location just deep to the dermis. After at least 1 to 2 mL of blood has been obtained, remove the needle from the artery. Apply firm pressure at the puncture site for a minimum of 3 to 5 minutes (see Fig. 20.4, step 5). If the patient is anticoagulated or has a coagulopathy, 10 to 15 minutes of pressure is required (Videos 20.2, 20.3, and 20.4).

Use of ultrasound is rapidly becoming a standard in procedures involving central vascular access. The same ultrasound-guided techniques are also being applied to arterial cannulation (Ultrasound Box 20.1.) Shiloh and colleagues reported 71% of blood has been obtained, remove the needle from the artery. Apply firm pressure at the puncture site for a minimum of 3 to 5 minutes (see Fig. 20.4, step 5). If the patient is anticoagulated or has a coagulopathy, 10 to 15 minutes of pressure is required (Videos 20.2, 20.3, and 20.4).

Use of ultrasound can be used with great success in the placement of peripheral and central intravenous lines, and it can also be used for the placement of arterial catheters.

First, identify the target vessel with ultrasound. Use a high-frequency transducer (10 to 12 mHz) to ensure proper resolution. The target vessel is typically easiest to identify in the transverse plane (with the indicator pointing toward the patient’s right side) (Fig. 20.US1). Arteries usually appear as rounded structures with thick walls, as opposed to veins which are ovoid. Depending on the size of the vessel, pulsations may be noted in the arteries. Application of color flow Doppler to a suspected artery will frequently demonstrate a pattern of pulsatile flow (Fig. 20.US2). Although arteries are usually thought to resist collapse from outside pressure (such as from applying pressure to the overlying transducer), smaller arteries in the periphery may show a degree of collapse. When seeking to identify the target vessel, multiple means of identification should be used before an attempt at cannulation.

Once the vessel has been identified, ultrasound can also be used to directly guide cannulation. Sterile technique should be maintained, particularly when accessing central arteries such as the femoral artery. A full description of the technique of ultrasound-guided venous access can be found in both the general ultrasound chapter (Chapter 66) and the central venous access chapter (Chapter 22) in this textbook. The principles of these procedures also apply to arterial cannulation, although a few points should be emphasized. When guiding access to the radial artery, a transverse approach is typically used because the vessel is small and difficult to visualize in the longitudinal approach. Additionally, it may be difficult to keep the image of the longitudinal vessel centered on the screen, especially when one physician is directing both the ultrasound and the needle.

Air in the sample will significantly increase \( \text{PO}_2 \) (mean increase, 11 mm Hg) after 20 minutes of storage, even if kept at \( 4^\circ \text{C} \). pH and \( \text{PCO}_2 \) are not significantly altered by air bubbles if the blood is stored at \( 4^\circ \text{C} \) for 20 minutes and no significant deterioration has occurred. \( 16,31 \) If blood is stored at room temperature for longer than 20 minutes, \( \text{PCO}_2 \) will increase.

Proper handling of the sample and rapid analysis are very important. When the needle is withdrawn, expel any air bubbles present in the syringe to avoid a false elevation in \( \text{PO}_2 \). \( 31 \) Remove the air neatly and easily by tapping the inverted syringe (needle pointing upward) to force any air to the top; then carefully and slowly depress the syringe plunger to push out the remaining air (see Fig. 20.4, step 6). A gauze pad or alcohol wipe may be used to collect any excess blood expelled with the syringe held upright and the plunger side down. Remove the needle and cap the syringe to ensure anaerobic conditions. Alternatively, many of the commercially available ABG kits come with an air bubble removal device (Filter-Pro [Smiths Medical, St. Paul, MN]) that allows the clinician to expel air bubbles from the sample and reduce potential exposure from the blood product.
and the pH will decrease, probably as a result of leukocyte metabolism. In a stored sample, PO₂ varies to such an extent that the change is unpredictable for chemical interpretation at 30 minutes, regardless of the storage method. High leukocyte or platelet counts, such as those seen in leukemic patients, may shorten acceptable storage intervals.12,13

Thus, ABG samples should always be kept on ice and analyzed within 15 to 20 minutes. Samples that cannot be analyzed within this time frame should be considered unreliable.

PERCUTANEOUS TECHNIQUE FOR ARTERIAL CANNULATION

Direct Over-the-Needle Catheter Cannulation

Placement of an angiocatheter directly into an arterial lumen in a manner similar to placement of an intravenous catheter is the most practiced and simplest method, but it is not always successful because of technical difficulties. The only routine site for this technique is the radial artery. Use of a catheter over a guidewire as per the Seldinger technique is strongly advised at most other sites.

Take time to ensure proper alignment of the desired site. Delays, complications, and inability to successfully cannulate an artery often occur as a result of failure to properly prepare the desired site and involved limb. An important preparatory step is to ensure that the target limb is secured flat and not rotated; any rotation could result in the desired artery being shifted from the expected anatomic position and making it more difficult to cannulate. For example, to adequately prepare the radial artery, immobilize the wrist and hand in mild dorsiflexion with some padding for support underneath the wrist (Fig. 20.5, step 1). Prepare the skin with sterile technique. Inject local anesthetic with a 25-gauge or smaller needle to achieve sufficient infiltration and to ensure a painless procedure. Subcutaneous infiltration of lidocaine or a similar anesthetic may also reduce vessel spasm at the time of arterial puncture.

Check the catheter assembly for proper movement and function. Alternatively, a 3-mL syringe with the plunger removed can be used as a blood reservoir. Advance the catheter toward the palpated artery at a comfortable angle for the operator, generally 30 to 45 degrees from the skin (see Fig. 20.5, step 2). Make a small incision with a No. 11 scalpel blade or a larger-bore needle to eliminate the problem of damage to the catheter from kinking on the skin. The tip of the needle is often perceived to pierce the artery, but successful puncture is confirmed by identifying a “flash” of arterial blood flow into the needle hub and reservoir. As the needle-catheter assembly advances through the skin toward the artery, the initial flash of arterial blood is obtained by the needle alone, which protrudes beyond the catheter. For this reason, the needle-catheter assembly should be lowered and advanced 2 mm forward to ensure that the tip of the catheter has cannulated the vessel, along with the needle (see Fig. 20.5, step 3). The position of the catheter within the vessel lumen is confirmed by continuous return of arterial blood. The catheter alone can now be advanced with care over the needle and into the artery (see Fig. 20.5, step 4). If the catheter fails to thread, it has not properly entered the vessel lumen and should not be forced to advance without confirmation of placement by active blood return.

When successful blood flow into the needle-catheter assembly has ceased, it may have pierced the backside of the arterial wall and may no longer be in the artery. This double-puncture method is useful for cannulating small vessels, yet it is not recommended as a routine procedure for inexperienced clinicians. If double puncture has occurred and blood has ceased to flow into the collection reservoir, do not remove the entire needle-catheter assembly. Instead, simply retract the needle slightly to determine whether blood flow into the catheter can be reestablished. If blood flow occurs, gently advance the catheter. If not, slowly withdraw the catheter until pulsatile blood flow reappears and then advance the catheter into the artery. It is important for the clinician to be aware of whether the tip of the needle or the catheter is the leading edge within the vessel.1

Once the catheter is fully advanced into the vessel lumen, maintain occlusive pressure on the proximal end of the artery to limit blood loss, and then remove the needle (see Fig. 20.5, step 5). Next attach narrow-bore, low-compliance pressure tubing to the catheter (see Fig. 20.5, step 6). Securely suture the apparatus to the wrist, and then apply an appropriate sterile dressing. Occasionally, difficulty will be encountered advancing the catheter into the lumen. The “liquid stylet” method may aid further passage of the catheter.34 Fill a 10-mL syringe with approximately 5 mL of sterile normal saline. Attach the syringe to the catheter hub, and aspirate 1 to 2 mL of blood to confirm intraluminal position. Then slowly inject the fluid from the syringe and advance the catheter behind the fluid wave.

Ultrasound guidance is now routinely used for both peripheral and central venous access and can also assist with arterial cannulation (see Ultrasound Box 20.1). Ultrasound-guided radial artery cannulation is associated with an increased first attempt success rate, fewer attempts prior to success, and a lower incidence of hematoma formation compared to the traditional method of palpation.35

With B-mode ultrasound, visualize the targeted artery in real-time by using a 7.5- to 10-MHz linear-array transducer. Differentiate the target artery from the adjacent vein by its pulsatility and noncompressibility under mild pressure by the transducer. Use color Doppler to further distinguish between arteries and veins. Either transverse or longitudinal views can be applied during arterial cannulation, but the transverse view is often more useful in smaller arteries. Once the catheter has entered the artery and it is confirmed by blood flow, place the ultrasound transducer on the field to free up the nondominant hand.16,37

The number of attempts and additional arterial punctures increases the size of the developing hematoma and the real risk for vessel wall damage, thrombosis, and even loss of arterial flow through the vessel. Despite the added trauma, there is no reported increase in complications when both walls, rather than one, are punctured in a single cannulation attempt.38–40

Guidewire Techniques for Arterial Cannulation

A modified Seldinger technique can often rescue a failed direct cannulation attempt with an over-the-needle catheter (Fig. 20.6). If the catheter has been placed in the arterial lumen with successful blood return, pass a properly sized guidewire through the catheter and into the artery. Advance the catheter over the guidewire and fully into the vessel. Be careful because stiffer guidewires, unlike most prepackaged ones, do not have a softer, more flexible end tip and the vessel wall may be
Arterial Cannulation: Over-the-Needle Catheter Technique

Immobilize the hand and wrist in mild dorsiflexion on a padded arm board. Prepare the skin with antiseptic, anesthetize, and apply a sterile drape.

Advance the needle into the artery at a 30° to 45° angle to the skin. Confirm arterial puncture by observing a flash into the needle hub.

Lower the angiocatheter and advance it 2 mm forward to ensure that the tip has cannulated the vessel. Confirm proper placement by observing continuous arterial blood return.

Carefully advance the catheter over the needle and into the artery. Do not force the catheter; if it fails to easily thread, it has not properly entered the vessel lumen. (See text for troubleshooting tips.)

Tamponade over the artery proximal to the tip of the catheter (to prevent blood loss), and remove the needle.

Attach the tubing from the pressure transducer to the catheter. Suture the catheter hub to the skin and cover with a sterile dressing, such as Tegaderm.

Figure 20.5 Arterial cannulation: over-the-needle catheter technique. Note that in this example the angiocatheter does not have a safety mechanism such as a retracting needle. This allows a syringe to be attached to the back of the hub as a blood reservoir. Such angiocatheters may be difficult to find in some institutions.
Arterial Cannulation: Guidewire Technique

1. Prepare the wrist and insert the needle as described in Fig. 20.5. Look for pulsatile blood return in the flash chamber.

2. Hold the catheter in place and remove the needle. Pulsatile blood should flow from the catheter. If not present, slowly back the catheter out, and observe for blood return.

3. Insert the guidewire through the catheter. It should advance freely and easily into the vessel. Do not force it.

4. Advance the catheter over the wire and into the artery. Remove the wire, attach the catheter hub to the transducer tubing, and secure it to the skin.

Figure 20.6 Arterial cannulation: guidewire technique.

damaged or even perforated with excessive force. Alternatively, catheter sets are available with an attachable, catheter-contained, wire stylet that permits a modified Seldinger technique for placement of the catheter. The over-the-needle catheter follows the self-contained guidewire during cannulation. Numerous commercially available sets feature styles of guidewire and reservoir attachments that are different from an over-the-needle catheter assembly. Most resemble the Arrow Arterial Catheterization Kit (Arrow International, Inc., Reading, PA) (Fig. 20.7; see also Fig. 20.2). These kits are extremely practical for smaller vessels, especially the radial, brachial, and axillary arteries, and have excellent success rates for first-time placement. Although some authors have suggested that guidewire-based techniques will improve arterial cannulation success rates in certain patients, studies have shown that success is more a function of operator experience and personal preference.

Seldinger Technique

The Seldinger technique for venipuncture is described in detail in Chapter 22. Overall success rates with the Seldinger, guidewire-directed technique are superior to those with direct arterial cannulation. A few available kits are designed specifically for cannulation of larger arteries, but single-lumen venous catheters with guidewires may be used if catheter size and length are appropriate for specific arteries (see the following section for guidelines). The guidewire technique should be used initially for critical patients.

Place the needle percutaneously into the arterial lumen, as described previously. Then place a guidewire through the needle into the vessel lumen, and remove the needle. Thread the catheter over the wire, and pull the wire out. Although most kits have vessel dilators, especially with larger catheter sizes, caution is advised. Dilate the tract only and not the artery to avoid unnecessary blood loss and excessive arterial injury.

Cutdown Technique for Arterial Cannulation

The cutdown technique is rarely used, but in certain circumstances it may be performed to obtain arterial access. With the increasing use of ultrasound-assisted catheter placement, this technique should seldom be required. Perform cannulation
Arterial Cannulation: Arrow Arterial Catheterization Kit

1. Ensure that the actuating lever is fully retracted, and then insert the needle into the artery. Monitor for blood flashback in the hub of the needle to confirm intraarterial placement.

2. Stabilize the needle and advance the guidewire into the vessel by using the actuating lever. When the lever reaches the reference mark (black arrow) on the device, the wire begins to exit the needle.

3. After the wire is fully inserted, advance the entire assembly 1–2 mm farther into the vessel. Firmly hold the needle in position, and advance the catheter over the wire and into the vessel. A slight rotating motion may be helpful.

4. Remove the needle and guidewire assembly and attach the transducer tubing to the catheter hub. Use the wing clip (arrow) to suture the catheter to the skin, and then cover with a sterile dressing.

Figure 20.7 Arterial cannulation: Arrow Arterial Catheterization Kit.

After direct visualization of the vessel. A cutdown can be performed on any artery but is most commonly reserved for distal lower limb arteries and, rarely, the brachial artery. After a site has been selected, prepare the overlying skin with an antiseptic solution. Using sterile technique, inject local anesthetic solution subcutaneously in a horizontal line 2 to 3 cm long and perpendicular to the artery. Omit this step if the patient is unconscious or otherwise anesthetized at the cutdown site.

Use a No. 10 or 15 scalp blade to incise the skin along the anesthetic wheal. Spread the underlying tissues parallel to the artery with a mosquito hemostat. Palpate the pulse repeatedly throughout the procedure to ensure proper positioning. Once the surrounding soft tissue has been retracted and after exposing approximately 1 cm of the artery, isolate the artery by passing two silk sutures underneath it with the hemostat. Strip away only enough perivascular tissue to expose the artery. Perivascular tissue will help limit bleeding at the time of catheter removal. Introduce an over-the-needle catheter device, such as the kind used in the percutaneous method, and introduce it through the skin just distal to the incision. Advance it into the surgical site (Fig. 20.8). Alternatively, use a modified Seldinger guidewire setup to catheterize the artery. Puncture the arterial wall with the tip of the needle, and thread the catheter into the vessel lumen. When this has been accomplished, remove the two silk sutures, which have only been used to control the vessel, and close the skin incision. Do not tie off the artery the way that a vein is tied off during a venous cutdown. Apply firm pressure over the cutdown site, as used after arterial puncture. If pressure is not applied, separation of the soft tissues during the procedure may allow considerable hemorrhage into the tissue.

Local Puncture Site and Catheter Care

Once the catheter has been placed successfully, advance it until the hub is in contact with the skin. Secure the catheter by fastening it to the skin with suture material. Silk (2-0) or nylon
Further secure the catheter and its connecting tubing with sterile sponges and adhesive tape. If the tubing becomes disconnected inadvertently, the patient can exsanguinate rapidly. Tie the loose ends of the suture around the catheter at its hub. Then, after laying two ties, place a second set of knots on the back portion without occluding the lumen by constriction (Fig. 20.9).

Another option to secure these lines is to apply commercially available sutureless securement devices. According to one study, sutured lines are associated with a 10% rate of catheter-related bloodstream infection. In comparison, lines that were secured with a sutureless method had an infection rate of less than 1% and eliminated the potential for accidental needlestick from suturing.43

After tying the catheter in place, apply a drop of antibiotic ointment to the puncture site44 and a self-adhesive dressing over the area. Certain transparent dressings containing chlorhexidine gluconate can retard the colonization of bacteria.45 Further secure the catheter and its connecting tubing with sterile sponges and adhesive tape. Make sure that all tubing connections are tight and secure. If the tubing becomes disconnected inadvertently, the patient can exsanguinate rapidly.

Fluid-Pressurized Systems

When successful arterial cannulation has been performed, attach the catheter to a pressurized fluid-filled system. A three-way stopcock can be interposed between the patient and the transducer for blood gas sampling and to allow flushing of the system. Flushing can be periodic or continuous at a rate of 3 to 4 mL/hr through a continuous-flow device. Most institutions use normal saline in place of heparinized solution to maintain patency. Use of a heparinized flush solution in pressurized arterial lines may result in greater long-term accuracy of pressure monitoring, but no real difference in catheter blockage has been reported, and this approach avoids heparin-related complications such as drug incompatibility, thrombosis, local tissue damage, and hemorrhage.46–50 For short-term setups as in the ED, saline is sufficient.

Blood samples are obtained easily from the arterial catheter system. Attach a syringe to the three-way stopcock and aspirate and discard the blood to clear the line. Studies examining the necessary discard volume of flushed blood solution have found considerable variation, depending on the volume of the system.51,52 Short lengths of tubing between the catheter and the aspiration port minimize the necessary discard volume. For a tubing length of 91 cm (36 inches), aspirate 4 to 5 mL; for a tubing length of 213 cm (84 inches), aspirate 8 mL.53 Attach a second syringe that has been heparinized, and aspirate 3 mL of blood to send for ABG analysis. If the blood is to be used for other tests, the second syringe does not need to be heparinized. Self-contained, nondetachable, blood sample withdrawing systems allow less blood wasting for sampling. Flush the stopcock and line to avoid clotting.

**SELECTION OF ARTERIES FOR CANNULATION**

**Radial and Ulnar Arteries**

The radial artery is most frequently used for prolonged cannulation. Widespread collateral flow is present in the wrist because of two major palmar anastomoses known as arches (Fig. 20.10). The superficial palmar arch lies between the aponeurosis palmaris and the tendons of the flexor digitorum sublimis. The arch is formed mainly by the terminal ulnar artery and the superficial palmar branch of the radial artery. The other major communication of these two vessels, the deep palmar arch, is formed by connections of the terminal radial artery with the deep palmar branches of the ulnar artery.54 Some collateral flow is almost always present at the wrist, with the deep arch alone being complete in 97% of 650 hand...
A false-positive result.60 Time permitting, performance of some compress the palmar arches between fascial planes and yield of the hand with wide separation of the digits, which may perform radial artery puncture. Be careful to avoid overextension If return of color takes longer than 5 to 10 seconds, do not of pallor 5 to 15 seconds after release of the radial artery.6,19,58,59 inadequate collateralization, is defined as the continued presence in infants and children when the radial arteries are obstructed for the procedure. The radial artery is more superficial as it moves closer to the wrist, the radial artery rests on the flexor digitorum superficialis muscle, the flexor pollicis longus muscle, the pronator quadratus muscle, and the radius bone.54 Isolate the pulsation of the artery on the palmar surface of the wrist. The radial artery is more superficial as it moves closer to the wrist. In this location, it provides a more consistent site for cannulation because of its fixation and decreased mobility. Dorsiflexing the wrist at approximately a 45-degree angle over a towel or sandbag and fixing the wrist to an arm board will also help isolate the artery.65 This degree of preparation should be considered standard when time for setup permits (see Fig. 20.5, step 1).38,39

Antegrade radial artery cannulation may be accomplished in infants and children when the radial arteries are obstructed and retrograde blood flow is observed during a failed cutdown attempt at standard retrograde arterial cannulation.66 In addition, displacement of perivascular interstitial fluid in neonates and bright light make the course of the artery visible. Then, under direct vision, cannulation of the artery becomes as easy as venous cannulation.65 Doppler ultrasound on selected patients with poor peripheral pulses may also facilitate percutaneous radial artery cannulation and minimize the number of punctures needed for placement.68

Figure 20.11 Allen test. Before puncturing the radial artery for cannulation, it is important to identify a competent ulnar artery should injury to the radial artery occur. This is not generally required, nor standard, for a single arterial puncture. (Adapted from Schwartz GR, editor: Principles and practice of emergency medicine, Philadelphia, 1978, Saunders, p 354. Reproduced by permission.)
The ulnar artery is seldom used because its smaller size makes it more difficult to puncture than the radial artery. At the wrist, the ulnar artery runs along the palmar margin of the flexor carpi ulnaris in the space between it and the flexor digitorum sublimis. Use caution because the artery runs next to the ulnar nerve as both pass into the hand just radial to the pisiform bone. Minimize any potential injury by approaching the ulnar artery from the radial side. Make the ulnar artery more accessible with dorsiflexion of the wrist.

**Brachial Artery**

Although it appears safe for arterial puncture, the brachial artery does not have the anatomic benefit of the collateral circulation that is found in the wrist. The brachial artery begins as the continuation of the axillary artery and ends at the head of the radius, where it splits into the ulnar and radial arteries. The preferred puncture site of the brachial artery is in or just proximal to the antecubital fossa. In this region, the artery lies on top of the brachialis muscle and enters the fossa underneath the bicipital aponeurosis with the median nerve, and on the medial side of the artery (Fig. 20.12). Both the radial and the axillary arteries are preferred over the brachial artery in the upper extremity. With the brachial artery, there is increased risk for ischemic complications from the reduction in collateral circulation, as well as the need to maintain the arm in extension for puncture and prolonged cannulation. Nonetheless, safe cannulation of the brachial artery has been demonstrated by some investigators. Bazaral and coworkers found only one minor thrombotic occurrence in a study of more than 3000 brachial artery catheterizations over a 3-year period in cardiac surgery patients. A longer catheter (10 cm) is required for the brachial artery so that sufficient length is available to traverse the elbow joint.

**Dorsalis Pedis Artery**

The dorsalis pedis artery continues from the anterior tibial artery and runs from approximately midway between the malleoli to the posterior end of the first metatarsal space, where it forms the dorsal metatarsal and deep plantar arteries. The lateral plantar artery, a branch of the posterior tibial artery, passes obliquely across the foot to the base of the fifth metatarsal. The plantar arch is completed at the point where the lateral plantar artery joins the deep plantar artery between the first and second metatarsals. On the dorsum of the foot, the dorsalis pedis artery lies in the subcutaneous tissue parallel to the extensor halluces longus tendon and between it and the extensor digitorum longus (Fig. 20.13).

Cannulate the artery in the midfoot region. Although this vessel is amenable to cutdown, the vascular anatomy of the foot is quite variable. This is of no consequence if a pulse can be palpated, but Huber, in his dissection of 200 feet, noted that the dorsalis pedis artery was absent in 12% of patients. In 16% of patients the dorsalis pedis artery provided the main blood supply to the toes. Although the dorsal pedis and posterior tibial arteries form similar collateral foot circulation as in the hand, the nature of advancing vascular disease makes this a more difficult cannulation, with higher complication rates than in the wrist. This site has its major utility in pediatric monitoring cases. Attempts to predetermine collateral flow with a modified Allen test using the posterior tibial and dorsalis pedis arteries is not as easily performed in the foot as in the hand, nor are there good data to prove its validity. Monitoring problems also exist with this artery. The pressure wave obtained with an electronic transducer attached to the dorsalis pedis artery will be 5 to 20 mm Hg higher than that of the radial artery and, in addition, will be delayed by 0.1 to 0.2 second.

**Femoral Artery**

The femoral artery is the second most commonly used vessel for arterial cannulation. Based on its ease of cannulation and
low record of complications, it has been called the vessel of choice for arterial access.73–77 Along with the axillary artery, the femoral artery more closely resembles aortic pressure waveforms than those from any other peripheral site. The femoral artery is the direct continuation of the iliac artery and enters the thigh after passing below the inguinal ligament. Arterial puncture must always occur distal to the ligament to prevent uncontrolled hemorrhage into the pelvis or peritoneum.78 The artery may be palpable easily midway between the pubic symphysis and the anterior superior iliac spine. The advantage of cannulating the artery at a site just distal to the inguinal ligament is that the artery can be compressed against the femoral head. Cannulation becomes more difficult the more distal the puncture site is from the inguinal ligament because the femoral artery splits into the superficial femoral and the deep femoral arteries. These arteries, especially the deep femoral, can be challenging to compress if bleeding needs to be controlled. One method of locating an appropriate arterial puncture site is to place the thumb and fifth finger on the pubic symphysis and the anterior superior iliac spine and locate the artery underneath the middle knuckle. When puncturing this vessel, be careful to avoid the femoral nerve and vein, which form the lateral and medial borders, respectively (Fig. 20.14).

A longer, larger-diameter catheter is required for accurate monitoring of the femoral artery because of its size and the relatively greater depth at which it lies. Only the Seldinger technique is recommended for this site, which enables placement of a 15- to 20-cm plastic catheter for prolonged monitoring. Avoid using catheter-through-the-needle or over-the-needle devices because of the vessel’s distance beneath the skin. Leakage around the catheter can occur with these devices as a result of the high arterial pressure and the loose fit of the cannula in the hole in the vessel wall. Regardless of the device used, enter the skin with the needle at an angle of approximately 45 degrees instead of the usual 15 to 20 degrees.

The extremely large ratio of arterial diameter to catheter diameter is thought to reduce the incidence of thrombosis, particularly total occlusion. However, occlusion has been reported with femoral cannulation for monitoring purposes.79 A commonly perceived disadvantage of this site is the increased possibility of bacterial contamination because of its proximity to the warm, moist groin and perineum; however, no studies have confirmed this hypothesis.80 The femoral area is inconvenient for any patient who is awake and mobile or for a patient who is able to sit in a chair. If the patient is that mobile, reconsider the risk-benefit ratio of invasive monitoring. Despite theoretical difficulties, some hospitals use femoral arterial lines almost exclusively, and the ICU nursing staff is often more comfortable caring for these lines than those at other sites.

**Umbilical and Temporal Arteries**

In neonates, arterial access can be accomplished for a short time through the umbilical artery (see Chapter 19). After this artery closes, the temporal artery provides a safe alternative. Prian described the use of the temporal artery and noted its accessibility and lack of clinical sequelae if it undergoes thrombosis.82 Use the cutdown method with a 22-gauge catheter after the artery’s course has been traced with an ultrasonic flow detector. Because of the increasing accuracy of ear oximeters and the use of capillary blood gas samples for determination of pH, prolonged arterial cannulation will become less frequent during infant care.

**COMPLICATIONS OF ARTERIAL CANNULATION**

Arterial cannulation is safe if care is taken to avoid complications. Most can be avoided by adhering to a few simple principles. Reported clinical sequelae of arterial puncture and cannulation range from simple hematomas to life-threatening infections and exsanguination. In addition, ischemia, arteriovenous fistula, and pseudoaneurysm formation are possible. The incidence of complications varies with the site selected, the method of cannulation, and the clinician’s level of skill and experience. Early detection of complications is greatly aided by enhanced vigilance.

It is difficult to compare complication rates at various sites because most published studies have primarily used only the radial artery. No studies have compared the approach and complication rates of arterial catheters in the ED with those placed in the ICU or operating room. In a large study spanning 24 months, 2119 ICU patients had arterial catheters placed at admission: 52% at the radial site and 45% at the femoral site. The most common complication was vascular insufficiency (4%), followed by bleeding (2.1%), and infection (0.6%). No difference was reported in infection rates at femoral versus radial sites.82 There are reports of complications from arterial puncture for procedures unrelated to cannulation, such as arteriography or simple puncture for blood sampling as routinely performed in the ED. In a study of 2400 consecutive cardiac catheterizations over a 12-month period, complications occurred in 1.6% of patients, including 17 needing vascular repair and 28 requiring transfusion.83
Hematoma formation at the puncture site is common. Zorab reported this complication in 50% of all catheterizations. Bruising was of minimal clinical significance in this report, but leakage can be dangerous when it occurs around the catheter or from the puncture site after the catheter is removed (Fig. 20.15). Compression neuropathies requiring surgical decompression have been reported after arterial puncture secondary to hematoma formation. The large amount of soft tissue surrounding the femoral artery makes bleeding in this area difficult to control. Large hematomas are not uncommon after femoral artery catheterization; indeed, Soderstrom and colleagues reported two cases of bleeding that required transfusion after femoral puncture. Though uncommon, a clinician should be aware of the potential development of a retroperitoneal hematoma, a morbid complication, following femoral artery cannulation. Suspect this complication in patients in whom hypovolemic shock with a falling hematocrit develops following cannulation. A large femoral hematoma is pictured here 3 days after iatrogenic injury.

Figure 20.15 Right groin hemorrhage with resulting hematoma. The injury resulted from a failed attempt to place a right femoral artery catheter via the Seldinger technique. Inadequate pressure after the procedure resulted in bleeding from the artery. A large femoral hematoma is pictured here 3 days after iatrogenic injury.

Radial artery cannulation without any long-term ischemic complications despite evidence of radial artery occlusion after decannulation in more than 25% of the patients. Serious complications after radial artery cannulation are extremely rare in the absence of contributing factors such as preexisting vasospastic arterial disease, previous arterial injury, protracted shock, high-dose vasopressor administration, prolonged cannulation, or infection. Prevention of bleeding complications can be accomplished with frequent careful inspection of the puncture site and the use of prolonged compression after removal of the catheter or needle. Maintain firm pressure for 10 minutes or longer after removing a peripheral artery catheter and longer after femoral cannulation or if the patient has received anticoagulants. Five minutes of pressure is sufficient after puncture for a blood gas sample in an individual with normal coagulation. Exsanguination may occur if the arterial line apparatus becomes disconnected. This is more common in an obtunded or combative patient, and restraints are often required for patients with indwelling arterial cannulas. Exsanguination should not occur if tight connections are maintained throughout the system and if frequent, careful inspections of both the circuit and the patient are made.

Meticulous attention to aseptic technique is necessary during insertion and catheter maintenance to minimize the risk for catheter-related infections. Serious infections rarely complicate arterial cannulation. Simple interventions can reduce the risk for serious catheter-related infection. Evidence supports the use of full-barrier precautions during catheter insertion, specialized nursing care, and newer-generation catheters with antiseptic hubs or antimicrobial-agent–impregnated catheters. The incidence of catheter-related infections increases with prolonged cannulation. Catheters placed via sterile technique have an extremely low rate of infection in the first 96 hours after placement. Catheters changed over a guidewire every 96 hours have an infection rate of approximately 10% at the radial and femoral sites.

Most infections begin locally at the puncture site and remain localized, although systemic sepsis has been reported. The radial and femoral sites have a similar incidence of complications; however, axillary cannulation seems to have a much higher incidence of infection (although no large studies of cannulation at this site exist). Arterial cannulas are more prone to infectious complications than other vascular catheters. Many mechanisms have been proposed for this increased incidence. The arterial pressure–monitoring system usually consists of a long column of fairly stagnant fluid and is subject to frequent manipulation. Stamm and coworkers found that patients were at greater risk for systemic infection if they had an arterial line and required frequent blood gas determinations than if they had the cannula alone. The sampling stopcock is a site of frequent bacterial contamination.

The risk for infection also increases as the duration of cannulation is prolonged. Older studies recommend that catheters be changed after 4 days if continued monitoring is necessary. In addition, Makai and Hassemer recommended changing the entire fluid-filled system, including the transducer chamber domes and continuous flow devices, every 48 hours. However, the risk for noninfectious complications increases with more frequent catheter and site changes. Therefore daily evaluation of the site is advised, and catheter change should not be mandatory until 7 to 8 days if the site remains clean.
Shinozaki and colleagues demonstrated a marked reduction in equipment contamination when the continuous flush device was located just distal to the transducer, as opposed to closer to the three-way stopcock used for sampling. This setup reduces the length of the static column of fluid between the sampling stopcock and the transducer. As mentioned previously, a drop of sodium hydroxide or antibiotic ointment applied to the puncture site decreases the incidence of local wound infection. This technique has drawn a great deal of criticism, however. The current standard is a clean, nonocclusive, dry dressing. An antibiotic- or silver-impregnated catheter is recommended for long-term placement.

Thrombosis of the vessel in which the cannula is placed is another frequently encountered problem. The incidence of thrombosis varies with the method used to determine the presence of clots. Bedford and Wollman found a greater than 40% occlusion rate when radial artery catheters were left in place for longer than 20 hours. All these occluded vessels eventually recanalized. Angiographic studies show deposition of fibrin on 100% of catheters left in place for longer than 1 day, although clinical evidence of ischemia secondary to occlusion by such thrombi is seen in less than 1% of cases in most studies. Most reports of nonangiographic catheterization involve the radial artery. Therefore it is difficult to compare the incidence of thrombosis at other sites, although during the 176 femoral catheterizations reported by Soderstrom and colleagues and Ersoz and associates, dorsalis pedis pulses were decreased in only two patients and no clinical signs of ischemia were noted. Larger catheter sizes, trauma during cannulation, and the presence of atherosclerosis have all been postulated to increase the incidence of thrombosis; however, conflicting studies abound.

Arterial spasm after puncture (usually following multiple attempts) can predispose to thrombus formation and can even lead to ischemic changes without fibrin deposition. Successful reversal of spasm with intraarterial lidocaine, reserpine, and phentolamine has been reported, but no reliable studies of their efficacy in this clinical situation have been published.

Thrombosis can be minimized by decreasing the duration of catheterization and proper flushing. Surgical embolectomy or thrombectomy is rarely required because the smaller vessels that are most likely to occlude usually have good collateral circulation. The larger femoral artery, which has poor collateralization, rarely occludes with catheterization when used for monitoring purposes.

Thrombosis can result in occlusion of the catheter. Time until occlusion of radial and femoral artery catheters has been compared. Radial cannulas became occluded at an average of 3.8 days, whereas femoral cannulas became occluded after 7.3 days. The importance of this comparison is minimal if the clinician follows infection prophylaxis guidelines and changes arterial catheters after 4 days.

A few less common complications are easily prevented. One that occurs only with the percutaneous catheter-through-the-needle method is catheter embolization. Once the catheter has been placed through the needle, it should never be pulled back because the end of the catheter may be sheared off by the sharp needle bevel. If this complication occurs, surgical removal of the tip of the catheter is necessary.

Skin necrosis is a complication of radial artery cannulation that involves an area of the volar surface of the forearm proximal to the cannula. Wyatt and colleagues believed this to be secondary to the poor blood supply in this area and thought that proper technique would decrease the incidence of necrosis.

One feared complication of indwelling radial and brachial arterial catheters is the occurrence of a cerebrovascular accident secondary to embolization from flushing of the catheters. As little as 3 to 12 mL of flush solution has been shown to reflux to the junction of the subclavian and vertebral arteries. A fatality caused by air embolism from a radial artery catheter has been reported and was re-created in a primate model. Although these animals were much smaller (7 kg) than an adult human, as little as 2.5 mL of air introduced at a relatively low flush rate was found to embolize in retrograde fashion to the brain. Cerebral embolization can be prevented with the use of continuous flush systems (3 mL/hr) and by ensuring the integrity of the tubing and transducer systems to prevent entry of air. In addition, small volumes (<2 mL) of intermittent flush solution should be used.

Complication rates also vary according to the method of arterial cannulation. Mortensen studied the three main techniques (discussed earlier in the section on Techniques), but unfortunately, most of his arterial cannulations were for angiographic purposes. The complications associated with prolonged cannulation time are therefore underrepresented. In Mortensen's series, cutdown arteriotomy exhibited the lowest incidence of complications (7.7%), whereas the Seldinger technique had a 17.7% incidence of complications. The complication rate with percutaneous cannulation was 11.3%. False passage of the guidewire, the catheter, or both was associated with increased intimal damage and complications. It is imperative that the wire or catheter be advanced only if no resistance is met.

Once the monitoring system is set up, manipulate it as little as possible. Perform any handling with aseptic technique. Change the tubing and other fluid-filled devices every 48 hours, and insert catheters into a vessel that provides the largest vessel-to-catheter ratio as possible.

If these principles are followed and the patient and system are carefully inspected at frequent intervals, complications of arterial puncture and cannulation can be minimized.

**INTERPRETATION**

An indwelling arterial catheter provides continuous blood pressure monitoring. The trend of a patient's pressure facilitates assessment of the effect of various therapeutic interventions. The absolute systolic and diastolic pressure measured will vary at different catheter sites, with higher peak systolic pressures measured at the periphery. The pressure will also be higher when measured at the distal end of the lower limb. A wide variance between direct arterial pressure and the pressure measured with a standard pneumatic cuff will always exist in some patients. Oscillometric blood pressure measurement can significantly underestimate arterial blood pressure. For this reason, regularly compare a cuff pressure with that obtained via invasive monitoring. Moreover, a change in their relationship may be the first indication of difficulty with the direct measuring system.

Waveform analysis may also provide an early indication of thrombosis in the arterial catheter. Many variables affect the waveform, including cardiac valvular disease and arteriosclerosis. Waveforms may vary tremendously among patients, but after an adequate monitoring system has been established,
a change in an individual's pressure wave is usually indicative of thrombosis or a malfunction in the monitoring system. A change in waveform may also indicate a change in the patient's cardiovascular status, such as a papillary muscle rupture. Before making a therapeutic decision based on an electronically generated number, recheck the patient's blood pressure with a pneumatic cuff. This device is less fallible than the electro-mechanical system.

Radial systolic arterial pressure poorly estimates the actual ascending aortic pressure, with more than 50% of cases reporting a difference in values of 10 to 35 mm Hg. Mean arterial pressure and diastolic pressure, in contrast, are highly accurate with greater than 90% of the values being within 3 mm Hg of aortic values. Longer catheters have also been used successfully from radial sites to more accurately reflect central aortic pressure for cardiac surgery patients.

An indwelling arterial cannula can provide valuable information about the hemodynamic status of a patient (through continuous pressure monitoring) and about the patient's respiratory and metabolic status (through intermittent sampling for ABG analysis and other blood tests). The PCO₂ and pH of the blood can be used to define four major groups of metabolic derangement: respiratory acidosis or alkalosis and metabolic acidosis or alkalosis. Rarely will a disorder be strictly classified into one of these groups; however, a simple chart such as that provided in the Appendix helps determine the relative effects of metabolic and respiratory influence on blood pH (see also discussion in the Appendix).

Adequacy of blood oxygenation can be determined from the measured Po₂ of arterial blood and the known concentration of oxygen that the patient is inspiring. To avoid iatrogenic complications of intensive care, one must be absolutely certain that the data are from an arterial sample that has been properly analyzed before basing one's treatment decisions on the numbers obtained. Not uncommonly, a venous sample is interpreted as though it were arterial. Furthermore, false readings may result if the sample is not free of air bubbles, not promptly chilled, or not analyzed within 20 to 30 minutes. Though still controversial, blood gas values that are not corrected for body temperature appear to be more appropriate for guiding therapy in hypothermic patients.

**CONCLUSION**

As intensive care knowledge and technology grow and develop, cannulation of the arterial system may decrease in frequency. Oximeters can determine the quality of blood oxygenation transcutaneously and are becoming more accurate and sophisticated. Electronic sphygmomanometers are being refined for continuous indirect blood pressure monitoring. As these devices improve and noninvasive sampling methods for clinically relevant electrolytes and physiologic markers are refined, the indwelling arterial cannula may in time become considered overly invasive. Despite improvements in noninvasive monitoring devices, the current need for frequent blood sampling for chemical and hematologic analysis remains an indication for its use in selected critically ill patients. Overzealous blood gas analysis may lead to iatrogenic anemia in the ICU. Multiple reports have documented the advantages of limiting frequent blood sampling.

Arterial puncture and cannulation are invaluable aids to the emergency and critical care clinician. Long-term catheterization is a safe procedure when the catheter is placed, maintained, and removed with care. The radial artery is the most favored location for puncture, but as more experience is gained and reported with femoral artery catheterization, the latter may become a more frequently used site. Selection of either site is associated with a low complication rate and should be determined by the skill of the clinician, the nursing team, and the relative convenience and comfort of the patient. Ultrasound guidance for arterial catheter placement is becoming the standard and should be considered in every instance to reduce the need for multiple puncture attempts.

**ACKNOWLEDGMENT**

The author would like to acknowledge the work of Drs. Dave Milzman and Tim Janchar, who were authors of this chapter in previous editions.

**REFERENCES ARE AVAILABLE AT** [www.expertconsult.com](http://www.expertconsult.com)
INTRODUCTION

Intravenous (IV) access is a mainstay of modern medicine. IV cannulation is a procedure (Videos 21.1 and 21.2) performed by a wide array of health care professionals, including physicians, nurses, physician assistants, phlebotomists, and emergency medical technicians. In the emergency department (ED), uncomplicated peripheral venous access is usually secured by a nurse or technician. In the United States, more than 25 million patients have peripheral IV catheters placed each year as vascular access for the administration of medications and fluids and the sampling of blood for analysis. IV access can usually be accomplished in less than 5 minutes.1–4 Despite their growing number, dedicated IV teams are very costly and not always cost effective.5,6 Moreover, in the ED setting, multiple providers may be called on to obtain IV access, thus making it an essential skill for both emergency physicians and nurses to master. Subtleties in technique are important and can be improved with practice; newer technologies such as ultrasound can assist providers in placing IV lines in even the most challenging situations.

HISTORICAL PERSPECTIVE

Bloodletting, or bleeding, dates to the time of Hippocrates. The ancient technique consisted of tying a bandage around the arm to distend the forearm veins, opening a vein with a sharp knife, and collecting the blood into a basin. In the Middle Ages, this was performed by barber-surgeons. In 1656, Sir Christopher Wren injected opium into dogs intravenously with a quill and bladder, thereby becoming the father of modern IV therapy.7 Blood transfusions also date back to the mid-1600s. The French physician Jean Denis is credited with the first successful transfusion by giving lamb’s blood to a 15-year-old boy.8,9

Originally, 16- to 18-gauge indwelling steel needles were used for IV infusions. In the 1950s the Rochester needle was introduced, which was a resinous catheter on the outside of a steel introducer needle. Because of increased comfort and mobility, plastic catheters have replaced indwelling metal needles and are now almost universal.7,10

INDICATIONS AND CONTRAINDICATIONS

Obtaining timely and adequate vascular access is a major priority during any resuscitation. In patients with normal perfusion, differences in delivery times for injections centrally versus peripherally are minimal, within seconds.11 During cardiopulmonary resuscitation (CPR), however, medications have been shown to reach the central circulation faster with central access than with peripheral venous access.12 A change in outcome, though, has not been demonstrated with the central administration of advanced cardiac life support drugs; hence,

### Peripheral Intravenous Access

#### Indications
- Venous blood sampling
- Intravenous fluid infusion
- Intravenous medication infusion
- Blood transfusion
- Intravenous contrast infusion

#### Contraindications
- Extremity with significant edema, burns, sclerosis, phlebitis, or thrombosis
- Ipsilateral radical mastectomy or fistula
- Overlying cellulitis

#### Equipment

- Tourniquet
- Alcohol pad
- IV catheter
- IV tubing
- Tape
- Saline flush
- Tegaderm
- Intravenous tubing (drip set)
- Intravenous fluid

#### Complications

**Early**
- Bruising
- Infiltration
- Air embolism

**Late**
- Phlebitis
- Infection
- Nerve damage
- Thrombosis

Review Box 21.1 Peripheral intravenous access: indications, contraindications, complications, and equipment.
Peripheral IV cannulation is the procedure of choice even during CPR because of the speed, ease, and safety with which it can be accomplished. In less critically ill patients, the role of IV therapy is more often debated and access is ultimately unnecessary in a large proportion of patients in whom it is obtained.\textsuperscript{13} In broad terms, IV access or therapy is needed in patients for whom IV medications are required or when oral therapy is inadequate (e.g., severe shock states), contraindicated (e.g., surgical emergencies), or impossible (e.g., intractable vomiting).

Saline or heparin locks are preferable when IV medications are needed and there are limited foreseeable fluid requirements. Saline locks cost less than a full IV fluid and tubing assembly and are especially helpful when vascular access is needed suddenly.\textsuperscript{14,15} Access to the catheter requires irrigation with a separate syringe and flush.

A peripheral IV central catheter (PICC) shares the attributes of both central and peripheral venous IV lines (see Chapter 24). A PICC is composed of a thin tube of biocompatible material with an attachment hub. It is inserted percutaneously, under ultrasound guidance by a dedicated PICC team, into a peripheral vein and then advanced into a large central vein, followed by radiographic confirmation of placement. PICCs are suitable for long-term vascular access for blood sampling, infusion of antibiotics and hyperosmolar solutions such as total parenteral nutrition, and infusion of certain chemotherapeutic agents. Insert a PICC line as soon as long-term access is anticipated.\textsuperscript{1}

Peripheral IV lines should not be placed in extremities with massive edema, burns, sclerosis, phlebitis, or thrombosis due to risk for extravasation or suboptimal volume flow. When practical, avoid placing an IV line in extremities on the same side as radical mastectomies, though they can be used when an urgent condition exists and other peripheral access is not possible. When feasible, cannulation at infected sites, such as through an area of cellulitis, and extremities with shunts or fistulas should be avoided because it may cause bacteremia or thrombosis. If possible, do not cannulate a vein over or distal to a recent fracture site on an extremity (Fig. 21.1). Veins that drain from an area affected by trauma or major vascular disruption (e.g., distal to a ruptured aorta) are also suboptimal because fluid or medications may not be delivered to the circulatory system.

Blood samples for laboratory analysis are usually drawn before IV cannulation to avoid contamination with IV fluid or medication. However, studies have shown that accurate basic electrolyte and hematologic values can be obtained with peripheral IV lines when infusions are shut off for at least 2 minutes, at least 3 mL of blood is wasted, and all tubes are completely filled to avoid inaccurate bicarbonate readings.\textsuperscript{16-18} By adopting these techniques, one can reduce the number of peripheral needlesticks, minimize trauma or sclerosis of the vein, and improve patient satisfaction.

**ULTRASOUND GUIDANCE AND TRANSILLUMINATION**

Though more commonly used with central venous access, ultrasound can also assist in the placement of peripheral lines. For IV placements that have been designated “difficult” after a certain number of attempts by nursing staff, use of ultrasound guidance increases the success rate and decreases the number of attempts necessary for successful cannulation in both adult and pediatric patients.\textsuperscript{19-22} One 2016 randomized controlled trial showed that ultrasound guidance is particularly helpful in patients with perceived difficult access. However, this same trial noted that patients with perceived easy access had more success with landmarks alone.\textsuperscript{23} The caliber of the vein identified on ultrasound is predictive of its ability to be cannulated. If no vessel is identified, cannulation is not usually possible.\textsuperscript{24,25} An additional issue with ultrasound-guided peripheral IV lines is their longevity. One study highlighted the high premature failure rate of ultrasound-guided peripheral lines.\textsuperscript{26} Ongoing studies are evaluating this concern. It is likely related to the depth of the veins being cannulated, the length and type of catheter used, and the angle of the catheter through soft tissue.

Other devices transilluminate the veins to increase their visibility. This appears to be especially helpful in infants, though little evidence exists evaluating their utility. One 2012 study showed an increase in successful first attempt rates in pediatric patients with difficult peripheral access.\textsuperscript{27} As emergency providers have become more comfortable with these advancing technologies, ultrasound-guided and illumination-assisted insertion of peripheral lines have increased.

**ANATOMY**

The success of cannulation depends on familiarity with the vascular anatomy of the extremities. In the upper extremity, the veins of the hands are drained by the metacarpal and dorsal veins, which connect to form the dorsal venous arch (Fig. 21.2). These sites are excellent for IV therapy and comfortably accommodate 22- and 20-gauge catheters. The venous supply of the wrist and forearm consists of the basilic vein, which courses along the ulnar portion of the posterior aspect of the forearm. It is often ignored because of its location but can easily be accessed if the patient’s forearm is flexed and the clinician stands at the head of the patient.\textsuperscript{28} On the radial side of the forearm, the cephalic vein is commonly known as the intern vein. Readily accessible, this vein can accommodate 22- to 16-gauge catheters. The median veins of the forearm course through the middle of the forearm, and the accessory cephalic

---

**Figure 21.1** Do not place peripheral intravenous (IV) access (long arrow) near or distal to a fracture in an extremity (short arrow). In this case an IV line for pain medication was placed before obtaining the radiograph. The scaphoid fracture was not suspected and the opposite arm had difficult access.
SECTION IV VASCULAR TECHNIQUES AND VOLUME SUPPORT

Safety
Universal precautions must be applied to all patients, especially in emergency care settings, in which the risk for exposure to blood is increased and the infection status of patients is largely unknown.30 One study showed that 11% of all hospital IV catheter injuries to health care workers occurred in the ED.31 Newer catheter devices have emerged that prevent inadvertent needle injuries (Fig. 21.4). The Protectiv IV Catheter Safety System (Smiths Medical, Minneapolis, MN) has a protective sleeve that encases the sharp stylet as it is retracted from the catheter. The needle of the Insyte Autoguard Shielded IV Catheter (Becton, Dickinson and Company, Franklin Lakes, NJ) is instantly encased inside a tamper-resistant safety barrel by pressing an activation button. The Saf-T-Intima IV catheter (Becton, Dickinson and Company), Punctur-Guard Safety Winged Set (Gaven Medical, Grand Island, NY), Vacutainer Safety-Lok (Becton, Dickinson and Company), Shamrock safety winged needle (Smiths Medical, Dublin, OH), and Angel Wing Safety Needle systems (Medtronic, Minneapolis, MN) are all

Veins on the radial aspect of the forearm are easily stabilized and accessible.

The antecubital veins consist of the medial cubital, basilic, and cephalic veins; these are often selected for catheters or blood drawing. IV placement here is easy, but mobility of the arm is restricted once the catheter is in place. The larger veins above the antecubital space, the cephalic and basilic veins, are often more difficult to see but can be accessed without difficulty if necessary.

The relevant lower extremity venous anatomy starts with the dorsal digital veins, which become the dorsal metatarsal veins and form the dorsal venous arch. The arch ultimately splits into the great saphenous vein, which travels up the medial aspect of the ankle, and the small saphenous vein, which courses laterally up the opposite side. These are the vascular structures most easily accessible for IV therapy.

The external jugular vein is formed below the ear and behind the angle of the mandible (Fig. 21.3). It then passes downward and obliquely across the sternocleidomastoid and under the middle of the clavicle to join the subclavian vein. It is important to note the presence of valves in the external jugular, usually approximately 4 cm above the clavicle, because they can significantly impede IV function.29

PREPARATION

Safety
Universal precautions must be applied to all patients, especially in emergency care settings, in which the risk for exposure to blood is increased and the infection status of patients is largely unknown.30 One study showed that 11% of all hospital IV catheter injuries to health care workers occurred in the ED.31 Newer catheter devices have emerged that prevent inadvertent needle injuries (Fig. 21.4). The Protectiv IV Catheter Safety System (Smiths Medical, Minneapolis, MN) has a protective sleeve that encases the sharp stylet as it is retracted from the catheter. The needle of the Insyte Autoguard Shielded IV Catheter (Becton, Dickinson and Company, Franklin Lakes, NJ) is instantly encased inside a tamper-resistant safety barrel by pressing an activation button. The Saf-T-Intima IV catheter (Becton, Dickinson and Company), Punctur-Guard Safety Winged Set (Gaven Medical, Grand Island, NY), Vacutainer Safety-Lok (Becton, Dickinson and Company), Shamrock safety winged needle (Smiths Medical, Dublin, OH), and Angel Wing Safety Needle systems (Medtronic, Minneapolis, MN) are all
Scalp veins are commonly used in neonates. In patients who have undergone radical mastectomy, avoid the arm on the same side as the surgery because IVs may impair circulation, affect flow, or lead to edema and other complications. The lower extremity veins can also be useful locations, especially in pediatric patients. In patients who may need IV access quickly, the lower extremity veins are appropriate for larger-bore IV lines. Avoid veins that are not resilient and feel hard or cordlike because they are often thrombosed. Deep, percutaneous antecubital venipuncture and external jugular vein cannulation are also options in patients with difficult veins or those who may need IV access quickly. An 18-gauge catheter in the antecubital fossa is the standard device for IV contrast-enhanced computed tomography (CT) studies such as pulmonary CT angiogram.

### Choosing the Catheter Gauge

The specific gauge of catheter to use depends on the clinical scenario (Fig. 21.5). The narrowest catheter typically used in adults is a 22 gauge, which is sufficient for the routine administration of maintenance fluids and antibiotics. A 20 or 18 gauge is necessary for the administration of blood products, and a 16-gauge needle is preferred in resuscitation settings when large amounts of fluid must be given quickly. A second IV line at a different location allows additional IV therapy and also acts as a backup line in critical resuscitations. An 18-gauge catheter in the antecubital fossa is the standard device for IV contrast-enhanced computed tomography (CT) studies such as pulmonary CT angiogram.

### Appropriate Site

Site selection depends largely on the expected duration of IV therapy, the patient’s activity level, and the condition of the extremities. When choosing a location to initiate IV access, the best place to start is the hand and then advance cephalad as necessary. Hand veins are appropriate for 22-gauge catheters. Cephalic, accessory, or basilic veins are ideal for larger-bore IV lines. Avoid veins that are not resilient and feel hard or cordlike because they are often thrombosed. Deep, percutaneous antecubital venipuncture and external jugular vein cannulation are also options in patients with difficult veins or those who may need IV access quickly. The lower extremity veins can also be useful locations, especially in pediatric patients. In patients who have undergone radical mastectomy, avoid the arm on the same side as the surgery because IVs may impair circulation, affect flow, or lead to edema and other complications. Scalp veins are commonly used in neonates.

### Adjuncts for Finding a Vein

Patients often have nonvisible and nonpalpable veins. A common method of increasing venous distention is to ask patients to open and close their fist. Lowering the arm below the level of the heart can also increase venous distention. Light tapping can likewise be effective, although heavy tapping may cause the vein to spasm. If these methods are inadequate, heat packs can be applied for 10 to 20 minutes to increase venous engorgement. This is particularly useful in the pediatric population.

Nitroglycerin ointment applied to the hands of patients with small-caliber veins has been shown to increase the diameter of the vein by two to six times and increase the rate of successful first-attempt cannulation. Once the tourniquet is applied to the wrist, apply a quarter inch of 2% nitroglycerin to a 2.5-cm² area, leave it on for 2 minutes, and then rub it off. Nitroglycerin alone can have adverse effects in neonates and premature babies, however. The combination of topical nitroglycerin with local anesthetics has been shown to increase success and decrease the pain of cannulation in children 1 to 11 years old. This technique is contraindicated in hypotensive patients.

In the late 1980s, several small studies demonstrated the potential use of a venous distention device, a cardboard mailing tube that was placed over the forearm with a sealed bulb at one end that caused a vacuum within the tube. Of the patients predetermined to be difficult to access, 90% were cannulated when this device was used. Reported complications were few and included petechiae and discomfort.

### Anesthesia

Though somewhat time-consuming, local anesthesia at the site should be considered part of routine care. Local anesthesia significantly decreases pain before cannulation. Anesthetics such as lidocaine or bupivacaine may be instilled just beneath the skin at the site of planned cannulation through a tuberculin (1-mL) syringe equipped with a 27- or smaller-gauge needle. Adding bicarbonate (e.g., buffered lidocaine), warming the solution to room temperature, instilling the solution slowly, and distracting the patient during injection all contribute to reducing pain. In the pediatric population, 2.5 g of EMLA (eutectic mixture of local anesthetics) can be applied topically over the site. Its main disadvantage is slow onset, with as long as an hour needed for induction of anesthesia before cannulation. Other options include ethyl chloride topical spray, which temporarily numbs the skin, and oral sucrose in infants.

### IV Assembly

Review Box 21.1 itemizes the materials necessary for IV cannulation. The procedure is detailed in Fig. 21.6. The first step is to prepare the IV fluids and tubing. Remove the cap from the IV tubing and the tab from the IV bag. Clamp the IV tubing shut and insert the spiked end into the IV fluid bag. Pinch the drip chamber and fill it halfway. Open the clamp slightly to flush the IV tubing. If saline locks are being used, flush them similarly before cannulation. To do this, attach the lock to a saline-filled syringe and push saline through it.

### Inspection and Positioning

After collecting and preparing the equipment and supplies, palpate the veins. Position the patient comfortably on a flat surface. Place a 1-inch-wide tourniquet on the upper part of the patient’s arm or forearm and pull it sufficiently tight to impede venous flow but not tight enough to compromise arterial flow. Place the tourniquet under the arm. Fold both ends of it above the arm and cross the ends. Pull the overlying end taut and tuck the middle portion below the underlying end to
**Peripheral Intravenous Access**

1. Apply tourniquet to arm.
2. Prepare the insertion site with an alcohol pad.
3. Insert the IV catheter with the bevel facing upward.
4. Advance the needle until a flash of blood is seen.
5. Advance the catheter over the needle until flush with the skin.
6. Attach the preflushed saline lock.
7. Cover the insertion site with Tegaderm (or similar) dressing.
8. Attach IV tubing to the saline lock for IV fluid administration.

**Figure 21.6** Peripheral intravenous (IV) access. IV lines placed in the dorsum of the hand are associated with the lowest infection rates from venous cannulation.
create a loop. After placement of the tourniquet, palpate the veins with the index and middle finger of one’s nondominant hand. The veins are soft, elastic, resilient, and pulseless.28

**Cannulation**

Wash your hands, don gloves (nonsterile is adequate), and clean the injection site with iodine, alcohol, or both. Iodine is a better antiseptic than alcohol and results in fewer infections.45 If using alcohol, allow it to dry on the surface of the skin. Stabilize the vein without contaminating the prepared site. One method is to position one’s thumb alongside the vein and pull down while the index finger is positioned more cephalad and pulls upward. Take the angiocatheter between the thumb and forefinger of the dominant hand. With the bevel up, angle the angiocatheter 10 to 30 degrees between the catheter and the vein and parallel to the vein. Puncture the vein. Once a flash of blood is seen, advance the catheter several millimeters more to ensure that the catheter has entered the vein and not just the wall. Avoid advancing too far and puncturing the posterior wall. Loosen the stylet and advance only the catheter. Use the fingers that were anchoring the vein to occlude the vein at the tip of the catheter to prevent extravasation of blood from the angiocatheter. Remove the needle; connect the saline lock, IV tubing, or syringe for phlebotomy; and release the tourniquet.28

Cannulation of the external jugular vein deserves a special note (Fig. 21.7). In patients with otherwise limited peripheral access, it can be cannulated as follows. Place patient in the Trendelenburg position to fill the external jugular vein. Rotate the head to the opposite side and prepare the area as described earlier. Take the cannula and align it in the direction of the vein with the point aiming toward the ipsilateral shoulder. Puncture midway between the angle of the jaw and the midclavicular line while lightly compressing the vein with the free finger above the clavicle. Proceed as described previously for cannulation.

**Anchoring the Device**

After the IV cannula has been connected to the saline lock or IV tubing, anchor the device (see Fig. 21.6). Use a half-inch-wide strip of tape, adhesive side up, under the hub of the catheter and fold it over in a bow-shaped manner. This will secure the catheter and prevent lateral movement. Clear polyurethane dressings can also be used with or instead of tape. Saline locks can be connected to needless hubs to prevent accidental needle injury. Then secure the loose saline lock or IV tubing with tape to prevent accidental dislodgement. Connect the IV tubing to the angiocatheter and anchor it. Alternatively, use a commercially available securing device. Sign and date the dressings to ensure timely dressing changes.7 As an option, topical antibiotics or iodophor ointment may be applied to the insertion site to prevent infection, though the efficacy of doing so is unproven.46

**Dressing**

It is not cost effective to continually redress peripheral venous catheters at periodic intervals. Sterile gauze or transparent, semipermeable, polyurethane dressings can be left in place until removal of the catheter without increasing infection rates, as long as the site is regularly evaluated.50 Securing techniques that use proprietary devices, such as the StatLock IV (Bard Access Systems, Inc., Salt Lake City, UT), a sterile, adhesive-backed anchor, and distal male Luer-tip extensions, may reduce complications by decreasing mobility and risk of dislodgement.31 Commonly used topical antimicrobial ointments have not been consistently proved to reduce the rate of peripheral catheter-related infection but have been associated with increased rates of antimicrobial resistance and Candida colonization. Such ointments are not harmful in the ED, but their routine use is not supported.

**Maintaining Patency**

An important component of IV care is maintaining patency with frequent flushing. Until recently, heparin solutions were used to flush catheters and maintain patency, but heparin can cause problems such as hemorrhage. Saline flushes are as effective as heparin in maintaining patency and preventing phlebitis but do not carry the risks of bleeding or heparin-induced thrombocytopenia.47–49

**Percutaneous Brachial Vein Cannulation**

Brachial vein cannulation is an option when attempts at peripheral IV access have failed or are contraindicated and may obviate the need for central venous access or surgical cutdown. Complications include brachial artery puncture, hematoma, and transitory paresthesias.
To cannulate the brachial vein, palpate the brachial artery in the antecubital fossa. Prepare the site in the usual manner and apply a tourniquet above the antecubital space. At a point immediately medial or lateral to the pulse, insert an angiocatheter with an attached syringe and advance it at a 45-degree angle while maintaining suction on the syringe. After entrance into the vein, continue 2 to 3 mm further to ensure cannulation. Advance the catheter and remove the needle as usual.52,53

**COMPLICATIONS**

Although IV placement is a common procedure, it is not without complications. Fortunately, morbidity is rarely severe. Phlebitis, infiltration, infection, nerve damage, air embolism, bruising, and thrombosis are the most common complications and rarely cause significant morbidity or fatality.

Phlebitis is a common complication after IV cannulation and is described as the presence of a palpable cord accompanied by warmth, erythema, tenderness, and induration over the involved vein (Fig. 21.8). Phlebitis necessitates removal of the catheter and replacement on another extremity. Avoiding IV placement in the lower extremities (where there is more often stagnant blood flow) and across joints (where motion traumatizes the venous wall) minimizes the incidence of IV catheter–related phlebitis.7 Other causes of phlebitis include IV infusion of potassium chloride, certain antibiotics (vancomycin, erythromycin), many cytotoxic chemotherapy agents, phenytoin, and any hyperosmolar solution (e.g., 50% dextrose solution).52,53 The role of in-line filters to prevent phlebitis is controversial. Particulates from reconstituted medications, degradation products, precipitates, glass from vials, and other foreign debris may all play a part in postinfusion phlebitis. In-line filters may therefore play a role in preventing phlebitis, but given their cost, risk of clogging, and paucity of evidence that they improve outcomes, these filters have not become routine.54

Even with the most pristine technique, there is approximately a 0.5% incidence of catheter-related bloodstream infection with peripheral IV catheters. IV devices facilitate infection by damaging epithelial barriers and thereby providing microorganisms direct access to the bloodstream.54 The risk for infection with peripheral venous catheters is higher in the lower extremity than in the upper extremity and higher in the wrist or upper part of the arm than in the hand. The most common infectious complication of peripheral IV access is self-limited cellulitis.56 The safety of maintaining peripheral IV lines for up to 72 hours before they are relocated has been established in a large, prospective study.50 With rates of clinically significant bacteremia lower than 0.5%, some argue that routine replacement of catheters is now no longer needed.57 Nonetheless, infection can be a costly and potentially devastating complication of IV therapy. Suppurative thrombophlebitis is extremely rare. It most frequently occurs in patients with thermal injury and long-term or lower extremity cannulation.55 Local signs of inflammation or suppuration are often absent and can occur 2 to 10 days after removal of the catheter.58 Treatment is immediate surgical excision of the entire length of the involved vein and tributaries. Though rare with peripheral IV catheters, intravascular device–related bloodstream infection may be an unrecognized cause of nosocomial infection. Peripheral IV catheters are most often associated with staphylococcal and candidal species.59 Infectious complications can be reduced significantly by hand washing, wearing gloves, preparing the site with iodine, and monitoring the site for signs of infection.7

Bruising is a common complication of IV therapy. Contrary to popular belief, flexing of the elbow after venipuncture does not prevent bruising in the antecubital site.60 Applying direct pressure immediately after decannulation is the most useful technique to prevent bruising.

Tissue or interstitial infiltration occurs when the catheter is dislodged from the vein during infusion. It is a common and usually relatively minor complication of IV therapy. Extravasation of certain infusions, such as hypertonic solutions, vasopressors, or chemotherapeutic agents, however, poses a significant risk for necrosis and skin sloughing when infiltration and extravasation occur. In extreme cases, skin grafting may be required.7 For extravasation of dopamine or epinephrine, local injection of antidotes such as phentolamine may be used to reverse the tissue damage.61

Nerve injury is rare after IV cannulation. Any peripheral nerve is potentially vulnerable to a needle-induced injury, and sequelae can range from a minor motor or sensory abnormality to complete paralysis. Nerve damage may be due to direct injury by the needle, intraneural microvascular damage from hematomas, or toxic effects of the agent injected.40 The first symptoms are usually pain, numbness, or paresthesia. Pain may persist for years and can be debilitating. Fortunately, most simple procedures do not result in nerve injury because nerves tend to roll or slide away from a needle. Like all procedures, knowledge of the relevant anatomy is essential. Should a patient complain of numbness or severe pain after a needle puncture, stop the injection immediately.61,54

Thrombosis and subsequent pulmonary embolism (PE) are commonly associated with centrally placed IV catheters.7 Though rare, thrombosis followed by clinically significant PE may occur in patients with peripheral IV lines if saline locks are not flushed or IV fluids run dry. Should this occur, aspirate the line. If the return fluid appears bloody, discard the syringe and then gently flush the saline lock and resume the infusion. If there is no bloody aspirate, use 2 to 3 mL of saline to gently flush the line. If resistance is encountered, stop flushing immediately because there is a risk for development of an embolism. Attempt IV insertion at another site.53

Venous air embolism is another significant, though exceedingly rare complication of peripheral IV access. Symptoms include chest pain, shortness of breath, sudden vascular collapse, cyanosis, and hypotension. If air embolism is suspected, place...
the patient in the left lateral decubitus Trendelenburg position. Invasive maneuvers include aspiration of air through a central venous catheter and even thoracotomy with direct aspiration from the heart (see Chapter 18). This complication can be prevented by eliminating air from the IV tubing before initiating therapy and not allowing IV lines to run dry. If air bubbles are present in an IV line, tap the tubing while holding it taut to allow the air to escape to the top. Similarly, curl the tubing around a pen or syringe to accomplish the same goal. If the air is near a Y-connector, one can use a needle and syringe to directly remove it. If all else fails and there is air between the Y-connector and the patient, disconnect the tubing and flush it.66

Recommendations of the Centers for Disease Control and Prevention for IV catheter care to prevent complications are as follows:
1. Record and date the time of catheter insertion in an obvious location near the insertion site.
2. Do not palpate the insertion site after the skin has been cleansed with antiseptic.
3. Palpate the insertion site for tenderness daily through an intact dressing.
4. Visually inspect the site if the patient reports tenderness.
5. Wash hands before and after palpating, inserting, replacing, or dressing any intravascular access site.
6. Replace dressings when they are damp, loose, or soiled.65

EXTRAVASATION OF MEDICATIONS AND VASOPRESSORS

Usually, infiltration of a vein is a relatively minor and common complication of IV therapy if only sterile fluid extravasates, even in large amounts. This often occurs when the catheter is dislodged from the vein during infusion. However, if the infusions consist of hypertonic substances, vasopressors, or chemotherapeutics, there is a significant risk for skin sloughing if infiltration and extravasation occur (Box 21.1). Pain at the infusion site or the alarm sounding on an infusion pump device requires inspection of the infusate site for extravasation. In extreme cases, grafting may be required for skin sloughing (Fig. 21.9).7 If dopamine, phenylephrine (Neo-Synephrine, Hospira, Inc., Lake Forest, IL), or norepinephrine extravasate, phentolamine may be used as an antidote to prevent ischemia locally; its use is encouraged as soon as extravasation is identified. Reversal of ischemia with phentolamine is a common technique, but its ability to totally reverse or prevent skin sloughing is not guaranteed. However, if infiltration of these vasopressors occurs, the authors suggest that it be used routinely. There are few downsides to this intervention, although hypotension is a theoretical side effect because phentolamine is an α-adrenergic antagonist. To inject phentolamine, place 5 mg in a vial and dilute with equal parts of saline (final form: 5 mg in 2 mL). For large areas, use two vials with the contents of each vial injected 10 minutes apart through a 25- to 27-gauge needle or a tuberculin syringe. If the IV line is still in place, inject approximately 1 mL of phentolamine through the catheter before it is removed; however, the IV line is often removed before this can be done. The entire area of skin blanching, or suspected area of extravasation, is injected with multiple small aliquots of the solution, approximately 0.25 to 0.5 mL each. The procedure may be repeated in 2 to 4 hours. Hyaluronidase is probably benign and has been suggested in the past to ameliorate some effects of extravasation of other solutions. Though it has been a common suggestion, its efficacy has not been well established, and the product is not readily available. Ice and heat have varying effects in counteracting fluid extravasation. Extravasation of IV contrast material is discussed in Chapter 36.

<table>
<thead>
<tr>
<th>Medications and Solutions That May Cause Tissue Injury When Extravasation Occurs in a Peripheral Vein</th>
<th>Aminophylline</th>
<th>Mithramycin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium chloride 10%</td>
<td>Mitomycin</td>
<td></td>
</tr>
<tr>
<td>Carmustine</td>
<td>Nafcin</td>
<td></td>
</tr>
<tr>
<td>Chloridiazepoxide</td>
<td>Neo-Synephrine (Hospira, Inc., Lake Forest, IL)</td>
<td></td>
</tr>
<tr>
<td>Colchicine</td>
<td>Nitroglycerin</td>
<td></td>
</tr>
<tr>
<td>Crystalline amino acids</td>
<td>Norepinephrine</td>
<td></td>
</tr>
<tr>
<td>4.25%/dextrose 10%</td>
<td>Parenteral nutrition solutions</td>
<td></td>
</tr>
<tr>
<td>Crystalline amino acids</td>
<td>Phenytoinb</td>
<td></td>
</tr>
<tr>
<td>4.25%/dextrose 25%</td>
<td>Potassium solutions</td>
<td></td>
</tr>
<tr>
<td>Dactinomycin</td>
<td>Propylene glycol</td>
<td></td>
</tr>
<tr>
<td>Daunorubicin</td>
<td>Renografin-60 (Bracco Diagnostics Inc., Monroe Township, NJ)</td>
<td></td>
</tr>
<tr>
<td>Dextrose 10%</td>
<td>Sodium bicarbonate 8.4%</td>
<td></td>
</tr>
<tr>
<td>Dextrose 50% in water</td>
<td>Sodium thiopental</td>
<td></td>
</tr>
<tr>
<td>Diazepam</td>
<td>Tetracycline</td>
<td></td>
</tr>
<tr>
<td>Dobutamine</td>
<td>Vasopressin</td>
<td></td>
</tr>
<tr>
<td>Dopamine</td>
<td>Vinblastine</td>
<td></td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>Vincristine</td>
<td></td>
</tr>
<tr>
<td>Epinephrine</td>
<td>Methchlorhethamine</td>
<td></td>
</tr>
<tr>
<td>Ethyl alcohol</td>
<td>Metaraminol</td>
<td></td>
</tr>
<tr>
<td>Methchlorhethamine</td>
<td>Vindesine</td>
<td></td>
</tr>
<tr>
<td>Phenytoin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sodium bicarbonate 8.4%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Potassium solutions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Propylene glycol</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Renografin-60 (Bracco Diagnostics Inc., Monroe Township, NJ)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sodium bicarbonate 8.4%</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sodium thiopental</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tetracycline</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vasopressin</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vinblastine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vincristine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methchlorhethamine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Metaraminol</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vindesine</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

4 Many medications and intravenous solutions will cause pain and occasionally skin sloughing if significant amounts extravasate into soft tissue. Thus, any complaint of pain during infusion or signs of tissue bulging should prompt an investigation for extravasation. Most extravasations have no specific therapy, so prevention is the only option. Phentolamine, injected subcutaneously to reverse vasoconstriction, is the most common technique, but its efficacy has not been well studied.

6 Use a maximum concentration of 2 mg/mL of saline or fosphenytoin solution to minimize this risk.

Figure 21.9 A, Infiltration of calcium chloride in an infant. Once this occurs, there is no treatment except débridement and possible skin grafting. Calcium gluconate will not cause such a reaction. Extravasation of hypertonic dextrose, phenytoin, and vasoconstrictors or vasopressors will cause similar necrosis. B, Full-thickness tissue injury from doxorubicin extravasation, not obvious until 7 to 10 days after the infusion.
Extravasation of chemotherapy solutions is particularly common and can produce full-thickness tissue sloughing. The patient may complain of pain and burning at the time of infusion, but skin sloughing may be delayed for many days. Table 21.1 lists possible antidotes and dosages for chemotherapy-induced extravasation injury. Results of these interventions vary.

**TABLE 21.1 Possible Antidotes for Extravasated Chemotherapeutic Agents**

<table>
<thead>
<tr>
<th>CHEMOTHERAPEUTIC AGENT</th>
<th>ANTIDOTE</th>
<th>DOSE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anthracycline</td>
<td>Dexrazoxane hydrochloride&lt;sup&gt;b&lt;/sup&gt;</td>
<td>First dose, inject the equivalent of 500 mg dexrazoxane intravenously over 1–2 hr, second dose at 24 hr, and third dose at 48 hr.</td>
</tr>
<tr>
<td>Mechlorethamine</td>
<td>Sodium thiosulfate</td>
<td>Multiple subcutaneous injections in and around the area of extravasation with a 25-gauge needle: 4 mL of 10% sodium thiosulfate + 6 mL water.</td>
</tr>
<tr>
<td>Vinca alkaloids (vincristine, vinblastine, and vinorelbine)</td>
<td>Hyaluronidase</td>
<td>Inject subcutaneously in and around the area of extravasation with a 25-gauge needle: 150 units (1 mL). For vinca alkaloids, apply local hot compresses.</td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>Granulocyte-macrophage colony-stimulating factor&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Inject subcutaneously in and around the area of extravasation with a 25-gauge needle.</td>
</tr>
<tr>
<td>Doxorubicin, daunorubicin, and mitomycin</td>
<td>Dimethyl sulfoxide (free radical scavenger)</td>
<td>Apply a 50%–70% solution topically qid for 14 days. Leave uncovered.</td>
</tr>
<tr>
<td>Mitomycin</td>
<td>Pyridoxine&lt;sup&gt;d&lt;/sup&gt;</td>
<td>Inject subcutaneously in and around the area of extravasation with a 25-gauge needle.</td>
</tr>
<tr>
<td>Nonspecific</td>
<td>Saline</td>
<td>Inject subcutaneously in and around the area of extravasation with a 25-gauge needle.</td>
</tr>
<tr>
<td>Nonspecific</td>
<td>Corticosteroids&lt;sup&gt;d&lt;/sup&gt;</td>
<td>Inject subcutaneously in and around the area of extravasation with a 25-gauge needle: hydrocortisone, 500 mg diluted in 500 mL saline.</td>
</tr>
</tbody>
</table>

<sup>a</sup>Many of these interventions are anecdotal and none are guaranteed to reverse or ameliorate tissue injury. Controversy surrounds the actual benefit, and no randomized prospective trials have been conducted for many of the suggested regimens. Also consider elevation and surgical débridement when necessary.

<sup>b</sup>Approved by US Food and Drug Administration for this indication.

<sup>c</sup>Not well studied, theoretical benefit.

<sup>d</sup>Results are variable; injury is not an inflammatory reaction.

Injury from extravasation of phenytoin can be minimized or avoided by using dilute solutions, no more than a 2-mg/mL concentration (1 g in 500 mL saline), or by using fosphenytoin instead of phenytoin. When possible, use calcium gluconate, not calcium chloride, in a peripheral IV line.

The bottom line is that most extravasated chemotherapy and other agents have no specific antidote or reversal agents.

### ULTRASOUND BOX 21.1: Peripheral Intravenous Access by Christine Butts, MD

Ultrasound-guided access is indicated when standard placement is difficult. This may include patients with no palpable or visible peripheral veins, history of intravenous drug use or multiple previous peripheral lines causing scarring or thrombosis, obesity, or previous surgeries causing distortion of the anatomy. Use of ultrasound to achieve peripheral intravenous access has been found to increase the rate of success, decrease both the time to placement and the number of attempts, and increase overall patient satisfaction. Nursing use of ultrasound to guide peripheral intravenous access has also shown promise in improving success and decreasing complications.<sup>2</sup>

There are two methods by which the ultrasound may be used to access peripheral veins. In the first method, ultrasound is used to simply evaluate the underlying anatomy. Frequently, veins are present superficially but cannot be seen or palpated from the surface of the skin. Apply the transducer to rapidly locate an ideal vein and cannulate it “blindly” in the typical fashion. The second method calls for ultrasound to be used to directly guide venous access. This method may be most practical when the veins are deeper within the tissues or adjacent to other more important structures.

Use a high-frequency (7.5-to 10-MHz) transducer to obtain the necessary resolution for evaluating the anatomy. Typically, for peripheral venous access, a sterile field is not necessary. However, it is important to clean the area with alcohol or chlorhexidine solution before the procedure. Use universal precautions. Apply a tourniquet when appropriate to assist in distending the veins and to make them easier to identify.
Multiple peripheral vessels are available for ultrasound-guided access. The basilic vein lies on the ulnar aspect of the forearm and the cephalic vein can be found on the radial aspect (Fig. 21.US1). The larger median cubital vein represents the junction of these two veins and lies in the antecubital fossa (Fig. 21.US2). The deep brachial vein is found on the median aspect of the distal end of the arm in the bicipital groove (Fig. 21.US3). The external jugular vein is found superficially in the neck and runs diagonally across the sternocleidomastoid muscle.

Distinguishing the artery from the vein may prove more challenging than with central vessels. Peripheral veins and arteries are smaller, and even arteries may collapse with pressure. Evaluate the veins before application of the tourniquet. Veins should collapse easily and in fact may collapse from only the pressure of applying the transducer to the skin. Placing the heel of the operator’s hand on the patient’s arm and then applying the transducer may decrease this effect. Arteries can be further identified by evaluating the Doppler flow pattern of the vessel. Even small arteries will have a biphasic, pulsatile flow pattern versus the steady low-amplitude venous pattern. Finally, venous augmentation can be used for confirmation. Color flow imaging is used over the vessel in question. Squeezing the arm distal to the area should cause a temporary flush of flow in the vein (Fig. 21.US4). The artery should not show any change in its typical pulsatile flow.

Once a location for cannulation has been chosen, scan the relevant area to locate the desired vessel. Once the vein has been identified, the cannulation procedure is identical for all peripheral veins. A transverse, or short-axis, approach is universally used because of the small size of the peripheral veins. Center the transducer over the target vessel. The depth of the image on-screen should be
to alter the final outcome. At most extravasation sites it may be best to avoid the empirical use of suggested treatments such as sodium bicarbonate, sodium thiosulfate, heparin, calcium gluconate, magnesium sulfate, lidocaine, cimetidine, diphenhydramine, and other chemical substances that are believed to inactivate drugs and reduce toxic effects on cells. In some experimental settings, these substances have made the necrosis and ulceration worse.54–57

A pitfall that may frustrate the sonographer is kinking or difficulty threading or advancing the catheter once a flash has been obtained. This frequently occurs in deeper vessels, such as the deep brachial. Using a longer, stiffer catheter [e.g., 1.75-inch Arrow twin catheter (Teleflex, Morrisville, NC) or the longer catheter from the Arrow arterial line kit] may help reduce this problem.

References

References are available at www.expertconsult.com
REFERENCES


Central venous access remains a cornerstone of resuscitation and critical care in both the emergency department (ED) and intensive care unit. Advanced hemodynamic monitoring, rapid infusion of fluid, placement of transvenous pacemakers, and administration of selected medications all require reliable central venous access. Central venous catheterization has also gained acceptance in the resuscitation and treatment of critically ill children (see Chapter 19). Traditionally, the subclavian vein (SV), internal jugular (IJ) vein, and femoral vein have provided reliable and easily obtainable vascular access through the use of identifiable anatomic landmarks. Over the past decade the increased availability of, and training and provider competence in bedside ultrasonography have had a significant impact on the standard approach to both peripheral and central venous catheterization. Ultrasound-guided central venous catheterization has improved success rates, reduced complication rates, decreased the time required to perform the procedure, and resulted in overall cost savings.

The various techniques described in this chapter each have inherent advantages and disadvantages, but all have a place in the practice of emergency medicine. Frequently, a clinician’s previous experience with a particular technique determines the preferred approach, but clinicians responsible for acute resuscitation of the ill and injured should master several of these techniques (Videos 22.1-22.8).

**HISTORICAL PERSPECTIVE**

In 1667, Lower placed the first known central venous catheter (CVC) into a human IJ vein for a blood transfusion from the carotid artery of a sheep. Modern central venous catheterization heralds back to at least 1928 when Werner Forssmann, a 25-year-old German surgeon, performed a venous cutdown on his own left antecubital vein, inserted a ureteral catheter to a distance of 65 cm, and then climbed several flights of stairs to the radiology suite to confirm that it terminated in the right atrium. Although the hospital fired Dr. Forssmann for not obtaining permission, he went on to win the 1956 Nobel Prize for his pioneering efforts.1,2

Duffy reported a large series of femoral, jugular, and antecubital vein catheterizations in 1949.3 Aubaniac developed subclavian venipuncture while working on French Army casualties between 1942 and 1952.4 His infraclavicular SV approach was refined by Keerl-Szanto in 1956, and the supraclavicular approach to the vein was first described by Yoffa in 1965.5,6 Aside from Duffy’s earlier work, Hermosura (1966) and English (1969) are generally credited with the scientific development of the percutaneous IJ approach.7

The most important advancement in modern CVC came in 1953 when the Swedish radiologist Sven Seldinger had the idea of advancing large catheters over a flexible wire that was inserted through a percutaneous needle.8,9 The role of central venous pressure (CVP) monitoring in the maintenance of optimal blood volume helped popularize central catheterization in the United States.10 This was accelerated by the advent of the pulmonary artery catheter, which was developed by Jeremy Swan and William Ganz in 1968.11 Swan, who was inspired by his observations of a sailing boat during a picnic with his children, developed a flow-directed balloon that allowed measurement of pulmonary artery pressure.12

**ANATOMY**

**SV System**

The SV begins as a continuation of the axillary vein at the outer edge of the first rib. It joins the IJ vein to become the innominate (sometimes referred to as the brachiocephalic) vein 3 to 4 cm proximally. The SV has a diameter of 10 to 20 mm, is approximately 3 to 4 cm long, and is valveless. After crossing over the first rib, the vein lies posterior to the medial third of the clavicle. It is only in this area that there is an intimate association between the clavicle and the SV. The costoclavicular ligament lies anterior and inferior to the SV, and the fascia contiguous to this ligament invests the vessel. Posterior to the vein and separating it from the subclavian artery is the anterior scalene muscle, which has a thickness of 10 to 15 mm. The phrenic nerve passes over the anterior surface of the scalene muscle laterally and runs immediately behind the junction of the SV and the IJ vein. The larger thoracic duct (on the left) and the smaller lymphatic duct (on the right) pass over the anterior scalene muscle and enter the SV near its junction with the IJ vein. Superior and posterior to the subclavian artery lies the brachial plexus. The dome of the left lung may extend above the first rib, but the right lung rarely extends this high (Fig. 22.1).

**IJ Vein**

The IJ vein begins just medial to the mastoid process in the jugular foramen at the base of the skull and is formed by the inferior petrosal sinus and the sigmoid sinus. It runs inferiorly and passes under the sternal end of the clavicle to join the SV and form the innominate or brachiocephalic vein. At the level of the thyroid cartilage, the IJ vein, the internal carotid artery, and the vagus nerve course together in the carotid sheath just deep to the sternocleidomastoid (SCM) muscle. Within the carotid sheath, the IJ vein typically occupies the anterior lateral position and the carotid artery lies medial and slightly posterior to the vein. This relationship is relatively constant, but studies have found that the carotid artery may overlap the IJ vein. Note that normally the IJ vein migrates medially as it nears the clavicle, where it may lie directly over the carotid artery. When using the most common (central) approach, the IJ vein tends to be more lateral than expected.14,15 Furthermore, in 5.5% of those studied, the IJ vein may even be medial to the carotid artery.16,17 The relationship between the IJ vein and the carotid artery also depends on head position. Excessive head rotation can cause the carotid artery to rotate over the IJ vein.18,19
Central Venous Catheterization

**Indications**
- Central venous pressure monitoring
- High-volume/flow resuscitation
- Emergency venous access
- Inability to obtain peripheral venous access
- Repetitive blood sampling
- Administering hyperalimentation, caustic agents, or other concentrated fluids
- Insertion of transvenous cardiac pacemakers
- Hemodialysis or plasmapheresis
- Insertion of pulmonary artery catheters

**Contraindications**
- Infection over the placement site
- Distortion of landmarks by trauma or congenital anomalies
- Coagulopathies, including anticoagulation and thrombolytic therapy
- Pathologic conditions, including superior vena cava syndrome
- Current venous thrombosis in the target vessel
- Prior vessel injury or procedures
- Morbid obesity
- Uncooperative patients

**Equipment** (contents of a typical central venous catheterization kit)

- Sterile drape
- Chlorhexidine
- 1% lidocaine without epinephrine
- 5-mL syringe (for anesthetic)
- 25- and 22-gauge needles (for anesthetic)
- Scalpel with a No.11 blade
- Dilator
- 18-gauge needle (for venipuncture)
- 5-mL syringe (for venipuncture)
- Triple-lumen catheter
- Guidewire
- Catheter clamp
- Silk suture

**Complications**
- Arterial puncture and hematoma
- Pneumothorax (subclavian and internal jugular approach)
- Hemothorax (subclavian and internal jugular approach)
- Vessel injury
- Air embolism
- Cardiac dysrhythmia
- Nerve injury
- Infection
- Thrombosis
- Catheter misplacement

Review Box 22.1 Central venous catheterization: indications, contraindications, complications, and equipment.
Anatomic landmarks for locating the vein include the sternal notch, the medial third of the clavicle, and the SCM. The two heads of the SCM and the clavicle form a triangle that is key to understanding the underlying vascular anatomy. The IJ vein can be located at the apex of the triangle as it courses along the medial head of the SCM and occupies a position in the middle of the triangle at the level of the clavicle before it joins the SV and forms the innominate vein. At the level of the thyroid cartilage, the IJ vein can be found just deep to the SCM.

Generally, the right IJ vein is bigger than the left IJ vein because of its connection to the SV and the right atrium. The IJ vein can be pulsatile, but in contrast to the aorta, these pulsations are not palpable. When visualized, however, the presence of venous pulsations can give an indication of patency of the IJ vein to the right atrium. The IJ vein also changes size with respiration. Because of the negative intrathoracic pressure at end-inspiration, blood in the IJ vein is actually drawn into the right atrium and the diameter of the IJ vein shrinks. In contrast, at end-expiration the increased intrathoracic pressure will limit return of blood to the right atrium and the diameter of the IJ vein will increase. Another unique characteristic of the IJ vein is its distensibility. The IJ vein will enlarge when pressure in the vein is increased, such as when flow of blood back to the right atrium is obstructed, as with thrombosis. This distensibility can be advantageous for the placement of central venous access. Use of a head-down (Trendelenburg) position or a Valsalva maneuver will increase the diameter of the IJ vein and thereby increase the likelihood of successful puncture (Fig. 22.2).

**Femoral Vein**

The femoral anatomy is less complex than that of the neck and shoulder and contains fewer vital structures. The femoral vein is most easily cannulated percutaneously in patients with a palpable femoral pulse. The femoral vein begins at the adductor canal (also known as Hunter's canal) and ends at the inferior margin of the inguinal ligament, where it becomes the external iliac vein. It is contained within the femoral triangle (inguinal ligament, medial border of the adductor longus, and lateral border of the sartorius muscle). Medially, the femoral vein abuts a robust system of lymphatics. Laterally, the vein is intimately associated with the femoral artery. The femoral nerve courses down into the leg just lateral to the femoral artery. These relationships from lateral to medial can be remembered with the mnemonic NAVEL (nerve, artery, vein, empty space, lymphatics). Note that as the femoral artery and vein course down the leg within the femoral sheath, their side-by-side relationship frequently rotates such that the femoral artery may lie on top of the vein. Therefore to avoid arterial puncture, keep cannulation attempts just under the inguinal ligament. When cannulating this vessel distal to the inguinal ligament, ultrasound guidance can be helpful to avoid arterial puncture (Fig. 22.3).

**INDICATIONS**

Central venous access has several clinical indications (see Review Box 22.1). If necessary, any central venous approach could be used for each of these situations. However, certain approaches offer advantages over others in specific clinical settings. The clinical indications are discussed in detail in the following sections.20–22

**CVP Monitoring and Oximetry**

For a period, pulmonary artery catheterization somewhat supplanted CVP monitoring; however, there is little evidence that this practice has any benefit with regard to patient mortality or quality of life. In the specific setting of resuscitation of patients in septic shock, CVP monitoring is an important component of “early goal-directed therapy” (EGDT).21,24 Continuous or episodic measurements of central venous oxygen saturation (ScVO_{2}) also have a prominent role in EGDT protocols for the aggressive treatment of septic shock.24 More recent evidence, however, does not demonstrate improvements
with severe sepsis or septic shock. As a result, there has been a de-emphasis on CVP monitoring in septic shock.

The rate of flow is proportional to the radius of the catheter because of the properties of Poiseuille's law, which states that the flow rate of fluid is inversely proportional to the radius of the catheter and directly proportional to the length. To illustrate, the gravity flow rate of saline through a peripheral 5-cm, 14-gauge catheter is roughly twice that through a 20-cm, 16-gauge CVC. Consequently, placement of large-bore peripheral catheters is generally the fastest method of volume loading.

**Delivery of High-Flow Fluid Boluses and Blood Products**

The advent of thermoregulating high-volume rapid infusers affords the advantage of using central venous catheterization in the setting of severe hemorrhagic shock or hypothermia. The available systems can infuse blood warmed to 37°C through an 8.5-Fr introducer sheath 25% more rapidly than through a 14-gauge peripheral intravenous (IV) line and up to 50% faster than through an 18-gauge peripheral IV line. The Level 1 Rapid Infuser (Smiths Medical, St. Paul, MN) and the Belmont FMS 2000 (Belmont Instrument Corporation, Billerica, MA) are examples of modern systems with infusion rates as high as 1500 mL/min. Massive air embolism was a concern with early rapid infusers, but safety precautions have now been engineered to prevent this. However, if the catheter is misplaced, fluid or blood can be rapidly infused into the thorax, mediastinum, or peritoneum with serious consequences.

**Emergency Venous Access and Inability to Achieve Peripheral Access**

The predictable anatomic locations of the subclavian and femoral veins and the speed with which they can be cannulated have prompted their use in cardiac arrest and other emergency situations. The need for a central line during cardiopulmonary resuscitation (CPR) is controversial. When achieved easily, central venous cannulation, especially via the IJ vein or SV route, is preferred over peripheral venous access because it provides a rapid and reliable route for the administration of drugs to the central circulation of patients in cardiac arrest. With resuscitation for aortic catastrophes or thoracoabdominal trauma, two CVCs, “one above and one below” the diaphragm, are often used.

Patients with a history of IV drug use, major burns, chronic disease, dehydration, or morbid obesity and those who require long-term access may have inadequate peripheral IV sites. Central venous cannulation may be indicated as a means of venous access in these patients even under nonemergency conditions. More recently with the use of ultrasound, deep brachial, axillary, and basilic vein cannulation may be attempted before central venous catheterization. This approach avoids the complications that can be associated with central venous access.

**Routine Serial Blood Drawing**

The potential complications of CVCs make them inappropriate solely for routine blood sampling. However, lines already in place may be used for this purpose if they are properly cleared of IV fluid. A 20-cm, 16-gauge catheter contains 0.3 mL of fluid, so at least this much must be withdrawn to avoid dilution of blood samples. Moreover, to avoid aspiration of crystalloid-diluted blood from the peripheral vein, it is advised that the IV line be turned off for at least 2 to 3 minutes before using the catheter for a blood draw. Because of the increased risk for infectious complications, air embolism, and venous back-bleeding, the IV tubing should not be repeatedly disconnected from the catheter hub. Interposition of a three-way stopcock in the IV tubing simplifies access and is an acceptable method for sampling blood in the intensive care setting, regardless of the IV site. The oxygen level in blood from the SV can be determined for guidance in EGDT for sepsis if one chooses not to place a continuous oximetric monitor.

Additionally, serial lactate levels may help guide early goal-directed resuscitation. With an imbalance in oxygen supply and consumption, tissue hypoperfusion and hypoxia lead to anaerobic metabolism. The final product of this process is lactate. Arterial lactate levels would best represent overall perfusion because such samples contain blood coming from the pulmonary veins, superior vena cava (SVC), and inferior vena cava (IVC). Peripheral lactate preferentially reflects perfusion and metabolism in the compartment from which the blood was drawn, but not overall perfusion. Arterial and central venous lactate correlate closely more than 96% of the time, whereas peripheral venous lactate and arterial lactate correlate only 87% of the time.19
Infusion of Hyperalimentation and Other Concentrated Solutions

Central venous hyperalimentation is safe and reliable. Use of the infraclavicular subclavian technique frees the patient’s extremities and neck; this procedure is therefore well suited for long-term applications. Hyperosmolar or irritating solutions with the potential to cause thrombophlebitis if given through small peripheral vessels are frequently infused through central veins. Examples are potassium chloride (>40 mmol/L), hyperosmolar saline, 10% calcium chloride (but not calcium gluconate, which can safely be given peripherally), 10% dextrose infusions, chemotherapeutic agents, and acidifying solutions such as ammonium chloride. Vasoactive substances (e.g., dopamine, norepinephrine) are best administered through a CVC because they may cause soft tissue necrosis if extravasation occurs in peripheral sites.

Central catheters, though safer than peripheral IV lines, are not immune to extravasation; indeed, fatal cases have been reported if the catheter becomes wedged up against the vessel wall, valves, or endocardium. Strategies to avoid this complication include delivering vesicant drugs only through the distal ports or reconfirming that the proximal port is safely in the vein by aspirating blood through it.

Other Indications

Other indications for central venous access include insertion of a pulmonary artery catheter or transvenous pacemaker, cardiac catheterization, pulmonary angiography, and hemodialysis. A pulmonary artery catheter can be valuable for determining fluid and hemodynamic status in the critically ill. Its widespread use in the 1980s and 1990s drew heavy criticism because data showing a benefit in patient-oriented outcomes were lacking. Pulmonary artery catheters have subsequently lost popularity and should be used only when the diagnostic benefits outweigh the potential risks.

Catheters such as the Uldall and Quinton devices can be inserted within minutes, thereby permitting emergency or short-term hemodialysis. However, these catheters are very large and relatively stiff and have been known to perforate the vena cava or atrial walls, with fatal outcomes. Extra caution should be applied during their insertion, possibly under ultrasound or fluoroscopic guidance.

CONTRAINDICATIONS

General contraindications to the various techniques of central venous access are presented in Review Box 22.1. Table 22.1 lists the general advantages and disadvantages with each approach. Most contraindications listed are relative and should be viewed in the context of the patient’s overall condition, urgency of need, and availability of alternative options for vascular access. Perhaps the only true absolute contraindication is insertion of catheters impregnated with antibiotics (most commonly tetracycline, rifampin, or chlorhexidine) if the patient has a serious allergy to the drug.

Local cellulitis and distorted local anatomy or landmarks preventing safe insertion are relative contraindications to any access route. Insertion of catheters through freshly burned regions, though somewhat challenging, is not associated with a higher incidence of infection until approximately 3 days after the burn, when bacterial colonization accelerates. One of the more commonly encountered impediments to CVC placement is morbid obesity. Surface landmarks in the neck are often obscured, and an abdominal pannus can block the femoral access site and consequently require deeper insertions and steeper angles. An ultrasound-guided

<table>
<thead>
<tr>
<th>TABLE 22.1 Advantages and Disadvantages of Central Venous Access Techniques</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TECHNIQUE</strong></td>
</tr>
<tr>
<td>IJ</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Femoral</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>SV, infraclavicular</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>SV, supraclavicular</td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

CVP, Central venous pressure; IJ, internal jugular; SV, subclavian vein.
IJ approach is safer under these circumstances. Insertion of another catheter on the same side as a preexisting one risks the complication of entrapment. Combativeness is an important factor in the decision to place a CVC because the risk for mechanical complications greatly increases in uncooperative patients. Sometimes it is best to sedate and intubate critically ill patients before attempting central venous catheterization. Other relative contraindications include conditions predisposing to sclerosis or thrombosis of the central veins, such as vasculitis, previous long-term cannulation, or illicit IV drug use via any of the deep venous systems.

Coagulopathy is a frequent concern surrounding insertion of a CVC, with the overall risk for clinically significant hemorrhage in these patients approximating 2%. A transfusion of fresh frozen plasma is commonly used to correct any existing coagulopathy. However, a 2005 review concluded that if good technique is used, correction of coagulopathy is not generally required before or during the procedure. Mumtaz and coworkers found that even in thrombocytopenic patients (platelet count <50 × 10^9/L), bleeding complications occurred approximately 3% of the time and were limited to bleeding at the insertion site; these complications were managed with additional sutures. Although the occasional patient may require a blood transfusion or replacement of clotting factors if a hemorrhagic complication arises, prophylactic correction of an abnormal international normalized ratio or platelet count before the procedure is not routinely necessary. Risk can be further reduced in coagulopathic patients with the use of ultrasound-guided placement techniques.

SV Approach

SV access is contraindicated in patients who have previously undergone surgery or sustained trauma involving the clavicle, the first rib, or the subclavian vessels; those with previous radiation therapy involving the clavicular area; those with significant chest wall deformities; and those with marked cachexia or obesity. Patients with unilateral deformities not associated with pneumothorax (e.g., fractured clavicle) should be catheterized on the opposite side. Subclavian venipuncture is not contraindicated in patients with penetrating thoracic wounds unless the injuries are known or suspected to involve the subclavian vessels or SVC. Generally, cannulate the vein on the same side as the chest wound to avoid the possibility of bilateral pneumothoraces. When (preexisting) SV injury is suspected, cannulate on the opposite side. Exercise greater caution when placing a CVC in the SV in coagulopathic patients because this vessel is not compressible. Formerly, subclavian venipuncture was not recommended for use in small children, but in experienced hands it has been demonstrated to be safe.

IJ Approach

Cervical trauma with swelling or anatomic distortion at the intended site of IJ venipuncture is the most important contraindication to the IJ approach. Likewise, the presence of a cervical collar is problematic. Although bleeding disorders are relative contraindications to central venous cannulation, the ultrasound-guided IJ approach is preferred over the SV route because the IJ site is compressible. However, prolonged compression of the artery to control bleeding could impair the cerebral circulation if collateral blood flow is compromised. In a study by Oguzkurt and colleagues, only minor bleeding complications occurred in less than 2% of patients after ultrasound-guided IJ vein catheterization. In the setting of severe bleeding diatheses, the ultrasound-guided femoral approach is an acceptable alternative. Ultrasound-guided IJ vein catheter placement is preferred in patients with abnormal anatomy from previous IJ vein trauma, small IJ vessels, and morbid obesity. Historically, carotid artery disease (obstruction or atherosclerotic plaque) is a relative contraindication to IJ vein cannulation because inadvertent puncture or manipulation of the artery could dislodge a plaque. If a preceding SV catheterization has been unsuccessful, the ipsilateral IJ route is generally preferred for a subsequent attempt. In this manner, bilateral iatrogenic complications can be avoided.

Femoral Approach

Contraindications to femoral cannulation include known or suspected intraabdominal hemorrhage or injury to the pelvis, groin, iliac vessels, or IVC. Additionally, avoid the femoral approach when known or suspected deep venous thrombosis is present. Palpation for femoral pulsations during CPR is difficult and the pulsations are often venous rather than arterial. Ultrasound-guided catheterization of the femoral vein during CPR is more successful, and it is less likely to incur inadvertent arterial puncture than the standard landmark-oriented approach.

PROCEDURE

The most commonly used method for central venous cannulation is the Seldinger (guidewire) technique, in which a thin-walled needle is used to introduce a guidewire into the vessel lumen. Seldinger originally described this technique in 1953 as a method of placing a catheter for percutaneous arteriography. The Seldinger technique is illustrated in Fig. 22.9. To obtain vascular access, insert a small needle into the intended vessel. Once the introducer needle is positioned within the lumen of the vessel, thread a wire through the needle and then remove the needle. The wire, now within the vessel, serves as a guide over which the catheter is inserted. Although the Seldinger technique involves several steps, it can be performed quickly once mastered. More importantly, this technique broadens the application of central venous cannulation by permitting the insertion of standard infusion catheters, multilumen catheters, large-bore rapid infusion systems, introducer devices, hemodialysis devices, and even peripheral cardiopulmonary bypass cannulas. Given this flexibility, the use of Seldinger-type systems is advantageous despite their greater cost.

Ultrasound guidance has revolutionized the cannulation of central veins. As with all anatomic structures in the human body, veins are highly variable in their location. Not surprisingly, research has demonstrated that the ability to see the internal structure’s location and proximity to other structures greatly increases the safety and success rate while decreasing the time required to perform the procedure. These advantages have been recognized by national organizations. In a report from the Agency for Healthcare Research and Quality (AHRQ), use of ultrasound guidance was listed as one of the top 10 ways to reduce morbidity and mortality. Many hospitals now require the use of ultrasound guidance for the placement of all CVCs.
The basic materials required for central venous cannulation are shown in Review Box 22.1 and are discussed here in further detail. The catheter may be a component in a guidewire system or may be of the over-the-needle variety (the other widely used method for catheter placement). Several types of CVC Seldinger-type prepackaged kits are commercially available and the variations in each kit are discussed in the next section.

**EQUIPMENT**

Preparation and organization of equipment ahead of time are imperative for a successful procedure. Most catheters now come from the manufacturer in convenient sterile kits. We strongly recommend stocking all additional equipment such as sterile gowns, gloves, and drapes in a dedicated “central line cart.” This is a fundamental part of the “bundling” practice that has been shown to reduce the search for supplies, improve compliance with full-barrier technique, and subsequently decrease catheter-related infections. Sterile barrier precautions with cap, face mask, sterile gown, and gloves should be used at all times during insertion of CVCs.

**Ultrasound**

Historically, many clinicians preferred to first locate the position of a central vein with a small exploratory or “finder” needle rather than directly cannulating the vein with a larger needle to accommodate a guidewire or catheter. This practice is less practical for the SV approach and has largely been replaced with the use of bedside ultrasound. Ultrasound-guided CVC placement allows the provider to survey the anatomy before the procedure, guide insertion of the needle into the correct vessel, and confirm placement of the catheter in the vessel (see Ultrasound Box 22.1).

**Needle**

Virtually any needle or catheter can be used to introduce a guidewire into a vessel, but there are advantages to using needles specifically designed for passage of a guidewire. These needles must be large enough to accommodate the desired wire, yet be as small as possible to minimize bleeding complications. The introducer needles provided with CVCs or introducer devices are usually thin walled to maximize lumen size relative to the overall needle diameter. If a needle that is not thin walled is used, choose a size that is 1 gauge smaller (larger bore) than that listed in Table 22.2. If unsure, simply test the equipment first to ensure compatibility.

Standard needles may have a uniformly straight-bore lumen throughout their length. A wire passing into a straight needle may encounter an obstacle at the proximal end. The proximal end of a Seldinger needle incorporates a funnel-shaped taper that guides the wire directly into the needle (Fig. 22.4). It is advisable to use a non–Luer-Lok or slip-tip type of syringe because the added twisting that is required to remove a Luer-Lok syringe from the introducer needle may dislodge a tenaciously placed needle. Safety syringe systems exist that permit passage of the wire without removal of the aspirating syringe by using a central tunnel in the barrel. This device incorporates a hollow syringe through which the guidewire can pass directly into the introducing needle without detachment. It also reduces the risk for air embolism, which can occur when the needle is open to air. It is not uncommon for the wire to become snagged at the junction of the safety syringe and the needle hub. In this case, simply remove the syringe and insert the wire directly.

**Guidewire**

Two basic types of guidewires are used: straight and J-shaped. Straight wires are for use in vessels with a linear configuration, whereas J-wires are for use in tortuous vessels. Both wires have essentially the same internal design (Fig. 22.5). The flexibility of the wire is the result of a stainless steel coil or helix that forms the bulk of the guidewire. Within the central lumen of the helix is a straight central core wire, called a mandrel, that adds rigidity to the steel coil. The mandrel is usually fixed at one end of the helix and terminates 0.5 and 3.0 cm from the other end to create a flexible or floppy tip. Wires are also available with two flexible ends, one straight and the other J-shaped. The flexible end of the guidewire allows the wire to flex on contact with the wall of a vessel. If the contact is tangential, as with an infracavicular approach to the SV, a straight wire is generally preferred. If the angle is more acute, as with an external jugular approach to the SV, a J-wire may be used. The more rounded leading edge of the J-wire provides a broader surface to manipulate within the vessel and decreases the risk for perforation. This is especially advantageous when attempting to thread a wire through a vessel with valves. Many guidewires also contain a straight safety wire that runs parallel to the mandrel to keep the wire from kinking or shearing.

| TABLE 22.2 Needle Sizes for Venous and Arterial Catheters<sup>a</sup> |
|-----------------------------|-----------------------------|
| **STANDARD FULL-LENGTH COIL** | **GUIDEWIRE CATHETER SIZE (Fr)** | **NEEDLE GAUGE<sup>b</sup>** |
| 3 | 21 |
| 4–4.5 | 20 |
| 5–6.0 | 20–19 |
| 6–8.5 | 19–18 |

<sup>a</sup>Any size of catheter from 3.0 to 8.5 Fr may be introduced with a 22-gauge needle if a solid wire (e.g., Cor-Flex, Cook Critical Care) is used.

<sup>b</sup>All needle gauges are for thin-walled needles only, the type supplied in central line kits.

**Figure 22.4** Introducing needles. A, Ordinary needle with a straight-bore lumen. B, Seldinger needle with a tapered lumen, which allows easy entry of the guidewire.
Catheters
A number of different catheter and introducer devices have been developed, and the method of passage into the vessel varies accordingly. The functions of catheters have become more sophisticated as well, most notably for continuous monitoring of SCVO₂ and cardiac output. Generally, one can place single-, double-, and triple-lumen catheters by sliding the catheter directly over a guidewire into the intended vessel (Fig. 22.4). Catheter insertion lengths are listed in Table 22.3. Larger catheters or devices without lumens can be introduced with a sheath-introducer system. Over-the-needle catheters can be introduced once intravascular placement is attained.

The Desilets-Hoffman–type sheath introducer became available in 1965 to aid in arteriography procedures that require many catheter changes. This device is commonly but incorrectly termed a Cordis, which is a proprietary trade name. The sheath-introducer unit includes two parts, an inner dilator and an outer sheath, as shown in Fig. 22.7B. The dilator is rigid with a narrow lumen to accommodate the guidewire. It is longer and thinner than its sheath and has a tapered end that dilates the subcutaneous tissue and vessel defect created by the needle. The sheath (or introducer catheter when used as a cannula for inserting Swan-Ganz catheters, transvenous pacemakers, or other devices) has a blunt end and is simply a large-diameter catheter.

Many modifications of the sheath exist, such as side arms and diaphragms to aid in the placement of devices without lumens. Care must be taken when using side-arm sets for rapid administration of fluid because some catheters are 8.5 Fr in diameter but have only a 5-Fr side arm. Some sets have a “single-lumen infusion catheter,” which performs the same function but is more easily secured to the sheath introducer. Selection of the appropriate diameter of introducer catheter should correspond to the indication for placement of a CVC. Generally, an 8.5-Fr catheter is used to facilitate placement
of a Swan-Ganz catheter and a 6.0-Fr catheter is used to facilitate transvenous placement of a pacemaker. If the introducer catheter is larger than required to support the intraluminal device, a leak may develop at the diaphragm insertion point.

Special catheters have been developed to prevent bacterial contamination and line sepsis.40,70,71 These catheters are impregnated with either antiseptics (silver sulfadiazine and chlorhexidine) or antibiotics (minocycline, rifampin, or cefazolin) to reduce bacterial colonization and microbial growth. Heparin-coated catheters are also available that prevent fibronectin binding, thereby inhibiting the formation of bacterial biofilm on the catheter’s surface. These catheters can decrease catheter-associated infection (CAI) significantly and are cost-effective when the prevalence of CAI is greater than 2%.81 They should be avoided in patients with a history of heparin-induced thrombocytopenia.72 Minocycline- and rifampin-impregnated catheters are currently considered to be the most effective.80,70 Other interventions that decrease central line infections include the use of full sterile barrier precautions,64,68 skin preparation with chlorhexidine solution,64,66,75 and placement by experienced physicians.64,74,76

Many different catheters are currently manufactured. Although this leads to great flexibility in choice and cost, it often results in confusion when a clinician is handed an unfamiliar catheter during an emergency. It is best to use one brand routinely and to ensure that all medical personnel are thoroughly familiar with its use.77

**TABLE 22.3** Formulas for Catheter Insertion Length Based on Patient Height and Approach

<table>
<thead>
<tr>
<th>SITE</th>
<th>FORMULA</th>
<th>IN SVC (%)</th>
<th>IN RA (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RSC</td>
<td>(Ht/10) – 2 cm</td>
<td>96</td>
<td>4</td>
</tr>
<tr>
<td>LSC</td>
<td>(Ht/10) + 2 cm</td>
<td>97</td>
<td>2</td>
</tr>
<tr>
<td>RIJ</td>
<td>Ht/10</td>
<td>90</td>
<td>10</td>
</tr>
<tr>
<td>LIJ</td>
<td>(Ht/10) + 4 cm</td>
<td>94</td>
<td>5</td>
</tr>
</tbody>
</table>

Ht, Patient height (in cm); LIJ, left internal jugular; LSC, left subclavian; RSC, right subclavian; RA, right atrium; RIJ, right internal jugular; SVC, superior vena cava.


**TECHNIQUE**

**Preprocedure Preparation**

When possible, discuss the procedure with the patient and obtain written informed consent. Place the patient and yourself in an appropriate position for the specific vessel being accessed. If available, perform an ultrasound survey to identify the patient’s anatomy, ensure vessel patency, and confirm the puncture site (Fig. 22.8). Ultrasound-guided CVC placement has been shown to decrease procedure times, as well as complication rates.49–52 Additionally, compliance with a central line bundling policy has been shown to significantly decrease central line–associated bloodstream infections (CLABSIs).61–66 Prepare and drape the puncture site while maintaining sterile technique, and observe universal precautions throughout the procedure (Fig. 22.9, steps 1 and 2). A gown, surgical cap, mask, eye protection, and sterile gloves should be worn throughout the procedure when possible. When performing ultrasound-guided placement of a CVC, ensure that a sterile transducer sheath and sterile transducer gel are used during the procedure (see Fig. 22.9, step 3). Using an assistant will prove valuable in patient preparation, maintenance of sterility, and handling of the equipment.

**Guidewire Placement With the Seldinger Technique**

When performing ultrasound-guided placement of a CVC, begin with an ultrasound survey of the target vein, surrounding structures, and venipuncture location, as shown in Fig. 22.8. Veins can easily be distinguished from the nearby artery by applying external pressure with the transducer. Veins collapse completely with pressure, whereas arteries may deform but do not usually collapse. Occasionally, the vein does not collapse with pressure. If this occurs, a thrombus may be present in the vein or the structure has been misidentified. If a suspected vein does not collapse with pressure, it is not an appropriate vessel for cannulation. If available, Doppler functions may also be helpful in the differentiation of veins and arteries. Select a venipuncture location where branching of the vein will allow...
nonpulsatile blood does not rule out arterial cannulation. If there are concerns about possible arterial puncture, either remove the introducer needle and draw a sample for blood gas analysis from the needle to compare with an arterial blood gas sample, or insert an 18-gauge single-lumen catheter over the wire and into the vessel because this step does not require the use of a dilator. The catheter can then be connected to a pressure transducer to confirm the presence of venous waveforms and venous pressure.

Introduce the flexible end of the guidewire into the hub of the needle (see Fig. 22.9, step 7). It may be easier to introduce the J-wire by advancing the plastic sleeve contained in the kit onto the floppy end of the wire to straighten the J-shape. This straightened end is then introduced into the needle hub. The guidewire should thread smoothly through the needle into the vessel without resistance. Do not force the wire if resistance is encountered, but remove it from the needle and reattach the syringe to aspirate blood and reconfirm intravascular needle placement. It is important for the wire to slip easily from the needle during removal. If resistance to removal of the wire is felt, the wire and needle should be removed as a single unit to prevent shearing of the wire and resultant wire embolism. It has been recommended by some that no wire should ever be withdrawn through the introducing needle. Although there are no clinical data to support this recommendation and newer wires are stronger and more resistant to shearing, it represents the safest course of action. The recommendation to remove the needle and wire as a unit is sometimes disregarded because of reluctance to abandon a potentially successful venipuncture. The clinician performing the procedure must use both caution and good judgment to determine the best course of action but the shortest path of the needle, will not obstruct passage of the catheter, and will not allow inadvertent puncture of other vital anatomic structures. (See Chapter 66 for additional information and descriptions of the ultrasound technique.)

Prepare the catheter for insertion by flushing each lumen with sterile normal saline. Anesthetize the insertion site with lidocaine or bupivacaine (see Fig. 22.9, step 5). Attach a small syringe to an introducing needle that is large enough to accommodate the guidewire. Insert the needle and syringe together. Slowly advance the needle into the vein and apply steady negative pressure on the syringe (see Fig. 22.9, step 6). When performing ultrasound-guided CVC placement, follow the needle trajectory in the soft tissue and observe penetration of the vessel. If the tip of the needle is not visualized at all times with ultrasound, the needle may be passed into structures other than the vein. The key concept in using ultrasound guidance for venous access is to visualize the tip of the needle at all times during cannulation (Fig. 22.10). Once the tip of the needle enters the vessel lumen, blood will be aspirated freely. Stabilize the needle hub to prevent movement of the needle and displacement of the tip from the vessel, and remove the syringe. This action can dislodge the needle tip and is the activity most associated with failure to pass a wire after the vein has initially been entered. The need to disconnect the syringe can be eliminated by use of the Arrow Safety Syringe (Teleflex, Morrisville, NC). After removing the syringe, cap the needle hub with your thumb before passing the guidewire to minimize the potential for air embolism.

Confirm that the blood flow is nonpulsatile. Bright red pulsatile blood is very suggestive of arterial puncture. Be aware that in shock states or marked arterial desaturation, dark, nonpulsatile blood does not rule out arterial cannulation. If there are concerns about possible arterial puncture, either remove the introducer needle and draw a sample for blood gas analysis from the needle to compare with an arterial blood gas sample, or insert an 18-gauge single-lumen catheter over the wire and into the vessel because this step does not require the use of a dilator. The catheter can then be connected to a pressure transducer to confirm the presence of venous waveforms and venous pressure.

Introduce the flexible end of the guidewire into the hub of the needle (see Fig. 22.9, step 7). It may be easier to introduce the J-wire by advancing the plastic sleeve contained in the kit onto the floppy end of the wire to straighten the J-shape. This straightened end is then introduced into the needle hub. The guidewire should thread smoothly through the needle into the vessel without resistance. Do not force the wire if resistance is encountered, but remove it from the needle and reattach the syringe to aspirate blood and reconfirm intravascular needle placement. It is important for the wire to slip easily from the needle during removal. If resistance to removal of the wire is felt, the wire and needle should be removed as a single unit to prevent shearing of the wire and resultant wire embolism. It has been recommended by some that no wire should ever be withdrawn through the introducing needle. Although there are no clinical data to support this recommendation and newer wires are stronger and more resistant to shearing, it represents the safest course of action. The recommendation to remove the needle and wire as a unit is sometimes disregarded because of reluctance to abandon a potentially successful venipuncture. The clinician performing the procedure must use both caution and good judgment to determine the best course of action but...
Central Venous Catheterization (Internal Jugular Approach)

1. Prepare the area with chlorhexidine solution. A gown, surgical cap, mask, eye protection, and sterile gloves should be worn throughout the procedure.

2. Apply a full-body, sterile drape. Meticulous attention must be paid to sterile technique to avoid iatrogenic infection.

3. Insert the ultrasound probe into a sterile sheath and use sterile ultrasound gel during the procedure. Enlist the help of an assistant in patient preparation and maintenance of sterility.

4. Identify the anatomic structures with ultrasound. The internal jugular vein (IJ) and carotid artery (CA) must be clearly distinguished from each other (see text for more details).

5. Anesthetize the tissues overlying the vein with local anesthetic. Here, the operator is using ultrasound guidance to ensure a proper entry site.

6. Insert the needle and syringe while slowly advancing and applying negative pressure to the plunger. Follow the needle trajectory with ultrasound until the vein is entered and blood enters the syringe (arrow).

7. Remove the syringe and advance the guidewire through the needle. Use the straightener (arrow) to facilitate entry of the J-wire into the hub. NEVER FORCE THE WIRE!

8. Once the wire has been inserted to the appropriate depth (see text for details), remove the needle (arrow). It is essential to always maintain a grip on the wire throughout the procedure (!).

Figure 22.9 Ultrasound-guided internal jugular central venous catheterization.

Continued
Central Venous Catheterization (Internal Jugular Approach)

9. Make an incision at the site of the wire to facilitate dilator and catheter passage. Make the incision the width of the catheter and extend it completely through the dermis.

10. Thread the dilator over the guidewire. The wire must always be protruding from the end of the dilator and firmly in your grasp (!). Advance the dilator several centimeters into the vessel and then remove.

11. Advance the catheter over the wire. It can be difficult to align the two pieces; hold the very end of the catheter and the wire to make this step easier.

12. Advance the catheter into the vessel. The guidewire will emerge from the distal port. It is essential that the guidewire protrudes from the hub and is grasped before catheter advancement (!).

13. Remove the wire. Cover the open port with your thumb (arrow) until the end cap is screwed on.

14. Flush all ports with saline.

15. Suture the catheter into place using nonabsorbable silk sutures. Several knots should be made to secure the line. Avoid making knots that place excessive pressure on the skin.

16. Clean the area around the catheter insertion site with chlorhexidine. Place a simple dressing, avoiding excessive amounts of gauze and tape.

Figure 22.9, cont’d
should not withdraw the guidewire against resistance. Manipulation
of the wire within an introducer needle should be done only with standard coil guidewires. Solid wires (such as Cor-Flex Wire Guides from Cook Critical Care) have a small lip at the point at which the flexible coil is soldered to the wire. This lip can become caught on the edge of the tip of the needle and shear off the coil portion of the wire. Solid wires must thread freely on the first attempt or the entire wire and needle assembly must be removed. Keep backup wires on hand.

Occasionally, a wire must be teased into the vessel; rotating the wire or needle often helps in difficult placements. If the wire does not thread easily, pull back slightly on the needle itself just before advancing the wire. This helps if the opening of the needle is abutting the vessel’s inner wall and blocking entry of the wire or if the vein is compressed by introduction of the needle. Changing wire tips from a straight wire to a J-wire or vice versa may also solve an advancement problem. If the inner lumen of a vessel is smaller than the diameter of the J, the wire will be prevented from returning to its natural shape and the spring in the coil will generate resistance. Any advantages of a J-wire will be negated if the wire fails to regain its intended shape. In this instance, it should be possible to introduce a straight tip without a problem. Alternatively, if the angle of entry of the needle into the vessel is more acute than suspected, a straight wire may not be able to bend appropriately as it encounters the vessel’s far wall. A J-tipped wire may be used and threaded in such a manner that the wire resumes its J-shape away from the far wall. All these maneuvers are performed with gentle free motions of the wire within the needle. If at any time the wire cannot be advanced freely, suspect improper placement and reevaluate the attempt.

If threading easily, advance the guidewire until at least one quarter of the wire is within the vessel. The further into the vessel the wire extends, the more stable its location when the catheter is introduced. However, advancing the guidewire too far may result in ventricular ectopy secondary to endocardial irritation, myocardial puncture leading to tamponade, or entanglement in a previously placed pacemaker, internal defibrillator, or IVC filter. In both the left and right IJ vein and infraclavicular SV approaches, fluoroscopic study during passage of the guidewire has determined the mean distance from skin to the SVC-atrial junction to be 18 cm. This distance has been recommended as the greatest depth of guidewire insertion for these approaches. It should be noted that 18 cm is not necessarily the appropriate final depth for the catheter being placed (see following discussion).

Cardiac monitoring may be helpful during the insertion of central lines. Any increase in premature ventricular contractions or a new ventricular dysrythmia should be interpreted as evidence that the guidewire is inserted too far and should be remedied by withdrawing the wire until the rhythm reverts to baseline. Usually, the procedure can be continued after a moment, with care taken to not readvance the wire. Persistent ventricular dysrythmias require standard advanced cardiac life support treatment and consideration of a new vascular approach.

Occasionally, a wire threads easily past the tip of the needle and then suddenly will not advance farther. If the introducer needle demonstrated free return of blood at the time of wire entry and the initial advancement of the wire met no resistance, the two options are to halt the procedure or seek confirmation of the wire position. The guidewire within the lumen of the vessel can be visualized and confirmed via cross-sectional and longitudinal views on ultrasound. Alternatively, the needle may be removed, the wire fixed in place with a sterile hemostat, and a radiograph taken to confirm the position of the wire.78,79 A freely advancing wire may suddenly stop once it is well within a vessel if the vessel makes an unsuspected bend or is being compressed or deviated by another structure, such as a rib or muscle. This seems especially common with the infraclavicular approach to the SV and can sometimes be remedied by a more lateral approach.

**Sheath Unit and Catheter Placement**

Once the wire is placed into the vessel, remove the needle in preparation for passage of the catheter (see Fig. 22.9, step 8). Proper positioning of the guidewire within the vessel lumen can be confirmed by cross-sectional and longitudinal ultrasound imaging (Fig. 22.11). This can be done at any point while inserting the wire to ensure that the correct vessel has been cannulated and that puncture of the posterior wall has not occurred. This technique can be quite useful when resistance is encountered while feeding the guidewire. A small skin incision is required at the site of the wire to widen the opening (see Fig. 22.9, step 9). Make the incision approximately the width of the catheter to be introduced and extend it completely through the dermis.
When placing soft multiple-lumen catheters, the tissue must be dilated from the skin to the vessel before placement of the catheter. Thread the guidewire through the distal opening of the rigid dilator until it extends through the proximal end of the dilator (see Fig. 22.9, step 10). The wire must always be visibly protruding from the end of the dilator or catheter during insertion to avoid inadvertent advancement of the wire into the circulation and potential loss of the wire. While maintaining control of the guidewire proximally, thread the dilator over the wire into the skin with a twisting motion. Advance the rigid dilator only a few centimeters into the vessel and then remove it. Once the dilator is removed, thread the soft catheter into position over the wire. Placement of multiple-lumen catheters requires identification of the distal lumen and its corresponding hub. Find the distal lumen at the very tip of the catheter. The corresponding hub is usually labeled “distal” by the manufacturer. If there is any confusion, inject a small amount of sterile saline through each hub until it is observed exiting the distal lumen. Once the distal hub is identified, remove its cover cap to allow passage of the guidewire. Place the catheter by threading the guidewire into the distal lumen and advancing it until it protrudes from the identified hub (see Fig. 22.9, step 11). It is imperative that the guidewire protrude from the catheter hub and that it be firmly grasped as the wire and catheter are advanced. If the wire does not protrude from the proximal end of the catheter, withdraw the wire at the skin entry point until it protrudes enough to be grasped. While maintaining control of the guidewire proximally, advance the assembly through the skin with a twisting motion until it is within the vessel. Grasp the unit at the junction of the sheath and dilator. This prevents the thinner sheath from kinking or bending at the tip or from bunching up at the coupler end. Keep the assembly intact and advance it through the skin to the hub. Once the catheter is placed, remove the wire and dilator from the sheath simultaneously (see Fig. 22.12, step 4). When removing the wire and dilator, the dilator must “unsnap” from the sheath unit and the wire must slip out easily. Once the single-lumen sheath-introducer catheter is placed correctly, it may be used to facilitate the placement of additional intraluminal devices such as a pulmonary artery catheter, transvenous cardiac pacemaker, or an additional multiple-lumen catheter. At times, critically ill patients who require initial large-volume resuscitation will later require multiple medications and therapies that dictate the need for a multiple-lumen catheter. An alternative method of placing a multiple-lumen catheter is to thread the catheter through a standard Desilets-Hoffman sheath-introducer system.

It is important to consider the depth of insertion of the catheter (see Table 22.3). The proper depth of catheter insertion is site specific (see later). After successful CVC placement, the catheter should be anchored to the skin with sutures (see Fig. 22.9, step 15). Each port should be immediately capped and flushed with a saline solution (see Fig. 22.9, step 14). The catheter insertion site should be dressed appropriately and all sharp implements disposed of in appropriate receptacles (see Fig. 22.9, step 16).

**Replacement of Existing Catheters**

In addition to placing new catheters, clinicians may use the guidewire technique to change existing catheters. Many patients with CVCs are seriously ill and will require subsequent monitoring of pulmonary artery wedge pressure, placement of a transvenous pacemaker, or insertion of a different catheter. The CVC that is initially inserted should have a lumen large enough to accept a guidewire and facilitate conversion to a different catheter. Clinicians may use the guidewire technique to change a single-lumen CVC to a triple-lumen catheter or a sheath-introducer set. Not all commercially available CVCs will accept a guidewire.

Replacement of an existing catheter begins with selecting a guidewire longer than either of the devices to be exchanged. Use meticulous aseptic technique. Insert the guidewire into the existing CVC until a few centimeters of wire protrudes from the proximal end. With one hand holding the wire securely, remove the catheter and wire as a single unit until the tip of the catheter just clears the patient's skin. Grasp the wire at
Central Venous Catheterization and Central Venous Pressure Monitoring

These catheters are not suitable for high-volume fluid resuscitation and they are too small for passage of a pacemaker lead. Once the clinical situation stabilizes, exchange this device for a larger central catheter via the Seldinger technique. Prepare the skin with chlorhexidine solution. Use a longer peripheral-type catheter (such as a 16-gauge, 5 1/4-inch angiocatheter) in an adult. Smaller-diameter devices, such as 20-gauge catheters, may be easier to pass but provide slower infusion rates. Attach the needle to a syringe and slowly advance it into the vein with steady negative pressure applied to the syringe. This may be difficult because of the longer length of the needle relative to the catheter. When using bedside ultrasound, follow the trajectory of the needle into the soft tissues and visualize penetration of the vessel. With over-the-needle catheters, the needle extends a few millimeters past the tip of the catheter. Return of blood will be obtained when the tip of the needle is in the vein, although the catheter may actually be outside the lumen. If the needle is withdrawn before the catheter is advanced, the tip of the catheter will remain outside the vein. It is therefore important to advance the needle a few millimeters after the venous flash is seen and then hold it steadily while advancing the catheter into the vein. Secure the catheter and verify its placement as detailed later in this chapter.

Insertion of the Sheath Introducer

The sheath introducer and dilator must be assembled prior to insertion. Some sheaths have a one-way valve that must be opened (by rotating the valve) before insertion of the dilator.

Grasp the guidewire as it protrudes from the sheath-dilator assembly. Advance the dilator and sheath as a unit over the wire. It is essential to grasp the guidewire as it protrudes from the dilator prior to advancing the catheter.

Open the one-way valve (if so equipped), and fully insert the dilator into the sheath.

After full insertion of the sheath, remove the dilator and guidewire simultaneously, and close the one-way valve (if so equipped).

Figure 22.12 Insertion of the sheath introducer. Insertion of a sheath introducer varies slightly from that for a triple-lumen catheter—the dilator and the catheter are inserted simultaneously as depicted. The remainder of the steps are analogous to those in Fig. 22.9. Once inserted, sheath introducers facilitate the placement of devices such as pulmonary artery catheters and transvenous pacemakers.

Over-the-Needle Technique

An optional method for cannulation is to place an over-the-needle catheter percutaneously. Over-the-needle devices (such as the Angiocath [Becton Dickinson, Franklin Lakes, NJ]) use a tapered plastic catheter that passes through the vessel wall into the lumen, with the tip of the needle being used as a guide. There are advantages with this system. The catheter does not pass through a sharp needle and there is less risk of shearing and resultant catheter embolization. Additionally, the hole made by the needle in the vessel wall is smaller than the catheter, thus producing a tighter seal. The IJ vein and SV via the supraclavicular approach are the most popular and appropriate approaches for this technique. These devices may be useful when rapid central venous access is required (e.g., in cardiac arrest). These catheters are not suitable for high-volume fluid resuscitation and they are too small for passage of a pacemaker lead. Once the clinical situation stabilizes, exchange this device for a larger central catheter via the Seldinger technique.

Prepare the skin with chlorhexidine solution. Use a longer peripheral-type catheter (such as a 16-gauge, 5 1/4-inch angiocatheter) in an adult. Smaller-diameter devices, such as 20-gauge catheters, may be easier to pass but provide slower infusion rates. Attach the needle to a syringe and slowly advance it into the vein with steady negative pressure applied to the syringe. This may be difficult because of the longer length of the needle relative to the catheter. When using bedside ultrasound, follow the trajectory of the needle into the soft tissues and visualize penetration of the vessel. With over-the-needle catheters, the needle extends a few millimeters past the tip of the catheter. Return of blood will be obtained when the tip of the needle is in the vein, although the catheter may actually be outside the lumen. If the needle is withdrawn before the catheter is advanced, the tip of the catheter will remain outside the vein. It is therefore important to advance the needle a few millimeters after the venous flash is seen and then hold it steadily while advancing the catheter into the vein. Secure the catheter and verify its placement as detailed later in this chapter.
SITE SELECTION

Subclavian Approaches

Subclavian venipuncture is the most frequently used means of achieving central venous access. The infraclavicular SV approach was the first popular means of central venous access and has been used widely for nearly half a century. It is useful in many clinical situations and relatively easy to learn. It is often the best approach in trauma because a cervical collar can interfere with the IJ technique. The supraclavicular SV approach may be preferable during CPR because it minimizes physical interference in chest compressions and airway management. In addition, the supraclavicular SV technique has been performed in the sitting position in patients with severe orthopnea. The subclavian route is associated with the lowest incidence of catheter-related infections and deep vein thrombosis, but is associated with the highest risk for pneumothorax. Finally, the left SV provides a more direct route to the SVC and is the preferred site for pacemaker placement and CVP monitoring.

IJ Approach

The IJ vein provides an excellent site for placement of a CVC. However, there is a 5% to 10% risk for complications, with serious complications occurring in approximately 1% of patients. Failure rate have been found to be 19.4% for landmark-placed IJ catheterization by a junior practitioner and 5% to 10% by a clinician with extensive experience. Despite its potential complications, the IJ vein is in many cases preferred over other options for central venous access. In contrast to the SV, arterial punctures are easier to control because direct pressure can be used, and the incidence of pneumothorax is lower. Hematoma formation is easier to diagnose because of the close proximity of the IJ vein to the skin. In addition, the right IJ vein provides a straight anatomic path to the SVC and right atrium. This is advantageous for passage of catheters or internal pacemaker wires to the heart. Disadvantages of IJ vein cannulation over other sites include a relatively high carotid artery puncture rate and poor landmarks in obese or edematous patients.

The IJ technique is useful for routine central venous access and for emergency venous access during CPR because the site is removed from the area of chest compressions. The differences in morbidity between the SV and the IJ vein approaches have probably been overstated. Catheter malposition is more frequent in the SV, but the risk for infection is probably slightly higher with IJ sites. The rate of arterial puncture is higher with IJ vein attempts, but the SV is not a compressible site. Though counterintuitive, the evidence available does not support a significant difference in the rate of pneumothorax and hemotorax. Although there may be a slight difference in complications between the two routes, in the absence of specific contraindications clinicians should use the technique with which they are most familiar. The rapid development of real-time ultrasound guidance may tip the scales toward the IJ vein as the preferred site.

Femoral Approach

Cannulation of the femoral vein for central venous access has become increasingly popular, especially for venous access, infusion ports, passage of transvenous pacemakers, and placement of pressure measurement catheters in critically ill patients. The relatively simple and superficial anatomy surrounding the femoral vein affords a rapid approach to the central venous system and avoids many of the more significant complications associated with cannulation of the IJ vein and the SV. These benefits are tempered somewhat by several long-term disadvantages, including higher infection rates and an increased risk for venous thrombosis. Other indications for ED femoral cannulation include emergency cardiopulmonary bypass for resuscitation purposes, charcoal hemoperfusion for severe drug overdoses, and dialysis access. The femoral area is less congested with monitoring and airway equipment than the head and neck area, and conscious patients who are still bedridden may turn their head and use their arms more freely without moving the central line. The femoral site is contraindicated in ambulatory patients who require central access.

SPECIFIC VESSEL ACCESS TECHNIQUES

If SV or IJ vein approaches are planned, prepare the skin in the area, including puncture sites for both the infraclavicular and supraclavicular SV and IJ vein approaches. This permits the clinician to change the site after an unsuccessful attempt without repeating the preparation or having to obtain an interval chest radiograph. Prepare the area, including the ipsilateral anterior aspect of the neck, the supraclavicular fossa, and the anterior chest wall 3 to 5 cm past the midline and the same distance above the nipple line. Prepare for femoral access by trimming groin hairs and applying chlorhexidine to cover an area the breadth of and extending 10 cm above and below the inguinal ligament. Each approach to central venous cannulation is described separately in the following sections. As with any invasive procedure, briefly describe the procedure to awake patients, and restate each step as it is about to be performed. After descriptions of the common approaches to the central veins, puncture site care, verification of placement, and other adjuncts to the procedure are summarized.

Infraclavicular Subclavian Approach

Descriptions of subclavian venipuncture often focus unduly on angles and landmarks. Indeed, recent studies have demonstrated that some traditional positioning maneuvers may actually hinder successful cannulation efforts.

Positioning

Place the patient supine on the stretcher with the head in a neutral position and the arm adducted at the side. Some authors have advocated various shoulder-, back-, head-, and arm-positioning maneuvers, but they take extra time and the help of an assistant and are often not helpful. We believe that the best position for almost all infraclavicular SV attempts is the neutral shoulder position with the arm adducted. Turning the head away may be helpful but is certainly not required if cervical injuries are suspected. Interestingly, Jung and colleagues found that in children, tilting the head toward the catheterization site improved catheter malposition rates. This technique has not been studied in adults.
In difficult cases, placing a small towel roll under the ipsilateral shoulder or having an assistant place caudal traction of approximately 5 cm on the extremity may also be helpful. Placing the patient in a moderate Trendelenburg position (10 to 20 degrees) decreases the risk for air embolism. The claim that this position distends the vein is controversial, but it may do so to a small favorable degree. If the Trendelenburg position is impractical, the SV approach is probably less affected than the IJ vein approach when resorting to a neutral or even an upright position.

Placing a pillow under the back is commonly recommended to make the clavicle more prominent, but as the shoulder falls backward, the space between the clavicle and the first rib narrows, thus making the SV less accessible. Significant compression of the subclavian vessels between these bony structures occurs as the shoulders retract, which can cause a "pinch off" of the catheter as it slides through the SV between the clavicle and the first rib.

**Venipuncture Site**

The right SV is usually selected first because of the lower pleural dome on the right and the need to avoid the left-sided thoracic duct. The more direct route between the left SV and the SVC is a theoretical advantage of left-sided subclavian venipuncture; however, there is no higher incidence of catheter malposition when the right infracavicular SV approach is used. In conscious patients, anesthetize the point of needle entry with 1% lidocaine. If possible, infiltrate the periosteum of the clavicle, found by touching the bone with the needle, to make the procedure less painful. Opinions vary regarding the best point of needle entry, more so than for the IJ or femoral approaches. With nonobese patients, look for the deltopectoral triangle, which is bounded by the clavicle superiorly, the pectoralis major medially, and the deltoid muscle laterally. The junction of the middle and medial thirds of the clavicle lies just medial to this triangle. Further medially, the vein lies just posterior to the clavicle and above the first rib, which acts as a barrier to penetration of the pleura. This protective effect is theoretically diminished when a more lateral location is chosen. However, when approaching the vein more medially, some clinicians have difficulty puncturing the SV, dilating the tissues, and passing the J-wire. Other recommended sites of approach include lateral and inferior to the junction of the clavicle and the first rib, with the needle aimed at this junction, and entry at the site of a small tubercle in the medial aspect of the deltopectoral groove. We recommend puncturing the skin at the lateral portion of the deltopectoral triangle via a shallow angle of attack.

**Needle Orientation**

Orient the bevel of the needle inferomedially to direct the wire toward the innominate vein rather than toward the opposite vessel wall or up into the IJ vein. Align the bevel of the needle with the markings on the barrel of the syringe to permit awareness of bevel orientation after skin puncture.

Before inserting the needle, place your left index finger in the suprasternal notch and your thumb at the costoclavicular junction (Fig. 22.13). These landmarks serve as reference points for the direction that the needle should travel. Orient the bevel of the needle inferomedially and aim the needle above and posterior to your index finger. Note: to avoid puncturing the lung or the subclavian artery, once the needle tip is under the clavicle keep the needle parallel to the clavicle, NOT directed posteriorly.

Return of pulsatile flow signifies arterial puncture and the needle should be withdrawn immediately. A single arterial puncture without laceration rarely causes serious harm. Using this technique eliminates the need to measure angles, to "walk" the clavicle or to concentrate excessively on maintaining the needle parallel to the chest wall. Avoid using sweeping motions of the tip of the needle to prevent unseen injuries.

**Unsuccessful Attempts**

Cannulation of the SV may not succeed on the first attempt. It is reasonable to try again, but after three or four unsuccessful attempts it is wise to move to a different anatomic approach or to allow a colleague to attempt the procedure. Use a new setup each time that blood is obtained because clots and tissue will clog the needle and mislead the clinician even if the vein has been entered successfully on subsequent attempts. If several attempts are made, inform the admitting clinician or anesthesiologist so that proper precautions are taken to identify subsequent complications. It is advisable to obtain radiographs of the chest even after unsuccessful attempts. If the initial puncture site was placed properly, use the same needle hole for subsequent attempts if possible for aesthetic reasons. If the SV route is unsuccessful on one side, attempt IJ vein catheterization on the same side rather than SV cannulation on the opposite side to avoid bilateral complications.

**Supraclavicular Subclavian Approach**

**Positioning**

The goal of the supraclavicular SV technique is to puncture the SV in its superior aspect as it joins the IJ vein. Insert the needle above and behind the clavicle, lateral to the clavicular head of the SCM muscle. Advance it in an avascular plane while directing it away from the subclavian artery and the dome of the pleura (Fig. 22.14). The right side is preferred.
because of the lower pleural dome, its more direct route to the SVC, and location of the thoracic duct on the left side. The patient’s head may be turned to the opposite side to help identify the landmarks.

**Needle Orientation**

After the area of the supraclavicular fossa has been prepared and draped, identify a point 1 cm lateral to the clavicular head of the SCM and 1 cm posterior to the clavicle. Alternatively, use the junction of the middle and medial thirds of the clavicle as the landmark for needle entry. Anesthetize the area with 1% lidocaine. If a 3-cm-long needle is used for anesthesia, it may also be used to locate the vessel in a relativelyatraumatic manner. The SV can almost always be located with this needle because of its superficial location and the absence of bony structures in the path of the needle. Advance a 14-gauge needle (or 18-gauge thin-walled needle) along the path of the scout needle. Apply gentle negative pressure with an attached syringe.

When seeking the SV, aim the needle so that it bisects the clavicosternomastoid angle and the tip points just caudal to the contralateral nipple. Orient the bevel medially to prevent the catheter from getting trapped against the inferior vessel wall. Point the tip of the needle 10 degrees above the horizontal. Successful vessel puncture generally occurs at a depth of 2 to 3 cm.

**Subclavian Ultrasound Technique**

Typically, puncture of the SV occurs at the point where the vein is coursing deep to the clavicle. However, with ultrasound, visualization of the SV can be difficult at this location because of interference with the overlying bone. Fortunately, more distally the vein lies farther away from the clavicle and chest wall. Hence, access to the SV typically occurs lateral to the curve of the clavicle bone, in the proximal axillary vein. As the vein moves laterally, the mean depth from the skin increases from 1.9 to 3.1 cm, whereas the distance from the rib cage to the vein increases from 1 to 2 cm. The arteriovenous distance also increases from 0.3 to 0.8 cm and there is less overlap of the artery and vein. Because the vein is not in close proximity to the clavicle, if a hematoma develops, manual pressure can be used to stop the bleeding. Furthermore, the axillary vein is farther from the chest wall and pleural surface, thus decreasing the possibility of pleural injury and subsequent pneumothorax. In clinical studies, ultrasound-guided axillary vein access had a first–needle pass success rate of 76% with successful placement in 96% of cases. Despite the use of ultrasound, however, the catheter malposition rate was unchanged at approximately 15%.

**IJ Approach**

**Positioning**

Position is critical for maximizing the success of blind (landmark technique) cannulation of the IJ vein. Place the patient in a supine position with the head down and turned approximately 15 to 30 degrees away from the IJ vein to be cannulated. Rotate the head slightly away from the site of insertion. Rotating the head more than 40 degrees has been shown to increase the risk for overlapping the carotid artery over the IJ vein. Occasionally, placing a towel roll under the scapula helps extend the neck and accentuate the landmarks. Stand at the head of the bed with all equipment within easy reach. This may involve moving the bed to the center of the room to allow a table or work surface to be placed at the head of the bed.

Ask the patient to perform a Valsalva maneuver just before inserting the needle to increase the diameter of the IJ vein. Alternatively, the patient can be asked to hum. Trendelenburg positioning, the Valsalva maneuver, and humming all increase the area of the vessel by approximately 30% to 40%. If the patient is unable to cooperate, coordinate the insertion with respiration because the IJ vein is at its largest diameter just before inspiration. In intubated patients this relationship is reversed because mechanical ventilation increases intrathoracic pressure at end-inspiration. External abdominal compression also helps distend the IJ vein.
Central Venous Catheterization and Central Venous Pressure Monitoring

Needle selection may vary depending on the central line kit used. The operator may choose from three approaches: central, posterior, and anterior (Fig. 22.15).

Central Route
This approach is favored by some who believe that the incidence of cannulation of the carotid artery is decreased and the cupula of the lung is avoided. First, palpate and identify the triangle formed by the clavicle and the sternal and clavicular heads of the SCM. Use a marking pen or a local anesthetic skin wheal to mark the lateral border of the carotid pulse, and perform all subsequent needle punctures lateral to this point. Aspirate before injecting anesthetic so that it is not injected into the carotid artery. Once infiltration is completed, use the needle to locate the IJ vein by aspirating blood into the syringe. Note the depth and angle of needle entry and use this as a mental guide to finding the IJ vein with the introducer needle. Typically, an 18-gauge 2.5-cm introducer needle attached to a syringe is used initially to puncture the IJ vein. However, needle selection may vary depending on the central line kit used. The operator may choose from three approaches: central, posterior, and anterior (Fig. 22.15).

Venipuncture Site
The right IJ vein provides a more direct route to the right atrium for transvenous pacing. The left IJ vein is often more tortuous and catheters must negotiate two 90-degree turns at the junction of the left IJ vein with the SV and at the junction of the SV with the SVC. However, if the right IJ vein is obstructed or scarred by previous access, the left IJ vein may be accessed with the same technique. Of note, the right IJ vein has been observed to be twice the size of the left IJ vein in 34% of normal adults.

Aspirate before injecting anesthetic so that it is not injected into the carotid artery or IJ vein. Once infiltration is completed, use the needle to locate the IJ vein by aspirating blood into the syringe. Note the depth and angle of needle entry and use this as a mental guide to finding the IJ vein with the introducer needle. Typically, an 18-gauge 2.5-cm introducer needle attached to a syringe is used initially to puncture the IJ vein. However, needle selection may vary depending on the central line kit used. The operator may choose from three approaches: central, posterior, and anterior (Fig. 22.15).

Central Approach
Insert needle at the apex of the triangle formed by the heads of the sternocleidomastoid muscle and the clavicle.

Entry angle = 30°.

Aim toward the ipsilateral nipple.

Note: estimate the course of the IJ vein by placing three fingers lightly over the carotid artery as it runs parallel to the vein. The vein lies just lateral to the artery, albeit often minimally so.

Anterior Approach
Insert needle along the medial edge of the sternocleidomastoid, 2-3 fingerbreadths above the clavicle.

Entry angle = 30° to 45°.

Aim toward the ipsilateral nipple.

Note: palpate the carotid artery during venipuncture. The artery may be slightly retracted medially.

Posterior Approach
Insert needle at the posterior (lateral) edge of the sternocleidomastoid, midway between the mastoid process and the clavicle.

Entry angle = 45°.

Aim toward the suprasternal notch.

Note: avoid the external jugular vein, which crosses the posterior SCM border. During needle advancement, apply pressure to the SCM to lift the body of the muscle. The vein is usually reached at a depth of 7 cm.

Figure 22.15 Approaches to the internal jugular vein. IJ, Internal jugular; SCM, sternocleidomastoid.
the course of the IJ vein by placing three fingers lightly over the course of the carotid artery as it runs parallel to the vein, using the fingers as a guide for needle placement. The vein consistently lies just lateral to the carotid artery, albeit often minimally so. Prolonged deep palpation of the carotid artery may decrease the size of the vein, so use the three-finger technique lightly to identify the course of the artery.

**Posterior and Anterior Routes**

In the posterior approach, make the puncture at the posterior (lateral) edge of the SCM, approximately midway between its origin at the mastoid process and its insertion at the clavicle. The external jugular vein courses in this area and can be used as a landmark, with the puncture occurring where the external jugular vein crosses the posterolateral border of the SCM. Be careful to not strike the external jugular vein. Advance the needle toward the suprasternal notch, just under the belly of the SCM, at an angle of approximately 45 degrees to the transverse plane. During advancement of the needle, apply pressure to the SCM in an effort to lift the body of the muscle. The vein is usually reached at a depth of 7 cm in an average-sized adult. Because the posterior approach occurs higher in the neck, there is less risk for hemothorax, pneumothorax, or carotid puncture. The benefits of the posterior approach are more dramatic in obese patients, with carotid puncture occurring in 3% of patients versus up to 17% with the anterior approach.

In the anterior approach, needle puncture occurs along the anterior or medial edge of the SCM approximately two to three fingerbreadths above the clavicle. Insert the needle at an angle of 30 to 45 degrees toward the ipsilateral nipple, away from the carotid pulse. If cannulation is unsuccessful, withdraw the needle to the skin and redirect it slightly toward the carotid artery.

Once the approach is chosen, slowly advance the needle toward the IJ vein. Create gentle negative pressure with the syringe while advancing the needle. Once blood is seen, stop advancing the syringe. Remove the syringe from the needle to determine whether the vessel is pulsatile. Be careful not to allow negative intrapleural pressure to draw air into the venous system through the open needle. Because the tip of the introducer needle is beveled, lateral motions of the needle tip may cause lacerations of the deep structures of the neck. It is therefore very important to remove the needle from the neck completely before any redirection of the needle.

Once cannulation of the IJ vein has been confirmed, remove the syringe from the needle and place a gloved digit over the needle hub to prevent air embolism. Insert a guidewire through the needle into the IJ vein and place the catheter using the Seldinger technique. Once the wire is inserted into the IJ vein, reduce the angle to the skin to make the needle nearly parallel to the vein. This allows a greater chance of directing the wire toward the heart. Do not let the guidewire extend into the right atrium. The average distance from the insertion site to the junction of the SVC, and right atrium is 16 ± 2 cm for the right IJ vein and 19 ± 2 cm for the left IJ vein. The spring wires supplied in kits are often much longer, up to 60 cm in length. If the full length of the wire is inserted, the wire could enter the right atrium or ventricle and cause myocardial irritability and subsequent dysrhythmias. Monitor cardiac rhythm during insertion of the spring wire to detect cardiac irritability. The distance that the catheter is introduced depends on the distance from the site of introduction to the junction of the SVC and right atrium. This distance will be shorter with the right IJ vein than with the left IJ vein.

**II Ultrasound Technique**

Cannulation of the IJ vein is an optimal location for the use of ultrason guidance. Whereas the landmark approach is associated with a complication rate of between 5% and 10% irrespective of the technique used or experience of the operator, with the use of ultrasound, the complication rate is significantly reduced. Even with novice users of ultrasound for IJ vein cannulation, first-attempt success is significantly increased when compared to the blind landmark technique (43% versus 26%). With experience, however, the first-attempt success rate improves to more than 75%. Use of ultrasound for placement of central lines in the IJ vein has also been shown to decrease overall catheter placement failures by 64%, reduce complications by 78%, and decrease the need for multiple catheter placement attempts by 40% in comparison to the standard landmark placement technique. The primary reason for the increased success rate is the variation in anatomy of the IJ vein relative to the carotid artery. The anatomy of the IJ vein has been shown to be aberrant in 9% to 19% of cases. Furthermore, the IJ vein may be unusually small (i.e., <0.5 cm) in up to 14% of patients. The IJ vein is thrombosed in up to 2.5% of some patient populations.

With the use of ultrasound there is no need for reliance on normal anatomy for cannulation. Therefore the IJ vein may be cannulated despite abnormal anatomy. Hence, cannulation may occur at the apex of the triangle, near the base at the junction with the innominate vein, or anywhere in between.

**Femoral Approach**

**Positioning and Needle Orientation**

Place the patient in the supine position for the femoral vein approach. This approach does not require any special positioning or tilting of the bed. Fully expose and thoroughly cleanse the area with a soap and water solution. Insert the needle at a 45-degree angle in a cephalic direction approximately 1 cm medial to this point and toward the umbilicus (Fig. 22.16). Palpate the femoral pulse two fingerbreadths beneath the inguinal ligament. Insert the needle at a 45-degree angle in a cephalic direction approximately 1 cm medial to this point and toward the umbilicus (Fig. 22.16). Palpate the femoral pulse two fingerbreadths beneath the inguinal ligament. Place the femoral pulse as a guide is ideal, it may not be palpable in an obese or hypotensive patient. A
more detailed understanding of the femoral landmarks can be used to guide cannulation attempts. On all but the most severely injured trauma patients with a disrupted pelvis (in which case a femoral approach would be contraindicated), the anterior superior iliac spine and the midpoint of the pubic symphysis are easily palpated. The line between these two bony references describes the inguinal ligament. When this line is divided into thirds, the femoral artery should underlie the junction of the medial and middle thirds. The femoral vein will lie approximately one fingerbreadth medial to this point. Alternatively, the vascular anatomy of the region can be evaluated and the line placed via ultrasound guidance.

**Venipuncture**

During advancement of the needle, maintain gentle negative pressure on the syringe at all times while the needle is under the skin. Direct the needle posteriorly and advance it until the vein is entered, as identified by a flash of dark, nonpulsatile blood. If the vessel is penetrated when the syringe is not being aspirated, the flash of blood may be seen only as the needle is being withdrawn. The femoral vein lies just medial to the femoral artery at the level of the inguinal ligament. It is closer to the artery than many clinicians appreciate. As the vein progresses distally in the leg, it runs closer to and almost behind the femoral artery.

**Femoral Ultrasound Technique**

Cannulation of the femoral vein under ultrasound guidance is very similar to that for the IJ vein. Using ultrasound, the common femoral vein, its junction with the saphenous vein, and the branches of the common femoral vein (i.e., the superficial and deep femoral veins) are easily identified. Typically, placement of the catheter should occur proximal to the bifurcation of the common femoral vein and preferably proximal to the junction with the saphenous vein.

**ANCHORING THE CENTRAL LINE**

After the CVC is placed, it will need to be anchored in place by one of three techniques: StatLock (Bard Medical, Covington, GA), suture, or staple (Fig. 22.17). The StatLock may not hold well in patients with oily skin but is excellent for older patients with thin skin. For suturing, one will need the sterile, nonabsorbable suture material (usually 2-0 silk) provided in the CVC kit. The straight suture needles found in many sets are awkward for many clinicians, so a curved needle with a driver may be helpful. To avoid a needlestick with the straight needle, pass the blunt end of the needle through the anchoring devices and pull the suture forward manually. Place the suture in the skin approximately a half centimeter from the catheter to anchor the central line in place. Several knots should be made to secure the line. Avoid making knots that place excessive pressure on the skin because this can lead to difficulty removing the knots and necrosis. Loose knots can lead to migration of the catheter and loss of access. Stapling a central line into place can be just as effective as suturing; however, the staples tend to fall out after a few days.

**Dressing**

Clean the area around the catheter insertion site with chlorhexidine, and then use a clear dressing (such as Tegaderm [3M, St. Paul, MN]) to cover up the insertion site of the catheter once secured (see Fig. 22.9, step 16). Apply a chlorhexidine patch (Biopatch [Ethicon, Somerville, NJ]) at the site where the catheter enters the skin (see Fig. 22.17E and F). Because dressings are inspected and changed periodically, place a simple dressing and avoid excessive amounts of gauze and tape. Take care to protect the skin against maceration.

**Assessing Line Placement**

Check all tubing and connections for tightness to prevent air embolism, loss of fluid, or bleeding. Before infusing IV fluids, lower the IV fluid reservoir to below the level of the patient’s right atrium and check the line for backflow of blood. Free backflow of blood is suggestive but not diagnostic of intravascular placement. Backflow could occur from a hematoma or hemothorax if the catheter is free in the pleural space. A pulsatile blood column may be noted if the catheter has been inadvertently placed in an artery. Less pronounced pulsations might also occur if the catheter is advanced too far and reaches the right atrium or ventricle. In addition, pulsations may be noted with changes in intrathoracic pressure as a result of respirations, although these pulsations should occur at a much slower rate than the arterial pulse. A final method of checking intravascular placement is to attach a syringe directly to the catheter hub and aspirate venous blood. It is also advisable to ensure that the catheter is easily flushed with a saline solution. This carries the additional benefit of removing air from the system. Radiographs are also always indicated to verify catheter location and assess for potential complications, except after routine femoral line placements. In an awake patient, infusing fluids
When evaluating the internal jugular (IJ) vein, the transducer (7.5 to 20 MHz) should be initially placed over the right or left side of the neck to evaluate the anatomy. An ideal initial location to begin is at the apex of the triangle formed by the two heads of the subclavian muscle (Fig. 22.US1). Placing the transducer over this area in the transverse orientation will enable the vessels to be located in cross section, where they can best be evaluated. The internal carotid artery and IJ vein will be seen as paired structures with anechoic central areas (Fig. 22.US2). The position of one relative to the other can be variable, but typically the IJ vein lies lateral and superficial to the carotid artery. Several characteristics of the IJ vein serve to distinguish it from the carotid artery. The IJ vein is typically more oval in shape (versus the rounded shape of the carotid artery), is thinner walled, and will compress with gentle pressure. Additionally, the size of the IJ vein will change with respiration and should be seen to increase in size with a Valsalva maneuver.

Complications can be reduced by several methods. First, ensure that the target vessel is indeed the vein and not the artery. Variant anatomy or variations in volume status (either depletion or overload) may make the vessels difficult to distinguish from one another. Confirmation should be attempted by noting multiple characteristics of the vessel (compressibility, shape, anatomic location, etc.). Once the vessel has been confirmed as the vein, the operator must take great care to ensure that the position of the tip of the needle is apparent at all times. Most complications occur when the tip of the needle is deeper or more medial than the operator realizes, thus placing it in proximity to other structures (e.g., lung, carotid artery). An extensive discussion of each approach can be found in the basic ultrasound chapter (see Chapter 66), and each approach has its drawbacks in determining position. In the transverse method, the angle of approach can be difficult to ascertain and cause the tip of the needle to be deeper than the operator realizes. Additionally, the tip of the needle may be difficult to follow. In the longitudinal approach, the medial to lateral orientation of the needle can be difficult to appreciate. Additionally, slight movements of the transducer may result in loss of the appropriate image. A combination of these two, or an oblique approach, may minimize these difficulties.

The femoral artery and vein lie together with the femoral nerve within a common sheath. They can be found at the level of the inguinal crease on the medial aspect of the thigh. Palpating the femoral pulse will also aid in localizing the vascular bundle. The transducer (7.5 to 10 MHz) should be placed in a transverse or slightly oblique orientation overlaying this area. Slightly rotating the thigh externally may facilitate this step. Classically, the artery is described to lie lateral to the vein. However, this is often not the case and multiple variations may be noted. The femoral artery and vein will appear as rounded anechoic structures (Fig. 22.US3). The femoral vein can be recognized

**Figure 22.US2** Transverse image of the carotid artery and internal jugular (IJ) vein. The IJ vein can be recognized by its oval or triangular shape and its thinner walls, and it will collapse with light pressure. Although both are rounded and contain anechoic (black) fluid, the IJ vein is slightly larger and more oval in shape. Though not evident in this image, slight pressure will cause collapse of the IJ vein. CA, Carotid artery.

**Figure 22.US3** Transverse image of the femoral artery and femoral vein. Similar to vessels in the neck, the femoral vein is more oval or triangular in shape. Though not evident in this image, slight pressure will cause collapse of the vessel. FA, Femoral artery; FV, femoral vein.

---

**IJ Vein**

When evaluating the internal jugular (IJ) vein, the transducer (7.5 to 20 MHz) should be initially placed over the right or left side of the neck to evaluate the anatomy. An ideal initial location to begin is at the apex of the triangle formed by the two heads of the subclavian muscle (Fig. 22.US1). Placing the transducer over this area in the transverse orientation will enable the vessels to be located in cross section, where they can best be evaluated. The internal carotid artery and IJ vein will be seen as paired structures with anechoic central areas (Fig. 22.US2). The position of one relative to the other can be variable, but typically the IJ vein lies lateral and superficial to the carotid artery. Several characteristics of the IJ vein serve to distinguish it from the carotid artery. The IJ vein is typically more oval in shape (versus the rounded shape of the carotid artery), is thinner walled, and will compress with gentle pressure. Additionally, the size of the IJ vein will change with respiration and should be seen to increase in size with a Valsalva maneuver.

Complications can be reduced by several methods. First, ensure that the target vessel is indeed the vein and not the artery. Variant anatomy or variations in volume status (either depletion or overload) may make the vessels difficult to distinguish from one another. Confirmation should be attempted by noting multiple characteristics of the vessel (compressibility, shape, anatomic location, etc.). Once the vessel has been confirmed as the vein, the operator must take great care to ensure that the position of the tip of the needle is apparent at all times. Most complications occur when the tip of the needle is deeper or more medial than the operator realizes, thus placing it in proximity to other structures (e.g., lung, carotid artery). An extensive discussion of each approach can be found in the basic ultrasound chapter (see Chapter 66), and each approach has its drawbacks in determining position. In the transverse method, the angle of approach can be difficult to ascertain and cause the tip of the needle to be deeper than the operator realizes. Additionally, the tip of the needle may be difficult to follow. In the longitudinal approach, the medial to lateral orientation of the needle can be difficult to appreciate. Additionally, slight movements of the transducer may result in loss of the appropriate image. A combination of these two, or an oblique approach, may minimize these difficulties.

**Femoral Vein**

The femoral artery and vein lie together with the femoral nerve within a common sheath. They can be found at the level of the inguinal crease on the medial aspect of the thigh. Palpating the femoral pulse will also aid in localizing the vascular bundle. The transducer (7.5 to 10 MHz) should be placed in a transverse or slightly oblique orientation overlaying this area. Slightly rotating the thigh externally may facilitate this step. Classically, the artery is described to lie lateral to the vein. However, this is often not the case and multiple variations may be noted. The femoral artery and vein will appear as rounded anechoic structures (Fig. 22.US3). The femoral vein can be recognized

---

**ULTRASOUND BOX 22.1: Central Venous Catheterization**

by Christine Butts, MD

**I} vein**

When evaluating the internal jugular (IJ) vein, the transducer (7.5 to 20 MHz) should be initially placed over the right or left side of the neck to evaluate the anatomy. An ideal initial location to begin is at the apex of the triangle formed by the two heads of the subclavian muscle (Fig. 22.US1). Placing the transducer over this area in the transverse orientation will enable the vessels to be located in cross section, where they can best be evaluated. The internal carotid artery and IJ vein will be seen as paired structures with anechoic central areas (Fig. 22.US2). The position of one relative to the other can be variable, but typically the IJ vein lies lateral and superficial to the carotid artery. Several characteristics of the IJ vein serve to distinguish it from the carotid artery. The IJ vein is typically more oval in shape (versus the rounded shape of the carotid artery), is thinner walled, and will compress with gentle pressure. Additionally, the size of the IJ vein will change with respiration and should be seen to increase in size with a Valsalva maneuver.

Complications can be reduced by several methods. First, ensure that the target vessel is indeed the vein and not the artery. Variant anatomy or variations in volume status (either depletion or overload) may make the vessels difficult to distinguish from one another. Confirmation should be attempted by noting multiple characteristics of the vessel (compressibility, shape, anatomic location, etc.). Once the vessel has been confirmed as the vein, the operator must take great care to ensure that the position of the tip of the needle is apparent at all times. Most complications occur when the tip of the needle is deeper or more medial than the operator realizes, thus placing it in proximity to other structures (e.g., lung, carotid artery). An extensive discussion of each approach can be found in the basic ultrasound chapter (see Chapter 66), and each approach has its drawbacks in determining position. In the transverse method, the angle of approach can be difficult to ascertain and cause the tip of the needle to be deeper than the operator realizes. Additionally, the tip of the needle may be difficult to follow. In the longitudinal approach, the medial to lateral orientation of the needle can be difficult to appreciate. Additionally, slight movements of the transducer may result in loss of the appropriate image. A combination of these two, or an oblique approach, may minimize these difficulties.

---

**Femoral Vein**

The femoral artery and vein lie together with the femoral nerve within a common sheath. They can be found at the level of the inguinal crease on the medial aspect of the thigh. Palpating the femoral pulse will also aid in localizing the vascular bundle. The transducer (7.5 to 10 MHz) should be placed in a transverse or slightly oblique orientation overlaying this area. Slightly rotating the thigh externally may facilitate this step. Classically, the artery is described to lie lateral to the vein. However, this is often not the case and multiple variations may be noted. The femoral artery and vein will appear as rounded anechoic structures (Fig. 22.US3). The femoral vein can be recognized

---

**Figure 22.US1** Placement of the ultrasound transducer at the apex of the triangle formed by the heads of the sternocleidomastoid.

**Figure 22.US2** Transverse image of the carotid artery and internal jugular (IJ) vein. The IJ vein can be recognized by its oval or triangular shape and its thinner walls, and it will collapse with light pressure. Although both are rounded and contain anechoic (black) fluid, the IJ vein is slightly larger and more oval in shape. Though not evident in this image, slight pressure will cause collapse of the IJ vein. CA, Carotid artery.

**Figure 22.US3** Transverse image of the femoral artery and femoral vein. Similar to vessels in the neck, the femoral vein is more oval or triangular in shape. Though not evident in this image, slight pressure will cause collapse of the vessel. FA, Femoral artery; FV, femoral vein.
by its thinner walls and slightly more oval shape, and it will collapse with gentle pressure. It will also typically increase in size when the lower part of the leg is squeezed. The vascular bundle may need to be followed inferiorly or superiorly to determine the most optimal location for puncture.

Complications can be reduced by several methods. First, ensure that the target vessel is indeed the vein and not the artery. Variant anatomy or variations in volume status (either depletion or overload) may make the vessels difficult to distinguish from one another. Confirmation should be attempted by noting multiple characteristics of the vessel (compressibility, shape, anatomic location, etc.). Once the vessel has been confirmed as the vein, the operator must take great care to ensure that the position of the tip of the needle is apparent at all times. The tip of the needle may be difficult to follow in the transverse approach and result in an inadvertent puncture of the posterior wall of the vessel. When the artery lies deep to the vein, arterial puncture or cannulation may result. In the longitudinal approach, the medial to lateral position of the needle may be difficult to appreciate and result in accidental arterial puncture. The oblique approach may minimize these difficulties.

Subclavian Vein
The subclavian vessels can be imaged from either a supraclavicular or an infraclavicular approach. For the supraclavicular approach, the transducer (7.5 to 10 MHz) is placed along the long axis of the clavicle on the superior aspect (Fig. 22.US4). It should be angled downward. In this view, the vessels should be seen in their long axis (Fig. 22.US5). The vein can be identified by its variation with respiration and change in size with the Valsalva maneuver. The vein can also be followed to identify the junction with the IJ vein, thereby offering further confirmation. In the infraclavicular approach, the transducer is placed beneath the clavicle at its most lateral aspect, in a sagittal (shown) or slightly oblique orientation, following the position of the clavicle (Fig. 22.US6). In this view, the vessels will be seen in cross section or a slightly oblique plane (Fig. 22.US7). The pleura may also be seen deep to the vessels as an echogenic vertical line that slides back and forth with respiration.
A longitudinal approach should be used in which the needle is introduced from the end of the transducer in either the infraclavicular or the supraclavicular approaches. This will enable a shallow angle to be used and thereby minimize the chance of damaging deeper structures such as the lung. Once a flash of blood has been obtained, the ultrasound transducer can be set aside and the procedure continued as described previously.

The subclavian artery and vein lie in close opposition to the pleura, so pneumothorax is a more common complication. Using a long-axis approach (in which the needle is introduced from the end of the transducer rather than from the middle) offers the advantage of visualizing the entirety of the needle in its course toward the vein. A shallow angle can be used, and the relationship of the needle to the pleura can also be appreciated.

**Securing a Central Venous Catheter**

A white rubber clamp is provided to secure the catheter when the full length is not needed. Twist open the pliable clamp and place it over the catheter at a site a few centimeters from the insertion site.

The rubber clamp is covered with a blue plastic fastener, and both the clamp and fastener are sutured to the skin to secure the catheter. The hub of the catheter is also sutured to the skin.

To avoid a needlestick, the blunt end of the needle is used to pass the suture through the holes of the fastening devices.

Alternatively, skin staples may be used. Tent the skin and pass the staples through the anchoring eyes.

This Biopatch is a chlorhexidine-containing hydrophilic covering placed at the site where the catheter enters the skin to deliver local antisepsis for 7 days.

A simple Tegaderm clear covering is then applied.

**Figure 22.17** Methods to secure a central venous triple-lumen catheter.
via a catheter tip positioned in the IJ vein may produce an audible gurgling or flowing sound in the patient’s ear.111

**Radiographs**

Following placement of lines involving puncture of the neck or thorax, listen to the lungs to detect any inequality of lung sounds suggestive of a pneumothorax or hemothorax. Obtain a chest film as soon as possible to check for hemothorax, pneumothorax, and the position of the tip of the catheter (Fig. 22.18). Because small amounts of fluid or air may layer out parallel to the radiographic plate with the patient in the supine position, take the film in the upright or semi-upright position whenever possible. In ill patients, a rotated or oblique projection on a chest radiograph may be obtained, and the clinician may be confused about the proper position of the catheter. In such cases, repeat the radiograph. A misplaced catheter tip is usually obvious on a properly positioned standard posteroanterior chest radiograph (Fig. 22.19), but occasionally, injection of contrast material may be required. For example, a catheter in one of the internal thoracic veins may simply appear more lateral than expected, but because of the close proximity of these veins and the SVC, malposition may not be appreciated by this subtle finding. Misplaced catheters should be repositioned or replaced.

Attention should also be given to the possibility of a retained guidewire. Although this complication is rare, if not specifically considered it can be overlooked by both clinicians and radiologists.112,113

A postprocedure radiograph is routine after initial placement, but radiographs are not always necessary for routine replacement of catheters over guidewires. If such patients are stable and hemodynamically monitored, radiography may be deferred safely in the absence of apparent complications or clinical suspicion of malposition.114,115

**Redirection of Misplaced Catheters**

Improper catheter tip position occurs commonly. It has been reported that only 71% of SV catheters are located in the SVC on the initial chest film. Complications of improper positioning include hydrothorax, hemothorax, ascites, chest wall abscess, embolization to the pleural space, and chest pain. More commonly, improper location yields inaccurate measurements of CVP or is associated with poor flow caused by kinking. An unusual complication attributable to improper tip position is cerebral infarction, which can occur following inadvertent cannulation of the subclavian artery.

Misdirection or inappropriate positioning of the tip of a CVC, when promptly recognized and corrected, is an inconsequential complication. Loop formation, lodging in small neck veins, tips directed caudally, and innominate vein position are common problems. Reposition misplaced catheters as soon as logically possible. If the catheter is being used for fluid resuscitation, the malposition may be tolerated for some time. If vasopressors or medications are infused, proper positioning of the tip of the catheter is more critical. A number of options are available to remedy malpositioning. One strategy is to insert a 2-Fr Fogarty catheter through the lumen of the central line and advance it 3 cm beyond the tip. Withdraw the entire
assembly until only the Fogarty catheter is in the SV. Inject 1 mL of air into the balloon and advance the Fogarty catheter. It is hoped that blood flow will direct the assembly into the SVC. Deflate the balloon and advance the central line over the Fogarty catheter, which is then withdrawn.

Another anecdotal strategy is to withdraw the catheter until only the distal tip remains in the cannulated vessel. This measurement is best appreciated by comparing the length of the indwelling catheter with another unused catheter. The clinician then simply readvances the catheter in the hope that it becomes properly positioned. Other manipulations with guidewires have been suggested, but reininsertion with another puncture is often required for the misplaced catheter to be positioned properly. This approach also decreases the risk for infection by avoiding the introduction of bacteria into the vessel from any nonsterile segment of the CVC.

SPECIAL CONSIDERATIONS FOR OTHER VESSELS

External Jugular Vein Approach

Central venous catheterization via the external jugular vein is time-consuming and often difficult. The difficulty in converting an external jugular catheter into a CVC frequently renders it a lower-yield clinical procedure. Use of the external jugular vein for achieving central venous access requires that a guidewire be used. After cannulation of the vein and intraluminal placement of the guidewire, advance the guidewire into the thorax by rotating and manipulating the tip into the central venous circulation. Advancement of the guidewire is the most difficult and time-consuming portion of the procedure, and the time requirement limits the usefulness of this technique in an emergency. A small-radius J-tipped wire, a distended vessel lumen, and exaggeration of patient head tilt, coupled with skin traction, may facilitate successful passage of the guidewire. Partially withdrawing the wire and twisting it 180 degrees before readvancing the tip may also be helpful.

Basilic and Cephalic Approaches

Passing a catheter into the central circulation is difficult via the basilic and cephalic routes, and failure is common. Insertion of a peripheral IV central catheter through these routes is often performed by specialized teams and is less suitable for emergency indications. The cephalic vein may terminate inches above the antecubital fossa or bifurcate before entering the axillary vein, while the basilic vein, located on the medial aspect of the antecubital fossa, is generally larger than the radially located cephalic vein. Furthermore, the basilic vein usually provides a more direct route for passage into the axillary vein, SV, and SVC.

Vascular Access in Cardiac Arrest

Immediate vascular access is required for resuscitation during cardiac arrest. Intraosseous access is a reasonable alternative to central venous access. Femoral CVCs are often used in this setting. The infraclavicular SV approach is also commonly used during cardiac arrest if logistics permit. The intuitive rationale for femoral CVC placement has been that much of the resuscitation activity, including chest compressions, occurs on the thorax, thus limiting the clinician’s ability to safely place a higher line. During cardiac arrest, the availability of drugs delivered to the central circulation may be slower via the femoral route than via supraclavicular SV or IJ vein infusions. Additionally, pulsations felt in the groin during CPR may be venous instead of arterial, and there is a high rate of unrecognized catheter malposition and arterial injury. To place a femoral catheter blindly (without ultrasound guidance or clear identification of the arterial pulse), divide the distance from the anterior superior iliac spine to the symphysis pubis into thirds. The artery typically lies at the junction of the medial and middle thirds and the vein is 1 cm medial to this location. Blind femoral central line insertions during arrest are less than optimal. The increasingly available intraosseous placement systems and bedside ultrasonography are commonly used to supplant such blind CVC placements during cardiac arrest and other emergencies that require immediate vascular access.

CVP MONITORING

CVP Measurement

Although described by Forssmann in 1931, it was not until the early 1960s that measurement of CVP became commonplace as a means of assessing cardiac performance and guiding fluid therapy. CVP measurement has been used as a guide for determination of volume status, fluid requirements, and investigation of tamponade, but its reliability has not been consistently demonstrated in the literature. Furthermore, advancements in our knowledge of complex hemodynamics (particularly during sepsis) and improvements in noninvasive dynamic imaging (ultrasound) have largely supplanted CVP monitoring in the critical care environment. Still, astute clinicians can maximize the usefulness of this diagnostic tool by understanding its basic principles, indications, and limitations.
Physiology

Simply stated, CVP is the pressure exerted by blood against the walls of the intrathoracic venae cavae. Because pressure in the venae cavae is generally within 1 mm Hg of right atrial pressure, CVP reflects the pressure under which blood is returning to the heart. Pressure in the central veins has two significant hemodynamic effects. First, CVP promotes filling of the heart during diastole, a factor that reflects right ventricular end-diastolic volume (preload). Second, CVP also represents the back-pressure of the systemic circulation, which opposes return of blood from peripheral venous circulation to the heart. CVP values are determined by a complex interaction of intravascular volume, right atrial and ventricular function, venomotor tone, and intrathoracic pressures.119,124,125,127

To measure CVP, the tip of a pressure-monitoring catheter is inserted into any of the great systemic veins of the thorax or into the right atrium.124,126 The femoral vein may also be used for CVP measurements as long as there is no abnormally increased abdominal pressure.10,127 The catheter is connected to a simple manometer or to an electronic pressure transducer interfaced with a monitoring system that is capable of calculating a mean pressure value and displaying pressure waveforms.124,126 When the catheter tip is placed in the right atrium the waveforms produced correlate with the cardiac cycle and create a typical wave pattern.

A common misconception is that CVP consistently reflects pressure in the left side of the heart. A different measurement, called pulmonary capillary wedge pressure (PCWP), is capable of reflecting left atrial pressure and corresponding changes in left ventricular pressure by transducing back-pressures from the left-sided circulation through the pulmonary capillary system. Additionally, flow-directed pulmonary artery catheters allow repeated calculations of PCWP, thus facilitating reliable estimation of left atrial pressures.126

Indications for and Contraindications to CVP Measurement

The five traditional major indications for monitoring CVP are:
1. Acute circulatory failure.
2. Anticipated massive blood transfusion or fluid replacement therapy.
4. Suspected cardiac tamponade.
5. Fluid resuscitation during goal-directed therapy in patients with severe sepsis.

The procedure is contraindicated when other resuscitative therapeutic and diagnostic interventions take priority over central venous access and CVP transducer setup and calibration, or in the setting of large vegetations on the tricuspid valve, SVC syndrome, or tumors or thrombus in the right atrium.

CVP monitoring is most helpful in patients without significant preexisting cardiopulmonary disease. Numerous studies highlight the unreliability of right-sided hemodynamic monitoring in patients with underlying cardiac or pulmonary disease.114,118

Review Box 22.2 Central venous pressure measurement: indications, contraindications, complications, and equipment.
Ultimately, however, the inconsistencies noted are not due to the failure of CVP monitors to reflect central hemodynamics. Rather, the discrepancies noted in the literature simply highlight the complex relationships between ventricular and vascular compliance, blood volume, and filling pressures in various disease states. As with pulmonary artery occlusion pressure measurements, the clinician is cautioned to be aware of the assumptions inherent in taking these measurements and to recognize the physiologic scenarios in which these assumptions do not hold true.

### Procedure

Although CVP may be determined with a manometry column assembled at the bedside (Fig. 22.20), the most common technique in practice is measurement with an electronic

---

**Measurement of Central Venous Pressure: Manometry**

1. Assemble the manometer as depicted above. When the stopcock is turned to direct flow of fluid to the patient, the manometer is bypassed. This is the position that is maintained to keep the catheter patent. Remember to always flush all tubing before connecting it to the patient’s central catheter.

2. To measure central venous pressure, first turn the stopcock to fill the manometer to 25 cm H₂O.

3. Next, open the stopcock to the patient and the manometer. Allow the column of water in the manometer to fall and stabilize before a reading is taken. Note that the zero mark must be horizontally aligned with the tricuspid valve (which is estimated as the midaxillary line in a supine patient).

---

*Figure 22.20* Measurement of central venous pressure with a manual manometer.
transducer interfaced to a bedside monitoring system (Fig. 22.21). Typical transducers include a nipple valve attached to a pressurized bag of saline to allow easy flushing of the system. The transducer is connected through the patient's central line with a length of tubing filled with saline. A three-way stopcock is placed between the patient and the transducer to simplify line flushing and calibration.

Flush all air bubbles from the system by opening the stopcock to air and flushing saline through the line. Do not flush air bubbles into the patient. This will dampen the CVP wave and potentially cause underestimation of venous pressure.

After the system has been flushed, place the stopcock (with the transducer still open to air) at the level of the patient's right atrium. Zero (calibrate) the monitor with the transducer at the level of the right atrium, which can be approximated on the skin surface as a point at the midaxillary line and the fourth intercostal space. Finally, set the stopcock so that the transducer is in continuity with the patient's venous catheter.

In spontaneously breathing patients, take readings at the end of a normal inspiration. If the patient is receiving positive pressure ventilation, the changes in CVP during the respiratory cycle are reversed: it rises with inspiration and decreases with expiration. In these patients, take readings near the end of expiration. Thus, during both normal and mechanical ventilation, the lowest reading is a useful estimate of mean CVP.

Errors in CVP Measurement

A number of extrinsic factors may alter the accuracy of the CVP reading (Box 22.1). In addition to the position of the patient, changes in intrathoracic pressure, malposition of the tip of the catheter, obstruction of the catheter, and failure to calibrate or zero the line may all adversely impact CVP measurements. Activities that increase intrathoracic pressure, such as coughing or straining, may cause spuriously high measurements. Make sure that the patient is relaxed and...
In the late stages of pregnancy (30 to 42 weeks), CVP is increased intrathoracic pressure (ventilator, straining, coughing) may result in inaccurate measurements that may be interpreted as a falsely elevated CVP measurement will be recorded. Recognition of a characteristic right ventricular pressure waveform on the patient's monitor should hopefully preclude this error. Such fluctuations may also occasionally be seen in appropriately positioned CVP lines when significant tricuspid regurgitation or atrioventricular dissociation (a cannon “a” wave) is present. Inaccurate low venous pressure readings are seen when a valve-like obstruction at the tip of the catheter occurs as a result of either clot formation or contact against a vein wall. Wave damping secondary to air bubbles in the transducer tubing also leads to faulty readings. Using poorly zeroed lines may result in inaccurate measurements that may be interpreted as a change in the patient's status when none has actually taken place. The transducer should be zeroed to the same level for every measurement.

**Interpretation of CVP Measurement**

Normal CVP values are as follows:

- Low: <6 cm H₂O.
- Normal: 6 to 12 cm H₂O.
- High: >12 cm H₂O.

In the late stages of pregnancy (30 to 42 weeks), CVP is physiologically elevated, and normal readings are 5 to 8 cm H₂O higher. A CVP reading of less than 6 cm H₂O is consistent with low right atrial pressure and reflects a decrease in the return of blood volume to the right heart. This may indicate that the patient requires additional fluid or blood. A low CVP reading is also obtained when vasomotor tone is decreased, as with sepsis, spinal cord injury, or other forms of sympathetic interruption.

A CVP reading falling within a normal range must be viewed in relation to the clinical scenario. A reading higher than 12 cm H₂O indicates that the heart is not effectively circulating the volume presented to it. This may occur in a normovolemic patient with underlying cardiac disease, such as left ventricular hypertrophy (with associated poor ventricular compliance), or in a volume-overloaded patient with a normal heart. High CVP can also be related to variables other than pump failure, such as pericardial tamponade, restrictive pericarditis, pulmonary stenosis, tricuspid regurgitation, pulmonary hypertension, and pulmonary embolism.

Changes in blood volume, vessel tone, and cardiac function may occur alone or in combination with one another; therefore it is possible to have a normal or elevated CVP in the presence of normovolemia, hypovolemia, and hypervolemia. Interpret the specific CVP values with respect to the entire clinical picture. The changes in CVP during an infusion are more important than the initial reading.

**Fluid Challenge**

Monitoring CVP may be helpful as a practical guide to fluid therapy. Serial CVP measurements provide a fairly reliable indication of the capability of the right side of the heart to accept an additional fluid load. Although PCWP is a more sensitive index of left heart fluid needs (and in some clinical situations measurement of PCWP is essential), serial measurement of CVP can nonetheless provide useful information.

A fluid challenge can help assess both volume deficits and pump failure. Although a fluid challenge can be used with either PCWP or CVP monitoring, only the fluid challenge for CVP monitoring is discussed here. Slight variations in the methodology of fluid challenge are reported in the literature. Generally, fluid boluses of 250 to 500 mL of crystalloid are administered sequentially and CVP is measured 10 minutes after each bolus. Repeat the fluid challenge until measurements indicate that adequate volume expansion has occurred. Discontinue the fluid challenge as soon as hemodynamic signs of shock are reversed or signs of cardiac incompetence are evident.

**Cardiac Tamponade**

In cardiac tamponade, pericardial pressure rises to equal right ventricular end-diastolic pressure. The pericardial pressure encountered in pericardial tamponade characteristically produces an elevated CVP. The degree of elevation in CVP is variable, and one must interpret measurements cautiously; CVP readings in the range of 16 to 18 cm H₂O are typically seen with acute tamponade, but elevations of up to 30 cm H₂O may be encountered. The precise CVP reading is often lower than one might intuitively expect, and it is not uncommon to encounter tamponade with a CVP of 10 to 12 cm H₂O. A normal or even low CVP reading may be seen if the tamponade is associated with significant hypovolemia. An excessive rise in CVP after fluid challenge may be more important than a single reading in the diagnosis of pericardial tamponade. Additionally, physician-operated point-of-care ultrasound has proven to be exceptionally sensitive for visualizing tamponade physiology when pericardial pressures have begun to impair right ventricular filling.

Excessive straining, positive pressure ventilation, agitation, inflation of pneumatic antishock garments, and tension...
pneumothorax may all increase intrathoracic pressure, produce a high CVP reading, and erroneously suggest the diagnosis of pericardial tamponade. Increases in vascular tone, as seen with the use of dopamine or other vasoppressors, may also elevate CVP and thus mimic tamponade and complicate estimations of volume.

**COMPLICATIONS**

The medical literature is replete with reports of CVC complications. Understanding the pathophysiology surrounding CVC complications helps clinicians anticipate, recognize, and manage complications should they arise and better educate patients and their families during the informed consent process.

More than 15% of patients who receive CVCs experience some type of complication, and complications occur despite pristine technique. This percentage is not surprising in view of the close proximity of vital structures, the complexity of patients’ medical conditions, and the exigent circumstances under which many of these procedures are often performed. The number of complications increases, especially thrombosis and line infection, with longer durations of indwelling catheters and increasing severity of illness. Common complications and the different approaches are summarized in Box 22.2 and can generally be categorized as mechanical, infectious, and thrombotic. Key complications and injuries by approach are discussed in the following sections.

The number of lumens does not directly affect the rate of catheter-related complications. One 3-year retrospective review of all central catheters placed in the ED (supraclavicular SV, IJ, and femoral lines) reported a mechanical complication rate of 3.4%, or 22 of 643 lines placed. Complications were defined as pneumothorax, hematoma, line misplacement, hemothorax, or any issue with the CVC (excluding infection or thrombosis) that required an inpatient consultation. In general, failure and complication rates increase as the number of percutaneous punctures increases. Historically, operator skill and experience have reliably predicted complication or success rates. It has previously been reported that clinicians who have placed more than 50 CVCs have less than half the complication rates of those who have fewer than 50 attempts.

Published complication rates vary in the literature and can now be classified according to whether ultrasound guidance was used during the procedure (Table 22.4). Studies have demonstrated that ultrasound-guided CVC placement techniques have improved success rates, reduced complication rates, and decreased time needed to perform the procedure.

---

**BOX 22.2 Complications of Central Venous Access**

<table>
<thead>
<tr>
<th>General</th>
<th>Infectious</th>
<th>SV and IJ Approaches</th>
<th>Neurologic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanical</td>
<td>Arterial puncture</td>
<td>Pulmonary Pneumothorax</td>
<td>Phrenic nerve injury</td>
</tr>
<tr>
<td>Puncture of an adjacent artery</td>
<td>Bloodstream infection</td>
<td>Hemothorax</td>
<td>Brachial plexus injury</td>
</tr>
<tr>
<td>Hematoma formation</td>
<td>Generalized sepsis</td>
<td>Hydrothorax</td>
<td>Cerebral infarct</td>
</tr>
<tr>
<td>Air embolus</td>
<td>Septic arthritis</td>
<td>Chylothorax</td>
<td>Femoral Approach</td>
</tr>
<tr>
<td>Pneumothorax</td>
<td>Osteomyelitis</td>
<td>Neck hematoma and tracheal obstruction</td>
<td>Intraabdominal</td>
</tr>
<tr>
<td>Pericardial tamponade</td>
<td>Cellulitis at the insertion site</td>
<td>Endotracheal cuff perforation</td>
<td>Bowel perforation</td>
</tr>
<tr>
<td>Catheter embolus</td>
<td>Thrombotic</td>
<td>Tracheal perforation</td>
<td>Psoas abscess</td>
</tr>
<tr>
<td>Arteriovenous fistula</td>
<td>Pulmonary embolism</td>
<td></td>
<td>Bladder perforation</td>
</tr>
<tr>
<td>Mural thrombus formation</td>
<td>Venous thrombosis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Large-vein obstruction</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dysrhythmias</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Catheter knotting</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Catheter malposition</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

IJ, Internal jugular; SV, subclavian vein.

---

**TABLE 22.4 Frequency of Complications Without and With Ultrasound Guidance**

<table>
<thead>
<tr>
<th>COMPLICATION</th>
<th>Without Ultrasound</th>
<th>With Ultrasound</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IJ</td>
<td>SV</td>
</tr>
<tr>
<td>Arterial puncture</td>
<td>6.3–9.4%</td>
<td>3.1–4.9%</td>
</tr>
<tr>
<td>Hematoma</td>
<td>&lt;0.1–2.2%</td>
<td>1.2–2.1%</td>
</tr>
<tr>
<td>Hemothorax</td>
<td>0%</td>
<td>0.4–0.6%</td>
</tr>
<tr>
<td>Pneumothorax</td>
<td>&lt;0.1–0.2%</td>
<td>1.5–3.1%</td>
</tr>
<tr>
<td>Infection (rate per 1000 catheter-days)</td>
<td>8.6</td>
<td>4</td>
</tr>
<tr>
<td>Thrombosis (rate per 1000 catheter-days)</td>
<td>1.2–3</td>
<td>0–13</td>
</tr>
</tbody>
</table>

IJ, Internal jugular; NA, not applicable; SV, subclavian vein.

Data from References 47, 78, and 129.
As a result, ultrasound guidance for CVC placement is recommended by the US Department of Health and Human Services. Reports by the AHRQ list ultrasound guidance for central vein cannulation as one of its most highly rated safety practices.60,131

Mechanical Complications

The most commonly reported mechanical complications are arterial puncture, hematoma, and pneumothorax. Inadvertent arterial puncture and hematoma formation are usually easily recognized and controlled with simple compression. Rarely, an artery is lacerated to such an extent that bleeding is significant and operative repair is necessary. In cardiac arrest, low-flow, or shock states, arterial puncture may not be obvious, and arterial cannulation and intraarterial infusions have occurred. This can lead to the development of ischemia or thrombosis of the artery and limb. When systolic blood pressure rises, arterial pulsations become more obvious. In critically ill patients, however, this complication may escape detection for some time. It has been reported that ultrasound-guided placement of IJ CVCs decreases the rate of arterial puncture to 1.4%.50

Though poorly studied, patients with coagulopathies may experience significant bleeding from CVC placement, especially if arterial puncture or laceration has occurred. Mumtaz and coworkers cited a 3% bleeding rate in coagulopathic patients who experienced only minor bleeding that could be controlled with digital pressure.47 Although central venous access procedures may be performed safely in patients with bleeding disorders without antecedent correction of the coagulopathy, caution is strongly urged. Areas amenable to arterial compression are preferred in these patients.47

Pneumothorax occurs in up to 6% of subclavian venipunctures and can also occur with the IJ approach.43,63,112 (Fig. 22.22A). Initially, the importance of this complication was minimized, but reports of fatalities caused by tension pneumothorax, bilateral pneumothorax, and combined hemothorax followed.48 One would expect a higher incidence of pneumothorax if the procedure is performed during CPR or positive pressure ventilation. A small pneumothorax can quickly become a life-threatening tension pneumothorax with positive pressure ventilation. Treatment of a catheter-induced pneumothorax is controversial, but not all patients will require formal tube thoracostomy. Some authors have reported that many stable outpatients exhibiting a pneumothorax after insertion of a CVC can be managed successfully by observation alone (60% in one series) or catheter (pigtail/Heimlich valve) aspiration, with large tube thoracostomy being reserved for refractory cases or emergency settings.112,113 Critically ill patients or those undergoing mechanical ventilation are more likely to require invasive treatment for a catheter-induced pneumothorax.

Hemothorax may occur after laceration of the SV or subclavian artery, puncture of the pulmonary artery, or intrathoracic infusion of blood (see Fig. 22.22B). Hydrothorax occurs as a result of infusion of IV fluid into the pleural space. Hydromediastinum is also possible. These are rarely serious complications, but fatalities have been reported. Surgical repair is occasionally required. Arteriovenous fistula formation has also been reported.114 Additional pulmonary complications include tracheal and endotracheal cuff perforation.

Air embolism is a very rare, but potentially life-threatening complication of central venous cannulation. Undoubtedly, minor and clinically inconsequential amounts of air enter the venous circulation during many cannulation procedures. Maintaining constant occlusion (with the operator’s finger) on all needles that are located in central veins can minimize this occurrence. A 14-gauge needle can transmit 100 mL of air per second with a 5–cm H2O pressure difference across the needle.98 Air embolism may occur if the line is open to air during catheterization or if it subsequently becomes disconnected. The recommended treatment is to place the patient in the left lateral decubitus position to relieve air bubble occlusion of the right ventricular outflow tract.99 If this is unsuccessful, aspiration with the catheter advanced into the

Figure 22.22 Pneumothorax and hemothorax. A chest radiograph should be taken routinely to assess the position of a central venous catheter introduced via the subclavian or internal jugular route. This confirms placement of the catheter. Chest radiography can also show potential complications of the procedure. A, Large right pneumothorax after right internal jugular catheterization. The catheter is still in place (large arrow), and the absence of lung markings on the right and the pleural reflection (small arrows) are readily apparent. B, Left hydropneumothorax after left subclavian venipuncture (the catheter was removed before this radiograph). Note the straight line of fluid (air-fluid level, black arrows) and no meniscus, indicating that a pneumothorax must also be present. The edge of the partially collapsed lung is difficult to appreciate. No clinician can place central venous catheters and fail to have at least some complications that are inherent to the procedure, regardless of even flawless technique.
right ventricle has been advocated. The incidence of septic complications varies from 0% to 25%. The frequency with which infectious complications occur is directly related to the attention given to aseptic technique during insertion and aftercare of the catheter. Femoral venous catheterization carries a greater risk for infection than subclavian catheterization. Merrer and associates reported the overall infectious complication rate from femoral and subclavian catheters to be 19.8% and 4.5%, respectively. Ultrasound-guided IJ CVC placement has resulted in a decrease in the rates of CLABSI. The exact mechanism by which ultrasound-guided CVC placement results in a lower risk for infection is unclear; it may be related to a reduced number of skin punctures. Organisms most commonly recovered from colonized femoral catheters are coagulase-negative staphylococci, Enterobacteriaceae, Enterococcus species, and Pseudomonas aeruginosa. CVCs cause an estimated 80,000 CLABSI and are implicated in up to 28,000 deaths per year in patients in the intensive care unit. The average cost has been estimated at $2.3 billion annually. The Centers for Disease Control and Prevention has recommended that central line bundling policies should be implemented to significantly decrease the incidence of CLABSI. This bundling policy includes five evidence-based interventions: (1) hand washing, (2) maximal barrier precautions, (3) chlorhexidine skin antisepsis, (4) optimal catheter site selection with avoidance of the femoral vein if possible, and (5) daily review of the necessity for the line and prompt removal of unnecessary lines.

**Thrombotic Complications**

Thrombosis and thrombophlebitis are significant risks associated with placement of a CVC. The risk for catheter-related thrombosis is directly related to the site of access. In one trial, catheter-related thrombosis was reported in up to 21.5% of patients with femoral CVCs and in 1.9% of patients with SV CVCs. For SV and IJ CVCs, it is important to determine that the tip of the catheter rests in the SVC, especially during the infusion of irritating or hypertonic solutions. The most common result of kinking is poor flow of IV fluids, although rare complications as severe as SVC obstruction syndrome have been seen. Thoracic duct laceration is a frequently discussed complication of left-sided subclavian venipuncture; however, it is extremely uncommon and has been reported only as a complication of IJ vein, not SV, cannulation.

Neurologic complications are extremely rare and presumably caused by direct trauma from the needle during venipuncture. Brachial plexus palsy and phrenic nerve injury with paralysis of the hemidiaphragm have been reported. Infusing hypertonic medications into the IJ vein via a malpositioned catheter may result in a variety of neurologic complications from retrograde perfusion of intracranial vessels. Again, this complication can be easily avoided by inspecting a postprocedure x-ray to confirm proper placement prior to utilization of a newly placed catheter.

**Infectious Complications**

Infectious complications include local cellulitis, thrombophlebitis, generalized septicemia, osteomyelitis, and septic arthritis.
IJ Approach

Many complications of IJ vein cannulation are similar to those of SV access. The incidence of complications appears to be higher with use of the left IJ vein than with the right.\textsuperscript{16} One common complication unique to the IJ approach is a localized hematoma in the neck.\textsuperscript{145} With the IJ approach, pressure can easily be maintained on the area of swelling, and most hematomas will resolve spontaneously. If puncture of the carotid artery is recognized and treated with compression, it rarely causes significant morbidity in the absence of marked atherosclerotic disease, although arteriovenous fistulas may occur after IJ vein puncture.\textsuperscript{134} Several neurologic complications unique to the IJ site of venipuncture have also been reported as a result of hematomas or direct injury. Such complications include damage to the phrenic nerves, iatrogenic Horner’s syndrome, trauma to the brachial plexus, and even passage of a catheter into the thecal space of the spinal canal.\textsuperscript{145} If the carotid artery is punctured, one may again attempt IJ vein or SV cannulation on the same side after appropriate, prolonged (15- to 20-minute) compression. The IJ vein valve is frequently damaged when cannulated, which often results in incompetence of the valve. The clinical significance of this, if any, is unknown.\textsuperscript{146}

Femoral Approach

Because vital structures in the neck and chest are not at risk, complications of femoral vein cannulation are generally less severe than those of other routes for central venous access. The most common immediate complications involve bleeding from damage to either the femoral artery or the femoral vein (Fig. 22.23). This can usually be managed with 10 to 15 minutes of direct pressure. Extra care should be taken in anticoagulated patients or after the administration of thrombolytic agents. In extreme cases when hemostasis cannot be achieved through direct pressure, a vascular surgeon should be consulted.

The peritoneum can also be violated with resultant perforation of the bowel. Bowel penetration is especially likely if the patient has a femoral hernia. Injury to the bowel is usually minimal and unlikely to require specific treatment. Nonetheless, the potential bacterial contamination of the femoral puncture site can pose a significant problem. Aspiration of air during placement of a femoral line necessitates removal of the catheter and reinsertion at another site. Other complications include muscular abscesses, infection of the hip joint, damage to the femoral nerve, and puncture of the bladder. Risk for these outcomes can be mitigated by strict aseptic technique, thorough assessment of landmarks, careful control of the needle’s depth, and the use of bedside ultrasound.

Two more complications merit special mention. The first is the increased risk for catheter infection. Presumably caused by anatomic association with the anogenital region, many studies have found that femoral lines become infected at significantly higher rates than IJ vein or supraclavicular SV lines do.\textsuperscript{64,65,68,135} Of note, some studies have failed to find a statistical difference, and it is unclear how much of the effect is due to the actual location of the line versus how it is placed and managed. The second is the incidence of deep vein thrombosis that is also increased in lines placed via the femoral route, which has been shown in the majority of studies,\textsuperscript{133,147} although the clinical significance of these clots has not been definitively addressed.

Figure 22.23 A femoral vein catheter is more prone to deep vein thrombosis and infection than a subclavian or internal jugular line, but it is a standard access route in the emergency department. Strict attention to sterile technique and limiting use to a few days will negate most of the negatives of this approach. A, Significant hemorrhage can occur after puncture of the femoral artery, but this area is readily compressed. The femoral route may be the approach of choice in a patient with an inadvertently placed arterial catheter who requires a central line. B, Bleeding from an inadvertently placed arterial catheter that was removed without adequate pressure in an anticoagulated patient.

TRAINING AND SIMULATION

CVC placement and ultrasound guidance techniques have a relatively steep learning curve. Simulation is recommended by the AHRQ to teach these techniques.\textsuperscript{60,141} Simulation training is independently associated with higher rates of correct needle insertion on the first attempt, as well as with higher successful CVC placement rates.\textsuperscript{131,146,148} There are many simulation models that can be used. Kendall and Faragher described a phantom model as an easy, inexpensive method for ultrasound-guided CVC placement training.\textsuperscript{150}

REFERENCES ARE AVAILABLE AT www.expertconsult.com


Management of critically ill or injured patients requires immediate and adequate vascular access, especially during trauma resuscitation, when rapid infusion of crystalloid or blood products may be necessary. Venous cutdown, a time-honored surgical technique, has largely been replaced by alternative methods of obtaining venous access, including intraosseous lines, the Seldinger technique, and ultrasound-guided central and peripheral venous cannulation. Nonetheless, venous cutdown still has a role as an emergency method of achieving vascular access when other techniques and equipment are unavailable, particularly in settings outside the United States.

First described by Keeley in 1940 and Kirkham in 1945, venous cutdown offered an alternative to venipuncture in patients with shock. Though no longer taught as a mandatory procedure in the Advanced Trauma Life Support course, venous cutdown is considered optional and continues to be taught at the discretion of the instructor. Realistically, percutaneous vascular access may be infeasible in a pulseless, hypovolemic, or anatomically scarred patient. With a thorough understanding of the anatomy, the procedure, and its potential complications, this mechanically simple procedure can be performed quickly and effectively.

**INDICATIONS**

Venous cutdown may be used as an alternative to venipuncture for critically ill patients in need of vascular access when less invasive options have been exhausted or are not available. Patients with severe shock, asystole, or pulseless electrical activity will lack palpable femoral pulses, thus making percutaneous femoral vein catheterization more difficult. Surface landmarks may be obscured and veins may be unusable in intravenous (IV) drug users, the extensively injured, or severely burned patients. Attempts at percutaneous venous cannulation may be complicated or even impossible in such patients. Venous cutdown (Video 23.1) and intraosseous routes (see Chapter 25) are both viable options in such scenarios.

**Children**

Venipuncture in small children poses a challenge in even the healthiest of patients, let alone those in extremis whose veins may be poorly visualized. Central vein catheterization, intraosseous line placement, or venous cutdown should be considered as an alternative means of emergency vascular access when other peripheral sites have been exhausted. The distal saphenous vein at the ankle is often recommended for venous cutdown in children given its large diameter and anatomic predictability at this location.

---

**Venous Cutdown**

**Indications**

As an alternative to venipuncture in critically ill patients in need of vascular access and in whom venipuncture may be difficult:
- Shock
- Asystole or pulseless electrical activity
- Sclerosed veins of intravenous drug abusers
- Extensive burn or other injury
- Small children

**Contraindications**

Absolute:
- When less invasive options exist for venous access
- Major blunt or penetrating trauma at proposed cutdown site

Relative:
- Overlying soft tissue infection
- Bleeding diathesis
- Immunocompromise
- Extremity injuries proximal to the site

**Complications**

- Transection of the vein
- Transection of the artery
- Bleeding
- Hematoma
- Phlebitis
- Sepsis
- Thrombus formation
- Injury to surrounding structures

**Equipment**

- Scalpel with No. 11 blade
- Curved hemostat
- Iris scissors
- 0-0 silk sutures
- Plastic venous dilator
- Large-bore intravenous catheter
- Intravenous tubing
- Tape

**Review Box 23.1** Venous cutdown: indications, contraindications, complications, and equipment.
Hypovolemic Shock

Initially popularized during the Vietnam War for rapid transfusion, venous cutdown has since been used for resuscitation of patients with profound hypovolemia. The flow rate of saline through a standard IV extension set cut to a length of 28 cm (12 inches) and inserted directly into the vein is 15% to 30% greater than through a 5-cm, 14-gauge catheter. This difference is larger if pressure is applied to the system. Moreover, the improvement in flow rate through large-bore lines is greater for blood than for crystalloid solutions because the viscous characteristics of blood impede its passage through small-bore tubing. A unit of blood can be transfused in as little as 3 minutes through IV extension tubing inserted directly into the vein. Consequently, large-bore lines placed by venous cutdown are an excellent mechanism for the treatment of severe hypovolemia.

CONTRAINDICATIONS

Venous cutdown is contraindicated when less invasive alternatives exist and when performing the procedure would cause excessive delay. Highly skilled clinicians may perform a cutdown in less than 60 seconds. However, multiple studies by Westfall, Rhee, Iserson, and their colleagues have indicated that on average, the procedure takes at least 5 to 6 minutes to complete. Use of the modified Seldinger technique described both by Shockley and Butzier, and by Klofas, has been shown to decrease that time by 22%. In general, the use of percutaneously inserted central venous catheters in either the subclavian, internal jugular, or femoral vein is preferable to a cutdown.

Absolute contraindications include major blunt or penetrating trauma involving the extremity on which the procedure is to be performed. Relative contraindications include vascular injury proximal to the cutdown site, overlying soft tissue infection, coagulopathies, compromised host defense mechanisms, and impaired wound healing. Other considerations include any previous saphenous vein harvest for coronary artery bypass or other vascular surgery proximal to the anticipated cutdown site. The indications for venous cutdown should be weighed against the potential complications.

ANATOMY AND SELECTION OF THE SITE

Knowledge of the relevant anatomy is imperative for success. Veins in both the upper and lower extremities may be used. The size and accessibility of the target vessel, along with the clinician’s experience and training, are the principal factors in selection of the site. There are four primary locations at which venous cutdown is performed: the great saphenous vein distally at the ankle and proximally at the thigh, the basilic vein above the elbow, and the cephalic vein below it. Brachial vein cutdown is no longer recommended as an emergency venous access route because of its time-consuming dissection and risk for neurovascular injury. The anatomy of individual vessels and their relative merits as cutdown sites are described in the following sections.

The Great Saphenous Vein

The great saphenous vein is the longest vein in the body, and it runs subcutaneously throughout much of its course (Fig. 23.1). It is most easily accessible at the ankle but may also be cannulated below the knee and below the femoral triangle. The great saphenous vein begins just anterior to the medial malleolus where it is a continuation of the medial marginal vein of the foot. The vein crosses 1 cm anterior to the medial malleolus and, together with the saphenous nerve, ascends along the anteromedial aspect of the leg. The saphenous nerve at this level is of relatively little clinical significance, for if transected, it causes sensory loss in only a small area along the medial aspect of the foot. At the ankle, the vessel can be exposed with minimal blunt dissection. The vein’s superficial, predictable, and isolated location has made the distal saphenous vein the traditional pediatric cutdown site. The saphenous nerve, ascending along the anteromedial aspect of the leg, has been shown to decrease that time by 22%. In general, the use of percutaneously inserted central venous catheters in either the subclavian, internal jugular, or femoral vein is preferable to a cutdown.

At the knee, the saphenous vein lies superficially and medially. A cutdown performed 1 to 4 cm below the knee and immediately posterior to the tibia has been described in the pediatric literature. This site is seldom used, however, because of its many disadvantages, including kinking of the line as the knee is flexed, and risk for injury to the saphenous branch of the genicular artery and the saphenous nerve. Note, the great saphenous vein is duplicated in the calf in 25% of the population and may be present on exploration. In the thigh, the saphenous vein begins on the medial aspect of the knee and crosses anterolaterally as it ascends toward the femoral triangle. Approximately 4 cm below the inguinal ligament and 3 cm lateral to the pubic tubercle, the saphenous
the proximal antecubital fossa. It is this predictability in anatomic location that makes the basilic vein an ideal site for venous cutdown in the upper extremity.

A more proximal cutdown site had previously been recommended to avoid the network of interconnecting veins at the level of the antecubital fossa. However, a closer association between the basilic vein and the medial cutaneous nerve in this segment may result in transection of the nerve and subsequent sensory loss on the ulnar side of the forearm.

The Cephalic Vein

This cephalic vein begins on the radial aspect of the wrist, crosses anteromedially, and ascends toward the antecubital fossa. In the forearm, it lies in close association with the lateral cutaneous nerve, which supplies sensory innervation to the radial aspect of the forearm (see Fig. 23.3). In the antecubital fossa, it lies subcutaneously and just lateral to the midline. It is at this level where the median cubital vein connects to the cephalic and basilic veins. The cephalic vein then ascends in the upper part of the arm over the lateral aspect of the biceps muscle and through the deltopectoral groove. Just below the clavicle, it pierces the clavipectoral fascia, becomes a deep structure, and enters the axillary vein.20

The basilic vein is a preferred site for venous cutdown in the upper extremity because of its predictable anatomic location. The size of this vein enables it to be located easily, even in hypotensive or hypovolemic patients. Superficially at this level there are no important associated structures, but the brachial artery and the median nerve are found deep to the basilic vein.

Veins of the dorsal venous network of the hand unite to form the cephalic and basilic veins, which travel along the radial and ulnar sides of the forearm, respectively (Fig. 23.3). At the midforearm level, the basilic vein crosses anterolaterally and then courses ventrally at the medial epicondyle. The medial cubital vein crosses over from the radial side of the arm to join the basilic vein just above the medial epicondyle. The basilic vein then continues proximally, where it occupies a superficial position between the biceps and pronator teres muscle. In this segment it lies in close proximity to the medial cutaneous nerve, which supplies sensation to the ulnar side of the forearm. At approximately midway in the upper part of the arm, the basilic vein perforates the deep fascia, where it joins the brachial vein and continues on into the axillary vein.20

The basilic vein is consistently found at the antecubital fossa, 2 cm above and 2 to 3 cm lateral to the medial epicondyle, on the anterior surface of the upper part of the arm. It is exposed through a transverse incision on the medial aspect of the proximal antecubital fossa. It is this predictability in anatomic location that makes the basilic vein an ideal site for venous cutdown in the upper extremity.

A more proximal cutdown site had previously been recommended to avoid the network of interconnecting veins at the level of the antecubital fossa. However, a closer association between the basilic vein and the medial cutaneous nerve in this segment may result in transection of the nerve and subsequent sensory loss on the ulnar side of the forearm.

The Cephalic Vein

This cephalic vein begins on the radial aspect of the wrist, crosses anteromedially, and ascends toward the antecubital fossa. In the forearm, it lies in close association with the lateral cutaneous nerve, which supplies sensory innervation to the radial aspect of the forearm (see Fig. 23.3). In the antecubital fossa, it lies subcutaneously and just lateral to the midline. It is at this level where the median cubital vein connects to the cephalic and basilic veins. The cephalic vein then ascends in the upper part of the arm over the lateral aspect of the biceps muscle and through the deltopectoral groove. Just below the clavicle, it pierces the clavipectoral fascia, becomes a deep structure, and enters the axillary vein.20

Venous cutdown is easily performed on the cephalic vein because of its large diameter and superficial location. In the forearm, it is important to avoid the lateral cutaneous nerve. A preferred location is in the antecubital fossa at the distal flexor crease. Cutdown on the cephalic vein at the wrist has also been reported, but the thin skin overlying the vein at this level usually permits simple percutaneous cannulation when the vein is available for cannulation.21 The cephalic vein may also be entered in the deltopectoral groove. However, the
slightly deeper position and possible interference with the performance of other procedures make this approach more difficult.

**EQUIPMENT**

The material required to perform a formal venous cutdown is shown in Review Box 23.1. Perhaps the most important piece of equipment and the most difficult to find is the vein dilator/lifter, a plastic device with a 90-degree angle that is used to facilitate entrance of a catheter into the cut vein. For pediatric patients, use a warming table, or radiant warmer, and a padded extremity board.

Choose a catheter based on the desired function of the venous line. When central venous pressure (CVP) needs to be monitored, choose a catheter long enough to reach the superior vena cava. Approximate this distance by aligning the catheter over the chest with the tip at the level of the manubriosternal junction. The average distance from the antecubital fossa to the superior vena cava is 54 cm in adult men. Lumen size is relatively unimportant when the line is inserted for monitoring CVP or to infuse drugs, but it is a critical factor in the treatment of hypovolemia. Short, large-bore catheters are preferred when fluid must be delivered rapidly. Silastic catheters, IV plastic tubing, or 5- or 8-Fr pediatric feeding tubes may be used as infusion catheters in older children and adults.

Tables 23.1 to 23.3 list the flow rates of various fluids through some commonly used catheter systems. It is essential to know the relative flow rates if maximal benefit is to be obtained from the time spent performing the cutdown. Excellent flow rates can be achieved by threading IV tubing directly into the vein or by using a 5-cm, 10-gauge IV catheter. Cut sterile tubing to the appropriate length and leave a slight bevel on the end to facilitate cannulation of the opened vein.11,25

In conscious patients, apply a local anesthetic before the procedure. Make a skin incision perpendicular to the course of the vein. A longitudinal incision, even though it decreases the risk of transecting neurovascular structures, may not provide sufficient exposure. Incise the skin through all its layers until subcutaneous fat bulges through the incision. Very carefully dissect the subcutaneous tissues bluntly by spreading them gently, with a curved hemostat parallel to the course of the vein and with the tips pointed downward. This is the most difficult and delicate portion of the procedure and may damage the vein and render it unable to be cannulated. Bleeding, however, is usually minimal unless the vein is nicked. Use a tissue spreader or a self-retaining retractor, if needed, to provide a wider field. Isolate the vein from the adjacent tissue and mobilize it for 1 to 3 cm.

For the standard venous cutdown technique, after mobilizing the vein, use a hemostat to pass proximal and distal silk ties under the vein for stabilization (Fig. 23.5).

Tie the distal ligature after initial placement, but leave the ends long for maneuvering the vein. As an option, use traction on the distal suture to control the vein, but do not tie it off. If the distal suture is tied, the vein will be sacrificed for future use. Leave the proximal ligature untied to maneuver the vein for insertion of the catheter or tubing and control of backbleeding (by lifting the sutures). Elevate the vein with a hemostat and stretch it flat. This provides good visualization, controls the vessel, and limits bleeding when the vessel is incised. Alternatively, place gentle traction on the proximal tie to control oozing around the puncture site. Using a No. 11 scalpel blade or a pair of iris scissors, incise through one third to one half the diameter of the vein at a 45-degree angle. If the incision is too small, the catheter may pass into a false channel in the adventitia. Conversely, if the incision is too large, the vein may tear completely and retract from the field buried within tissue.26 If desired, make a longitudinal incision in the vein to avoid transecting the vessel, but realize that this technique makes it more difficult to identify the lumen. Be aware that some bleeding will normally occur after the vein has merely been nicked on the surface. To perform a mini-cutdown at this point, puncture the vein with an IV catheter and introducer needle and do not make an incision in the vein.

Before introducing the cannula into the vein, make a bevel in the cannula at a 45-degree angle. This is unnecessary if the cannula has a tapered tip. Make the bevel short, but be careful to not make it sharp to avoid piercing the posterior wall or otherwise damaging the vein. If using the rounded tip of a feeding tube, it may be more difficult to introduce but can be advanced less traumatically. If using an IV cannula, introduce it directly through the skin incision or through a separate stab incision.

Threading the catheter into the vein is often the most difficult and time-consuming portion of the procedure. Difficulty threading has several causes. The lumen may have been incorrectly identified, or a false passage into the adventitia may have been created. This can be difficult to recognize because a catheter can easily pass between layers of the vessel wall and never reach the lumen of the vein. Other causes include penetrating the posterior vessel wall, getting stuck in a valve, or using a catheter that is too large to cannulate the vein.

Use of a plastic venous dilator can help identify and elevate the vessel lumen. Thread the small, pointed tip of the dilator into the vein to expose the lumen before advancing the tip of

---

**Table 23.1** Comparative Average Flow Rates (mL/min) for Tap Water

**Table 23.2** Comparative Average Flow Rates (mL/min, 200-mm Hg Pressure) for Red Blood Cells

**Table 23.3** Comparative Average Flow Rates (mL/min)

---

**TECHNIQUE**

The technique of venous cutdown is essentially the same regardless of the vessel cannulated (Fig. 23.4). Prepare the skin around the incisional area with an antiseptic solution and then cover it with sterile drapes. Place a tourniquet proximal to the cutdown site to help visualize the vein. For children, immobilize the lower part of the leg or elbow (depending on the cutdown site) on a padded board before beginning the procedure.
**TABLE 23.1** Comparative Average Flow Rates (mL/min) for Tap Water

<table>
<thead>
<tr>
<th>CATHETER</th>
<th>PRESSURE, 200 mm Hg (95% CI)</th>
<th>GRAVITY (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Central Venous Catheters</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>USCI 9-Fr introducer</td>
<td>566 (±16)</td>
<td>247 (±2)</td>
</tr>
<tr>
<td>ID, 0.117 inch; length, 5.5 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>USCI 8-Fr introducer</td>
<td>540²</td>
<td>243 (±5)</td>
</tr>
<tr>
<td>ID, 0.104 inch; length, 5.5 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deseret Angiocath</td>
<td>341 (±6)</td>
<td>157 (±6)</td>
</tr>
<tr>
<td>Gauge, 14; length, 5.25 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deseret Angiocath</td>
<td>195 (±4)</td>
<td>91 (±2)</td>
</tr>
<tr>
<td>Gauge, 16; length, 5.25 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deseret subclavian jugular catheter</td>
<td>142 (±4)</td>
<td>54 (±3)</td>
</tr>
<tr>
<td>Gauge, 16; length, 12 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Peripheral Venous Catheters</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intravenous tubing</td>
<td>500 (±21)</td>
<td>222 (±4)</td>
</tr>
<tr>
<td>ID, 0.12 inch; length, 12 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Argyle Medicut</td>
<td>484 (±8)</td>
<td>194 (±5)</td>
</tr>
<tr>
<td>Gauge, 14; length, 2 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deseret Angiocath</td>
<td>405 (±2)</td>
<td>173 (±4)</td>
</tr>
<tr>
<td>Gauge, 14; length, 2 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vicra Quick-Cath</td>
<td>—</td>
<td>167 (±1)</td>
</tr>
<tr>
<td>Gauge, 14; length, 2.25 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Argyle Medicut</td>
<td>353 (±4)</td>
<td>151 (±3)</td>
</tr>
<tr>
<td>Gauge, 16; length, 2 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deseret Angiocath</td>
<td>231 (±1)</td>
<td>108 (±1)</td>
</tr>
<tr>
<td>Gauge, 16; length, 2 inches</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vicra Quick-Cath</td>
<td>—</td>
<td>108 (±1)</td>
</tr>
<tr>
<td>Gauge, 16; length, 2 inches</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

¹Mean of three trials with a hydrostatic pressure head of 1 m.
²Ninety-five percent confidence interval not calculated because all three trials resulted in 11.1 seconds for 100-mL flow.

CI, Confidence interval; ID, internal diameter.

### TABLE 23.2 Comparative Average Flow Rates (mL/min, 200–mm Hg Pressure) for Red Blood Cells

<table>
<thead>
<tr>
<th>CATHETER</th>
<th>DILUTED PRBCs, Hct 45% (95% CI)</th>
<th>DILUTED PRBCs, Hct 45%, THROUGH A BLOOD WARMER (95% CI)</th>
<th>PRBCs, Hct 65% (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Central Venous Catheters</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>USCI 9-Fr introducer ID, 0.117 inch; length, 5.5 inches</td>
<td>343 (±21)</td>
<td>218 (±26)</td>
<td>124 (±2)</td>
</tr>
<tr>
<td>USCI 8-Fr introducer ID, 0.104 inch; length, 5.5 inches</td>
<td>324 (±23)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Deseret Angiocath Gauge, 14; length, 5.25 inches</td>
<td>210 (±7)</td>
<td>171 (±9)</td>
<td>63 (±6)</td>
</tr>
<tr>
<td>Deseret Angiocath Gauge, 16; length, 5.25 inches</td>
<td>125 (±4)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td><strong>Peripheral Venous Catheters</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intravenous extension tubing ID, 0.12 inch; length, 12 inches</td>
<td>312 (±1)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Argyle Medicut Gauge, 14; length, 2 inches</td>
<td>287 (±21)</td>
<td>192 (±15)</td>
<td>96 (±6)</td>
</tr>
<tr>
<td>Deseret Angiocath Gauge, 14; length, 2 inches</td>
<td>257 (±11)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Argyle Medicut Gauge, 16; length, 2 inches</td>
<td>220 (±5)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Deseret Angiocath Gauge, 16; length, 2 inches</td>
<td>158 (±14)</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

CI, Confidence interval; Hct, hematocrit; ID, internal diameter; PRBCs, packed red blood cells.

### TABLE 23.3 Comparative Average Flow Rates (mL/min)

<table>
<thead>
<tr>
<th>CATHETER</th>
<th>TAP WATER AT 200 mm Hg</th>
<th>DILUTED PRBCs AT 200 mm Hg</th>
<th>TAP WATER, GRAVITY</th>
<th>DILUTED PRBCs, BLOOD WARMER, 200 mm Hg</th>
<th>PRBCs AT 200 mm Hg</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Central Venous Catheters</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>USCI 9-Fr introducer ID, 0.117 inch; length, 5.5 inches</td>
<td>566 (±16)</td>
<td>343 (±21)</td>
<td>247 (±2)</td>
<td>218 (±26)</td>
<td>124 (±2)</td>
</tr>
<tr>
<td>USCI 8-Fr introducer ID, 0.104 inch; length, 5.5 inches</td>
<td>540a</td>
<td>324 (±23)</td>
<td>243 (±5)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Deseret Angiocath Gauge, 14; length, 5.25 inches</td>
<td>341 (±6)</td>
<td>210 (±7)</td>
<td>157 (±6)</td>
<td>171 (±9)</td>
<td>63 (±6)</td>
</tr>
<tr>
<td><strong>Peripheral Venous Catheters</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intravenous extension tubing ID, 0.12 inch; length, 12 inches</td>
<td>500 (±21)</td>
<td>312 (±1)</td>
<td>222 (±4)</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Argyle Medicut Gauge, 14; length, 2 inches</td>
<td>484 (±8)</td>
<td>287 (±21)</td>
<td>194 (±5)</td>
<td>192 (±15)</td>
<td>96 (±6)</td>
</tr>
<tr>
<td>Argyle Medicut Gauge, 16; length, 2 inches</td>
<td>353 (±4)</td>
<td>220 (±5)</td>
<td>151 (±3)</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

CI, Confidence interval; ID, internal diameter; PRBCs, packed red blood cells.

---

9 Ninety-five percent confidence interval not calculated because all three trials resulted in 11.1 seconds for 100-mL flow.
After applying a tourniquet and cleansing the skin, make a skin incision perpendicular to the course of the vein.

Bluntly dissect, isolate, and mobilize the vein.

Use a hemostat to isolate the vein and pass silk ties under it, proximal and distal to the proposed cannulation site.

Incise the vein while retracting the proximal ligature. Lift the proximal untied suture to control backbleeding.

Use the venous dilator to lift the flap and then advance the catheter into the vein. Attach intravenous tubing to the catheter.

Tie the proximal silk suture around the vein and catheter. Remove the proximal suture and suture the skin. If the distal suture has not been tied, remove it before suturing the skin.

Tie the distal suture only. Optionally, you can apply traction on the distal untied suture to control bleeding and remove it after cannulation.

Figure 23.4 Venous cutdown. (Adapted from Custalow CB: Color atlas of emergency department procedures, Philadelphia, 2005, Saunders.)
the catheter. Alternatively, bend a sterile 20-gauge needle at a 90-degree angle to serve as a venous dilator or elevator. A vein dilator is useful for very small veins, such as in pediatric cutdowns, but is generally unnecessary in adults. To thread large catheters in adults, grasp the proximal edge of the vessel with small forceps or a mosquito hemostat. Apply countertraction and advance the catheter (Fig. 23.6). Never force a catheter that will not advance easily.

Once the catheter is advanced into the lumen, backbleed any air from the cannula and connect it to IV tubing. Tie the proximal ligature around both the vessel and the cannula (Fig. 23.7). If the distal suture has not been tied, remove it. If it has been tied, cut the ends of the suture near the knot. Remove the tourniquet, affix the catheter to the skin, and close the incision. Apply antibiotic ointment at the point where the catheter passes through the skin, and dress the wound. In an emergency situation one may delay skin closure if necessary and simply wrap the wound with sterile dressing. Loop the IV tubing under the outer layers of the dressing to minimize the risk of pulling the cannula out if the external IV line is inadvertently tugged.

**Mini-Cutdown**

The mini-cutdown is an alternative method designed to preserve the vein and bypass the time-consuming step of inserting a catheter into the vein.²⁷ It is preferred if time is limited. Basically, a deep vein is cannulated through an incision under direct vision with a standard peripheral IV catheter. Use a skin incision and blunt dissection to locate the vessel. Once identified, puncture the vein under direct vision with a percutaneous venous catheter. Introduce the needle through either the skin incision or a separate stab incision. If an over-the-needle device (e.g., Angiocath [Becton Dickinson, Franklin Lakes, NJ], Medicut [Medtronic, Minneapolis, MN]) is used, withdraw the needle and discard it. With a through-the-needle device,
Modified Cutdown Technique

Shockley and Butzier described a further modification in which a guidewire, dilator, and sheath system is inserted after standard cutdown and venotomy. To perform this modification set up the guidewire, dilator, and sheath system before making the skin incision. Once the vein has been incised, insert the end of the guidewire followed by the dilator and sheath. Remove the wire and dilator while leaving the sheath. Ligatures are not usually needed with this technique. These authors found that when performed by novices, this technique saved more than 2 minutes in comparison to the standard technique. Moreover, in the event of a transection, there was an increased vein salvage rate. Klofas used a similar technique at the distal saphenous vein. He also developed a model for teaching the modified technique with wood, gauze, cast padding, and tape.

To remove catheters inserted by cutdown, cut the skin stitches holding the catheter in place and then withdraw the catheter. Control backbleeding from the proximal venous end by applying a simple pressure dressing.

COMPLICATIONS

Complications of venous cutdown include: local hematoma, infection, sepsis, phlebitis, embolization, wound dehiscence, and injury to associated structures. An indirect but significant complication is deterioration of an unstable patient during a time-consuming attempt at cutdown. Documentation of complications and their frequency in the literature has been sparse. Bogen reported a 15% complication rate in 234 cases. Infection and phlebitis each occurred at a rate of 4%. Infectious complications may result from the introduction of pathogens during placement of the line, transcutaneous invasion along the course of the cannula, or deposition of blood-borne organisms on the tip of the catheter. A clear correlation exists between the incidence of infectious complications and the length of time that a catheter is left in place. Moran and colleagues found that the infection rate rose from 50% to 78% when a catheter was left in place for more than 48 hours. Druskin and Siegel, studying a mixed population of patients who had undergone cutdown and others who had catheters inserted percutaneously, found that the incidence of culture-positive catheter tips rose from 0% to 52% after 48 hours. In the study by Moran and coworkers, Staphylococcus albus was the predominant organism that was isolated, but organisms more commonly thought of as pathogenic (Staphylococcus aureus, Enterococcus spp, and Proteus spp) were isolated with greater frequency from cutdowns that had been in place for longer periods. Rhee and coauthors reported a 1.4% infection rate and one episode of cellulitis after 73 cutdown attempts. All catheters were removed within 24 hours.

Some evidence indicates that the rate of infectious complications decreases when a broad-spectrum topical antibiotic ointment is applied to the cutdown site. Moran and colleagues found an infectious complication rate of 18% when topical polymyxin B-neomycin-bacitracin (Neosporin) was used as opposed to a 78% rate in a placebo-treated group. In this study, topical antibiotic use resulted in only a moderate decrease (from 53% to 37%) in the incidence of phlebitis, but a significant decrease (from 86% to 14%) in the incidence of phlebitis associated with positive cultures. This suggests that phlebitis is often a chemical or an irritative process rather than the
result of infection. Whatever the cause, the incidence of phlebitis is clearly related to the duration of catheterization. Early catheter removal is a key factor in the prevention of both phlebitis and the infectious complications of venous cutdown. This is especially true of lines inserted during emergency resuscitative treatment. Such lines should be removed as soon as the patient’s condition permits and alternative access routes are in place.

Proper attention to details of the surgical technique will limit the occurrence of minor complications such as local hematoma, abscess, and wound dehiscence. One can avoid injury to associated structures by selecting a site where the vein is well isolated, and the physician feels most comfortable performing the procedure.

ACKNOWLEDGMENT

The authors and editors wish to sincerely thank Patricia Lanter and Justin Williams for their contributions to this chapter in previous editions.
REFERENCES

4. Committee on Trauma, American College of Surgeons: Advanced trauma life support instructor manual, Chicago, 2008, American College of Surgeons.
Indwelling Vascular Access Devices: Emergency Access and Management

Scott H. Witt, Christine M. Carr, and Diann M. Krywko

Indwelling vascular access devices (IVADs) provide routes for short- and long-term infusion of antibiotics, antifungal agents, hyperalimentation fluids, chemotherapeutic agents, blood products, analgesics, and anesthetic agents. In addition, they provide access for lifesaving procedures such as hemodialysis (HD) and plasmapheresis. As of 2013, nearly 422,000 patients were receiving HD therapy, with 3 to 5 million central venous catheters being placed yearly in patients in the United States.1,2 For the purpose of this chapter, all implanted devices and intermediate- to long-term catheters for vascular access are considered IVADs. Arteriovenous (AV) fistulas and AV grafts are included because of their similarities to IVADs.

HISTORICAL PERSPECTIVE

A major advance that ultimately led to the development of several types of indwelling catheters was the introduction of Silastic (polymerized silicone rubber) in 1948 by the Dow Corning Corporation. This biocompatible material is an ideal substrate for intravenous (IV) catheters because it is chemically inert, antithrombogenic, rigid at room temperature, and pliable at body temperature. In 1973, Broviac and coworkers used this material to develop an indwelling right atrial (RA) catheter for total parenteral nutrition (TPN).3 In 1979, Hickman and colleagues reported their experience with a catheter that used this material to develop an indwelling right atrial (RA) catheter for total parenteral nutrition (TPN).4 In 1979, Hickman and colleagues reported their experience with a catheter that could be used for blood products and drug therapy in bone marrow transplant recipients.4 A totally implantable vascular access device (TIVAD) was described by Fortner and Pahnek in 1976.5 Since that time, TIVADs have become a mainstay of treatment in oncology patients, improving quality of life by allowing less painful IV access and permitting unrestricted mobility.

Temporary access for HD via an external AV shunt was pioneered by both Quinton and Scribner and their coworkers in 1960.6,7 This original shunt was composed of a loop of tubing lying on the volar aspect of the forearm that connected the radial artery to a wrist vein. Although it provided effective dialysis, it was associated with a high rate of infection, thrombosis, and restriction of patient activity. Brescia and colleagues then introduced the peripheral subcutaneous autogenous AV fistula in 1966.8 This Brescia-Cimino internal fistula used a side-to-side anastomosis to connect the radial artery to the cephalic vein in the nondominant hand. Erben and associates described the routine use of percutaneous cannulation of the subclavian vein for HD in 1969.9 Finally, in 1979, Uldall and coworkers reported the development of a single-needle, subclavian HD catheter.10

INDWELLING VASCULAR ACCESS DEVICES (IVADS)

IVADs are typically chosen based on the least invasive, smallest catheter, with the lowest risk for complications, which will last as long as the length of therapy that is anticipated.11 Length of therapy is often the major consideration when choosing a device. Long-term IVADs consist of cuffed, tunneled RA catheters, and implantable ports (Video 24.1). Medium-term IVADs include midline catheters (lasting weeks) and peripherally inserted central catheter (PICC) lines (lasting months). Short-term devices (see Chapter 21) include short peripheral IV, subcutaneous (butterfly), and percutaneous, nontunneled central catheters. Additional IVADs include those used for dialysis, in addition to AV fistulas and grafts.

Cuffed, Tunneled Right Atrial (RA) Catheters (Broviac, Hickman, Hemocath, Leonard, Raaf)

Several cuffed, tunneled RA catheters are available, each with differences tailored to specific applications (Fig. 24.1A). The Broviac (Bard Peripheral Vascular, Tempe, AZ) is an all-Silastic, single-lumen catheter, with a 1.0-mm internal diameter (ID). It is 90 cm long with a thin intravascular segment (55 cm). The Hickman (Bard), also a Silastic catheter, has a 1.6-mm ID. It allows more frequent blood sampling without jeopardizing luminal patency.12 Single-, double-, and triple-lumen variations exist. Hemocath/Permacath has the largest bore of the RA catheters, with an ID of 2.2 mm. Quinton Instrument Company (Bothell, WA) manufactures these catheters for HD, plasmapheresis, long-term nutritional support, and pain control. The main advantage of these catheters is that they can be used immediately as a bridge to longer-term devices. They are not recommended for long-term access in patients undergoing dialysis if an AV fistula or graft (Video 24.2) is possible, as long-term dialysis with tunneled catheters has been associated with an increased risk for death, a five- to tenfold increase in the risk for infection, and a decreased likelihood of adequate dialysis.13

Nonemergency insertion of RA catheters is typically done in an operating room or interventional radiology suite. The device is introduced via the arm, upper anterior chest wall, or neck, and tunneled subcutaneously to enter the superior vena cava (SVC) system via the cephalic, subclavian, internal, or external jugular vein (Fig. 24.2), advancing the distal tip of the flexible catheter to the distal SVC or into the mid-RA area. The subcutaneous tunnel isolates the venous puncture site from the skin and decreases the potential for bacterial contamination. Dacron cuffs (one near the venous entrance site and one near the skin exit site) anchor the catheter and are believed to inhibit colonization of the SVC by skin organisms.14 However, no study has been able to support this belief. Advantages of an RA catheter include ease of insertion and use, minimal interference with patient activity, low incidence of major complications or unintended dislodgment, ease of removal, and potential repair via a kit. Disadvantages include the need for regular maintenance and the potential for unacceptable cosmesis.
The cutdown technique offers potential speed (mean placement time, 15 minutes), safety (negligible risk for pneumothorax), and avoidance of early and late complications. Placement is typically in the nondominant arm, with the portal in the upper part of the arm or chest, unless a vein is occluded or radiation therapy is planned on that side.

Disadvantages of this type of device include increased cost, the need for a specific non-coring Huber access needle, and the small gauge (20 to 22) of the access needle, which limits fluid infusion rates. Typical TIVADs do not have the capability to withstand the pressure required for power injection. PowerPort by Bard is an implantable port that allows for venous access and power injections required for contrast-enhanced computed tomography scans.

**Groshong Catheter**

In contrast to the Broviac and Hickman catheters, the Groshong (Bard) catheter has slit-like openings just proximal to the end of the intravascular portion of the catheter (see Fig. 24.1B). This feature obviates the need to use a heparin lock (saline may be used). In addition, external catheter clamping is not necessary. Disadvantages are its high cost and requirement for pressurized infusion systems.

**TIVADs/Ports**

*(Port-A-Cath, Proport, Infuse-A-Port, Mediport)*

Since 1983, implanted ports have become the mainstay for long-term cancer therapy. TIVADs are tunneled RA catheters, but they differ from Broviac and Hickman catheters in that they have a subcutaneous titanium or plastic portal with a self-sealing septum (Fig. 24.3) that may be accessed by puncturing a specially designed needle (90-degree angled Huber needle) through intact skin (Fig. 24.4). Cosmetically, they are superior to external tunneled catheters, require less maintenance, and afford patients greater freedom of movement and activities, such as swimming or bathing.

TIVADs may be inserted on an outpatient basis under local anesthesia via a subcutaneous tunnel or open cutdown.
vessels. PICCs (Fig. 24.5) are made of two substances, either polyurethane (Intracath) or silicone (Intrasil), are radiopaque, and measure 50 to 60 cm in length, with an outside diameter of 2 to 7 Fr. The catheter may have a single- or double-lumen configuration and can be open- or close-ended or valved (e.g., Groshong). An open-ended PICC cannot prevent feedback of blood into the catheter and therefore must be flushed one or more times daily with heparinized saline. The Groshong valve reduces backup of blood into the catheter and therefore requires flushing as infrequently as once a week.

Selection of the device should be based on the number of lumens necessary for therapy. Selection of the access site depends on many factors including: the suitability of target vessels, body habitus, handedness, ability to manage self-care, comorbid conditions, desired infusion rate, number and compatibility of concurrent infusions, infusate characteristics, and the estimated duration of therapy. Infusate that is hyperosmolar (e.g., TPN) or vesicant requires rapid dilution. Accordingly, the tip must be in the SVC, where the estimated flow rate is 2000 mL/min. PICC lines are most frequently placed in the superficial veins proximal to the antecubital fossa (usually the basilic or cephalic vein) (Fig. 24.6). However, they may also be placed via a transhepatic or translumbar approach when the SVC is thrombosed or occluded.17

Advantages of PICC lines include usefulness in a wide variety of clinical situations, ease of placement, and ease of use and maintenance. They do not require surgical insertion and may be placed in an outpatient setting. A PICC line is an excellent vehicle for medium- to long-term IV therapy. With proper care, PICCs can remain in place for long periods, even months to years.18 To remove a PICC line, simply withdraw it from the vein, and apply pressure and a sterile bandage.

**Midline Peripheral Catheters**

Midline catheters are often confused with PICC lines. They are also placed peripherally in the superficial veins of the antecubital fossa or upper part of the forearm. They differ from PICCs in that they are peripheral, not central catheters. Midlines are typically shorter (20 cm), with the tip terminating near the axillary vein. Placement above the axillary vein results...
Hemodialysis (HD) IVADs

Vascular access for dialysis, which is often referred to as the Achilles heel of patients with end-stage kidney disease, remains problematic. From the moment that the first access is created, an ongoing process is started that will end with the loss of all access sites if the patient survives long enough. Clinical practice guidelines of the National Kidney Foundation—Disease Outcomes Quality Initiative (NKF-DOQI) recommend early construction of an AV fistula and avoidance of catheters for permanent or prolonged vascular access, except in rare circumstances where AV graft or fistula formation is not feasible because of a lack of acceptable anatomic sites or limited life expectancy of the patient. However, close to 70% of patients in the United States begin dialysis with a central catheter because a well-developed AV fistula is not available at the time of the initial catheter requirement.

Temporary Dialysis Catheters (Quinton, Mahurkar, Tessio, Vascath, Uldall)

Temporary vascular access catheters (Fig. 24.7) are used for emergency HD or for temporary HD access if a more permanent dialysis route (AV fistula or graft) is not available, or has recently been placed and has not matured yet. The advantage of tunneled catheters is the ability to provide immediate access or temporary access while a more permanent structure matures, but this carries an increased long-term risk for infection, dysfunction, and central venous stenosis. The majority of bacteremia episodes in patients undergoing HD are caused by HD catheters, with an approximate 20% to 25% risk over the average duration of use.

These large-bore catheters allow the necessary blood flow rate of 300 mL/min for dialysis. The Quinton catheter has two ports, one to deliver the patient's blood to the dialysis machine, and another to return the blood to the patient's circulation (see Fig. 24.2B). These catheters are placed in a central vein, either the internal jugular, subclavian, or femoral. The right internal jugular approach is preferred, even if permanent access is to be created on the right side, because it has the lowest thrombosis rate. The NKF-DOQI recommends avoiding the subclavian vein unless no other options exist or the ipsilateral arm has no more permanent access sites. For patient comfort a special 180-degree catheter can be used (Fig. 24.8). There are two avenues to place this catheter: percutaneous or surgical. Emergency percutaneous placement may be performed by the emergency clinician at the bedside. Using sterile technique and after injection of a local anesthetic, insert the catheter by following the same procedure for placing a central line into one of the central veins.
Indwelling Vascular Access Devices: Emergency Access and Management

is a direct subcutaneous anastomosis of an artery and vein without prosthetic material and is the preferred means of vascular access for HD (see Fig. 24.9). The minimum time for fistula maturation is at least 1 month, much longer than required for graft cannulation.

Historically the percentage of patients with AV fistulas fell well below the recommended goal, with most patients receiving AV grafts or long-term vascular access catheters. In recent years, however, there has been an increased focus on fistula creation. The goal of chronic HD vascular access is to provide safe, effective, and repeated easy access to the circulation. The three principal types of access are native AV fistulas, synthetic grafts, and double-lumen tunneled cuffed catheters (described previously). Fistulas and grafts are collectively termed shunts. It may be difficult to distinguish a fistula from a graft by gross inspection alone. Hints to identification include location and shape. Grafts are rarely placed in the forearm, which is the preferred site for fistulas. Fistulas tend to be more tortuous and serpentine, whereas synthetic grafts are straighter or C shaped.

HD access types differ in failure rates, patency, complications, and other morbidity. Both grafts and fistulas are subject to vascular perturbation and integrity issues from the high flow rates and repeated access. Grafts are subject to pseudoaneurysms when there is a breach in the integrity of the graft and they are more likely to rupture. Fistulas, also subjected to bulging of the vessel walls, may form a true aneurysm. Dilated veins in a fistula can simulate an aneurysm. Both types of vascular deformities can rupture. Multiple defects may require a replacement access.

Both fistulas and grafts must mature before they can be used for HD, a process that may take several weeks to several months. HD is often performed via a Quinton catheter during this hiatus, so a patient in the emergency department (ED) with both a catheter and a shunt either has a nonfunctioning fistula or graft, or an access site that is still immature.

Complications common to both grafts and fistulas are thrombosis, infection, steal syndrome, venous hypertension, bleeding (Video 24.3), seromas, and aneurysms (Fig. 24.9). Overall, grafts are more likely to experience infection and thrombosis requiring thrombectomy or require other types of access intervention.

Arterial Venous (AV) Fistulas

In general, fistulas are preferred over grafts because of superior long-term patency and lower complication rates. An AV fistula
years there has been a concerted effort to increase the utilization of AV fistulas, most notably the National Kidney Foundation's Fistula First Breakthrough Initiative. The most recent data from the Dialysis Outcomes and Practice Patterns Study (DOPPS) show that from 2002 to 2011 AV fistula use in the United States almost doubled (32% to 62%), and AV graft use fell by more than 50%. Use of an autogenous fistula is associated with the longest period of patency with relative freedom from thrombotic and infectious complications.

Fistulas have a 15% rate of primary failure, defined as being unable to provide reliable access as a result of early thrombosis or failure to mature. However, once a fistula matures, long-term patency is high (48% of AV fistulas versus 14% of AV grafts at 5 years), with low infection rates relative to grafts.

An autogenous AV fistula is constructed by anastomosing an artery to a vein (see Fig. 24.9), preferably a nearby one. Various configurations are possible, but AV fistulas are typically an end-to-side, vein-to-artery anastomosis. The radial-cephalic (Brescia-Cimino) fistula in the forearm is the most frequently used (Fig. 24.10), with the brachial-cephalic, brachial-basilic, and rarely the proximal part of the thigh being alternatives. Over time, the venous portion of the shunt is subjected to high pressure, and the fistula becomes arterialized (hypertrophied and dilated), which renders it suitable for repeated vascular access. Full epithelialization of the shunt does not occur for 3 to 6 months, thus necessitating anticipatory placement as renal function worsens.

AV Grafts

If a forearm radiocephalic fistula cannot be constructed or has failed, an AV bridge graft using a donor vein or synthetic material is a well-accepted alternative. Several synthetic materials are used for grafts. Grafts can usually be cannulated earlier than fistulas, often within weeks of placement. Polytetrafluoroethylene (PTFE) is most commonly used, but takes 3 weeks to mature. An available polyurethane graft (Vectra [Bard]) has the ability to be accessed within 24 hours. A standard graft is 6 to 8 mm in diameter and usually positioned in a U-shaped subcutaneous tunnel in the forearm. The graft is attached by end-to-side anastomoses to the brachial artery and antecubital vein. If no suitable antecubital vein is available, a straight bridge graft between the brachial artery and either the axillary or the basilic vein is often used. Multiple configurations are possible (Fig. 24.11). A jump graft between opposite extremities with creation of a loop across the chest or anastomosis of the axillary artery to the iliac vein is a possibility, if all other sites have been exhausted.

When compared with AV fistulas, AV grafts have a significantly higher incidence of thrombosis, infection, pseudoaneurysm formation, and limb loss. The risk for infection with HD grafts is approximately 10% over prolonged use and 1% with HD fistulas. Graft infection requires complete excision to eradicate an infection of the foreign material, whereas fistula infections may resolve with IV antibiotic use. AV grafts have a low primary patency rate (31% at 1 year according to one study). However, they have a low incidence of aneurysm formation and are comparatively easy to revise. The estimated life span of an AV graft in clinical practice is often less than 2 years.

ACCESSING IVADS IN THE ED

When IV access is required in patients with IVADs, standard methods of peripheral access should be attempted first to preserve the life span of the vascular access device and avoid complications. However, IVADs, AV fistulas, and grafts may be accessed as a last resort in emergency situations for phlebotomy and infusion of medications and fluids. Because of the complications of infection and catheter malfunction, dislodgment, and fracture, only personnel with the requisite knowledge...
and skill should access IVADs if feasible. When IVADs are accessed, ensure antisepsis throughout the procedure. Assuming that proper access methods are used to prevent infection, the greatest risk is sludging in the catheter with resultant occlusion.

**Accessing Long-Term Venous Access Catheters**

To access the catheter (with the exception of Groshong catheters, which have backflow valve protection), first clamp it to prevent air embolism. Patients usually carry their own clamps, but a hemostat with teeth will suffice in an emergency. In this case, wrap sterile tape or tubing around the teeth of the hemostat. Remove the cap of the catheter and withdraw any mobile clots with a syringe. Remove approximately 3 to 5 mL of blood and then attach a 10-mL syringe to a single-dose vial of sterile normal saline. Smaller syringes generate greater amounts of pressure for infusion, which can lead to increased intraluminal pressure and rupture of the catheter. Inject 3 to 5 mL of solution and then again withdraw it to ensure patency. Flush again with saline. More pronounced infusion might be necessary to ensure the patency of Groshong catheters.

To accomplish phlebotomy, withdraw the dead space solution, reclamp it, and use a separate syringe to remove the desired amount of blood. If clots are withdrawn, continue withdrawing blood until it is clot free. Inject bolus medications and infuse IV solutions through the catheter, and clamp it whenever it is unattached. Do not administer medications concurrently that are known to be incompatible when mixed (e.g., calcium and bicarbonate), even through separate ports of multilumen catheters. Deliver a 5-mL normal saline flush between medications through a 10-mL syringe. On completion of either blood withdrawal or medication infusion, inject 3 to 4 mL of saline to flush it and then inject 3 to 5 mL of heparin (1000 Units/mL). Clamp the line and reposition the cap. Note that 1000 Units/mL of heparin should be used; less concentrated solutions may promote clotting. Do not inject larger amounts because it can systemically heparinize the patient. Groshong catheters need not be flushed with heparin but instead may be flushed briskly with 5 to 10 mL of saline. Multilumen central catheters have one port for each lumen, so access each one in the same manner. After antisepsic preparation, gain access either by inserting a needle or a syringe into the protective cap or by removing the cap entirely. Flush with 5 mL normal saline or sterile water, and verify backflow before all subsequent procedures. Perform phlebotomy through the proximal lumen to prevent mixing with medications being delivered through the other ports. Deliver IV infusions in similar fashion, and inject a normal saline flush between medications. Following the procedure, flush 3 to 5 mL of heparin (1000 Units/mL) through each port.

**Accessing TIVADs**

The procedure for accessing TIVADs is unique because these devices are not external. Instead, a circular reservoir (cylinder) lies subcutaneously on the anterior chest wall. First, palpate the cylinder and then prepare the overlying skin with povidone-iodine solution. Fill a 10-mL syringe with normal saline and attach it to connecting tubing. Attach this to a 19- to 22-gauge, 90-degree tapered (Huber) needle. The Huber needle is a specialized needle designed for use with TIVADs to prevent damage to the portal septum. It has a 90-degree bend with a slightly curved tip and the opening on the side rather than on the end. Most importantly, the Huber is a noncoring needle. This avoids damage to the Silastic septum and allows up to 2000 punctures. Do not access the TIVAD with a standard needle unless an arrest is in progress and a Huber needle is not immediately available. Apply a clamp to the connecting tubing whenever the system is open. Expel the air and insert the Huber needle through the septum of the reservoir. Insert the needle slowly and steadily through the diaphragm until it contacts the back of the reservoir. Be aware that although incomplete perforation of the septum will block flow, substantial pressure may also damage the back of the device and bend the tip of the needle. Remove the clamp slowly and inject 5 mL of solution to ensure patency. If patency is not easily demonstrated, consider using alteplase (recombinant tissue plasminogen activator) as a fibrinolytic agent for catheter occlusion.

Once the solution has been injected, apply gentle negative pressure to demonstrate backflow of blood. Stabilize the Huber needle by building a 4- × 4-inch gauze pad about the needle and further reinforce it with 2.54-cm (1-inch) silk tape. First, remove 8 to 9 mL of blood with a separate syringe and waste it, and then perform phlebotomy through the extension tubing. If necessary, deliver IV solutions through extension tubing, but remember that the rate of flow will be limited by the smaller radius of the Huber needle. Deliver a 5-mL sterile normal saline flush between medications. Complete the procedure with a 3- to 5-mL heparin (1000 Units/mL) flush and remove the Huber needle.

**Accessing AV Fistulas, Grafts, and HD Catheters**

AV fistulas, grafts, and temporary dialysis catheters are placed in patients who require HD, and they represent the sole access for that purpose. Consequently, routine use of these sites for phlebotomy and fluid administration is strongly discouraged. In fact, venipuncture in the same extremity as a patent AV shunt is not recommended, except for the veins in the dorsum of the ipsilateral hand. However, when standard IV access cannot be obtained under emergency circumstances, fistulas, grafts, and catheters may all be used to administer IV solutions and medications (Fig. 24.12). Though avoided whenever possible, the option to access these sites is based on clinical judgement by the clinician. If possible, ascertain patency of the fistula by noting a bruit and palpable thrill, although these signs may not be appreciable if the patient is in extremis. Prepare the area overlying the fistula with antisepsic solution and access the fistula with the smallest gauge needle that is appropriate. Puncture 1 to 2 cm from the end of the anastomosis nearest the venous side and avoid aneurysmal sites. Access AV shunts similarly by placing the smallest needle possible in the catheter and bridging the arterial and venous circulations. Apply local pressure for at least 5 minutes after the procedure is completed and monitor subsequently for hemorrhage. Access the Uldall and Mahurkar catheters in much the same way that multilumen central catheters are accessed. Remove or inject through the retaining cap on each arm. Up to 5000 units of heparin are present within the two lumens, thus it is imperative to aspirate before administering fluid or medications. After use, flush each catheter arm with 10 mL of normal saline and instill 1.5 mL of heparin solution (1000 Units/mL) into each one.
Infection is the most common complication leading to removal of IVADs and potentially the most serious. Infection can also result in shunt thrombosis, rupture, or massive hemorrhage. It is estimated that more than 250,000 cases of nosocomial IVAD-related bacteremia occur annually in the United States with an associated mortality rate greater than 10%. Infectious complications include endocarditis, septic arthritis, pulmonary emboli, osteomyelitis, and spinal epidural abscess. The definitions of IVAD-related infections are not uniform, which makes it difficult to compare the results of various investigations. However, it is generally agreed that risk factors for infection include the site of insertion, duration of catheter placement, type of catheter, and patient factors. Most studies show a lower incidence of infection for TIVADs than for external systems, presumably because of lack of direct access by cutaneous organisms. Rates of IVAD infection tend to be highest in the first 3 months after insertion, with skin flora being most common. The rate of infection decreases significantly and reaches a plateau after 5 to 6 months.

Clinical findings are unreliable in the diagnosis of infection secondary to IVADs. Fever, rigors, and an elevated white blood cell count are nonspecific, whereas purulent drainage at the insertion site is specific but not sensitive. Therefore evaluation beyond physical examination alone is essential when infection is suspected, including Gram stain of purulent material at the site, blood cultures, and culture of catheter segments. In addition, transesophageal echocardiography is indicated if valvular vegetations are suspected by blood cultures positive for *Staphylococcus aureus*, persistent bacteremia or fungemia after removal of the catheter, or lack of clinical improvement.

Draw two sets of blood for culture, with one set through the catheter itself and one from a peripheral site. Positive blood cultures for *S. aureus*, coagulase-negative staphylococci, or Candida species, in the appropriate patient setting and in the absence of another identifiable source of infection, should increase suspicion for catheter-related bloodstream infection. Similarly, if cultures drawn from the IVAD become positive 2 or more hours before peripherally drawn cultures, it suggests the patient has a catheter-related bloodstream infection. Infusate-related bloodstream infection is uncommon and defined as isolation of the same organism from both the infusate and separate percutaneous blood cultures, along with no other source of infection. When this diagnosis is suggested, cultures
of the infusate should be performed in addition to catheter and peripheral blood cultures.12

When a catheter-related infection is suspected, an important management decision is required to determine whether to remove the catheter. Removal is recommended in patients who have a catheter placed solely for short-term use. In patients with long-term catheters, removal of the catheter is advised in the setting of severe sepsis, suppurrative thrombophlebitis, endocarditis, or persistent positive blood cultures (longer than 72 hours) while on appropriate antibiotics, as well as if the infection is caused by *S. aureus*, *Pseudomonas aeruginosa*, fungi, or mycobacteria. For patients who require long-term vascular access for survival or have no other vascular access options, salvage of the catheter may be attempted if the infecting agent is not *Bacillus*, *Micrococcus*, or *Propionibacterium*, in addition to the previously listed organisms. If the decision is made to leave the catheter in place, obtain blood for repeated cultures and remove the catheter promptly if they remain positive following 72 hours of appropriate antibiotic therapy.36

If an IVAD is removed, determination of the offending organism is made by culturing a portion of the catheter. Place the catheter in a sterile container and send it to the laboratory for culture, sensitivity, and Gram stain. The most widely used laboratory technique for the clinical diagnosis of catheter-related infection is the semiquantitative method, in which a segment of the catheter is rolled across the surface of an agar plate and colony-forming units (CFUs) are counted after incubation overnight. Quantitative culture of the catheter segment requires either flushing the segment with broth or vortexing in broth, followed by serial dilutions and surface plating on blood agar. A yield of 15 CFUs or more from a catheter by semiquantitative culture or a yield of 102 or more from a catheter by quantitative culture with accompanying signs of local or systemic infection is indicative of a catheter-related infection.31

The process of HD requires several connections to the HD catheter, thereby increasing the risk for infection. The rate of bacteremia in HD patients attributed to the vascular access site varies from 48% to 73%. The incidence is highest when central venous dialysis catheters are used, whereas native AV fistulas carry the lowest risk for infection.37 As with other IVADs, if infection is suspected, initiate empirical antibiotic therapy based on epidemiologic and patient factors, followed by narrowed-spectrum therapy after isolation and determination of sensitivities. The most common infecting organism is *S. aureus.*

**Antimicrobial Therapy**

Pending the results of culture, the initial choice of an antibiotic is empirical and depends on the clinical setting, site of infection, type of device, host factors (e.g., immunocompromised state), severity of illness, and whether the device has been left in situ. If an IVAD-related bloodstream infection is suspected, vancomycin should be initiated empirically as coagulase-negative *Staphylococcus* is the most common infectious organism and the prevalence of methicillin-resistant *S. aureus* continues to increase.38 This should be followed by a semisynthetic penicillin or other appropriate antibiotics as guided by sensitivity studies. If the patient is severely ill, septic, in an immunocompromised state, or has a femoral catheter in place, additional coverage for gram-negative bacilli should be initiated.39 Institute a third-generation cephalosporin unless the organism is extended-spectrum β-lactamase positive, in which case a carbapenem is more appropriate.40 If there is concern for *P. aeruginosa*, initiate therapy with a fourth-generation cephalosporin (e.g., cefepime) or carbapenem, with or without an aminoglycoside. In addition to bacterial pathogens, fungemia should be suspected in septic patients who are receiving TPN; those with prolonged use of antibiotics; those with a history of a hematoologic malignancy, or a bone marrow or solid organ transplant; or if a femoral catheter is in place.41 If fungemia is suspected or confirmed, initiate an antifungal medication and remove the catheter. An echinocandin medication (e.g., micafungin, caspofungin) is the recommended empirical treatment of suspected fungal infection.39 Although there are no compelling data to specify a duration of antibiotic therapy, expert recommendation is to treat *S. aureus* infection for 14 days, gram-negative bacilli for 7 to 14 days, and candidal infection with antifungal therapy for 14 days after the first negative blood culture. Complicated infections (e.g., those with suppurrative thrombophlebitis, endocarditis, or other similar infections) require 4 to 6 weeks of antibiotic treatment.39

If an exit site infection is suspected, any drainage from the site should be cultured in addition to obtaining blood cultures. For uncomplicated exit site infections (those lacking signs of systemic infection, positive blood cultures, or purulence) initial treatment consists of topical antimicrobials. If the infection does not resolve with topical antimicrobials, systemic antibiotics should be administered and consideration given to removing the catheter. Patients with a tunnel infection or port abscess should have the catheter removed and receive systemic antibiotics for 7 to 10 days.39

If the IVAD is retained, consider antibiotic lock therapy (ALT). ALT is a therapeutic option if the goal is to salvage the catheter. Most infections in tunneled catheters originate in the hub and spread to the catheter lumen. A biofilm can form and make eradication of the organism difficult because systemic antibiotics cannot achieve therapeutic levels there. ALT involves filling the catheter hub and lumen with a higher concentration of antibiotics and leaving them in place for extended periods. It is recommended that dwell times not exceed 48 hours before reinstilling the antibiotic solution.39,40 If it is a femoral catheter, dwell time should not exceed 24 hours. ALT should always be used in conjunction with systemic antibiotics, and the recommended duration of treatment is 7 to 14 days.39

**Prophylactic Measures**

**Antibiotic Prophylaxis During Initial Line Insertion**

Prophylaxis with vancomycin or teicoplanin during central line insertion has not demonstrated a benefit in reducing catheter-related bloodstream infections. Based on the limited data available, Centers for Disease Control and Prevention (CDC) guidelines currently recommend against giving vancomycin prophylactically because it is an independent risk factor for the acquisition of vancomycin-resistant enterococci and staphylococci, with reduced susceptibility to glycopeptides.31 Rather than using antibiotic prophylaxis, focus efforts on interventions designed to discourage the emergence of antimicrobial resistance such as maximal barrier precautions.42

**Impregnated Catheters**

Using antimicrobial- or antiseptic-impregnated catheters or silver-impregnated collagen cuffs may be an effective intervention to reduce IVAD-related bloodstream infection. A 2013 metaanalysis favored impregnated catheters in reducing central line–associated bloodstream infections (CLABSIs), but it was
acknowledged that the overall methodology of the studies included was poor. Nonetheless, the CDC currently recommends the use of these devices if the catheter is anticipated to remain in place for more than 5 days and if the institution’s CLABSI rate is not decreasing after implementation of a comprehensive strategy to reduce catheter-associated infections.

Routine Line Changing
Despite the incidence of infection and the potential complications, routine changing of IVADs is not recommended. Cobb and coworkers found that replacement of IVADs every 3 days did not prevent infection and that doing so over a guidewire actually increased the risk for bloodstream infection.

Thrombus Formation
It has been estimated that 2% to 42% of IVADs are associated with deep venous thrombosis (DVT). Risk factors for catheter-related DVT include the composition, diameter, and position of the IVAD; elevated intraluminal pressure; turbulent blood flow; vascular calcification; endothelial injury; and increased levels of fibronectin. Polyurethane and silicone catheters have a lower rate of IVAD-related DVT than do polyethylene or Teflon-coated catheters. A catheter with an external diameter of less than 2.8 mm has a lower incidence of DVT. Incorrect placement of the IVAD in the SVC, as opposed to the junction of the SVC and right atrium, results in a higher incidence of catheter-related DVT. The majority of these thromboses are asymptomatic. The incidence of clinically overt DVT has been found to range from 0.3% to 28.3%, with most studies finding the incidence to be less than 5%. The first 6 weeks after catheter insertion presents the greatest risk for thromboembolic complications.

Most HD access failures (80%) are related to thrombosis, with greater than 90% of these thromboses associated with venous outflow stenosis. Histologically, hyperplasia of the endothelium and fibromuscular vessel wall occurs. This may continue to the point of complete occlusion, and prevent infusion or aspiration from the access site. Early detection may be enhanced by maintaining a high index of suspicion and noting prolonged bleeding after withdrawal of the cannula. Clinically, an absence of a thrill or change in the bruit on examination indicates occlusion of the fistula or graft. Prompt consultation with a vascular surgeon for declotting of the thrombosed access and correction of any underlying stenosis is indicated.

Catheter Occlusion
Occlusion of the catheter or low flow can be caused by the following: improper positioning, kinking, or compression of the catheter; intraluminal thrombi; extraluminal thrombi; fibrin deposits at the tip of the catheter; or intraluminal precipitation of infusate. Pinch-off syndrome occurs when the line (most often a PICC) is compressed between the clavicle and the first rib. It is manifested clinically as difficulty injecting that is posturally related. Overzealous withdrawal of the syringe will collapse the catheter and cause occlusion. Chest radiographs may reveal scalloping of the catheter. Maneuvers to facilitate flow of a potentially occluded catheter include the Valsalva maneuver, the reverse Trendelenburg position, slight tension on the catheter, placement of the catheter more laterally, IV hydration, and extension of the arms above the head. If these measures are unsuccessful, the occlusion may be caused by clot formation. The clinician should be familiar with local institutional recommendations for thrombolysis in the setting of recent IVAD occlusion attributed to a fibrin plug. Admission and consultation are usually required, and maneuvers to address clotted or occluded catheters are not a standard ED procedure, but may be performed under proper circumstances and according to local guidelines. Alteplase has been shown to be both effective and safe in treating central venous catheter occlusions caused by a clot. If attempted, first withdraw any mobile clot by aspiration with a syringe. Instill 2 mL of alteplase (1 mg/mL) and allow it to dwell for up to 120 minutes. If it remains occluded, the dose may be repeated once.

Embolization
A serious but uncommon complication of IVAD use is embolization with air, a catheter fragment, or thrombotic emboli. Air emboli develop when the lumen of the catheter is left open to air, or the catheter is fractured, perforated, or cut. The one-way valve in the Groshong catheter prevents this complication. Be careful to maintain a closed system by clamping the catheter appropriately to prevent delivery of air into the venous circulation. Should the externalized portion of a catheter be damaged, immediately place an appropriate clamp between the damaged portion and the skin. Be suspicious for air embolism if an open or damaged catheter is found in association with tachypnea and hypotension. Place the patient in a left lateral decubitus and Trendelenburg position to reduce ventricular outflow obstruction by air pockets. Initiate supportive measures, including high-flow oxygen. If this is unsuccessful, attempt aspiration with the catheter advanced into the right ventricle, if possible. Cardiothoracic surgery consultation and emergency thoracotomy to aspirate air (see Chapter 18) may be warranted.

Embolization of a catheter fragment is a potentially life-threatening complication that can cause acute dyspnea, palpitations, atypical chest pain, hypoxia, and atrial fibrillation. As with any foreign-body embolus sequelae include: sepsis, lung abscess, dysrhythmias, vascular or cardiac perforation, and sudden death. Catheter fragments may be identified radiographically, or potentially by ultrasonography, and may be removed either surgically or by intravascular retrieval methods.

Embolization of thrombi with potentially lethal sequelae may occur during routine flushing and injection of solutions through the catheter. Anderson and coworkers prospectively evaluated the size and frequency of catheter thrombi in 43 patients by aspirating after a urokinase flush. Clots were noted in 40 of 43 subjects and in 153 of 508 total specimens. Thrombi varied in size from small fragments to 5 cm in length. Burns and McLaren reported one case of a hemodynamically significant pulmonary embolus associated with PICC placement in a 77-year-old man.

Hemorrhagic Complications
Bleeding may be seen with any indwelling IVAD or AV shunt and is a potentially fatal complication. Between 2000 and 2006 over 1600 dialysis patients in the United States died from vascular access hemorrhage. It is often related to mechanical trauma, transient or preexisting thrombocytopenia, uremia or drug-induced platelet dysfunction, and infection. Repeated
cannulation of the fistula or graft weakens the wall of the device. Many fistulas will develop aneurysms from repeated use; synthetic grafts can develop pseudoaneurysms from frequent punctures (Fig. 24.13). Severe life-threatening hemorrhage from this high-pressure system may occur and necessitates rapid control. The primary goal should be to control bleeding and prevent exsanguination; however, take care to minimize damage to the IVAD that would prevent future use. Use full universal precautions because there is an increased risk for communicable disease in this patient population (2.4% for hepatitis B virus and 7.4% for hepatitis C virus in the United States). When the bleeding stops, observe the patient for 2 hours for evidence of rebleeding and to detect shunt thrombosis. Treatment of hemorrhage can be divided into mechanical modalities and correction of coagulopathy.

**Mechanical**

Literature addressing acute hemorrhage in patients with IVADs is clearly lacking. Many of the techniques discussed are anecdotal examples used by emergency clinicians, nephrologists, and vascular surgeons informally polled at our institutions. Individual institutional guidelines, techniques, and preferences may vary; therefore discussion with consultants is recommended when possible.

**Approach to Bleeding Complications**

It is important to identify the specific bleeding site, which is challenging when hemorrhage is massive (Fig. 24.14). This is most easily accomplished by (1) applying concomitant digital pressure over the feeding and draining vessels of the shunt above and below the bleeding site (see Fig. 24.17), or (2) applying a pneumatic blood pressure cuff or tourniquet distal to the fistula or graft to impede distal-to-proximal arterial flow, unless it is a loop graft, in which case it is applied above (proximal to) the device. The most easily controlled bleeding is typically that occurring immediately after dialysis, when the bleeding is from dialysis access puncture sites. Spontaneous bleeding between dialysis treatments often signifies more serious problems, such as infection, or mechanical problems with the access site. High pressure from venous stenosis secondary to long-term dialysis complicates the control of bleeding (Fig. 24.15).

**Direct Pressure**

Begin management of bleeding with direct pressure. Always use sterile technique to lessen the chance of infecting a bleeding shunt. With gloved hands, apply direct pressure with fingers and a sterile gauze bandage. For bleeding AV fistulas and grafts secondary to cannulation, apply pressure...
focally over the site of cannulation. For tunneled catheters, apply pressure over the site of vascular entry of the catheter, not the subcutaneous exit site. Note that this is not possible with subclavian IVADs. Hold the pressure for a minimum of 5 minutes and then reevaluate for hemostasis. Holding direct pressure for extended periods or maintaining more diffuse pressure may cause thrombosis and ultimate shunt failure, which is a known and unfortunate risk. Wrapping the site with an elastic bandage for a few minutes is acceptable, but longer application in this broad manner may lead to the entire graft clotting.

**Dialysis Clamps**

Special self-retaining dialysis clamps may be applied over a specific bleeding site (Fig. 24.16). Be careful that the clamp does not slip off the bleeding site because the surface area of the clamp is small. Place sterile gauze or topical thrombin-impregnated gauze under the clamp. Ten to fifteen minutes may be required to stop the bleeding, but it is best to avoid frequent checks during the first 5 to 8 minutes to avoid disrupting hemostasis.

**Thrombogenic and Dermal Adhesive Agents**

If venous oozing fails to stop with direct pressure, oxidized cellulose hemostatic agents (Oxycel [Becton Dickinson, Franklin Lakes, NJ], Surgicel [Ethicon, Somerville, NJ]), or a skin adhesive such as 2-octyl-cyanoacrylate, may be used to achieve hemostasis. Apply topical hemostatic agents directly over the site of bleeding and hold them in place with gloved hands and a gauze bandage or clamp. If utilizing a skin adhesive, apply the adhesive over the site of bleeding while also applying direct pressure proximally. After achieving hemostasis the patient should be observed to ensure rebleeding does not occur.

**Vasoconstrictive Agents**

Subcutaneous injection of 2 to 4 mL of lidocaine (2%) with epinephrine to form a wheal around the site may decrease bleeding by both vasoconstriction and local pressure. This may be used in conjunction with chemical cautery.

**Chemical Cautery**

Silver nitrate (AgNO₃), a mildly caustic and hemostatic agent, may stop residual bleeding. The agent needs to be protected from light until just before use. When ready, remove the wooden stick from the container and black plastic wrap and gently
apply the gray-tipped end directly on the site. Do not apply it aggressively because this can result in dissolution or dislodgment of the formed clot.

Once the bleeding is controlled, rebleeding is infrequent. It is prudent to observe the patient for 30 to 45 minutes and have the patient walk around and use the arm gently to ensure that bleeding will not recur when discharged. If no bleeding recurs, discharge is appropriate unless precluded by other conditions.

Suture
If direct pressure and previously mentioned techniques fail to achieve hemostasis, suturing may be necessary. Inject the bleeding site subcutaneously with lidocaine with epinephrine. Place a figure-of-eight or horizontal mattress suture with 4-0 nonabsorbable polypropylene or nylon (noncutting needle) at the site of bleeding (Fig. 24.17). Be careful to suture as superficially as possible to prevent damage to the underlying graft or fistula. The patient may require a venogram for evaluation of shunt patency before use. Remove the suture in a few days.

Coagulopathy
Control of hemorrhage may require treatment of an underlying coagulopathy. HD patients are generally considered to be at increased risk for bleeding because of uremic platelet dysfunction. The incidence of major bleeding in these patients is increased with the concomitant use of anticoagulant medications.58 Perform laboratory testing for evaluation as follows: complete blood count, blood urea nitrogen, prothrombin time, international normalized ratio (INR), partial thromboplastin time, and other tests depending on the patient's medical history or signs. Consult nephrology, vascular surgery, or both if possible before reversing therapeutic anticoagulation given the potential for shunt thrombosis and the need for close follow-up.

Uremic Platelet Dysfunction
If platelet dysfunction is suspected because of uremia, administer desmopressin (DDAVP) or cryoprecipitate to control the hemorrhage.30 Adult dosing of DDAVP to control uremic bleeding is 0.3 µg/kg intravenously as a single dose or every 12 hours. Onset occurs in 1 to 2 hours, and its duration is 6 to 8 hours after a single dose. Disadvantages include high cost and adverse reactions, including anaphylaxis, water intoxication or hyponatremia, and thrombotic events (rare). Use of DDAVP may be most appropriate if the platelet count and standard coagulation profile is otherwise normal.

Heparin-Associated Coagulopathy
Heparin is routinely administered during dialysis, and although there is no standardized dosing there are common regimens, which include providing an initial bolus (25 to 30 IU/kg) followed by intermittent, lower dose boluses (500 to 2000 IU) or an initial bolus followed by a continuous infusion.59 Heparin is usually stopped approximately 1 hour before the end of dialysis, and hence heparin-related bleeding is unusual in the ED. If uncontrolled bleeding occurs in the context of recent heparin use, reversal of heparin's effect with protamine sulfate may be necessary. Administering 1 mg of protamine sulfate will effectively reverse the anticoagulant effect of 100 units of unfractionated heparin. Protamine is less efficacious in the reversal of low-molecular-weight heparins. Nonetheless, current recommendations are to administer 1 mg of protamine for every 1 mg of low-molecular-weight heparin that the patient received.60

Warfarin-Associated Coagulopathy
The use of warfarin predisposes patients to potentially serious bleeding complications. In an actively bleeding patient with an elevated INR, immediate reversal of the medication's effect can be achieved with fresh frozen plasma or prothrombin complex concentrates (PCCs). PCCs have the advantage of requiring less total volume to achieve reversal of anticoagulation and therefore less risk for volume overload.60 Vitamin K should also be considered for reversal, following warfarin-associated bleeding guidelines.

Catheter Displacement or Malposition
Catheter displacement may occur accidentally secondary to patient movement, iatrogenically, or both. The catheter should be firmly secured with sutures, sterile tape strips (with care taken to avoid direct contact with the catheter itself), and premanufactured devices.18 Even with these measures disruption of placement may still occur. With regard to TIVADs, malposition of the intravascular portion may occur as a result of incorrect initial positioning or secondarily from forceful flushing, neck flexion, obesity, severe coughing, emesis, or upper extremity movement. Malposition of the port body may occur intentionally or by unintentional manipulation of the port (twiddler’s syndrome). Obtain a computed tomography scan of the chest to diagnosis this condition. Surgical intervention is usually necessary.61

Catheter Fracture
Fracture of an IVAD can occur either subcutaneously or in the externalized portion.62 It may take place with certain physical activities that involve repetitive and excessive motion.

Figure 24.17 To suture a bleeding dialysis shunt, instruct an assistant to apply proximal and distal pressure to stem the flow of blood and then inject the site with lidocaine with epinephrine. Place a figure-of-eight (shown) or horizontal mattress suture as superficially as possible to prevent damage to the underlying graft or fistula.
such as golfing, swimming, and weight lifting. Subcutaneous fractures cause pain and swelling and require removal of the line. Fractures in the externalized portion may be repaired with commercially available kits and do not always require replacement.

**Steal Syndrome**

Vascular steal syndrome is an uncommon but serious complication of AV fistulas and grafts, which is difficult to predict and may lead to access failure. It occurs because of preferential flow through the low-resistance fistula at the expense of the distal circulation. The syndrome is manifested by classic arterial insufficiency symptoms with exacerbation during dialysis: pain, pallor, numbness, motor weakness, and diminished or absent pulses distal to the shunt. Steal syndrome must be differentiated from diabetic or uremic neuropathy because it may lead to the development of irreversible neuromuscular dysfunction, tissue necrosis, and amputation. Prompt recognition and correction of hand ischemia leads to increased salvage and use of functioning shunts. Steal syndrome usually requires ligation of the AV access with placement of a new access device in the opposite extremity.

**AFTERCARE INSTRUCTIONS**

Before release from the ED instruct the patient regarding proper catheter care to prolong the device’s lifetime, and to decrease morbidity and mortality. Patients may bathe and swim normally after maturation of the site; they should avoid direct pressure on the reservoir and report bruising or bleeding immediately. Long-term venous access catheters and multilumen catheters should be dressed in sterile fashion and observed daily for bleeding and signs of infection, including fever, pain, redness, swelling, and purulent drainage. IVADs should be gently flushed on a routine basis. Heparin flushes are essential to prevent thrombosis. Tunneled (e.g., Hickman) and nontunneled (e.g., PICC) catheters require flushing twice weekly with 5 mL of heparin (10 Units/mL). TIVADs require flushing with heparin every 4 weeks and after use. Generally, Groshong catheters are flushed with 5 mL of saline once weekly. Mahurkar and Uldall catheters are flushed during dialysis. Mahurkar catheters are also used for apheresis, in which case they are treated three times a week with normal saline and heparin, as outlined earlier. Patients should not allow phlebotomy or infusion into an extremity with an IVAD, with the exception of phlebotomy in the ipsilateral dorsum of the hand, nor should they have blood pressure measured in that extremity. Clinicians should be knowledgeable about indwelling devices and complications. They should practice sterile technique, use care in handling, and avoid use unless an emergent situation requires access of the IVAD.

**Acknowledgment**

We gratefully acknowledge the work of Cemal B. Sozener, MD, and Pino Colone, MD, authors of this chapter in previous editions of this text.
REFERENCES


Establishing vascular access in critically ill and injured patients is central to the practice of emergency medicine. Moreover, placing an intravenous (IV) catheter in an acutely ill child can be one of the most challenging and frustrating procedures that a clinician can be called on to perform. Children have small peripheral vessels that collapse during shock, and their higher proportion of body fat makes visualization and palpation of peripheral vessels difficult. These factors often result in prolonged attempts and high failure rates.

In a review of vascular access success rates in children in cardiac arrest, the average time needed to establish percutaneous peripheral IV access was 7.9 ± 4.2 minutes, with only a 17% success rate. Some authors advocate the use of central venous lines in children, but such lines are also difficult to place and are associated with risk for arterial injury, infection, and pneumothorax. Alternative routes of drug administration, such as the endotracheal and rectal routes, are not reliable in patients in shock or cardiac arrest.

Peripheral IV access can also be difficult in adults, including those who are obese, have burn injuries, are volume-depleted, or are in shock. This difficulty is compounded in the prehospital or military setting, where environmental factors and the need for rapid transport can challenge even the most skilled practitioners.

Intraosseous (IO) access (Videos 25.1 to 25.5) can provide rapid, lifesaving intravascular access in challenging environments and in both pediatric and adult patients. All drugs and fluids that are delivered through a peripheral or central venous line can also be administered through an IO cannula. Drug and fluid dosing is the same as for IV administration. The serum concentrations and onset of action of IO administered medications during cardiopulmonary resuscitation (CPR) are comparable to those achieved with IV administration. Per animal studies, the IO route is as effective as central venous administration and may be superior to the peripheral IV route during cardiac arrest.

The American Heart Association, the American Academy of Pediatrics, and the American College of Surgeons all recommend IO access in children in emergency situations when IV access is not immediately possible. The latest edition of the advanced trauma life support course of the American College of Surgeons also notes that IO access using specially designed equipment is an important option in adult trauma patients.

IO access is often faster than IV access, and the success rate after failed IV attempts is high. In a retrospective study of pediatric cardiac arrest patients, time to IO access was significantly shorter than time to IV access. In a similar review of intravascular access during pediatric cardiac arrest, Brunette and Fischer noted that when compared with central venous access and venous cutdown, IO access and IV access were faster, and the success rate for IO access was 83% versus 17% for IV access. IO access can also be a rapid and successful technique in adults. Because of their success under difficult battlefield conditions, the US military has adopted the use of IO infusion devices.

IO access is not always successful. In a 5-year review of prehospital IO needle placement, success rates were higher in children younger than 3 years (85%) than in older children or adults (50%). The main causes of failure were errors in identification of landmarks and bending of the needles. Better needle design and new devices have helped overcome problems with bone penetration (see later section on Equipment and Setup).

HISTORICAL PERSPECTIVE

One of the earliest references describing the IO route is attributed to Drinker and colleagues, who examined the circulation of the sternum in 1922 and suggested it as a site for transfusion. The route was not used clinically until 1934 when Josefson, a Swedish clinician, injected liver concentrate into the sternum of 12 adult patients with pernicious anemia and reported that all 12 improved. Subsequently the technique became widespread in Scandinavian countries. In 1944 British physician Hamilton Bailey described the utility of sternal IO access in blackout conditions in London during World War II.

In 1940 the technique was introduced to American clinicians by Tocantins and associates, who described a series of animal and clinical studies demonstrating that fluid is rapidly transported from the medullary cavity of long bones to the heart. They recommended using the manubrium in older children and adults and the upper part of the tibia or lower part of the femur in children 3 years or younger. Over the next two decades, thousands of cases of IO infusion of blood, crystalloid substances, and drugs were reported. The procedure was more commonly used in children because of the difficulty in achieving other forms of IV access. Nevertheless, during the 1940s IO infusion was also used extensively in adults, and a sternal puncture kit for bone marrow infusion was a common component of emergency medical supplies during World War II. This resulted in more than 4000 reported cases of successful IO access in wounded soldiers. During this time relatively few complications were reported despite the fact that the needles were often left in place for 24 to 48 hours. In 1947 Heinild and coworkers reviewed 982 cases of IO infusion and reported only 18 failures and 5 cases of osteomyelitis. None of the cases of osteomyelitis occurred in patients who received isotonic solutions.

With the introduction of plastic catheters and improved cannulation techniques, the need for IO infusion as an alternative route for IV access diminished, and the technique was all but abandoned. In the mid-1980s James Orlowski brought about a renaissance in the use of IO access for pediatric resuscitation. While traveling through India during a cholera epidemic, he observed emergency health care workers using IO access to deliver fluids and medications. In 1984 he wrote an editorial, “My Kingdom for an Intravenous Line,” in which he advocated the use of IO access during pediatric resuscitation. After Orlowski’s editorial, others began promoting the use of IO access to allow rapid drug delivery during CPR in children. In 1988 IO techniques were adopted by the American Heart Association and included in the pediatric advanced life support guidelines. Since then the technique has become widespread throughout the United States and is recognized
as an accepted alternative to IV access in pediatric emergencies, and, increasingly, in neonatal and adult emergencies. In addition, the safety, ease, and effectiveness of the technique have led to its use in prehospital emergency care.\textsuperscript{7,23–26}

**ANATOMY AND PHYSIOLOGY**

Long bones are richly vascular structures with a dynamic circulation. They are capable of accepting large volumes of fluid and rapidly transporting fluid or drugs to the central circulation. The bone, like most organs, is supplied by a major artery (nutrient artery). The artery pierces the cortex and divides into ascending and descending branches, which further subdivide into arterioles that pierce the endosteal surface of the stratum compactum to become capillaries. The capillaries drain into medullary venous sinusoids throughout the medullary space, which in turn drain into a central venous channel (Fig. 25.1).

The medullary sinusoids accept fluid and drugs during IO infusion and serve as a route for transport to the central venous channel, which exits the bone as nutrient and emissary veins.\textsuperscript{27} The medullary cavity thus functions as a rigid, noncollapsible vein, even in the presence of profound shock or cardiopulmonary arrest.\textsuperscript{28} Radiographic studies have demonstrated that radiopaque dye spreads only a few centimeters in the medullary space before being transported to the venous system.\textsuperscript{29}

Almost every drug and fluid commonly used during resuscitation has been reported upon in clinical and preclinical IO studies. Essentially, all medications and fluids that might be delivered via a venous line can also be administered through the IO route. Dosing of IO medications and fluids is the same as for IV administration. Medications and fluids that have been administered through IO infusion are listed in Box 25.1. Crystalloid infusion studies in animals have demonstrated that infusion rates of 10 to 17 mL/min may be achieved with gravity infusion and rates as high as 42 mL/min with pressure infusion.\textsuperscript{30–32} In a swine model of hemorrhagic shock, Neufeld and colleagues\textsuperscript{31} found that the IO delivery rate of crystalloid was similar to that with both peripheral and central venous administration. IO crystalloid infusions have also been shown to produce a significant increase in blood pressure in a hemorrhagic shock model in rabbits.\textsuperscript{32}

In small animals (7 to 8 kg), the size of the marrow cavity is the rate-limiting factor, whereas in larger animals (12 to 15 kg), the size of the needle determines the flow.\textsuperscript{33,34} Blood under pressure can be infused approximately two-thirds as fast as crystalloid fluids.\textsuperscript{33} However, in a swine model evaluating IO infusion for mild therapeutic hypothermia, IO infusion of ice-cold saline was not as efficacious as intravenous infusion.\textsuperscript{35}

Comparisons of IO and IV infusion of drugs have demonstrated that the drugs reach the central circulation by both routes in similar concentrations and at the same time.\textsuperscript{14,36} This holds true even during CPR, during which sodium bicarbonate...
BOX 25.1 Medications and Fluids That Can Be Administered Intraosseously

<table>
<thead>
<tr>
<th>MEDICATIONS</th>
<th>FLUIDS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenosine</td>
<td>Mannitol</td>
</tr>
<tr>
<td>Antibiotics</td>
<td>Morphine</td>
</tr>
<tr>
<td>Antitoxins</td>
<td>Naloxone</td>
</tr>
<tr>
<td>Anesthetic agents</td>
<td>Pancuronium</td>
</tr>
<tr>
<td>Atracurium besylate</td>
<td>Phenobarbital</td>
</tr>
<tr>
<td>Atropine</td>
<td>Phenytoin</td>
</tr>
<tr>
<td>Calcium chloride</td>
<td>Propanol</td>
</tr>
<tr>
<td>Calcium gluconate</td>
<td>Sodium bicarbonate</td>
</tr>
<tr>
<td>Contrast media</td>
<td>Succinylcholine</td>
</tr>
<tr>
<td>Dexamethasone</td>
<td>Thiopental</td>
</tr>
<tr>
<td>Diazepam</td>
<td>Vecuronium</td>
</tr>
<tr>
<td>Diazoxide</td>
<td></td>
</tr>
<tr>
<td>Digoxin</td>
<td></td>
</tr>
<tr>
<td>Dobutamine</td>
<td>Dextrose solutions</td>
</tr>
<tr>
<td>Dopamine</td>
<td>Sodium chloride solutions</td>
</tr>
<tr>
<td>Ephedrine</td>
<td>Lactated Ringer’s solution</td>
</tr>
<tr>
<td>Epinephrine</td>
<td></td>
</tr>
<tr>
<td>Heparin</td>
<td>Blood and blood products</td>
</tr>
<tr>
<td>Insulin</td>
<td>Packed red blood cells</td>
</tr>
<tr>
<td>Levartemol</td>
<td></td>
</tr>
<tr>
<td>Lidocaine</td>
<td>Plasma</td>
</tr>
<tr>
<td>Lorazepam</td>
<td></td>
</tr>
</tbody>
</table>


has been shown to provide greater buffering capacity when administered by the IO route than by the peripheral IV route. Early IO access and infusion of epinephrine improved 24-hour survival in a swine model of ventricular fibrillation versus delayed IV administration. IO infusion of iodinated computed tomographic contrast material has also been reported as being successful, as has scorpion antivenin. In a porcine model of septic shock, IO and IV infusions had similar peak concentrations. Intraosseous infusion of intralipid reversed bupivacaine toxicity in a rat model. Cyanide toxicity reversal with hydroxycoalbumin has been shown to be equally efficacious between the IO and IV routes in a porcine model. Intraosseous administration has been successfully used for rapid sequence intubation. Intraosseous injection of iodinated computed tomography contrast agent in an adult blunt trauma patient has also been shown to be successful.

Voelckel and associates demonstrated that bone marrow blood flow responds to both the physiologic stress of hemorrhagic shock and vasopressors given during resuscitation after hypovolemic cardiac arrest in dogs. After successful resuscitation, bone marrow blood flow decreased after high-dose epinephrine but was maintained after high-dose vasopressin. These findings in animal models suggest the need for pressurized IO infusion techniques during hemorrhagic shock and certain drug therapies.

INDICATIONS

When children or adults need immediate resuscitation and IV access cannot be achieved quickly or reliably, the IO route provides a rapid and effective means of administering drugs, fluid, and blood. Once the patient has been stabilized, percutaneous peripheral or central intravascular access may be achieved.

Obtaining venous access can be a difficult task even under the best circumstances. This difficulty is compounded during high-stress situations or low-flow states such as cardiac arrest. Studies have shown that IO devices provide a rapid and effective means of fluid and drug administration during pediatric CPR. This is also true during resuscitation of critically injured infants and children in whom IO infusion of blood, colloids, or crystalloids (or any combination thereof) may be lifesaving. IO infusion is also beneficial in the management of children with other medical conditions, including those with respiratory distress, neurologic insults, dehydration, and sepsis. The widespread use of IO devices has led to improved prehospital vascular access and a marked reduction in critical transport with failed vascular access.

IO access is not commonly used in infants, but it is recommended as an alternative for medication and crystalloid administration when venous access is not readily obtained. IO infusion has been used with success in both premature and term infants. In one study, 30 IO lines were placed in 20 preterm and 7 full-term neonates with a variety of illnesses (e.g., respiratory distress syndrome, perinatal asphyxia, congenital cardiac anomalies) in whom conventional venous access had failed. All survived resuscitation with no long-term effects from IO line placement. Gestational age ranged from 32 to 41 weeks and birth weight ranged from 515 to 4050 g. In 2000 Abe and coworkers reported on the speed and ease of establishing newborn emergency vascular access by using turkey bones and plastic infant legs to simulate IO access and fresh umbilical cord to simulate umbilical venous catheterization. They demonstrated that for individuals who do not perform newborn resuscitation frequently, IO placement was easier and quicker than umbilical venous catheterization. A comparison of umbilical venous versus IO access in a simulated model of neonatal resuscitation showed that IO access was faster with no difference in error rate or perceived difference in ease of use.

IO infusion is also indicated for adult patients in whom attempts at peripheral or central venous access have been unsuccessful. This may include adult patients with burns, trauma, shock, dehydration, or status epilepticus. Intraosseous placement for resuscitation has been shown to be significantly faster than central line placement, with a significantly lower failure rate. Multiple sites including the iliac crest, femur, proximal and distal ends of the tibia, radius, clavicle, and calcaneus may be used. Of these, the tibia may be less desirable because red marrow is replaced by less vascular yellow marrow or fat by the fifth year of life. In contrast, the sternum has been advocated as the best site to establish IO access in adults because it is large and flat and can readily be located. In addition, the sternum’s cortical bone is thin (1 to 2 mm) and the marrow space relatively uniform (6 to 11 mm).

There has also been renewed interest in IO access by the US military. In addition to logistic constraints that limit the volume of isotonic crystalloid fluids available to resuscitate injured soldiers, hypotension, environmental and tactical conditions, and the presence of mass casualties can lead to excessive delay in obtaining vascular access. The Army Institute for Research has compared several IO infusion devices, including the FAST-1 IO Infusion System (PYNG Medical Corporation, Richmond, British Columbia, Canada), the Bone Injection...
insertion through areas of cellulitis, infection, or burns should also be avoided. Patients with right-to-left intracardiac shunts (e.g., tetralogy of Fallot, pulmonary atresia) may be at higher risk for fat or bone marrow embolization.69,70

### EQUIPMENT AND SETUP

The following is a review of products currently available for IO infusion. Information regarding the use of these products is limited, and few prospective studies have compared IO needles or devices in clinical practice. Until more information becomes available, practitioners are encouraged to review the products available and choose those that best meet their needs.

**IO Needles (Fig. 25.2)**

Needles used for IO access range in size from 13- to 20-gauge and must be sturdy enough to penetrate bone without bending or breaking. They must also be long enough to reach the marrow cavity. Standard needles used for drawing blood or administering medications are not adequate for IO infusions; they are not sturdy enough to penetrate bone and do not have a stylet to prevent bone from plugging the lumen. A cadaver study of IO puncture suggested that nonstyletted needles (2.5-cm, 18-gauge phlebotomy needles and 7.6-cm, 14-gauge IV needles) enter the marrow space successfully only approximately half the time.71 In the past, an 18-gauge spinal needle was commonly used in children younger than 12 to 18 months. This needle, although readily available in most emergency departments (EDs), often bends, is too long for rapid infusion of fluid, and has a greater risk for occlusion from clotted blood.72 Very small “butterfly” needles have been used with success in preterm infants.73

**Bone Marrow Aspiration Needle**

Bone marrow aspiration needles can be used for IO access range in size from 13- to 20-gauge and must be sturdy enough to penetrate bone without bending or breaking. They must also be long enough to reach the marrow cavity. Standard needles used for drawing blood or administering medications are not adequate for IO infusions; they are not sturdy enough to penetrate bone and do not have a stylet to prevent bone from plugging the lumen. A cadaver study of IO puncture suggested that nonstyletted needles (2.5-cm, 18-gauge phlebotomy needles and 7.6-cm, 14-gauge IV needles) enter the marrow space successfully only approximately half the time.71 In the past, an 18-gauge spinal needle was commonly used in children younger than 12 to 18 months. This needle, although readily available in most emergency departments (EDs), often bends, is too long for rapid infusion of fluid, and has a greater risk for occlusion from clotted blood.72 Very small “butterfly” needles have been used with success in preterm infants.73

**Illinois Sternal/Iliac Aspiration Needle**

The Illinois Sternal/Iliac Aspiration Needle (Monojet, Division of Sherwood Medical, St. Louis, MO) was designed for bone marrow aspiration but can be used for IO infusion. The needle

In addition to serving as a route for fluid administration, the IO needle may be used to obtain blood for typing, cross-matching, and determining blood chemistry in the marrow cavity. Serum electrolyte, blood urea nitrogen, creatinine, glucose, and calcium levels are very similar to those in samples obtained from an IO aspirate.64–66 Blood gas values obtained from the IO site were similar to those obtained from central venous sites during steady and low-flow states in one animal model.67 Brickman and colleagues68 demonstrated that bone marrow aspirates obtained from an IO needle in the iliac crest could be used reliably to type and screen blood for transfusion. A complete blood cell count may not be reliable because it reflects the marrow cell count rather than the cell count in the peripheral circulation. Furthermore, the aspirated blood usually clots within seconds, even if placed in a tube that contains heparin.

### CONTRAINDICATIONS

Relatively few contraindications to IO infusion exist. Osteoporosis and osteogenesis imperfecta are associated with a high potential for fracture; therefore unless absolutely necessary, the procedure should be avoided when these diagnoses are known. A fractured bone should be avoided because as fluid is infused, it increases intramedullary pressure and forces fluid to extravasate at the fracture site. This may slow the healing process, cause nonunion of the bone, or lead to a compartment syndrome. Similar extravasation of fluid can occur through recent IO puncture sites placed in the same bone. Hence, recent previous use of the same bone for IO infusion represents a relative contraindication to IO line placement. Needle
is available in both 16- and 18-gauge. It has an adjustable plastic sleeve to prevent the needle from penetrating through the opposite bony cortex. However, its long shaft and poorly designed handle make it prone to dislodgement during transport and other procedures.

**Jamshidi Disposable Sternal/Iliac Aspiration Needle**
Like the Illinois Sternal/Iliac Aspiration Needle the Jamshidi Disposable Sternal/Iliac Aspiration Needle (Cardinal Health, Dublin, OH) was designed for bone marrow aspiration, but it has a shorter shaft and smaller handle, which makes it easier to use. It comes in either 15- or 18-gauge and also features an adjustable plastic sleeve to prevent overpenetration. Once inserted the needle protrudes approximately 2 inches from the skin, which increases the risk for accidental dislodgement. In a study using a turkey bone model participants rated the Jamshidi needle easier to use than the Cook IO needle.74

**Cook IO Needle**
The Cook IO Needle (Cook Critical Care) is specifically designed for IO insertion and infusion. It comes in a variety of sizes from 18- to 14-gauge and can be inserted to a depth of 3 to 4 cm. It has a detachable handle, which reduces the risk of it being dislodged, and a depth marker to help ensure proper placement.

**Sur-Fast Needle**
The Sur-Fast Needle (Cook Critical Care) is also specifically designed for IO insertion and infusion. It has a threaded shaft that helps secure the needle in the bone and a detachable handle that may be reused with multiple needles. In a study by Jun and associates,75 the Sur-Fast IO needle had a success rate similar to that of a standard bone marrow aspiration needle.

**IO Devices**

**FAST-1 Intraosseous Infusion System (Fig. 25.3)**
The FAST-1 Intraosseous Infusion System (PYNG Medical) uses an impact-driven device designed for sternal placement only. The FAST-1 has been used successfully by both military and prehospital care providers.56,76 In one prehospital study, flow rates of 80 and 150 mL/min were obtained by using gravity and a pressure bag, respectively.77 In a prospective prehospital evaluation, the mean time to successful placement was 67 seconds.78 The device has a series of stabilizing probes that help maintain good contact with the sternum and serve as the depth control mechanism for insertion of the needle. These probes use the surface of the manubrium rather than the patient’s skin to ensure the proper depth of insertion. Once the device is positioned against the sternum, additional pressure triggers the release of a hollow needle into the medullary space. The needle comes preconnected to IV tubing. The handle is automatically released from the stylet and infusion tubing once the needle has met its preset depth. Removal of the needle requires a threaded tool provided with the device. The FAST-1 is larger and heavier than other IO devices and once triggered cannot be reused.

**Bone Injection Gun—BIG**
The BIG (PerSys Medical, Houston, TX) is another spring-loaded, impact-driven device that comes in both pediatric and adult sizes. (Fig. 25.4) Like the FAST-1 system, this device is designed for single use only. An advantage of the BIG is the ability to adjust the depth of insertion, which allows it to be used at different sites (e.g., tibia, humerus). However, if the device is not carefully stabilized before and during insertion, incorrect placement can easily occur. In addition, there is the potential for operator and patient injury if the device is accidentally triggered or mistargeted.79 In a prospective, prehospital setting the BIG device was shown to have a 71% success rate in children and a 73% success rate in adults.79 In another prehospital study, 91% of 181 patients had successful insertion on the first attempt.80

**EZ-IO Device (Fig. 25.5)**
The EZ-IO Device (Teleflex, San Antonio, TX) is a handheld, battery-powered device that drills an IO needle to the appropriate depth in the IO space. The EZ-IO device allows the operator
SECTION IV  VASCULAR TECHNIQUES AND VOLUME SUPPORT

Placement can be achieved in less than 10 seconds in the vast majority of patients, with first-time successful insertion rates ranging from 77% to 97%. In the United States, the EZ-IO has been approved for use at two anatomical sites, the proximal tibia and the humeral head. The EZ-IO has been shown to have a first pass success rate in the prehospital setting of 84% to 92%.

NIO Device
The New Intraosseous (NIO) device (PerSyS Medical) is a Food and Drug Administration (FDA)-approved device that is spring-loaded and used primarily for tibial insertion (Fig. 25.6). It can also be used in the humerus. It is lightweight, does not require batteries, and is a self-contained unit designed for a single use and ease of deployment. There is some early published evidence that suggests that it has comparable successful deployment rates for both tibial and humeral placement as the EZ-IO (Teleflex).

Comparative Trials Between Devices
The effectiveness, speed, and complication rates of manually placed versus mechanical drill-assisted IO catheters by emergency medicine resident physicians were compared in a 2013 study. The mechanical placement was shown to be significantly faster (3.66 vs. 33.57 seconds, \( P = 0.01 \)), the Jamshidi 15G needle, the FAST-1, and the BIG were compared in a helicopter-based emergency service system. In this randomized controlled trial, The Jamshidi needle and the FAST-1 had similar success and complication rates. However, the Jamshidi needle was placed significantly faster than the FAST-1. Frascone and colleagues conducted two sequential prospective nonrandomized trials to compare the FAST-1 with the EZ-IO, both in the prehospital setting. Of 178 IO insertions evaluated, 64 of the 89 FAST-1 insertions were successful as were 78 of the 89 EZ-IO insertions (72% vs. 87%). During the 5-year study period, 11 technical complications were recorded with the EZ-IO and 25 with the FAST-1. There were no differences in provider comfort or provider assessed device performance between the two devices. The BIG has also been compared to the EZ-IO in the adult population in two randomized trials. Twenty patients were treated with the BIG and 20 with the EZ-IO, with success rates in the first attempt of 85% and 90%. Mean procedure time was 2.2 min ± 1.0 for the BIG, versus 1.8 min ± 0.9 for the EZ-IO. Five technical problems were reported with the BIG, none with the EZ-IO. In five patients who were treated with the BIG, the stylet became stuck within the cannula and could only be removed with a clamp. In a large multi-year retrospective analysis of a trauma database, rates of success on first attempt were significantly higher with the EZ-IO compared with the manual needle and the BIG. Overall success rates were 50% with the manual needle, 55% with the BIG, and 96% with the EZ-IO.

Simulation Studies Comparing Devices
In a simulation cadaver model laboratory with US military providers, four IO devices were compared: the FAST-1, the BIG, the hand-driven threaded Sur-Fast needle, and the Jamshidi needle. High success rates were recorded for all four devices (94%–97%), but the BIG had the fastest deployment time. Frascone and colleagues conducted two sequential prospective nonrandomized trials to compare the FAST-1 with the EZ-IO, both in the prehospital setting. Of 178 IO insertions evaluated, 64 of the 89 FAST-1 insertions were successful as were 78 of the 89 EZ-IO insertions (72% vs. 87%). During the 5-year study period, 11 technical complications were recorded with the EZ-IO and 25 with the FAST-1. There were no differences in provider comfort or provider assessed device performance between the two devices. The BIG has also been compared to the EZ-IO in the adult population in two randomized trials. Twenty patients were treated with the BIG and 20 with the EZ-IO, with success rates in the first attempt of 85% and 90%. Mean procedure time was 2.2 min ± 1.0 for the BIG, versus 1.8 min ± 0.9 for the EZ-IO. Five technical problems were reported with the BIG, none with the EZ-IO. In five patients who were treated with the BIG, the stylet became stuck within the cannula and could only be removed with a clamp. In a large multi-year retrospective analysis of a trauma database, rates of success on first attempt were significantly higher with the EZ-IO compared with the manual needle and the BIG. Overall success rates were 50% with the manual needle, 55% with the BIG, and 96% with the EZ-IO.
paramedics, the BIG had the highest success at first pass (91%) versus the EZ-IO (82%) and the Jamshidi needle (47%).

**PROCEDURE**

**Sites for IO Needle Placement**

The patient’s age and size are the two most important factors when choosing the best site for needle penetration. In infants and children younger than 6 years, the proximal end of the tibia is the preferred site, followed by the distal ends of the tibia and femur. Other sites such as the clavicle and humerus have been used, but neither has gained popularity. In adults, the distal part of the tibia has been the most common site for IO access. However, with the introduction of spring-loaded and drill devices, IO locations once reserved only for children are now also potential sites in adults. In addition, the FAST-1 System makes the sternum a simple and effective location for IO access in adults.

**Proximal Tibia**

The tibia is a large bone with a thin layer of overlying subcutaneous tissue that allows landmarks to be palpated readily. Insertion here does not interfere with airway management or CPR. This is one of the most common infusion sites. On the proximal end of the tibia, the broad, flat, anteromedial surface is used, and the tibial tuberosity serves as a landmark. The site of IO cannulation is approximately 1 to 3 cm (2 finger widths) below the tuberosity (Fig. 25.7A). This location is far enough away from the growth plate to prevent damage, but it is in an area where the bone is still soft enough to allow easy penetration with the needle. In adults, penetrating the thick bone in the proximal end of the tibia is much more difficult and requires a 13- to 16-gauge needle. A spring-loaded device such as the BIG or a battery-powered drill such as the EZ-IO can make penetration much easier and allows the use of smaller-gauge needles. In a comparison of first-attempt success between tibial and humeral IO insertion during out-of-hospital cardiac arrest, tibial placement was significantly more successful (90%) than humeral placement (60%) with a lower rate of needle dislodgement.

**Distal Tibia**

The distal end of the tibia, although a preferred site in adults, may also be used in children. The cortex of the bone and the overlying tissue are both thin. The site of needle insertion is the medial surface at the junction of the medial malleolus and the shaft of the tibia, posterior to the greater saphenous vein (see Fig. 25.7B). The needle is inserted perpendicular to the long axis of the bone or 10 to 15 degrees cephalad to avoid the growth plate.

**Distal Femur**

The distal portion of the femur is occasionally used as an alternative site in children, but because of thick overlying muscle and soft tissue it is more difficult to palpate bony landmarks (see Fig. 25.7C). If chosen, the needle should be inserted 2 to 3 cm above the femoral condyles in the midline and directed cephalad at an angle of 10 to 15 degrees from the vertical.

**Humerus**

The proximal end of the humerus is a relatively new option for IO access, but it is well tolerated and easily accessed. The close proximity of the greater tubercle of the humerus to the heart provides rapid infusion of medication and fluid into the general circulation. The humerus has been shown to be an effective site relative to peripheral or central access in a prospective resuscitation model. A study evaluating flow rates through all three sites in a porcine model showed that the proximal humerus had almost double the flow rate than either the tibial or distal femur site. In a retrospective study in a prehospital setting of cardiac arrest, IO insertion into the humerus had a higher success rate than into the proximal tibia. However, in another study first pass success for humeral placement in a paramedic performed prehospital population...
during cardiac arrest was over 90%. The patient’s arm should be placed such that the hand is resting on the patient’s umbilicus, with some adduction. Medial rotation of the humerus will allow better identification of the proximal humeral head. The humerus is most easily palpated at the insertion point for the deltoid muscle, between the biceps and triceps muscles. This point is approximately midway along the length of the arm. Palpation of the bone requires pressure because of overlying structures. The surgical neck can be located by palpating up the length of the humerus until the clinician feels a notch or groove. The appropriate insertion site is approximately 1 cm above the surgical neck for most adults (see Fig. 25.12). A study evaluating the humeral head approach during tactical situations was shown to have a 96% first pass success rate using an EZ-IO, and a 90% success rate using a manual driver.102

**Sternum**

The sternum has several advantages over peripheral bones but is rarely used in the ED. Its advantages include a large, relatively flat body that can be readily located; retention of a high proportion of red marrow, which allows rapid transfer of infused fluids and drugs to the central circulation; and thinner, more uniform cortical bone overlying a relatively uniform marrow space. In addition, the sternum is less likely to be fractured in major trauma.**76** Introduction of the FAST-1 System, which allows safe and effective penetration of the sternum, has led to increased use and popularity of sternal IO insertion in adults.

**Other Sites**

Other sites, including the clavicle and calcaneus, can be used as alternatives, but these sites are less popular.

**Site Preparation**

To prepare the proximal end of the tibia or distal end of the femur for IO insertion, place a small support such as a towel roll behind the knee. Cleanse all insertion sites with chlorhexidine, povidone-iodine, or an alcohol-based antibacterial solution. If the patient is conscious, anesthetize the skin and periosteum.

**Manual Needle Insertion (Fig. 25.8)**

Before insertion stabilize the site with the free hand (i.e., the hand not holding the IO needle) and use it to identify the landmarks. For example, during the proximal tibia insertion, stabilize the proximal end of the tibia with the thumb and index finger of the free hand and use them to palpate the tibial tuberosity (the main bony landmark for proximal tibia insertion). During insertion avoid injury to this hand by keeping it out of the plane of insertion and clear of the puncture site. Direct the IO needle perpendicular (90 degrees) to the bone’s long axis and slightly caudal (60 to 75 degrees). Directing the needle slightly caudal helps avoid penetration of the growth plate. Advance the needle with a twisting or rotating motion (but not a rocking motion) to drive it into the bone and to puncture the cortex. Once the cortex has been penetrated, there will be a sudden decrease in bony resistance and a “crunchy” feeling as the needle enters the marrow cavity. Penetration of the inner cortex usually occurs at a depth of approximately 1 cm.

Aspirate for blood or marrow contents (or both) to confirm correct placement; however, aspiration may not always produce blood even with a properly placed IO needle. Other signs of correct placement include the needle’s ability to remain upright without support and to have free-flowing fluid without signs of extravasation into surrounding tissue. Note that some resistance to flushing is expected because the bone marrow is not distensible.103,104

Once proper placement is confirmed, secure the needle and tubing with tape. Fasten the leg to an appropriately sized leg board to further stabilize a lower extremity insertion site in infants and small children. Protect the needle from accidental dislodgement by cutting the bottom out of a plastic cup and taping and bandaging the cup in place over the device. Commercially made shields are also available for this purpose. Remove the IO needle as soon as IV access has been secured, and apply a sterile dressing over the site. Control excessive bleeding by applying direct pressure over the site for 5 minutes.105

**USE OF SPECIFIC IO DEVICES**

**FAST-1 (Fig. 25.9)**

The FAST-1 device (PYNG Medical) was designed specifically to penetrate the sternum.62,106 It is prepackaged with alcohol and iodine and comes with a protective dressing to hold the device in place. It also includes a threaded tip remover for easy removal of the metal tip and infusion tubing (see instruction video at: [http://www.pyng.com/fast1/](http://www.pyng.com/fast1/)).

After disinfecting the skin site over the sternum, place the target patch over the midline of the manubrium with the hole in the middle of the target approximately 1.5 cm below the sternal notch. Next, place the FAST-1 introducer in the center of the target zone. The introducer has a “bone cluster” of needles that form a circle. These needles “sense” the cortex of the sternum and help ensure proper needle depth. Once the introducer is in position over the target zone, apply pressure to the handle to release the inner needle located in the center of the bone cluster. This needle has a small metal tip that is preconnected to plastic infusion tubing. After release, the central IO needle will advance 5 mm beyond the circular cluster of needles, stop at the bony cortex, and position the metal tip at the cortex-medullary junction. At this point withdraw the handle so that only the plastic infusion tube is left protruding from the insertion site. Marrow aspiration and rapid flow of fluid help verify the appropriate position. Attach the plastic dome to the target patch via Velcro fasteners and secure the tubing in place. Removal of the infusion tube requires the use of a threaded-tip remover, which is included. The tube can also be removed by directly pulling it; however, the metal tip is sometimes left behind and must be extracted through a small incision.61

**BIG (Fig. 25.10)**

The BIG (Waismed) incorporates a loaded spring to facilitate penetration of the bone (see instruction video at: [http://www.waismed.com/BIGmovie.html](http://www.waismed.com/BIGmovie.html)). To adjust the depth of insertion remove the safety pin from one end and turn the other end clockwise or counterclockwise to reduce or increase needle depth, respectively. Place the BIG firmly against the skin perpendicular or slightly caudal to the long axis of the bone. Fire the gun by applying palmar force on the back of the unit and pulling on the flanges with the middle and ring fingers. Confirm placement by aspirating marrow, flushing with the same syringe, and observing flow through the IV
Manual Intraosseous Needle Insertion

1. Cleanse the skin with antiseptic.

2. Stabilize the leg and identify the tibial tuberosity with your free hand. Insert the needle perpendicular or slightly caudad with a twisting or rotating motion.

3. Remove the stylet once the cortex is penetrated (feel for a decrease in resistance and a “crunchy” feeling).

4. Aspirate blood and/or marrow to confirm the correct position. If the patient is awake, very slowly infuse 2–5 mL of 2% lidocaine a few minutes prior to infusion. Flush the needle with 10 mL of saline to prime the infusion.

5. Connect the IV tubing to the needle.

6. Secure the IO needle in place with tape.

7. Protect the needle from dislodgement with a plastic cup in which the bottom has been cut out, and then tape and bandage if in place over the needle.

Figure 25.8 Manual intraosseous (IO) needle insertion. IV, intravenous.
tubing. Slide the slotted safety pin into the needle to maintain stability. To remove the needle, rotate it back and forth using the small clamps provided with the unit. Dress the site in a manner determined by the care provider.

**EZ-IO Needle (Figs. 25.11 and 25.12)**

This battery-operated “drill” can drive the IO needle through thick bone with relative ease. The EZ-IO kit comes with a battery-operated drill and an IO needle with a stylet. The EZ-IO AD (adult) comes with a 15-gauge, 25-mm IO needle for use in patients heavier than 40 kg, and the EZ-IO PD (pediatric) comes with a 15-gauge, 15-mm needle for use in patients lighter than 39 kg. To operate the drill, insert the needle into the driver tip and make sure that it is securely seated onto the drill. Remove the safety cap from the needle and position the drill perpendicular (or slightly caudad) to the insertion site. While applying gentle pressure, squeeze the trigger to penetrate the skin. When the tip of the needle comes in contact with the bone, at least 5 mm of the IO catheter should be visible. If not, the overlying soft tissue may be too deep for the needle to enter the marrow cavity. To penetrate the bone, continue to squeeze the trigger while applying steady downward pressure until a sudden “give” or “pop” occurs, which signals entry into the medullary space. Too much pressure on the device can cause the drill to stall and prevent the needle from penetrating the cortex.

After entry into the marrow cavity, attach the EZ-Connect extension set provided with the EZ-IO kit and aspirate blood and bone marrow contents to confirm correct placement. Once catheter placement has been checked, fluids or medications can be infused. Avoid attaching syringes and IV tubing directly to the IO needle because this can enlarge the hole in the cortex and result in extravasation of fluid. Secure
while holding the needle stabilizer against the insertion site. Continue holding the needle stabilizer in place and pull up the stylet to remove.

**NIO Needle (Fig. 25.13)**

Place the device approximately 2 cm medial and 1 cm proximal to the tibial tuberosity. Unlock the NIO by rotating the cap 90 degrees in either direction. Place the dominant hand over the cap and press the device against the patient. While pressing down on the device with the palm, pull trigger wings upwards with fingers. Gently pull the NIO up in a rotating motion while holding the needle stabilizer against the insertion site. Continue holding the needle stabilizer in place and pull up the stylet to remove.

**COMPLICATIONS**

**Technical Difficulties**

Technical difficulties are the most common complications, but they decrease as familiarity with the technique increases.

**Figure 25.10** Use of the Bone Injection Gun (BIG) (Waismed, Yokenam, Israel).
Connect the appropriate needle set to the driver (pink 15 mm = patients 3–39 kg; blue 25 mm = patients > 40 kg; yellow 45 mm = proximal humerus, patients with excessive tissue over the site).

Cleanse the site with an antiseptic agent per institutional protocol.

Remove the power driver from the needle set while stabilizing the catheter hub, and then remove the stylet by turning it counterclockwise.

Attach the EZ-Connect tubing to the Luer-Lok adapter on the catheter hub. Confirm placement by aspirating blood and/or marrow contents.

Position the driver at the insertion site with the needle at a 90-degree angle to the surface of the bone. Gently pierce the skin with the needle set and advance needle without using the trigger yet, until the needle set tip touches the bone.

Penetrate the bone cortex by now squeezing the driver’s trigger and applying gentle, consistent, steady downward pressure. Allow the driver to do the work.

Remove the power driver from the needle set while stabilizing the catheter hub, and then remove the stylet by turning it counterclockwise.

Attach the EZ-Connect tubing to the Luer-Lok adapter on the catheter hub. Confirm placement by aspirating blood and/or marrow contents.

If the patient is responsive to pain, consider slowly administering plain 2% lidocaine for anesthetic effect before flushing with 10 mL of saline. Begin the infusion with a pressure bag.

Remove the device within 24 hours. Connect a sterile Luer-Lok syringe to the catheter hub, and rotate clockwise while pulling straight up. Avoid rocking the needle on removal.

Figure 25.11 Use of the EZ-IO device (Teleflex, San Antonio, TX).
The most common mistake is to place excessive pressure on the needle during insertion and force it entirely through the bone and out the other side (Fig. 25.15). Minimize this risk by using appropriate landmarks and keeping the needle perpendicular to the long axis of the bone. In addition, hold the needle with the index finger approximately 1 cm from the bevel. When this finger touches the skin, the needle should be in the marrow cavity and no further pressure needs to be applied. Some IO needles have a mark 1 cm from the bevel (e.g., Cook IO Needle), whereas others have a special guide or mechanism to ensure proper insertion and depth of penetration (e.g., Illinois Sternal/Iliac Aspiration Needle). If available, use of these adjuncts will also help prevent overpenetration.
At times the needle appears to be in the marrow cavity, but blood or bone marrow cannot be aspirated and fluids do not flow freely. This may follow incomplete penetration of the bone or overpenetration into the opposite cortex. Incomplete penetration usually results in extravasation of fluids and can be corrected by replacing the stylet and slowly advancing the needle until successful aspiration of marrow contents and free flow of fluids occur. Penetration into the opposite cortex generally results in little or no flow. If overpenetration is suspected, pull the needle back 1 to 2 mm and check for free flow of fluids.

To ensure flow, rapidly inject 10 mL of saline into the marrow. This is a painful procedure in awake patients, but failure to initially flush the compartment is a common reason for inadequate flow. A pressurized bag system is suggested if large volumes of fluid are administered. Flush each dose of medication with 3 to 5 mL of saline. Fluids that initially flowed freely may stop flowing if the needle becomes clogged by blood clots or bone spicules. Flush the needle frequently with 3 to 5 mL of saline to help avoid this problem. If none of these maneuvers results in free flow of fluid, remove the needle and attempt insertion in the opposite extremity or another site. This helps avoid fluid extravasation through the hole that is left after removing the needle.

Extravasation of fluid is less common but may be associated with a number of adverse events. Extravasation may be caused by fluids being infused under excessive pressure and with prolonged use of an IO site. As noted previously, extravasation may also result from incomplete needle penetration or penetration through the opposite cortex. Even when an IO needle has been positioned properly, fluid can leak out through holes made by previous IO attempts or through an insertion site that was made too large by “rocking” during insertion or from an improperly secured needle that becomes loose with movement. Interestingly, the type of needle used does not appear to influence extravasation rates. Regardless of the cause, if extravasation occurs, remove the needle quickly and apply pressure to the site. If left unchecked, extravasation can lead to a number of adverse events (see later section on Soft Tissue and Bony Complications). In addition, though not directly harmful, extravasation of fluid through multiple cortical defects from previous IO attempts has been associated with lower serum levels of infused drugs.

### Soft Tissue and Bony Complications

#### Infection

In the past, concerns about infection led clinicians to shy away from using the IO route. Although the potential for infection is real, its actual incidence is low. A literature review of more than 4000 cases from 1942 to 1977 found a 0.6% infection rate. Although most of the affected access sites were not placed under emergency conditions, the needles were often left in place for 1 to 2 days, thus increasing the likelihood of infection. A survey of more than 1000 US and foreign medical schools found that the incidence of infection for IO needles placed in emergency conditions was less than 3%. The most common infection is cellulitis at the puncture site, which usually responds well to antibiotics. Osteomyelitis is less common but also usually responds well to antibiotics. Heinild and coworkers reported three cases of osteomyelitis in 25 patients who received infusions of undiluted 50% dextrose in water. More recently, Platt and coworkers reported a case of fungal osteomyelitis and sepsis secondary to an IO infusion device. A case of tibial osteomyelitis with IO abscess has also been reported.

In addition to infection, inflammatory reactions of the bone may be seen. Such reactions are most common when hypertonic or sclerosing agents are used and may produce an elevation of the periosteum on plain radiographs or a positive bone scan (Fig. 25.16). Unlike patients with osteomyelitis from bacteria, a child with a sterile inflammatory reaction should not appear ill. One hypertonic sclerosing drug that may be used during cardiac arrest is sodium bicarbonate. Heinild and coworkers reported 78 cases of bicarbonate infusion with no complications. Animal studies have reported a decrease in cellularity with edema and destruction of some cells, but these changes are temporary and resolve completely in a few weeks. A case of systemic fibrinolysis through IO vascular access in a patient with ST-elevation myocardial infarction has also been reported.

#### Skin Sloughing

Skin sloughing and myonecrosis have been reported secondary to extravasation of infused fluids and medications. This occurs if fluid or drugs extravasate from the puncture site into...
CHAPTER 25  Intraosseous Infusion

it is appropriate to take follow-up radiographs of patients who have undergone IO needle attempts or placement. Cortical defects may be seen on radiographs for up to 40 days after injection.130

**Fat Embolism**

Fat embolism is another potential complication of IO insertion.3,72 However, this condition is rare and has been reported only in adult patients.65 Animal studies have found no changes in blood gases during IO infusion and limited evidence of fat globule collections in the lungs.69,70 In a swine cardiac arrest model, there was no difference in the risk for fat emboli in pigs that had an IO line inserted versus those receiving IV medications.131,132 Because the marrow in infants and children is primarily hematopoietic, this potential complication is unlikely to occur in this population.

**Pain With Infusion**

Most patients undergoing IO infusion will not be in a condition to sense pain, but infusion into the bone marrow can be quite painful. Slowly infusing 2 to 5 mL of 2% preservative-free lidocaine (0.5 mg/kg), over 1 minute before flushing and infusion of medications and fluids, has been suggested to relieve pain in awake patients. The duration of analgesic effect is approximately 1 hour and may be repeated as needed. A maximum dose of lidocaine per hour is 200 mg. Systemic opioids are also helpful. Medications intended to remain in the medullary space, such as local anesthetics, must be injected very slowly until the desired anesthetic effect is achieved. A 2010 study suggests that lidocaine is efficacious in attenuating the pain related to IO infusion.133

**TRAINING**

Procedural training can be performed on simulation models, animal bones (e.g., chicken or turkey legs), or cadavers. IO insertion techniques are part of the curriculum of the pediatric advanced life support, advanced pediatric life support, and advanced trauma life support courses. Training typically consists of an hour of didactics followed by a hands-on session. Novice users have achieved success rates of 93% to 97% for manual and battery-powered IO placement in a variety of simulated settings.5,22,48,55,79,134,135 Battery-powered IO placement has also been shown to have high success rates after training.

**DURATION OF USE**

IO cannulation is temporary vascular access and should be replaced with a venous line as soon as possible. IO cannulation may be left in place during transport and for up to 24 hours, but local edema, infiltration, and an increased risk for osteomyelitis are seen after 24 hours.

**Acknowledgment**

The author would like to sincerely thank Rachael Stanley, MD, for her contribution to this chapter in previous editions.
REFERENCES


Alternative Methods of Drug Administration

Solomon Behar

The rapid administration of lifesaving, pain-relieving, and sedative medications lies at the core of the practice of emergency medicine. The intravenous (IV) route is usually the delivery method of choice. However, there are circumstances in which vascular access is either not available or contraindicated. Thus, emergency providers need to have a working knowledge of alternative routes of drug administration. This chapter describes the endotracheal (ET), intranasal (IN), rectal, and subcutaneous routes. Intraosseous (IO) access is covered in Chapter 25.

ENDOTRACHEAL (ET) ADMINISTRATION OF MEDICATION

Certain drugs can be delivered simply, rapidly, and effectively to the central circulation by way of the ET tube. Because the efficacy of ET medication administration is not clear, this method is best reserved for situations in which a patient’s condition warrants immediate pharmacologic intervention but more conventional means of drug delivery, such as IV and IO, are not readily available. Such circumstances frequently arise in the prehospital or cardiac arrest setting. Knowledge of the appropriate drugs and dosages that can be delivered effectively by this route may prove to be lifesaving.

Historical Perspective

ET drug administration dates to 1857, when Bernard demonstrated that the lung could rapidly absorb a solution of curare. In this historical experiment, he instilled a fatal solution into the upper respiratory tract of dogs by way of a tracheostomy. Over the following decades, other investigators expanded this work and demonstrated that solutions containing salicylates, atropine, potassium iodide, strychnine, and chloral hydrate were also rapidly absorbed from the lung and excreted in the urine after injecting aqueous solutions into the tracheas of experimental animals. The use of intrapulmonary medication for the treatment of lung disease gained further acceptance when studies demonstrated that inhaling epinephrine mist dramatically relieved the symptoms of asthma.

In the late 1930s and 1940s, several important observations were made concerning ET drug therapy: (1) penicillin delivered by the ET route demonstrated a depot effect, which resulted in therapeutic blood levels that lasted twice as long as those with intramuscular (IM) injection; (2) various diluents mixed with penicillin affected both the rate and the degree of absorption from the lungs; and (3) higher serum drug levels were attained with direct ET drug administration than with aerosolized administration. In the 1950s it was noted that drugs delivered endotracheally were absorbed much more rapidly than those applied to the posterior part of the pharynx. Drugs applied locally to the larynx and trachea were absorbed rapidly and even resulted in blood levels significant enough to cause adverse anesthetic reactions.

Endotracheal Medication Administration

**Indications**
- Cardiac arrest with no peripheral or intraosseous access

**Contraindications**
- Peripheral or intraosseous access

**Complications**
- Depot effect of drug
- Transient hypoxemia
- Loss of catheter or needle down endotracheal tube

**Equipment**
- Bag-valve mask
- Medication (epinephrine, lidocaine, atropine, naloxone, flumazenil)
- Syringe (10-20mL adult, 5mL pediatric)
- Diluent (saline or water)
- Endotracheal tube
- Long small-bore catheter or 8-Fr feeding tube (pediatric)
- 18-gauge needle
- 18-gauge spinal needle
- IV adapter lock

Review Box 26.1 Endotracheal medication administration: indications, contraindications, complications, and equipment. Gloves, mask, and eye protection are also necessary equipment for endotracheal medication administration. See text for further explanation on medication.
In 1967, Redding and coworkers\textsuperscript{7} studied the use of ET administration as a route of drug delivery in a canine model of cardiopulmonary arrest. They administered epinephrine by the IV, intracardiac, and intratracheal routes to resuscitate dogs that had undergone both respiratory and circulatory arrest secondary to hypoxia. They then evaluated the effectiveness of the epinephrine after administration of the drug by all three of these routes. Their study revealed that all three routes of drug administration were equally effective in restoring the circulation of dogs in hypoxia-induced cardiac arrest, again demonstrating that the ET route of drug delivery provides effective access to the systemic circulation.

In the late 1970s, Roberts, Greenberg, and colleagues\textsuperscript{8–11} studied ET drug delivery in a series of laboratory experiments and clinical applications of ET epinephrine. Since that time, a number of important animal and human studies, as well as case reports, have been published in which the various aspects of ET drug administration were investigated. These studies have addressed (1) the appropriate dose of drug to administer; (2) the effect of the drug solution’s volume; (3) the effect of different diluent solutions; (4) the role of different ET drug delivery techniques; and (5) the effects of hypoxia, hypotension, shock, and cardiopulmonary arrest on the absorption, distribution, and efficacy of endotracheally administered drugs.

**Recommendations for ET Drug Delivery**

ET drug delivery is not the delivery method of choice if other routes are available. The American Heart Association (AHA) recommends that if IV access is not available, IO access should be obtained.\textsuperscript{12} The AHA makes specific recommendations regarding the use of ET drug delivery for cardiac resuscitation (Table 26.1).\textsuperscript{12–14}

**Appropriate Dose**

Much of the existing literature is controversial and contradictory at times in regard to proper dosing of ET medications.\textsuperscript{10,15–19} The ET dose of a medication should be at least equal to the IV dose of the same drug when given for the same indication, but most studies agree that higher doses are needed when administering drugs via the ET route. For advanced cardiac life support (ACLS) medications in adults, the AHA recommends a dose that is 2 to 2.5 times the usual IV dose when administered via an ET.\textsuperscript{12,20–22} Studies using both normotensive and cardiac arrest canine models have shown that epinephrine doses of 0.01 mg/kg administered via ET produced serum levels approximately one tenth of that produced when the same dose is given intravenously.\textsuperscript{9,23,24} These studies suggested increasing the ET epinephrine dose to 0.05 to 0.1 mg/kg of the 1:1000 dilution of epinephrine, and are the basis for the 2010 AHA recommendation (upheld in a 2015 update\textsuperscript{25}) to use this tenfold increased dose when administering ET epinephrine to pediatric patients.\textsuperscript{11} One notable exception of epinephrine concentration is in the neonatal population (the first month of life) in which the AHA recommends using the 1:10,000 (not 1:1000 that is used in pediatric patients and adults) concentration of epinephrine using a dose of 0.01 to 0.03 mg/kg.\textsuperscript{26}

ET drug delivery is associated with a “depot” effect, with ET drugs being “stored” and released slowly over time, similar to a continuous IV drip. This presumably occurs as a result of local vasoconstriction and lymphatic storage of the drug\textsuperscript{6} or pooling in lung tissue because of poor lung perfusion.\textsuperscript{27} With ET epinephrine use, the depot effect can produce post-resuscitative dysrhythmias, hypertension, and tachycardia together with resultant increased myocardial oxygen demand. Given these conflicting data, it seems reasonable in adults to start with a dose 2.0 to 2.5 times the usual IV dose. If this is ineffective, higher doses may be used subsequently.

**Volume for a Single Dose**

For ET drugs, the AHA recommends a total volume of 10 mL in adults,\textsuperscript{12} 5 mL in pediatric patients,\textsuperscript{13} and 1 mL in neonates.\textsuperscript{14} Animal and human studies demonstrate that diluting medications in higher volumes of liquid increases blood levels of the drug, but at the expense of decreases in partial pressure of arterial oxygen (PaO\textsubscript{2}).\textsuperscript{28–31} Data and volume recommendations in the setting of multiple doses of the drug are lacking. It may be difficult or impossible to limit the total volume of the drug solution to 10 mL when using prefilled syringes. Prefilled syringes of epinephrine contain 1 mg in 10 mL (1:10,000). Giving 2.5 times the IV/IO dose requires the administration of 20 to 25 mL. It is possible to obtain epinephrine 1:1000 (1 mg/mL) and dilute it to a total volume of 5 to 10 mL, but the higher concentration of epinephrine may not be readily available during a cardiac arrest code. Likewise, prefilled syringes of atropine contain 1 mg in 10 mL (0.1 mg/mL). Prefilled syringes of lidocaine contain 20 mg/mL (100 mg/5 mL). A dose of 2 to 2.5 times the IV/IO dose could easily amount to 20 mL or more total volume in an obese patient.

**Appropriate Diluent**

Both normal saline and distilled water may be used as diluents for ET drug administration, but it remains unclear which is preferred. Normal saline may produce less pulmonary dysfunction than distilled water, but distilled water appears to deliver a greater amount of drug than when the drug is mixed in normal saline.\textsuperscript{29,30,32,33}

---

**TABLE 26.1** American Heart Association Guidelines for Endotracheal Drug Administration

<table>
<thead>
<tr>
<th>GUIDELINE</th>
<th>ADULT\textsuperscript{a}</th>
<th>PEDIATRIC\textsuperscript{b}</th>
<th>NEONATAL\textsuperscript{c}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medication dose</td>
<td>2–2.5 times the recommended intravenous dose</td>
<td>Epinephrine, 0.1 mg/kg (10 times the recommended intravenous dose) Atropine, 0.04–0.06 mg/kg Lidocaine, 2–3 mg/kg</td>
<td>Epinephrine (1:10,000) 0.05–0.1 mg/kg (class indeterminate)</td>
</tr>
<tr>
<td>Total volume to instill</td>
<td>10 mL</td>
<td>5 mL</td>
<td>1 mL</td>
</tr>
<tr>
<td>Diluent</td>
<td>Normal saline or distilled water</td>
<td>Normal saline</td>
<td>Normal saline</td>
</tr>
</tbody>
</table>
Technique for ET Drug Delivery

Techniques for ET drug administration include direct instillation into the proximal end of the ET tube, administration via a catheter that extends just beyond the distal tip of the ET tube, deep endobronchial administration using a longer catheter, administration via ET tube monitoring ports, administration with equipment developed specifically for ET atomized drugs, and injection through the side of the ET tube with a needle. Several studies have indicated that the use of a catheter or feeding tube may not be needed to enhance the drug's effectiveness, including neonates. In studies of patients with normal perfusion, some support “deep bronchial” ET drug administration, whereas others do not. Some suggest that drug absorption with direct instillation into the ET tube is inconsistent during cardiopulmonary arrest, and one study found no difference in plasma epinephrine levels when epinephrine was instilled during apnea versus instillation during the ventilator inspiratory cycle. Given these conflicting studies, use of a catheter to enhance deep pulmonary delivery seems reasonable. However, if a catheter is not readily available, direct injection into the ET tube appears to be acceptable.

Effects of Hypoxia, Hypotension, and Cardiopulmonary Arrest

Despite concerns that medications might not be absorbed in states of hypoxia or low blood flow, the data available reveal the opposite to be true. In a hemorrhagic shock model, Mace demonstrated that higher plasma lidocaine levels were obtained via the ET route during shock than during non-shock states. In a lamb model, when epinephrine was administered endotracheally, higher plasma epinephrine levels were achieved during hypoxia-induced low pulmonary blood flow than during baseline, normal pulmonary blood flow. Finally, plasma lidocaine levels rose earlier when lidocaine was administered endotracheally to dogs that were hypoxemic than to dogs that were not.

Indications

ET drug therapy is indicated when emergency pharmacologic intervention is needed and other access, either IV or IO, is not available. Specific indications for the delivery of a drug endotracheally are the same as those for IV and IO administration. However, only a limited number of emergency drugs can be given safely by the ET route. Medications that are appropriate for ET administration based on animal and human studies include epinephrine, atropine, lidocaine, and naloxone. Diazepam has also been shown to be effective, but with the possibility of a medication (or diluent) induced pneumonitis. The AHA has removed diazepam from its list of medications that can be given safely via the ET route.

Experimental studies of vasopressin, midazolam, fluoromazenil, and propranolol in animal models suggest that these medications may also be effective when administered endotracheally, but no clinical studies in humans have been conducted to verify these findings. Based on a study by Wenzel and colleagues, the 2010 ACLS guidelines added vasopressin to the list of cardiac resuscitation drugs that can be administered via the ET route. However, vasopressin was removed from the 2015 ACLS out-of-hospital cardiac arrest guidelines due to equivalency of response to epinephrine and a want to simplify arrest algorithms. As midazolam is approved for IM use, it seems unlikely that ET administration would be frequently used. Palmer demonstrated that therapeutic blood levels of flumazenil were obtained within 1 minute after ET delivery of 1 mg of the drug diluted in 10 mL of saline, which is 10 times the recommended IV dose of 0.1-0.2 mg aliquots.

Contraindications

At present, the only true contraindication to the ET delivery of an appropriate drug is the presence of another form of access to the systemic circulation through which the needed drug can be delivered rapidly and effectively. Specific medications have been shown to be ineffective or unsafe when given via the ET route. Sodium bicarbonate and amiodarone have direct deleterious effects on the lungs and isoproterenol have not been shown to attain adequate serum levels despite high ET doses given.

Equipment

The patient must first be endotracheally intubated. In studies in which the recommended ET tube doses of medications were administrated by Combitube (Kendall-Sheridan, Argyle, NY) or laryngeal mask airway (LMA North America, San Diego, CA), absorption of drugs was found to be subtherapeutic. A Combitube, when placed in the esophagus (requiring medications to travel out the side holes to reach the trachea), needs 10 times more epinephrine than that used with an ET tube to obtain the same serum concentration and hemodynamic effects. Presumably, a Combitube that enters the trachea directly would function equivalently to an ET tube, but no studies have been done to support this assumption.

The equipment listed here is that required to perform any of the four different techniques described. This equipment is suggested for the ideal situation; at no time should drug delivery be delayed while searching for the “perfect” piece of equipment.

1. Manual bag ventilation device capable of delivering a fraction of inspiratory oxygen (FiO₂) of at least 50%. When ET drug delivery is indicated, the patient’s condition almost always warrants supplemental oxygen. Although the technique may not result in any significant deterioration in respiratory function, it is still advisable to administer additional oxygen after drug delivery. Use the bag ventilation device to also deliver several rapid insufflations immediately after drug delivery to assist in delivery of the drug distally, where it may be absorbed more rapidly and effectively.

The priorities of drug administration via the ET route must be balanced against the potential deleterious effects that such rapid insufflation might have on hemodynamics and cerebral perfusion. Excessive hyperventilation of victims of out-of-hospital cardiac arrest is common and associated with poor outcomes.

2. A fine-bore catheter or special ET tube designed to deliver the drug at or beyond the distal end of the ET tube. For adults, select a catheter that is at least 8 Fr in size and 35 cm (14 inches) in length. It should be long enough to protrude past the distal end of the ET tube. The diameter of the catheter should be large enough to allow rapid delivery of 10 mL of solution. In children, the distance to the end of the ET tube is usually equal to three times the circumference of the tube in centimeters measuring from the lip. So for example, in a child intubated with a 4.0 ET tube, make sure the catheter tip lies at least 12 cm distal to where the
ET tube enters the mouth. Several different types of tubes and catheters commonly available in the emergency department (ED) can be used for this purpose:

a. A 16-gauge central venous pressure or cutdown catheter. Because most are only 30 cm in length, the proximal end of the ET tube should be shortened so that the catheter can protrude past the end.

b. An 8- or 10-Fr polyethylene pediatric feeding tube (e.g., Argyle, St. Louis). These tubes are much longer than needed, so cut them to reduce dead space. Luer-Lok ends fit onto the proximal end of the tube. For neonates, use a 5-Fr feeding tube with a syringe and an IV adapter.

c. An 8-Fr (or larger) pediatric pulmonary suction catheter without the control port. Because this catheter is designed to extend past the tip of the ET tube, it is an ideal length. However, with some brands it is difficult to attach a syringe or IV adapter lock after the suction control port is removed.

Alternatively, some ET tubes are made with built-in ports that allow the instillation of drugs without removing the bag ventilation device.

3. An IV adapter lock. This can be placed as needed onto the proximal end of the irrigation lumen of the Hi-Lo Jet Tracheal Tube (Nellcor, Pleasanton, CA) or on the catheters described previously to convert them for use with prefilled syringes. This adapter is generally unnecessary if a standard syringe is used.

4. A 10 to 20 mL syringe, preferably a Luer-Lok type, large enough to deliver the desired volume of drug solution plus an additional 5 mL of air. Most of the medications now prescribed for emergency situations come in prefilled syringes. This type of apparatus does not usually allow one to draw up diluent or an additional volume of air to empty the syringe of solution. In addition, depending on the manufacturer and model, some prefilled syringes have either needles or a needleless system that may require an IV adapter lock to use them for ET injection.

5. Diluent solution. Keep an adequate volume of diluent available, such as normal saline or distilled water.

6. Medication to be instilled.

7. An 18- or 19-gauge needle to draw up the medication and inject it. Use an 18-gauge, 8.9-cm (3.5-inch) spinal needle for direct instillation of medications into the proximal end of the ET tube.

8. Alcohol wipes to clean the vials and injection ports.

9. Gloves, mask, and eye protection. After instillation, the solution often refluxes out of the ET tube, which makes blood and body fluid precautions critical.

**Procedure**

The procedure of choice is one that will deliver the medication to the patient in the least amount of time. Secure the ET tube before instilling medications endotracheally to prevent the tube from being expelled if the patient coughs. Inflate the cuff of the tube, if present.

**Direct Instillation Into the ET Tube**

While the patient is being ventilated, draw up the desired drug into a syringe (or use a prefilled syringe) (Fig. 26.1, step 1). Dilute the drug to a final volume of 10 mL (adults), 5 mL (children), or 1 mL (neonates) with normal saline or distilled water. Attach an 18- or 19-gauge needle. Some authors recommend using an 8.9-cm (3.5-inch) spinal needle. If using a prefilled syringe, draw up an appropriate volume of diluent in a second syringe so that the total instillation volume (drug plus diluent) equals 10 mL (adults), 5 mL (children), or 1 mL (neonates). Attach an 18- or 19-gauge needle which will be used to flush the ET tube after instillation of the drug.

Interrupt the connection between the proximal end of the ET tube and the bag ventilation device. Insert the needle of the syringe into the proximal opening of the ET tube (see Fig. 26.1, step 2). Hold the proximal end of the needle with one hand to prevent loss of the needle into the tube. Inject the drug solution rapidly and forcefully. If using a prefilled syringe, flush the tube immediately with the diluent in a second syringe. If the patient makes an effort to cough, place a thumb over the opening of the ET tube to prevent expulsion of the solution. Reattach the bag ventilation device and deliver five rapid insufflations.

**Use of a Catheter**

Draw the plunger back to add 5 mL of air to the liquid in the syringe. If the drug to be delivered is in a prefilled syringe, place an IV adapter lock on the catheter if necessary to accommodate the syringe needle or needleless tip. Attach the syringe to the catheter at this time or once the catheter has been placed within the ET tube. In addition, draw up the appropriate volume of diluent (normal saline or distilled water) plus 5 mL air into a second syringe to flush the catheter after instillation of the drug from the prefilled syringe. The air flush presumably forces out any medication adhering to the walls of the catheter’s lumen. Rehan and colleagues determined that, when using a catheter in a neonatal model, more medication was delivered with an additional air flush than without an air flush.

Disconnect the proximal end of the ET tube from the bag ventilation device. Place the catheter into the lumen of the ET tube in such a manner that the distal end of the catheter extends approximately 1 cm beyond the ET tube (see Fig. 26.1, step 3). Hold the proximal ends of the catheter and ET tube at all times during the procedure. If it has not already been done, attach the syringe to the catheter. Inject the drug solution rapidly and forcefully through the catheter into the trachea followed by the 5 mL of air needed to flush the catheter of any remaining drug solution (see Fig. 26.1, step 4). If using a prefilled syringe, use the second syringe to promptly flush with the diluent and air. Immediately remove the syringe and catheter from the ET tube. Reconnect the bag ventilation device with supplemental oxygen to the ET tube and deliver five rapid insufflations.

**Use of ET Tubes With Irrigation and Drug Delivery Lumens**

The following tubes have built-in ports:

1. ET tubes designed for bronchoscopy (e.g., Hi-Lo Jet Tracheal Tube, Nellcor, Pleasanton, CA) (Fig. 26.2A). These tubes have two additional ports, one for monitoring or irrigation (opaque lumen) and one for jet ventilation (transparent lumen). They are available in only uncuffed sizes. The major disadvantage of this ET tube is the need to be familiar with the specific ports before use. If one has never seen the tube previously, determining which port is used for irrigation could prove to be time-consuming. In addition, the port requires placement of an IV adapter lock or Luer-Lok to use a prefilled syringe.
2. ET tube with a side port (ETSP; e.g., EMT Emergency Medicine Tube, Nellcor, Pleasanton, CA) (see Figs. 26.1, step 5, and 26.2B). This tube is designed specifically for ET drug administration but is available in cuffed sizes only. The instillation lumen opens into the tube at the Murphy eye (a hole approximately 1 cm from the end of the tube). The injection port has an IV adapter lock, which makes it amenable to use with prefilled syringes. The ETSP is not available in pediatric sizes. In one study comparing the administration of lidocaine via the ETSP with administration through the proximal end of the standard ET tube, serum lidocaine measurements never reached therapeutic levels in the ETSP group, in contrast to the ET group and IV control group.65

3. Uncuffed tracheal tube with a monitoring lumen (Nellcor, Pleasanton, CA). This tube contains a separate monitoring lumen in the wall of the tube that opens inside the distal tip. A three-way stopcock with a Luer-Lok adapter provides access to the monitoring lumen. The major disadvantage of this ET tube is the need to be familiar with the additional port.

The advantage of all these specialized ET tubes is that they eliminate the need to disconnect the bag ventilation device and the ET tube.

Figure 26.1 Endotracheal administration of medication.
produce adverse effects. Administration of epinephrine during cardiopulmonary resuscitation (CPR) has been noted in case reports to produce prolonged hypertension, tachycardia, and arrhythmias after the return of a perfusing rhythm.10,23 It appears that these side effects are related to the depot effect, in which larger doses of drugs administered endotracheally are released slowly over time. In addition to epinephrine, atropine and lidocaine also exhibit a depot effect when administered endotracheally.58 No serious long-term sequelae, however, have been reported to result from this effect.

A potential concern with ET drug therapy is a transient decrease in arterial oxygen content during or after drug delivery. If total volumes are maintained between 5 and 10 mL in adults, the effect on pulmonary function appears to be minimal. Supplemental oxygen should always be administered to improve oxygenation and offset any transient drop in arterial oxygen content that might develop.

One unusual complication of ET epinephrine is that, during CPR, some of the instilled medication may be expelled and enter the eye. This will produce a fixed dilated pupil, simulating brain death or, if unilateral, brain herniation.

**Injection Through the Wall of the ET Tube**

This method of drug delivery has not yet been evaluated scientifically but has been used clinically.66,67 As with ET tubes with drug delivery lumens, this technique requires no interruption of the connection between the bag ventilation device and the ET tube (see Fig. 26.1, step 6). Placing an IV adapter lock on the needle allows it to be left inserted in the ET tube for use with additional medications.67

**Use of the ET Atomizer**

The Mucosal Atomizer Device-Endotracheal Tube (MADett, LMA North America, San Diego, CA) is an L-shaped port that attaches to both the bag-valve-mask device (BVM) and endotracheal tube (ETT) and allows uninterrupted ventilation during drug administration. The catheter should be inserted until the black depth marker is at the 26-cm mark on the ETT.

**Complications**

Reported complications of ET drug therapy are rare, in part because of the infrequent use of this technique. Because most patients who receive ET drug therapy are in cardiopulmonary arrest or are otherwise critically ill, it is difficult to ascertain whether an adverse outcome is the result of the therapy.

With regard to the techniques of ET drug administration, no serious complications have been reported. A theoretical complication is loss of a needle or catheter down the ET tube, which can be prevented by holding the catheter or needle while instilling the drug.

After ET drug administration, the well-described systemic effects of drugs administered in emergency situations may...
Intranasal Medication Administration

**Indications**
- Pain relief
- Anxiolyis/sedation
- Seizure control
- Narcotic overdose

**Contraindications**
- Abnormal nasal anatomy
- Nasal trauma

**Complications**
- Side effects of the specific medication
- Local nasal irritation
  - For epinephrine: Fixed dilated pupil(s) if expelled into eye(s) during CPR

**Equipment**

---

**Narcotic Overdose**

The Food and Drug Administration approved in November 2015 IN naloxone for use in opioid overdose. Drugs used routinely in the ED that have been studied intranasally include the opioid antagonist naloxone, the benzodiazepine midazolam, the sedative-hypnotic ketamine, the opioids fentanyl and sufentanil, and most recently the α2-agonist sedative dexmedetomidine. Most of these studies have focused on the pediatric patient population as an alternative to IV medication. The patient does not need to be developmentally able to “sniff” in order to enjoy the benefits of IN medication, but only to have the mucosal surface coated in the medication, making its use in the very young patient feasible.

**Seizures**

Bhattacharyya and colleagues studied IN midazolam versus rectal diazepam in pediatric seizure patients. Midazolam (0.2 mg/kg) delivered intranasally, was found to have a significantly more rapid onset of seizure cessation with minimal effect on respiration and oxygen saturation. IN midazolam is now available on the market for pediatric and adult patients.
used in both the prehospital setting and the ED to deliver antiepileptic medication until an IV line can be initiated. A systematic review and metaanalysis comparing non-IV midazolam (IN, buccal, IM routes) and per rectum (PR) diazepam showed non-IV midazolam is as effective and safe as intravenous or rectal diazepam in terminating early status epilepticus. In a network analysis, IN midazolam was better than buccal midazolam, sublingual lorazepam, PR diazepam, and IM midazolam at stopping seizures at 10 minutes, and for seizures lasting more than 1 hour.

**Sedation**

IN midazolam (0.3 mg/kg) compared favorably against buccal and oral routes of midazolam use in children requiring procedural sedation, exhibiting a faster onset of sedation, achieving adequate sedation in a greater proportion of patients, and more parents of pediatric patients stating that they would choose the same regimen again. Patients may experience local irritation of the nasal mucosa due to the acidity of midazolam. Local irritation is amplified when drops are used to instill the midazolam. Premedication with IN lidocaine may decrease the local irritation associated with midazolam.

Ketamine (5 mg/kg) in combination with midazolam (0.3 mg/kg) administered intranasally has been used as an effective anesthesia induction agent with a rapid onset of sedation. Compared with IN midazolam (0.2 mg/kg) as a premedication agent, IN ketamine (5 mg/kg) had a longer time to onset and was associated with more side effects than midazolam. For dissociative levels of sedation, doses of 7 to 9 mg/kg of IN ketamine have been necessary to attain this state. The volume of medication required to reach these higher doses often exceeds the nasal cavity capacity to absorb the medication, rendering IN ketamine for dissociative sedation impractical.

Dexmedetomidine, an α2-agonist with sedative, anxiolytic, and analgesic properties can be given intranasally (2–3 mcg/kg) to sedate children for nonpainful procedures. It has not been well studied for painful procedures. Onset of action is 10 to 25 minutes and its duration of action is 40 to 100 minutes (measured as ability to be discharged). Overall side effects of this medication are rare (bradycardia and hypotension) and have little effect on respiratory drive. There is a case report of vasovagal syncope and bradycardia in an 11-year-old child who had received IN dexmedetomidine at a dose of 2.4 mcg/kg.

**Pain Management**

Fentanyl is widely used for the management of acute pain. IN fentanyl has been shown to have desirable pharmacokinetics and high bioavailability in the treatment of acute pain. Foster and colleagues demonstrated that in adults, IN fentanyl (75 to 200 μg) had only a slight, non-statistically significant lag in the onset of analgesia but equivalent pain reduction to the IV form of fentanyl.

IN fentanyl has been shown to be as effective as IV morphine and had a similar side effect profile. IN fentanyl can be used for pain control in patients prior to initiating an IV line or as a noninvasive alternative to IV or IM injection. Onset of action of IN fentanyl is 2 to 5 minutes, lasting at least 60 minutes. Pediatric dosing is safe and most effective when given at a range of 1 to 3 mcg/kg. There is even a case report of IN fentanyl being successfully used to abort a blue “tet spell” in an infant with the cyanotic congenital heart condition Tetralogy of Fallot. Combination IN therapy using sufentanil (0.5 mg/kg) and ketamine (0.5 mg/kg) is effective in providing analgesia for acute pain. Among adult patients, an average dose of 1 mg/kg of IN ketamine effectively reduced acute pain in just over one half of patients. Effective adult dosing of IN ketamine has not been well studied.

**Contraindications**

There are no absolute contraindications to the intranasal administration of medication with the exception of medication allergy. Abnormal nasal anatomy or increased mucus production may reduce absorption and necessitate repeated dosing.

**Equipment**

The only equipment necessary is the desired medication, a needle to draw the medication from the vial, a 1-mL syringe, and an atomizing device if available.

**Procedure**

Two methods to deliver drugs to the nasal mucosa can be used: drops or aerosol. Nasal drops require a cooperative patient and correct positioning to enhance drug delivery. Atomization of medication is much preferred to drops, in that much of the drug is lost to the environment by dripping out the nose or into the throat and then swallowed. This can result in metabolism of the drug in the liver through first-pass metabolism. Nasal atomization has been found to be the optimal method of enhancing mucosal coverage and increasing plasma concentrations of IN medications. Regardless of the method, the relatively small volume of the nasal cavity limits the volume of medication to approximately 1 mL per naris. If the volume is greater than 1 mL, split the dose and instill half into each naris. Use concentrated forms of medications to decrease the overall volume.

**Nasal Drops**

Draw up the appropriate dose and volume of medication. Position the patient properly to enhance delivery to the mucosa and prevent runoff or swallowing. Instill the drops slowly to prevent runoff.

One position is to place the patient on the back with the head down and nose pointing up. Slowly instill the drops into each naris along the nasal septum and allow the medication to flow into the turbinates. A second position is the lateral decubitus position with the head angled downward. Use pillows or towels to elevate the body at the shoulders if necessary. Instill the drops into the naris that is “up” so that the medication runs along the nasal septum and turbinates. An alternative, though more uncomfortable position is to place the patient on the knees with the head down and the vertex parallel to the bed, essentially in a position similar to starting a forward roll. Instill the medication against the septum and let it flow to the turbinates.

**Nasal Atomization**

Many commercial devices are available for home and outpatient use to deliver medications intranasally. The Mucosal Atomization Device (LMA North America, San Diego, CA) is small, easy to use, and attaches to nearly any syringe (Fig. 26.5). Atomization of midazolam was found to achieve higher plasma concentrations than nebulized midazolam. Atomization results
in very fine particles that distribute over the surface and can be absorbed more readily.

Draw up an appropriate volume of medication into a syringe with an additional amount to accommodate for the dead space of the device (0.1 mL). Insert the device approximately \( \frac{1}{4} \) to \( \frac{1}{2} \) inch into the vestibule. Rapidly depress the plunger. If the total volume is greater than 1 mL, repeat this in the other naris (Fig. 26.6).

An obvious advantage of an atomizer is that proper positioning is not necessary and the medication is instilled rapidly. This can result in less runoff and requires less cooperation from the patient.

**Complications**

Aside from the side effects of the medications (allergy, nausea, apnea, etc.), there are few to no complications when using IN medications. As mentioned previously, midazolam has been noted to cause short-term local irritation. A single case of anosmia following long-term use of IN ketamine has been reported.\(^ {109} \)

**Nebulized Naloxone**

Naloxone has been administered via nebulization, as an alternative to more traditional routes, to reverse opioid intoxication.\(^ {110} \) The benefit of nebulized naloxone is that intravenous access is not required, the reversal effect may be prolonged or continued as long as the nebulizer is used, and there may be a more gradual, but often not complete, reversal of opioid intoxication, reversing opioid effects without producing acute

---

**Rectal Administration Of Medication**

**Indications**

Drug administration when more desirable routes are unavailable or impractical:
- Children frightened of intravenous catheterization
- Patients who refuse parenteral drug administration
- Patients with nausea/vomiting or inability to swallow

**Contraindications**

Immunosuppression
Severe thrombocytopenia or coagulopathy
Active gastrointestinal bleeding
Diarrahea
Chronic anorectal problems (fissures, hemorrhoids, fistulas, etc.)

**Complications**

Erratic absorption
- Delayed, prolonged, or unusually rapid
- Local trauma
- Pain

---

**Equipment**

- Syringe with medication
- Water-based lubricant
- Medication suppository
- 16- or 18-gauge intravenous catheter
- 6- or 8-French pediatric feeding tube

---

**Review Box 26.3** Rectal medication administration: indications, contraindications, complications, and equipment.
withdrawal. Most reports are anecdotal experience or case reports, and there are little data on the specific use. Weber and colleagues\textsuperscript{111} reported success with nebulized naloxone in approximately 80% of spontaneously breathing patients with suspected opioid intoxication when paramedics administered the medication. The procedure may be applicable to nonapneic patients and can be used by both prehospital and ED clinicians. Empirical dosing is 2 to 4 mg of naloxone in 3 mL of saline, delivered by an oxygen driven nebulizer, such as those used to deliver aerosolized β-agonists to asthmatics. Rescue IN, IM, or IV naloxone should be available for those not responding appropriately.

**RECTAL ADMINISTRATION OF MEDICATION**

Medications may be administered rectally when other more preferable routes are not available. This most often occurs when IV access is impractical or impossible, the medication is not suitable for IM administration, or other non-parenteral routes are unavailable or less desirable. In these situations, rectal administration may offer an alternative method of drug delivery. The major drawback to rectal drug administration is unpredictable and often erratic drug absorption.

Drug absorption from the rectum is a simple diffusion process across the lipid membrane. In general, the rate of absorption rises with increasing lipid solubility of the drug and, when applicable, with increased rate of drug release from its carrier (e.g., time to liquefaction of suppository preparations). Other factors affecting transmucosal rectal absorption include the volume of liquid, concentration of the drug, length of the rectal catheter (i.e., site of rectal drug delivery), presence of stool in the rectal vault, pH of the rectal contents, rectal retention of the drug or drugs administered, and differences in venous drainage within the rectosigmoid region.\textsuperscript{18}

**Anatomy and Physiology**

The rectum is the terminal portion of the large intestine; it begins at the confluence of the three taeniae coli of the sigmoid colon and ends at the anal canal. In adults, the anal canal is approximately 5 cm in length and the rectum is approximately 10 to 15 cm in length. In children, the size of the rectum varies with age (Table 26.2).

<table>
<thead>
<tr>
<th>AGE</th>
<th>DIAMETER (cm)</th>
<th>LENGTH (cm)</th>
<th>VOLUME (mL)</th>
<th>LENGTH TO INSERT CATHETER (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 mo</td>
<td>1.5</td>
<td>3</td>
<td>7</td>
<td>1.5</td>
</tr>
<tr>
<td>3 mo</td>
<td>3.0</td>
<td>6</td>
<td>42</td>
<td>3</td>
</tr>
<tr>
<td>1 yr</td>
<td>3.5</td>
<td>7</td>
<td>67</td>
<td>3.5</td>
</tr>
<tr>
<td>2 yr</td>
<td>4.0</td>
<td>8</td>
<td>100</td>
<td>4</td>
</tr>
<tr>
<td>6 yr</td>
<td>4.5</td>
<td>9</td>
<td>143</td>
<td>4.5</td>
</tr>
<tr>
<td>10 yr</td>
<td>5.0</td>
<td>12</td>
<td>235</td>
<td>6</td>
</tr>
</tbody>
</table>


The rectum has two main routes of venous drainage. The superior rectal vein drains into the portal circulation (by way of the inferior mesenteric vein), whereas the middle and inferior rectal veins drain into the caval system (by way of the inferior iliac vein). This pattern of venous drainage has a significant impact on the peak serum concentrations achieved by rectally administered drugs. Drugs administered high in the rectum (the area drained by the superior rectal vein) are carried directly to the liver via the portal vein and are subject to first-pass metabolism. In contrast, drugs administered low in the rectum are delivered systemically into the inferior vena cava, thereby avoiding first-pass elimination in the liver.

**Indications and Contraindications**

Rectal drug administration is indicated when more desirable routes are unavailable or impractical. For example, IV access in small children may be very difficult to obtain and frightening to the child. In these situations, the option of rectal administration may outweigh the benefits of IV drug delivery. Patients who refuse parenteral drug administration may also benefit from rectal delivery, as will those with nausea and vomiting or an inability to swallow.

There are few, if any, absolute contraindications to rectal drug administration. Rectal administration should be avoided in immunosuppressed patients, in whom even minimal trauma could lead to abscess formation, and in patients with severe thrombocytopenia or coagulopathy to avoid difficult-to-control bleeding.\textsuperscript{28,112} Patients with active lower gastrointestinal bleeding and those with severe diarrhea are not generally good candidates for rectal drug administration. Finally, patients with a variety of acute or chronic anorectal problems such as fissures, hemorrhoids, or perianal abscesses or fistulas may not tolerate rectal drug administration.

**Equipment**

For medications in suppository form, a water-soluble lubricant (e.g., Surgilube, HR Pharmaceuticals, Inc., York, PA) will reduce the discomfort associated with insertion. For liquid and gel formulations, use an appropriately sized syringe attached to a small (e.g., 6- or 8-Fr) rubber feeding tube or an 18- or 16-gauge plastic IV catheter with the needle removed. A specialized rectal catheter, the Macy Catheter (Hospi Corp, Newark, CA) (Fig. 26.7), is a device that offers a rectal route for administration of medications or fluids via rectal mucosal absorption. The catheter is a thin silicone tube 14-Fr in diameter with a 15-mL balloon at the tip, sized to allow secure retention, yet also provide for ready elimination in the event of need for defecation.
It features multiple exit ports for fluid and medication passage, and an internal one-way check valve to prevent backflow of fluids.

**Procedure**

**Suppositories**

Place adults and large children in a lateral recumbent position on the stretcher or examination table. Flex the upper leg at the knee and hip, but keep the lower leg extended. Infants, because of their small size, can be placed in almost any position. Place the lubricated suppository at the rectal opening and gently push it into the rectum toward the umbilicus until the gloved index finger has been inserted approximately 7.5 cm in adults, 3.5 cm in children, or 1.5 cm in infants (see Table 26.2). To help prevent expulsion of the suppository, do not allow the patient to get up for approximately 10 to 15 minutes after insertion.

Most suppositories have an apex at one end (pointed end) and taper to a blunt base at the other end. For ease of insertion, manufacturers recommend inserting the tapered end first. However, in 1991, Abd-El-Maeboud and colleagues found that inserting suppositories blunt end first resulted in greater retention within the rectum and a lower expulsion rate. Nonetheless, these findings have not been corroborated and have been challenged by nursing educators as insufficient evidence on which to base clinical practice.

**Liquids and Gels**

Position the patient as described for insertion of a suppository. Draw up the desired dose of medication into an appropriately sized syringe attached to a 6- or 8-Fr rubber feeding tube (adults and large children) or an 18- or 16-gauge IV catheter (infants and small children) with the needle removed (Fig. 26.8). The goal is to deposit the drug in the low to mid-portion of the rectum to avoid first-pass elimination by the liver. For adults, insert the rubber feeding tube approximately 7.5 to 10.0 cm and slowly inject the drug. In children, catheter depth varies with age (see Table 26.2). When administering rectal medication to infants and young children, be sure to squeeze the buttock cheeks closed after withdrawing the catheter to prevent expulsion of the medication. A 3-inch piece of tape placed across the buttocks also works well and frees the clinician to perform other duties.

**Medications**

A variety of medications can be administered rectally. In emergency medicine practice the most common medications given rectally are analgesics and antipyretics, sedative-hypnotic agents, anticonvulsants, antiemetics, and cation exchange resins (e.g., Kayexalate).

**Analgesics and Antipyretics**

Acetaminophen is frequently administered rectally in children for both fever and pain. Common reasons for rectal administration include refusal to take the medication orally, vomiting, and altered mental status. Acetaminophen is commercially available in suppository form and is easy to obtain and administer. Studies comparing oral and rectal administration of acetaminophen have demonstrated equal antipyretic effectiveness.

Rectal dosing of acetaminophen is the same as oral dosing (Table 26.3).

Though not commonly administered rectally, aspirin, nonsteroidal antiinflammatory drugs (NSAIDs) including indomethacin and diclofenac (diclofenac suppositories are not available in the United States), and morphine also come in rectal formulations and can be very useful in a variety of ED encounters. For example, aspirin is commonly administered rectally to adults with symptoms of a transient ischemic attack, an acute stroke, or an acute coronary syndrome who may have an impaired swallowing mechanism or are too unstable to take medication orally. Like acetaminophen, the oral and rectal doses of aspirin are similar (see Table 26.3). Rectal NSAIDs and morphine may be an alternative for patients discharged from the ED who require ongoing analgesia but cannot tolerate oral medications. Rectal doses of indomethacin and morphine are shown in Table 26.3.

**Sedative-Hypnotic Agents**

Sedative-hypnotic agents, including midazolam, methohexital, and thiopental may be administered rectally in children requiring sedation in the ED. This occurs most often in pediatric patients in whom IV access may be problematic or for procedures that require only minimal sedation in conjunction with the use of local anesthetics (see Chapter 33 for a detailed discussion of sedation and analgesia). Rectal administration of methohexital and thiopental is particularly useful for non-painful procedures such as sedating children before advanced imaging studies. The major drawback of rectal administration is an inability to titrate these medications to the desired level of sedation.

The rectal doses of midazolam, methohexital, and thiopental in children are 0.25 to 0.5 mg/kg, 25 mg/kg, and 5 to 10 mg/kg, respectively (see Table 26.3). To prepare a solution of methohexital for rectal administration, add 5 mL of sterile water or saline to a 500-mg vial of methohexital and mix well; this provides a methohexital solution of 100 mg/mL.

**Anticonvulsants**

Anticonvulsants are generally administered orally or intravenously in the ED. However, there may be situations, such as in actively seizing patients, in which oral or IV administration is impossible or unlikely to be accomplished in a reasonable
<table>
<thead>
<tr>
<th>CLASS</th>
<th>RECTAL DOSE/FREQUENCY</th>
<th>ONSET</th>
<th>DURATION</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Analgesics/Antipyretics</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acetaminophen</td>
<td>Adults: 325–650 mg q4–6 hr or 1000 mg q6–8 hr (max, 4 g/day) Children: 10–15 mg/kg q4–6 hr (max, 2.6 g/day)</td>
<td>&lt;1 hr</td>
<td>4–6 hr</td>
<td>Studies have demonstrated equal antipyretic efficacy with rectal and oral administration of acetaminophen</td>
</tr>
<tr>
<td>Aspirin</td>
<td>Adults: 300–600 mg q4–6 hr (max, 4 g/day) Children: 10–15 mg/kg q4–6 hr (max, 4 g/day)</td>
<td>&lt;1 hr</td>
<td>4–6 hr</td>
<td>Contraindications are the same as for oral administration (e.g., children with chickenpox or recent flulike symptoms)</td>
</tr>
<tr>
<td>Indomethacin</td>
<td>Adults: 25–50 mg/kg per dose q8–12 hr (max, 200 mg/day) Children ≥2 yr: 1–2 mg/kg per day q6–12 hr (max, 4 mg/kg per day) Children &gt;14 yr: see adult dosing</td>
<td>≈30 min</td>
<td>4–6 hr</td>
<td>Indocin suppositories only come in 50-mg doses, which may exceed pediatric dosing needs</td>
</tr>
<tr>
<td>Diclofenac</td>
<td>N/A</td>
<td></td>
<td></td>
<td>Not available in a rectal formulation in the United States</td>
</tr>
<tr>
<td>Morphine</td>
<td>Adults: 10–20 mg q3–4 hr Children (≥6 mo and &lt;50 kg): 0.15–0.2 mg/kg q3–4 hr</td>
<td>≈30 min</td>
<td>3–4 hr</td>
<td>Rectal absorption may vary widely; close observation for respiratory depression is mandatory in infants and children</td>
</tr>
<tr>
<td>Sedative Hypnotics</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Midazolam</td>
<td>Adults: seldom given rectally Children: 0.25–0.5 mg/kg</td>
<td>10–30 min</td>
<td>60–90 min</td>
<td>Because these drugs cannot be titrated when given rectally, close monitoring for oversedation is mandatory</td>
</tr>
<tr>
<td>Methohexital</td>
<td>Adults: seldom given rectally Children: 25 mg/kg</td>
<td>5–15 min</td>
<td>60 min</td>
<td>Prepare a solution of methohexital for rectal administration by adding 5 mL of sterile water or saline to a 500-mg vial and mix well</td>
</tr>
<tr>
<td>Thiopental</td>
<td>Adults: 3–4 g/dose Children: 5–10 mg/kg per dose</td>
<td>5–15 min</td>
<td>60 min</td>
<td></td>
</tr>
<tr>
<td>Anticonvulsants</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diazepam</td>
<td>Adults: 10 mg; may repeat once Children: 0.5 mg/kg; then 0.25 mg/kg in 10 minutes (max, 20 mg)</td>
<td>2–10 min</td>
<td>30 min</td>
<td>Dosing guidelines are for status epilepticus; the adult rectal delivery system contains 4 mL (20 mg) of diazepam gel with a 6-cm tip; the pediatric rectal delivery system contains 2 mL (10 mg) of gel with a 4.4-cm tip (see text for details)</td>
</tr>
<tr>
<td>Antiemetics</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prochlorperazine</td>
<td>Adults: 25 mg Children &gt;9 kg: 0.4 mg/kg per day q6–8 hr (not recommended in children &lt;10 kg)</td>
<td>≈60 min</td>
<td>12 hr</td>
<td>Central nervous system effects similar to those with other routes of administration for both prochlorperazine and promethazine</td>
</tr>
<tr>
<td>Promethazine</td>
<td>Adults: 12.5–25 mg q4–6 hr Children &gt;2 yr: 0.25–1 mg/kg per day q4–6 hr (max, 25 mg/dose)</td>
<td>≈30 min</td>
<td>4–6 hr (up to 12 hr)</td>
<td></td>
</tr>
<tr>
<td>Cation Exchange Resin</td>
<td>Sodium polystyrene sulfonate (Kayexalate) Adults: 30–50 g q6hr Older children: 1 g/kg per dose q2–6 hr Young children/infants: calculate the dose (1 mEq K+/g of resin)</td>
<td>2–24 hr</td>
<td>6 hr</td>
<td>Avoid solutions containing sorbitol; see text for details regarding rectal administration</td>
</tr>
</tbody>
</table>

*The onset of action of rectally administered medications may vary widely because of erratic absorption and first-pass elimination in the liver. ED, Emergency department; q, every.
period. In these cases, rectal administration may be an effective alternative.\textsuperscript{122} Studies have demonstrated that diazepam, because of its high lipid solubility, is rapidly absorbed from the rectum and can quickly halt seizures.\textsuperscript{125,126} Lorazepam has much lower lipid solubility and is not recommended for rectal use.

Diazepam is commercially available in a gel formulation that is preloaded in a rectal delivery system (Diastat AcuDial). However, the undiluted parenteral formulation can also be used. The preloaded rectal delivery system is available for both pediatric and adult use. The adult device contains 4 mL (20 mg) of diazepam gel and has a 6-cm tip for rectal administration. This device is designed to deliver set doses of 10, 12.5, 15, 17.5, and 20 mg of diazepam. Two pediatric devices are available: one contains 0.5 mL of a 5-mg/mL gel, and the other contains 2 mL of a 5-mg/mL gel. The latter is designed to deliver set doses of 5, 7.5, and 10 mg of diazepam. Both pediatric devices have a 4.4-cm tip for rectal administration. The recommended dose of diazepam rectal gel for treating actively seizing children and those in status epilepticus is 0.5 mg/kg, followed by 0.25 mg/kg in 10 minutes if needed (maximum dose, 20 mg); the adult dose is 10 mg, which may be repeated once (see Table 26.3).

**Antiemetics**

Antiemetics are frequently used in the ED for patients with nausea and vomiting. In most cases, these medications are given intravenously. Oral dissolving tablets (ODT) antiemetics such as ondansetron have largely replaced the use of rectal antiemetics as this route is preferred by most patients. However, for patients with mild symptoms who do not otherwise require IV access and for discharged patients expected to have ongoing nausea and vomiting (e.g., patients with a viral gastroenteritis), rectal administration is an alternative. Rectal administration may also be used as the initial treatment of ED patients with active vomiting in whom IV access will be delayed, and ODT antiemetics have failed, such as those requiring central venous access.

Historically, the two most common antiemetics administered rectally in the ED are prochlorperazine and promethazine. Both come in suppository formulations, which makes rectal administration easy. Prochlorperazine requires a higher dose when given rectally, whereas promethazine dosing is the same regardless of the route of administration (see Table 26.3).

**Cation Exchange Resin**

The most commonly available cation exchange resin is sodium polystyrene sulfonate (Kayexalate). In the gut, sodium polystyrene sulfonate absorbs potassium and releases sodium. Each gram of resin may bind as much as 1 mEq of potassium and release 1 to 2 mEq of sodium. In the ED, sodium polystyrene sulfonate is frequently given to patients with hyperkalemia as a temporizing measure before dialysis.

Sodium polystyrene sulfonate may be given orally or rectally as a retention enema. Oral dosing is more effective if intestinal motility is not impaired. Common reasons for rectal administration include an inability or refusal to swallow (the oral solution is not very palatable), vomiting, and altered mental status. The resin comes in two forms: a powder that must be reconstituted and a premixed suspension containing sorbitol.

Before drug administration, perform a cleansing enema with warm tap water. Prepare a sodium polystyrene sulfonate enema by dissolving 50 g of the resin in 100 to 150 mL of tap water warmed to body temperature. In adults, administer the resin emulsion via a 6- or 8-Fr rubber feeding tube placed approximately 20 cm from the rectum with the tip well into the sigmoid colon. Retain the enema in the colon for at least 30 to 60 minutes and for several hours if possible. Once retention is complete, irrigate the colon with 50 to 100 mL of a non-sodium–containing fluid. The recommended dose of sodium polystyrene sulfonate for rectal administration in adults is 30 to 50 g every 6 hours. For children, the dose is 1 g/kg per dose every 2 to 6 hours. One may also calculate the dosage in neonates based on the exchange ratio of 1 mEq K⁺/g of resin (see Table 26.3).

**Complications**

Complications specific to rectal administration include erratic absorption and local trauma. Absorption of a rectally administered drug may be delayed or prolonged, or uptake may be almost as rapid as though the agent were administered intravenously.\textsuperscript{28} Unusually rapid absorption resulted in the death of a 7½-month-old child receiving rectal morphine for postoperative pain.\textsuperscript{129} In addition to erratic absorption, peak serum concentrations may vary considerably depending on how much of the drug undergoes first-pass elimination in the liver. These factors make rectal drug administration less desirable in most cases when parenteral administration is possible.

Insertion of a suppository or catheter into the rectum may cause mild pain and mucosal irritation, which is usually well tolerated by most patients. Similarly, bleeding from local trauma is usually of no clinical consequence, except in patients with a clinically significant coagulopathy or severe thrombocytopenia.\textsuperscript{112} Development of a perianal or perirectal abscess is a theoretical concern in neutropenic patients, and thus rectal drug administration is best avoided in such patients.

**OTHER ALTERNATE ROUTES OF ADMINISTRATION OF MEDICATION**

There is a limited but growing body of evidence demonstrating the benefits of rehydration of patients (usually the elderly\textsuperscript{130–132} or very young\textsuperscript{133,134} in the emergency department using a subcutaneous route called “hypodermoclysis” (Box 26.1). Historically, hypodermoclysis in humans was used in the elderly in non-emergency situations for rehydration. The advent of human recombinant hyaluronidase has made it possible to give faster rates of IV fluid infusion making its use in the ED feasible. The most common practical use of this method is in the setting of dehydration when IV access is unobtainable, though theoretically, medications may also be given via this route. Potential advantages of subcutaneous rehydration compared with IV rehydration are that it expedites and simplifies parenteral access, requires fewer staff resources, and is potentially less distressing to parents and patients.\textsuperscript{131}

**Anatomy and Physiology**

Hypodermoclysis uses human recombinant hyaluronidase to break up hyaluronan, a mucopolysaccharide found in the intercellular matrix of connective tissue that normally prevents flow of fluids across the subcutaneous layer of tissue. This transiently increases permeability of the connective tissue matrix, allowing movement of subcutaneously administered fluids (and medications) into the surrounding tissues, with subsequent systemic absorption via the capillary beds.
Indications and Contraindications

Subcutaneous hydration is indicated in the moderately dehydrated patient in whom IV access is not easily obtainable, and in whom enteral routes (oral [PO]/nasogastric [NG]) of rehydration have failed. This method should not be employed in the severely dehydrated patient in hypovolemic shock, where IO or central venous access should be considered. Do not use hypodermoclysis if an enteral route (PO, NG) is tolerated.

Equipment

An IV solution bag, a tube with a drip chamber, a small-gauge (21 or smaller) long-tube butterfly needle, povidone-iodine solution or alcohol skin preparation, and a sterile occlusive dressing.

Procedure

Hypodermoclysis sites include the abdomen, upper chest, above the breast, over an intercostal space and the intrascapular area, the lateral thighs, the abdomen, and the outer aspect of the upper arm (Fig. 26.9). In uncooperative adult patients or

**Figure 26.9** Anatomic sites usable for hypodermoclysis. (From Bruno VG: Hypodermoclysis: a literature review to assist in clinical practice, *Einstein* 13(1):122–128, 2015.)
Assemble fluid and tubing. Prime line with selected fluid and 150 units of hyaluronidase. Swab the site with povidone-iodine skin or alcohol preparation solution. Inject local anesthetic into insertion site. Pinching the skin between fingers, insert a 21-gauge (or smaller) needle, bevel up, into subcutaneous tissue at a 45- to 60-degree angle. Administer 150 units of human recombinant hyaluronidase into the subcutaneous space. Secure needle and tubing with occlusive dressing. Adjust fluid drip rate as desired.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


INTRODUCTION

Autologous blood transfusion, or autotransfusion, is the collection and reinfusion of a patient’s own blood for volume replacement. Preoperative blood banking and intraoperative cell salvage techniques have increased in a multitude of surgical specialties. Autotransfusion in the emergency department (ED) is usually limited to patients with severe, traumatic hemothorax and clinically significant blood loss.

Autotransfusion is usually performed in trauma centers or in EDs with high trauma volume. Though not a uniform standard of care for emergency clinicians, it is applicable to any ED. As the procedure requires familiarity with the equipment, continuing education, and quality control, it would be counterproductive to institute the procedure in a hospital that has a low trauma census or in a setting in which it will be used infrequently enough that staff education issues are problematic. Clinicians practicing in more austere environments with a paucity of clinical resources, such as combat and disaster zones, may find that the procedure’s benefits outweigh its risks.

BACKGROUND

Reports of autotransfusion can be found as early as 1818 when Blundell, an English practitioner, reinfused shed blood after witnessing a woman exsanguinate from uterine hemorrhage. In 1886, Duncan published the first known human account of autotransfusion in which he reinfused shed blood in a patient with a traumatic amputation without any notable ill effects. In 1917, Elmendorf published a description of the first case of autotransfusion in a patient with traumatic hemothorax. The discovery of ABO blood typing at the turn of the century and the institution of blood banks in the 1930s led to the almost exclusive use of allogeneic (homologous) blood up

Autotransfusion

Indications
- Hemothorax containing >1500 mL
- Hemothorax with an immediate need for transfusion and insufficient allogeneic blood available
- Hemothorax with an urgent need for blood and the patient’s religious beliefs prohibit the use of banked blood

Contraindications
- Coagulopathy or disseminated intravascular coagulation
- Possibility of malignant cells in the salvaged blood
- Active infection
- Gross contamination of pleural blood from gastrointestinal contents

Complications
- Hematologic
  - Decreased platelet count
  - Decreased fibrinogen level
  - Increased fibrin split products
  - Prolonged prothrombin time
  - Prolonged partial thromboplastin time
  - Red blood cell hemolysis
  - Elevated plasma free hemoglobin
  - Decreased hematocrit
- Nonhematologic
  - Bacteremia
  - Sepsis
  - Microembolism
  - Air embolism
  - Renal insufficiency

Equipment

Chest drainage and autotransfusion system.
(Atrium Ocean ATS shown above, Atrium Medical Corporation, Hudson, NH)

Review Box 27.1 Autotransfusion: indications, contraindications, complications, and equipment.
to and following World War II. During the 1960s and 1970s, cardiopulmonary bypass surgery and combat trauma experience during the Vietnam War generated extensive data regarding intraoperative retrieval of large quantities of blood for reinfusion.16–17 This revitalized interest, coupled with increased experience in surgical, trauma, and combat situations, has thus initiated a new era in autotransfusion.

ANATOMY

Hemothorax refers to a collection of blood within the pleural space. Severe hemorrhage is more often associated with laceration of vessels on the inside of the chest wall, such as the internal mammary and intercostal arteries. Blunt trauma and penetrating trauma are by far the most common causes of hemothorax. Spontaneous hemothorax can occur secondary to intrathoracic malignancy, pulmonary infarction, bullous emphysema, virulent pulmonary or mediastinal infection, vascular malformation, and endometriosis of the pleura (catamenial).18

PATHOPHYSIOLOGY

The pathophysiologic sequelae of hemothorax include both hemodynamic instability and respiratory compromise. The pleural cavity can accommodate more than 50% of the total blood volume, so clinically significant intrathoracic blood loss can occur with minimal external signs of bleeding. The clinician should suspect hemothorax in the setting of chest trauma with clinical signs of hypovolemia. Physical examination may demonstrate decreased breath sounds and reduced tactile fremitus. Imaging during the initial resuscitation period is usually limited to supine chest radiographs, which may show haziness in the affected lung field as blood layers in the posterior pleural space (Fig. 27.1). Ultrasound is also useful in the initial evaluation of hemothorax (Fig. 27.2).20 Tube thoracostomy is the treatment of choice for acute hemothorax, with thoracotomy and video-assisted thoracic surgery performed as indicated for severe or ongoing hemorrhage.

ADVANTAGES

Shed blood from traumatic hemothorax is immediately available for rapid transfusion. The blood is normothermic and compatible, which avoids the risk of allergic reaction or infection from transfusion transmissible diseases. There are numerous transfusion transmissible diseases, including viruses such as human immunodeficiency virus (HIV) and hepatitis, bacteria, parasites, and most recently reported, variant Creutzfeldt-Jakob disease.21–22 Although the risk for transfusion transmissible diseases has decreased dramatically in developed countries, it is still very problematic worldwide.23 Immunologic transfusion reactions and posttransfusion sepsis continue to be risks as well.24 In trauma patients, allogeneic transfusions have been shown to be an independent risk factor for infection,25–28 which may be dose dependent and independently associated with increased morbidity and mortality.28–30 In patients whose religious convictions (e.g., Jehovah’s Witness) prohibit transfusions with homologous blood, reinfusion of autologous blood that does not involve blood storage may be an acceptable alternative.31

A recent multicenter study by Rhee and colleagues32 compared outcomes in 136 patients receiving autotransfused blood from a traumatic hemothorax to an equal number receiving only allogeneic transfusions. There was no difference between the groups in mortality or in-hospital complications (sepsis, disseminated intravascular coagulation [DIC], acute lung injury, or renal insufficiency). Despite its retrospective design, this study offers support for the practice of autotransfusion based on patient-oriented outcomes.

Autologous blood provides societal benefits by preserving the limited stores of banked blood and reducing the cost of medical care.32–35 Adias and colleagues compared the direct cost of banked blood with the cost of autologous blood transfusion and found substantial savings with autotransfusion.16

Box 27.1 summarizes the advantages of autotransfusion.

INDICATIONS

In general, all victims of severe trauma, whether blunt or penetrating, should be considered potential candidates for

Figure 27.1 Hemothorax secondary to a gunshot wound. Note the haziness over the right hemothorax with the bullet seen in the right upper lobe. In this supine radiograph, the volume of the hemothorax may not be fully appreciated. (From Marx JA: Rosen’s emergency medicine, ed 7, St. Louis, 2009, Mosby.)

Figure 27.2 Ultrasonographic appearance of hemothorax. Blood in the pleural cavity appears anechoic (arrow) and is easily seen interspersed between the collapsed lung and diaphragm.
BOX 27.1 Advantages of Autotransfusion

1. Immediately available—no delay in crossmatching and no storage required
2. Blood compatibility and allergic reaction not an issue
3. Autologous blood usually nonthermothermic
4. No risk for transfusion transmissible disease
5. No risk for hypocalcemia or hyperkalemia
6. Decreased risk for acute respiratory distress syndrome
7. Higher levels of 2,3-diphosphoglycerate than in banked red blood cells
8. Decreased use of banked blood; more available for subsequent patients in need
9. Decreased cost of medical care
10. May be acceptable to religions opposed to homologous blood transfusions
11. May be a valuable alternative to banked blood in developing countries where infected donor blood is a problem

CONTRAINDICATIONS

In some situations, emergency autotransfusion may pose more risk than benefit. Coagulopathy and DIC are important contraindications. Others include active infection, gross contamination, and the possibility of malignant cells in the salvaged blood. With direct autotransfusion of pleural blood through a microfilter, the risk of reinfusing tumor cells is unknown. However, work using perioperative cell salvage suggests that the risk for dissemination of malignant disease is minimal. Concerns regarding gross contamination from concomitant gastrointestinal tract injury have also been disputed. An early study of autotransfusion before cell washing was implemented found that despite reinfusion of massively contaminated blood, 17 of 25 patients survived without evidence of septic complications.

Work on contamination involving cell salvage systems incorporates a cell-washing step. Several investigators believe that reinfusion of limited amounts of contaminated blood from the peritoneal cavity may now be accomplished with acceptable risk. Nonetheless, the current consensus is that exsanguinating hemorrhage without available homologous blood is the only acceptable indication for autotransfusion when there is recognized intestinal contamination.

Concurrent use of systemic antibiotics is advised. Autotransfusion should be performed within 4 to 6 hours from the time of injury, although it has been performed after this period in combat situations without significant complications.

EQUIPMENT AND MATERIAL

Blood Filters

In-line filtration is used routinely during reinfusion of blood products to reduce the danger of microembolization and resultant pulmonary insufficiency. The relationship between the presence of microaggregates and the development of acute respiratory distress syndrome is controversial. However, most investigators advise the use of a micropore filter with pore size recommendations ranging from 20 to 170 µm. The majority of investigators believe that a pore size of 40 µm minimizes the risk for microembolization without undue elevation in filtration pressure. The manufacturers of the commercial devices discussed in this chapter also recommend at least a 40-µm filter size.

Vacuum Suction

The level of vacuum suction should be limited to minimize hemolysis of red blood cells. The precise level at which clinically significant hemolysis occurs is uncertain, and there is wide variation (5 to 100 mm Hg) in recommendations. Several commercial products recommend a starting vacuum pressure of 20 mm Hg.

Anticoagulation

Blood retrieved from the pleural and abdominal cavities frequently will not clot because it is devoid of fibrinogen. This is believed to be because moderate rates of bleeding allow time for de fibrination by contact with the serosal surfaces and by mechanical agitation from respiratory and cardiac movement. For this reason, some recommend simple reinfusion through a filter without any anticoagulant. However, wounds of the great vessels may bleed at a rate that allows coagulable blood to enter the collection reservoir and clot the entire system.

In the ED setting where autotransfusion is indicated for rapid blood loss, anticoagulation is recommended.

The level of vacuum suction should be limited to minimize hemolysis of red blood cells. The precise level at which clinically significant hemolysis occurs is uncertain, and there is wide variation (5 to 100 mm Hg) in recommendations. Several commercial products recommend a starting vacuum pressure of 20 mm Hg.

Anticoagulation

Blood retrieved from the pleural and abdominal cavities frequently will not clot because it is devoid of fibrinogen. This is believed to be because moderate rates of bleeding allow time for de fibrination by contact with the serosal surfaces and by mechanical agitation from respiratory and cardiac movement. For this reason, some recommend simple reinfusion through a filter without any anticoagulant. However, wounds of the great vessels may bleed at a rate that allows coagulable blood to enter the collection reservoir and clot the entire system.

In the ED setting where autotransfusion is indicated for rapid blood loss, anticoagulation is recommended.

Anticoagulants

Citrate compounds, an alternative to heparin, are frequently recommended for autotransfusion. Citrates bind with the calcium ion and prevent conversion of protein fibrinogen into insoluble fibrin, which would cause the blood to clot. Because citrate binds only with calcium, it anticoagulates just the blood in which it is dissolved. Once the anticoagulated blood is infused, citrate is rapidly metabolized by the liver. Early studies used acid citrate dextrose (ACD). More recently, citrate phosphate dextrose (CPD) has been used because it necessitates less volume as an anticoagulant and results in less acidosis than ACD.

Rarely, excessive use of CPD can cause cardiac dysrhythmias. Moreover, use of insufficient or outdated CPD may result in clotting of collected blood. Baldan and colleagues described the use of simple autotransfusion in more than 200 patients with penetrating chest injuries and recommended using half-dose citrate phosphate dextrose adenine solution. Although anticoagulant therapy and dosage recommendations are at the...
discretion of the physician, commercially available devices do offer some general guidelines (Table 27.1).

**Historical Techniques Using Standard ED Equipment**

Before purpose-built systems were available, autotransfusion was possible by using common items available in most EDs. These techniques have not been as widely tested and are more appropriate for use in dire situations, such as may occur in a disaster or on the battlefield.

Symbas described a simplified collecting system involving a standard chest tube bottle. More than 400 patients were autotransfused via this method with no adverse effects attributable to the procedure. After insertion of a chest tube, blood was collected into a standard bottle containing 400 mL of normal saline, with suction maintained at 12 to 16 mm Hg. The blood collected in the chest bottle can be reinfused in one of two possible ways as shown in Fig. 27.3. Improvements on his historical technique, such as the addition of anticoagulation and blood filters, should be considered.

**Autotransfusion Units**

All the commercially available autotransfusion systems (ATSs) are based on the same three-stage system (called the three-bottle system) for collection of pleural fluid (Fig. 27.4). Currently used devices incorporate the three-bottle system into one unit.

![Figure 27.3](http://www.atriummed.com/PDF/Red%20Handbook.pdf)

**Figure 27.3** Historical technique of autotransfusion for traumatic hemothorax. **A**, Blood is collected into a sterile blood collection bottle with 400 mL of normal saline. **B**, After blood is collected, the chest tube is disconnected from the collection bottle. **C**, Blood is infused directly from the blood collection bottle whilst a new chest tube drainage bottle is connected to the chest tube, or **D**, blood collected from the chest tube is transferred into a sterile blood bag. **E**, Blood transferred into the blood bag is infused into the patient. (From Symbas PN: Extraoperative autotransfusion from hemothorax, Surgery 84:722, 1978.)

**TABLE 27.1  ACD-A and CPD Dosage Recommendations**

<table>
<thead>
<tr>
<th>BLOOD VOLUME EXPECTED</th>
<th>FOR 1:7, ADD THIS AMOUNT OF ACD-A</th>
<th>FOR 1:20, ADD THIS AMOUNT OF ACD-A</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low = 140–250 mL</td>
<td>20–35 mL</td>
<td>7–12 mL</td>
</tr>
<tr>
<td>Incremental volume over 250 mL</td>
<td>1 mL for each 100 mL of collected blood</td>
<td>5 mL for each 100 mL of blood collected</td>
</tr>
<tr>
<td>Medium = 250–500 mL</td>
<td>40–70 mL</td>
<td>12.5–25 mL</td>
</tr>
<tr>
<td>High = 500–1000 mL</td>
<td>70–140 mL</td>
<td>25–50 mL</td>
</tr>
</tbody>
</table>

**CPD DOSAGE RECOMMENDATIONS**

Anticoagulant CPD solution can be added at the discretion of the physician at a control dosage of 14 mL of CPD solution to 100 mL of blood collected (70 mL CPD/500 mL blood).

Dosage ratios are approximate. Note that 35 mL of ACD-A provides a 1:7 ratio of ACD-A to blood for low volume and a 1:20 ratio for high-volume blood and may be a good starting point if it is unclear how much blood will accumulate. ACD-A, Acid citrate dextrose anticoagulant; CPD, citrate phosphate dextrose. From A personal guide to managing chest drainage autotransfusion, Atrium Medical Corporation, pp 26, 27. http://www.atriummed.com/PDF/Red%20Handbook.pdf.
Stage 1 is the collection of pleural fluid (blood). Stage 2 is the water seal stage, in which water acts as a one-way valve (water seal) that allows air to be sucked out of the pleural space but not leak back in. Stage 3 is the suction control stage, which is essentially a safety stage so that if the degree of suction suddenly increases beyond the set point, the system will pull air from the atmosphere instead of inappropriately increasing suction in the pleural cavity. Historically, water was (and still frequently is) used in the suction control stage.

The disadvantage of using water in stages 2 or 3 is that if the device is knocked over, loss of the water seal and spillage of water from the suction control stage can occur. Newer designs have solved this problem, and two systems currently available do not use water (Atrium Express [Atrium Medical Corporation, Hudson, NH] and Pleur-evac Sahara [Teleflex, Morrisville, NC]). These newer devices use a one-way valve instead of a water seal in stage 2 and a pressure regulator instead of water in the suction control stage (stage 3).

To collect blood for autotransfusion, a collection bag can be connected beforehand in series with the chest drainage system. With this arrangement, blood goes from the patient to the blood collection bag to the chest drainage system. When the blood bag is filled, it is removed and replaced with a new blood collection bag. If the bag overfills, the fluid will spill into the usual chest fluid collection system. Eventually, when no more blood needs to be autotransfused, the blood collection bag is taken out of the series and the chest tube drainage system functions as usual. Because blood goes directly into the blood collection bag, anticoagulant needs to be added to the blood collection bag before the blood flows into it.

Blood may also be drawn directly from a chest drainage system. Some devices have an additional port at the bottom called an ATS access line. This is referred to as a self-filling ATS. Not all devices permit withdrawal of blood directly from the chest drainage system. In devices without this option, any blood that goes into the chest drainage system is not available for transfusion. The advantage of the ATS access line is that autotransfusion need not be anticipated before the chest tube collection system is connected to the chest tube. Because blood first goes to the collection chamber of the chest drainage device, anticoagulant needs to be added to the collection chamber before blood enters it.

Some models allow these chest drainage systems to be connected directly to the patient to provide continuous autotransfusion. In this setup, a blood filter with intravenous (IV) blood tubing is hooked up to an IV infusion pump. The blood can then be continuously reinfused as it collects in the chest drainage system; it goes from the blood collection chamber out the ATS access line and is pumped into the patient with an IV infusion pump in a closed system (Fig. 27.5). Other models offer continuous infusion directly from a blood collection
bag and not the chest drainage device. The blood simply goes from the chest tube into the blood collection bag, and a port on the bag is accessed to return the blood back to the patient.

**PROCEDURE FOR AUTOTRANSFUSION**

Whether in-line, self-filling, or continuous techniques are chosen depends on the chest tube drainage system used. Some systems allow more than one technique to collect blood for autotransfusion. The specific port locations and connections may vary slightly among the different designs, but the general concepts are relatively universal. See Figs. 27.6 and 27.7 for a description of several different systems from Atrium and Pleur-evac.

The procedure described here is for in-line blood collection and infusion using the Atrium Ocean model; however, all Atrium models have similar connectors. The procedure using Pleur-evac systems is also similar (Video 27.1).

**In-Line Blood Collection and Infusion Procedure**

1. Place the autotransfusion blood bag onto the chest drain via the attached flexible hanger.
2. Close both autotransfusion blood bag clamps and remove the protective caps over the autotransfusion bag connectors.
3. Close the patient's chest tube clamp.
4. Separate the connectors between the patient's chest tube and the Ocean chest drain by depressing the connector lock.
5. Insert the male patient chest tube connector into the female autotransfusion bag connector.
6. Insert the male autotransfusion bag connector into the female Ocean chest drain connector.
7. Open the clamps in the following order:
   a. Open the autotransfusion bag clamp going to the Ocean chest drain.
   b. Open the autotransfusion bag clamp going to the patient's chest tube.
   c. Open the patient's chest tube clamp to resume drainage. (Blood will then flow from the patient's chest tube into the autotransfusion blood bag.)
8. When the blood bag is filled (600 mL maximum), close the patient's chest tube clamp and both autotransfusion blood bag clamps.
9. Disconnect the autotransfusion blood bag connection with the Ocean chest drain connection first.
10. Disconnect the autotransfusion blood bag from the patient's chest tube connector.
11. Insert the male patient chest tube connector into the female Ocean chest drain connector.
12. Open the patient's chest tube clamp so that drainage may resume into the Ocean chest drain device.

---

**Figure 27.6** Pleur-evac chest drainage systems. (Courtesy Teleflex Medical, Research Triangle Park, NC.)
13. Connect the male and female autotransfusion bag connectors to each other.
14. Connect and pre-prime the blood filter and the IV blood tubing with sterile saline (spike the blood tubing into the filter).
15. Invert the autotransfusion bag so that the spike port at the bottom points upward.
16. Remove the tethered cap with sterile technique.
17. Insert the saline-primed filter spike into the autotransfusion bag spike port with a firm twisting motion.
18. Return the autotransfusion bag to an upright position and place it on a standard IV pole.
19. Open the filtered air vent located on top of the autotransfusion bag first, and then open the IV tubing clamp.
20. Evacuate all the remaining air from the IV line and attach it to the patient to begin the infusion.
21. For gravity infusion, leave the air vent at the top of the autotransfusion bag open.
22. For pressure infuser application, the filtered air vent must remain closed (the maximum ATS bag infuser pressure is 150 mm Hg).

**COMPLICATIONS**

In general, complications from autotransfusion are clinically insignificant if proper technique is followed. Complications may be categorized as hematologic and non-hematologic (see Review Box 27.1).

**Hematologic Complications**

The degree to which autotransfusion contributes to the development or exacerbation of coagulopathy in an actively bleeding patient has still not been conclusively determined. Broadie and coworkers found that blood collected from trauma victims during thoracotomy or laparotomy had markedly elevated prothrombin, partial thromboplastin, and thrombin times, as well as absent fibrinogen. The dilemma is that severely injured trauma patients who are candidates for autotransfusion are more likely to have other independent risk factors for coagulopathy, such as hypothermia and acidosis. Horst and colleagues studied 154 trauma patients who underwent intraoperative autotransfusion and found that patients who received more than 15 units of autologous blood or more than 50 units of combined autologous and banked blood were coagulopathic but also more severely injured, hypothermic, and acidotic. Although coagulation test abnormalities following the infusion of salvaged blood have been interpreted as evidence of DIC, these changes are probably the result of infusion of fibrin degradation products and do not represent consumptive coagulopathy. In general, even though coagulopathy should be anticipated, it has not proved to be clinically important when volumes of autotransfused blood remain below 1500 to 2000 mL in adult patients. When reinfused volumes exceed 3500 mL, laboratory evidence of dilutional coagulopathy may become evident. When volumes of autotransfused blood are greater than the patient’s total blood volume, animal studies suggest that the risk for a true consumptive coagulopathy is increased.

Recommendations regarding the volume of autologous blood that should trigger the infusion of plasma or platelets range from 25% of the total blood volume (approximately 1250 mL in a 70-kg adult) to 3500 mL (Box 27.2). Others advise reliance on laboratory tests and clinical findings rather than a volumebased protocol. Prudent clinical judgment dictates application of the more liberal guidelines for replacement therapy in patients with extensive hepatic injury, intractable shock, or ongoing loss requiring immediate surgical intervention.

Hemolysis occurs with autotransfusion, in part because of prolonged exposure of cells to the serosa of the traumatized body cavities. Hemolysis also results from mechanical damage during collection and reinfusion or from excessive exposure to air-fluid interfaces. The hematocrit falls in direct proportion to the quantity of blood transfused, with the average decline being 10% to 20%. However, nontraumatized red blood cell survival has been reported to be normal in all cases studied.

**Nonhematologic Complications**

The theoretical risk for sepsis after the administration of potentially contaminated blood always exists within the nonsterile environment of the typical ED. Experience has shown this risk to be minimal after autotransfusion from an isolated hemothorax, and there is no evidence suggesting that routine prophylaxis with systemic antibiotics is beneficial. Collected blood is generally transfused as soon as the collection bag is full. The American Association of Blood Banks guidelines allow a 6-hour interval between collection and reinfusion. The age of collected blood, however, should be calculated from the time of injury (see Box 27.2). Blood reinfused after this 6-hour period should be considered potentially hazardous.

The risk for microemboli secondary to platelet aggregation and fat emboli has largely been eliminated by the use of micropore filters. During reinfusion of collected blood there is usually a mild increase in screen filtration pressure, indicative of the formation of microemboli trapped by the filter. There has been no clinical evidence of pulmonary insufficiency or elevation of the alveolar-arterial oxygen gradient that might be attributed to the passage of microemboli beyond micropore filter systems.

As with all IV infusions, improper technique, such as applying pressure to air-containing systems, may lead to air embolism. This uncommon but potentially fatal complication has been associated with ATSs that use automated roller pump units in which the aspirate reservoir was inadvertently allowed to run dry. Air embolism with gravity or with a manually assisted technique is rare.

Infusion of large quantities of unwashed blood that contains hemolyzed red blood cells may contribute to renal failure, particularly in patients with preexisting renal insufficiency. Elevated plasma free hemoglobin is a consistent finding in

---

**BOX 27.2 General Autotransfusion Information**

1. Use each liner bag only once.
2. Insert a new filter for each autotransfusion bag used.
3. To minimize risk for bacterial overgrowth, blood collected must be reinfused within 6 hours from the time of injury.
4. After reinfusing a total of 3500 mL (approximately 7 units) of autologous blood, it has been suggested that 1 unit of fresh frozen plasma be given for every 2 units (approximately 1000 mL) of autotransfused blood.
5. If some or all the collected blood becomes clotted in the liner bag, the blood should be discarded.
6. To reduce the risk for air embolism, remove all the air from the bag with the collected blood before hanging it for reinfusion.

*The American Association of Blood Banks suggests no more than a 6-hour shelf life between collection and reinfusion. The age of blood collected should be calculated from time of injury. Blood reinfused after this time should be considered hazardous.*
patients who have undergone autotransfusion. In the past it was believed that elevated levels of free hemoglobin following hemolytic transfusion reactions caused renal failure by its precipitation and obstruction of renal tubules. However, more recent evidence suggests that the mechanism in this setting is independent of free hemoglobin but rather is the result of an antigen-antibody–induced intravascular coagulation that when compounded by vasoconstriction and hypotension, leads to renal ischemia. Even though renal failure as a direct consequence of autotransfusion has not been reported, transient elevations in serum creatinine do occur. In the presence of shock and systemic acidosis, acute tubular necrosis remains a potential complication. The clinician must weigh the urgency of blood transfusion against the timely availability of an alternative source.

RESOURCES

Brief educational videos are available online for Atrium products at http://www.atriummed.com/EN/Chest_Drainage/edu-files/ATS-video1.asp and show in detail the step-by-step processes to initiate autotransfusion with their devices. They also have a 24-hour help line to assist with any questions during the setup or use of their devices (800-528-7486). Pleur-evac has some limited information available online (http://www.teleflexmedical.com) and can be contacted during regular business hours (866-246-6990, 8 AM to 7 PM EST) to answer questions.
CHAPTER

REFERENCES
1. Schaff HV, Hauer JM, Brawley RK: Autotransfusion in cardiac surgical
2. Yoshiba F: [Autologous transfusion for patients with increased risk of
massive and/or critical bleeding—indication, efficacy and limitations].
Masui 60:40–46, 2011.
3. Tavare AN, Parvizi N: Does use of intraoperative cell-salvage delay recovery
4. Shantikumar S, Patel S, Handa A: The role of cell salvage autotransfusion
in abdominal aortic aneurysm surgery. Eur J Vasc Endovasc Surg 42:577–584,
2011.
5. Ralph CJ, Sullivan I, Faulds J: Intraoperative cell salvaged blood as part
of a blood conservation strategy in caesarean section: is fetal red cell
saver on patients undergoing open abdominal aortic aneurysm surgical
7.	Hansen E, Seyfried T: [Cell salvage]. Anaesthesist 60:381–389, quiz 390,
2011.
8. Ahmed AM, Riye MH, Baldan M: Autotransfusion in penetrating chest
war trauma with haemothorax: the Keysaney Hospital experience. East
Chir Trans 9:56–92, 1818.
Munch Med Wochenschr 64:36, 1917.
unit under varying conditions of contamination. Preliminary report. Am
16. Klebanoff G: Early clinical experience with a disposable unit for the
20.	McEwan K, Thompson P: Ultrasound to detect haemothorax after chest
21.	Llewelyn CA, Hewitt PE, Knight RS, et al: Possible transmission of
variant Creutzfeldt-Jakob disease by blood transfusion. Lancet 363:417–421,
2004.
transfusion in a PRNP codon 129 heterozygous patient. Lancet 364:527–529,
2004.
24.	Madjdpour C, Heindl V, Spahn DR: Risks, benefits, alternatives and
indications of allogenic blood transfusions. Minerva Anestesiol 72:283–298,
2006.
with infections in trauma patients in a dose-dependent manner. Am Surg
increases the risk of postoperative bacterial infection: a meta-analysis.
the first 24 hours after trauma is associated with increased systemic
inflammatory response syndrome (SIRS) and death. Surg Infect (Larchmt)

27   Autotransfusion

499.e1

shock severity, is associated with worse outcome in trauma. J Trauma
transfusion leads to less allogeneic transfusions and is safe. J Trauma Acute
33. Autologous blood transfusions. Council on Scientific Affairs. JAMA 256:
and alternative methods of minimising perioperative allogeneic blood
transfusion: a systematic review and economic model. Health Technol Assess
38. O’Riordan WD: Autotransfusion in the emergency department of a
39.	Hughes DB, Ullery BW, Barie PS: The contemporary approach to the
40. Victorino G, Wisner DH: Jehovah’s Witnesses: unique problems in a
.pdf.
42. The use of autologous blood. The National Blood Resource Education
43. Kruskall MS: Autologous blood transfusion and related alternatives to
allogenic transfusion. In Hillyer CD, Silberstein LE, Ness PM, et al,
editors: Blood banking and transfusion medicine, Philadelphia, 2003, Churchill
Livingstone.
44. Roostar L: Clinical pictures of penetrating chest injuries: infusion therapy
and haemotransfusion. In Gunshot Chest Injuries, Tartu, Estonia, 1996,
Tartu University Press, p 91.
45. Griswold RA, Ortner AB: The use of autotransfusion in surgery of the
47. Özmen V, McSwain NE, Jr, Nichols RL, et al: Autotransfusion of potentially
culture-positive blood (CPB) in abdominal trauma: preliminary data from
50. Timberlake GA, McSwain NE, Jr: Autotransfusion of blood contaminated
by enteric contents: a potentially life-saving measure in the massively
51.	Upton DA, Proehl JA: General principles of autotransfusion. In Proehl
54. Barriot P, Riou B, Viars P: Prehospital autotransfusion in life-threatening
57.	Mattox KL, Espada R, Beall AC: Performing thoracotomy in the emergency
58. Wright CB, Solis RT: Microaggregation in canine autotransfusion. Am
61. Von Koch L, Wilson Defore W, Mattox KL: A practical method of
_drains/edu-ats.asp.


Transfusion of blood components (red cells, white cells, platelets, whole plasma, or plasma fractions) is commonplace in the emergency department (ED). Annually in the United States, 15 million blood donations take place and 14 million units of red blood cells (RBCs) are transfused.1 Although acute hemorrhage is the most common emergency indication for blood transfusion, more nonemergency transfusions and blood component therapy now occur in the ED as a result of the general migration of health care away from inpatient settings. Technical advances have made component therapy directed at specific acute and chronic pathologic conditions practical, safe, and affordable.

BACKGROUND

RBC Antigens and Antibodies

The first documented transfusion took place in the early 1600s, and sporadic advancement in transfusion medicine occurred over the next three centuries, mainly dabbling in cross-species whole blood transfusions. It was not until the early 1900s that the Austrian Karl Landsteiner found that an individual’s serum reacted with the red cells of some but not all other individuals, thereby discovering the red cell antigen-antibody system.

RBC membranes contain a series of glycoprotein moieties, or antigens, that give the cell an individual identity. Two different genetically determined antigens, type A and type B, occur on the cell surface. Any individual may have one, both, or neither of these antigens. Because the type A and type B antigens on the surface of the cell make the RBC susceptible to agglutination, these antigens are termed agglutinogens. The presence or absence of agglutinogens is the basis for the ABO blood group classification, and the blood types are named accordingly as A, B, or AB. Blood type O contains neither the A nor the B agglutigen. These blood type antigens are represented in Fig. 28.1.

Within the first year of life, antibodies begin to form against the standard red cell agglutinogens that are not present in the individual patient. These agglutinins are γ-globulins of the immunoglobulin (Ig) M and IgG types and are probably produced by exposure to agglutinogens in food, bacteria, or exogenous substances other than blood transfusions. In the absence of type A agglutinogens (blood types B and O), anti-A antibodies, or agglutinins, spontaneously develop in the plasma. Similarly, in the absence of type B agglutinogens (blood types A and O), anti-B antibodies develop. When both A and B agglutinogens are present (blood type AB), no agglutinins are formed. Blood groups and their genotypes and constituent agglutinogens and agglutinins are shown in Fig. 28.1.

The reaction between red cell antigens and the corresponding agglutinins results in red cell destruction when noncompatible blood types are mixed. As many as 300 different red cell antigens have been identified, but clinically the A and B antigens are most important; severe, potentially fatal agglutination can occur with the first transfusion of ABO-incompatible blood. The Rh system is likewise very important because there is a chance that transfusion of Rh-positive blood to an Rh-negative patient will result in the formation of Rh antibodies. These antibodies are capable of causing severe hemolysis following a second exposure to the Rh antigen. Of the 40 antigens in the Rh system, D is the most antigenic, but others can also stimulate the production of antibodies in recipients lacking the antigen and thus complicate future transfusions. Other antigen systems in which antibodies could potentially cause hemolytic reactions are the Kell (K and k alleles), Duffy (Fy<sup>+</sup> and Fy<sup>−</sup>), Kidd (Jk<sup>+</sup> and Jk<sup>−</sup>), and MNS (M and N and closely linked S and s) systems. Other antigen systems are rarely of clinical importance in transfusion therapy, except in certain patient populations who may require repeated transfusions, such as those with sickle cell anemia.

Crossmatching

Compatibility testing, or crossmatching, involves mixing the donor’s RBCs and serum with the serum and RBCs of the recipient to identify the potential for a transfusion reaction. The end point of all crossmatches is the presence of RBC agglutination (either gross or microscopic) or hemolysis. Testing is performed immediately after mixing, after incubation at 37°C for varying times, and with and without an antiglobulin reagent to identify surface immunoglobulin or complement. Each unit of blood product, when properly crossmatched, can be administered with the expectation of safety. Full crossmatching takes approximately 45 minutes to complete.

Types of RBC Preparations

Whole Blood

The normal blood volume of a healthy adult and healthy child is approximately 70 mL/kg and 80 mL/kg, respectively. Though intuitively an ideal transfusion agent, whole blood is seldom used except for autologous transfusions (e.g., autotransfusion) and for exchange transfusions. Whole blood is not indicated for the treatment of hypovolemic shock, which can be treated effectively with a combination of crystalloids (e.g., lactated Ringer’s [LR] solution, 0.9% sodium chloride), colloids (e.g., plasma protein, albumin), and packed RBCs (PRBCs). It is also not indicated for the correction of thrombocytopenia, replacement of coagulation factors, or treatment of anemia.2

The incidence of transfusion reactions following transfusion with whole blood is approximately 2.5 times greater than that with PRBCs.3 Whole blood contains antigenic leukocytes and serum proteins, which carry higher risk (approximately 1%) for an allergic reaction. Nevertheless, warm fresh whole blood has seen increased popularity in military settings and has been proposed as an alternative to component therapy for civilian use in massive transfusion protocols. Other methods of whole blood transfusion, including autologous transfusion in massive hemothorax, are being explored as promising alternatives to minimize reaction to whole blood transfusion.4
CHAPTER 28 Transfusion Therapy

Transfusion Therapy

In cases of massive transfusion (>10 units), there is a significant risk for metabolic and respiratory acidosis, as well as hypocalcemia, which can reach life-threatening levels. Although underlying illness or injury obviously plays a major role in the cause of death, the overall mortality of patients requiring massive PRBC transfusions is approximately 60%.7,8

Transfusion of PRBCs is indicated to provide additional oxygen-carrying capacity and expansion of volume. Packed cells are most commonly used to treat acute hemorrhage and anemia that is not amenable to nutritional correction. When treating acute hemorrhage, PRBCs are usually given: (1) if the hemoglobin level falls below established critical levels for that particular given patient population (see the section on Transfusion Thresholds), (2) after rapid crystalloid infusion fails to restore normal vital signs, or (3) concurrently with crystalloid infusion in the treatment of obvious life-threatening blood loss.

Specially prepared or screened types of red cells are listed in the following sections. Their indications for use are presented in Box 28.1.

Washed RBCs

After centrifugation, red cells can be washed to further remove leukocytes, platelets, microaggregates, and plasma proteins.

Packed Red Blood Cells

PRBCs are prepared by centrifugation and removal of most of the plasma from citrated whole blood. One unit of PRBCs contains the same red cell mass as 1 unit of whole blood at approximately half the volume and twice the hematocrit (55% to 80%) in 250 mL of volume.8 One unit of PRBCs raises the hematocrit approximately 3% in an adult or increases the hemoglobin level of a 70-kg individual by 1 g/dL. In children, there is an approximate rise in hematocrit of 1% for each 1 mL/kg of packed cells. For example, if 5 mL/kg of PRBCs is transfused, the hematocrit will rise by approximately 5%. Actual changes depend on the state of hydration and the rate of bleeding. Because most of the plasma has been removed, PRBCs cause fewer transfusion and allergic reactions than whole blood does.

PRBCs contain less sodium, potassium, ammonia, citrate, hydrogen ions, and antigenic protein than whole blood does. This may offer advantages in patients with reduced cardiovascular, renal, or hepatic function. The rate of urticaria is still relatively high at 1% to 3% of transfusions, but the incidence of adverse reactions to packed cells is approximately one third of that noted with whole blood. The benefit of increased hemoglobin must be weighed against the potential for volume, electrolyte, and acid-base imbalances following PRBC administration. In cases of massive transfusion (>10 units), there is a significant risk for metabolic and respiratory acidosis, as well as hypocalcemia, which can reach life-threatening levels. Although underlying illness or injury obviously plays a major role in the cause of death, the overall mortality of patients requiring massive PRBC transfusions is approximately 60%.7,8

Transfusion of PRBCs is indicated to provide additional oxygen-carrying capacity and expansion of volume. Packed cells are most commonly used to treat acute hemorrhage and anemia that is not amenable to nutritional correction. When treating acute hemorrhage, PRBCs are usually given: (1) if the hemoglobin level falls below established critical levels for that particular given patient population (see the section on Transfusion Thresholds), (2) after rapid crystalloid infusion fails to restore normal vital signs, or (3) concurrently with crystalloid infusion in the treatment of obvious life-threatening blood loss.

Specially prepared or screened types of red cells are listed in the following sections. Their indications for use are presented in Box 28.1.

Washed RBCs

After centrifugation, red cells can be washed to further remove leukocytes, platelets, microaggregates, and plasma proteins.

Figure 28.1 Blood group types and their associated antibodies and antigens. (From Abbas AK: Cellular and molecular immunology, ed 7, Philadelphia, 2011, Saunders.)


**BOX 28.1 Indications for the Use of Specialty Red Blood Cells for Transfusion**

<table>
<thead>
<tr>
<th>IRRADIATED</th>
<th>Neonates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients with hematologic malignancies</td>
<td></td>
</tr>
<tr>
<td>Stem cell–transplant patients</td>
<td></td>
</tr>
<tr>
<td>Directed donations from family members</td>
<td></td>
</tr>
<tr>
<td>HLA-matched platelets</td>
<td></td>
</tr>
<tr>
<td>Patients with cellular immune deficiency</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>WASHED</th>
<th>Recipients with a history of severe allergic transfusion reactions</th>
</tr>
</thead>
<tbody>
<tr>
<td>IgA-deficient patients</td>
<td></td>
</tr>
<tr>
<td>Paroxysmal nocturnal hemoglobinuria</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>LEUKOCYTE REDUCED</th>
<th>Multiple-transfused patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multiparous females</td>
<td></td>
</tr>
<tr>
<td>Cancer patients undergoing chemotherapy</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CMV AND EBV SERONEGATIVE</th>
<th>Seronegative patients who are currently pregnant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Premature or low–birth-weight infants</td>
<td></td>
</tr>
<tr>
<td>Bone marrow or organ transplant recipients</td>
<td></td>
</tr>
<tr>
<td>Immunosuppressed patients</td>
<td></td>
</tr>
</tbody>
</table>

CMV, *Cytomegalovirus*; EBV, *Epstein-Barr virus*; HLA, *human leukocyte antigen*; IgA, immunoglobulin A; RBCs, red blood cells.


Washing reduces the titer of anti-A and anti-B, thereby permitting safer transfusion of type O PRBCs into non-O recipients.

**Leukocyte-Reduced RBCs**

Leukocyte-reduced blood products contain less than 5 × 10⁶ leukocytes/unit, whereas standard RBC units contain 1 to 3 × 10⁹ leukocytes. Reduction can be performed at the time of collection, in the transfusion laboratory, or at the bedside during transfusion. Leukocyte-reduced products are used to decrease the likelihood of febrile reactions, immunization to leukocytes, and transmission of disease. Currently, approximately 60% to 75% of the U.S. blood supply is leukoreduced.⁹ Several groups advocate the use of 100% leukocyte-reduced blood products because of the many adverse transfusion reactions that are associated with leukocytes. Non–leukocyte-reduced products are virtually the exclusive method of transmission of several viruses, including human T-lymphotropic virus 1 and 2, *Epstein-Barr virus* (EBV), and *cytomegalovirus* (CMV). Additionally, they help reactivate and disseminate CMV and human immunodeficiency virus (HIV). Moreover, increased rates of bacterial contamination and postoperative and line infections have been associated with the use of non–leukocyte-reduced products. Furthermore, leukocytes lead to human leukocyte antigen (HLA) alloimmunization, which results in increased graft rejection and platelet refractoriness.¹⁰

**Irradiated RBCs**

Blood products can be irradiated to reduce the risk for graft-versus-host disease (GVHD) in susceptible patients. Irradiation destroys the donor lymphocytes’ ability to respond to host foreign antigens. Box 28.1 lists the indications for use of irradiated PRBCs.

**Infectious Complications of Transfusions**

Though relatively uncommon, transmission of infectious diseases is the transfusion-related complication most feared by the lay public. Transmission of a wide variety of infectious diseases has been reported, but modern screening methods have sharply reduced the frequency of transmission. Viral illnesses remain the most problematic.

Between 1985 and 1999, 694 deaths associated with transfusion were reported to the Food and Drug Administration (FDA). Seventy-seven (11.1%) of these deaths were caused by bacterial contamination. However, sepsis is an uncommon occurrence because both the citrate preservative and refrigeration kill most bacteria. Concern over sepsis is responsible for the practice of completing transfusions within 4 hours and returning unused blood products to the blood bank refrigerator for future use only if they have been unrefrigerated for less than 30 minutes. Both gram-negative and gram-positive organisms are transmitted, with gram-negative virulence being more commonly associated with mortality. A prospective observational study found that the rate of nosocomial infections was significantly higher in patients receiving blood transfusion. Leukoreduction did not significantly reduce the rate of infection.¹¹ A 2001 multicenter study by the Centers for Disease Control and Prevention further evaluated the risk for bacterial contamination in the blood pool. The results showed the rate of bacterial sepsis to be much lower than previously thought. Only 0.21 cases and 0.13 deaths per million red cell transfusions occurred. The rate was slightly higher for platelet transfusions, with 10 cases and 2 deaths per million transfusions.¹² Mandatory screening of platelets for bacterial contamination began in 2004 and has further reduced the rate of reported death.

Syphilis may theoretically be transmitted by transfusion, but both refrigeration and citrate markedly reduce the survival of *Treponema pallidum*. Thus, transmission is only a concern with fresh blood or platelet transfusions. The incubation period for syphilis transmitted by transfusion is 4 weeks to 4 months, and the initial clinical manifestation is commonly a rash. No cases of transfusion-transmitted syphilis have been recognized for many years.¹³,¹⁴

The risk for parasitic infection via transfusion is exceedingly low (<1 per 1,000,000), although prospective blood product donors who have been to an endemic region within 12 months or treated with malarial prophylaxis within 3 years are prohibited from blood donation. Those with a history of babesiosis or Chagas disease are permanently barred. Donors with a history of Lyme disease may donate if they are symptom free and have undergone a complete course of treatment.

Viruses are the organisms most likely to be transmitted by transfusion and are the agents with the greatest potential to cause serious disease. They include CMV, EBV, HIV, West Nile virus (WNV), and the hepatitis viruses.

Most blood products have the potential to transmit hepatitis. Routine testing of blood donors for hepatitis C virus (HCV) has occurred since 1991, but the initial screening tests were relatively inaccurate. Since April 1999, the use of nucleic acid amplification
testing (NAAT) to detect HCV RNA has been mandatory. This test has essentially eliminated false positives and has a sensitivity of greater than 99%.15,16 The American Association of Blood Banks (AABB) reported the risk for transmission of HCV to be less than 1 per 1,000,000 transfusions. The incubation period for HCV is 2 to 12 weeks following parenteral infusion. The reported risk for transmission of hepatitis B virus (HBV) is higher at 1 per 137,000 transfusions.

Both CMV and EBV may cause a mononucleosis-like syndrome 2 to 6 weeks after a transfusion. Indications for CMV- and EBV-negative preparations are listed in Box 28.1. Alternatively, leukocyte-reduced products can help protect against CMV and EBV.

The AIDS epidemic has affected transfusion therapy profoundly. In the United States, 3% of AIDS cases have been linked to blood products. The estimated likelihood of transmitting HIV through transfusion is 1 in 1,900,000. Currently, NAAT is used to detect HIV in blood. Because the test detects genetic material in lieu of antibody to the virus, it has significantly reduced the window period during which infection is undetected. Other methods of reducing transmission, including techniques to kill the virus in collected samples (viral inactivation) and the use of blood component substitutes, are being investigated.

Efforts to reduce the risk for transmission of HIV to the general population receiving blood products began early in the course of the epidemic and have had considerable success. Voluntary deferment of blood donation by high-risk groups was encouraged beginning in 1983, and formal screening of all blood products commenced in 1985.

Transmission of WNV by blood transfusion was first documented in the United States in 2002.17,18 Since 2003, universal screening for WNV by investigational NAAT occurs on all blood donations. From 2003 to 2005, 1,400 potentially infectious donations were removed from the blood pool. Since that time, however, multiple cases of transfusion-associated transmission of WNV have been confirmed. This residual risk for transmission is due to blood units with low levels of viremia. Public health authorities continue to look for ways to eliminate this risk from the blood pool.19,20

Emerging infectious risks to the blood supply are always under investigation. Blood-transmitted infections under current surveillance include parvovirus B19, dengue virus, and the prions that cause Creutzfeldt-Jakob disease. Although a viremic phase of human herpesvirus-8, avian flu (H5N1), H1N1, and Lyme disease has been well documented, no cases of transmission through transfusion have been noted.21

A summary of infection risks associated with red cell transfusion is presented in Table 28.1.

### Transfusion Reactions

Transfusion reactions can be divided into two phases: acute and chronic. The vast majority of transfusion reactions occur proximate to or concurrently with the administration of red cells. If the ED is the site for a nonemergency transfusion and the patient is otherwise stable enough for discharge, it is a common practice for the patient to be released shortly after the transfusion is completed.

#### Acute Reactions

**Allergic**

The most common manifestation of a minor allergic transfusion reaction is urticaria; however, wheezing and angioedema can also be observed. The allergic response is due to the presence of atopic substances that interact with antibodies in the donor or recipient plasma, but the severity is not dose related.6 Whenever a transfusion reaction is suspected, the first step in management is to stop the transfusion. Treatment is the same as for other allergic reactions and includes antihistamines, corticosteroids, and intramuscular (IM) or intravenous (IV) epinephrine if needed. For mild reactions (e.g., those limited to skin findings), the transfusion can be resumed once treatment has been given.

**Anaphylactic**

The reported incidence of transfusion-associated anaphylaxis is 1 in 20,000 to 50,000. Anaphylaxis occurs most commonly in IgA-deficient patients who have IgA-specific antibodies of the IgE class.21 Manifestations of an anaphylactic transfusion reaction include shock, hypotension, angioedema, dyspnea, bronchospasm, and laryngospasm. The symptoms are typically rapid in onset and begin within seconds to minutes of starting the transfusion. If this type of reaction occurs, the transfusion

---

**Table 28.1 Estimated Risks of Transfusion per Unit of Packed Red Blood Cells in the United States**

<table>
<thead>
<tr>
<th>RISK</th>
<th>RATE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major allergic reactions</td>
<td>1/100</td>
</tr>
<tr>
<td>Anaphylaxis</td>
<td>1/20,000–50,000</td>
</tr>
<tr>
<td>Anaphylactic shock</td>
<td>1/500,000</td>
</tr>
<tr>
<td>Hemolytic reaction (minor)</td>
<td>1/6000</td>
</tr>
<tr>
<td>Hemolytic reaction (fatal)</td>
<td>1/100,000 allergic reactions</td>
</tr>
<tr>
<td>Death from sepsis (RBC)</td>
<td>1/5 million</td>
</tr>
<tr>
<td>Death from sepsis (platelets)</td>
<td>1/500,000</td>
</tr>
<tr>
<td>Parasitic infections (Lyme, malaria, Chagas)</td>
<td>&lt;1/million—data lacking</td>
</tr>
<tr>
<td>Hepatitis C</td>
<td>&lt;1/million</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>1/140,000</td>
</tr>
<tr>
<td>Parvovirus, Creutzfeldt-Jakob disease</td>
<td>Extremely rare—data lacking</td>
</tr>
<tr>
<td>HTLV 1/2 infection</td>
<td>1/200,000</td>
</tr>
<tr>
<td>HIV infection</td>
<td>1/2 million</td>
</tr>
<tr>
<td>West Nile virus</td>
<td>Extremely rare—data lacking</td>
</tr>
<tr>
<td>CMV/Epstein-Barr</td>
<td>Rare—data lacking</td>
</tr>
<tr>
<td>Acute lung injury</td>
<td>1/500,000</td>
</tr>
<tr>
<td>Graft-versus-host disease</td>
<td>Extremely rare—data lacking</td>
</tr>
<tr>
<td>Immunosuppression</td>
<td>Unknown</td>
</tr>
<tr>
<td>Syphilis</td>
<td>No cases reported currently</td>
</tr>
</tbody>
</table>

CMV: Cytomegalovirus; HIV: human immunodeficiency virus; HTLV: human T-cell lymphotropic virus; RBC: red blood cell.
must be stopped immediately. Treatment includes airway management as necessary, IM or IV epinephrine, fluids, corticosteroids, and antihistamines, followed by appropriate supportive care and continued close observation. If a transfusion is still required, the patient needs to be pretreated with corticosteroids and antihistamines 30 to 60 minutes before the transfusion. Alternatively, or in addition, washed cellular products can be used.

**Febrile (Nonhemolytic) Reactions**
A febrile, nonhemolytic reaction is defined as an increase in temperature of 1°C or higher during or within 6 hours of the transfusion. The mechanism for this type of reaction is most commonly attributed to an interaction between recipient antibodies and donor leukocytes.25 This stimulates the release of cytokines such as interleukin-1, which ultimately produces a febrile response. Although this type of reaction is not life-threatening, it is difficult to distinguish from more serious transfusion reactions. Accordingly, all patients with a fever attributable to a transfusion must have the transfusion stopped. Symptoms can be treated with acetaminophen or nonsteroidal antiinflammatory drugs. There is no role for antihistamines in the treatment of this type of reaction. Although controversy exists, premedication with antipyretics and antihistamines may prevent these transfusion reactions.24

**Acute Hemolytic Reactions**
An acute hemolytic reaction is usually the result of donor-recipient major ABO incompatibility. This in turn is most commonly the result of blood product misassignment related to clerical error. Hemolytic transfusion reactions are estimated to occur once per every 6000 blood units transfused, with a fatality rate of 1 per every 100,000 units transfused.

When incompatible blood is given, the result may range widely from no effect to death. If the recipient does not have antibodies (naturally occurring or acquired) directed against the foreign RBC antigen received, there will be no immediate reaction, but antibodies to the infused blood may develop within weeks, thus limiting the safety of subsequent transfusions from the same donor or same antigenic type. If the recipient’s serum has preformed antibodies directed against the donor RBCs (e.g., an incompatibility in the major crossmatch), the recipient will begin to hemolyze the donor cells within seconds or minutes. In most cases of major crossmatch reactions, RBCs of the donor blood are agglutinated and hemolyzed. It is rare for transfused blood to produce agglutination of the recipient’s cells because the plasma portion of the donor blood becomes diluted by the plasma of the recipient. This reduces the titer of the infused agglutinins to a level too low to cause significant agglutination. Because the recipient’s plasma is not diluted to any significant degree, the recipient’s agglutinins can react with donor cells. The end result of antigen-antibody incompatibility is red cell hemolysis. Occasionally this occurs immediately, but more often the cells first agglutinate. They are then trapped in small vessels and become phagocytized over a period of hours to days and release hemoglobin into the circulatory system.24 Clinical manifestations of acute hemolysis include chills, fever, tachycardia, abdominal pain, back pain, hypotension, fainting, and a feeling of “impending doom.”23 Derived from the liberation of intracellular material associated with hemolysis, vasoactive substances may cause hypotension and shock; other substances may precipitate disseminated intravascular coagulation and high-output cardiac failure. Acute renal failure may also result. The presence of hemoglobinemia and hemoglobinuria is essential in making the diagnosis. A decrease in hematocrit and haptoglobin or an increase in lactate dehydrogenase may also be seen.

Treatment of an acute hemolytic reaction begins with immediate cessation of the transfusion. The blood bank should be alerted immediately because a second patient is now at risk for receiving the wrong product. Resuscitation and supportive care along with close monitoring of laboratory values are essential. A sample of blood from the recipient needs to be obtained for a direct antiglobulin test, plasma-free hemoglobin, and repeated type and crossmatch. Urine can also be tested for free hemoglobin. Renal function and electrolytes should be monitored for evidence of renal failure and hyperkalemia. Dialysis is occasionally required.25,26 Fluid resuscitation and diuresis with normal saline are recommended to maintain urine output above 100 to 200 mL/hr. LR solution should be avoided because calcium can precipitate clotting.

**Drug-Induced Hemolysis**
Drug-induced hemolysis is not a transfusion reaction per se; however, it can be indistinguishable from an acute hemolytic reaction in patients receiving blood transfusions. In this case, both autologous and transfused cells are affected. A patient’s serum can react with red cells in the presence of certain drugs. Two examples of drugs that can cause this type of reaction are cefotetan and ceftriaxone.

**Transfusion-Related Acute Lung Injury**
Transfusion-related acute lung injury (TRALI) refers to noncardiogenic pulmonary edema occurring during or shortly after the transfusion of blood products. A leading cause of transfusion-related mortality and morbidity, TRALI has been reported to occur in as many as 3% of patients receiving transfusions.27,28 TRALI appears to be associated with components from female plasma; preferential distribution of male plasma by the American Red Cross has recently decreased its incidence.29

The potential for TRALI is one reason why some authorities are reluctant to transfuse high ratios of plasma to PRBCs in massive transfusion protocols. TRALI is thought to result from the activation of recipient neutrophils in the lung and the production of vasoactive mediators, which leads to increased pulmonary capillary permeability and leakage. Initial symptoms include respiratory distress, hypoxia, hypotension, fever, and bilateral pulmonary edema; however, the spectrum of TRALI can also include more milder reactions.30,31

Treatment of TRALI is supportive and includes supplemental oxygen, endotracheal intubation, and cardiovascular support as necessary. Diuresis and corticosteroids are not effective.12,33

**Delayed Reactions**
**Delayed Hemolytic Transfusion Reactions**
Even when major and minor crossmatches indicate compatibility, delayed hemolytic transfusion reactions can occur days to weeks after transfusion. This is due to antibody production by either the donor or recipient B cells in response to exposure to antigens on red cells. Usually seen in patients who have had multiple transfusions or in multigravida women, these reactions may be unavoidable without complete RBC antigen typing, a procedure occasionally indicated for recipients of repeated transfusions. An incompatibility in the minor crossmatch does not usually result in a serious reaction, although the recipient’s
red cells can be hemolyzed if the titer of the antibody is sufficiently large. Fortunately, 90% of transfusions are now given as PRBCs, which contain a very small volume of plasma, thus minimizing the chance of a transfusion reaction occurring as a result of donor sensitization.

The signs and symptoms of a delayed hemolytic reaction include low-grade fever, a decrease in hemoglobin, mild jaundice, a positive direct antiglobulin test, and elevation of lactate dehydrogenase.

Treatment of a delayed hemolytic reaction is not needed unless there is evidence of brisk hemolysis. In the case of brisk hemolysis, treatment consists of fluids, antigen-negative (type O) blood transfusions, or red cell exchange.

Graft-Versus-Host Disease
GVHD is a transfusion complication most commonly associated with allogeneic hematopoietic cell transfusions. However, it can occur whenever immunologically competent lymphocytes are transfused, especially in immunocompromised hosts. Donor lymphocytes engraft in the recipient and then attack host tissue. Symptoms are typically observed 7 to 14 days after the transfusion and include fever, rash, and diarrhea. Hepatitis and marrow aplasia also occur. GVHD is often fatal; failure of the host’s marrow leads to overwhelming infection or bleeding. The use of gamma-irradiated cellular components prevents this complication by making the donor lymphocytes incapable of proliferating.

Posttransfusion Purpura
In rare cases, profound thrombocytopenia can develop 1 to 3 weeks after a transfusion associated with an antibody response to a platelet antigen. A probable pathophysiologic mechanism for this is the production of low-affinity antibodies that cross-react with autologous platelets. Eventually, as the immune response matures, the low-affinity antibody is eliminated and the thrombocytopenia resolves spontaneously. Only patients at risk for bleeding or hemorrhage need to be treated. Treatment consists of high-dose immune globulin, plasmapheresis, or platelet transfusion.

Miscellaneous Transfusion Issues

Transfusion Thresholds
PRBCs are a precious commodity. Guidelines to limit transfusions to those that are absolutely necessary have set transfusion thresholds or “triggers.” A liberal transfusion trigger is approximately 10 g/dL of hemoglobin, whereas restrictive thresholds are set at 7 to 8 g/dL. The limits for restrictive thresholds stem from the finding that aerobic metabolism can still occur at hemoglobin concentrations as low as 5 g/dL.

Clinically, however, almost all patients show signs of physiologic stress at hemoglobin concentrations of less than 6 g/dL.

It is at this level that patients will begin to reliably demonstrate the symptoms and signs of anemia: dyspnea on exertion or even at rest; pallor, particularly of the palms and mucus membranes; and resting tachycardia.

The Transfusion Requirements in Critical Care study compared a strategy of restrictive transfusion triggers with conventional, more liberal triggers. The authors concluded that a restrictive strategy appears to be at least as effective as a more liberal strategy, with the possible exception of patients with coronary insufficiency. In trauma patients, more liberal use of blood has also been questioned. One of the risks associated with restrictive transfusion thresholds is an increased incidence of infection. Nonetheless, mortality, cardiac events, and length of hospital and intensive care unit (ICU) stay appear to be unaffected by more restrictive thresholds. More recently, in a study of patients with acute upper gastrointestinal (GI) bleeding, Villanueva and colleagues found that using a transfusion threshold of 7 g/dL versus 9 g/dL resulted in decreased incidence of further bleeding, decreased adverse events, and increased survival at 6 weeks. No single criterion should be used as an indication for red cell component therapy. Multiple factors related to the patient’s clinical status and oxygen delivery needs should be considered. Current practice focuses on the needs of the individual patient. Particularly close attention should be paid to the subset of patients at risk for coronary ischemia, with more liberal triggers possibly being applied to these patients. In a U.S.-based study of Medicare patients with acute myocardial infarction, Wu and colleagues found RBC transfusions to be beneficial in elderly patients when hematocrit values were lower than 33%. In the setting of severe sepsis a more conservative threshold of 10 g/dL may also be appropriate.

In contrast, new data from pediatric ICU settings suggest that adopting a threshold of 7 g/dL imparts no worse clinical outcome and may result in benefits in long-term mortality and morbidity. Similar findings were documented in pediatric postsurgical patients.

In addition to cost and transmitted infections, there is a risk for systemic inflammatory response syndrome with transfusion. This syndrome is closely associated with diminished organ function and mortality in critically ill adult patients and its
incidence increases with the administration of more than 4 units of PRBCs. Red cell administration is also associated with an increased risk for life-threatening acute respiratory distress syndrome and multiple-organ failure. This has been linked to immunologic alterations and their effects on plasma cytokine and cytokine receptor concentrations. In patients receiving more than 15 units of PRBCs, levels of both interleukin and soluble tumor necrosis factor were elevated. It remains unclear whether the use of leukocyte-reduced PRBCs can mitigate the expected inflammatory response.

The judicious and restricted administration of PRBCs appears to be clinically founded in the majority of patients. By implementing a restrictive transfusion strategy, the probability of a patient requiring blood can be decreased by 42% and the volume of PRBCs transfused can be decreased by 0.93 units per patient.

Another area of study focuses on the projected need for administration of PRBCs in any given patient. Knowing which patients will probably need blood based on initial findings can be helpful in resource allocation and determination of the need for crossmatching. Studies have correlated the base deficit (BD) with the need for transfusion by using a cutoff of –6 mEq/L. Patients with a BD larger than –6 mEq/L have a 72% chance of requiring blood, whereas those with BDs smaller than –6 mEq/L have only an 18% chance of requiring PRBCs. More elaborate scales have been proposed based on easily assessable parameters. The emergency transfusion score is a point system based on systolic blood pressure, the presence of free fluid on focused assessment with sonography for trauma (FAST), an unstable pelvic ring fracture, advanced patient age, admission from the scene, motor vehicle collision, or a fall as being predictors of future transfusion requirements.

It is impossible to define strict PRBC transfusion recommendations in the ED setting. Such decisions must be made in real time after considering multiple factors, some of which may not be known at the time. In summary, definitive data are lacking at this time and no dogmatic recommendations can possibly apply to all ED scenarios. As a general concept, however, the likelihood of a transfusion being of benefit is high when the patient's hemoglobin level is less than 6 to 7 g/dL and low when it is greater than 10 g/dL. The correct strategy is unclear when the hemoglobin level is between 7 and 10 g/dL. Continued blood loss of varying degrees renders transfusion strategies even more obscure. The elderly and those with cardiovascular or respiratory disease may not tolerate anemia as well as those without these parameters.

**Massive Transfusion**

Massive transfusion is loosely defined. In the 1970s it was considered to be the transfusion of more than 10 units of blood to an adult, equivalent to 1 blood volume, within 24 hours. Historically, massive transfusion was associated with dismal survival rates ( <10%). As blood banking technology and storage methods have improved, the mortality associated with massive transfusions has decreased significantly. There is no clear physiologic threshold to define a massive transfusion. Mortality in patients receiving fewer than 3 units is currently approximately 10%, in those receiving 6 to 9 units it is approximately 20%, and in those receiving 10 or more units it is greater than 50%. Alternative triggers for initiation of a massive transfusion protocol are now being proposed, including elevation of the international normalized ratio (INR) and BD, in addition to hemoglobin. Despite the challenges of treating the expected posttransfusion inflammatory and immunologic complications, patients requiring massive transfusions can have good outcomes.

**Transfusion Coagulopathy**

Pathologic hemostasis occurs following massive blood transfusions. Coagulopathy and subsequent uncontrolled bleeding are major contributors to trauma-related deaths. Although such abnormalities rarely develop within the time frame of the initial resuscitation in the ED, an understanding of the problem can lead to a more thoughtful approach to transfusion practices and the anticipation of potential problems. Significant alterations in blood and blood products occur during storage. Moreover, in patients who are given a transfusion equal to 2 blood volumes, only approximately 10% of the original elements remain. The development of transfusion coagulopathy is multifactorial; important factors include tissue injury, acidosis, the duration of shock, and hypothermia, in addition to activation, consumption, and dilution of coagulation factors.

Transfusion coagulopathy is also related in part to dilution of the recipient’s platelets by transfused blood devoid of functioning platelets. Dilutional thrombocytopenia is a well-recognized complication of massive transfusion, and a platelet count should be obtained routinely if more than 5 units of blood are transfused.

Disseminated intravascular coagulation (from a hemolytic reaction) may play a secondary role in posttransfusion bleeding. Factors V and VIII are labile in stored blood and absent in packed cells. Fibrinogen is relatively stable in stored blood but is absent in packed cells. A deficiency of most clotting factors, especially factors V and VIII and fibrinogen, occurs with massive transfusions. This deficiency probably occurs on a “washout” (i.e., dilutional) basis, although the dynamics are poorly understood. Replacement of these factors may be required. Specific assays for the individual factors are available, but it is more practical to measure the prothrombin time (PT), partial thromboplastin time (PTT), and fibrinogen levels. Plasma has been used to correct clotting factor abnormalities secondary to dilution from massive transfusions, but its effectiveness has not been firmly established. Cryoprecipitate has also been used to replace factor VIII and fibrinogen, but it is rarely required because plasma contains some fibrinogen. Fresh frozen plasma (FFP) should be infused to correct the coagulopathy as indicated by clotting studies. Cryoprecipitate may be required if fibrinogen levels fall below 100 mg/dL despite the use of plasma. Although blood component therapy can be based on measured coagulopathy parameters, as a general guide 1 to 2 units of plasma for each 5 to 6 units of blood may be given empirically.

Numerous massive transfusion protocols exist. Traditionally, transfusion-related coagulopathies have been evaluated and treated as per laboratory indicators, but rapid or massive transfusions do not allow equilibration or timely laboratory analysis. Although this approach is quite acceptable in most patients, the aim of transfusion protocols is to prevent transfusion-related coagulopathy before it occurs. Obviously one cannot simply continue to transfuse only PRBCs to patients experiencing significant blood loss, and some combination or ratio of plasma to PRBCs to platelets should be adopted. The ideal combination is, however, not known with certainty, and it is largely dependent on the underlying need for transfusion.
and specific patient characteristics. Most protocols recommend a plasma-to-PRBC ratio of 1:1.5 or 1:1.8.67 Other protocols also include empirical platelet administration and use of an RBCs to platelets to FFP ratio of 1:1 : 1.68 A 2015 study, the Pragmatic, Randomized Optimal Platelet and Plasma Ratios (PROPPR) trial, compared the transfusion strategy of plasma, platelets and RBCs in a 1 : 1 : 1 ratio versus a 1 : 1 : 2 ratio. In patients with major trauma and severe bleeding they found no significant differences in mortality at 24 hours or 30 days using either strategy. They did note in the 1 : 1 : 1 group, “more patients achieved hemostasis and fewer experienced death due to exsanguination by 24 hours.”69

At this time, however, no specific transfusion ratio has been proven superior. Strict adherence to any protocol must be balanced against the risk for multisystem organ failure and infection associated with high doses of platelets and plasma. All protocols recommend warming of blood and blood products because hypothermia occurs quickly during massive transfusions and can contribute to further coagulopathy.

Severe Trauma and Coagulopathy
A transfusion coagulopathy often develops in individuals who are injured during military combat and receive transfusions because of widespread tissue trauma. The U.S. military has advocated an approach to transfusion therapy that includes prompt initiation of 1 : 1 : 1 resuscitation ratios with RBCs, prethawed universal-donor AB plasma, and apheresis platelets, with conversion to fresh whole blood as soon as it can be obtained. This ratio has not been universally adopted by civilian hospitals.

Emergency Transfusions
In an emergency, three alternatives to fully crossmatched blood exist. The preferred substitute is type-specific blood with an abbreviated crossmatch. The abbreviated crossmatch includes ABO and Rh compatibility. In addition, the recipient’s serum is screened for unexpected antibodies, and an immediate “spin” crossmatch is performed at room temperature. This abbreviated crossmatch requires approximately 30 minutes. Many institutions are now using this procedure as their standard crossmatch for most patients. The safety and utility of the type-specific abbreviated crossmatch have been demonstrated repeatedly, with transfusion reactions occurring only rarely.70,71 Brickman and coworkers demonstrated that bone marrow aspirates obtained with an intraosseous (IO) needle can also be used for crossmatching.72

The second preference for an alternative to fully crossmatched blood is type-specific blood that is only ABO and Rh compatible, without screen or immediate spin crossmatch. The patient’s ABO group and Rh factor can be determined within 2 minutes, and in an emergency, typing of the blood group and the Rh factor is all that is necessary before transfusion. Type-specific blood that has not been crossmatched has been used in numerous military and civilian series without serious consequences. While the type-specific blood is being transfused, the antibody screen and crossmatch are carried out in the laboratory. The transfusion should be stopped if an incompatibility is found.

A third alternative to fully crossmatched blood is group O blood, although type-specific blood is generally preferable.67 Commonly, this is available at the point of care and has the advantage of being immediately available in cases of severe shock with ongoing bleeding. Thus, despite the theoretical preference for type-specific blood in emergency situations, type O is often a reasonable and practical alternative.

One may transfuse both Rh-positive and Rh-negative group O packed cells into patients who are in critical condition. It is a common misconception that patients who are Rh negative will have an immediate transfusion reaction if given Rh-positive blood. There is no particular advantage in determining the Rh factor because preformed, naturally occurring anti-Rh antibodies do not exist. Theoretically, individuals who are Rh negative may become sensitized either through pregnancy or by previous transfusions, and a delayed hemolytic transfusion reaction will result if Rh-positive blood is transfused. However, this scenario is very rare and is of little significance when compared with life-threatening blood loss. Sensitization to the Rh factor is most problematic for Rh-negative women of reproductive age.74,75 Any sensitized patient may experience a transfusion reaction if exposed again to Rh-incompatible blood. However, significant subsequent transfusion reactions with Rh-incompatible blood in men sensitized to the Rh factor are very rare. Many advise routine use of the more widely available type O Rh-positive packed cells in all patients in whom the Rh factor has not been determined, except in females of childbearing age, for whom Rh sensitization may be an important consideration. Once resuscitated with Rh-positive packed cells, patients may receive their own type without a problem. Because individuals with type O Rh-negative blood represent only 15% of the population and the blood may be in short supply, it is reasonable to save type O Rh-negative blood for Rh-negative females of childbearing potential and to use type O Rh-positive packed cells routinely as the first choice for emergency transfusions. In a study of emergency blood needs, Schmidt and colleagues reported 601 units of blood into 262 untyped patients, including 8 Rh-negative women, before the blood type was determined.76 No acute hemolytic reactions occurred and no women were sensitized. A non–emergency-based study found the rate of Rh sensitization in Rh-negative recipients receiving Rh-positive blood to be approximately 8% and this figure may be reduced if Rh immune globulin is given after transfusion.77,78 Thus, prophylaxis with Rh immune globulin is recommended only for Rh-negative women of childbearing potential receiving Rh-positive blood.

Rh immune prophylaxis with Rh(D) human immune globulin (RhoGAM) is also indicated for Rh-negative pregnant women who may be bearing Rh-positive children and may have fetomaternal transplacental hemorrhage, including bleeding in early pregnancy, such as spontaneous or elective abortion, ectopic pregnancy, and other potential causes of antepartum hemorrhage such as trauma. RhoGAM suppresses the immune response of Rh-negative women to Rh-positive RBCs and is effective when given up to 72 hours after exposure to fetal erythrocytes. Standard doses are 50 µg for women up to 12 weeks of pregnancy and 300 µg in the second and third trimester. In the setting of fetal-maternal transfusion greater than 15 mL (usually only in the third trimester when fetal blood volume becomes more substantial), higher doses may be necessary. In such circumstances, the correct dose may be calculated by quantitative testing for fetal erythrocytes in the mother’s blood (Kleihauer-Betke test).79

If non-crossmatched blood is transfused, the laboratory should receive a plain (e.g., without a serum separator) red-topped tube of venous blood as soon as possible to begin a formal crossmatch procedure. Whenever possible, this sample should be drawn before any blood is transfused.
Metabolic Disturbances

RBCs undergo metabolic, biochemical, and molecular changes during storage that are collectively known as the erythrocyte “storage lesion.” These changes are generally subtle but can be measured: a decrease in levels of 2,3-diphosphoglycerate (2,3-DPG), pH, and intracellular potassium and a concomitant increase in supernatant potassium.

Theoretically, citrate salts, which are the usual anticoagulants in donor blood, may combine with ionized calcium in plasma and produce hypocalcemia and rarely hypocalcemic-related cardiovascular depression. In clinical practice the hemodynamic consequences of citrate-induced hypocalcemia are minimal, although the QT interval may be prolonged on the electrocardiogram with citrate infusion. Supplemental calcium administration is not usually necessary, even during massive blood replacement, as long as circulating volume is maintained because the liver is able to remove citrate from the blood within a few minutes. Alterations to this recommendation may be necessary in patients with severe liver disease. If calcium replacement is deemed necessary by clinical judgment, 10 to 20 mL of calcium gluconate may be given intravenously, via a different vein, for each 500 mL of blood transfused. If calcium chloride is used, only 2 to 5 mL per 500 mL of blood should be given. Calcium chloride may be preferable in patients with abnormal liver function, such as those with bleeding esophageal varices, because citrate metabolism is decreased, which results in slower release of ionized calcium. Care must be taken to avoid administering too much calcium and inducing hypercalcemia, ideally by monitoring the ionized calcium concentration.

Directed and Autologous Donations

The system of “directed donations” by which friends or family members may donate blood for a specific individual has been proposed in response to concerns about the transmission of infectious disease. At this time, directed donation systems are in place in some institutions but the practice has not been widely supported. Directed donations probably do not decrease the risk for infectious disease transmission and may disrupt the normal anonymous blood donor system and thus leave fewer units available for other needy patients.

Although autologous donations are of limited clinical applicability in emergencies, they are commonplace in elective surgery. It has been suggested that up to 10% of the blood supply could be provided through this mechanism. However, current studies show that the peak of autologous donations represented less than 2% of the total blood collections and this number is declining. Applications at this time include elective cardiac, gynecologic, orthopedic, and vascular surgery. Benefits include avoidance of exogenous blood-borne disease and sensitization. An individual can donate 1 unit of blood weekly until 3 days before surgery. Because blood can be stored for up to 35 days, donations usually begin 5 weeks before needed. The blood donor will require iron supplements and must maintain a hemoglobin level higher than 11 g/dL.

RBC Substitutes

Concerns over infection, availability, storage difficulties, and risk for transfusion reactions have fueled interest in the development of blood substitutes. The ideal blood substitute should: (1) deliver oxygen efficiently, (2) require no compatibility testing, (3) cause few or no side effects, (4) have prolonged storage qualities, (5) persist in the circulation, and (6) be affordable. Blood substitutes can be categorized as synthetic emulsions and hemoglobin-based oxygen carriers (HBOCs) or “stromal-free” hemoglobin solutions.

HBOCs can be derived either from humans (typically from outdated human PRBCs) or from animals, notably bovine. HBOCs are created by lysing red cells, extracting the hemoglobin, and then chemically cross-linking single hemoglobin molecules to create a larger molecule, one less likely to cause impairment in renal function by obstruction of the renal tubules.

Early HBOCs had a propensity to cause renal injury, but this has markedly improved in subsequent generations. Advantages of HBOCs over packed red cells include a much longer shelf life (1 to 4 years) and no need for crossmatching.

In some cases HBOCs are used to provide a bridge therapy for patients who cannot receive transfusions but in whom erythropoiesis-stimulating agents may be effective in 2 to 4 days. An important disadvantage is a much shorter circulating half-life (1 day versus 31 days).

Perfluorocarbons are carbon-fluorine compounds that have a high oxygen- and carbon dioxide–dissolving capacity. They are not miscible with water and must therefore be emulsified for administration. Perfluorocarbons are totally synthetic, essentially limitless in supply, chemically stable, and harbor no risk for infection. However, they are limited by a short half-life, with degradation occurring within 48 hours of administration. Trials to date have focused mainly on their adjunctive use with standard transfusion therapy. Preliminary studies are examining earthworm and marine worm hemoglobins, as these creatures survive and oxygenate without RBCs. This work is in the very early stages but may prove promising in the future.

Other potential blood analogues in the investigation phase include biodegradable micelles and hemerythrin, an oligomeric protein responsible for transport of oxygen in the marine invertebrate. Alternative sources for red cell therapy include stem cells and placental umbilical cord blood as another stem cell analogue to create greater supplies of transfusion-safe blood.

OTHER BLOOD PRODUCTS

The individual blood components are discussed in the following sections. A summary of the dosages and characteristics of each component is provided in Table 28.2.

Platelet Concentrates

Platelet concentrates are prepared by rapid centrifugation of platelet-rich plasma. Platelets are obtained by single-donor apheresis or from pooled random-donor whole blood units. Platelets obtained by single-donor apheresis have the advantage of exposing the recipient to only one donor. This reduces the risk for exposure to many different donors and confers a lower risk for transfusion-transmitted disease and other complications. HLA-matched platelets may be used when HLA antibodies develop as a result of repeated random-donor platelet transfusions. Platelet concentrates contain most of the platelets from 1 unit of blood in 30 to 50 mL of plasma.

Confusion sometimes arises about how to order platelets. Platelets are typically ordered by the “pack.” Many institutions use a “6-pack” composed of 6 individual units of platelets.
The issue of prophylactic platelet transfusion remains controversial. Spontaneous bleeding rarely occurs if the platelet count is greater than 10,000 to 20,000/mm³. Even in the event of surgery or trauma, excessive bleeding is uncommon in patients whose platelet count exceeds 50,000/mm³. It is generally recommended that active hemorrhage be treated by platelet transfusion if the platelet count is lower than 50,000/mm³, but prophylactic transfusion may be withheld safely until the count is lower than 20,000/mm³, and more recent data suggest that this threshold can be lowered to less than 10,000 or even 5000/mm³.

Patients with idiopathic thrombocytopenic purpura (ITP) may be severely thrombocytopenic. Despite low platelet counts in ITP, it is uncommon to see spontaneous bleeding. Because of the presence of antiplatelet antibodies in these patients, transfused platelets may last only a short time (minutes to hours) before being removed from the circulation. Nonetheless, patients with ITP and life-threatening bleeding or severe head trauma should be given platelets at two to three times the normal dose.

Platelet transfusion should be avoided in patients with thrombotic thrombocytopenic purpura (TTP), a rare microangiopathic hemolytic anemia, because platelets will result in further microemboli. TTP may be suspected in patients with both anemia and thrombocytopenia but not leukopenia. Clinical

<table>
<thead>
<tr>
<th>BLOOD COMPONENT</th>
<th>DOSE</th>
<th>NOTES</th>
<th>EXPECTED RESULTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Platelets</td>
<td>1 unit/10 kg or 1 apheresis pack or a 6 (4)-pack of pooled platelets</td>
<td>ABO and Rh compatible is preferred but in an emergency, is not necessary. 30–50 mL/unit = 180–300 mL/6-pack Available in 5–15 min</td>
<td>Each unit should raise the platelet count by 5000–10,000, so a 6-pack or 1 apheresis unit should raise the count by 30,000–60,000.</td>
</tr>
<tr>
<td>Fresh frozen plasma</td>
<td>15 mL/kg or = 1000 mL or 4 units in a 70-kg patient</td>
<td>ABO compatibility is desirable but not required in an emergency, and Rh compatibility is never required. 200–250 mL/bag Should begin with 4 units minimum to correct an elevated INR Not possible to get an INR &lt;1.5 with FFP</td>
<td>Each unit raises all coagulation factors by 2%–3% in average-sized adults.</td>
</tr>
<tr>
<td>Cryoprecipitate</td>
<td>10–20 bags, depending on the indication</td>
<td>ABO compatibility is desirable but not required in an emergency, and Rh compatibility is never required. Hypofibrinogenemia &lt;100 mg/dL: give 10 bags, 15–20 mL/bag Fibrinolytic-induced bleeding: give 10–12 bags Factor VIII deficiency when specific factor therapy is not available: 1 bag/5 kg</td>
<td>With fibrinolytic-induced bleeding, recommended doses may help correct bleeding. In factor VIII deficiency, the recommended dose will raise factor VIII to 50% of normal.</td>
</tr>
</tbody>
</table>

ASA, Acetylsalicylic acid; FFP, fresh frozen plasma; INR, international normalized ratio; ITP, idiopathic thrombocytopenic purpura.
features often include fluctuating neurologic symptoms and signs, jaundice, renal dysfunction, and fever. Laboratory confirmation includes the identification of intravascular hemolysis on a peripheral blood smear (e.g., schistocytes), as well as other laboratory evidence of hemolysis. Treatment consists of plasma exchange, although emergency treatment may begin with an infusion of FFP.106

Limited retrospective studies suggest an increase in morbidity and possibly increased mortality in patients with traumatic intracranial hemorrhage (ICH) who are taking platelet inhibitors such as aspirin and clopidogrel, in comparison to controls who are not taking antiplatelet agents.27 One ex vivo study involving 11 healthy volunteers found that complete normalization of platelet function could be obtained in patients taking platelet inhibitors by giving 10 to 15 random-donor units (two to three 4- or 6-packs) or 2 to 3 single-donor apheresis units.29 Another in vitro study (2015) looked at reversal of the effects of a three-drug antiplatelet therapy (aspirin, clopidogrel, and vorapaxar [Aralez Pharmaceuticals, Mississauga Ontario, Canada]) via the administration of platelets. In this study, the administration of a larger amount of platelets (up to 10 random-donor units) was able to reverse some of the antiplatelet effects of this three-drug combination.29 Unfortunately, to date there have been no in vivo studies looking at outcomes in patients receiving platelet infusions for antiplatelet-associated bleeding.

At this time, no established guidelines exist for managing bleeding catastrophes in patients on antiplatelet agents, and the AABB states: “The AABB cannot recommend for or against platelet transfusion for patients receiving antiplatelet therapy who have intracranial hemorrhage (traumatic or spontaneous).”100 Given the devastating nature of ICH in patients taking antiplatelet agents, the benefits of large-volume platelet transfusions may outweigh the risks; however, this is still unclear (Table 28.3).

Crossmatching is unnecessary for platelet transfusion, but the donor and the recipient should be ABO and Rh compatible. Note that platelet concentrates contain enough RBCs to sensitize an Rh-negative individual. In an emergency situation, if ABO-compatible platelets are not available, unmatched platelets can be transfused. This will reduce the number of platelets available from the transfusion but otherwise does not cause the same reaction that is expected with an incompatible red cell transfusion. Most institutions have a policy that limits the amount of incompatible platelets that can be given.101 With massive blood transfusions there may be a diluting effect of the platelet count, resulting in thrombocytopenia. When more than 10 units of blood are transfused, the platelet count must be routinely evaluated and platelets must be replaced accordingly. Clinically significant platelet depletion rarely occurs if less than 15 units of blood (or 1.5 to 2 times the blood volume) has been transfused.102

Each 4 to 6 units of platelets contains 250 to 350 mL of plasma (=1 unit of FFP), which includes coagulation factors and may reduce the requirement for FFP. Platelets may be infused rapidly (1 unit/10 min) with the use of specialized platelet filters.

**Fresh Frozen Plasma**

FFP is prepared by separating plasma from the cellular components of single-donor whole blood, followed by rapid freezing and storage at 18°C or lower. Freezing preserves the soluble coagulation factors of the contact activation (intrinsic) and tissue factor (extrinsic) clotting systems, including the labile factors V and VIII. FFP also contains fibrinogen, though not as much as cryoprecipitate does. Plasma stored for 3 months retains approximately 60% of the normal factor VIII activity and has a shelf life of up to 1 year. Thawed solvent- or detergent-treated plasma stored at 4°C for 6 days still contains sufficient coagulant activity of factors II, V, VII, VIII, IX, XI, and XII; fibrinogen; antithrombin; protein C; and von Willebrand factor (vWF) antigen and can be safely administered.103 Ideally, transfused plasma should be compatible with the recipient’s ABO group. Rh compatibility is not considered essential.104 Each unit of FFP has a volume of approximately 200 to 250 mL. The INR of a unit of FFP is approximately 1.5. Transfusion of even very large amounts of FFP into a patient with an elevated INR will not correct the INR to below 1.5. It is not clinically useful to give FFP to patients with an INR lower than 1.7.105

Because each unit of FFP increases levels of all coagulation factors by 2% to 3% in an average-sized adult, it is useful in clinical scenarios involving depletion of clotting factors. FFP may also be valuable in patients with hereditary or acquired clotting abnormalities, such as a deficiency of factor II, V, VII, X, XI, or XIII; von Willebrand syndrome; hemophilia A (factor VIII deficiency); hemophilia B (factor IX deficiency); or hypofibrinog enemia. However, because of the large volume of FFP that is generally required, its effectiveness is limited in these inherited diseases with severe clotting abnormalities. Specific factor replacement, when available, is always preferred over FFP.

<table>
<thead>
<tr>
<th>TABLE 28.3</th>
<th>Summary and Dosage of Reversal Agents for Platelet Inhibitors</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>AGENT</strong></td>
<td><strong>DOSE</strong></td>
</tr>
<tr>
<td>Platelet transfusion</td>
<td>2 to 3 units of pooled platelets or 2 to 3 apheresis packs</td>
</tr>
<tr>
<td>DDAVP (Desmopressin)</td>
<td>0.3 µg/kg IV</td>
</tr>
</tbody>
</table>

*DDAVP, 1-Deamino-(8-µ-arginine)-vasopressin; IV, intravenous.*
FFP is also indicated for clotting factor deficiencies resulting from massive blood replacement. However, pathologic hemorrhage after massive transfusions is often caused by thrombocytopenia rather than by a depletion of clotting factors. More aggressive FFP replacement formulas are becoming commonplace, rather than the accepted 1 unit of FFP for every 5 to 6 units of PRBCs (see earlier section on Massive Transfusion).

Although FFP can be used for rapid reversal of serious acute bleeding from warfarin anticoagulants or for prophylaxis before surgery or an invasive procedure, the trend is moving toward the use of prothrombin complex concentrates (PCCs) for this indication. When used, the timing of FFP administration is important. In a study on warfarin-related ICH, each 30-minute delay in administering the first dose of FFP translated into a 20% lower chance of reversing the patient's coagulopathy within the first 24 hours.106 In an emergency situation, 10 to 15 mL/kg of FFP will effect a rapid reversal of the vitamin K-dependent factors II, VII, IX, and X. As a general rule of thumb, in a life-threatening situation 4 units of FFP should be considered a starting dose. It is appropriate to use FFP to treat the acquired deficiency of multiple factors, such as seen in patients with severe liver disease, disseminated intravascular coagulation, or vitamin K depletion, and for plasma exchange in those with TTP or hemolytic-uremic syndrome. It is not appropriate for volume expansion or enhancement of wound healing.

Reactions to FFP include fever, chills, and allergic responses, and recipients have a risk for HIV infection and hepatitis similar to the risk with whole blood. FFP should be infused rapidly after thawing because the clotting factors are labile and rapidly lost.

One less common but very serious complication of the administration of blood products containing plasma (FFP and platelets) is TRALI. It is believed to be an immune-mediated process and can occur in up to 1 in 5000 transfusions containing plasma. It carries a mortality of 6% to 9%. Although it is more commonplace, rather than the accepted 1 unit of FFP for every 5 to 6 units of PRBCs (see earlier section on Massive Transfusion), although FFP can be used for rapid reversal of serious acute bleeding from warfarin anticoagulants or for prophylaxis before surgery or an invasive procedure, the trend is moving toward the use of prothrombin complex concentrates (PCCs) for this indication. When used, the timing of FFP administration is important. In a study on warfarin-related ICH, each 30-minute delay in administering the first dose of FFP translated into a 20% lower chance of reversing the patient's coagulopathy within the first 24 hours.106 In an emergency situation, 10 to 15 mL/kg of FFP will effect a rapid reversal of the vitamin K-dependent factors II, VII, IX, and X. As a general rule of thumb, in a life-threatening situation 4 units of FFP should be considered a starting dose. It is appropriate to use FFP to treat the acquired deficiency of multiple factors, such as seen in patients with severe liver disease, disseminated intravascular coagulation, or vitamin K depletion, and for plasma exchange in those with TTP or hemolytic-uremic syndrome. It is not appropriate for volume expansion or enhancement of wound healing.

Reactions to FFP include fever, chills, and allergic responses, and recipients have a risk for HIV infection and hepatitis similar to the risk with whole blood. FFP should be infused rapidly after thawing because the clotting factors are labile and rapidly lost.

One less common but very serious complication of the administration of blood products containing plasma (FFP and platelets) is TRALI. It is believed to be an immune-mediated process and can occur in up to 1 in 5000 transfusions containing plasma. It carries a mortality of 6% to 9%. Although it is more common with plasma products, it can also occur with RBC transfusions because of the small amount of residual plasma in PRBCs.107,108 It was described earlier in the section on Transfusion Reactions.

Start by giving 4 units of FFP if the PT is greater than 1.5 to 1.7 times normal or the activated PTT is greater than 1.5 times the top normal value.109 Each 5 to 6 units of platelets contain the equivalent of 1 unit of FFP, so concomitant platelet infusions may lower the requirements for FFP. In critically ill patients with acute hemorrhage and suspected coagulopathy (e.g., end-stage liver disease), it is appropriate to begin empirical treatment before the laboratory values are known.

**Cryoprecipitate**

Cryoprecipitate is prepared from single-donor plasma by gradual thawing of rapidly frozen plasma. This process causes precipitation of proteins rich in fibrinogen, as well as factor VIII. Each unit of cryoprecipitate typically yields 100 to 250 mg of fibrinogen, 80 to 100 units of factor VIII, and 50 to 60 mg of fibronectin.110 Cryoprecipitate is a plasma product and therefore requires ABO compatibility (see earlier discussion in the section on Fresh Frozen Plasma). The volume of each bag unit is approximately 15 to 18 mL.

Cryoprecipitate is indicated for the treatment of patients with fibrinogen deficiency, congenital afibrinogenemia, dysfibrinogenemia, and factor XIII deficiency and in some patients with hemophilia A or von Willebrand's disease.111,112 It can also be used as a second-line treatment to correct a deficiency in coagulation factor VIII (in hemophilia A) when factor VIII concentrates are not readily available. Because cryoprecipitate contains no factor IX, it is of no value in the treatment of factor IX deficiency (hemophilia B).

Mild deficiencies in factor VIII are defined as 10% to 30% of normal activity and severe deficiencies as less than 3% of normal activity. When treating bleeding, the goal depends on the site and severity of hemorrhage, but in general one should aim for at least 50% of normal factor VIII activity. For life-threatening hemorrhage, aim for 100% activity. The amount of cryoprecipitate required to correct coagulation defects ranges from 10 to 20 units/kg for minor bleeding, such as hemarthrosis, to 50 units/kg for control of bleeding in surgery or trauma. Guide specific replacement by laboratory assay of factor VIII activity. The half-life of factor VIII in plasma is 8 to 12 hours.

One bag of cryoprecipitate per 5 kg of body weight will raise the recipient's factor VIII level to approximately 50% of normal. In a 70-kg patient, this equals 14 bags. The large number of units that must be given increases the chance of exposure to blood-borne diseases. Factor VIII concentrate is a better choice because of improved methods of viral inactivation and the availability of factor VIII prepared with recombinant DNA technology.

The vWF in cryoprecipitate degrades during storage, thus leading to variable amounts in each bag. It can be used for life-threatening bleeding in patients with von Willebrand's disease but only when the proper preparations of concentrated vWF (Humate-P [CSL Behring GmbH, Marburg, Germany]) are not available.

Cryoprecipitate may be required to correct significant hypofibrinogenemia (<100 mg/dL). A typical adult dose of approximately 10 bags of cryoprecipitate raises the fibrinogen level by up to 1 g/L (60 to 100 mg/dL). In cases of severe bleeding after the use of a fibrinolytic agent such as tissue plasminogen activator, cryoprecipitate can be used to help control the bleeding. A consensus on dosing has not been reached, but many sources recommend between 10 and 12 bags.113

**Specific Factor Therapy**

A summation of the dosages and characteristics of each factor appears in Table 28.4.

**Factor VII**

Activated recombinant factor VII (rFVIIa) is a DNA product which is structurally similar to human serum factor VIIa. It has been approved by the FDA for the prevention and treatment of bleeding in patients with hemophilia A or B who have inhibitors to factors VIII and IX, as well as prevention and treatment of bleeding in patients with congenital factor VII deficiency. It works by binding to the surface of activated platelets, which then activate factor X by using the tissue factor pathway (formerly known as the extrinsic pathway). This obviates the need for either factor VIII or IX. Activated factor X then complexes with factor Va, which leads to thrombin burst and clot formation. Factor VII has a half-life of 2.7 hours. A thromboembolic rate of 1% to 2% has been reported.114 A large meta-analysis of 35 randomized trials showed that arterial clots were more common than venous clots and the risk appears to increase with age.115
Increasingly, activated factor VII is being used to control bleeding in patients who are not hemophiliacs.\textsuperscript{116} One multicenter study found that of 701 patients receiving factor VII, 92% were for off-label uses.\textsuperscript{117} However, a previous well-designed systematic review of the use of rFVIIa for the prevention and treatment of bleeding in patients without hemophilia denoted an increased arterial clot risk while failing to show a mortality benefit.\textsuperscript{118} Several randomized, controlled trials have investigated the use of rFVIIa in specific settings, including ICH, GI bleeding, and trauma. Dosing has varied markedly from 5 to 400 µg/kg. A significant consideration when contemplating the use of factor VII is cost. At an average cost per dose of $5000 (80 µg/kg), this can be a limiting factor, especially when some studies use protocols consisting of eight sequential doses.\textsuperscript{119}

**Hemophilia**

Although factor VII is being used increasingly in nonhemophilic patients, its original indication was for hemophiliacs in whom factor VIII inhibitors had developed and who were having acute bleeding events. Study dosages in these patients were generally higher than for off-label use (100 to 300 µg), but the drug was effective in controlling bleeding episodes, with an acceptably low rate of thromboembolic events.\textsuperscript{120,121} Its use as a more general hemostatic drug remains unproven, and use outside its current approved indications and clinical investigations should be avoided.

**Intracranial Hemorrhage**

ICH is a predictor of poor survivability and neurologic function in a patient who has undergone an acute stroke. The risk for hemorrhage expansion within the first 24 hours is between 20% and 40% in these patients. The risk of hematoma expansion is doubled in patients on vitamin K antagonists (warfarin).\textsuperscript{122} This would make it seem logical to attempt to rapidly reverse the anticoagulant effects of warfarin and this strategy is recommended in national and international guidelines. Many older studies acknowledged that reversal of elevated INR did not necessarily translate into improved survival or neurologic outcomes. One study by Parry-Jones and colleagues suggested that reversal of warfarin with PCC, FFP, or both demonstrated improvement in 30-day mortality compared to no reversal.\textsuperscript{123} However, the study authors acknowledge the presence of many confounding variables that may have affected the results. Interestingly, no single strategy to reverse warfarin has been conclusively shown to improve clinical outcome better than any other strategy.\textsuperscript{124}

In 2005, Mayer and associates\textsuperscript{124} published a double blind, placebo-controlled trial that evaluated the use of rFVIIa for acute ICH. They randomized 399 patients to placebo or to 40 µg/kg, 80 µg/kg, or 160 µg/kg of rFVIIa within 1 hour of a baseline computed tomography (CT) scan of the head. CT was repeated in 24 hours. The primary outcome measured was the percent change in volume of ICH from the initial to the repeat scan. The study showed a 29% increase in the volume of ICH in the placebo group versus 16%, 14%, and 11% in the treatment groups, respectively. Ninety-day outcomes were also evaluated and showed a 69% rate of death or severe disability in patients in the placebo group and 55%, 49%, and 54% in the rFVIIa groups, respectively. When looking at mortality alone, the rate was 29% in the placebo group and 18% in the rFVIIa groups combined. A subsequent study of 841 patients by the same investigators also showed a reduction in growth of the hematoma with rFVIIa, but in contrast to the initial study, no improvement in mortality or functional outcome occurred.\textsuperscript{125} Studies of traumatic ICH have yielded similar findings. In a retrospective study by Stein and coworkers\textsuperscript{126} of 63 patients with severe traumatic brain injury and coagulopathy at admission, 29 patients who received rFVIIa were compared with 34 patients who received FFP. Time to surgical intervention was less in the rFVIIa group; however, there was no difference between the groups at discharge with respect to neurologic outcome or mortality. In a prospective, dose escalation study of factor VII in patients with traumatic ICH, Narayan and colleagues\textsuperscript{127} found no differences in mortality rates or ICH volume in the placebo and rFVIIa groups. A

---

**TABLE 28.4 Managing Elevated International Normalized Ratios**

<table>
<thead>
<tr>
<th>CONDITION</th>
<th>ACTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>INR above the therapeutic range but &lt;5.0 AND No bleeding</td>
<td>Lower or omit the dose, monitor more frequently, and resume at a lower dose when the INR is therapeutic; if only minimally above the therapeutic range, dose reduction may or may not be required.</td>
</tr>
<tr>
<td>INR ≥5.0 but ≤10 AND No significant bleeding</td>
<td>Omit the next 1 or 2 doses, monitor more frequently, and resume at a lower dose when the INR is in a therapeutic range. Consider the use of vitamin K, 1–2.5 mg orally, particularly if at increased risk for bleeding. Note that INR correction with vitamin K in this situation may result in prolonged warfarin resistance for the next 3–5 days.</td>
</tr>
<tr>
<td>INR &gt;10 AND No significant bleeding</td>
<td>Hold warfarin therapy and give a higher dose of vitamin K (5–10 mg orally) with the expectation that the INR will be reduced substantially in 24–48 hrs. Monitor more frequently and use additional vitamin K if necessary. Resume therapy at a lower dose when the INR is therapeutic.</td>
</tr>
<tr>
<td>Serious or life-threatening bleeding at any elevation of INR</td>
<td>Hold warfarin therapy and give vitamin K (10 mg by slow intravenous infusion), AND 4-factor prothrombin complex concentrate or fresh frozen plasma.</td>
</tr>
</tbody>
</table>

INR, International normalized ratio.

reasonable conclusion to the use of rFVIIa in ICH is that although it may limit the size of the ICH, there does not appear to be any relevant effect on clinical outcomes and therefore it should not be considered for routine use in this setting.128

Trauma
Several investigators have examined the use of factor VII in the setting of trauma. A study by Boffard and coworkers129 in 2005 enrolled 301 ED patients with major trauma who required at least 8 units of PRBCs. The treatment group received three doses of rFVIIa immediately after the eighth unit of blood had been given. The doses were repeated at hours 1 and 3. The primary outcome measured was the total transfusion requirement and subgroup analysis was performed for blunt and penetrating trauma. For blunt trauma, there was a small decrease in the transfusion requirement (7.0 units in the treatment group versus 7.5 in the placebo group) and a decrease in the number of patients needing massive transfusion (14% of the treatment group versus 33% of the placebo group). In the penetrating trauma group the differences were not statistically significant. In the entire cohort, no differences were found in mortality at 48 hours or 30 days in comparison to placebo. Thromboembolic events were uncommon (4%) and did not differ between the treatment and placebo groups. Raoaikady and colleagues130 evaluated a group of 48 patients with traumatic pelvic fracture who were scheduled for surgical repair. Patients were randomized to a dose of 90 µg/kg of rFVIIa or placebo at the time of first incision. No significant difference was found in the primary outcome measure of transfusion requirement. Other groups have found that the administration of factor VII can favorably affect the subsequent need for other blood products, specifically PRBCs, cryoprecipitate, and platelets; however, mortality is not significantly affected.131,132 A 2008 Cochrane review of seven trials involving 1214 individuals (687 patients receiving rFVIIa) concluded that there is no advantage or disadvantage of rFVIIa over placebo in any of the studied outcomes.133 A more recent article by Hauser and colleagues demonstrated that fewer units of blood needed to be transfused in trauma patients who received rFVIIa; however, they similarly found that there was no decrease in mortality in these patients when compared to placebo.134 At the time of this writing, rFVIIa cannot be recommended as a general hemostatic agent in trauma patients.

Factor VIII Concentrate
Human Antihemophilic Factor
Factor VIII extracted from pooled human plasma produces a concentrated stable product with a shelf life of up to 2 years. Significantly more concentrated than cryoprecipitate and available for home use, factor VIII concentrate was a major breakthrough in the treatment of hemophilia. Unfortunately, the presence of viruses in the donor pool contributed to the high prevalence of hepatitis and HIV infection in hemophiliacs who used earlier versions of this product. Newer products (Alphanate [Grifols Biologicals, Los Angeles, CA], Hemofil [Baxter Healthcare Corporation, Deerfield, IL], Humate-P [CSL Behring GmbH, Marburg, Germany], Koate-DVI [Kedrion Biopharma Inc., Fort Lee, NJ], Monoclate-M [Baxter Healthcare Corp.], and Monoclate-P [CSL Behring LLC, Kankakee, IL]) are produced with one or more methods to reduce viral contamination. These methods have markedly reduced the risk for viral transmission, especially lipid-encapsulated viruses (HIV, HBV, HCV). To date there have been no reports of transmission of these viruses with the newer preparations.

Recombinant Antihemophilic Factor
Since the gene for production of factor VIII was discovered in 1984, research into recombinant genetics has aimed to provide a safer product that will theoretically be more readily available and less expensive to produce. Two recombinant DNA-derived factor VIII preparations (Recombinate [Shire Pharmaceuticals Lexington, MA], Kogenate [Bayer Pharma AG, Berlin, Germany]) were approved by the FDA in 1993. More recent introductions include Bioclate (Baxter Healthcare Corp.), Helixate, Helixate FS (CLS Behring) and Kogenate FS (Bayer Pharma). These genetically engineered products have hemostatic activity equivalent to that of plasma-derived factor VIII and minimal risk for viral contamination. Because some of these products are prepared from human albumin and other animal proteins, there is a potential for transmission of viruses. Products such as Helixate FS and Kogenate FS are prepared without human albumin, which should eliminate the possibility of viral contamination. Though costlier, they are a better choice for young patients with a newly diagnosed condition who have not already been exposed to hepatitis or HIV.135

Factor VIII Concentrate
Administration of 1 unit of factor VIII concentrate per kilogram of body weight should increase factor VIII activity by 2%. The dosage should be individualized according to the severity of bleeding, the known deficiency of factor VIII activity, and the presence of factor VIII antibodies. Factor VIII levels should be increased to 20% to 40% of normal levels for minor bleeding (e.g., small joint), 40% to 60% for moderate bleeding (e.g., large joint, neck, oral cavity), and 60% to 100% for life-threatening bleeding (e.g., intracranial, intrabdominal, pharyngeal).

Antibodies develop in up to 15% of factor VIII recipients. Administration of massive doses of factor VIII has proved to be beneficial in overwhelming the endogenous antibody response. In addition, immunosorbent techniques to remove the antibody have met with some success. The general use of immunosuppressants and plasmapheresis has also had limited success. In patients with inhibitors, rFVIIa and factor VIII inhibitor-bypassing activity (FEIBA) have proved to be very useful and effective hemostatic agents.

Long-Acting Factor VIII
A long-lasting antihemophilic factor has recently been approved for use in adults and children with hemophilia A. It is a fusion of factor VIII bound to the Fc region of human immunoglobulin G1 (IgG1), which binds to the neonatal Fc receptor. This means that it is not degraded as fast as regular factor VIII and lasts longer in circulation. The trade name for this new preparation is Eloctate (Biogen Idec, Inc, Cambridge, MA).

Factor VIII Inhibitor-Bypassing Activity
FEIBA is a product derived from pooled human plasma that contains factors II, IX, and X, each mainly nonactivated, and factor VII mainly in the activated form. It promotes coagulation by bypassing the need for factors VIII and IX. FEIBA is vapor heated to achieve greater than 10 logs of reduction in all target viruses and its safety profile is favorable.136 FEIBA is used to treat bleeding episodes in hemophiliac patients with antibodies to factor VIII and is generally efficacious in this
with liver and heart disease or those who are pregnant or at risk for thrombotic complications exists, especially in patients ing, nausea, vomiting, and an occasional allergic reaction. The synthetic analogue of the natural hormone arginine)-vasopressin (DDAVP) has been found to stimulate absorption of 1-Deamino-(8-arginine)-Vasopressin

A synthetic analogue of pituitary vasopressin, 1-deamino-(8-D-arginine)-vasopressin (DDAVP) has been found to stimulate the release of von Willebrand's factor from vascular endothelium. Although a synthetic analogue of the natural hormone vasopressin, DDAVP administration has no impact on hemodynamics. vWF carries factor VIII in the blood and leads to a rise in factor VIII levels in some mild hemophiliacs. Treatment of mild hemophiliacs with 0.3 mg/kg intravenously over a 15-minute period has been recommended when avoidance of the inherent risks of the factor VIII concentrate is desired. DDAVP has also been used in patients with mild von Willebrand's disease, congenital platelet dysfunction, acquired platelet dysfunction due to uremia, or intake of drugs such as acetylsalicylic acid, clopidogrel, and other antiplatelet agents. Responsiveness to DDAVP varies from patient to patient.

Factor IX Concentrate

Factor IX is prepared from pooled human plasma and is available as a lyophilized powder (AlphaNine SD [Grifols Inc., Los Angeles, CA] Mononine [CSL Behring]). Factor IX is also available in recombinant technology as BeneFix (Wyeth Pharmaceuticals Inc., Philadelphia, PA).

Historically, the use of factor IX concentrate has carried a very high risk for transmission of viral hepatitis. However, improved donor screening and new methods of viral reduction have substantially reduced the risk for transmission of viruses. As with factor VIII concentrate, the risk for transmission of HIV and hepatitis is very low with current human-derived products. Recombinant factor IX is not derived from human products and carries no risk for viral transmission.

Factor IX is indicated for the treatment of bleeding episodes in hemophilia B patients with a severe deficiency of factor IX. Administration of 1 unit/kg body weight will increase the factor IX concentration by approximately 1%. High levels of factor IX are not required to control bleeding. Levels should be increased to 15% to 25% of normal for mild to moderate bleeding and to 25% to 50% of normal for more serious bleeding or before major surgery.

Prothrombin Complex Concentrate

PCC is a plasma-derived concentrate of nonactivated clotting factors II, VII, IX, and X. These preparations undergo a process of viral inactivation to reduce the risk for viral transmission. PCCs are divided into two groups: three-factor preparations and four-factor preparations. This nomenclature can be confusing because both preparations contain four factors; however, the three-factor preparations have very low amounts of factor VII, whereas the four-factor preparations contain clinically relevant amounts. Some PCC products also contain protein C and protein S. These are added to balance the procoagulant effect of the concentrated clotting factors. The two most commonly available three-factor products are Prophine Nine SD (Grifols Inc.) and Bebulin VH (BDI Pharma, Inc., Columbia, SC). Four-factor preparations have been used in Europe for many years. The first four-factor PCC was approved in April 2013 by the FDA for use in the United States, specifically for the reversal of warfarin-related coagulopathy. The product is called Kcentra (CSL Behring) in the United States and Beriplex P/N (CSL Behring) in Europe. In addition to factors II, VII, IX and X, Kcentra contains proteins S and C and heparin.

In the ED, PCCs can play a very important role in the reversal of anticoagulation from warfarin (see hereafter). One concern with these preparations is a possible prothrombotic effect. A systematic review of 14 studies (460 patients) found only seven thrombotic complications: three strokes, two myocardial infarctions, and two deep venous thromboses. This translates to a 1.5% risk for clotting with PCC administration.

Two more recent reviews comparing four-factor PCCs to FFP in the reversal of warfarin-induced coagulopathy found similar rates of thrombotic events in both groups.

Warfarin-Induced Coagulopathy: Reversal Strategies

Elevated INRs may be encountered fortuitously or in patients with trauma or serious medical conditions. Guides to approaching and treating such patients are found in Table 28.4. Reflex reversal of an elevated INR should be avoided, especially in the absence of bleeding. Even minor bleeding can be tolerated in lieu of losing the beneficial effects of anticoagulation in selected patients (e.g., those with mechanical heart valves, left ventricular assist devices [LVADs], or active clots). In the presence of significant trauma or serious hemorrhage, however, any warfarin effect should be reversed.

Warfarin works as a vitamin K antagonist by inhibiting vitamin K epoxide reductase, decreasing available vitamin K, and thereby blocking synthesis of the active forms of the vitamin K–dependent coagulation factors II, VII, IX and X, along with proteins S and C. The factors that are made under the influence of warfarin are biologically inactive. Over a few days, the body's remaining stores of active factors are used up and replaced by the inactive forms creating a coagulopathy. Because protein S and C synthesis is relatively quickly blocked, the antithrombotic effect of these proteins is likewise curtailed, creating a transient hypercoagulable state while the warfarin is slowly developing its anticoagulation effects.

There are two parts to warfarin reversal: sustained reversal via the administration of vitamin K and “immediate” reversal using agents described in the following section.

Vitamin K, (Phytonadione)

The first step in the reversal of warfarin is to replete active vitamin K by administering vitamin K according to the recommended dosage and route guidelines outlined in Table 28.5. This provides a sustained reversal effect. Vitamin K can be given either orally or intravenously. Oral use is universally safe and should be used when possible. The IV preparation can be given orally to titrate small amounts, if needed. Although IV administration has been implicated in anaphylactoid-like systemic reactions, the incidence of these reactions is very small and should not preclude its use when the oral route is unavailable. Additionally, administration of the IV dose over at least 20 minutes may help to mitigate such reactions. Some institutions may elect to extend default administration times to 30 to 60 minutes as a further precautionary measure. The best treatment for a systemic reaction to intravenous administration of vitamin K is unknown, but general supportive measures for anaphylactoid reactions seem to work well. Because of erratic absorption patterns, subcutaneous administration of vitamin K is no longer recommended.

Tables are available in only 5-mg strength.
Reduction in the INR begins within 1 to 2 hours when vitamin K is given intravenously, with peak effects seen in 12 to 14 hours. Oral doses generally have an onset of effect in 6 to 10 hours, with a peak effect in 24 to 48 hours. Correction to within the normal range is generally achieved in these time frames if hepatic function is normal and a sufficiently large dose is given. Reversal of anticoagulation with vitamin K can be sustained for several days, making reanticoagulation with warfarin difficult. This is often referred to as warfarin resistance. This situation can happen with a patient on warfarin who has an artificial valve or an LVAD and has an acute GI bleed. Temporary reversal of anticoagulation may be necessary awaiting definitive correction of the GI bleeding source, followed by the rapid reintroduction of anticoagulation to protect the valve or LVAD. In these cases it may be desirable to withhold the vitamin K while attempting to achieve a short-term reversal with FFP or PCC.

The next step in the rapid correction of an elevated INR is to replace the missing clotting factors (II, VII, XI, and X). Traditionally, FFP has been used for this purpose. FFP is effective but requires a significant volume and takes time to thaw and prepare for administration. PCCs have been used in Europe and other parts of the world for many years to reverse undesirably high INR values. Studies have demonstrated the effectiveness of PCCs in normalizing elevated INRs within 15 minutes after administration without the complication of excessive volume or a delay in time to administration.

**Fresh Frozen Plasma**

FFP in doses of 10 to 15 mL/kg (with a minimum starting dose of 4 units) seems to effectively reverse coagulopathy, as measured by a decreasing INR. Issues related to volume expansion and the time required to obtain FFP have been discussed earlier. If PCCs are not readily available, FFP should be administered as quickly as possible in the coagulopathic, bleeding patient. Good judgment must be used when deciding to wait for the type and crossmatch, thawing, and administration of FFP in a critically sick, bleeding patient who requires transfer to another facility that can provide definitive treatment for the source of bleeding.

**Prothrombin Complex Concentrates**

Studies comparing the use of PCCs and FFP have failed to show significant benefits with regard to survival. The rapidity of administration, low volume required, and quick correction of the INR are compelling reasons to choose PCCs over FFP when managing an acutely bleeding patient. Dosing ranges for PCCs have varied. One study suggested a standard absolute dose for patients with an elevated INR (500 units for INR >5 and 1000 units for INR >5). This study used four-factor preparations only available outside the United States; the same authors found suboptimal reversal effects with three-factor preparations. Other studies have suggested using “unit/kg” dosing, with the amount varying according to the degree of INR elevation. In a recent article, Klein and colleagues suggested using a fixed dose of 1500 international units (IU) of four-factor PCC. In their study, the authors evaluated 39 patients and found a high rate of successful INR reversal with no thrombotic adverse events within 7 days of PCC administration.

**Recombinant Factor VII (NovoSeven)**

Another strategy to reverse warfarin-related coagulopathy is to bypass the need for the missing factors by creating a very large thrombin burst (e.g., by directly activating factor X). rFVIIa has been used off-label for this purpose. Although rapid normalization of the INR is achieved, sustained correction requires additional doses because of the short half-life of rFVIIa.

### TABLE 28.5 Summary and Dosage of Reversal Agents for Warfarin

<table>
<thead>
<tr>
<th>AGENT</th>
<th>DOSE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin K</td>
<td>1–10 mg PO or IV</td>
<td>SC delivery is no longer used. Reduciton in the INR begins within 1–2 hr when given IV with peak effects seen in 12–14 hr. Oral doses generally have onset of effect in 6–10 hr with a peak effect in 24–48 hr.</td>
</tr>
<tr>
<td>Prothrombin Complex Concentrate 3-Factor (Profilnine), 4-Factor (Kcentra)</td>
<td>Strategy 1: INR- and weight-based dosing INR 2–4: 25 IU/kg by IV push INR ≥4–6: 35 IU/kg by IV push INR &gt;6: 50 IU/kg by IV push Strategy 2: INR-based dosing INR &lt;5: 500 units; INR ≥5: 1000 units Strategy 3: Fixed dose 1500 IU</td>
<td>Multiple dosing strategies INR-based dosing is most effective with 3-factor preparations. Absolute dosing strategies should not be used with 3-factor PCCs. Any of the 3 strategies can be used with 4-factor PCCs. These authors’ institution uses strategy 1: weight- and INR-based dosing.</td>
</tr>
<tr>
<td>aPCC (FEIBA)</td>
<td>INR &lt;5: 500 units; INR &gt;5: 1000 units</td>
<td>May be more thrombogenic than nonactivated PCC and should be reserved for use when 4-factor PCC is not available</td>
</tr>
</tbody>
</table>

*Most of the dosing is off-label and is derived from various studies using these agents for unapproved conditions. Other dosing regimens may exist.

1FDA approved for the reversal of warfarin-related bleeding.

aPCC, Activated prothrombin complex concentrate; FEIBA, factor VIII inhibitor-bypassing activity; INR, international normalized ratio; IV, intravenous; PCC, prothrombin complex concentrate; PO, oral; SC, subcutaneous.
Although the INR can be corrected rapidly, some studies have called into question the effects of rFVIIa on actual clinical reversal of bleeding.\textsuperscript{10,15} Additionally, outcome studies have failed to demonstrate a significant mortality benefit in patients with ICH on warfarin who received rFVIIa.\textsuperscript{128} At present, there is not enough data to support the routine use of rFVIIa for reversal of warfarin-associated bleeding. One study found FEIBA to be more effective in reversing the INR, activated PCC carries the risk of being more thrombogenic than nonactivated PCCs. With the four-factor PCC agent Kcentra now approved by the FDA for reversal of warfarin-associated bleeding, FEIBA should only be considered when a four-factor PCC is not available. A comparison of FFP, PCC, and factor VII is provided in Table 28.4. Dosing of agents used to reverse warfarin can be found in Table 28.5. Table 28.6 lists advantages and disadvantages of these reversal agents.

### Table 28.6 Correction of the International Normalized Ratio in Warfarin-Anticoagulated Patients

<table>
<thead>
<tr>
<th>REVERSAL AGENTS</th>
<th>ADVANTAGES</th>
<th>DISADVANTAGES</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCC</td>
<td>1. Onset within 10 min</td>
<td>1. The Bebulin VH product (but not Profilnine) contains heparin—this would not be appropriate when heparin-induced thrombocytopenia is suspected.</td>
</tr>
<tr>
<td>United States: Profilnine SD, Bebulin VH (3-Factor PCC), Kcentra\textsuperscript{4} (4-Factor PCC)</td>
<td>2. Low risk for viral contamination</td>
<td>2. Kcentra is a 4-factor PCC, FDA approved for the reversal of warfarin anticoagulation. It contains factors II, VII, IX, X; proteins C and S; albumin; and heparin.</td>
</tr>
<tr>
<td></td>
<td>3. Effectively and immediately decreases INR</td>
<td>3. Clotting events have been reported.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4. Duration of 12–24 hr may be an advantage or disadvantage. The coadministration of vitamin K will prevent a rebound of anticoagulation.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5. There is no significant evidence that the use of PCCs is more effective than FFP or that it improves clinical outcomes.</td>
</tr>
<tr>
<td>Activated PCC (FEIBA)</td>
<td>1. Onset within 10 min</td>
<td>1. May be more thrombogenic than nonactivated PCCs</td>
</tr>
<tr>
<td></td>
<td>2. Effective and rapid lowering of an elevated INR</td>
<td>2. There is no significant evidence that the use of PCCs (activated or nonactivated) is more effective than FFP or that they improve clinical outcomes.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Clotting events have been reported, especially at doses &gt;1 mg.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4. Short duration of 6 hr</td>
</tr>
<tr>
<td>rFVIIa</td>
<td>1. A recombinant product does not carry risks associated with blood products.</td>
<td>1. New data suggest that although the INR may be reversed in warfarin anticoagulation, clinical bleeding is not significantly affected. It is NOT recommended for the reversal of warfarin coagulopathy.</td>
</tr>
<tr>
<td></td>
<td>2. Onset within 10 min</td>
<td>2. Some clotting events have been reported, especially at doses &gt;1 mg.</td>
</tr>
<tr>
<td></td>
<td>3. Storage at room temperature</td>
<td>3. Cost: approximately $1200 for a 1-mg dose: 90 µg/kg in a 70-kg patient, \approx $7560</td>
</tr>
<tr>
<td>FFP</td>
<td>Cost: approximately $65/unit</td>
<td>4. Short duration of 6 hr</td>
</tr>
<tr>
<td></td>
<td>15 mL/kg with 250 mL/unit in a 70-kg adult = $273. Clinician familiarity.</td>
<td>1. High volume of fluid may be undesirable for heart failure</td>
</tr>
<tr>
<td></td>
<td>Available everywhere</td>
<td>2. Slow onset</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3. Frozen—must be thawed over a 30-min period</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4. Risk for TRALI</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5. Its use is rapidly being replaced by the use of PCCs.</td>
</tr>
</tbody>
</table>

\textsuperscript{4}FDA approved for the reversal of warfarin-related bleeding. 
FDA, Food and Drug Administration; FEIBA, factor VIII inhibitor-bypassing activity; FFP, fresh frozen plasma; INR, international normalized ratio; PCC, prothrombin complex concentrate; rFVIIa, recombinant factor VIIa; TRALI, transfusion-related acute lung injury.

### Reversal of the Novel Anticoagulants (NOACS)

With the introduction of new anticoagulants, direct factor Xa inhibitors (rivaroxaban [Xarelto, Janssen Pharmaceuticals Inc., Titusville, NJ], apixaban [Eliquis, Bristol-Myers Squibb Company, Princeton, NJ], edoxaban [Savaysa, Daiichi Sankyo Inc., Parsippany, NJ], and direct factor II inhibitors (dabigatran [Pradaxa, Boehringer Ingelheim Pharmaceuticals Inc., Ridgefield, CT]), there are concerns about reversal during an acute bleeding episode. Because these inhibitors act at key points in the coagulation cascade, they may be difficult to reverse.

#### Direct Factor II Inhibitors (Dabigatran)

Although dabigatran is removed by dialysis, such intervention is rarely practical in an acutely bleeding patient. Until the fall of 2015, recommendations for dabigatran reversal were largely based on case reports, animal models, and human volunteer studies. At the time, no reversal agent had been approved...
specifically for direct factor II inhibitors. In cases of life-threatening bleeding in patients receiving direct factor II inhibitors, the off-label use of PCC, FEIBA, or activated factor VIIa (or any combination) were all tried. FEIBA seemed to reverse the effects of dabigatran in healthy volunteers, whereas nonactivated PCCs appeared to have little effect. This led the American Society of Hematology to recommend FEIBA as the first-line agent for life-threatening bleeding from dabigatran.

Idarucizumab (PraxBind [Boehringer Ingelheim Pharmaceuticals Inc.]), a monoclonal antibody fragment to dabigatran, was introduced in the fall of 2015. It has an affinity for dabigatran that is 350 times higher than that of thrombin. In a phase I trial of healthy volunteers, idarucizumab rapidly and effectively corrected abnormal coagulation test results without significant adverse events. In a 2015 uncontrolled trial of patients on dabigatran with a serious bleeding event or the need for an urgent procedure, idarucizumab reversed the effects of dabigatran in minutes in 88% to 98% of the patients studied. Importantly, the effectiveness of idarucizumab in this study was measured by reversal of abnormal clotting parameters, not by clinical cessation of bleeding. There is currently no evidence that the reversal of dabigatran with idarucizumab will lead to improved patient outcomes. Notwithstanding, in the absence of a specific and clearly safe alternative, patients on dabigatran with life-threatening bleeding should be considered potential candidates for idarucizumab treatment. Definitive recommendations for the use of idarucizumab await additional studies.

The fibrinogen level should be checked in patients with dabigatran-associated bleeding. If fibrinogen is less than 200 mg/dL, two pools of cryoprecipitate should be given.

Dosing for agents used to reverse bleeding in patients on NOACs is listed in Table 28.7.

### Table 28.7 Summary and Dosage of Reversal Agents for Anti–Factor II Agents

<table>
<thead>
<tr>
<th>AGENT</th>
<th>DOSE</th>
<th>NOTES</th>
</tr>
</thead>
<tbody>
<tr>
<td>FEIBA</td>
<td>25–100 units/kg (50 units/kg is used by many institutions for this indication)</td>
<td>May be more thrombogenic than nonactivated PCC</td>
</tr>
<tr>
<td>Idarucizumab</td>
<td>5 g, provided as 2 separate vials each containing 2.5 g/50 mL</td>
<td>The only FDA-approved antidote to dabigatran-related bleeding</td>
</tr>
<tr>
<td>Cryoprecipitate</td>
<td>2 pools</td>
<td>If fibrinogen is &lt;200 mg/dL, give 2 pools cryoprecipitate</td>
</tr>
</tbody>
</table>

*FDA, Food and Drug Administration; FEIBA, factor VIII inhibitor-bypassing activity; PCC, prothrombin complex concentrate.*

Direct Factor Xa inhibitors (Rivaroxaban, Apixaban, Edoxaban)

Direct factor Xa inhibitors also present a challenge to reversal in the setting of acute hemorrhage. A variety of strategies using rFVIIa, PCC, and FEIBA have been tried in the treatment of these patients. In an ex vivo study of healthy volunteers, rFVIIa reversed abnormal coagulation studies in patients who were given rivaroxaban. rFVIIa failed, however, to reverse active bleeding in a rivaroxaban animal model. At the present time, rFVIIa is not recommended as a reversal agent for direct factor Xa inhibitors.

In contrast, FEIBA was effective in reversing abnormal coagulation studies in rivaroxaban-treated healthy volunteers and seemed to be effective at reversing bleeding in two animal models. As a result, FEIBA is an acceptable agent to reverse the effects of direct Xa inhibitors. Four-factor PCCs also seem effective at reversing abnormal coagulation parameters in healthy humans taking these medications. Although the effectiveness is suggested in animal bleeding models, there have been no studies evaluating the effect of PCCs on clinical bleeding in humans receiving factor Xa inhibitors. According to the Hemostasis and Thrombosis Research Society, the use of four-factor PCCs seems to be a reasonable approach to reversing the effects of direct Xa inhibitors. The risks and benefits must be thoroughly weighed when choosing to reverse anticoagulation with agents that could cause thrombosis.

A 2015 study published in the New England Journal of Medicine (NEJM) looked at a novel reversal agent, andexanet alfa. This drug, which is scheduled for release in the fall of 2016, is capable of reversing the effect of all direct factor Xa inhibitors (rivaroxaban, apixaban, and edoxaban) and indirect factor Xa inhibitors (low-molecular-weight heparins and fondaparinux). Andexanet alfa acts as an inactive factor Xa decoy that binds both direct and indirect factor Xa inhibitors and makes them unavailable to inhibit native factor Xa. In humans, the drug has been shown to reverse markers of anticoagulation, including anti–factor Xa activity. The NEJM study had two different arms; each looked at andexanet alfa reversal of either apixaban or rivaroxaban compared to a placebo. Both arms of the study first looked at a bolus dose of andexanet alfa and then a bolus followed by a continuous 2-hour infusion. The end point was measurement of anti–factor Xa activity and thrombin generation. In the apixaban group (33 participants), anti–factor Xa activity was reduced by 94% in the andexanet alfa–treated group (n = 24) compared to 21% among those who received a placebo (n = 9). Thrombin generation was fully restored in 100% of the treatment group and 11% of the placebo group. In the rivaroxaban group (41 participants) a 92% reduction in anti-Xa activity was seen in the andexanet alfa–treated group (n = 27), compared to 18% in the placebo group (n = 14). Thrombin generation was 96% restored. In this small study, there were no serious adverse events and no cases of inappropriate thrombosis. Larger studies are currently underway to further evaluate the effectiveness and safety of andexanet alfa.

Although the theoretical benefit of a safe and specific reversal agent such as andexanet alfa seems promising, there are no data as yet to show that there will be any actual beneficial effects or improved outcomes in human patients.

Dosing for agents used to reverse bleeding in patients on anti–Xa agents can be found in Table 28.8.

### Collection and Storage of Blood Products

Whole blood is collected from donors into 500-mL plastic bags containing 63 mL of citrate-phosphate-dextrose with a
resultant hematocrit of 35% to 40%. Immediately after collection, sophisticated techniques permit separation of the whole blood into various components and fractions. Blood components such as FFP, PRBCs, granulocytes, and platelets are prepared from a single donor, separated, and transfused as single units. Minor blood fractions, including albumin, γ-globulin, cryoprecipitate, and fibrinogen, are often pooled from multiple donors. Within 24 hours, blood is essentially devoid of normally functioning platelets and some clotting factors, especially the labile factors V and VIII. Separation into individual components permits specialized storage and transfusion techniques designed to optimize the survival and availability of each component.

As is true of whole blood, PRBCs can be stored for up to 21 days, although newer preservatives such as ADSOL (adenine, dextrose, saline, mannitol, and water) may allow 49-day storage. Minor blood fractions, including albumin, γ-globulin, cryoprecipitate, and fibrinogen, are often pooled from multiple donors. Within 24 hours, blood is essentially devoid of normally functioning platelets and some clotting factors, especially the labile factors V and VIII. Separation into individual components permits specialized storage and transfusion techniques designed to optimize the survival and availability of each component.

The individual blood components are discussed in the following sections. A summary of the dosages and characteristics of each component is provided in Table 28.8. A “type and hold” or “type and screen” (no crossmatch) request alerts the blood bank to the possibility that a blood product will be required for the patient so that appropriate units can be acquired and kept on hand. A type and crossmatch procedure takes 45 minutes and restricts a unit of blood to a specific patient. In the ED, a crossmatch procedure should be considered for a blood product only if the adult patient: (1) manifests shock, (2) has symptoms of anemia (usually associated with a hemoglobin level <10 g/dL) in the ED, (3) has a documented loss of 1000 mL of blood, or (4) requires a blood-losing operation immediately (e.g., thoracotomy). A type and hold can be requested for all other situations in which a blood transfusion is considered possible during the patient’s care; a desirable ratio of units crossmatched to units transfused can thus be achieved. Hooker and colleagues found that the empirical trigger of prehospital hypotension (systolic blood pressure <100 mm Hg) was a useful discriminator for ordering early crossmatched blood.

The number of units requested for a crossmatch procedure is determined by the size of the patient, the response of the patient to the injury and subsequent emergency treatment, and the presence of ongoing blood loss (e.g., arterial or massive GI bleeding).

Red cell preparations for transfusion are not routinely tested for the presence of sickle hemoglobin. Donors with sickle trait are not excluded, and blood with sickle trait can be safely given to almost every patient because occlusion of blood flow caused by intravascular sickling would occur only in extreme conditions of acidity, hypoxia, or hypothermia, which are unlikely to be compatible with life. Nonetheless, when a transfusion is being performed in infants and patients with known sickle cell anemia, the blood bank should be alerted, and a “sickle preparation”

### ORDERING OF BLOOD

When it has been decided that a patient needs a transfusion and the patient’s condition is stable enough, ask the patient or relatives about any previous transfusion reactions and whether the patient abides by any religious prohibitions to transfusion.
should be requested for donor blood to avoid the infusion of blood with sickle trait into such patients. Patients receiving multiple transfusions for sickle cell disease are at increased risk for alloimmunization. Directed donation may be helpful to decrease this risk.165,166

**Blood Request Forms**

Proper identification of the patient and the intended unit of blood is critical. Transfusion of an incorrect unit is a potentially fatal error. Most transfusion reactions are attributable to clerical error, and the vast majority of such errors occur at the bedside (e.g., not in the blood bank).167 Just before administering the blood, the nurse or clinician (or both) must check the identity of the numbered labels. In addition, the blood bank laboratory slip must identify the patient by name and number and contain the identification number of the unit of blood.

Usual procedures require a separate blood bank request form for each unit of RBCs or whole blood ordered. A number of units of FFP, cryoprecipitate, and platelet concentrates may be ordered on one form with proper identification (depending on individual blood bank procedures).

**Blood Products for Jehovah’s Witnesses**

There are approximately 1.2 million Jehovah’s Witnesses in the United States. Based on the religious belief that the Bible prohibits blood or blood product transfusion (Acts 15:28–29), devout Jehovah’s Witnesses do not accept transfusions of whole blood, packed cells, white blood cells, platelets, plasma, or autologous blood.168 Since 1961, willing acceptance of a blood transfusion by an unrepentant member has been grounds for expulsion from the religion.169,170 An individual’s decision to abandon the teachings of the church and accept blood products can lead to congregational disciplinary actions, including disfellowshipping, a term for formal expulsion and shunning. Some members may permit the infusion of albumin, clotting factor solutions, plasma expanders, or intraoperative autotransfusion.171 Although no guidelines for the administration of blood products to Jehovah’s Witnesses are absolute, certain recommendations can be made. Even though a transfusion may be necessary to save a patient’s life and would otherwise be considered standard care, administration of blood or blood products in the face of refusal can legally be considered battery. In an awake and otherwise competent adult, courts have ruled that clinicians cannot be held liable if they comply with a patient’s directive and withhold lifesaving blood administration after refusal of transfusion when specific and detailed information of the consequences of such an omission in treatment are provided. The issue becomes clouded when patients are incompetent, unconscious (most Jehovah’s Witnesses carry cards informing medical personnel of their religious beliefs), or minors.

In the absence of specific directives to the contrary, it is prudent to administer blood products to patients who are unconscious, judged to be incompetent adults, or minors.172 Explicit documentation of the intent of the clinician to preserve life, coupled with an accurate description of discussion of the issue with the patient or the family and clarification of the patient’s mental capacity is mandatory. Furthermore, emergency legal assistance (e.g., court orders, appointment of a temporary guardian) should be sought immediately with rapid judicial resolution.173 Various clinical techniques to maximize oxygen delivery and minimize oxygen consumption should be used.

Examples include limited blood drawing, the use of high-dose erythropoietin and nutritional support with aggressive iron supplementation, hypothermia, volume expansion, sedation, oxygen, and the use of synthetic hemoglobin substitutes.174,175 Hyperbaric oxygen therapy can dissolve sufficient oxygen to sustain life in the absence of hemoglobin and may be another option when blood cannot be used.176–178 It can be used in a pulsed fashion (3 to 4 hr/day) to reduce the oxygen debt until hemoglobin can rise to adequate levels.

**ADMINISTRATION OF BLOOD PRODUCTS**

**IV Transfusions**

Do not open a unit of blood unless a free-flowing IV access line has been established in a large-bore vein. Use a 14- to 16-gauge IV catheter if possible, both to minimize hemolysis and to ensure rapid infusion of fluid for the treatment of hypovolemia or hypotension. When a large quantity of blood must be given rapidly, administer it by means of a high-flow infusion system if possible. The purpose of a large-bore infusion line is defeated if blood is piggybacked with an 18- to 20-gauge needle through a side port in the infusion tubing. For an elective transfusion, however, blood may be given through a smaller lumen. Combining hemodilution (250 mL of saline with 1 unit of PRBCs) and pressurization can safely increase the flow rate through 20- and 22-gauge catheters severalfold.179 No significant hemolysis occurs when small (21-, 23-, 25-, and 27-gauge) short needles are used for the transfusion of fresh blood or packed cells in infants and children and when the maximum rate of infusion is less than 100 mL/hr.180 For rapid infusion, however, connect the blood administration tubing directly to the infusion catheter. Monitor the infusion site for infiltration, infection, or local reactions.

If the patient already has a suitable IV line in place, flush the system with a solution of normal saline before administering the blood. Do not use other IV fluids because of the risk for hemolysis or aggregation (e.g., with 5% dextrose in water).181,182 Do not place any medications in the unit of blood or infusion line for the same reasons.

Errors associated with blood transfusions have serious consequences and frequently result in severe transfusion reactions and death. The Joint Commission on Accreditation of Healthcare Organizations has set forth guidelines in an attempt to reduce these errors, including the following:

- **Training personnel in procedures and recognition of reactions.**
- **Revising staffing models.**
- **Redesigning methods to identify patients, attain patient and blood verification, gain consent, and process multiple samples.** It was strongly suggested that room numbers be discontinued as a method of identifying patients and use of a unique identification band be considered for patients receiving blood.
- **Enhancing technical and computer support of the process.**
- **Discontinuing the use of refrigerators for multiple blood units.**

Many EDs have moved to a two-nurse mandate for checking patient and blood unit identities before administration. As the risk for infectious complications associated with transfusion medicine gradually declines, medical errors have supplanted them as the most serious cause of transfusion mishaps.183
Intraosseous Transfusions

Although IV administration is by far the most common route of blood and blood component therapy, IO administration appears to be safe and effective. Administration rates via an IO route are generally slower (on average 21 mL/min versus 35 mL/min by the IV route) but appear to be metabolically and hemodynamically equivalent. In animal models, no evidence of fat embolism and lung or kidney inflammation was noted after IO-PRBC infusion. Administration of IO liposome-encapsulated hemoglobin (one of the forms of HBOC therapy) is also effective and may have additional advantages over the IV route. Similarly, limited studies on the administration of other blood products via the IO route have been promising.

Filters

Infuse all blood and blood products through an appropriate filter, such as those supplied in-line in blood administration tubing sets. In the past, filtration was required merely to keep the IV line from becoming blocked by clots. Adverse consequences from infusing unfiltered blood products have since been recognized. Debris consisting of clots and aggregates of fibrin, white blood cells, platelets, and intertwined RBCs (ranging in size from 15 to 200 µm) accumulates progressively during the storage of blood. The usual filter, made of a single layer of plastic with multiple 170-µm pores, traps larger particles yet allows the rapid infusion of 2 to 3 units of blood before flow is obstructed. Purified components of blood plasma can be safely administered through a filter with pores as fine as 5 µm.

It has been suggested that microaggregates of debris that can pass through a 170-µm filter may in part contribute to the syndrome of “shock lung” seen after the transfusion of many blood units in patients suffering from severe trauma and hemorrhage. Some clinicians therefore recommend using a microaggregate blood infusion filter with a mesh pore size of 40 µm when multiple units of blood are administered to trauma victims, patients with compromised pulmonary function, and neonates. Microaggregate filters tend to become blocked, impede the rate of infusion more quickly, and are not commonly required in the emergency setting. A significant number of platelets are removed by microaggregate filters, and some clinicians advise against using these filters when platelet packs are infused. Others believe that although platelets are removed with the microaggregate filters, the trapped platelets can be removed with a saline flush without any significant loss.

Replace standard filters after 2 to 3 units of blood product have been administered. Change microaggregate filters after each unit.

Rate of Infusion

One unit of whole blood can be safely administered to a hypotensive patient at a rate of at least 20 mL/kg per hr. In the setting of hypovolemic shock and continued hemorrhage, there is no limit to the transfusion rate. Multiple units may be transfused simultaneously, even under pressure. Use of a rapid transfuser device can assist in the rapid administration and warming of blood (Fig. 28.3). In stable patients, administer 1 unit of whole blood (500 mL) over approximately a 2-hour period (3 to 4 mL/kg per hr). After this time RBCs begin to lose metabolic activity. In addition, the unit of blood, which is an excellent culture medium, is likely to become contaminated if bacteria and fungi are allowed to grow at room temperature. Give packed cells at approximately the same rate. Give plasma products more rapidly; in a patient without cardiovascular compromise, administer FFP at a rate faster than 15 to 20 min/unit because the coagulant activity begins to deteriorate rapidly after thawing. In patients with severe anemia and congestive heart failure, administer a rapidly acting diuretic, such as furosemide at the onset of transfusion to prevent circulatory overload.

If a transfusion of blood must be interrupted or delayed for any reason, return the remainder of the blood unit to the blood bank. Refrigerators in the ED or on the hospital unit should not be used to store blood products unless they are specifically equipped and authorized to do so.

For patients in hemorrhagic shock, administer blood through two large-bore catheters, at different sites if necessary. Usually, gravity provides a sufficient pressure gradient if the unit is raised above the patient to increase the rate of infusion when the clamps are wide open. Using a pressure pump makes the infusion quicker. Do not use a standard sphygmomanometer cuff wrapped around a unit of blood to create increased infusion pressure because the nonuniform application of pressure could burst the plastic bag containing the blood component.

If desired, dilute PRBCs with normal saline (0.9% without dextrose) before infusion simply by opening the clamps on the upper tubes of the Y-infusion set and leaving the lower (recipient end) clamps closed. Although it is generally agreed that LR solution should not be mixed with blood because of possible clot formation, multiple studies have challenged this belief.
and results have shown that small amounts of LR solution (up to 150 mL) are safe in the clinical setting.\textsuperscript{[190,191]} Furthermore, larger volumes of LR solution have not been found to produce microaggregate clots in laboratory settings when using AS-3–preserved PRBCs, a typical additive nutritive solution containing sodium chloride, sodium citrate, sodium phosphate, citric acid, dextrose, and adenine.\textsuperscript{[192]}

Diluting blood, while increasing its volume, will allow more rapid infusion by decreasing blood viscosity, which is dependent on hematocrit. Alternatively, add approximately 200 mL of normal saline directly to the bag of PRBCs to bring the hematocrit in the blood bag to approximately 45%.

Rewarming

Blood is stored at approximately 4°C to maintain cellular integrity and prevent the growth of microorganisms. Blood products usually passively warm to 10°C by the time that they are administered to the patient unless administered under pressure. The adverse effects of hypothermia on cardiac conduction and flow rates are evident when rapid administration of a large volume of blood is performed without prewarming.

Various mechanisms have been used to warm blood to 35°C to 37°C. An ideal blood warmer should allow liberal flow rates while preventing thermal hemolysis of blood cells. Commonly used devices include those with bath coils that allow a plastic tube to reside in a closely regulated warm water bath and dry heat devices that allow blood to circulate through flat, thin bags sandwiched between aluminum blocks that contain electric heating elements. Both devices have relatively low flow rates and suboptimal thermal clearance.\textsuperscript{[6]} Immersion of blood bags in warm water baths is safe but is considered imprecise and slow. Despite some evidence of their safety, the use of microwave heating devices is not recommended by the Association of Blood Banks because of concerns of hemolysis.\textsuperscript{[193]} Herron and colleagues\textsuperscript{[194]} advocated keeping PRBC temperature below 50°C because they noted significant hemolysis beginning at 51°C to 53°C.

Rapid admixture warming is a promising alternative technique.\textsuperscript{[195]} Mix the unit of whole blood with an equal amount of normal saline that has been preheated to 60°C to 70°C. Once mixed, administer the product to the patient; this gives a resultant delivery temperature of approximately 35°C. This technique combines dilution of blood product and warming into one step. Regardless of the rewarming technique used, warming refrigerated blood to body temperature decreases its viscosity twofold to threefold and avoids venous spasm, thus facilitating transfusion.

Monitoring

During the first part of the transfusion of any blood product, carefully monitor the patient for evidence of a transfusion reaction. Look for signs and symptoms such as hives, chills, diarrhea, fever, pruritus, flushing, abdominal or back pain, tightness in the chest or throat, and respiratory distress. A potentially life-threatening acute hemolytic transfusion reaction in a patient who has previously received transfusions may differ clinically from a minor allergic reaction only by its effects on the patient’s pulse and blood pressure. Treat an allergic reaction (hives, itching) to leukocytes or plasma proteins by administering an antihistamine (but not in the blood infusion line), and stop the transfusion.

Stop the transfusion immediately when the following signs are encountered: an increase in pulse rate, a decrease in blood pressure, respiratory symptoms, chest or abdominal discomfort, or a sensation of “impending doom.” Administer normal saline to maintain blood pressure and urine output. Send samples of urine and blood to the laboratory to verify the presence of free hemoglobin. In addition, send the blood bank a clotted sample of blood to reassess for the presence of an immune reaction. If the blood bank concludes that the reaction is a nonhemolytic allergic response, premedicate with antihistamines (diphenhydramine or hydroxyzine) and antipyretics before the next transfusion. Alternatively, use washed cells.

If a hemolytic transfusion reaction is suspected, treat the patient vigorously and promptly.\textsuperscript{[196]} Most morbidity and mortality are due to hypotension and shock, with subsequent cardiovascular instability, renal insufficiency, respiratory manifestations, or hemorrhagic complications of disseminated intravascular coagulation. Infuse crystalloid or vasoressors to treat the hypotension directly, if required. Determine the volume and rate of infusion by blood pressure response. Treat symptomatically with acetaminophen, a warming blanket, antihistamines, and inhaled or subcutaneous β-agonists for bronchospasm or subglottic edema.

If an acute hemolytic transfusion reaction occurs, alkalinize the urine with IV sodium bicarbonate to prevent the precipitation of free hemoglobin. Force diuresis with mannitol to maintain urine output at 50 to 100 mL/hr. The benefit of alkalization and diuresis in preventing acute renal failure is uncertain, although the use of these techniques is commonly advocated.\textsuperscript{[197]} After shock is controlled, assess hemostasis, respiratory function, renal function, and cardiac function to help direct therapy for complications; disseminated intravascular coagulation may require the administration of plasma, platelets, or fibrinogen, and acute tubular necrosis may complicate fluid management.

Delayed, or “late,” hemolytic transfusion reactions may occur days or even weeks after the transfusion of RBCs. Such reactions are characterized by falling hemoglobin levels, jaundice, hemoglobinemia, and indirect hyperbilirubinemia.\textsuperscript{[198]} This complication is usually self-limited and is not life-threatening. Therapy is symptomatic, but future attempts at crossmatching for transfusions may be difficult because of the presence of RBC antibodies. Individuals thus affected should wear identification tags or bracelets to alert medical personnel that previous transfusion reactions have occurred.

On completion of a transfusion, make an entry in the patient’s record to indicate the type and volume of the transfusion and the presence or absence of any reaction. The progress note, the transfusion record sheet, or the transfusion laboratory slip can be used for this purpose and should be signed and dated by the clinician, in accordance with hospital policies.

Emphasize to the patient and family how critically important any blood transfusion is to the patient’s care. Suggest that the family consider arranging for replacement donations of units of blood to afford future patients the luxury of an ample, available supply of blood products.

Conclusion

Some general guidelines and caveats regarding the transfusion of blood products in the setting of acute blood loss are outlined in Box 28.2.
The total volume of blood circulating in the body is approximately 7% of ideal body weight in adults and 8%-9% in children. A 70-kg adult, therefore, has a total blood volume of approximately 5 L.

**DEFINITION OF MASSIVE BLOOD LOSS**
Loss of total blood volume (10 units in a 70-kg adult) in 24 hr or loss of 50% of total blood volume over a 3-hr period.

**GENERALIZED GUIDE TO TRANSFUSION OF RBCS**
- Rarely transfuse if the hemoglobin level is >10 g/dL; usually transfuse if the level is <6 g/dL. In general, the aim is to keep the hemoglobin level at approximately 7–8 g/dL and the hematocrit approximately 30%–35% to sustain hemostasis and oxygen delivery.
- In critically ill patients, a restrictive transfusion policy of administering RBCs if the hemoglobin level drops below 7 g/dL and maintaining levels in the 7–9 g/dL range is at least as beneficial as more liberal criteria, except perhaps for patients with acute coronary ischemia.

**GENERAL GUIDE TO PLATELET TRANSFUSION**
Aim for a platelet count of 50,000/mL or higher in actively bleeding patients, but lower thresholds may be safe, especially in the absence of bleeding.

**FFP**
Fibrinogen levels may fall to a critical level (<1 g/dL) after 150% has been lost. Other labile clotting factors fall to 25% activity after approximately 200% blood loss. Transfusing 1 unit of FFP per every 5–6 units of RBCs has been traditional teaching, but decisions are best based on coagulation profiles. Preliminary evidence indicates that an FFP/RBC ratio of 1:1 may be more beneficial in patients with exsanguinating hemorrhage.

**CRYOPRECIPITATE**
If FFP does not raise fibrinogen levels to more than 1 g/dL, consider cryoprecipitate. Use in the ED is supported, but not mandated.

**PROTHROMBIN COMPLEX CONCENTRATES**
Profilnine SD and Bebulin VH are both 3-factor PCCs. They contain insignificant amounts of factor VII. Kcentra is a 4-factor PCC. FEIBA (factor VIII inhibitor-bypassing activity) contains factor VII, mostly in the activated form and factors II, IX, and X in the mostly nonactivated form.

**RECOMBINANT FACTOR VIIA**
No strict guidelines or proven benefits exist. May consider if:
1. No heparin/warfarin effect remains.
2. Surgical control of bleeding is not possible.
3. Bleeding continues despite adequate replacement of other coagulation factors (FFP, platelets, cryoprecipitate).
4. Acidosis is corrected.
   - Use in the ED is supported but not mandated, nor proven to be clinically effective.

ED, Emergency department; FFP, fresh frozen plasma; INR, international normalized ratio; RBCs, red blood cells.


**ACKNOWLEDGMENTS**
Thanks to the following emergency medicine pharmacists for their help in review of this chapter:
Erin Reichert, PharmD, BCPS, Specialty Practice Pharmacist, Emergency Medicine; Andrew M. North, PharmD, BCPS, MBA, Specialty Practice Pharmacist, Emergency Medicine.

REFERENCES ARE AVAILABLE AT www.expertconsult.com


Deleted in review.


Local and Topical Anesthesia
Douglas L. McGee

Local anesthetic agents are important tools used in the everyday practice of emergency medicine. This chapter describes the mechanism of action, the nuances of clinical use, and the adverse reactions to anesthetics that are commonly used in the emergency department (ED). Detailed technical guidance for the performance of topical and infiltrative local anesthesia is provided.

BACKGROUND

The first local anesthetic was cocaine, an alkaloid in the leaves of the *Erythroxylon coca* shrub from the Andes Mountains. Early Incan society used cocaine for invasive procedures, including cranial trephination. In 1884, Koller used topical cocaine in the eye and was credited with the introduction of local anesthesia into clinical practice. In the same year, Zenfel used a topical solution of alcohol and cocaine to anesthetize the eardrum, and Hall introduced the drug into dentistry. In 1885 Halsted demonstrated that cocaine blocked nerve transmission, thereby laying the foundation for nerve block anesthesia. The search for alternatives to cocaine led to synthesis of the benzoic acid ester derivatives and the amide anesthetics used today. It was not until the 1960s that detailed understanding of the physiochemical properties, mechanism of action, pharmacokinetics, and toxicity of these agents emerged.

PHARMACOLOGY AND PHYSIOLOGY

Chemical Structure and Physiochemical Properties

Most useful local anesthetic agents share a basic chemical structure:

Aromatic segment—Intermediate chain—Hydrophilic segment

Subtle variations in this basic structure determine each agent's main physiochemical properties: the negative log of dissociation constant (pK_a), the partition coefficient (a measurement of lipid solubility), and the degree of protein binding. Each of these properties determines the drug's potency, onset, and duration of action. However, physiochemical properties are not the sole determinant of clinical activity; other factors influence the drug's effect. The intermediate chain between the aromatic and the hydrophilic segments is either an amino-ester or an amino-amide; these chemical structures form the basis for the two main classifications of local anesthetics. Common ester-type agents include procaine, chloroprocaine, cocaine, and tetracaine. Common amide-type agents include articaine, lidocaine, mepivacaine, prilocaine, bupivacaine, and etidocaine. Different biochemical pathways metabolize each class. Esters are hydrolyzed by plasma pseudocholinesterase. Cocaine, an ester, is also partly metabolized by N-demethylation and nonenzymatic hydrolysis. Individuals with pseudocholinesterase deficiency may have a greater potential for cocaine toxicity if large doses are used, although this has not been an issue when cocaine is used clinically as an anesthetic. Amides are metabolized in the liver by enzymatic degradation. Local anesthetics are poorly soluble weak bases combined with hydrogen chloride to produce the salt of a weak acid. In solution, the salt exists both as uncharged molecules (nonionized) and as positively charged cations (ionized). The nonionized form is lipid soluble, which enables it to diffuse through tissues and across nerve membranes. The ratio of nonionized to ionized forms depends on the pH of the medium (vial solution or tissue milieu) and on the pK_a of the specific agent. The pK_a is the pH in which 50% of the solution is in the uncharged form and 50% is in the charged form. When the pH of the solution or tissue is less than the pK_a, more of the drug is ionized. When the pH increases, more of the drug is in the nonionized form. Because the nonionized form of drug can diffuse through tissues and nerves, manipulating the pH of the solution can alter a drug's diffusion properties.

Local anesthetics are available in single-dose vials or ampules and in multidose vials, with and without epinephrine. Most solutions have a pH higher than 5. Multidose vials contain methylparaben (MPB), an antibacterial preservative. Local anesthetics premixed with epinephrine also contain an antioxidant (sodium bisulfite or sodium metabisulfite) to prevent deactivation of the vasoconstrictor. These solutions must be adjusted to a more acidic pH, approximately 3.5 to 4.0, to maintain the stability of epinephrine and its antioxidant. These properties as they relate to the amide group are depicted in Table 29.1.
is closed, thereby preventing these ions from moving along their concentration gradient (out → in). Although potassium can leave the cell to follow its concentration gradient (in → out), the need to maintain electrical neutrality inside the cell prevents it from completely doing so. Potassium is in equilibrium between the concentration gradient and the electrochemical gradient, thus creating the negative resting potential.

### TABLE 29.1 pH and Additives of Amide Local Anesthetics

<table>
<thead>
<tr>
<th>SOLUTION CONTENT</th>
<th>pH (RANGE)</th>
<th>PRESERVATIVE (METHYLPARABEN)</th>
<th>ANTIOXIDANT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plain, single dose</td>
<td>4.5–6.5</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Plain, multidose</td>
<td>4.5–6.5</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Commercial epinephrine, single dose</td>
<td>3.5–4.0</td>
<td>−</td>
<td>+</td>
</tr>
<tr>
<td>Commercial epinephrine, multidose</td>
<td>3.5–4.0</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Prepared epinephrine, single dose</td>
<td>4.5–6.5</td>
<td>−</td>
<td>−</td>
</tr>
</tbody>
</table>

**Figure 29.1 Schwann Cell Sheath of Unmyelinated (Left) and Myelinated (Right) Nerve Fibers.** (Reproduced with permission from Wildsmith JAW: Peripheral nerve and local anesthetic drugs, Br J Anaesth 58:692, 1986.)

**Figure 29.2 Axon Membrane.** ECF, Extracellular fluid; ICF, intracellular fluid. (Reproduced with permission from Wildsmith JAW: Peripheral nerve and local anesthetic drugs, Br J Anaesth 58:692, 1986.)

**Figure 29.3 Cross-Section of a Peripheral Nerve.** (Reproduced with permission from Wildsmith JAW: Peripheral nerve and local anesthetic drugs, Br J Anaesth 58:692, 1986.)

### Nerve Structure and Impulse Transmission

#### Functional and Structural Components of a Peripheral Nerve

The functional nerve unit includes the nerve axon and its surrounding Schwann cell sheath. The Schwann cell (Fig. 29.1) may surround several unmyelinated axons or a single myelinated nerve fiber and form a myelin sheath. Junctions between sheaths along the axon, called nodes of Ranvier, contain sodium channels necessary for depolarization. As myelin sheath thickness increases from autonomic to sensory to motor fibers, the nodes of Ranvier are spaced farther apart. The most important structure affecting transmission of nerve impulses is the axon membrane (Fig. 29.2). The membrane consists of a double layer of phospholipids into which are embedded protein molecules that serve as channels containing pores for the movement of ions in and out of the cell. Most pores have a filter, or gate, that controls ion-specific movement and a sensory mechanism that opens or closes the gate. Bundles of nerve fibers (Fig. 29.3) are embedded in the endoneurium, which is made of collagen fibrils, and they are surrounded by a cellular layer, the perineurium. The perineurium functions as a diffusion barrier and maintains the composition of extracellular fluid around the nerve fibers. Surrounding the entire structure is the outer layer of a peripheral nerve, the epineurium, which is composed of areolar connective tissue.

#### The Nerve Impulse and Transmission

The inside of a nerve fiber, or axoplasm, is negative (−70 mV) at rest in comparison to the outside. This resting potential is the net result of the differences in ionic concentration on each side of the axonal membrane and the forces that tend to maintain that difference. Specifically, there is a surplus of sodium extracellularly and potassium intracellularly. The sodium channel
The sodium channel opens when a nerve is stimulated. Sodium ions enter slowly at first until a critical threshold is reached. They then enter the cell rapidly, along the electrochemical and concentration gradients, and cause depolarization. The influx of sodium is halted when the membrane potential reaches +20 mV, but potassium continues to move out of the cell and repolarizes it until the resting potential is reached. When the excitation process has been completed and the nerve cell is electrically quiet, the relative excess of sodium inside the cell and potassium outside the cell is readjusted by the adenosine triphosphate–dependent sodium-potassium pump.

Depolarization of a portion of the nerve causes a current to flow along the adjacent nerve fiber. This current makes the membrane potential less negative and actuates the sensor to open the next sodium channel. The action potential cycle is repeated, thereby propagating the impulse. Nerve conduction is essentially unidirectional because the sodium channel is not only closed but inactivated as well, and delayed closure of specific potassium channels prevents the critical threshold from being reached in the segment just depolarized. Impulses travel continuously down the axon in unmyelinated nerve fibers. In myelinated fibers, current flows from node to node and depolarizes intervening segments at once. This saltatory conduction causes a faster rate of impulse transmission in myelinated fibers.

**Mechanism of Action**
How local anesthetic agents block nerve conduction depends on the active form of the agent and the specific physiologic and cellular activity.

**The Active Form**
Anesthetic solutions contain uncharged and charged forms. The concentration of the uncharged form increases in more alkaline milieus. Only this uncharged lipid-soluble form can cross tissue and membrane barriers. Once the uncharged drug is through a barrier, it reequilibrates into uncharged and charged forms in a proportion dependent on the prevailing pH. Because local anesthetics are more effective in alkaline solutions, it was originally thought that the uncharged form was responsible for conduction blockade. Alkaline solutions are currently believed to be more effective because of increased penetration through tissue barriers, but it is the cationic charged form that is responsible for the actual neuronal blockade.

**The Physiologic and Cellular Basis for Neuronal Blockade**
Prevention of sodium influx across the nerve membrane forms the physiologic basis for conduction blockade. Local anesthetics slow sodium influx, thereby decreasing the rate of rise and amplitude of depolarization. If sufficient anesthetic is present and the firing threshold is not reached, the action potential is not formed. With no action potential, no impulse is transmitted and conduction is blocked, which results in local anesthesia.

How anesthetic agents prevent sodium influx is still not completely understood. It is believed that the cationic charged form blocks the action potential from inside the membrane; the agent enters the sodium channel from the axoplasmic side and binds to a receptor.\(^4,5\) This “specific receptor” theory is well accepted and is considered the predominant mechanism in preventing influx of sodium. However, this theory cannot account for the action of benzocaine and other neutral com-

**Table 29.2 Activity Profile With a Primary Physiochemical Determinant**

<table>
<thead>
<tr>
<th>AGENT</th>
<th>ONSET: pK(_a)</th>
<th>POTENCY: LIPID SOLUBILITY</th>
<th>DURATION: PROTEIN BINDING</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetracaine</td>
<td>Slow</td>
<td>8</td>
<td>Long</td>
</tr>
<tr>
<td>Procaine</td>
<td>Slow</td>
<td>1</td>
<td>Short</td>
</tr>
<tr>
<td>Chloroprocaine</td>
<td>Fast</td>
<td>1</td>
<td>Short</td>
</tr>
<tr>
<td>Lidocaine</td>
<td>Fast</td>
<td>2</td>
<td>Moderate</td>
</tr>
<tr>
<td>Mepivacaine</td>
<td>Fast</td>
<td>2</td>
<td>Moderate</td>
</tr>
<tr>
<td>Prilocaine</td>
<td>Fast</td>
<td>2</td>
<td>Moderate</td>
</tr>
<tr>
<td>Bupivacaine</td>
<td>Moderate</td>
<td>8</td>
<td>Long</td>
</tr>
<tr>
<td>Etidocaine</td>
<td>Fast</td>
<td>4–6</td>
<td>Long</td>
</tr>
</tbody>
</table>
Adding sodium bicarbonate to raise the pH of the anesthetic solution (a technique that decreases pain on injection) yields a higher concentration of the uncharged lipid-soluble form and decreases onset time. Increasing the total dose by using a higher concentration of the same volume or a greater volume of the same concentration also shortens onset time. For most procedures performed in the ED, the time of onset of most agents is short enough that manipulation to achieve shorter times is unnecessary.

**Potency**

The lipid solubility of an anesthetic is a primary physicochemical factor determining potency. The drug’s partition coefficient, not the concentration of the lipid-soluble form determined by the pK_a or pH, confers its lipid solubility. Because the nerve membrane is lipid, lipophilic anesthetics pass more easily into the cell and few molecules are needed to block conduction. The degree of protein binding of an anesthetic primarily determines its duration of action. Agents that bind more tightly to the protein receptor remain in the sodium channel longer (see Tables 29.2 and 29.3). Like potency, the duration of action is reduced by the vasodilation produced by local anesthetics. Prilocaine, which is less protein bound than lidocaine, has a longer duration of action because of its lesser degree of vasodilation. The duration of action also varies with the mode of administration. It is shorter when agents are applied topically.

The duration of action may be prolonged by several methods. Increasing the dose, usually by increasing the concentration, prolongs the duration to limits imposed by toxic effects. Raising the pH of the anesthetic solution has also been shown to prolong duration.\(^6\)\(^7\) The most practical way to increase duration is to use solutions that contain epinephrine.\(^8\) Epinephrine causes vasoconstriction, decreases systemic absorption, and allows more drug to reach the nerve. The effect of epinephrine varies according to the agent. Anesthetics that intrinsically produce more vasodilation (e.g., procaine, lidocaine, mepivacaine) benefit more from epinephrine’s vasoconstrictive action. The long-acting, highly lipid-soluble agents (e.g., bupivacaine, etidocaine) are less affected because they are substantially taken up by extradural fat and released slowly. In fact, lidocaine with epinephrine may be effective for as long as bupivacaine without epinephrine. Generally, most ED procedures can be accomplished quickly before the anesthesia wears off regardless of which drug is selected. Choose agents with a long duration of action when the procedure is lengthy or if postoperative analgesia is desired.

**TOPICAL ANESTHESIA**

Local anesthetic agents may be applied topically to mucous membranes, intact skin, and lacerations. There are sufficient differences among these sites to merit a separate discussion of each one. Topical anesthesia of the eye is discussed in Chapter 62.

**Mucous Membranes**

**Agents and Uses**

Effective anesthesia of the intact mucous membranes (not intact skin) of the nose, mouth, throat, tracheobronchial tree,
esophagus, and genitourinary tract may be provided by several anesthetics (Box 29.1). Tetracaine, lidocaine, and cocaine are the most effective commonly used agents (Table 29.4 and Box 29.1). Benzocaine (14% to 20%) is commonly used for intraoral or pharyngeal anesthesia (Fig. 29.4). The anesthesia produced is superficial and does not relieve pain that originates from submucosal structures. The onset of action may be slow, which limits its usefulness in urgent situations (such as passing a nasogastric tube). Agents applied topically can be absorbed systemically, and concentrated topical agents can cause toxicity.

Tetracaine solution is an effective and potent topical agent with a relatively long duration of action. It is used in concentrations from 0.25% to 1% with a recommended maximum adult dose of 50 mg. In overdose, it has the disadvantage of severe cardiovascular toxicity without any preceding central nervous system (CNS) stimulatory phase.

Lidocaine is also an effective topical agent that is marketed in a variety of forms (solutions, jellies, and ointments) and concentrations (2% to 10%). The 10% form is most effective, and minimal topical anesthesia is achieved with less potent concentrations. Lidocaine is commonly used as the 2% viscous solution prescribed for inflamed or irritated mucous membranes of the mouth and pharynx. Patient misuse of viscous lidocaine, by repeated self-administration, can lead to serious toxicity. Topical lidocaine provides an adequate duration for most procedures, with the maximum safe dose being 250 to 300 mg.

Cocaine is an effective, but potentially toxic topical agent that is applied to the mucous membranes of the upper respiratory tract. Although it is an ester, hepatic metabolism occurs, as does hydrolysis by plasma pseudocholinesterase. Absorption is enhanced in the presence of inflammation. Cocaine is the only anesthetic that produces vasoconstriction at clinically useful concentrations, hence its popularity for treating epistaxis.

### BOX 29.1 Common Local and Topical Anesthetics Used in the ED

- Benzocaine spray will produce transient anesthesia of the mucous membranes. Rarely, it can precipitate methemoglobinemia in standard doses. Anbesol (Pfizer Inc., New York, NY) is a popular over-the-counter benzocaine anesthetic for dental problems, such as teething. Topical anesthesia of mucous membranes can be obtained by applying a gel mixture of 10% lidocaine, 10% prilocaine and 4% tetracaine (Profound, Steven’s Pharmacy Costa Mesa, CA, www.stevensrx.com) and is preferred by editors.
- EMLA cream (lidocaine and prilocaine) will produce anesthesia of intact skin, but it must be in place for approximately 60 minutes to provide significant benefit.
- EMLA-Max, now known as LMX4 (Ferndale Laboratories Inc, Ferndale, MI), is another topical lidocaine preparation with a more rapid onset of action.
- “Magic mouthwash” contains equal parts of diphenhydramine elixir, Maalox (Novartis, East Hanover, NJ), and 2% viscous lidocaine. Each 5-mL teaspoon contains less than 50 mg of lidocaine. It is swished, held in the mouth for 1 to 2 minutes, and expectorated.
- Lidocaine (2%) may be used intraorally, but repeated use may produce systemic toxicity, especially in children. Each 5-mL teaspoon contains 100 mg of lidocaine. It should not be swallowed but, instead, expectorated after holding it in the mouth for a few minutes. Viscous lidocaine is not useful for acute pharyngitis. Systemic narcotics are preferred if the pain is severe.

**ED**, Emergency department; **EMLA**, eutectic mixture of local anesthetics.

### TABLE 29.4 Practical Agents for ED Use: Mucosal Application

<table>
<thead>
<tr>
<th>AGENT</th>
<th>USUAL CONCENTRATION (%)</th>
<th>Maximum Dosage*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Adult (mg)</td>
</tr>
<tr>
<td>Tetracaine</td>
<td>0.5</td>
<td>50</td>
</tr>
<tr>
<td>Lidocaine</td>
<td>2–10</td>
<td>250–300</td>
</tr>
<tr>
<td>Cocaine</td>
<td>4</td>
<td>200</td>
</tr>
</tbody>
</table>

*These are conservative figures; see text for explanations.

The lower dosage should be used for a maximum safe dose when feasible.

The 10% cocaine solution is best avoided because of minimal additional clinical benefit and the potential for coronary vasoconstriction in patients with coronary artery disease.

**ED**, Emergency department.
This major advantage is offset by its susceptibility to abuse and toxic potential. The toxic effects are due to direct stimulation of the CNS and blockade of norepinephrine reuptake in the peripheral nervous system. Cocaine should not be administered to patients who are sensitive to exogenous catecholamines or who are taking monoamine oxidase (MAO) inhibitor antidepressants. Clinical manifestations of toxicity include CNS excitement, seizures, and hyperthermia. Central and peripheral effects of hypertension, tachycardia, and ventricular arrhythmias may also be seen. Acute myocardial infarction has been reported after topical application. Cocaine is commonly used as a 4% solution with a maximum safe dose of 200 mg (2 to 3 mg/kg). A 10% solution is available, but this concentration adds little to the topical effect while enhancing the potential for toxicity. Coronary vasoconstriction may occur with doses as low as 2 mg/kg applied to the nasal mucosa. Although the clinical effect of this vasoconstriction is usually benign and without electrocardiographic changes, topical cocaine should be used cautiously in patients with known or suspected coronary artery disease. Dyclonine offers advantages over other topical anesthetic agents. It is a ketone derivative without an ester or amide linkage and may be used in patients who are allergic to the common anesthetics. Extensive experience with the topical preparation has shown it to be effective and safe. Dyclonine is marketed in 0.5% and 1% solutions, often in sore throat preparations, with a maximum adult recommended dose of 300 mg.

Benzocaine is an ester that is marketed in its neutral form in 14% to 20% preparations (Cetacaine [Cetylite Industries, Inc., Pennsauken, NJ], Americaine [Celltech Pharmaceuticals, Rochester NY], Hurricane) (see Fig. 29.4). Its low water solubility prevents significant penetration of the mucous membranes, thus reducing systemic toxicity if applied to intact mucosa. However, it is not a potent anesthetic and has a brief duration of action. It is more allergenic than other topical agents. Benzocaine is usually dispensed in an admixture with other therapeutic ingredients and is clinically effective only at relatively high (>14%) concentrations. Benzocaine is available as a nonprescription gel and liquid (6.3% to 20% Anbesol [Pfizer Inc., New York, NY], for example) and is used for a variety of maladies, including ear pain, mouth pain, and teething. It is commonly used by dentists to produce mucosal anesthesia before intraoral nerve blocks (see Chapter 30). Adriani and Zepernick recommended this agent for lubricating catheters, airways, endotracheal tubes, and laryngoscopes and reported only one adverse reaction (methemoglobinemia) in their experience with approximately 150,000 patients. Methemoglobinemia occurs rarely after mucosal absorption of benzocaine used repeatedly for teething infants and after standard doses of benzocaine sprays used in endoscopic procedures.

An excellent topical preparation is a combination of lidocaine, prilocaine, and tetracaine, especially useful for dental mucosa anesthesia. Topical gel mixtures of 2.5% lidocaine and 2.5% prilocaine (eutectic mixture of local anesthetics [EMLA]) are commonly used on intact skin but have also been used on mucous membranes. EMLA is more effective than 20% benzocaine when applied to the oral mucosa before needle injection for dental anesthesia. One study demonstrated that pain was reduced more quickly with EMLA than with benzocaine when applied to the buccal mucosa.

As with infiltrated anesthesia, toxic reactions to topically applied anesthetics correlate with the peak blood levels achieved and not necessarily with the dose administered. Systemic absorption of a topical agent is more rapid, with a higher level being achieved than with the same dose given by infiltration. The total dose of a topical anesthetic should be considerably less than that used for infiltration at a given site. Fractionating the total dose into three portions administered over a period of several minutes effectively reduces peak blood levels. Inadvertent suppression of the gag reflex, combined with difficulty swallowing, may lead to aspiration, an important potential adverse reaction to topical anesthesia of the nose, mouth, and pharynx. Infections from drug solutions in multidose vials for topical anesthesia of the larynx and trachea have not been substantiated.

**Technique and Precautions**

A commonly used “magic mouthwash” for the topical treatment of painful gingivostomatitis in children is often prescribed by emergency clinicians and pediatricians. There has been little scientific study of the preparation and it is not available commercially, but it has been used safely for decades. It consists of equal parts viscous lidocaine (2%), Maalox as a binder, and diphenhydramine elixir. Corticosteroids or nystatin is often added when the mixture is used to treat chemotherapy-induced mucositis. However, a 2004 review found no evidence that magic mouthwash is effective in treating this condition. The creamy mixture is swished around the mouth and expectorated, or painted on specific lesions with a cotton swab. Packing an area with a cotton ball soaked with this mixture is another option. Repeated doses or swallowing of the elixir can produce systemic toxicity, so careful instruction should be given to limit use of the solution to every few hours. This combination would be theoretically less toxic than simply using topical lidocaine.

Emergency clinicians often prescribe 2% viscous lidocaine (20 mg/mL) for patients with pharyngitis, stomatitis, dental pain, or other inflammatory or irritative lesions in the oropharynx. Although this intervention is widely used and generally safe, the common misconception that topical anesthesia is totally innocuous may result in poor patient instructions and serious consequences. Topical lidocaine is helpful for painful mouth lesions, but is of little practical value for acute pharyngitis, for which systemic analgesics are usually a better option. Seizures and death from topical lidocaine have been reported when excessive repeated doses have been administered. Toxic blood levels may occur because the anesthetic effect of viscous lidocaine lasts for only 30 to 60 minutes and patients with recurrent pain may either ignore or be ignorant of the safe dosing interval of 3 hours and medicate themselves more frequently. Patients tend to increase each dose to obtain greater relief, and inflammation may increase systemic absorption. In addition, painful oral lesions may last for several days.

Children are at higher risk for the rare toxicity of oral lidocaine. When compared with adults, children may exhibit increased lidocaine absorption, decreased clearance, and a longer half-life. Continued medication use allows lidocaine and its major metabolites monoethylglycinexylidide (MEGX) and glycinexylidide (GX) to accumulate. Both MEGX and GX are produced from the hepatic metabolism of lidocaine and are excreted in urine. They possess anesthetic and arrhythmic activity and have the potential for CNS toxicity. Although these metabolites are less potent than lidocaine, their elimination half-lives are considerably longer. Several investigators regard MEGX and GX to be the cause of CNS toxicity with prolonged topical use of lidocaine. The length of time...
that viscous lidocaine is retained in the mouth and whether the excess is expectorated or swallowed also affect the blood level produced. Expectorating the medication after swishing it in the mouth produces much lower blood levels than when it is swallowed. It seems logical that the most hazardous mode of administration would be to retain the solution in the mouth “until absorbed.”

Clearly explain the proper way to use viscous lidocaine and inform patients not to dose themselves ad libitum. Note that a 2% solution contains 20 mg/mL, or 100 mg per standard teaspoon (5 mL). The recommended maximum adult dose is 300 mg (15 mL of a 2% solution) no more frequently than every 3 hours. When possible, instruct the patient to decrease the dose by using direct cotton swab application. When gargled or swished in the mouth, limit application time to 1 to 2 minutes, and instruct the patient to expectorate excess solution. Limit use to 2 or 3 days, especially if swallowing the solution is necessary to obtain relief. Prescribe lower doses for patients at risk for decreased clearance (see later section on Systemic Toxic Reactions). Doses for children are prescribed at 3 mg/kg. Because infants cannot expectorate well, do not use viscous lidocaine for minor oral irritation and teething. Recommend that no food be eaten for 1 hour after application because anesthesia of the oropharynx can interfere with swallowing and cause aspiration. Special note should be made of the over-the-counter availability of benzocaine, commonly used for toothaches and teething. A gel or liquid (Anbesol, Pfizer) is available in 6.3% to 20% formulations. When used repeatedly in the oral cavity on irritated tissue, systemic toxicity, including methemoglobinemia, may occur.

Lidocaine 4% solution can be atomized with a standard nebulizer device commonly used for delivering asthma medications and inhaled by the patient before insertion of a nasogastric tube. This method effectively anesthetizes the nasopharyngeal and oropharyngeal tissues, thereby easing the pain of tube insertion. Lidocaine (4%) and cocaine (4%) atomized by a single-use mucosal atomizer, compared with lidocaine gel (4%), demonstrated equal reductions in nasal pain scores during the insertion of a nasogastric tube in healthy study participants, although they indicated that overall discomfort was less with lidocaine gel injected into the nose.

Intact Skin

Agents and Uses

The stratum corneum provides a cutaneous barrier that prevents the commonly marketed aqueous solutions (acid salts) from producing anesthesia, but saturated solutions of the bases of local anesthetics are effective on intact skin. When applied topically to abraded skin, most anesthetic agents produce peak blood levels similar to those resulting from infiltration in 6 to 10 minutes.

Lidocaine Cream and Patch

In 1974, Lubens and coworkers used 30% lidocaine cream, saturated on a gauze pad and adherent to an elastic patch, for a myriad of procedures including minor operative procedures (e.g., excision of lesions, incision and drainage of abscesses), lumbar puncture, venipuncture, and allergy testing. Today, lidocaine is available in a 5% patch, and remains one of the most commonly used topical compounds. Recent literature regarding its effectiveness, however, has been mixed. In a study by Cheng, a lidocaine patch applied at the site of rib fractures in patients taking oral pain medication decreased pain scores when compared to a placebo patch. In contrast, a meta-analysis of 251 patients in 5 trials compared lidocaine patches to placebo patches for acute or postoperative pain, but failed to conclusively demonstrate that a lidocaine patch decreases pain intensity, reduces opioid consumption, or reduces hospital length of stay.

EMLA Cream, ELA-Max, and Tetracaine Base Patch

Various topical anesthetics have been suggested to decrease the pain of venipuncture or injections and to provide topical anesthesia for painful skin abrasions and lesions. These agents have been studied extensively and are safe, but they are not practical in many ED settings because of their slow onset of action and inadequate efficacy. However, their activity profiles make them more applicable than 30% lidocaine cream to emergency medicine. Tetracaine base is available as a solution, a gel, and a patch preparation. It is effective in crossing the lipid-rich barrier of the stratum corneum because it is highly lipophilic. EMLA was approved in the United States in 1992. It contains 2.5% lidocaine and 2.5% prilocaine in a unique oil-and-water emulsion, yielding 5% EMLA. The mild lipophilic and hydrophilic properties of the component drugs are greatly increased when mixed together, thereby allowing absorption through intact skin. ELA-Max and ELA-Max5, now known as LMX 4 and LMX 5 respectively (Ferndale Laboratories Inc, Ferndale, MI), are topical lidocaine anesthetic creams with a more rapid onset of action than EMLA cream. ELA-Max is a 4% concentration, and 5% ELA-Max is marketed as an anorectal cream that may benefit patients undergoing painful rectal procedures. Neither product has prilocaine, as is found in EMLA cream, and neither has US Food and Drug Administration approval for pain relief before painful injections or intravenous (IV) insertion, but both have such potential.

Tetracaine base seems to offer the advantage of being able to achieve effective anesthesia with a shorter application time and a longer duration. For tetracaine and EMLA preparations, onset, depth of anesthesia, duration, and blood levels vary directly with application time, use on thinner or inflamed skin, and larger doses. These preparations exhibit a reservoir effect. The drug is deposited in the stratum corneum and continues to diffuse along its concentration gradient, even after it is removed from the skin.

Tetracaine base, lidocaine cream, and EMLA can be useful in the ED for providing anesthesia for many procedures: venous cannulation, venipuncture, or any needle insertion, including preinfiltration anesthesia and lumbar puncture; a variety of minor surgical procedures; and anesthetizing the tympanic membrane and external auditory canal. EMLA has also been used effectively for débridement of ulcers. EMLA cream applied to wound edges prior to local wound infiltration with 1% lidocaine decreased the pain of laceration repair compared with infiltrative anesthesia alone.

EMLA and 5% lidocaine cream are equally effective in reducing the pain of IV insertion. Luhmann and colleagues demonstrated that 5% lidocaine cream applied for 30 minutes under an occlusive dressing was as effective as infiltrated buffered lidocaine before IV catheter insertion in children. Obviously, infiltrative administration of buffered lidocaine requires skin puncture, but its onset of anesthesia is almost immediate. When time is not an issue, topical creams may be an acceptable alternative to infiltrated anesthesia or no anesthesia at all. Approximately a 60-minute interval after application is required...
for these preparations to provide optimal topical analgesia to the intact skin for such procedures as venipuncture. Early cutaneous placement of these agents (e.g., over common IV sites while the patient is being triaged) is important for practical ED use.

Ethyl Chloride and Trichloromonofluoromethane and Dichlorodifluoromethane (Fluori-Methane) Sprays
These topical agents are often used for limited skin incisions (e.g., drainage of small abscesses), trigger point injections, joint aspiration, or injection of bursitis or tendinitis. These agents evaporate quickly from the skin and cool it to the point of freezing. Anesthesia is effective and immediate, but drawbacks include its short duration (only up to 1 minute), potential pain on thawing, and possibly lowered resistance to infection and delayed healing. Highly volatile ethyl chloride spray is flammable. Ethyl chloride has been studied in children to reduce the pain of venipuncture, but the results are mixed. Given their short duration of action and the time needed to perform pediatric venipuncture, these preparations have limited use for this purpose.

Technique
Lidocaine Cream
This 30% cream is saturated on a gauze pad that is adherent to an elastic patch and placed over the area to be injected or incised. The high concentration of anesthetic and an occlusive patch are needed to achieve effective skin penetration. The duration of action varies with the application time. A 45-minute application time is needed for most procedures. To achieve a topical anesthetic duration of 30 minutes, a 2-hour application is necessary.

Tetracaine Base Patch and EMLA Cream
Because both agents demonstrate a reservoir effect, anesthesia may increase or begin minutes after removal of the drug. A precise description of application times and duration is not possible. Tetracaine base requires a minimum of 20 to 30 minutes of application time to produce several hours of anesthesia. EMLA requires an application time of 1 to 2 hours for a reported duration of 30 minutes to several hours. Occlusive dressings seem to increase penetration of EMLA whenever the cream is used. Patches are more convenient and cause no loss of effectiveness.

EMLA dosing is based on the amount of cream applied, not on the amount of anesthetic. Each gram of EMLA cream contains 25 mg of lidocaine and 25 mg of prilocaine. Dosages are given in grams of cream, not milligrams of anesthetic. In general, apply EMLA as a thick layer over intact skin under an occlusive dressing for approximately 1 hour before a procedure. Application of a thick layer approximates to 1 to 2 g/10 cm². For minor procedures such as needle insertions, apply 2.5 g of EMLA over 20 to 25 cm² for at least 1 hour. For more painful procedures, apply approximately 2 g of cream per 10 cm² for at least 2 hours. The maximum application area (MAA) determines the appropriate total dose applied. Base the MAA on the patient’s weight as follows: up to 10 kg, MAA = 100 cm²; 10 to 20 kg, MAA = 600 cm²; more than 20 kg, MAA = 2000 cm².

Ethyl Chloride and Fluori-Methane Sprays
Collect all the equipment required and make all preparations needed to immediately perform the desired procedure. Invert the bottle 25 cm from the skin and spray a stream along the proposed incision site until the area turns white and hard. Make the incision or local anesthetic injection immediately or during actual spraying of the agent for optimal results because the effect is fleeting. Some clinicians use these vapocoolant sprays to decrease the pain of injection of a more traditional local anesthetic such as lidocaine.

Iontophoresis
Anesthetic agents may be drawn into the skin without needles by electrical current applied through electrodes in a process called iontophoresis. Lidocaine with epinephrine administered via iontophoresis provides adequate anesthesia before venipuncture in pediatric patients and is superior to EMLA in providing cutaneous anesthesia. Iontophoresis is not widely used in emergency medicine but may be another alternative to applied anesthetic agents.

Microneedle Pretreatment
Recently developed technology known as microneedle pretreatment may improve the efficacy of cutaneously applied topical anesthetics. A functional microarray of fine needles painlessly perforates the stratum corneum to facilitate the penetration of applied medications without using traditional needle injections. A 2010 study demonstrated that perforation of the skin of healthy volunteers with a microneedle to which dyclonine was applied resulted in decreased time to anesthesia and a greater degree of pain reduction than did application of dyclonine to nonperforated skin. Microneedling followed by the application of EMLA prior to facial laser resurfacing, and lidocaine 2.5%/prilocaine 2.5% prior to full face fractional microneedling for atrophic acne scars was superior than either topical anesthetic applied alone. This emerging technology may improve the practical application of anesthetics to intact skin in the ED.

Jet Injection
Jet injection of anesthetic through intact skin may overcome some of the limitations imposed by the cutaneous application of anesthetics without needle infiltration. A jet injector is a device containing carbon dioxide gas that rapidly forces a plunger to expel the drug through a small orifice applied over intact skin. Medication penetrates the epidermis to a depth of 5 to 8 mm in 0.2 second and causes the drug to be rapidly dispersed through the skin. Safety is improved because no needles are used to penetrate the skin. Several studies have demonstrated that jet injection of anesthetics provides more rapid anesthesia than topical application of anesthetic agents and is preferred by patients over no anesthesia at all for painful procedures such as laceration repair, lumbar puncture, IV insertion, and arterial blood gas sampling.

Complications
General adverse reactions to anesthetic agents are discussed in the Complications section later in this chapter. Tetracaine base is quite safe, with a low blood concentration after proper use. In one study, cutaneous erythema developed at the site of application in approximately 25% of patients. This vasodilatory effect may actually be an advantage when starting IV lines or performing venipuncture. EMLA is also quite safe. Although it has a high rate of local skin reactions, they are mild and transient, with disappearance 1 to 2 hours after removal of the cream. Despite the reported successful
Lacerations

Background

In 1980, Pryor and colleagues reported their experience with a topical anesthetic solution (tetracaine-adrenalin-cocaine [TAC]) for wound repair. The original formula, used in most subsequent studies, consists of a solution of 0.5% tetracaine, 1:2000 epinephrine (adrenaline), and 11.8% cocaine. Traditionally, anesthesia is produced by firmly applying a solution-saturated gauze pad or cotton ball directly to the laceration for 10 minutes. The resulting loss of cutaneous sensation is centered about the area of application.

Topically applied anesthetics are often covered with an occlusive dressing to contain the medication. Covering the medication with an occlusive dressing may dramatically increase drug absorption and the amount of metabolites. Oni and associates demonstrated that 4% lidocaine cream applied to volunteers with an occlusive dressing resulted in three times the level of serum lidocaine and double the level of MEGX when the same dose was applied to volunteers without occlusion. Use caution when occluding topical anesthetics over intact skin because markedly elevated serum levels of the drug may be reached.

Indications and Contraindications

Use of TWA is generally restricted to young children with wounds less than 5 cm in length, in whom the delay in application of an anesthetic is acceptable and proper application can be ensured (Fig. 29.5). TWA containing vasoconstricting agents is not generally used in structures without a collateral blood supply (e.g., digits, tip of the nose, pinna of the ear, penis), although there is no evidence that this is unsafe. When some wound preparation is desired before anesthesia, remove large use of EMLA cream on skin ulcers, Hansson and associates described an increase in bacterial growth, infection, and inflammation when used in experimental wounds. Methemoglobinemia resulting from the metabolites of prilocaine may occur with EMLA. The risk for clinically significant methemoglobinemia seems remote when EMLA is used properly. It is contraindicated in any infant younger than 3 months and in infants between 3 and 12 months of age who are currently taking methemoglobinemia-inducing drugs (nitrites, sulfonamides, antimalarials, phenobarbital, and acetaminophen). The risk for adverse effects is increased in patients with anemia, respiratory or cardiovascular disease, and deficiencies in glucose-6-phosphate dehydrogenase or methemoglobin reductase. Prolonged inhalation of ethyl chloride spray may produce general anesthesia, coma, or cardiorespiratory arrest. Ethyl chloride is also flammable, thus precluding its use with electrocautery.

Use caution when occluding topical anesthetics over intact skin because markedly elevated serum levels of the drug may be reached.

Figure 29.5 Topical Wound Anesthesia

A, The topical anesthetic (in this case, lidocaine-epinephrine-tetracaine gel) is applied to a cotton ball or gauze pad. B, The anesthetic is held in place with an occlusive dressing such as Tegaderm. C, After a 30-minute application period, vasoconstriction of local tissue can be readily observed (arrow), and effective anesthesia has been achieved. D, Local wound repair, including additional infiltrative anesthesia if necessary, can proceed with little if any patient discomfort.
debris and clotted blood to allow the appropriate application of TWA and then finish wound preparation once the wound is properly anesthetized.

TWA appears to be less effective on the trunk and extremities than on the face and scalp and less effective than lidocaine infiltration in these areas. The 10- to 20-minute onset time may be unacceptable in a busy ED. Two other often mentioned drawbacks may be more theoretical than real. Vasoconstrictor-induced higher infection rates (see later section on Complications) have not been clearly demonstrated. The argument that the necessary 10-minute application period is time-consuming and takes valuable nursing time is partially offset by using the child’s caretaker or adhesive paper tape alone to hold the solution in place. It is not necessary for anyone to “hold” the medicine in place when gel is used.

Some EDs still stock topical anesthetic gels that contain cocaine. Cocaine use is complex because of cost and federal regulations requiring storage in a locked cabinet and maintenance of separate written records of its use. In view of the evidence that mixtures without cocaine are efficacious, avoid the potential complications related to cocaine that do not have the potential for abuse that cocaine does. The practicality of cocaine-containing TWA has also been questioned.

**Agents and Effectiveness**

**TAC and Related Mixtures**

Three clinical trials directly compared TAC with infiltrated lidocaine. Without specifying wound location, Pryor’s group found equal anesthetic effect. Complete anesthesia with TAC was achieved in 82% to 86% of patients as compared with 83% to 92% for subcutaneous lidocaine. The remaining patients obtained partial anesthesia. Hegenbarth and associates and Anderson and coworkers demonstrated results similar to those of Pryor’s group. Hegenbarth and associates found TAC to be equal to lidocaine only on the face and scalp and inferior to it at other locations. In contrast, other studies have confirmed excellent rates of effectiveness, especially on the face and scalp. TAC is more effective than its component drugs alone. On the face and scalp, TAC was found to be superior to tetracaine alone, although on nonfacial areas, both produced equally poor results. TAC was found to be more effective than cocaine alone and more effective than a tetracaine-epinephrine solution in the same dosage ratio.

In 1990, Bonadio and Wagner showed that an epinephrine-cocaine solution (1:2000 epinephrine and 11.8% cocaine) was equal to TAC in effectiveness. Bonadio and Wagner also found that half-strength TAC (0.25% tetracaine, 1:4000 epinephrine, 5.9% cocaine) achieved excellent results in patients with dermal lacerations of the face, lip, and scalp. Smith and Barry compared three strengths of TAC and found equal effectiveness among them; they recommended the lowest-strength cocaine formulation (1% tetracaine, 1:4000 epinephrine, 4% cocaine). Ernst and colleagues found similar effectiveness to TAC when a slightly different lidocaine-epinephrine-tetracaine (LET) solution (4% lidocaine, 1:2000 epinephrine, 1% tetracaine) was used.

TAC has also been compared with EMLA when placed in a wound for 60 minutes before wound repair. In a study of 32 wounds, supplemental anesthesia was required less often with EMLA. The non–cocaine-containing formulations are generally considered less toxic and have advantages in terms of reduced cost and avoidance of controlled substance precautions during storage.

**LET and Related Solutions**

Many variations of mixtures that do not contain cocaine have been studied; these include LET, lidocaine-epinephrine, tetracaine-phenylephrine, tetracaine-lidocaine-phenylephrine, bupivacaine-norepinephrine, and prilocaine-phenylephrine. Schilling and associates found that a LET solution (4% lidocaine, 1:1000 epinephrine, 0.5% tetracaine) was as effective as TAC. LET gel preparations are at least as effective as LET solutions. Singer and Stark determined that EMLA or LET gel placed in a wound on initial evaluation and before infiltration with lidocaine reduced the pain of infiltration, thereby allowing an essentially painless injection of lidocaine. Gauze soaked with lidocaine 2% without vasoconstrictive agents placed in a wound was not effective in eliminating the need for infiltrative anesthesia prior to suturing.

**Technique and Dosage**

Because the topical mixtures noted earlier (especially TAC) are not innocuous anesthetics, pay close attention to the technique of application and the recommended maximum dose. There is no uniformly accepted application technique, composition of components, or concentration of components.

Generally, apply topical solutions such as TWA to the wound in a gravity-dependent position and carefully fill the wound cavity. After 3 minutes, place a single 2 x 2-cm gauze pad or cotton ball saturated with TWA on the wound. Tape or hold the pad firmly in place for 15 to 20 minutes. The person holding the gauze should wear latex examination gloves to minimize the risk for cutaneous absorption of the solution. The average dose of TWA mixture for most wounds is 2 mL.

Gaufberg and coauthors described “sequential layer application” (SLA) of topical lidocaine with epinephrine (TLE) to deeper wounds that might not typically be amenable to the use of TWA. Cotton soaked with TLE was applied to the wound surface and 2 mm of surrounding skin for 10 to 15 minutes and then removed. A second similarly soaked piece of cotton was packed deeper into the wound for 10 to 15 minutes. Deep wounds were treated with a third piece of TLE-soaked cotton placed deeper into the wound. Anesthesia was presumed when 3 mm or more of pallor was seen on all wound margins. The group receiving lidocaine by SLA was compared with a group receiving infiltrated 2% buffered lidocaine with epinephrine. Both groups experienced a similar degree of anesthesia during suturing. Mean time to anesthesia with SLA was 29 minutes versus 5 minutes for the infiltration group. Despite the longer time in the SLA group, the authors argued that the additional time is worthwhile because the painful injection, risk for hollow-bore needle injury, and tissue distortion with infiltration were eliminated. If time permits, the SLA technique may be useful for wounds that would have otherwise required infiltrated anesthesia.

Hegenbarth and associates estimated the maximum safe dose of full-strength TAC to be 0.09 mL/kg based on the known maximal safe dose of infiltrated tetracaine, the mucosal application of cocaine, and an estimate of absorption of solution onto the applicator. The key to safety is to avoid TAC on mucosal surfaces or areas in which sniffing or swallowing may accidentally occur. Topical mucosal anesthesia is discussed elsewhere in this chapter.

Prepare an epinephrine-cocaine gel by adding 0.15 g of methylcellulose to 1.5 mL of epinephrine-cocaine solution. Stir the mixture thoroughly for a minute or two until a gel consistency is obtained. After sterile preparation, place the
wound in a gravity-dependent position and apply the gel with a cotton-tipped swab to coat the entire wound cavity and margins. Allow the wound to stand for 15 to 20 minutes and thoroughly wash the wound cavity to remove the gel. In Bonadio and Wagner’s study, the average dose used was 0.35 mL of gel containing only 40 mg of cocaine. Again, other agents are equally efficacious and safer; select these over cocaine-containing mixtures whenever possible.

Complications
Most adverse events associated with TWA occur with mixtures that contain cocaine. Lidocaine toxicity is a well-recognized, adverse event when higher doses are used (see later section on Systemic Toxic Reactions), but lidocaine toxicity has not been reported with TWA mixtures.

Mucosal application may rarely lead to significant systemic toxicity; fatalities have been reported after application. Even after nonmucosal TAC use, cocaine levels appear in the blood and cocaine metabolites appear in the urine in the majority of patients, but tetracaine does not. Gel formulations of TAC tend to stay in the wound and thus reduce the risk of solution runoff onto mucosal surfaces. There is no need to use a gauze pad to apply the medication in a gel formulation or to hold it in place. Because the entire applied dose will stay in the wound, a lower dose can be used. Gel also provides more uniform application to tissues, which improves the anesthetic effect. In tissues containing end-arteries, ischemia caused by vasoconstrictors may occur. Avoid TAC on the digits, the tip of the nose, the penis, and the pinna of the ear. Use TAC with caution on patients with coronary artery disease, uncontrolled hypertension, seizures, or peripheral vascular disease. Patients with decreased plasma cholinesterase levels are theoretically at increased risk for systemic toxic effects, but this potential risk is of little clinical concern.

INfiltrATION ANESTHESIA

Background
Injection of an anesthetic agent directly into tissue before surgical manipulation is known as infiltration anesthesia (Fig. 29.6). Field block anesthesia is also considered a form of infiltration anesthesia, particularly as the agents, concentrations, and recommended maximum dosages are the same. To create a field block, inject a field of anesthetic around the operative site. Make the injection proximal to or surrounding the area that you plan to manipulate. Combine infiltrative anesthesia with procedural sedation (see Chapter 33) to reduce anxiety or motion.

Indications and Contraindications
Infiltration anesthesia is indicated when good operative conditions can be obtained with this technique. It may be used for the majority of minor surgical procedures, such as excision of skin lesions and suturing of wounds. Infiltration anesthesia is considered quicker and safer than nerve block and general anesthesia. Local infiltration can provide hemostasis, both by direct distention of tissue and by the concurrent use of epinephrine.

A disadvantage of local infiltration over nerve blocks is that a relatively large dose of the drug is needed to anesthetize a relatively small area. For extensive wounds, the amount of anesthetic required may risk systemic toxicity. The maximum allowable volume can be increased by adding epinephrine, using a lower concentration of the anesthetic agent, or both (Table 29.5). When large volumes are anticipated and a nerve block is anatomically feasible, the nerve block is preferred. Avoid using infiltration for large procedures in small children and in apprehensive patients, especially those with previous adverse reactions to the medications (whether vasovagal or otherwise). Local infiltration distorts the tissues that will be incised or repaired, which makes it undesirable in areas requiring precise anatomic alignment (e.g., some lip repairs).

![Figure 29.6 Infiltration anesthesia for incision and drainage of a pilonidal abscess. Minimize pain by using a small needle (ideally 30-gauge if injecting through intact skin or 25- to 27-gauge if going through the cut edges of a wound), buffering the anesthetic with sodium bicarbonate, warming the anesthetic to body temperature, and injecting slowly in the subdermal plane.](image)

<table>
<thead>
<tr>
<th>TABLE 29.5 Maximum Allowable Volume (Adults)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>AGENT</strong></td>
</tr>
<tr>
<td>-----------------</td>
</tr>
<tr>
<td>Lidocaine</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Bupivacaine</td>
</tr>
<tr>
<td>Lidocaine-epinephrine</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Bupivacaine-epinephrine</td>
</tr>
</tbody>
</table>

*These are quite conservative figures for infiltration anesthesia; see text for explanation.
*Also see Table 29.9. A 0.5% solution is 0.5 g/100 mL = 500 mg/100 mL = 5 mg/mL. A 1% solution is 1 g/100 mL = 1000 mg/100 mL = 10 mg/mL.
*Some physicians recommend 400 mg as the maximum safe dose for bupivacaine.
Choice of Agent

The local anesthetic agents most frequently used for infiltration are 0.5% to 1% lidocaine, 0.5% to 1% procaine, and 0.25% bupivacaine (Table 29.6). Higher concentrations are of no additional benefit. Lidocaine is most commonly used because of its excellent activity profile, low allergenicity, low toxicity, user familiarity, and ready availability. Procaine is useful for patients who are allergic to amide anesthetics. Some clinicians prefer bupivacaine because of its prolonged duration. Bupivacaine may also be preferred when postoperative analgesia is desired, for prolonged procedures, for dental anesthesia, or even for short procedures that may be interrupted in a busy ED.

A comparison of equianesthetic doses of lidocaine and bupivacaine for infiltration anesthesia (Table 29.7) revealed that the duration of action is the major difference between the two agents. For the majority of ED procedures it is not necessary to extend the duration of anesthesia beyond 1 hour, which makes plain lidocaine a logical anesthetic choice. However, patients experience a moderate amount of pain after repair of a laceration when the lidocaine wears off in approximately 1 hour. Bupivacaine reduces the pain after repair of lacerations for at least 6 hours. This benefit of a prolonged duration of anesthesia must be weighed against the hazards of injury to the mucous membranes or an unprotected limb or the annoyance of prolonged numbness in patients who have undergone simple surgical procedures.

A prolonged duration of anesthesia can also be achieved by adding epinephrine, sodium bicarbonate, or both to lidocaine. Epinephrine provides excellent wound hemostasis and slows systemic absorption. This latter property decreases the peak blood level, reduces the potential for a toxic reaction, and allows a greater volume of agent to be used for extensive lacerations. The major disadvantage of epinephrine is theoretical damage to host defenses, but it is generally clinically inconsequential (Box 29.2). Bicarbonate added to the anesthetic just before injection decreases the pain of administration. Bupivacaine, if used with due caution, is safe and easy to use. The deciding factors are many, but some logical choices are as follows:

- For a wound with excessive bleeding, use lidocaine with epinephrine and sodium bicarbonate.
- For an apprehensive patient, use lidocaine with sodium bicarbonate.
- For anticipated prolonged postprocedure pain, use bupivacaine.

Equipment

The pain of injection is reduced with the use of small-gauge needles. Ideally, a 30-gauge needle is used if the injection is...
made through skin. If the injection is made through the cut edges of the wound, a 25- to 27-gauge needle suffices. A small-gauge needle slows the rate of injection and reduces the rate and pain of tissue distention. A 10-mL syringe is recommended both for its ease of handling and for the relatively slow rate of injection that it allows.

**Technique**

Once an agent has been chosen, proper administration technique minimizes pain, prevents spread of bacteria, and avoids intravascular injection. Buffering, temperature manipulation, and careful infiltration also reduce the pain of injection.

**Buffering**

Raising the pH of an anesthetic by adding sodium bicarbonate decreases pain dramatically, whereas lowering the pH by adding epinephrine increases pain. Buffering is probably the best way to reduce the pain of local anesthetic injections, and its routine use is highly recommended. It is probable that pH is not the sole factor in producing pain because the pain produced by various agents does not correlate strictly with the pH. Sodium bicarbonate probably works by increasing the ratio of nonionized to ionized molecules, which either renders the pain receptors less sensitive or causes more rapid diffusion of solution into the nerve and a shorter time to the onset of anesthesia.

To alkalize lidocaine, add 1 mL of sodium bicarbonate (8.4%, 1 mmol/mL) to every 10 mL of anesthetic solution. As the pH of the solution is raised, the anesthetic becomes unstable and has a decreased shelf life. It was initially recommended that buffered lidocaine be prepared just before use to avoid precipitation and degradation, but buffered lidocaine retains its effectiveness for 1 week and refrigeration may further increase its shelf life. Bicarbonate may be combined with plain lidocaine for both infiltrative anesthesia and digital nerve blocks. In one volunteer study, sodium bicarbonate was effectively combined with lidocaine and epinephrine.

Sodium bicarbonate can be added to bupivacaine, but the solution tends to precipitate as the pH rises. The clinical effect of such precipitation is unclear, but if it occurs, it is probably prudent to use another solution prepared with less buffer.

**Temperature Manipulation**

Warming an anesthetic to body temperature (37°C to 42°C) reduces the pain of infiltration, but warming may not reduce injection pain as much as buffering with sodium bicarbonate does. Bartfield and colleagues found that lidocaine warmed to 38.9°C was more painful than room-temperature buffered lidocaine during intradermal injection. Brogan and associates, using lidocaine warmed to 37°C, found the warmed lidocaine and room-temperature buffered lidocaine to be equivalent during wound infiltration. Neither study found a synergistic effect with combined warming and buffering. Martin and coworkers found that warmed (37°C) lidocaine was less painful than buffered lidocaine. Anesthetic solutions can be warmed in a baby food warmer with thermostat temperature control or in an IV solution warmer. Warming is not believed to adversely affect the shelf life of the local anesthetic.

Locally cooling the area to be infiltrated may provide additional pain relief. Leff and colleagues demonstrated that patients receiving local infiltrative anesthesia for repair of an inguinal hernia had less pain when the incision area was cooled before infiltration. Patients were randomized to two groups and had a 1000-mL bag of saline placed over the inguinal area for 5 minutes. One group used saline bags at room temperature, the other group used saline bags cooled to 4°C. A significant decrease in pain perception was found in the group in which the inguinal area was cooled before lidocaine infiltration. Cooling of wounds in the ED before infiltration has not been studied.

**Injection**

The pain from injection of local anesthetics is primarily a result of skin puncture (which can be minimized with small-gauge needles) and subcutaneous injection. Although patients fear the needle, there is little perceived pain merely from the needle’s presence in subcutaneous tissue. Place the injection in subdermal tissue to minimize needle puncture pain and the tissue distention that occurs with intradermal placement. Place the needle “up to the hub” and inject while withdrawing along the just-created subdermal tunnel to minimize tissue distention. After an initial injection, instead of totally withdrawing the needle from the tissue, redirect it along another path to lessen the number of skin punctures. Slowly inject the smallest volume necessary to reduce pain. Hamelin and coworkers demonstrated that blinded volunteers receiving digital nerve block with 2 mL of lidocaine and epinephrine (1:100,000) experienced less pain when the solution was injected over 60 seconds rather than over 8 seconds.

Because the patient barely feels a needle placed subcutaneously and skin puncture is often quite painful, make all wound

**Box 29.2 Epinephrine Use**

**ADVANTAGES**

1. Prolongs the duration
2. Provides hemostasis
3. Slows absorption:
   - Decreases the agent’s toxic potential
   - Allows an increased dose
4. Increases the level of blockade

**DISADVANTAGES**

1. Impairs host defenses—increases the incidence of infection
2. Delays wound healing
3. Do not use for:
   - Areas supplied by end-arteries
   - Patients “sensitive” to catecholamines
4. Toxicity—catecholamine reaction

*Based on laboratory studies and is of unknown clinical importance.

*For example, in patients taking monoamine oxidase inhibitors.

Precipitation varies directly with the concentration of bupivacaine and the time since mixture. Cheney and coworkers showed that 0.05 mL of 8.4% sodium bicarbonate (measured in a tuberculin syringe) could be mixed with 10 mL of 0.5% bupivacaine without precipitation. The goal of using bupivacaine is to prolong the duration of anesthesia; this effect can also be accomplished somewhat by using buffered lidocaine (plain or with epinephrine).

In view of the amount of published literature demonstrating that adding sodium bicarbonate to lidocaine before infiltrative anesthesia reduces the pain of injection without adverse effects, lidocaine should almost always be buffered when infiltrated in the ED; there is no practical reason to avoid the addition of bicarbonate.
injections through the wound edge and not through the skin (Fig. 29.7). Spread of infection beyond the wound margin has not been demonstrated clinically with this technique. Some clinicians choose to inject the anesthetic through intact skin in patients with a grossly contaminated wound. Bierman described a technique of patient distraction by applying light pressure to alternate sides of the wound with one’s fingers and repeated ambiguous questioning about feeling the light pressure rather than the ongoing wound injection. Ask school-aged children to count backward or to say the “ABCs” to distract them during injection. Inject 1% lidocaine through intact skin with the needle bevel facing upward because this is less painful than injecting with the needle bevel facing downward.

To prevent a systemic toxic reaction, avoid giving an intravascular injection. For infiltration anesthesia with a small-gauge needle, however, aspiration is usually unnecessary unless the injection is deeper than the subcutaneous area or the area to be injected contains many large vessels.

**SPECIAL CONSIDERATIONS**

**Hematoma Block**

Hematoma block (Video 29.1) has been used for many years to provide anesthesia for fracture reduction, particularly of the distal end of the forearm and hand (Fig. 29.8). Its popularity has waned somewhat because of the fear (unproven and theoretical) of introducing infection at the fracture site and its limited efficacy. Although several studies have shown hematoma block to be safe, the anesthesia that it provides is not as good as that provided by a Bier block (see Chapter 32). Nevertheless, there are several reasons to consider this technique.

**Figure 29.7** A, Except in the setting of gross contamination, wounds should be anesthetized by inserting the needle through the cut edges, not through the intact skin. Patients often will not feel a 25-gauge or smaller needle passed into subcutaneous tissue when it is advanced slowly through the cut edge. However, pain generally occurs with distention of tissue by the anesthetic, and hence injection should be slow and deliberate. B, If a wound is grossly contaminated, the anesthetic may be introduced through the intact skin. The operator should limit the number of needlesticks. The needle is first introduced at a point in line with the wound and beyond the wound edge (1), and while the anesthetic is slowly injected, the needle is advanced to include one entire side of the wound (if possible) to a point well past the opposite end of the wound. If the needle is not long enough to encompass the entire wound, the skin is painlessly punctured at a midway point that has already been anesthetized (2). The other side may be anesthetized by passing the needle through the area already infiltrated by the first injection (3) to make the skin puncture painless. A 3.8-cm (1.5-inch) 27-gauge needle is a good choice. C, Inject local anesthetic through subcutaneous tissue, not intact skin.

**Figure 29.8** Hematoma block. This technique is useful for reducing fractures of the distal end of the forearm and hand. Here it is being used before reduction of a distal radius (Colles) fracture. To perform the block, slowly inject 5 to 15 mL of plain 1% lidocaine into the fracture cavity and around the adjacent periosteum. It may require 10–15 minutes for anesthesia to occur.
lower dose of the anesthetic agent is required for a hematoma block than for a Bier block (see Chapter 32). A hematoma block is particularly useful when a Bier block and general anesthesia are contraindicated.

Prepare the skin over the fracture site with antiseptic solution and insert the needle into the hematoma, as confirmed by aspirating blood. Slowly inject 5 to 15 mL of plain 1% lidocaine or 5 to 10 mL of plain 2% lidocaine into the fracture cavity and around the adjacent periosteum. Larger fractures require larger volumes of local anesthetic. Adequate anesthesia occurs in approximately 5 to 10 minutes and may last for several hours. A common error is to attempt the procedure too soon after injection. Do not perform this procedure through dirty skin or with open fractures.

**Intraarticular Anesthesia and Analgesia**

Findings from the history and physical examination of an acutely traumatized joint, such as the knee, often underestimate the severity of the injury. Instillation of 5 mL of 1% lidocaine after joint aspiration may help relieve pain and facilitate an examination, but its use is not routinely recommended. As spasm and apprehension are often not relieved by local anesthesia, the information gained from this maneuver does not usually influence the ED treatment plan. Intraarticular anesthesia of the knee has no effect on gait pattern or joint proprioception. Postprocedure weight bearing may be allowed without fear of producing or increasing injury if otherwise indicated. Intraarticular anesthesia may enhance elbow use after aspiration of a hematoma associated with a radial head fracture. The technique of administration is analogous to arthrocentesis (see Chapter 53). Intraarticular lidocaine has been effective in facilitating reduction of a shoulder dislocation. Animal experiments have demonstrated chondrotoxicity when local anesthetics are continuously infused into a joint. Although continuous joint infusion of anesthetic may be harmful when used in a postoperative setting, there is no evidence that a single injection of intraarticular anesthetic in the ED is harmful.

Morphine may be injected directly into joints for postoperative pain relief and potentially provide prolonged analgesia after reduction of fractures. Theoretically, there are local opioid receptors in joints that are capable of being stimulated by rather small doses of morphine to provide relief for up to 24 hours with a single dose. There may also be some systemic absorption of the intraarticular morphine that contributes to analgesia, but adverse systemic opioid effects are not seen with this technique. Data are sparse on the exact mechanism and optimal doses. A period of 3 to 6 hours may be required for analgesia to reach its maximum effect. In a systematic review, Gupta and colleagues concluded that postoperative intraarticular morphine injected into the knee joint at doses of 2 to 4 mg provides analgesia for up to 24 hours. There is a wide variability in efficacy, which may be dose related. Though studied primarily for postoperative use after knee surgery, a similar concept may be intuitively applied to traumatic joint pain, but this has not been studied adequately.

**Intrapleural Anesthesia**

**Indications**

Intrapleural anesthesia introduces a local anesthetic into the pleural space (i.e., between the parietal and visceral pleura) through an epidural catheter. The anesthetic can also be introduced through a previously placed chest tube. The technique can provide relief for several conditions, primarily postthoracotomy pain, postcholecystectomy pain, and most importantly for emergency clinicians, posttraumatic chest pain (e.g., rib fractures, pneumothorax, hemothorax). This procedure is not only useful for pain relief but also facilitates turning, coughing, and deep breathing. Several studies have demonstrated improved respiratory mechanics when intrapleural anesthesia is used. Though not unanimous, most studies show that intrapleural anesthesia is effective in providing analgesia. Concern has been raised that intrapleural anesthesia may create a level of anesthesia below the umbilicus and make posttraumatic abdominal examinations unreliable. Until this issue is clarified, it seems prudent to rule out intraabdominal injury before intrapleural anesthesia is used.

**Technique**

If a chest tube is in place, it is preferable to inject anesthetic into the pleural space through the chest tube. Clamp the tube for 10 to 15 minutes to allow the anesthetic to diffuse. When the tube cannot be taken off suction or if no tube is present, inject the local anesthetic percutaneously. With the patient in the lateral position (with the affected side up), place a 16-gauge Tuohy needle 8 to 10 cm from the posterior midline in the eighth intercostal space. Angle the needle 30 to 40 degrees with respect to the skin and aim medially, with the bevel up and directed just above the rib. After perforating the posterior intercostal membrane (felt as a distinct resistance), remove the stylet and attach a well-wetted, air-filled glass syringe to the Tuohy needle. Advance the needle until it enters the pleural space, which is denoted by the plunger being drawn down the syringe as a result of the negative pressure created during inspiration. Remove the syringe and introduce an epidural catheter 5 to 6 cm into the pleural space. Remove the Tuohy needle, obtain a chest radiograph to confirm proper position, and secure the catheter.

The most commonly used anesthetic and dose is 20 mL (0.3 mL/kg) of 0.5% bupivacaine. A dose repeated every 8 hours is safe and effective. Alternatively, infuse 0.25% bupivacaine at 0.1 to 0.2 mL/kg per hour after a bolus has been administered. The solution presumably diffuses from the pleural space “back” through the parietal pleura and the intercostal muscle to reach the intercostal spaces, where it blocks the intercostal nerves. The level of anesthesia can extend from T2 to T12 and involve the skin, chest, abdominal wall, and potentially the viscera if the visceral afferent fibers are blocked at the sympathetic chain in the paravertebral gutter.

Though not yet a consistently proven or a completely standardized technique, intrapleural anesthesia offers promise for patients and is a valuable procedure for the emergency clinician to master.

**COMPLICATIONS**

**Local Anesthetic Effect on Wounds**

**Wound Healing**

Local anesthetics produce cytotoxic effects on cell structure and function in a dose- and time-related manner. These effects, at doses well below those used clinically, involve fibroblasts more than nervous tissue. Local anesthetics impair mitochondrial function and fibroblast proliferation and hasten apoptosis in...
vitro; the clinical relevance of these effects to the ED use of local anesthetics is likely to be minimal.\textsuperscript{103,104} Collagen synthesis is inhibited by lidocaine and bupivacaine.\textsuperscript{105} Morris and Tracey\textsuperscript{106} found that lidocaine in increasing concentrations progressively reduces the tensile strength of wounds. Epinephrine added to 1% and 2% concentrations of lidocaine further reduced tensile strength, but when epinephrine was added to distilled water or to 0.5% lidocaine, it had little effect. Several conclusions may be clinically relevant. Although it may delay the onset of anesthesia, 0.5% lidocaine solution, without epinephrine, may be best for maintaining wound strength.

Eriksson and associates\textsuperscript{107} found that lidocaine reduces the inflammatory response in wounds by decreasing the number of white cells and their metabolic activity. Although an inflammatory response may be beneficial in a contaminated wound, it can be detrimental in a sterile wound because of tissue toxicity created by the release of superoxide anions, lysosomal enzymes, thromboxanes, leukotrienes, and interleukins. The clinical relevance of this is unknown. None of the concerns mentioned earlier should prohibit the use of standard anesthetics or epinephrine when their use is otherwise indicated.

**Wound Infection**

Though not generally appreciated, it has long been known that local anesthetics possess antimicrobial activity in vitro. Lidocaine and procaine demonstrate concentration-dependent inhibition of culture growth of most gram-negative organisms.\textsuperscript{108-110} Gram-positive isolates are also significantly affected by lidocaine and, to a lesser extent, by procaine. Lidocaine inhibits the growth of common nosocomial pathogens, including *Enterococcus faecalis*, *Escherichia coli*, *Pseudomonas aeruginosa*, and several strains of methicillin-resistant *Staphylococcus aureus* and vancomycin-resistant enterococci.\textsuperscript{111} Administering anesthetics before obtaining material for culture, including injecting a joint before arthrocentesis, may give false-negative culture results. To avoid this problem, if possible inject the skin at the injection site and along the needle tract but not into the actual joint space until after a synovial fluid specimen has been obtained for culture. Berg and coworkers\textsuperscript{112} demonstrated that lidocaine administered before tissue biopsy of chronic wounds did not affect the culture results when the exposure time before culture was less than 2 hours. This effect is also significant when anesthetic ointments are applied before culture. EMLA cream applied before culture demonstrates powerful antimicrobial properties.\textsuperscript{113} Furthermore, it has been shown that adding sodium bicarbonate to lidocaine greatly enhances its inhibitory effect on bacteria.\textsuperscript{114} Although local anesthetics can interfere with culture testing, several studies have shown that local anesthetics, by themselves, do not appear to alter the incidence of wound infection.\textsuperscript{113,114}

Epinephrine appears to exert a deleterious effect on host defenses, at least in animal models. Studies of infiltrated and topically applied epinephrine solutions in contaminated animal wounds show an increased potential for infection.\textsuperscript{113,114} Epinephrine-induced vasoconstriction may contribute to tissue hypoxia, retard the killing of *S. aureus* by leukocytes, and reduce leukocyte migration into the tissue.\textsuperscript{115} Most clinical studies using topical anesthetics with vasoconstrictor properties (e.g., TAC mixtures) do not demonstrate a significant rise in infection rates.\textsuperscript{30,31,32,27} The concerns mentioned earlier should not prohibit the use of epinephrine for wound preparation when its use is otherwise appropriate.

**Local Injuries**

Injury may result from the direct application of an anesthetic agent to a nerve or from passage of a needle through soft tissue structures. Factors implicated in transient or persistent neuropathy include acidic solutions, additives, the agent itself, needle trauma, compression from hematomas, and inadvertent injection of neurolytic agents. Born\textsuperscript{116} described a series of 49 wrist and metacarpal blocks with bupivacaine in which significant neuropathy developed in eight patients. He postulated that damage occurred from trapping the drug in a confined space and recommended that whenever bupivacaine is used in this situation, it should be low in concentration and volume. Infection, hematomas, and broken needles are other local problems that can be avoided by using proper technique. Erroneous needle placement can also produce complications such as pneumothorax during a brachial plexus or intercostal block.

**Use of Epinephrine With Local Anesthetics**

Epinephrine in conjunction with local anesthetics prolongs the duration of anesthesia and produces a temporary hemostatic effect, but its inclusion in digital block solutions has traditionally been discouraged because of the belief that it can lead to ischemia and necrosis. Areas of special concern include the digits, penis, tip of the nose, and earlobe. Although tissue ischemia and sloughing have been reported with concentrations of 1:20,000, current practice involves concentrations in the range of 1:100,000 to 1:200,000 and the use of submaximal doses. Several authors suggest that epinephrine-containing solutions can be safely injected into the fingers without adverse sequelae.\textsuperscript{117-120}

A study of more than 3000 cases of elective injection of low-dose epinephrine (≤ 1:100,000) into the hand and fingers failed to identify a single case of digital tissue loss, and phentolamine (Regitine, Novartis Pharmaceuticals Corp., East Hanover, NJ) was not required to reverse vasoconstriction.\textsuperscript{117} Of 1111 procedures reported by Chowdhry and coauthors\textsuperscript{121} involving digital blocks with 1% lidocaine plus epinephrine (1:100,000) at a dose ranging between 0.5 and 10 mL (average, 4.3 mL), no patient in the epinephrine group exhibited digital ischemia or experienced nerve injuries or unusually delayed wound healing. Current data support the use of epinephrine, when correctly applied, for the performance of digital blocks of the fingers and toes.

The use of phentolamine, which produces postsynaptic α-adrenergic blockade, is recommended for clinically significant vasoconstrictor-induced tissue ischemia. This medication is usually given by local infiltration, in the area where epinephrine has been injected, at a dose of 0.5 to 5.0 mg diluted 1:1 with saline. If local infiltration is ineffective because of tension within a tissue compartment or if the area of vasoconstriction is large, give phentolamine by the intraarterial route.\textsuperscript{122}

**Systemic Toxic Reactions**

Although systemic toxic reactions occur in only 0.1% to 0.4% of patients after the administration of local anesthetics, they are the most frequent serious adverse reactions encountered (Table 29.8).\textsuperscript{123} After the administration of a local anesthetic, some of the drug reaches its intended target and some is absorbed quickly into the systemic circulation. Peak blood levels are generally produced within 30 minutes. Many vagal reactions, nonspecific anxiety reactions, and sensitivity to preservatives...
Dose and Concentration

The larger the total dose, the higher the peak blood level. It is uncertain whether increasing the concentration while maintaining the total dose by decreasing the volume affects the serum level.

Addition of Epinephrine

Epinephrine produces vasoconstriction and reduces systemic absorption, thereby resulting in lower peak blood levels. Occasionally, the apprehension, tachycardia, or palpitations induced by epinephrine can be incorrectly interpreted by both the clinician and patient as an "allergic" reaction.

Specific Drugs

The more potent agents are more toxic on a milligram-to-milligram basis. Because anesthetics are used in equipotent doses (e.g., 1 mg of bupivacaine vs. 4 mg of lidocaine), they are approximately equitoxic. The blood levels achieved by a particular agent depend on the agent's absorption, distribution, and clearance from the circulation. Agents with high lipid solubility and lower protein binding (etidocaine > bupivacaine > lidocaine > mepivacaine) tend to become sequestered in tissue and have slower absorption and lower blood levels. Agents with a greater volume of distribution or faster clearance (etidocaine > lidocaine > mepivacaine > bupivacaine) also produce lower blood levels. Together, these effects produce margins of safety for each anesthetic, with etidocaine having the greatest safety margin, followed by bupivacaine, which is equal to or better than lidocaine.

Esters are difficult to measure in blood because of their rapid hydrolysis by pseudocholinesterase. As a group, toxicity have been attributed to "allergies" or to systemic toxicity to local anesthetics. Patients may also demonstrate systemic reactions to hidden allergens that may mimic a systemic reaction, such as anaphylactic reactions to the latex in surgical gloves.

High Blood Levels

Systemic toxic reactions result from high blood levels of local anesthetic. Several factors are important in producing high blood levels, including the site and mode of administration, rate of administration, dose and concentration, addition of epinephrine, specific drug, clearance, maximum safe dosage, and inadvertent intravascular injection.

Site and Mode of Administration

In comparing the routes of administration for a given dose, the intravascular route produces the highest levels, followed by topical mucosal application and then infiltration. The more vascular the site, the more systemic absorption that occurs and the higher the level obtained. The following blocks are arranged in decreasing order of systemic absorption: intercostal, caudal, epidural, brachial plexus, and subcutaneous. It follows that the site of administration is an important variable in determining the safe dose of an anesthetic. For example, 400 mg of lidocaine may produce a nontoxic blood level with subcutaneous abdominal wall infiltration but produce a toxic level when used for an intercostal nerve block.

Rate of Administration

A more rapid IV injection will produce a higher blood level than a slower injection. A single topical application leads to a higher level than a dose that is fractionated over time.

---

### Table 29.8 Differentiating Systemic Adverse Reactions

<table>
<thead>
<tr>
<th>FINDINGS</th>
<th>TOXIC REACTIONS</th>
<th>ALLERGY</th>
<th>VASOVAGAL</th>
<th>EXCESS CATECHOLAMINES, ANXIETY*, (ENDOGENOUS), VASOCONSTRICTOR (EXOGENOUS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Relatively specific signs and symptoms</td>
<td>Metallic taste</td>
<td>Acute rhinitis</td>
<td>Syncope&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>Headache</td>
</tr>
<tr>
<td></td>
<td>Tongue numbness</td>
<td>Pruritus&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td>Hypertension&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Drowsiness</td>
<td>Dermatitis</td>
<td></td>
<td>Palpitations</td>
</tr>
<tr>
<td></td>
<td>Nystagmus</td>
<td>Urticaria&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td>Apprehension&lt;sup&gt;b,c&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Slurred speech</td>
<td>Facial swelling</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Seizures&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Laryngospasm</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Coma</td>
<td>Bronchospasmb</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Respiratory arrest&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Overlapping signs and symptoms</td>
<td>Paresthesia</td>
<td>Light-headedness</td>
<td>Light-headedness</td>
<td>Paresthesia&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Light-headedness</td>
<td>Tachycardia&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Tachycardia</td>
<td>Light-headedness&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Tinnitus</td>
<td>Hypotension&lt;sup&gt;b&lt;/sup&gt;</td>
<td>(early)</td>
<td>Tinnitus&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Tremor</td>
<td>Cardiac arrest</td>
<td>Bradycardia&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Tremor</td>
</tr>
<tr>
<td></td>
<td>Tachypnea</td>
<td>Nausea and vomiting</td>
<td>Hypotension&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Tachypnea&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Tachycardia (early)</td>
<td>Dyspnea</td>
<td>Diaphoresis</td>
<td>Tachycardia&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Bradycardia&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td>Nausea and vomiting</td>
</tr>
<tr>
<td></td>
<td>Hypotension&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
<td>Dyspnea</td>
</tr>
<tr>
<td></td>
<td>Cardiac arrest</td>
<td></td>
<td></td>
<td>Diaphoresis</td>
</tr>
</tbody>
</table>

<sup>a</sup>Anxiety reaction, including hyperventilation syndrome.

<sup>b</sup>Denotes common and significant reactions.

<sup>c</sup>Vasovagal syncope occurs with the patient upright; any loss of consciousness in the recumbent position implies a severe toxic or anaphylactic reaction.

<sup>d</sup>Although apprehension is classically associated with anxiety and vasoconstrictor reactions, milder toxic and allergic reactions may cause patient apprehension.
is inversely proportional to the rate of hydrolysis. Tetracaine is slowly hydrolyzed and is most toxic. Chloroprocaine is quickly hydrolyzed and is the least toxic. Procaine falls between the two.

Clearance

The liver metabolizes amides, with the clearance rate being a function of hepatic blood flow and the extraction capacity of the liver. Decreased hepatic blood flow, produced by norepinephrine, propranolol, or general anesthesia, slows clearance and potentially raises drug blood levels. Decreased drug extraction, associated with congestive heart failure, cirrhosis, or hypothermia, may produce a higher blood level. Hypovolemia, which decreases hepatic flow, does not raise blood levels because it causes an offsetting decrease in absorption.

Decreased clearance of esters and an increased risk for toxicity occur in patients with either low levels or an atypical form of pseudocholinesterase. Low levels occur in various disease states, including severe liver disease and renal failure, and in pregnancy. Atypical pseudocholinesterase is an inherited trait, and its presence reduces the hydrolysis rate of procaine to a greater extent than low levels do.

There are significant differences between pediatric and adult drug distribution and metabolism. Neonates exhibit both reduced levels of pseudocholinesterase and reduced hepatic metabolism, thus increasing the risk for toxicity. In older children, the effects of increased hepatic metabolism and a relatively larger volume of distribution increase their tolerance for higher doses.

Because lidocaine is metabolized in the liver by cytochrome P-450 enzymes, drugs that inhibit these enzymes may slow lidocaine clearance and increase the risk for lidocaine toxicity. Although the effect of ciprofloxacin and erythromycin on infiltrated lidocaine has not been studied, these drugs decrease the metabolism of lidocaine and increase the concentration of its major metabolites when lidocaine is injected intravenously.\textsuperscript{124-126}

The clinical effect of this previously discussed phenomenon is unknown and probably of little consequence in ED wound care.

Maximum Safe Dosage

The maximum safe dose of a drug may be defined as the dose that produces a blood level of the drug just below the toxic level (Table 29.9). One maximum dose of an anesthetic agent appropriate for all patients and all conditions cannot be stated. A maximum safe dose cannot be based solely on the weight of a patient. In an adult, peak blood levels do not correlate well with weight because the volume of drug distribution is relatively constant.\textsuperscript{127,128} As an approximation, Arthur and McNicol\textsuperscript{129} recommended that maximum dosages for children be based on weight. Plain lidocaine may be used in doses of up to 4.5 mg/kg, and the addition of epinephrine allows a maximum dose of 7 mg/kg. Bupivacaine is not recommended for children younger than 12 years, although it is commonly used without adverse consequences. Furthermore, the dose should be modified according to the site and mode of administration.

Maximum safe doses as stated on package inserts should be used only as guidelines because most of them are derived from animal experiments and are based on absorption data only. Levels vary with the site of administration, use of a vasoconstrictor, and to some extent, the health of the patient. Levels can often be exceeded safely when the drug is admin-

<table>
<thead>
<tr>
<th>TABLE 29.9 Calculation of Anesthetic Doses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anesthetic solutions are marketed with the drug concentration expressed as a percentage (e.g., bupivacaine 0.25%, lidocaine 1%). To ascertain the strength of a solution in milligrams per milliliter, consider the following:</td>
</tr>
<tr>
<td>A 1% solution is prepared by dissolving 1 g of drug in 100 mL of solution.</td>
</tr>
<tr>
<td>Therefore 1 g/100 mL = 1000 mg/100 mL = 10 mg/mL.</td>
</tr>
<tr>
<td>Note: To calculate the strength from the percentage quickly, simply move the decimal point one place to the right, as follows:</td>
</tr>
<tr>
<td>0.25% = 2.5 mg/mL e.g., bupivacaine</td>
</tr>
<tr>
<td>0.5% = 5 mg/mL e.g., tetracaine</td>
</tr>
<tr>
<td>1% = 10 mg/mL e.g., lidocaine</td>
</tr>
<tr>
<td>2% = 20 mg/mL e.g., viscous lidocaine</td>
</tr>
<tr>
<td>4% = 40 mg/mL e.g., cocaine</td>
</tr>
<tr>
<td>5% = 50 mg/mL e.g., lidocaine ointment</td>
</tr>
<tr>
<td>20% = 200 mg/mL e.g., benzocaine</td>
</tr>
</tbody>
</table>

When combined in an anesthetic solution, epinephrine is usually in a 1:100,000 or a 1:200,000 dilution. For example:

1:100,000 concentration of epinephrine = 1 g/100,000 mL = 1000 mg/100,000 mL =
1 mg /100 mL or 0.01 mg/mL.
1 mL of 1:1000 epinephrine = 1 mg.
0.1 mL of 1:1000 epinephrine in 10 mL of anesthetic solution = 1:100,000 dilution = 0.01 mg/mL.
0.1 mL of 1:1000 epinephrine in 20 mL of anesthetic solution = 1:200,000 dilution = 0.005 mg/mL.

Some examples detailing epinephrine content:

<table>
<thead>
<tr>
<th></th>
<th>1:100,000</th>
<th>1:200,000</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 mL</td>
<td>0.050 mg</td>
<td>0.025 mg</td>
</tr>
<tr>
<td>10 mL</td>
<td>0.100 mg</td>
<td>0.050 mg</td>
</tr>
<tr>
<td>20 mL</td>
<td>0.200 mg</td>
<td>0.100 mg</td>
</tr>
</tbody>
</table>

Therefore 50 mL of 1% lidocaine with epinephrine 1:200,000 contains 500 mg of lidocaine and 0.25 mg of epinephrine.

Inadvertent Intravascular Injection

Most toxic reactions are caused by inadvertent intravascular injection of anesthetics whose doses were calculated for their intended extravascular sites. For example, lidocaine, 300 mg, is a safely infiltrated dose that would probably cause toxicity if directly injected into the bloodstream.

Anesthetics that are injected intravascularly must pass through the lungs before they reach other organs. Lung tissue sequesters a significant amount of drug, which lowers the arterial

blood concentration. Anesthetics that bypass the lungs, in cases of inadvertent injection into the carotid or vertebral arteries or in patients with intracardiac right-to-left shunts, can produce CNS toxicity at low doses. Intravenous injections into subcutaneous end-arteries about the head or neck are capable of retrograde flow into the cerebral circulation if the injection pressure exceeds arterial pressure. Because the blood volume in the brain is only approximately 30 mL at any given moment, even 1 mg of lidocaine injected into the carotid artery can theoretically produce toxic concentrations. Patients with low cardiac output or hypovolemia and preferential cerebral blood flow may suffer enhanced CNS toxicity.

**Host Factors**

Four factors tend to lower the body’s systemic tolerance to local anesthetic agents: hypoxia, acid-base status, protein binding, and concomitant drug use.

**Hypoxia**

It was initially thought that an overdose of a local anesthetic produces CNS stimulation and subsequent intracellular hypoxia, which then became the key precipitant for all toxic manifestations of the drug. It is now known that hypoxia may enhance anesthetic toxicity but is not the primary factor.

**Acid-Base Status**

Although studies of metabolic alkalosis have produced conflicting results, acidosis, particularly respiratory acidosis, can increase toxicity. The elevated CO₂ produced by respiratory acidosis crosses the blood-brain barrier, where it may act directly on the receptor and indirectly by lowering intracellular pH. The lower pH causes more of the drug to ionize, thereby furthering the block in the sodium channel and increasing the potential for toxicity.

**Protein Binding**

The concentration of unbound drug relates more closely to toxic effects than does the total drug concentration (bound plus unbound) as measured in the blood. The amount of α-acid glycoprotein (AAG), the major plasma protein responsible for binding local anesthetics, is considerably decreased in neonates in comparison to adults. Arthur and McNicol⁹ implied that the low AAG levels in neonates are responsible for the increased toxic potential. Tucker and colleagues¹⁰ listed several disease states that alter AAG levels and protein binding but questioned whether they lead to changes in free drug concentration in vivo.

**Concomitant Drugs.** For years, barbiturates were used to prevent and treat local anesthesia–induced seizures. Barbiturates were found to worsen anesthetic-induced apnea and cardiovascular depression. CNS depressants are used with caution when concern exists for local anesthetic toxicity. CNS stimulants have been shown to increase anesthetic-induced excitability and are avoided. Mixtures of local anesthetics have an additive effect on toxicity. If two drugs are used at half strength, they produce the same degree of toxicity as though each were used alone at normal strength. As discussed previously, drugs that slow metabolism by inhibiting hepatic enzymes may increase the risk for toxicity.

**Recognition of CNS Toxicity**

The earliest manifestation of systemic toxicity is CNS stimulation resulting from blockade of inhibitory synapses. CNS depression follows and is produced by direct depression of the medulla, although hypoxia may play a role. The signs and symptoms are dose related. Potential signs and symptoms of CNS toxicity, in progressing order, are numbness of the tongue, light-headedness, tinnitus, visual disturbances, muscle twitching, convulsions, coma, and apnea. Drowsiness, commonly seen with lower doses of lidocaine, is not associated with bupivacaine or etidocaine. Tetracaine may produce apnea or cardiovascular toxicity without CNS manifestations.

**Recognition of Cardiovascular Toxicity**

Moderate blood concentrations of local anesthetics produce slight increases in cardiac output, heart rate, and arterial pressure because of the effects of direct peripheral vasodilation and CNS stimulation. At concentrations generally well above CNS toxicity levels, local anesthetics cause direct myocardial depression, hypotension, and bradycardia, perhaps leading to cardiovascular collapse. These agents also slow electrical conduction and lead to reentry phenomena and various supraventricular and potentially lethal ventricular dysrhythmias, especially with bupivacaine and etidocaine.

**Prevention of Toxicity**

Knowledge of factors contributing to toxicity guides preventive measures. Avoid esters in patients with an atypical form or a quantitative deficiency of pseudocholinesterase. Use amides with caution in patients with severe liver disease or congestive heart failure. Pay attention to maximum safe dosages based on the site, technique, use of epinephrine, and patient status. Add epinephrine when possible to decrease the rate of drug absorption at vascular sites. Reduce the drug concentration by saline dilution to increase the volume for administration when a large area must be infiltrated. Frequently aspirate in areas of high vascularity, even though a negative aspiration may not prevent IV administration.¹¹ Slow infiltration is safer and less painful.

**Treatment of Systemic Toxicity**

Local anesthetics should not be administered without the ability to recognize and treat a toxic reaction, including having all necessary equipment and drugs readily available and being knowledgeable in their use. Despite taking all possible precautions, toxic reactions still occur, and close observation of the patient allows early detection and treatment.

Providing proper oxygenation and ventilation at the earliest sign of a reaction is the cornerstone of treatment. Encourage patients who are alert to moderately hyperventilate to lower the pressure of carbon dioxide (PCO₂) and raise the seizure threshold. Intubation with high-flow oxygen and hyperventilation is performed for patients who cannot adequately ventilate. Initiate IV access and monitor vital signs and cardiac rhythm closely.

Seizures are generally self-limited but are treated if they persist or prevent adequate ventilation. Because respiratory depression secondary to toxicity may follow, low-dose lorazepam or an ultrashort-acting barbiturate (thiopental or sodium methohexitol) is preferred. Intubate the patient to ensure an effective airway and prevent further lactic acidosis if the seizures persist. If toxicity is caused by an ester, especially if there is an associated pseudocholinesterase problem, succinylcholine will compete with the anesthetic for the pseudocholinesterase and may increase the toxicity of both compounds.
Treat hypotension and bradycardia with fluids, leg elevation, α- and β-agonists (epinephrine, ephedrine, or dopamine), or atropine as the need dictates. Although lidocaine (with diazepam pretreatment) has been shown to be effective for bupivacaine-induced ventricular dysrhythmias, strong theoretical and experimental evidence indicates that bretylum is more effective. However, until bretylum becomes available again, amiodarone is a reasonable alternative. High doses of atropine and epinephrine can be successful in correcting pulseless idioventricular rhythm. Cardiopulmonary resuscitation is institued when necessary.

Intravenous Lipid Emulsion
Animal studies, case reports/small series, and personal opinion have advocated the use of 20% lipid emulsion intravenously as a remarkable antidote to resuscitate bupivacaine- and mepivacaine-related cardiac arrest, a situation that is usually fatal (see https://lipidrescue.com). Rosenblatt and associates described the IV injection of 100 mL of 20% Intralipid (Baxter Healthcare Corp., Deerfield, IL; formulation used for hyperalimentation) followed by an infusion (0.5 mL/kg per min over a 2-hour period) and related this intervention to successful resuscitation in a scenario that appeared hopeless. Picard considers lipid emulsion a “crucial antidote” that should be available when local anesthetics are used for peripheral nerve blocks. A large collaborative workgroup of leading toxicologists recommends Intralipid (Baxter) infusion during cardiac arrest associated with bupivacaine but strikes a neutral posture for cardiogenic arrest due to other local anesthetics. Lipid emulsion therapy is typically administered intravenously but bupivacaine-induced toxicity is reversed in rats when lipid emulsion is administered by the intraosseous route. Intraosseous infusion may be a reasonable alternative to IV infusion when IV access is not available, though this route of administration in humans is of unproven benefit. Lipid emulsion therapy in otherwise hopeless situations of cardiac arrest secondary to local anesthetic overdose is supported (Box 29.3). It appears prudent and intuitive to initiate this antidote before cardiac arrest when significant local anesthetic toxicity is identified.

Allergic Reactions

Allergic Agents

True allergic reactions are rare and account for only 1% to 2% of all adverse reactions, but they are important to recognize because of their serious potential. Ester solutions (procaine, tetracaine) that produce the metabolite para-aminobenzoic acid (PABA) account for the majority of these reactions. Amide solutions (lidocaine, bupivacaine) are rarely involved, and usually the preservative MPB, which is structurally similar to PABA, is responsible. Although pure esters and pure amides do not cross-react, amides may appear to do so if multidose vials containing MPB are used. Patients may manifest an allergic response on first contact with a local anesthetic because of previous sensitization to these agents. MPB is found in creams, ointments, and various cosmetics, and PABA is an ingredient in many sunscreen preparations. Patients who are latex sensitive may have an allergic reaction incorrectly attributed to the local anesthetic.

Cell-mediated delayed reactions manifesting as dermatitis are rare; it is immediate hypersensitivity that most concerns the emergency clinician. A spectrum of signs and symptoms may occur, from rhinitis and mild urticaria to bronchospasm, upper airway edema, or anaphylactic shock. Onset may be immediate and occur even during administration of the agent. Treat anaphylaxis in the usual manner.

The more frequent problem facing emergency clinicians is a patient who claims to have a past history of allergy to local anesthetics. Most patients assume that any adverse reaction to a local anesthetic procedure is an allergy. Because allergy is rarely the case, a careful history and a review of previous records, if available, are crucial in evaluating these patients. Procaine, trade name Novocaine (Hospira, Inc, Lake Forest, IL), was commonly used in dentistry, and many patients who experienced many types of reactions in the dentist's office, rarely true allergy, state they are allergic to Novocaine. Procaine is no longer used in dental practice, but it is the local anesthetic in intramuscular penicillin preparations (procaine penicillin G).

Attempts to uncover the actual cause of the past reaction and the specific agent involved are often fruitless. Ask about the exact signs and symptoms, technique of administration, amount of drug used, and how the patient was treated. If an allergic reaction cannot be ruled out and the drug previously used is known, use an agent from the other class (whether amide or ester). Lidocaine from a dental cartridge does not contain MPB, and if this were the allergenic source, an ester agent could be used. However, if lidocaine from a multidose vial is implicated, do not use an ester because MPB may cross-react with PABA. In this case, it may be safer to use an amide without MPB or to choose an alternative (see later). In most cases the allergen is an ester, and the patient can safely be given an amide without MPB. Single-dose ampules of 1% lidocaine without MPB, readily obtainable from a resuscitation cart, can be used for this purpose.

Uncertainty often exists regarding the specific agent involved, and the clinician must choose an alternative approach to local anesthesia. If the wounds are extensive and the risk is acceptable, procedural sedation (see Chapter 33) or general anesthesia may be used. Conversely, if minimal pain is expected and the procedure is short (e.g., one or two sutures or staples in the scalp), no anesthesia may be required. These methods may be useful, but the degree of anesthesia produced is frequently not sufficient. Antihistamines injected into a wound have been used successfully for many years and represent a good alternative. Local anesthetic efficacy is found in varying degrees with all

---

**BOX 29.3 Preliminary Strategies for Lipid Emulsion Rescue Therapy in Patients With Severe Local Anesthetic Toxicity**

Intravenous lipid emulsion (ILE) is available as Liposyn II (Hospira, Inc., Lake Forest, IL) (20%) or Intralipid (Baxter Healthcare Corp., Deerfield, IL) (20%).

Protocol: Infuse 20% ILE intravenously. Administer a bolus injection of 1.5 mL/kg* over a period of 1 to 2 minutes. It may be repeated 1 to 3 times every 5 minutes. Follow with a continuous infusion at 0.25 mL/kg per min for 30 to 120 minutes.

Increase the rate of infusion to 0.5 mL/kg per min in patients with declining blood pressure.

*Can be extrapolated to other tissues. Continue other resuscitative efforts, including cardiopulmonary resuscitation.

*For the initial dose, withdraw 100 mL from a 500-mL bag/bottle and inject with a syringe. Then attach to an infusion pump. From Brent J: Poisoned patients are different—sometimes fat is a good thing, Crit Care Med 37:1157, 2009; and https://lipidrescue.com.
antihistamines. Ketamine anesthesia may be a useful alternative in some situations and is commonly used in children.

**Diphenhydramine and Benzyl Alcohol**

Several studies have demonstrated that 1% diphenhydramine (Benadryl, Johnson & Johnson, New Brunswick, NJ) is as effective as 1% lidocaine for infiltrative anesthesia.\(^{138-140}\) As long as diphenhydramine is not used at concentrations greater than 1%, potential problems of skin necrosis or significant sedation are rare. Dilute the standard 5% parenteral form to a 1% concentration for subcutaneous injection (1 mL of drug to 4 mL of saline). The duration of action of diphenhydramine is shorter than that of lidocaine but appears to be adequate for most procedures. The injection pain of diphenhydramine exceeds that of lidocaine but can be diminished by reducing the concentration to 0.5%. At this lower concentration, the effectiveness of this agent on facial wounds is lost.\(^{131}\) The addition of epinephrine to 0.5% diphenhydramine results in a more painful solution with a shorter duration of action than a standard buffered lidocaine with epinephrine solution.\(^{132}\) Benzyl alcohol (0.9%) with epinephrine (1:100,000) compares favorably with diphenhydramine as an effective local anesthetic. This appears to be a useful alternative to diphenhydramine when lidocaine cannot be used but is of shorter duration than diphenhydramine.\(^{143-145}\)

**Skin Testing**

Skin testing and progressive subcutaneous challenge doses deserve special mention because they appear to be logical and well-studied approaches. However, intradermal skin testing with local anesthetics is controversial and often of no practical benefit in the ED. False-positive results are frequently produced by local release of histamine in response to needle trauma, tissue distention, or preservatives in the solution.\(^{146}\) In addition, a high incidence of false-negative results can occur. It is questionable whether these low-molecular-weight drugs or their allergenic metabolites are ever capable of eliciting positive responses.\(^{147}\) Other disadvantages of skin testing include its time-consuming nature and potential hazard when even minute traces of an allergen may precipitate a serious reaction. Subcutaneous challenge testing in graduating doses has been advocated and may well eliminate many false responses, but it does not eliminate the problems of time and hazard. Swanson,\(^{148}\) recognizing that allergy to pure lidocaine is extremely rare, recommended 0.1 mL as a single intradermal skin test. Although his approach eliminates the time disadvantage, intradermal placement can still produce false responses. It would seem more reasonable to give this test dose subcutaneously while exercising due caution in the unlikely event that a patient exhibits a serious reaction.

**Summary of Anesthetic “Allergy” Management**

Generally speaking, the optimal approach to a patient with a presumed anesthetic allergy is to determine the specific anesthetic agent associated with a presumed allergic reaction and then use a preservative-free agent from the other class (see earlier discussion). If the agent is unknown, use an antihistamine or give 0.1 mL of preservative-free lidocaine as a subcutaneous test dose and proceed with the full dose if no reaction occurs within 30 minutes. Given the studies mentioned earlier, prudent choices would seem to be diphenhydramine (Benadryl, Johnson & Johnson) or benzyl alcohol. Epinephrine (1:100,000) can be added to both these drugs to prolong the duration of action. Ketamine anesthesia is also an alternative.

---

**Catecholamine Reactions**

Anxiety and vasoconstrictor (epinephrine) reactions are discussed together because each produces similar manifestations caused by elevated catecholamine levels. These relatively common disorders are difficult to distinguish from each other and are not generally serious.

Excess catecholamine levels produce tachycardia, palpitations, hypertension, apprehension, tremulousness, diaphoresis, tachypnea, pallor, and on occasion, anginal chest pain. Thus, catecholamine excess may resemble the CNS stimulation phase of local anesthetic toxicity.

Catecholamine reactions are not usually caused solely by exogenous epinephrine because if it is used in its optimal concentration (1:200,000), the maximum safe dose (0.25 mg) is rarely exceeded. However, many patients produce significant endogenous catecholamines because of anxiety about the anesthetic approach or upcoming procedure. In this case, even the addition of small amounts of epinephrine could trigger a catecholamine reaction. Therefore patient preparation includes proper explanation and reassurance to decrease anxiety. Exercise caution with patients who have hyperthyroidism, hypertension, or atherosclerotic cardiovascular disease, although these conditions do not contraindicate the judicious use of epinephrine-containing anesthetics. Do not give epinephrine-containing anesthetics to patients taking MAO inhibitors.

Treatment of a catecholamine reaction includes stopping further drug administration, observing the patient closely, and administering α- or β-antagonists or benzodiazepine agents, if necessary, to combat severe reactions.

**Vasovagal Reactions**

It is not standard to monitor patients (cardiac, pulse oximetry) during routine local anesthesia procedures. Vasovagal reactions are, however, common, especially in dental procedures (reported incidence, 2% to 3%), during which the patient is generally in an upright position. To limit vasovagal reactions related to local anesthesia in the ED, do not draw up medication in a syringe in front of the patient, and inject only when the patient is supine (Figs. 29.9 and 29.10). The patient initially experiences anxiety when a triggering event, commonly the
sight or sensation of needle insertion, causes loss of sympathetic tone and an increase in vagal tone. The resultant hypotension and bradycardia may lead to syncope. Address the patient’s anxiety and administer the injections with the patient recumbent as useful preventive measures. Cardiac monitoring may help identify the onset of vagally induced bradycardia when suggested by the past history. Lay the patient supine and elevate the legs. Rarely, atropine is required. Should a patient lose consciousness while in a recumbent position, consider diagnoses other than vasovagal syncope, although significant bradycardia and even complete heart block may accompany a vagal reaction in a supine patient.

**SUMMARY**

Emergency medicine cannot be practiced without the use of local anesthetic agents. Their effectiveness when applied topically or by infiltration makes them extremely adaptable to many clinical circumstances. A working knowledge of commonly used agents is necessary to ensure the safe administration of these medications. Aim specific efforts at maximizing the drugs’ anesthetic effects while minimizing the pain of administration and risk for toxicity.

**Figure 29.10** A, What’s wrong in this picture? Almost everything! The patient is sitting upright and directly observing the procedure. This may lead to a vasovagal event, and should he fall from the bed, significant injury may occur. Additional problems include improper positioning of the overhead light (behind the physician’s back), lack of privacy (open curtain and door), equipment tray placed too far away, and a hunched-over posture of the physician. B, A much better approach. Note that the patient is supine, the bed is raised, the light and equipment tray are in a better position, the curtain is closed, and a family member is present (but sitting to avoid a “bystander vagal event”) to distract the patient during the procedure. The clinician could also be seated for comfort.
544.e2

SECTION

V   Anesthetic and Analgesic Techniques

56. Bonadio WA, Wagner V: Efficacy of tetracaine-adrenaline-cocaine topical
anesthetic without tetracaine for facial laceration repair in children.
57. Bonadio WA, Wagner V: Adrenaline-cocaine gel topical anesthetic for
58. Schaffer D: Clinical comparison of TAC anesthetic solutions with and
59. Smith SM, Barry RC: A comparison of three forms of TAC (tetracaine,
Adrenalin, cocaine) for anesthesia of minor lacerations in children. Pediatr
60. Ernst AA, Marvez-Valls E, Nick TG, et al: LAT (lidocaine-adrenalinetetracaine) versus TAC (tetracaine-adrenaline-cocaine) for topical
1995.
61. Zempsky WT, Karasic RB: EMLA versus TAC for topical anesthesia of
(Adrenalin) and cocaine (TAC) versus lidocaine, epinephrine and tetracaine
(LET) for anesthesia of lacerations in children. Ann Emerg Med 25:
lacerations: a randomized trial of lidocaine-epinephrine-tetracaine solution
64. Singer AJ, Stark MJ: LET versus EMLA for pretreating lacerations: a
lidocaine hydrochloride gauzes in acute traumatic wounds in triage, a
66.	Gaufberg S, Walta M, Workman T: Expanding the use of topical anesthesia
67. Altieri M, Bogema S, Schwartz RH: TAC topical anesthesia produces
levels following application of topical anesthesia in children. Ann Emerg
70. Bartfield JM, Homer PJ, Ford DT, et al: Buffered lidocaine as a
and epinephrine used for local anesthesia. J Dermatol Surg Oncol 17:
as a local anesthetic for simple laceration repair. Ann Emerg Med 19:
versus buffered lidocaine in local wound infiltration. J Emerg Med 10:
74. Bartfield JM, Ford DT, Homer PJ: Buffered versus plain lidocaine for
75. Peterfreund RA, Datta S, Ostheimer GW: pH Adjustment of local
anesthetic solutions with sodium bicarbonate: laboratory evaluation of
76. Cheney PR, Molzen G, Tandberg D: The effect of pH buffering on
reducing the pain associated with subcutaneous infiltration of bupivacaine.
77. Bainbridge LC: Comparison of room temperature and body temperature
78. Waldbillig DK, Quinn JV, Stiell IG, et al: Randomized double-blind
controlled trial comparing room temperature and heated lidocaine for
and buffering on pain of infiltration of lidocaine. Acad Emerg Med 2:
80. Brogan GX, Jr, Giarrusso E, Hollander JE, et al: Comparison of plain,
warmed, and buffered lidocaine for anesthesia of traumatic wounds. Ann
81. Martin S, Jones JS, Wynn BN: Does warming local anesthetic reduce
82. Leff DR, Nortley M, Dang V, et al: The effect of local cooling on pain
perception during infiltration of local anaesthetic agents, a prospective
83. Hamelin ND, St. Amand H, Lalonde DH, et al: Decreasing the pain
of finger block injection: level II evidence. Hand (N Y) 8:67–70,
2013.

84. Kelly AM, Cohen M, Richards D: Minimizing the pain of local infiltration
anesthesia for wounds by injection into the wound edges. J Emerg Med
85. Bartfiled J, Sokaris S, Raccio-Robak N: Local anesthesia for laceration:
pain of infiltration inside vs outside the wound. Acad Emerg Med 5:100–104,
1998.
86. Bierman SF: Painless wound injection through use of a two-finger
bevel position on pain for subcutaneous lidocaine injection. J Perianesth
89. Cobb AG, Houghton GR: Local anaesthetic infiltration versus Bier’s
90. Newman AP: Meniscal and ligamentous injuries of the knee. Top Emerg
the knee after intra-articular anesthesia. Am J Sports Med 11:258–261,
1983.
92. Holdsworth BJ, Clement DA, Rothwell PN: Fractures of the radial
head—the benefit of aspiration: a prospective controlled study. Injury
intraarticular bupivacaine infusion: an experimental model investigating
epinephrine, and preservatives found in local anesthetics containing
peripheral analgesic effects of intraarticular morphine. Anesth Analg
96. Karmakar MK, Critchley LAH, Ho AMH, et al: Continuous thoracic
paravertebral infusion of bupivacaine for pain management in patients
blunt trauma patients with unilateral rib fractures: a randomized trial.
98. Knottenbelt JD, James MF, Bloomfield M: Intrapleural bupivacaine
analgesia in chest trauma: a randomized double-blind controlled trial.
with multiple rib fractures: comparison with epidural block. J Emerg
100. Pond WW, Somerville GM, Thong SH, et al: Traumatic splenic rupture
101. Stromskag KE, Minor B, Steen PA: Side effects and complications related
1990.
102. Engdahl O, Boe J, Sandstedt S: Plasma concentrations and hemodynamic
changes after repeated interpleural injections of bupivacaine-epinephrine.
and suppresses mitochondrial functions of human neutrophil in vitro.
fibroblasts to local anaesthetics impairs cell growth. Clin Exp Immunol
108. Schmidt RM, Rosenkranz HS: Antimicrobial activity of local anesthetics:
109. Feldman JM, Chapin-Robertson K, Turner J: Do agents used for epidural
110. Parr AM, Zutman DE, Davidson JS: Antimicrobial activity of lidocaine
against bacteria associated with nosocomial wound infection. Ann Plast
and lidocaine in wound tissue culture biopsies for culturing. Wound Repair
112. Thompson KD, Welkyj S, Massa MC: Antibacterial activity of lidocaine
in combination with a bicarbonate buffer. J Dermatol Surg Oncol 19:


Emergency physicians encounter patients with dental pain, facial lacerations, and other injuries of the head on a regular basis. The assessment of these injuries can be both time consuming and painful. Thankfully, regional anesthesia for these conditions is easy to instill, reliable, and safe. Techniques such as nerve blocks can be used for patient comfort during repair of lacerations of the face, ear, forehead, and, particularly, the lips. A block avoids direct infiltration into the actual laceration, which preserves the anatomy in these delicate areas and thus allows better cosmetic repair.

Nerve blocks can also be used to give patients with dental pain almost immediate relief. They can be used to treat dentalgia caused by infection, fracture, or dry socket. They decrease the need for narcotic drugs. The ability of an emergency physician to master these blocks leads to fast and efficient patient care as well as improved patient satisfaction (Table 30.1).

ANATOMY OF THE TRIGEMINAL NERVE (CRANIAL NERVE V)

Sensation to the face and head region is supplied primarily by cranial nerve V (CN V), which is also known as the trigeminal nerve, due to its three large branches: V1, the ophthalmic nerve; V2, the maxillary nerve; and V3, the mandibular nerve (Fig. 30.1A). This is the largest of the 12 cranial nerves, originating at the upper portion of the pons. CN V contains motor neural fibers that control the muscles of mastication. This chapter focuses on its sensory function (see Fig. 30.1B).

The ophthalmic nerve, V1, courses through the cavernous sinus and exits through the supraorbital fissure. It then gives off three branches: the frontal, lacrimal, and nasociliary (see Fig. 30.1C). These nerves and their branches innervate the eye, orbit, forehead, and portions of the nose.1

The maxillary nerve, V2, exits through the foramen rotundum and gives off several branches, including the infraorbital nerve and the posterior, middle, and anterior superior alveolar nerves (see Fig. 30.1C). These nerves innervate the face, lip, maxillary teeth, and mucosa.1

The mandibular nerve, V3, exits through the foramen ovale into the infratemporal fossa. It then splits into many branches that supply the dura and muscles of mastication (motor function), as well as the buccal branch and the auriculotemporal, lingual, and inferior alveolar branches (see Fig. 30.1C). These branches supply sensation to the skin on the side of the head, the auricle of the ear, the tongue, the mucosa and skin of the cheek, the mandibular teeth, and the lower lip.1

EQUIPMENT FOR FACIAL NERVE BLOCKS

The equipment needed to perform facial nerve blocks is very minimal, and should be available in any emergency department (ED) (Review Box 30.1). Medications include common anesthetics such as lidocaine or bupivacaine, with or without epinephrine. The practitioner should always be aware of the maximum doses of anesthetic agents being utilized, although most blocks will require much less than the maximum dosing. Standard 25- to 27-gauge needles are used, with 3-ml syringes for anesthesia and dental aspirating syringes for aspiration of blood. Cotton pledgets and applicators are necessary for hemostasis.
27-gauge needles are recommended. Using a smaller needle can lead to false-negative aspiration in the case of an arterial puncture. Some of the equipment listed in Review Box 30.1 might not be necessary for all blocks, and some can be substituted. For instance, a tongue depressor is recommended for use as a retractor, but any other acceptable instrument or even a finger can be used for this purpose.

GENERAL RECOMMENDATIONS

Some general recommendations apply to nearly all types of blocks (Fig. 30.2). These procedures have very few contraindications. One of them involves infected tissue: the needle should never be inserted through infected tissue because this approach could result in inoculation of deep tissue with bacteria. In addition, patients with an allergy to the anesthetic agent should not undergo these procedures unless an alternative agent is available. Although not an absolute contraindication, coagulopathy might present a higher risk for hematoma and bleeding complications, so, as for any procedure, the risks should be considered and might outweigh the benefits.

When performing many of these blocks, inserting the needle with the bevel facing toward the bone allows the anesthetic to be injected as close to the nerve or bone as possible and increases the likelihood that the procedure will be successful. During and after injection, particularly when working in areas near the lip, slight exterior pressure and massage not only help the anesthetic diffuse to its target but also help prevent ballooning of the lip or facial tissue. Finally, it is generally recommended that the full length of the needle should not be inserted into the mucosa when performing an intraoral injection, so that it can be retrieved if it breaks.

Application of a topical anesthetic before insertion of the needle is not a necessary step prior to inducing any of the intraoral blocks, but it can increase patient comfort (Fig. 30.3). To apply a topical anesthetic, first dry the tissue with gauze and then apply a topical mucosal anesthetic, for instance, viscous lidocaine. Another technique is to soak a piece of gauze with the anesthetic and then place it over the dry mucosa. Spray anesthetics such as cetacaine are popular and effective alternatives to lidocaine-soaked gauze.

Supraperiosteal Injection

**Indication**

Supraperiosteal injection is generally used to achieve anesthesia of individual maxillary teeth, but it can be used for any tooth. This block works well if the anesthesia is needed for only one or two teeth, and it can be very helpful to alleviate pain associated with a simple toothache. As noted previously, care should be taken to avoid inserting the needle through any infected tissue.

**Anatomy**

The nerves for each individual tooth enter at the apex of that tooth and are protected by the bone supporting the tooth. Supraperiosteal injection is designed to anesthetize a single tooth (Fig. 30.4A).

**Approach**

Begin by applying a topical anesthetic, as previously described. Once the anesthetic has had time to take effect, retract the lip until the tissues are taut (down and out for maxillary teeth and up and out for mandibular teeth). Insert the needle at the mucobuccal fold, with the bevel facing the tooth. The needle needs to be inserted only a few millimeters (see Fig. 30.4B). Intraarterial injection is unlikely in this block, but you should aspirate prior to slowly injecting 1 or 2 mL of anesthetic. Apply slight pressure and massage as previously discussed. Because the anesthetic needs to penetrate the bone, a few minutes are needed for anesthesia to occur.

**Complications**

Complications with this injection are very rare. The risk of inoculating deeper tissues with bacteria is possible if the injection goes through infected tissue. Anesthesia can fail if the injection is too high, too low, or too far away from the nerve.

Posterior Superior Alveolar Nerve Block

**Indication**

The posterior superior alveolar nerve (PSAN) block can be used to provide anesthesia for the first through the third maxillary molars. However, it is likely that only partial anesthesia of the first molar will be achieved due to its innervation with accessory nerves, particularly at the mesiobuccal root of this molar.

### TABLE 30.1 Nerve Blocks and Indications

<table>
<thead>
<tr>
<th>NERVED BLOCKED OR AREA INJECTED</th>
<th>AREA ANESTHETIZED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supraperiosteal infiltration</td>
<td>Individual teeth</td>
</tr>
<tr>
<td>Posterior superior alveolar nerve</td>
<td>Second and third molars and partial anesthesia of first molar</td>
</tr>
<tr>
<td>Middle superior alveolar nerve</td>
<td>First and second premolars, as well as partial anesthesia of first molar</td>
</tr>
<tr>
<td>Anterior superior alveolar nerve</td>
<td>Central incisor to the canine</td>
</tr>
<tr>
<td>Infraorbital nerve</td>
<td>Midface region, skin of lateral nose, lower eyelid, and MSA and ASA regions</td>
</tr>
<tr>
<td>Inferior alveolar nerve</td>
<td>All ipsilateral mandibular teeth, lower lip, and chin</td>
</tr>
<tr>
<td>Mental nerve</td>
<td>Mucosa and skin of lower lip and chin</td>
</tr>
<tr>
<td>Scalp block</td>
<td>Skin of the scalp within the blocked area</td>
</tr>
<tr>
<td>Greater and lesser occipital nerve</td>
<td>Occipital region of the scalp</td>
</tr>
<tr>
<td>Ophthalmic nerve</td>
<td>Skin of the forehead and scalp (as far back as lambdoid suture), upper eyelid</td>
</tr>
</tbody>
</table>

ASA, Anterior superior alveolar; MSA, middle superior alveolar.
Figure 30.1 A, Distribution of the areas innervated by the three major branches of the trigeminal nerve. B, Cutaneous branches of the trigeminal nerve and their exit points from the skull. C, Branches of the trigeminal nerve. BR, Branch; n, nerve. (A, Borrowed from Henry Vandyke Carter-Henry Gray *Anatomy of the human body* 1918, Gray's anatomy, Plate 784. B and C, Adapted from Eriksson E, editor: *Illustrated handbook in local anesthesia*. Philadelphia, 1980, Saunders.)
**Head and Neck Regional Anesthesia: General Technique**

1. Confirm relevant local anatomy (in this example, the infraorbital nerve block is depicted).
2. Retract the lip and dry the mucous membranes with gauze or a cotton pledget.
3. Apply a topical anesthetic, such as Benzo-Jel, on a cotton-tipped applicator (shown here), or use Hurricaine spray (not depicted).
4. Insert the needle, aspirate the syringe to exclude intravascular placement, and then inject the appropriate amount of local anesthetic (usually about 2 mL).

**Figure 30.2** Head and neck regional anesthesia: general technique depicting the infraorbital nerve block.

**Anatomy**

The PSAN branches from the maxillary branch of the trigeminal nerve and travels inferiorly just before the maxillary nerve enters the infraorbital groove. The nerve descends along the posterior lateral portion of the maxillary tuberosity and gives off branches to the second, third, and partially the first maxillary molars (Fig. 30.5A).

**Approach**

Two techniques for creating a PSAN block have been well described in the literature. The traditional method begins similar to the other blocks, that is, by applying a topical anesthetic to the mucosa. The insertion point for the needle is just distal to the root of the second molar, at the height of the mucobuccal fold (see Fig. 30.5B). For this block, because of the posterior and medial location of the nerve complex, insert the needle in an upward, inward, and posterior direction (toward the maxillary tuberosity), approximately 45 degrees in each direction. Ultimately, the needle should be inserted approximately 15 mm and certainly no more than 25 mm. The operator should not feel any resistance while inserting the needle and should not encounter bone. If resistance or bone is felt, withdraw the needle nearly and then redirect it. Once the appropriate depth is reached, aspirate and, if negative, slowly inject 1 to 3 mL of anesthetic.

The second technique involves a curved 24-mm needle, to approach the posterior maxillary surface. Again, anesthetize the mucosa with a topical anesthetic. The insertion point for the needle is more posterior than the traditional approach, just distal to the third molar, at the corner of the posterior lateral portion of the maxilla and directed along the posterior maxilla. Insert the needle 10 to 12 mm from the initial insertion point along the posterior wall of the maxilla, and orient it just slightly medially. After aspirating, inject 1 to 3 mL of lidocaine.

**Figure 30.3**

A, Local anesthesia: basic setup for intraoral application using an aspirating dental syringe. B, Topical mucosal anesthesia can make the injection nearly painless. Swab the gauze-dried mucosa with the topical agent or have the patient hold cotton swabs soaked in the agent, and wait for 1 to 3 minutes.
Based on a literature review, Singla and Alexander recommended insertion to approximately 15 mm; however, this is lower than many recommendations and up to 25 mm, no deeper, may be required. The further the needle is inserted, the greater the risk of complications.2

Middle Superior Alveolar Nerve Block

Indication
The middle superior alveolar nerve (MSAN) block can be used to anesthetize the lateral upper lip and the first and second premolars simultaneously, as well as the mesiobuccal root of the first molar. This technique can be especially useful if the lateral lip is not anesthetized by an infraorbital nerve block.

Anatomy
The MSAN descends posteriorly to anteriorly along the lateral maxillary wall as an extension of the maxillary nerve. The landmark needed for this procedure is the junction between the first molar and the second premolar (Fig. 30.6). A subset...
of patients have an anatomic variant in which the MSAN does not exist.\(^1\)

**Approach**

Begin as you would for any of the other intraoral block procedures, by applying topical anesthetic. Retract the upper lip and pull the tissues taut. Direct the needle toward the intersection of the first molar and the second premolar at the mucobuccal fold, and insert it 5 to 10 mm at a 45-degree angle posteriorly, with the bevel facing the bone, in an effort to place as much anesthetic near the nerve as possible (see Fig. 30.6B). Slowly inject 1 to 3 mL of anesthetic while massaging the tissue.

**Complications**

Complications of this procedure are few: they are the same complications inherent to all procedures, such as bleeding. Because this is an area with low vascularity, hematoma formation and arterial aspiration are rare. For patients with the anatomic variation of absence of the MSAN, this block will obviously fail, so an infraorbital nerve block, an anterior superior alveolar nerve (ASAN) block, or a PSAN block will be necessary, depending on the region needing anesthesia.

### Anterior Superior Alveolar Nerve Block

**Indication**

Depending on the area of injury or pain, multiple areas of the maxilla might require anesthesia. If anesthesia from the central incisor to the canine is required, the ASAN can be blocked.

**Anatomy**

The ASAN is a distal branch from the infraorbital nerve. As the infraorbital nerve travels down the anterior maxilla, it gives off the ASAN, which travels to innervate the area from the central incisors to the canine of the ipsilateral side. The landmark needed is the apex of the canine (Fig. 30.7A).

**Technique**

Begin by applying a topical anesthetic, as for the blocks described previously. Then, at the height of the mucosal reflection toward the apex of the canine, insert the needle at a 45-degree angle, to a depth of 5 mm (see Fig. 30.7B). Inject 1 to 2 mL of anesthetic slowly.
Complications
Complications associated with this procedure are rare, unless the needle is inserted too far. The infraorbital nerve could be blocked, depending on the final depth of the needle. The block could be unsuccessful if the needle insertion is too superficial for the anesthetic to reach the nerve.

Anterior Middle Superior Alveolar Block

Indications
This approach is indicated when anesthesia is needed from the central incisor to the bicuspid. It can be used for greater palatine anesthesia as well as nasopalatine, anterior superior alveolar (ASA), and middle superior alveolar (MSA) nerve coverage, with the goal of avoiding multiple injections and avoiding anesthetizing the lip unnecessarily by incidental infiltration of anesthetic into the surrounding soft tissue.

This block uses the palatal nutrient pores to distribute anesthesia through the palate to the ASA and the MSA simultaneously. Small pores in the palate allow diffusion of the anesthetic after injection into the palate.

Approach
All the blocks described earlier in this chapter began on the buccal surfaces to anesthetize the maxillary teeth. The injection for the anterior middle superior alveolar block is placed through the hard palate at the intersection of an imaginary line dividing the premolars at a point halfway between the midpalatine crest and the free gingival margin (Fig. 30.8).6 The anesthetic must be injected very slowly—no faster than 0.5 mL/min for 4 to 5 minutes—to avoid pain and discomfort and to allow diffusion. Lee and colleagues found that computer-assisted delivery of the anesthetic was more successful than the traditional clinician-controlled approach, due to the slow pace that is necessary.7

Complications
Few complications are associated with this injection because of low vascularity in the area involved. The main challenges presented by this approach are pain, the time needed for the injection, and the time until the block takes effect. As noted previously, Lee and colleagues had greater success with computer-assisted delivery of the anesthetic (which is not available in most EDs) than with administration by a clinician, but failure rates were still high for both techniques, suggesting that this block is not reliable for inducing anesthesia.8

Infraorbital Nerve Block

Indications
The ability to induce an infraorbital nerve block is a necessary skill for an emergency medicine provider. This block can be achieved with an intraoral or extraoral approach. It anesthetizes the midface from the lateral upper lip to the lower eyelid, including the nasal skin but not the mucosa (Fig. 30.9A). This block is helpful in wound care and abscess drainage.

Anatomy
The infraorbital nerve is the terminal branch of the mandibular branch of the trigeminal nerve. The second division exits the cranium from the foramen rotundum and ultimately enters the face through the infraorbital canal. The infraorbital foramen is located on the inferior border of the infraorbital ridge on a vertical (sagittal) line with the pupil when the patient stares straight ahead (see Fig. 30.9B). The middle and superior alveolar nerves are also in close proximity to the main trunk of the infraorbital nerve near the foramen.

Intraoral Approach
Due to the location of the foramen, an intraoral approach is preferred. Provide local anesthetic to the buccal and gingival mucosa, as previously described. Place one finger extraorally on the foramen. This has two purposes: it gives the practitioner a tangible feeling of the needle, and the pressure will help direct the deposited anesthetic from pushing upward into the lower eyelid. Retract the cheek and enter the mucosa above the second premolar. Aim the needle toward the infraorbital foramen, approximately 2 cm from the mucosal surface, taking care not to enter the foramen (see Fig. 30.9C). Once the needle is in the proper position, inject 2 to 3 mL of anesthetic and gently massage the tissue overlying the area to promote distribution of the anesthetic. The alternative method is to perform a field block, which is less precise and anesthetizes a larger area. This technique can be used if the patient has no palpable landmarks or if a wider area of anesthesia is necessary.

Extraoral Approach
The infraorbital foramen can also be approached from an extraoral route (Fig. 30.10). This approach requires external preparation of the skin. Use the same landmarks as those used to locate the infraorbital foramen as described in the intraoral approach. The needle can be felt as it passes through the skin, the subcutaneous tissue, and the quadratus labii superioris muscle. After injection, firmly massage the infiltrated tissue for 10 to 15 seconds. It is usually visibly swollen. Place a finger under the eye to limit edema.

Complications
If the needle is angled posteriorly, it might enter and anesthetize the orbit. The risk of such positioning can be limited by placing the finger on the foramen and palpating the needle below the orbital rim. Injection into, as opposed to near, the foramen

Figure 30.8 For the anterior middle superior alveolar nerve block, inject anesthetic through the hard palate at the intersection of an imaginary line dividing the premolars at a point halfway between the midpalatine crest and the free gingival margin (arrow).
Regional anesthesia of the inferior alveolar nerve (IAN) should be in every emergency provider’s armamentarium. Dental complaints have been the cause of an increasing number of ED visits. If the pain or an injury involves the mandibular teeth or the lower lip and chin, the IAN block will be a useful tool. Examples of its application are for facial or dental trauma, dry socket, periapical abscess, and pericoronitis. The IAN block provides anesthesia to all the teeth on that side of the mandible and desensitizes the lower lip and chin via blockade of the mental nerve. The IAN can be anesthetized using the classic approach, the Gow-Gates technique, or Akinosi-Vazirani methods.

Anatomy
The IAN is the largest of the branches of the mandibular nerve (V3). It provides sensation to all the lower teeth and lip; the central incisors, lateral incisors, and the buccal aspect of the molar teeth may also receive sensory innervation. The nerve descends, covered by the external pterygoid muscle, and passes between the ramus of the mandible and the spheno-mandibular ligament to enter the mandibular canal. It is accompanied by the inferior alveolar artery and vein, proceeds along the mandibular canal, and innervates the teeth. At the mental foramen, the nerve bifurcates into an incisive branch, which continues forward to supply the anterior teeth. It gives off a side branch, the mental nerve, which exits from the mental foramen to supply the skin.

Approach
The patient can be seated either in a chair or upright with the head firmly against the back of the stretcher, so that when the mouth is opened, the body of the mandible is parallel to the floor. The easiest approach for the provider is with the patient seated or slightly reclined. Depending on the practitioner's handedness and comfort level, the best approach is to be positioned on the contralateral, or opposite, side of the nerve you are trying to block, as this will give better visual access to the targeted area. The best approach to regional

![Figure 30.9 Infraorbital nerve block. A, Distribution. B, Anatomy. C, Insert the needle directly above and parallel to the upper second premolar toward the infraorbital foramen. Placing a finger over the foramen helps to direct and determine the depth of insertion.](image)

![Figure 30.10 Infraorbital nerve block, external approach. While injecting, place a finger under the eyelid to minimize lid swelling.](image)
quick jerk of the head when an anxious patient first feels the needle.

To begin the technique, palpate the retromolar fossa with the index finger or thumb. With this maneuver, the greatest depth of the anterior border of the ramus of the mandible (the coronoid notch) can be identified (Fig. 30.11A). With your thumb in the patient’s mouth, resting in the retromolar fossa, and your index finger placed externally behind the ramus at the same height as the thumb, retract the tissues toward the buccal (cheek) side, and then visually locate the pterygomandibular triangle (see Fig. 30.11C). This technique also moves your finger safely away from the tip of the needle.

When topical anesthesia has been achieved, hold the syringe parallel to the occlusal surfaces of the teeth, angled so that its barrel lies between the first and second premolars on the opposite side of the mandible (see Fig. 30.11B). Failing to achieve the required angle is the most common reason for failure of this nerve block. If a large-barrel syringe is used, the corner of the mouth might hamper efforts to obtain the proper angle; therefore a 5- or 10-mL syringe with a 25-gauge needle should be sufficient. If you have difficulty achieving the necessary angle, you can carefully bend the 25-gauge needle approximately 30 degrees (Fig. 30.12). This modification presents the risk of needle fracture and its inherent risks; an alternative is to use one of the other blocks described in the following sections. Puncture inside the pterygomandibular triangle at a point 1 cm above the occlusal surface of the molars.

anesthesia is to use a topical anesthetic over the approach area to increase patient compliance. Place a gauze pad soaked with topical lidocaine in the area of the IAN while preparing the rest of the supplies, or use a cotton swab with viscous lidocaine. Despite the use of topical anesthesia, be ready for an unexpected
the needle position is suboptimal. In children, the angulation is not parallel to the occlusal surfaces of the teeth, so the barrel of the syringe should be held slightly higher because the mandibular foramen is lower.

One of the benefits of an IAN block is its ability to affect the other nerves in relative proximity. For example, one can anesthetize the lingual nerve, which innervates the anterior two thirds of the tongue, by placing several drops of anesthetic solution in that area while slowly withdrawing the syringe and needle. This technique consistently blocks the lingual nerve because of the proximity of the two nerves. The area that is anesthetized can also be extended by blocking the long buccal nerve by injecting 0.5 cc of anesthetic just distal and buccal to the last mandibular molar. Supplementing the IAN block with both lingual and buccal nerve blocks helps anesthetize any aberrant fibers that are innervating the teeth. Shortly after a successful injection, the patient will report tingling in the lower lip; however, 3 to 5 minutes is usually required to achieve complete anesthesia.

Complications
This nerve block reportedly has a 15% to 20% failure rate. As with any block, the complications are mainly related to deposition of the anesthetic in inadvertent areas; for the IAN block, misplacement most commonly affects the parotid gland, anesthetizing the facial nerve. This is a relatively benign complication that causes temporary facial paralysis (similar to Bell's palsy). It affects the orbicularis oculi muscle, making the patient unable to close the eyelid. Should this occur, protect the eye with an eye patch or tape until the local anesthetic has worn off (2 to 3 hours for 1% lidocaine). The duration of this complication is elongated to 3 to 6 hours if anesthesia is induced with bupivacaine (perhaps longer if it is used in combination with epinephrine).

Gow-Gates Block
Because the IAN block is frequently unsuccessful, other techniques have been developed to facilitate anesthesia in this region. The Gow-Gates and Akinosi-Vazirani techniques for regional anesthesia can augment or replace the IAN block. Introduced in the 1970s, the Gow-Gates mandibular block is an acceptable technique for achieving mandibular anesthesia, especially if traditional approaches fail (Fig. 30.13A). It involves the deposition of anesthetic at the lateral aspect of the anterior condylar head, in an attempt to anesthetize the trunk of the mandibular branch of the trigeminal nerve as it exits the foramen ovale (see Fig. 30.13B).

Approach
The key to this technique is patient compliance because the patient has to be able to open his or her mouth wide enough to allow the angle of approach that is necessary to achieve this block (see Fig. 30.13C). The optimal plane is parallel to an imaginary line drawn from the corner of the mouth to the intratragal notch. This is different from the plane used with the classic IAN block. Place a finger inferior to the tragus of the ear as the landmark for the lateral aspect of the anterior condyle (which is now sitting at the eminence). Insert the needle opposite the second molar with the barrel of the syringe between the opposite lower premolars and the corner of the mouth (similar to the inferior alveolar approach). With this plane maintained, advance the needle until it reaches the
condylar neck. The goal is to head below the insertion of the lateral pterygoid muscle, and it uses extraoral landmarks (tragus of the ear and corner of the mouth).\textsuperscript{9} Withdraw the needle, aspirate, and inject 1 to 2 mL of anesthetic.\textsuperscript{8,9}

**Complications**
The positive vascular aspiration rate with the Gow-Gates procedure is reported to be 1.6%, compared with 3.5% to 22% for the conventional IAN block technique.\textsuperscript{6} If the needle is directed more toward the medial aspect of the anterior condylar process, it will be in the proximity of the pterygoid plexus of veins and the sympathetic plexus of the internal carotid artery. If the sympathetic plexus is anesthetized, the patient will experience a temporary version of Horner syndrome.

**Akinosi-Vazirani Block**
The Akinosi-Vazirani block can be used to anesthetize the IAN, including its mental and lingual nerve branches. A benefit of this block is that it can be performed when the patient’s mouth is closed. This can be very helpful for the emergency physician because the condition necessitating the block can cause trismus, so opening the mouth wide enough for the classic or Gow-Gates techniques might not be possible.

**Approach**
With the patient’s mouth closed, distract the buccal surface from the teeth, which will allow palpation of the bony landmarks. The main difference with this procedure and the IAN or Gow-Gates block centers on the fact that you do not want to hit bone with the needle; instead, you are aiming for the pterygomandibular space. The syringe should be positioned at the height of the gingival junction of the maxillary teeth and parallel to the occlusal surfaces of the teeth (Fig. 30.14). If bone is contacted, simply redirect the needle slightly toward the teeth, which should allow you to enter the desired space. The temporalis muscle is also in this area, and care should be taken to not inject the anesthetic into it.

**Mental Nerve Block**
The mental nerve block is an easy and useful form of regional anesthesia and should be considered for all lower lip laceration repairs, especially those involving the vermilion border (Fig. 30.15). The mental nerve can be blocked with a complete IAN block or alone where it exits the mental foramen. An intraoral or extraoral approach can be used. An isolated block of the mental nerve is achieved by infiltrating local anesthetic where the nerve exits its bony foramen on the mandibular surface.

**Anatomy**
The mental nerve is a continuation of the IAN and innervates the mucosa and skin of the lower lip on the ipsilateral side of the mandible, with limited crossover of midline fibers. The nerve emerges from the mental foramen below the second premolar (Fig. 30.16 \textit{A} and \textit{B}). Generally, the foramen will be just medial to the pupil of the eye (when the patient stares straight ahead) along a sagittal plane.

**Approaches**
Like the infraorbital nerve, the mental nerve block can be performed via an intraoral or extraoral approach. As the area being anesthetized is very sensitive and limiting scarring is a priority, the intraoral approach should be used when possible. Before using either approach, identify the mental foramen by palpation approximately 1 cm inferior and anterior to the second premolar. As with the IAN block, packing a gauze pad soaked in viscous lidocaine into the gingival sulcus while preparing the remainder of the supplies is a good way to ensure topical anesthesia. Approach the mental foramen at approximately a 45-degree angle with a 1.3-cm (\(\frac{1}{2}\)-inch), 25- or 27-gauge needle on a 3-mL syringe, and infiltrate the area adjacent to the foramen with 1 to 2 mL of local anesthetic (see Fig. 30.16C). Avoid inserting the needle into the mental nerve foramen because the injection of liquid into that site can produce neuropraxia. Infiltrate around the foramen to provide anesthesia of the lower lip and soft tissue of the chin. For lacerations over the midline of the lip, administer anesthetic over the mental foramina bilaterally. This practice anesthetizes the crossover fibers and will ensure midline anesthesia.
Scalp Anesthesia

Scalp blocks provide surgical anesthesia for repair of lacerations, drainage of superficial abscesses, and exploration and débride-ment of wounds. In general, the larger nerves innervating the scalp are blocked best when they are induced near the exit of the nerve from the skull. This is because the nerves branch and spread superficially very quickly, so it becomes difficult to achieve regional anesthesia.

Anatomy

As shown in Fig. 30.17, the scalp receives its nerve supply from branches of the trigeminal nerve (CN V) and the cervical plexus. The forehead is supplied by the supraorbital and supratrochlear nerves, both of which are branches of the ophthalmic division of the trigeminal nerve. The temporal region receives its nerve supply from the zygomaticotemporal nerve (a V2 branch nerve) and the temporomandibular and auriculotemporal nerves (V3 branch nerves). The posterior aspect of the scalp is innervated by the great auricular and the greater, lesser, and least (third) occipital nerves. The nerves that supply the posterior aspect of the scalp originate from the cervical plexus.

Topographically, the nerves and vessels of the scalp are located in subcutaneous tissue above the epicranial aponeurosis (Fig. 30.18). From this level, they divide into small branchesgenerously infiltrating this area in a fanlike motion will block multiple sensory nerves.

Complications

The main complication of the mental nerve block stems from damaging the mental nerve where it exits the foramen, by direct needle trauma or pressure from the injection, causing neuropraxia.
that extend to the deeper layers (epicranium and periosteum).\textsuperscript{10,11} The scalp is a very vascular structure, so take care to provide hemostasis with vasoconstrictive agents while simultaneously monitoring the systemic effects of the chosen agents.

**Approaches**

A complete scalp block can be accomplished by individually blocking each nerve that supplies the scalp, but this approach is time-consuming, difficult, and rarely necessary. Because the nerves on the scalp are located superficially, especially above the eyebrows to the vertex of the skull, a local anesthetic can be used around the wound’s edges in most cases. Injection of local anesthetic to deeper levels is necessary only if bone is to be removed. Injection of a local anesthetic only in the deeper layers without subcutaneous infiltration results in an unsuccessful block and a greater amount of bleeding during surgical intervention.

In preparation for the procedure, clip hair or use sterile petroleum gel to slick hair away from the injured area. Prepare the skin with an antiseptic solution, and raise a skin wheal at any point along the skin with a 1.3-cm (\(\frac{1}{2}\) -inch), 25-gauge needle. Insert a 7.6-cm (3-inch), 22-gauge needle through the skin wheal and into the subcutaneous tissue. Advance it along the scalp circumferentially to surround the wound. Inject 0.5\% to 1\% lidocaine or 0.125\% to 0.25\% bupivacaine with epinephrine (1:200,000). Use epinephrine, unless there is a contraindication to it, to provide vasoconstriction and prevent excessive blood loss and absorption of local anesthetic. The total dose of the local anesthetic should not exceed the recommended dose for that particular agent. The first 10 to 15 minutes after the injection is the most critical period to monitor for local anesthetic toxicity.

**Greater and Lesser Occipital Nerve Block**

This relatively simple block may be useful in the ED for treating occipital neuralgia and tension headaches. For occipital neuritis, a long-acting corticosteroid such as methylprednisolone (20 to 40 mg) can be combined with a local anesthetic (see Chapter 52).

**Anatomy**

The posterior aspect of the head is supplied by the posterior rami of the cervical nerves. Two important branches of these nerves are the greater and lesser occipital nerves. The greater occipital nerve becomes superficial on each side at the inferior border of the obliquus capitis inferior muscle and runs superiorly toward the vertex over this muscle. The nerve is located medial to the occipital artery. The lesser occipital nerve is located approximately 2.5 to 3.5 cm lateral and 1 to 2 cm caudal to the greater occipital nerve (Fig. 30.19 A).

**Approach**

It is not usually necessary to shave or clip the scalp before performing greater and lesser occipital nerve blocks. The greater occipital nerve is best blocked at the nuchal line, which extends from the middle of the external occipital protuberance to the mastoid process. The nuchal line is located within the insertion sites of the trapezius muscle and the semispinalis muscles. At this site, the greater occipital nerve is just medial to the occipital artery.

First, palpate the occipital artery. Next, attach a 3.8-cm, 23- to 25-gauge needle to a syringe containing 5 mL of local anesthetic. Insert the needle into the skin (see Fig. 30.19 B). After obtaining paresthesia at the vertex, inject 5 mL of local anesthetic solution. Block the lesser occipital nerve with a fanlike injection of a local anesthetic solution, 2.5 to 3.5 cm lateral and 1 cm caudal to the point described for the greater occipital nerve.

This procedure is not usually associated with any complications. Intraarterial injections can be avoided by careful aspiration.

**Ophthalmic (V1) Nerve Block**

The lateral and medial branches of the supraorbital, supratrochlear, and infratrochlear nerves can be blocked by percutaneous local injection at the point where they emerge from the superior aspect of the orbit. Anesthesia of the forehead and scalp is achieved as far posteriorly as the lambdoid suture. This block can be used for débridement or topical treatment of burns or abrasions and for delicate lacerations of the upper eyelid. It is also ideal for removing small pieces of glass embedded in the forehead from a windshield injury (Fig. 30.20). Although this block is used infrequently, it is easily performed and not associated with serious side effects. It should be
Nerve blocks about the head and neck are relatively painless when induced carefully and slowly after topical mucosal anesthesia (for intraoral approaches) or local skin anesthesia when anesthesia of the forehead or the anterior aspect of the scalp is desired.

**Anatomy**

The subtle supraorbital notch, which is in line with the pupil (when the patient is staring straight ahead), can be palpated along the superior orbital rim (Fig. 30.21B). This landmark is the site of injection for blockade of the supraorbital nerves. The supratrochlear nerve is found 0.5 to 1.0 cm medial to the notch. This nerve innervates the midline and should be blocked bilaterally for wounds or procedures that require midline anesthesia. The infratrochlear nerve is not usually blocked but is found in the most medial aspect of the superior orbital rim.

**Approach**

With the patient in the supine position, hold a finger or a roll of gauze firmly under the orbital rim to avoid ballooning of anesthetic into the upper eyelid and raise a skin wheal over the lateral border of the upper orbital ridge. This should be approximately 0.5 cm from the supraorbital notch. Extend a 3.8-cm, 23- to 25-gauge needle attached to a syringe containing 5 mL of local anesthetic. Inject 1 to 3 mL of anesthetic in the area medial to the supraorbital notch (see Fig. 30.21C), and continue to withdraw the needle. Place another 1 to 2 mL of anesthetic in the area of the supraorbital notch as well. This effectively places a line of anesthetic solution along the orbital rim laterally to medially to ensure a block of all the branches of the ophthalmic nerve. Paresthesia in the form of an electric shock sensation over the forehead indicates a successful block.

**Complications**

Hematoma formation or swelling of the eyelid may occur but requires only local pressure to limit expansion. Occasionally, ecchymosis of the periorbital region will appear the next day, and the patient should be warned of this possibility.

**CONCLUSION**

Nerve blocks about the head and neck are relatively painless when induced carefully and slowly after topical mucosal anesthesia (for intraoral approaches) or local skin anesthesia...
(for extraoral blocks and approaches). They can alleviate patient apprehension, minimize wound margin misalignment, and contribute to better patient outcomes. The blocks should be considered in all cooperative patients who require regional anesthesia in the head and neck.

**Acknowledgments**

We would like to thank James T. Amsterdam and Kevin P. Kilgore for writing the original chapter and developing many of the images.

We thank Linda J. Kesselring, MS, ELS, for her copyediting, formatting, and organization. 

REFERENCES


Regional Anesthesia of the Thorax and Extremities
John J. Kelly and Jason Younga

Virtually every peripheral nerve can be blocked at some point along its course from the spine to the periphery (Videos 31.1 to 31.14), but digital nerve blocks (e.g., fingers and toes) are more commonly used than proximal blocks. Other common applications include femoral blocks for fractures of the femur, ankle blocks for foot injuries and infections, intercostal blocks for rib fractures, and wrist blocks for injuries to the palm.

The preparation, technique, choice of anesthetic, precautions, and complications are similar for all nerve blocks and are described in general in the following sections. The clinician is encouraged to use the same basic techniques and precautions for all nerve blocks. Specific precautions unique to a particular nerve block are included with the description of that block. Obvious precautions, such as aspiration before injection when the needle is in close proximity to a vascular structure, are not restated to avoid redundancy.

GENERAL CONCEPTS

Indications
For most lacerations and injuries seen in the emergency department (ED), local infiltrative anesthesia is adequate and more efficient than using a nerve block (see Chapter 29). Patients who require extensive repair and anesthesia of the entire extremity are often referred to a specialist, who may prefer to examine an unanesthetized limb. A nerve block is indicated when it will provide advantages over other techniques. Scenarios in which this requirement is met include the following:

- When distortion from local infiltration hampers closure (e.g., facial wounds) or compromises blood flow (e.g., fingertip)
- When anesthesia is required over a large area and multiple injections would be painful or when the large amount of anesthetic needed for local infiltration exceeds the recommended dose
- When a nerve block is the most efficacious form of treatment, such as an intercostal block for treating a rib fracture in a patient with chronic obstructive pulmonary disease (COPD)
- When local infiltration of the wound would be more painful than a regional nerve block, such as in the plantar surface of the foot or the palm of the hand
- When the block is performed to decrease pain during reduction of a finger or toe dislocation
- When extensive limb surgery or manipulation is required (e.g., extensive tendon repair) and other options are not available

Preparation
A brief history, including drug allergies (particularly to local anesthetics), medications, and systemic illnesses, should be taken from the patient. Peripheral vascular, heart, and liver disease may increase the risk for severe complications. Therefore information about the existence of these diseases should also be sought.

Instructions
Explain the procedure to the patient, including the pain of needle insertion, paresthesias, and possible complications that may occur. Discuss the potential need for additional anesthetic or alternative procedures if the nerve block fails. Be sure that the patient understands that the additional administration of an anesthetic is part of the normal procedure rather than an attempt to correct an improperly performed nerve block. It is not standard to obtain written informed consent for nerve blocks performed in the ED.

Equipment
For most nerve blocks performed in the ED, the following equipment is required: latex gloves, an antiseptic solution (e.g., chlorhexadine), a 10-mL syringe, an 18-gauge needle for drawing the anesthetic from the vial, and a 3.75-cm, 25- or 27-gauge needle for the nerve block. Note that the needle sizes given in text are general recommendations, but for the majority of blocks, a 25-gauge needle is ideal. In addition, keep standard resuscitation equipment for advanced cardiac life support readily available any time that local anesthetic agents are given.

Choice of Anesthetic
Factors influencing the choice of anesthetic agent for nerve blocks are similar to those for local infiltration (see Chapter 29 for extensive discussion). In general, most nerve blocks are done for the repair of painful traumatic injuries that are likely to cause pain long after the repair is completed. In such cases, select the anesthetic with the longest duration of action to maximize the patient’s analgesia. Buffering the anesthetic is strongly encouraged to lessen the pain of infiltration (see Chapter 29). For most of the blocks described in this chapter, buffered 0.25% bupivacaine is suggested as the anesthetic of choice, but equal volumes of 1% lidocaine with epinephrine can be substituted to provide faster onset of relief. The use of epinephrine on end-organ areas has traditionally been discouraged (e.g., tip of the nose, peripheral ear pinna, distal end of the penis), although the theoretical risk is unsubstantiated in clinical practice. Literature from 2010 describes the use and confirms the safety of lidocaine with epinephrine (1:100,000 concentration) for digital blocks. It would be prudent to avoid epinephrine-containing anesthetics in injuries involving vascular compromise or for those patients with peripheral artery disease. Higher concentrations of lidocaine (≤2%) or bupivacaine (0.5%) are commonly used for large nerves. Ropivacaine is another anesthetic with a rapid onset and a long duration of action (several hours). It has been reported to have fewer cardiotoxic and central nervous system effects than bupivacaine. Take care to avoid exceeding the recommended dosages of the anesthetic chosen.

Positioning the Patient
When possible, perform nerve blocks with the patient in the supine position to minimize the vasovagal syncope that may
Preparation of the Area to Be Blocked

To limit the incidence of infection, prepare the field in aseptic fashion before needle puncture. Allow the antiseptic solution to dry fully to achieve its maximal antibacterial effect. Sterile drapes and gloves are not routinely required but may be considered in addition to aseptic skin preparation for the initiation of blocks that (1) are close to large joints, vessels, and nerves; (2) are located in inherently contaminated areas of the body (e.g., groin, perineum); or (3) require simultaneous palpation of the underlying structures while injecting.

Choosing the Nerves to Block

Successful anesthesia requires appropriate knowledge of the relevant anatomy. Most areas to be anesthetized have overlapping sensory innervation and therefore require two or more nerves to be blocked. In addition, the cutaneous distribution of the various peripheral nerves differs slightly from patient to patient. Use a liberal margin of error when determining which nerves supply the desired area of anesthesia.

Locating the Nerve

When locating a nerve to be blocked, approach it from a site with easily identifiable anatomic landmarks. The best sites are those with good structural landmarks (e.g., prominent bones or tendons) immediately next to the nerve. For example, the four digital nerves are reliably found at the 2, 4, 8, and 10 o’clock positions around and just superficial to the proximal phalanx, whereas the median nerve lies between the palpable palmaris longus and flexor carpi radialis tendons at the proximal wrist crease. Nerves that course adjacent to easily palpable arteries, such as in the axilla and groin, are also easy to locate and are good sites for performing nerve blocks. Nerves that do not have adjacent structural or vascular landmarks are much more difficult to block.

Blocking nerves with good structural or vascular landmarks is straightforward: palpate the landmarks and follow the course of the nerve in relation to these landmarks. After visualizing the anatomy in the mind’s eye, insert the needle in close proximity to the nerve.

Blocking nerves with poor landmarks, such as the radial nerve at the elbow, requires skill, practice, and some degree of luck. To increase the likelihood of successfully blocking these nerves, consider using ultrasound-guided techniques (see Ultrasound Box 31.1).

Nerve Stimulator

A nerve stimulator is commonly used by anesthesiologists but has never gained popularity among emergency clinicians. Nevertheless, it helps locate nerves that do not have adjacent structural or vascular landmarks, which greatly increases the chances of successfully blocking these nerves.

Ultrasound

Use of ultrasound to identify injection sites for peripheral nerve blocks has gained popularity. Ultrasound has been used successfully to locate and block nerves in the neck (e.g., interscalene, axillary, and brachial plexus), lower extremity (e.g., femoral and saphenous nerve blocks), upper extremity (e.g., radial, ulnar, and median nerve blocks at the elbow and forearm), and the lumbar plexus. Ultrasonographic guidance negates the effects of anatomic variability, provides real-time needle guidance, and allows the operator to visualize the “spread” of local anesthetic. The use of ultrasound is associated with superior success rates with fewer attempts, less time to perform the block, and fewer complications compared to anatomic nerve blocks or use of nerve stimulators.

There are two basic ultrasound techniques used to facilitate nerve blocks: the in-plane and out-of-plane techniques. Using the in-plane technique, the needle is inserted at the side of the probe and advanced toward the target. With this technique, the entire needle is visualized as it traverses the plane of ultrasound. With the out-of-plane technique, the needle enters the skin away from the probe and is aimed at the plane of sound. As the needle moves toward the target, only the needle tip is visualized. Detailed descriptions of ultrasound-guided nerve blocks can be found later in this chapter (see Ultrasound Box 31.1).

Paresthesia

A common technique to ensure that the tip of the needle is in close proximity to the nerve is to elicit a paresthesia. Touching and mechanically stimulating the nerve with movement of the needle tip produces a tingling sensation or jolt known as a paresthesia, and it is felt along the distribution of the nerve. In practice, the jolt of a true paresthesia is often difficult to distinguish from the “ouch” of a pain-sensitive structure. When blocking proximal nerves at the elbow or axilla, the paresthesia travels far enough away from the injection site that it can be reliably distinguished from locally induced pain. Paresthesias at the level of the hand and wrist are more difficult to distinguish from pain. In both cases, paresthesia is a subjective feeling that requires intelligent and cooperative patients to understand what they are expected to feel and to remain relaxed and attentive so that they can distinguish an “ouch” from a jolt. Before the procedure, a simple explanation of what the patient should or may feel will facilitate cooperation. Although eliciting paresthesias is generally reliable in demonstrating that the needle is close to its target, some authors believe that it may theoretically increase the rate of complications as a result of mechanical trauma or intraneural injection.

Once the paresthesia is elicited, it is important to withdraw the needle 1 to 2 mm before injecting the anesthetic. If a paresthesia persists, stop the injection and reposition the needle.

Injecting the Anesthetic

Strive to ensure that the anesthetic agent is not inadvertently injected into a vessel or nerve bundle. In practice, a misplaced intravascular injection is usually of minimal consequence, but small amounts of epinephrine may cause systemic symptoms such as tachycardia or anxiety. Intraarterial injection, theoretically, is more dangerous than intravenous injection. Either way, aspirate the syringe to check for blood before injection. If no blood is aspirated, inject the anesthetic while observing the extremity for blanching, which suggests intravascular injection. If blanching occurs, reposition the needle before further injection.

Nerve bundle injection has the potential to cause nonspecific nerve injury. Severe pain or paresthesia during injection or
resistance to depressing the plunger suggests the possibility of intraneural placement of the needle. If any of these problems occur, immediately stop injecting and reposition the needle.

The onset and duration of anesthesia are both greatly influenced by how close the injected anesthetic is to the nerve. Onset occurs within a few minutes if the anesthetic is in immediate proximity to the nerve. Onset takes longer or may not occur at all if the anesthetic must diffuse more than 2 to 3 mm, which underscores the importance of locating the nerve before the injection.

More anesthetic is required if it must diffuse a long distance to the nerve. A range of suggested volumes of anesthetic is given with each nerve block description. For blocks in which a definite paresthesia is elicited or a nerve stimulator or ultrasound is used, the minimal recommended amount of anesthetic suffices. For blocks of smaller nerves, paresthesias are often not easily elicited, and the anesthetic must be injected in the general vicinity of the nerve. For these blocks or when doubt exists about the proximity of the needle to the nerve, larger amounts of anesthetic are recommended. This point cannot be emphasized strongly enough. The difference between a successful and an unsuccessful block may be merely an additional 2 mL of anesthetic. When in doubt, err on the high side of the recommended dosage. When blocking large nerves, many clinicians also opt for 2% lidocaine rather than the 1% solution.

With most blocks, the onset of anesthesia occurs in 2 to 15 minutes, depending on the distance that the anesthetic must diffuse to the nerve and the type of anesthetic used. Wait 30 minutes before deciding that a block is unsuccessful.

Complications and Precautions

Complications may result from peripheral nerve blocks but are rare in clinical practice. Most cannot be prevented by even perfect technique. General precautions include measures to minimize nerve injury, intravascular injection, and systemic toxicity. No actual statistics exist on the complication rate from nerve blocks performed by emergency clinicians, but they are extremely rare in clinical practice. Theoretically, infrequently performed blocks, blocks that require high doses of anesthetic, and blocks close to major vascular structures are more likely to have complications.

Nerve Injury

Nerve injury is rare but can occur secondary to (1) chemical irritation from the anesthetic, (2) direct trauma from the needle, or (3) ischemia as a result of intraneural injection. Overall, the incidence of serious neuronal injury is rare and occurs in 1.9 per 10,000 blocks. Given that placement of a nerve block is a blind procedure, nerve injuries do not necessarily represent an error in technique.

Chemical neuritis from the anesthetic is the most common nerve injury. The patient may complain of pain and varying degrees of nerve dysfunction, including paresthesia or motor or sensory deficit. Most cases are transient and resolve completely. Supportive care and close follow-up are the mainstays of treatment. Emergency clinicians should not exceed the recommended doses and concentrations of anesthetic (Table 31.1). In general, buffered lidocaine 1% or 2% or buffered bupivacaine 0.25% or 0.5% is safe for nerve blocks performed in the ED.

Direct nerve damage can be minimized by proper needle style, positioning, and manipulation. Use a short, beveled needle and keep the bevel parallel to the longitudinal axis of the nerve. Sharp pain or paresthesia indicates that the needle is close to or in the nerve. Avoid excessive needle movement when the tip of the needle is in contact with the nerve. If a 25-gauge needle is used, physical damage to a nerve should be minimal, even when directly touched by the tip of the needle. A 27-gauge needle is theoretically attractive, but its small size may limit aspiration testing and it may bend or break when attempting to block deep nerves.

Intraneural injection may rarely cause nerve ischemia and injury. Elicitation of a paresthesia or severe pain suggests that the needle has made contact with the nerve. When a paresthesia is elicited, withdraw the needle 1 to 2 mm before injecting the anesthetic. If the paresthesia occurs during injection, stop the injection and reposition the needle. Most neurons are

---

**TABLE 31.1 Recommended Volumes of Anesthetic for Various Nerve Blocks**

<table>
<thead>
<tr>
<th>NERVE</th>
<th>VOLUME (mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axillary</td>
<td>20–30&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Elbow</td>
<td></td>
</tr>
<tr>
<td>Ulnar</td>
<td>5–10&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Radial</td>
<td>5–15&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Median</td>
<td>5–15&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Wrist</td>
<td></td>
</tr>
<tr>
<td>Ulnar</td>
<td>5–15&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Radial</td>
<td>5–15&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Median</td>
<td>3–5&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Hip</td>
<td></td>
</tr>
<tr>
<td>Femoral</td>
<td>10–20&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Three-in-one</td>
<td>25–30&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Knee</td>
<td></td>
</tr>
<tr>
<td>Tibial</td>
<td>5–15&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Peroneal</td>
<td>5–10&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Saphenous</td>
<td>5–10&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Ankle</td>
<td></td>
</tr>
<tr>
<td>Posterior tibial</td>
<td>5–10&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Deep peroneal</td>
<td>3–5&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Saphenous, sural, and superficial peroneal</td>
<td>4–10&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Intercostal</td>
<td>5–15&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Hand</td>
<td></td>
</tr>
<tr>
<td>Metacarpal and web space</td>
<td>2–4&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Finger</td>
<td>1–2&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Foot</td>
<td></td>
</tr>
<tr>
<td>Metatarsal</td>
<td>10–15&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Web space</td>
<td>3–5&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Toe</td>
<td>2–5&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup>Anesthetic: 1% lidocaine or 0.25% bupivacaine (both with epinephrine).

<sup>b</sup>Anesthetic: 1% lidocaine or 0.25% bupivacaine (both without epinephrine).
surrounded by a strong perineural sheath through which the nutrient arteries run lengthwise. Injection directly into a nerve sheath may increase pressure within the nerve and compress the nutrient artery. Impaired blood flow results in nerve ischemia and subsequent paralysis. Intraneural injection is often heralded by severe pain, which worsens with further injection and may radiate along the course of innervation. The operator may notice difficulty depressing the plunger of the syringe. If the tip of the needle is in proper position, slow injection of the anesthetic should be minimally painful, and the anesthetic should go in without resistance.

**Intravascular Injection**

Intravascular injection may rarely result in both systemic and limb toxicity. Inadvertent intravascular injection produces high blood levels of the anesthetic. Exercise care when administering large amounts of anesthetic in close proximity to large blood vessels.

Intraarterial injection of anesthetics with epinephrine may cause peripheral vasospasm and further compromise injured tissue. Intravascular anesthetic is not toxic to the limb itself, although it may produce transient blanching of the skin by displacing blood from the vascular tree. Epinephrine, however, can cause prolonged vasospasm and subsequent ischemia if it is injected into an artery. This is especially worrisome when anesthetizing areas with little collateral circulation, such as the toes, fingers, penis, and tip of the nose. Severe epinephrine-induced tissue blanching or vasospasm may be reversed with local or intravascular injection of phentolamine (see extensive discussion in Chapter 29).

Vasospasm associated with the epinephrine in anesthetic solutions is rare, but experience in related clinical situations can help guide therapy. Roberts and Krisanda used a total of 5 mg of phentolamine infused intraarterially to reverse arm ischemia following 3 mg of epinephrine inadvertently administered into the brachial artery during cardiac resuscitation. Accidental injection of epinephrine from an Epi-Pen (Mylan, Canonsburg, PA) will produce an area of vasoconstriction. This is usually self-limited and requires no intervention, but it may take a few hours to be complete. However, digital ischemia from inadvertent epinephrine autoinjection (Epi-Pen) has been successfully treated both by proximal “digital block” with 2 mg of phentolamine and by local infiltration at the ischemic site with 1.5 mg of phentolamine.

The route of phentolamine administration should be guided by the clinical situation. Phentolamine must reach the site of vasospasm. Local infiltration may be effective for ischemia in a single toe or finger, whereas arterial injection has the advantage of delivering the medication directly to the arteries exhibiting spasm. For larger areas of involvement or in instances in which local infiltration is ineffective, use intraarterial injection. A dose of 1.5 to 5 mg appears to be effective in most cases, although a total of 10 mg may be used for local infiltration. Phentolamine, 5 mg, can be mixed with 5 mL of either normal saline or lidocaine. The small volume of the distal pulp space may limit the volume of the infiltration dose to 0.5 to 1.5 mL in the fingertip. Larger volumes and dosages can be used with proximal infiltrations. For intraarterial infusion of the radial artery at the wrist or the dorsalis pedis at the ankle, dosages of 1.5 to 5 mg of phentolamine are suitable. Slow infusion or graded dosages of 1 mg may provide enough phentolamine to reverse the ischemia without excessive systemic effects such as hypotension.

**Hematoma**

Hematoma formation may result from arterial puncture, particularly during blocks in which a major blood vessel is being used as a landmark to locate the nerve (e.g., axillary or femoral artery). Direct pressure for 5 to 10 minutes usually controls further bleeding. Use of small-gauge needles (e.g., 25- to 27-gauge) also helps minimize bleeding from a punctured artery. A minor coagulopathy is not a contraindication to a nerve block.

**Infection**

Infection is rare and can be minimized by following aseptic technique and using the lowest possible concentration of epinephrine. Injection should be made through noninfected skin that has been antiseptically prepared. Injection through a site of infection may spread the infection to adjacent tissues, fascial planes, and joints.

**Systemic Toxicity**

The incidence of systemic toxicity with local anesthetics has diminished significantly in the past 30 years. Interestingly, peripheral nerve blocks have been reported to have the highest incidence of systemic toxicity. Allergic reactions account for only 1% of untoward reactions (see Chapter 29).

**Limb Injury**

Injury to the anesthetized limb can result if the patient is permitted to use the limb, is advised to use heat or cold application, or performs wound care before the anesthesia has worn off. With major nerve blocks, do not release the patient from the ED until sensation and function have returned.

With minor blocks, the patient may be sent home but should be properly cautioned. Advise the patient to avoid ischemia-producing compression dressings (e.g., elastic bandages) because the anesthetized area may not sense impending problems.

**SPECIFIC NERVE BLOCKS**

**Intercostal Nerve Block**

Blocking the intercostal nerves produces anesthesia over an area of their cutaneous distribution (Fig. 31.1) and provides considerable temporary pain relief for patients with rib contusions or fractures. Rib fractures are typically quite painful and cause the patient to try to splint respirations to avoid excessive movement of the injured site. The resulting hypventilation, atelectasis, and poor expectoration may cause hypoxia or lead to pneumonia. This is particularly true in patients with preexisting pulmonary disease and minimal respiratory reserve, in whom further impairment of function may cause significant respiratory compromise. Theoretically, anesthetizing injured ribs eases pain and facilitates deep breathing and coughing. Unfortunately, no controlled studies have compared intercostal blocks and oral analgesics in patients with the types of rib fractures that are commonly managed on an outpatient basis. However, studies do suggest that intercostal blocks may be superior to analgesics in patients who have undergone thoracotomy. In these studies, those receiving intercostal nerve blocks had better results on pulmonary function tests, greater oxygenation, and earlier ambulation and discharge than did those receiving opioid analgesics.
There are several arguments against the routine use of intercostal nerve blocks in the ED. First, rib fractures are often tolerated well by young patients, who usually require minimal oral analgesics. Second, these blocks have a relatively short duration of action. The typical duration of action of a long-acting anesthetic with epinephrine is 8 to 12 hours. However, it should be noted that patients often experience partial analgesia for up to 3 days, a period that cannot be attributed to direct action of the anesthetic on the nerve. Perhaps the anesthetic reduces muscle spasm and the associated cycle of pain.

Finally, a wrongly perceived high incidence of pneumothorax and unsuccessful blocks deters many clinicians from performing intercostal nerve blocks in the ED. The true incidence of pneumothorax after intercostal nerve blocks is very low and not significant enough to prohibit the procedure. Moore reported that in more than 10,000 individual rib blocks performed, the incidence of pneumothorax was less than 0.1%.\textsuperscript{20} However, Shanti and associates reported that the incidence of pneumothorax was 1.4% for each individual intercostal nerve blocked.\textsuperscript{21} If more than one nerve requires blockade, the incidence of pneumothorax may be greater. The suggested approach to discussing intercostal blocks is to give patients the facts with regard to the duration of analgesia and possible complications and then allow them to decide on the method for themselves. Frequently, they prefer oral analgesics initially but may return for further relief of pain, at which time they are more amenable to a nerve block.

**Anatomy**

Each thoracic nerve exits the spine through the intervertebral foramen, which lies midway between adjacent ribs (Fig. 31.2A). It immediately gives off the posterior cutaneous branch, which supplies the skin and muscles of the paraspinal area. The intercostal nerve then continues around the chest wall and gives off lateral cutaneous branches at the midaxillary line. These branches are the sensory supply to the anterior and posterior lateral chest wall.

The intercostal nerve runs with the vein and artery in the subcostal groove. The vein and artery lie above the nerve and are somewhat protected by the rib during a nerve block. Posteriorly, the nerve is separated from the pleura and lungs by the thin intercostal fascia. When blocking the nerve in the posterior aspect of the back, particular care must be taken to avoid puncture of the thin fascia and underlying lung. Fortunately, most rib fractures occur in the anterior or lateral portion of the rib and can be blocked in the posterior axillary line, where the internal intercostal muscles lie between the nerve and the lung’s pleura and provide a buffer for minor errors in needle placement. Note that blocking the nerve in this area will anesthetize the entire course of the intercostal nerve because it is blocked before the cutaneous branches are given off.

**Technique**

To achieve adequate analgesia for most rib fractures, the lateral cutaneous branch needs to be anesthetized. Therefore perform blocks between the posterior axillary and midaxillary line at a point proximal to the origin of this branch (see Fig. 31.2A, arrow). Explain the procedure and its benefits and its risks, including potential pneumothorax, systemic toxicity, and ineffective block, before proceeding.

Use buffered 0.25% bupivacaine in a 10-mL syringe with a 3.75-cm, 25-gauge needle. Prepare the area to be injected in the usual aseptic manner. Use the index finger of the nondominant hand to retract the skin at the lower edge of the rib cephalad and pull it up and over the rib (see Fig. 31.2B). With the syringe in the opposite hand, puncture the skin close to the tip of the finger that is retracting the skin over the rib. Keep the syringe at an 80-degree angle to the chest wall with the needle pointing cephalad, and rest the hand holding the syringe on the chest wall for stability. In this position, the depth of needle penetration is well controlled. Slowly advance the needle until it comes to rest on the lower border of the rib. The bone should be felt through the tip of the needle.

At this point, release the skin retracted over the rib. As the skin returns to its natural position, the shaft of the needle will become perpendicular to the chest wall and the tip of the needle will be at the inferior margin of the rib. Shift the syringe from the dominant hand to the index finger and thumb of the nondominant hand. Rest the middle finger of the same hand against the shaft of the needle and exert gentle pressure on the shaft to “walk” the needle off the lower edge of the rib. Again, keep the palm of the hand planted firmly on the chest wall to ensure control of the needle. With the help of the dominant hand, slowly advance the needle 3 mm. Aspirate to be sure that the needle has not penetrated a blood vessel, and then inject 2 to 4 mL of anesthetic while carefully moving the needle in and out 1 mm to ensure that the compartment containing the nerve between the internal and external intercostal muscles is penetrated. This may also serve to minimize intravascular injection. Repeat the procedure on the two ribs above and below to ensure that the overlapping innervation from adjacent nerves is blocked.

Although the procedure just discussed seems extensive, it takes 1 to 2 minutes to perform once the operator is familiar with the technique, and three to five intercostals can be blocked in 10 minutes total.

**Precautions**

Initially place the needle at the lower edge of the rib. If it contacts the rib above this point, it cannot be walked off the lower edge of the rib at the proper angle. If it is inserted too low, over the intercostal space, it may be advanced through
**Intercostal Nerve Block**

**Anatomy**

Nerve blocked here

**Technique**

1. **Retract the skin**

2. **Introduce the needle with the skin retracted (touch the rib with the tip of the needle)**

3. **Skin traction released for needle advancement and infiltration**

The anesthetizing needle is advanced until it touches the rib, an obvious sensation to the operator. The needle is walked down the inferior portion of the rib until it is felt to drop off the bone. The needle is advanced a few millimeters and a generous amount of the anesthetic is deposited (2 to 4 mL per rib). Penetration too deep risks pneumothorax.

**Note the posterior area where the intercostal block is usually performed (arrow at the angle of the rib/posterior axillary line); the anesthetic will also block the lateral cutaneous, lateral mammary, and anterior cutaneous branches.**

**Note that the vein and artery lie under the inner portion of the rib, which offers them protection from the anesthetizing needle.**

the pleura and into the lung before the operator realizes that it is too deep. Before inserting the needle, it is always prudent to estimate the depth of the bone. If the bone is not encountered by this depth of insertion, reevaluate the position of the needle. Even after the needle has been properly walked off the edge of the rib, take care to not puncture the pleura and lung. The depth of the intercostal groove in which the nerve runs is 0.6 cm posteriorly and diminishes to 0.4 cm anteriorly.

Because the incidence of pneumothorax is low, a chest radiograph is not routinely required after this procedure. Observe asymptomatic patients for 15 to 30 minutes and instruct them to return if problems arise. If the patient has symptoms of pneumothorax (e.g., cough, a change in nature of the pleuritic pain, or shortness of breath), obtain a chest film before discharge.

If the clinician inadvertently causes a pneumothorax, treatment depends on its size. Many pneumothoraces from this procedure are small and require no specific intervention. Those smaller than 20% may be observed for 6 hours. During this time, administer a high concentration of oxygen to help decrease the size of the pneumothorax. If the pneumothorax does not enlarge, the patient may be released home with arrangements for close follow-up. Needle or catheter aspiration of larger pneumothoraces may be all that is needed. A chest tube is necessary if this method fails (see Chapter 10).

Nerve Blocks of the Upper Extremity

The upper extremity is supplied by the brachial plexus. The nerve roots and its branches—primarily the median, radial, ulnar, and musculocutaneous nerves—can be blocked at the level of the interscalene muscles, axilla, elbow, wrist, hand, or fingers. Nerve blocks at the axilla and elbow are seldom used in the ED, but interscalene nerve blocks are becoming more popular. Nerve blocks of the wrist are performed occasionally in the ED, but interscalene nerve blocks are becoming more popular. Nerve blocks of the wrist are performed occasionally before painful procedures or for repair of injuries to the hand. Metacarpal and digital blocks are used frequently to treat fractures, lacerations, and infections of the fingers.

Interscalene Nerve Block

An interscalene nerve block anesthetizes the nerve roots from the upper (C5 and C6) and middle (C7) trunks of the brachial plexus, providing anesthesia to the shoulder and proximal arm. The lower trunk of the brachial plexus (C8 and T1) is blocked approximately 40% of the time, resulting in inconsistent anesthesia to the medial aspects of the forearm and hand (i.e., the territory supplied by the ulnar nerve). In the ED, an interscalene nerve block may be ideal for shoulder dislocations or complex lacerations of the upper extremity.

Anatomy and Technique

The brachial plexus is formed by the ventral rami of the lower cervical and upper thoracic nerve roots and supplies cutaneous and muscular innervation to the shoulder and proximal arm. The trunks of the brachial plexus pass between the anterior and middle scalene muscles where they can be blocked at the level of the cricoid cartilage (i.e., C6) (Fig. 31.3).

To perform an interscalene block, position the patient supine with the head turned away from the side of the block. Identify the sternal notch, the sternal and clavicular heads of the sternocleidomastoid muscle, and the clavicle. The anterior and middle scalene muscles lie posterior to the clavicular head of the sternocleidomastoid muscle at the level of the cricoid cartilage (C6). The safest and most successful technique used to perform an interscalene block utilizes ultrasound guidance and is described in detail later in this chapter (see Ultrasound Box 31.1).

Complications

Interscalene nerve blocks using typical volumes of local anesthetic frequently cause diaphragmatic paralysis by blocking the phrenic nerve. This may result in dyspnea, hypoxemia, or hypercapnia that may not be tolerated by patients with underlying pulmonary dysfunction (e.g., COPD). Using smaller volumes of local anesthetic will reduce the incidence of phrenic nerve palsy, but it will also reduce the likelihood of a successful block. Spread of local anesthetic to surrounding neural tissue may cause a Horner syndrome from stellate ganglion block or hoarseness from a recurrent laryngeal nerve block. Epidural and total spinal anesthesia have also been reported. Rarely, pneumothorax due to the proximity of the pleura may occur.

Nerve Blocks at the Elbow

The median, ulnar, and radial nerves can be blocked at the elbow to provide anesthesia to the distal end of the forearm and hand (Fig. 31.4). For most injuries extensive enough to require a nerve block at the elbow, all three nerves must be blocked for successful anesthesia because of the variable and overlapping innervation of the forearm. Furthermore, injuries to the proximal and middle aspects of the forearm may require additional circumferential subcutaneous field blocks of the lateral, medial, and posterior cutaneous nerves.

Ulnar Nerve: Anatomy and Technique (Fig. 31.5A)

The ulnar nerve can be palpated in the ulnar groove on the posteromedial aspect of the elbow between the olecranon and the medial condyle of the humerus. This nerve supplies innervation to the small finger, the ulnar half of the ring finger, and the ulnar aspect of the hand.

With the elbow flexed, insert buffered 0.25% bupivacaine via a 3.75-cm, 25-gauge needle 1 to 2 cm proximal to the ulnar groove and advance the needle parallel to the course of the nerve. The tip of the needle should come to rest close to the proximal end of the groove. Do not block the nerve in the
Cross section of the elbow looking cephalad, right arm. The radial nerve (1), median nerve (2), and ulnar nerve (3) are demonstrated.

Figure 31.4

Radial nerve: Anatomy and Technique (See Fig. 31.5B)

The radial nerve and sensory branch of the musculocutaneous nerve run together in the sulcus between the biceps and brachioradialis muscles on the anterolateral aspect of the elbow. The block produces anesthesia of the lateral dorsal of the hand and the lateral aspect of the forearm.

Palpate the sulcus in which the nerve runs between the sharp border of the biceps muscle and the medial border of the brachioradialis muscle in the antecubital fossa just proximal to the skin crease of the elbow. Having the patient flex the elbow to 90 degrees and isometrically contract and relax these muscles will help define their borders. Puncture the skin with a 3.75-cm, 25-gauge needle halfway between the muscles, or 1 cm lateral to the biceps tendon, at a point 1 cm proximal to the antecubital crease and inject 5 to 15 mL of anesthetic. As with the radial nerve block, the use of ultrasound may improve success rates of median nerve blocks at the elbow and forearm (see Ultrasound Box 31.1).25-27 If a nerve stimulator is used, flexion of the small and ring fingers signals proximity to the nerve.

Median nerve: Anatomy and Technique (See Fig. 31.5C)

The median nerve runs medial to the brachial artery in the anteromedial aspect of the elbow. Block of this nerve anesthetizes the index, middle, and radial portion of the ring finger and the palmar aspect of the thumb and lateral part of the palm.

Palpate the brachial artery in the flexed arm at the elbow just proximal to the antecubital crease and medial to the prominent biceps tendon. Once the anatomy is defined and marked in the flexed arm, extend the arm to 30 degrees and insert buffered 0.25% bupivacaine via a 3.75-cm, 25-gauge needle slightly medial to the artery and perpendicular to the skin to the depth of the artery, approximately 2 to 3 cm, and inject 5 to 15 mL of anesthetic. As with the radial nerve blocks, the use of ultrasound may improve success rates of median nerve blocks at the elbow and forearm (see Ultrasound Box 31.1).25-27 A nerve stimulator can also facilitate the process and produces flexion of the wrist and index finger. Most commonly, median nerve blocks are performed at the wrist.

Nerve Blocks at the Wrist

The median, ulnar, and radial nerves may be blocked at the wrist to provide anesthesia to the hand. Most extensive injuries and procedures for which a wrist nerve block could be used can also be managed by local infiltration or a digital block. When compared with direct infiltration, wrist block anesthesia may have a slow and unreliable onset and can require more time to take effect if all three nerves are to be blocked. There are several circumstances, however, in which wrist nerve blocks are more advantageous than other types of blocks or anesthesia.

Diffuse lesions that may be difficult to anesthetize with local infiltration can easily be anesthetized with a wrist block. Deep abrasions with embedded debris, commonly the result of “road burn” from bicycle and motorcycle crashes, can be cleaned and debrided painlessly after a nerve block at the wrist. Hydrofluoric acid burns, which require treatment with numerous subcutaneous injections of calcium gluconate, and thermal burns that require extensive debridement are better tolerated after a wrist nerve block. Wrist blocks are also advantageous in a severely swollen and contused hand, in which small amounts of anesthetic injected locally may increase tissue pressure and produce further pain. Finally, deep lacerations of the palm are very painful to anesthetize with local infiltration and will also benefit from a wrist block.

When compared with nerves in the axilla and elbow, the nerves in the wrist are more easily located anatomically and can be blocked more reliably. All three nerves lie in the volar aspect of the wrist near easily palpated tendons. A nerve stimulator is not necessary, but may be useful in locating the nerves, particularly when one is learning how to perform these blocks.

The anatomy and technique for blocking each nerve follow. Note that the median nerve lies in the midline and deep to the fascia and the ulnar and radial nerves lie on their respective sides and have branches that wrap around dorsally. Blocking all three nerves at the wrist requires a block that when viewed end-on, roughly resembles a horseshoe straddling a horseshoe stake (Fig. 31.6).

Median Nerve: Anatomy and Technique (Fig. 31.7A)

In the wrist, the median nerve lies just below the palmaris longus tendon or slightly radial to it between the palmaris longus and flexor carpi radialis tendons. Both tendons are easily palpated, but the palmaris longus may be absent in up to 20% of patients, in which case the nerve is found...
Nerve Blocks at the Elbow

Ulnar Nerve
Distribution

Anatomy and Technique
With the elbow flexed, insert the needle 1 to 2 cm proximal to ulnar groove, and advance toward the groove parallel to the course of the nerve. Deposit 5 to 10 mL of anesthetic.

Radial Nerve
Distribution

Anatomy and Technique
Insert the needle on the volar surface of the elbow, 1 cm proximal to the antecubital crease, midway between the brachioradialis and biceps muscles. Deposit 5 to 15 mL of anesthetic.

Median Nerve
Distribution

Anatomy and Technique
Palpate the brachial artery in the flexed arm, proximal to the antecubital crease and medial to the biceps tendon. Then with arm flexed at 30 degrees, inject 5 to 15 mL of anesthetic slightly medial to the artery.

Figure 31.5 Nerve blocks at the elbow.
approximately 1 cm in the ulnar direction from the flexor carpi radialis tendon. The nerve lies deep to the fascia of the flexor retinaculum, but at a depth of 1 cm or less from the skin. The superficial position of the median nerve at the wrist is emphasized because a major cause of failure of this block is to instill the anesthetic too deep.

The palmaris longus tendon is located by having the patient make a fist with the wrist flexed against resistance (Fig. 31.8). Insert a 3.75-cm, 25-gauge needle perpendicular to the skin on the radial border of the palmaris longus tendon just proximal to the proximal wrist crease. Advance the needle slowly until a slight “pop” is felt as the needle penetrates the retinaculum and a paresthesia is produced. If no paresthesia ensues, it may be elicited in a more ulnar direction under the palmaris longus tendon. If a paresthesia is still not elicited, deposit 3 to 5 mL of anesthetic in the proximity of the nerve at a depth of 1 cm under the tendon. Although the nerve is surprisingly close to the skin, it is better to err slightly on the deep side of the retinaculum and continue depositing anesthetic as the needle is withdrawn because the retinaculum is an effective barrier to a successful nerve block from superficially injected anesthetic.

Radial Nerve: Anatomy and Technique (See Fig. 31.7B)
The radial nerve follows the radial artery into the wrist but gives off sensory nerve branches proximal to the wrist. These branches wrap around the wrist and fan out to supply the dorsal and radial aspect of the hand.

Nerve block here requires an injection in close proximity to the artery and a field block that extends around the dorsal aspect of the wrist. Insert a 3.75-cm, 25-gauge needle immediately lateral to the palpable artery at the level of the proximal palmar crease. At the depth of the artery, inject 2 to 5 mL of anesthetic. Distribute an additional 5 to 6 mL of anesthetic subcutaneously from the initial point of injection to the dorsal midline. Withdraw the needle and reposition it to complete the block. Withdrawing the needle and repositioning it to a site that has already been anesthetized help decrease the discomfort of numerous needlesticks. Ultrasound can also be used at the level of the wrist to identify the precise location of the radial artery.

Ulnar Nerve: Anatomy and Technique (See Fig. 31.7C)
The ulnar nerve follows the ulnar artery into the wrist, where they both lie deep to the flexor carpi ulnaris tendon. The flexor carpi ulnaris tendon is easily palpated just proximal to the prominent pisiform bone by having the patient flex the wrist against resistance. At the level of the proximal palmar crease, the artery and nerve lie just off the radial border of the flexor carpi ulnaris tendon; however, the nerve lies between the tendon and the artery and deep to the artery, which makes it difficult to approach the nerve from the volar aspect of the wrist without involving the artery.

A nerve block of the ulnar nerve can be carried out by two different approaches: lateral and volar. The lateral approach may be easier because of the reason stated previously. For the lateral approach, insert a 3.75-cm, 25-gauge needle on the ulnar aspect of the wrist at the proximal palmar crease and deposit a wheal of anesthetic horizontally under the flexor carpi ulnaris tendon. Then direct the needle toward the ulnar bone at a point deep to the flexor carpi ulnaris tendon and inject 3 to 5 mL of anesthetic solution as the needle is withdrawn. Like the radial nerve, cutaneous nerves branch off the ulnar nerve, wrap around the wrist, and supply the dorsum of the hand. Block these branches by subcutaneously injecting 5 to 6 mL of anesthetic from the lateral border of the flexor carpi ulnaris tendon to the dorsal midline. Another advantage of the lateral approach is that the dorsal branches can be blocked from the same injection site. As mentioned with the radial nerve, ultrasound can easily identify the ulnar artery at the level of the distal radius and ulna.

Nerve Blocks of the Digits
The digital nerve block is one of the most useful and most used blocks in the ED. Indications for choosing it include repair of finger lacerations and amputations, reduction of fractures and dislocations, drainage of infections, removal of fingernails, and relief of pain (e.g., from a fracture or burn).


**Nerve Blocks at the Wrist**

**Median Nerve**

*Distribution*

- Locate the palmaris longus tendon. Insert the needle on the radial side of the tendon just proximal to the volar wrist crease. Feel for a “pop” as the needle penetrates the retinaculum, and inject 3 to 5 mL of anesthetic.

**Radial Nerve**

*Distribution*

- First inject 2 to 5 mL of anesthetic immediately lateral to the radial artery at the level of the proximal palmar crease (not shown). Then inject another 5 to 6 mL from the initial injection point to the dorsal midline.

**Ulnar Nerve**

*Distribution*

- Insert the needle on the ulnar aspect of the wrist at the proximal palmar crease, under the flexor carpi ulnaris tendon. Inject 3 to 5 mL of anesthetic. Next, deposit 5 to 6 mL of anesthetic from the lateral border of the flexor carpi ulnaris to the dorsal midline (not shown).

*Figure 31.7* Nerve blocks at the wrist.
Regional Anesthesia of the Thorax and Extremities

The principal nerves supplying the finger are the palmar digital nerves (also called the common digital nerves). These nerves originate from the deep volar branches of the ulnar and median nerves, where they branch in the wrist. The palmar digital nerves follow the artery along the volar lateral aspect of the bone, one on each side, and supply sensation to the volar skin and interphalangeal joints of all five digits (Fig. 31.10). In the middle three fingers, these nerves also supply the dorsal distal aspect of the finger, including the fingertip and nail bed. Although many clinicians routinely block both sets of digital nerves, in the presence of normal anatomy, only the volar (palmar) branches must be blocked to obtain adequate anesthesia of the middle three fingers distal to the distal interphalangeal joint.

**Technique**

The digital nerves can be blocked anywhere in their course, including sites in the finger, in the web space between the fingers, and between the metacarpals in the hand. There are a variety of approaches to the nerves, including the dorsal and palmar approaches and the web space approach. Each has its merits, and the technique is similar at each level.

The dorsal approach has the advantage of thinner, less pain-sensitive skin than encountered with volar approaches. The hand can be held firmly and flat on the table to prevent withdrawal. The disadvantage is that two injections are needed with this approach to block both volar digital nerves.

The dorsal approach can be used in the dorsum of the hand at the metacarpals, just proximal to the finger webs at the proximal end of the proximal phalanx or distal to the web.
After injecting one side of the finger, pull the needle back slightly (without removing it) and redirect it across the top of the digit to anesthetize the skin on the opposite side (see Fig. 31.12B). Completely withdraw the needle and reinsert it at the site that was just anesthetized, and continue the block as described earlier. The presumed advantage of this method is that it minimizes the pain of the second skin puncture. However, because this technique requires the needle to be placed across the dorsal aspect of the finger, it increases the risk for extensor tendon puncture and trauma.

The palmar and web space approaches can be used most successfully for the middle three fingers when only a single puncture is required to block both volar nerves. This technique takes advantage of the anatomic fact that only the volar digital nerves must be blocked to obtain anesthesia of the total finger (except the proximal dorsal surface). If the thumb or fifth finger must be anesthetized, the dorsal branches must also be blocked to obtain anesthesia of the fingertip and fingernail area (see Fig. 31.11).

The palmar approach requires an injection in the palm, which is more painful than an injection in the dorsal skin. Insert the needle directly over the center of the metacarpal head and slowly inject the anesthetic as the needle is advanced to the bone (see Fig. 31.12C). At this point, withdraw the needle 3 to 4 mm and redirect it slightly to the left and right of center to block both digital nerves without withdrawing the needle. To be successful, a palpable soft tissue fullness should be appreciated. The technique requires 4 to 5 mL of anesthetic.

With the web space approach, hold the patient’s hand with your thumb and index finger over the dorsal and volar surface of the metacarpal head, respectively. Use your third finger to separate the patient’s fingers to expose the web space while your fourth and fifth fingers support the patient’s finger being anesthetized (see Fig. 31.12D). Insert the needle into the web space and inject 1 mL of anesthetic. Slowly advance the needle until it is next to the lateral volar surface of the metacarpal head and inject additional anesthetic. Withdraw the needle slightly and redirect it across the midline of the metacarpal head to the opposite digital nerve. Use the index finger to palpate “a fullness” as the anesthetic is injected. Again, firm massage of the injected area for 15 to 30 seconds enhances diffusion of the anesthetic through the tissue to the nerves. When needed, redirect the needle to the adjacent finger without withdrawing it to block both fingers with a single puncture (Fig. 31.13).

**Alternative Techniques**

**Jet Injection Technique**

Jet injection for a digital nerve block can be used effectively and is less painful than standard needle techniques. The technique described by Ellis and Owens uses 0.15 mL of 1% lidocaine delivered by a jet injector at 2600 psi. Make three injections in the lateral aspect of the proximal phalanx: the first, midway between the volar and dorsal surfaces; the second, dorsal to this; and the third, volar. Administer a combined total of 0.45 mL to each side of the phalanx at the 2, 3, and 4 o’clock positions, and 8, 9, and 10 o’clock positions in relation to the bone.

Advantages of jet injection are a less painful injection and avoidance of “needle phobia,” particularly in children. Potential disadvantages include lacerations, which may occur with tangential injection. Holding the injector perpendicular to the skin avoids this problem. Thick skin associated with older age,

**Figure 31.11** For the index, middle, and ring fingers, the volar digital nerves provide sensation not only to the volar surface of the fingers but also to the dorsal surface distal to the distal interphalangeal joints. Thus, to obtain anesthesia of the distal portion of these three fingers, only the volar nerves need be blocked. Note that for the thumb and little finger, both the dorsal and volar nerves need to be blocked to obtain distal anesthesia.

**Figure 31.12** A variation of the dorsal approach is performed as follows. With the web space approach, hold the patient’s hand with your thumb and index finger over the dorsal and volar surface of the metacarpal head, respectively. Use your third finger to separate the patient’s fingers to expose the web space. Insert a 3.75-cm, 25- or 27-gauge needle at the web space, just distal to the knuckle at the lateral edge of the bone (Fig. 31.12A). Once the tip of the needle is subdermal, it usually contacts the bone. At this point, create a skin wheal by injecting 0.5 to 1 mL of anesthetic without epinephrine. This serves to block the dorsal digital nerve and provide anesthesia at the injection site. Pass the needle lateral to the bone and toward the palmar surface until the palmar skin starts to tent slightly. Withdraw the needle 1 mm and inject 0.5 to 1.5 mL of anesthetic. This procedure is repeated on the opposite side of the finger. The result is a circumferential band of anesthetic at the base of the finger. Firm massage of the injected area for 15 to 30 seconds enhances diffusion of the anesthetic through the tissue to the nerves.

A variation of the dorsal approach is performed as follows. After injecting one side of the finger, pull the needle back slightly (without removing it) and redirect it across the top of the digit to anesthetize the skin on the opposite side (see Fig. 31.12B). Completely withdraw the needle and reinsert it at the site that was just anesthetized, and continue the block as described earlier. The presumed advantage of this method is that it minimizes the pain of the second skin puncture. However, because this technique requires the needle to be placed across the dorsal aspect of the finger, it increases the risk for extensor tendon puncture and trauma.

The palmar and web space approaches can be used most successfully for the middle three fingers when only a single puncture is required to block both volar nerves. This technique takes advantage of the anatomic fact that only the volar digital nerves must be blocked to obtain anesthesia of the total finger (except the proximal dorsal surface). If the thumb or fifth finger must be anesthetized, the dorsal branches must also be blocked to obtain anesthesia of the fingertip and fingernail area (see Fig. 31.11).

The palmar approach requires an injection in the palm, which is more painful than an injection in the dorsal skin. Insert the needle directly over the center of the metacarpal head and slowly inject the anesthetic as the needle is advanced to the bone (see Fig. 31.12C). At this point, withdraw the needle 3 to 4 mm and redirect it slightly to the left and right of center to block both digital nerves without withdrawing the needle. To be successful, a palpable soft tissue fullness should be appreciated. The technique requires 4 to 5 mL of anesthetic.

With the web space approach, hold the patient’s hand with your thumb and index finger over the dorsal and volar surface of the metacarpal head, respectively. Use your third finger to separate the patient’s fingers to expose the web space while your fourth and fifth fingers support the patient’s finger being anesthetized (see Fig. 31.12D). Insert the needle into the web space and inject 1 mL of anesthetic. Slowly advance the needle until it is next to the lateral volar surface of the metacarpal head and inject additional anesthetic. Withdraw the needle slightly and redirect it across the midline of the metacarpal head to the opposite digital nerve. Use the index finger to palpate “a fullness” as the anesthetic is injected. Again, firm massage of the injected area for 15 to 30 seconds enhances diffusion of the anesthetic through the tissue to the nerves. When needed, redirect the needle to the adjacent finger without withdrawing it to block both fingers with a single puncture (Fig. 31.13).

**Alternative Techniques**

**Jet Injection Technique**

Jet injection for a digital nerve block can be used effectively and is less painful than standard needle techniques. The technique described by Ellis and Owens uses 0.15 mL of 1% lidocaine delivered by a jet injector at 2600 psi. Make three injections in the lateral aspect of the proximal phalanx: the first, midway between the volar and dorsal surfaces; the second, dorsal to this; and the third, volar. Administer a combined total of 0.45 mL to each side of the phalanx at the 2, 3, and 4 o’clock positions, and 8, 9, and 10 o’clock positions in relation to the bone.

Advantages of jet injection are a less painful injection and avoidance of “needle phobia,” particularly in children. Potential disadvantages include lacerations, which may occur with tangential injection. Holding the injector perpendicular to the skin avoids this problem. Thick skin associated with older age,
Palpate the flexor tendon in the palm proximal to the metacarpophalangeal joint. Introduce a 25-gauge needle attached to a 3-mL syringe at a 45-degree angle and advance it toward the tendon sheath while maintaining constant slight pressure on the plunger of the syringe (Fig. 31.14). When the sheath has been entered, the anesthetic should flow freely. If it does not, it is presumed that the tendon has been entered. If this happens, withdraw the syringe slowly (while keeping slight pressure on the plunger) until anesthetic flows smoothly and easily. Inject a total of 2 mL of anesthetic solution (smaller volumes should be used in children). After the needle is

### Transthecal Digital Block Technique

A transthecal block is performed by making a single injection into the flexor tendon sheath, which produces rapid and complete finger anesthesia. It was first described by Chiu in 1990, who noted rapid finger anesthesia after injection treatment of a trigger finger. Cadaver studies suggest that the injected fluid diffuses out of the tendon sheath and around the phalanx and all four digital nerves.

### Digital Nerve Blocks

#### Dorsal Approach

1. Insert the needle at the web space, just distal to the knuckle at the edge of the bone. Once the needle tip is subdermal, inject 0.5 to 1 mL of anesthetic to block the dorsal digital nerve.

2. Advance the needle along the bone toward the palmar surface until the palmar skin begins to tent. Inject another 0.5 to 1 mL of anesthetic to block the volar digital nerve.

3. Repeat steps 1 and 2 on the opposite side of the finger. The result is a circumferential band of anesthetic around the base of the finger. Firmly massage the area for 30 seconds to enhance diffusion of the anesthetic.

#### Dorsal Approach—Alternative Method

1. Block both the dorsal and volar digital nerves on one side of the finger as described above. Do not fully remove the needle after blocking the volar nerve.

2. Without removing the needle, redirect it across the top of the finger to anesthetize the skin on the opposite side. After injecting the opposite side, remove the needle.

3. Insert the needle at the site that was anesthetized in step 2, and block the other side of the finger. The presumed benefit of this technique is that it minimizes pain at the second skin puncture site.

---

**Figure 31.12** Digital nerve blocks.

*Continued*
Digital Nerve Blocks, Cont’d

Palmar Approach

1. Insert the needle directly over the center of the metacarpal head and slowly inject anesthetic while advancing the needle to the bone.
2. Withdraw the needle 3 to 4 mm (without completely removing it) and slightly angle it to the right or left of center to block one of the volar digital nerves.
3. Repeat on the other side of the digit. To be successful, a palpable soft tissue fullness should be appreciated. Usually 4 to 5 mL of anesthetic is required.

Web-Space Approach

1. Hold the patient’s hand with your thumb and index finger over the metacarpal, and spread the fingers to expose the web space.
2. Insert the needle into the web space and inject 1 mL of anesthetic. Slowly advance the needle until it is next to the volar surface of the metacarpal head and inject additional anesthetic.
3. Withdraw the needle slightly and redirect it across the midline of the metacarpal head to the opposite digital nerve. Inject additional anesthetic at this location. You will be able to feel the tissue distention by the anesthetic with your thumb, but be careful to avoid passing the needle through the skin and puncturing your thumb.

Complications and Precautions
The injection should go in smoothly, without resistance during injection. Although the finger is forgiving of transient pressure from excessive anesthetic, if the injection site becomes excessively tense, digital perfusion may be compromised. Even if epinephrine-containing solutions are used for a digital block in otherwise healthy individuals without peripheral vascular disease, it is unlikely that serious ischemic injury will occur. However, Hill and colleagues found the technique to be “clinically equal” to traditional digital blocks. Other authors have stated that the traditional digital block is easier to perform and produces less pain during and after injection. Theoretically, the technique may increase risk for injury to the tendon.

Significant vasoconstriction generally lasts less than 60 minutes, within the time interval for which an ischemic tourniquet can safely be used in the same area. However, if the entire digit remains blanched for more than 15 minutes, it is prudent to reverse the α-agonism of epinephrine with phentolamine (see Chapter 29). Using a pulse oximeter on the affected finger may help quantitate the degree of ischemia.
The small size of the digital arteries and nerves makes intravascular or intraneural injection less likely. Inadvertent intravascular injection may cause digital ischemia from vasospasm or displacement of blood out of the capillary bed by the anesthetic. Blanching of the finger as the anesthetic is injected suggests intravascular injection. If this is observed, immediately discontinue the injection. Usually, the ischemia is transient and self-resolving, and serious complications are rare. Massage or topical application of nitroglycerin paste may be attempted if the ischemia persists.37

Commonly, the digital nerve is lacerated or damaged by the initial injury to the finger. Careful evaluation using two-point discrimination should be performed to determine the extent of nerve injury before blocking the nerve. Even if nerve injury is questionable, it should be documented in the chart, and the patient should be advised of the injury before the nerve block. Careful evaluation and patient education should prevent misconceptions about the cause of the nerve injury. Although most isolated digital nerve injuries are not debilitating, they heal slowly and can be annoying to the patient. Digital nerve injury proximal to the distal interphalangeal joint may be repaired surgically. Nerve repair may be undertaken immediately when specialty consultation is available or be delayed after initial simple closure.

**Figure 31.13** Web space block techniques. Both volar digital nerves are blocked with one injection. A, For a metacarpal head block from the palmar surface, bending the needle 30 to 45 degrees allows easier access to the digital nerves without the syringe getting in the way. Make sure the needle is advanced far enough to block the side opposite the needle entrance, in this case to where the gloved thumb is positioned. B, Adjacent fingers can be blocked with only one needle puncture in the web space. After blocking the middle finger, withdraw the needle slightly, and without exiting the skin, redirect it to the metacarpal head of the index finger (arrow).

**Figure 31.14** Transthecal block. The flexor tendon sheath is entered volarly just proximal to the metacarpophalangeal joint. With the use of a 25-gauge needle on a 3-mL syringe, the fluid should flow easily. Inject 2 to 3 mL of anesthetic and apply pressure to the proximal end of the tendon sheath.

Nerve Blocks of the Lower Extremity

In the lower extremities, nerve blocks can be performed in the groin (e.g., femoral nerve block), ankles, popliteals, metatarsals, and toes. Femoral nerve block is the least often performed but is an effective method for providing analgesia to ED patients with femoral neck fractures. Ankle, metatarsal, and digital blocks are used frequently to treat ingrown toenails, foreign bodies, fractures, and lacerations of the forefoot and toes.

**Femoral Nerve Block (Three-In-One Block)**

A femoral nerve block provides significant analgesia to the proximal end of the femur and complete analgesia to the femoral shaft. It has been used to supplement anesthesia for a variety of surgical procedures on the anterior aspect of the thigh and knee and to provide postoperative analgesia after hip surgery. In the ED, this block is primarily used to provide analgesia for patients with hip fractures.38-40 It can be especially helpful in the management of elderly patients and those with respiratory compromise or poor pulmonary reserve, in whom high doses of opiate analgesics may be problematic.38

The three-in-one nerve block may be used to block the femoral, obturator, and lateral femoral cutaneous nerves with a single injection. The femoral nerve runs down the thigh in a fascial sheath that is continuous with the nerve sheath that contains all three nerves more proximally. If a large amount of local anesthetic is injected into this sheath, it will track proximally, medially, and laterally and thereby block all three nerves and provide more complete analgesia of the femoral neck and hip joint. The technique for performing both a femoral and three-in-one nerve block is identical except that the three-in-one block requires a larger volume of local anesthetic (25 to 30 mL vs. 20 mL).

Because the femoral and three-in-one nerve blocks are technically similar and associated with the same potential complications, and the fact that the three-in-one nerve block provides better analgesia of the femoral neck and hip joint, it
would seem logical to choose the three-in-one over the femoral nerve block for these patients. In clinical practice, however, the lateral femoral cutaneous nerve is less likely to be blocked than the femoral nerve, and the obturator nerve is frequently left unblocked despite proper technique. Nevertheless, because of potentially better analgesia obtained with the three-in-one nerve block and no clinically significant downside in comparison to a femoral nerve block, the remainder of the section will focus on performing the three-in-one nerve block.

**Anatomy (Fig. 31.15A)**

The femoral nerve is formed from the posterior branches of L2-L4 and is the largest branch of the lumbar plexus. The nerve emerges from the psoas muscle and descends between the psoas and iliacus muscles. It passes under the inguinal ligament in the groove formed by these muscles lateral to the femoral artery and divides into anterior and posterior branches. The anterior branches innervate the anterior aspect of the thigh, and the posterior branches innervate the quadriceps muscle and continue below the knee as the saphenous nerve to provide sensory innervation from the medial side of the calf to the medial malleolus.

The lateral femoral cutaneous nerve arises from the second and third lumbar nerve roots. The nerve emerges from the lateral border of the psoas muscle and travels under the iliac fascia, across the iliac muscle, and under the inguinal ligament 1 to 2 cm medial to the anterior superior iliac spine. It branches into anterior and posterior branches 7 to 10 cm below the anterior superior iliac spine. The anterior branch innervates the skin over the anterolateral aspect of the thigh to the knee, whereas the posterior branch of the nerve innervates the lateral part of the thigh from the greater trochanter to the middle of the thigh.

The obturator nerve arises from the anterior divisions of L2-L4. It descends through the fibers of the psoas muscle and emerges from its medial border near the brim of the pelvis. It then passes behind the common iliac arteries and runs along the lateral wall of the lesser pelvis, above and in front of the obturator vessels to the upper part of the obturator foramen. Here, it enters the thigh through the obturator canal and divides into an anterior and a posterior branch. The obturator nerve is responsible for sensory innervation of the skin of the medial aspect of the thigh and motor innervation of the abductor muscles of the lower extremity.

**Technique (See Fig. 31.15B)**

Place the patient in a supine position and prepare the skin overlying the femoral triangle in the usual fashion. Palpate the femoral artery 1 to 2 cm distal to the inguinal ligament. Inject a wheal of lidocaine 1 to 2 cm lateral to this point. Advance the needle at a 45° to 60° angle to the skin until (1) a “pop” and sudden loss of resistance are felt, (2) paresthesia is elicited, or (3) the needle pulsates laterally. Inject 25 to 30 mL of anesthetic. If proximity to the nerve is uncertain, inject in a fanlike distribution lateral to the femoral artery.

---

**Figure 31.15** The femoral nerve/“three-in-one” block.
Anatomy and Technique

The fascia iliaca block is an alternative to the femoral and three-in-one nerve blocks that more reliably blocks the lateral cutaneous femoral nerve. It has comparable effectiveness to the three-in-one nerve block, but is easier to perform because it does not depend on deposition of local anesthetic in close proximity to the nerves. Instead, it relies on diffusion of the anesthetic along the fascial plane to block the femoral and lateral cutaneous nerves. The fascia iliaca block can be performed using an anatomic approach or with ultrasound guidance.

A nerve block of the foot is better tolerated by the patient than local infiltration in all but the most minor procedures; it is the method of choice for treating injuries (e.g., lacerations, foreign bodies) of the sole. The skin of the sole is thicker and more tightly bound to the underlying fascia by connective tissue septa than is skin in other parts of the body. Puncturing this skin can be difficult and is always quite painful. The fibrous septa can limit the amount and spread of anesthetic. If large amounts of anesthetic are injected, the volume of injected substance quickly exceeds the space available, which can lead to painful distention of the tissue and circulatory compromise of the microvasculature. Local infiltrative anesthesia is adequate for treating minor injuries to the dorsum of the foot in which only small amounts of anesthetic are needed. However, for more extensive procedures such as incision and drainage, extensive wound care, and foreign body removal, an ankle block is better tolerated.

Nerve Blocks of the Ankle

Nerve block of the five nerves of the ankle—the deep peroneal (anterior tibial), posterior tibial, saphenous, superficial peroneal (musculocutaneous), and sural nerves—provides anesthesia to the foot. Of all the nerve block techniques described, these are the most technically difficult and most prone to failure. Depending on the desired area of anesthesia, one or more of the five nerves are blocked. These blocks can be used during operative procedures and repair of injuries to the foot. They are particularly useful in providing anesthesia to the sole of the foot for repair of lacerations and removal of foreign bodies.
Block the posterior tibial nerve in the medial aspect of the ankle between the medial malleolus and the Achilles tendon. Palpate the tibial artery just posterior to the medial malleolus. The injection site is 0.5 to 1.0 cm superior to this point. If the artery is not palpable, use a point 1 cm above the medial malleolus and just anterior to the Achilles tendon.

Insert a 3.75-cm, 25-gauge needle at a 45-degree angle to the mediolateral plane (the needle is almost perpendicular to the skin), just posterior to the artery. At the estimated depth of the artery (approximately 0.5 to 1.0 cm deep), wiggle the needle slightly to produce a paresthesia. If a paresthesia is easily palpated by having the patient dorsiflex the big toe. The deep peroneal nerve (also called the anterior tibial nerve) supplies the web space between the big and second toes. In the ankle, it lies under the extensor hallucis longus tendon. The saphenous nerve runs superficially with the saphenous vein between the medial malleolus and the tibialis anterior tendon, which is prominent when the patient dorsiflexes the foot. The saphenous nerve supplies the medial aspect of the foot near the arch.

The posteriorly located nerves are the posterior tibial and sural nerves (see Fig. 31.17B). The sural nerve runs subcutaneously between the lateral malleolus and the Achilles tendon and supplies the lateral border, both volar and dorsal, of the foot. The posterior tibial nerve runs with the posterior tibial artery, which can be palpated between the medial malleolus and the Achilles tendon. It lies slightly deep and posterior to the artery.

The posterior tibial nerve is one of the major nerve branches to the foot. After passing through the ankle, it branches into the medial and lateral plantar nerves, which supply sensation to most of the volar aspect of the foot and toes and motor innervation to the intrinsic muscles of the foot.

**Technique**

A complete nerve block of the foot requires blocking three subcutaneous nerves and two deeper nerves (Fig. 31.18). Once familiar with the anatomy, an experienced clinician can anesthetize all five nerves quickly by placing subcutaneous band blocks around 75% of the ankle circumference and two deep injections: one next to the palpable posterior tibial artery and the other under the extensor tendon of the big toe.

The five nerves of the foot are commonly blocked in combinations of two or more. Small procedures clearly within the distribution of one nerve may require only a single nerve block. However, overlap of the nerves’ sensory distribution frequently necessitates blocking a number of nerves for adequate anesthesia. Nerve block of the sural and posterior tibial nerves together anesthetizes the bottom of the foot and is the most useful combination. Using anatomic landmarks, nerve blocks of the ankle have a 95% success rate, making ultrasound unnecessary in most cases.45

**Figure 31.17** Anatomy and distribution of the sensory nerves of the lower part of the leg and foot.

**Figure 31.18** A complete nerve block of the foot requires blocking three superficial nerves (sural, saphenous, and superficial peroneal) and two deep nerves (posterior tibial and deep peroneal). *AT*, Anterior tibial; *EHL*, extensor hallucis longus.

**Posterior Tibial Nerve** (Fig. 31.19A)

Block the posterior tibial nerve in the medial aspect of the ankle between the medial malleolus and the Achilles tendon. Palpate the tibial artery just posterior to the medial malleolus. The injection site is 0.5 to 1.0 cm superior to this point. If the artery is not palpable, use a point 1 cm above the medial malleolus and just anterior to the Achilles tendon.

Insert a 3.75-cm, 25-gauge needle at a 45-degree angle to the mediolateral plane (the needle is almost perpendicular to the skin), just posterior to the artery. At the estimated depth of the artery (approximately 0.5 to 1.0 cm deep), wiggle the needle slightly to produce a paresthesia. If a paresthesia is
**Posterior Tibial Nerve Blocks at the Ankle**

**Distribution**

Palpate the tibial artery posterior to the medial malleolus. Insert the needle 1 cm superior to this point, perpendicular to the skin. At a depth of 1 cm, inject 3 to 5 mL of anesthetic. See text for additional details.

**Anatomy and Technique**

![Anatomy and Technique](image1)

**Sural Nerve Nerve Blocks at the Ankle**

**Distribution**

Palpate the Achilles tendon and lateral malleolus. Inject 3 to 5 mL of anesthetic subcutaneously in a band between the Achilles tendon and lateral malleolus, about 1 cm superior to the malleolus.

**Anatomy and Technique**

![Anatomy and Technique](image2)

**Superficial Peroneal Nerve Blocks at the Ankle**

**Distribution**

Palpate the extensor hallucis longus tendon and the lateral malleolus. Inject 4 to 10 mL of anesthetic subcutaneously in a band between the tendon and the malleolus.

**Anatomy and Technique**

![Anatomy and Technique](image3)

*Figure 31.19 Nerve blocks at the ankle.*
elicited, inject 3 to 5 mL of anesthetic. If no paresthesia is produced, advance the needle inward, again at a 45-degree angle, until it hits the posterior aspect of the tibia. Withdraw the needle approximately 1 mm and inject 5 to 7 mL of anesthetic while slowly withdrawing the needle another 1 cm. A rise in temperature of the foot, because of vasodilation from loss of sympathetic tone, may herald a successful block.

Sural Nerve (See Fig. 31.19B)
Block the sural nerve on the lateral aspect of the ankle between the Achilles tendon and the lateral malleolus. Inject 3 to 5 mL of anesthetic subcutaneously in a band approximately 1 cm above the lateral malleolus between the Achilles tendon and the lateral malleolus.

Superficial Peroneal Nerves (See Fig. 31.19C)
Block the superficial peroneal nerves on the anterior aspect of the ankle between the extensor hallucis longus tendon and the lateral malleolus by subcutaneously injecting 4 to 10 mL of anesthetic in a band between these landmarks.

Deep Peroneal Nerve (See Fig. 31.19D)
Block the deep peroneal nerve anteriorly beneath the extensor hallucis longus tendon. The injection site is 1 cm above the base of the medial malleolus between the extensor hallucis longus and anterior tibial tendons. Palpate the tendons by having the patient dorsiflex the big toe and foot, respectively. Create a lidocaine wheal at the injection site. Direct the needle approximately 30 degrees laterally and under the extensor
Anatomy
Each toe is supplied by two dorsal and two volar nerves, which are branches of the major nerves of the ankle. The dorsal digital nerves are the terminal branches of the deep and superficial peroneal nerves. The volar nerves are branches of the posterior tibial and sural nerves.

The location of the nerves in relation to the bones varies with the site of the foot. In the toes, the nerves lie at the 2, 4, 8, and 10 o’clock positions in close relationship to the bone. In the proximal part of the foot, the nerves run with the tendons and are not in close relationship with the bones (Fig. 31.20).

Technique
The digital nerves can be blocked at the metatarsals, interdigital web spaces, or toes. The bones of the foot can be palpated easily from the dorsum and are used as the landmarks for estimating the location of the nerves. Proximally, the nerves’ relationship to the bones is less consistent, which makes definitive needle placement and successful block less reliable. In the toes, the position of the nerves is more consistent; however, minimal subcutaneous tissue space is available for the injected solution. At the web space, the nerves are located in close relationship to the bone, and ample space is available for injecting the anesthetic; hence, for most procedures the web space is the preferred site for a digital nerve block.

The technique for toe and metatarsal blocks is similar (Fig. 31.21). All four nerves supplying each toe are usually blocked because of their sensory overlap. Perform the block from the dorsal surface, where the skin is thinner and less sensitive. Start by placing a 1-mL skin wheal dorsally between the metatarsal bones. Advance the needle until the volar skin tents slightly, and inject 2 mL of anesthetic as the needle is withdrawn. Without removing it, redirect the needle in a different volar direction, and repeat the procedure. Deposit a total of 5 mL of anesthetic in a fanlike pattern in each metatarsal space. Again, because of sensory overlap, two or more spaces need to be anesthetized for each toe to be blocked.

For a web space block, select a site on the dorsum just proximal to the base of the toe. Insert a 3.75-cm, 27-gauge needle attached to a 10-mL syringe at the lateral edge of the bone and place a subcutaneous wheal between the skin and the bone with 0.5 to 1.0 mL of anesthetic. This serves to block the dorsal nerve and minimize pain at the needle insertion site. Advance the needle just lateral to the bone toward the sole until the needle tents the volar skin slightly. Withdraw the needle 1 mm and inject 0.5 to 1.0 mL of anesthetic. As the needle is withdrawn, inject another 0.5 mL to ensure a successful block. Repeat the procedure on the opposite side of the toe. In this manner, two columns of anesthetic are placed on each side of the toe in the area through which the four digital nerves run. A total of 2 to 4 mL of anesthetic is used. For blocks done in the toe itself, the procedure is the same, but smaller amounts of anesthetic (e.g., <2 mL) are used because of the limited subcutaneous space and fear of vascular compression. Alternative techniques using a single injection site, as described for the finger, can be performed.

Complications and Precautions
Complications of lower extremity nerve blocks are similar to those associated with nerve blocks performed in the upper extremity and include intravascular injection, local anesthetic toxicity, nerve trauma, hematoma formation, and failure of the block.

The precautions that apply to the hand and fingers apply to the foot and toes. Ischemic complications can be avoided by paying attention to changes in the skin during the injection. Blanching heralds possible intravascular injection or vascular compression. If the skin blanches, halt the procedure and reevaluate the position of the needle and the amount and content of the injected solution. The total volume of anesthesia should not exceed the recommended amount. The literature
Nerve Blocks of the Toes

**Web Space Block**

1. Insert the needle on the dorsum just proximal to the base of the toe. Advance to the edge of the bone and inject 0.5 to 1 mL of anesthetic to block the dorsal nerve.

2. Advance the needle along the bone toward the sole until the needle slightly tents the volar skin. Withdraw the needle 1 mm and inject 0.5 to 1 mL of anesthetic.

3. Repeat on the other side of the toe. A total of 2 to 4 mL of anesthetic is typically used.

**Alternative Technique**

1. Block the dorsal and volar nerves on one side of the toe as described above. Do not completely remove the needle after anesthetizing the volar nerve.

2. Redirect the needle (without completely removing it) across the dorsal surface of the toe. Inject anesthetic on the dorsal surface of the opposite side of the toe, and then remove the needle.

3. Insert the needle through the newly anesthetized region, and block the dorsal and volar nerves on this side of the toe. This method minimizes pain felt at the second injection site.

**Figure 31.21** Nerve blocks of the toes.

---

**ULTRASOUND BOX 31.1: Nerve Blocks of the Thorax and Extremities**

Regional nerve blocks are typically performed by identifying anatomic landmarks and blindly injecting anesthetic agents. Nerve stimulators may be used to identify larger nerves and ensure proper placement of anesthetics. However, these techniques are not infallible and improper placement of anesthetic may result. Additionally, some nerve blocks, such as the scalene block, may be avoided because of concern regarding adjacent anatomic structures. Use of ultrasound allows the clinician to identify the nerve in question, as well as to directly guide the application of anesthetic. Furthermore, nearby structures such as arteries or veins can be identified and avoided, thereby offering the operator greater confidence in performing more advanced blocks. Despite a limited number of randomized controlled trials, preliminary evidence seems to support the use of ultrasound, especially with regard to patient safety.1

Although each nerve block will require a slightly different approach, similar principles apply. Use a high-frequency transducer (10 to 12 MHz or higher) to ensure the proper resolution and to identify the structures in question (Fig. 31.US1). Gather equipment as described earlier in this chapter. Sterile technique is not typically required for peripheral nerve blocks but should be used when accessing larger, more central structures such as the femoral nerve.

Peripheral nerves have a characteristic appearance when viewed by ultrasound and are usually easily identified, especially in the transverse orientation. They are hyperechoic (white) in appearance and are generally round or oval, although some may also appear more triangular (Fig. 31.US2). In larger nerves, the individual fascicles may be visible, especially when viewed with higher frequency. Nerve trunks (such as those used for scalene blocks) appear as rounded objects with a hypoechoic (darker gray) center (Fig. 31.US3). They may resemble blood vessels so take care to ensure that they are distinguished. This can be done by applying color flow Doppler and noting the absence of blood flow.
Once the nerve has been visualized, the “in-plane” technique is often the most useful to guide the needle to the selected area. Place the transducer in the transverse or slightly oblique plane relative to the nerve. Adjust the transducer so that the nerve is further away from the entry point of the needle (Fig. 31.US4). This will ensure that when the needle is inserted under ultrasound guidance, it can be “followed” as it advances toward the nerve in question. Once the image has been obtained, introduce the needle from the end of the transducer (Fig. 31.US5). Again, the entry point should be away from the nerve. Once the needle tip or needle is seen on screen, advance it toward the nerve. Once the tip of the needle is seen adjacent to the nerve, inject anesthetic under direct ultrasound guidance. The best results are usually obtained by injecting anesthetic in a pattern that surrounds the nerve in a concentric manner. This can be achieved by injecting under ultrasound guidance and then repositioning the tip of the needle.

Once the nerve has been visualized, the “in-plane” technique is often the most useful to guide the needle to the selected area. Place the transducer in the transverse or slightly oblique plane relative to the nerve. Adjust the transducer so that the nerve is further away from the entry point of the needle (Fig. 31.US4). This will ensure that when the needle is inserted under ultrasound guidance, it can be “followed” as it advances toward the nerve in question. Once the image has been obtained, introduce the needle from the end of the transducer (Fig. 31.US5). Again, the entry point should be away from the nerve. Once the needle tip or needle is seen on screen, advance it toward the nerve. Once the tip of the needle is seen adjacent to the nerve, inject anesthetic under direct ultrasound guidance. The best results are usually obtained by injecting anesthetic in a pattern that surrounds the nerve in a concentric manner. This can be achieved by injecting under ultrasound guidance and then repositioning the tip of the needle.

---

**ULTRASOUND BOX 31.1: Nerve Blocks of the Thorax and Extremities—cont’d**

**Figure 31.US1** High-frequency transducer.

**Figure 31.US2** Ultrasound image of a peripheral nerve (arrow), in this case the median nerve. Peripheral nerves are characterized by a brightly echogenic (white) texture and appear slightly fibrillar.

**Figure 31.US3** Ultrasound image of a nerve trunk (arrow), in this case a brachial plexus trunk. Nerve trunks appear similar to vascular structures, with a hypoechoic (light gray) area surrounded by a hyperechoic (white) wall. Applying color flow will aid in distinguishing the two.

**Figure 31.US4** Ultrasound image demonstrating movement of the transducer to place the target nerve away from the anticipated entry point of the needle. The path of the needle is approximated by the arrow.

**Figure 31.US5** Introducing the needle along the long axis of the transducer. Use of this technique will allow the sonographer to directly follow the course of the needle as it travels toward the nerve.

---

Continued
under ultrasound guidance until the nerve has been surrounded. The needle can also be inserted from the midpoint of the transducer, although this technique may cause more difficulty in following the tip of the needle.

Detailed descriptions of the anatomy and technique of the individual nerve blocks can be found throughout this chapter. However, it is important to discuss the nerve blocks typically performed under ultrasound guidance because the landmarks differ slightly.

**Interscalene Block**

The interscalene nerve blocks focuses on the trunks of the brachial plexus, specifically C5, C6, and C7. Blocking these trunks will provide anesthesia to most of the shoulder and upper extremity and spare the medial aspects of the arm and hand (these are innervated by the C8 and T1 nerve roots). This block is ideal for shoulder dislocations or complex lacerations of the upper extremity.

The trunks can be found grouped together in the neck and are typically easily identified with ultrasound. Because a number of critical structures are located near these trunks, using ultrasound to guide the injection will offer the physician increased confidence in the procedure, as well as increased success in the block. Several methods are described in the literature for localizing the nerve trunks. One of the most straightforward is to use the surrounding anatomy. Begin by placing the transducer, in the transverse orientation, lateral to the trachea at the level of the thyroid cartilage (Fig. 31.US6). Move the transducer laterally until the internal jugular vein and carotid artery are visualized (Fig. 31.US7). Once these vessels are seen, continue moving the transducer slightly laterally until the muscle bellies of the anterior and middle scalenes are visible. The border between the muscles may be subtle; however, shifting the transducer to a slightly oblique plane may help to better distinguish the anatomy. The trunks of the brachial plexus will be seen as rounded structures lying between the muscle bellies (Fig. 31.US8). They typically have an echogenic (white) border with a hypoechoic (dark gray) to anechoic (black) center. As noted previously, the nerve trunks can resemble blood vessels, so take care to evaluate the target structures before inserting the needle.
ULTRASOUND BOX 31.1: Nerve Blocks of the Thorax and Extremities—cont’d

The femoral nerve is easily identified in the femoral triangle accompanying the femoral vein and artery. A high frequency linear transducer is best suited for identification of these structures in most patients and should be placed transversely (or at a slight oblique angle) at the level of the inguinal crease. The vascular structures may be deeper than expected. Typically, the vein is the largest and most medial vessel, and compresses with gentle pressure. The femoral artery is typically more round, thicker walled, and less compressible. It is typically lateral to the vein. The femoral nerve is usually seen as a triangular, hyperechoic (white) structure.

FOREARM NERVE BLOCKS

Although blockade of the nerves that innervate the hand and wrist is typically performed at the wrist by using anatomic landmarks, it can also be performed under direct ultrasound guidance in the forearm. The median, radial, and ulnar nerves are easily identified with ultrasound, and direct visualization of anesthetic produces excellent results. Placing the transducer on the middle of the forearm in the transverse orientation will allow rapid identification (Fig. 31.US10). The median nerve is found in the center of the forearm, surrounded by the muscle belly. It is brightly echogenic (white) and usually has an oval or slightly triangular appearance (Fig. 31.US11). The radial nerve can be found toward the radial aspect of the forearm, adjacent to the radial artery and vein (Fig. 31.US12). The ulnar nerve is similarly found on the ulnar aspect of the forearm, adjacent to the ulnar artery and vein (Fig. 31.US13). The radial and ulnar nerves may appear smaller but have a similar echogenic, slightly triangular shape. Once the nerves have been visualized, the block can proceed as described previously.

LOWER EXTREMITY BLOCKS

The femoral nerve is easily blocked with ultrasound and gives anesthesia to the anterior thigh, knee, and hip joint. It is ideal for femur fractures and can be helpful in controlling pain in patients with comorbidities that make more aggressive systemic pain management difficult.

Figure 31.US9 Image of the brachial plexus trunks again highlighted in red. In this image, the potential space in the sheath around the trunks is highlighted in blue. Placement of an anesthetic agent in this potential space ensures the intended effect on the nerve trunks with minimal spread to adjacent structures.

Figure 31.US10 Placement of the transducer over the middle of the forearm to localize the median nerve. The transducer can then be shifted slightly to the right or left to locate the ulnar and radial nerves.

Figure 31.US11 Ultrasound image of the median nerve. Although the median nerve does not travel with an associated artery, it is easily located in the middle of the forearm and is characterized by its triangular appearance. Its hyperechoic (white) structure is highlighted by the arrow in this image.
structure lateral to the femoral artery (Fig. 31.US14). The vessels and nerve are bounded superiorly by the fascial planes of the fascia lata.

Once the nerve has been identified, advance a needle using the in-plane technique to infiltrate anesthesia to the surrounding area. Take care to keep track of the needle tip at all times to avoid injecting medication into the artery or vein. An amount of 15 to 20 cc of anesthesia is usually sufficient to achieve anesthesia to the femoral nerve. If the nerve is not immediately identified, but the femoral artery and vein are clearly seen, injecting a small amount of anesthesia to the area lateral to the femoral artery may cause the nerve to become more visible. Once the needle tip penetrates the fascial plane created by the fascia lata and is near the area where the nerve would be expected, placement of a small amount (approximately 1 to 2 cc) of anesthetic may make the nerve more visible. This may make the physician more confident in placing larger amounts of medication.

A variation of the femoral nerve block, the 3-in-1 block, utilizes a larger volume of medication in an attempt to obtain a greater area of anesthesia. The 3-in-1 block gives anesthesia to the area supplied by the femoral nerve, but also to the lateral thigh (lateral femoral cutaneous nerve) and the medial thigh (obturator nerve). These three nerves arise from the lumbar plexus and travel proximally within the same fascial compartment. The 3-in-1 block proceeds as described previously, but utilizes a volume of 20 to 30 cc of medication.

To perform the fascia iliaca block using ultrasound guidance, insert the needle in-plane just inferior to the inguinal ligament and guide it under the fascia iliaca, which lies just superior to the vessels. After aspiration, inject 30 to 40 cc of local anesthetic.

The tibial and common peroneal nerves can be blocked in the popliteal fossa to provide anesthesia to the distal part of the calf, ankle, and foot. Both are easily identified in the popliteal fossa, where they exist as separate structures. The transducer should be placed transversely in the popliteal fossa, and the popliteal artery should be sought (Fig. 31.US15). It is easily identified as a pulsatile, rounded, anechoic structure. Once the artery has been identified, the tibial nerve can be found slightly superficial (“high and outside”) to the artery (Fig. 31.US16). As with other peripheral nerves, the tibial nerve appears as an echogenic, rounded structure. Once the tibial nerve has
ULTRASOUND BOX 31.1: Nerve Blocks of the Thorax and Extremities—cont’d

Figure 31.US15 Placement of the ultrasound transducer in the transverse plane in the popliteal fossa to enable visualization of the tibial and peroneal nerves.

Figure 31.US16 Ultrasound image of the tibial nerve (arrow) viewed in transverse orientation. The popliteal vein can be seen just below the nerve and serves as a landmark for locating the nerve.

Figure 31.US17 Ultrasound image of the tibial and peroneal nerves. The peroneal nerve (arrow) is typically slightly more superficial than the tibial nerve (arrowhead).

has been found, it should be followed slightly proximally until the common peroneal nerve is identified. It is similar in appearance to the tibial nerve but is seen more superficially (Fig. 31.US17). Once both nerves have been identified, apply anesthetic under direct ultrasound guidance as described earlier.

Reference:

suggests that epinephrine-containing anesthetics are safe for digital nerve blocks, but some clinicians opt for epinephrine-free alternatives because of the theoretical risk for ischemic complications (see subsection on Complications and Precautions and Chapter 29).

As with upper extremity nerve blocks, note any neural or vascular injuries before the injection. The close proximity of these structures to the skin and bones means that they are frequently injured. Deficits, even if questionable, should be documented in the record and brought to the attention of the patient before performance of the nerve block.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
Clinical use of intravenous regional anesthesia (IVRA) has been well established as a safe, quick, and effective alternative to general anesthesia in selected cases requiring surgical manipulation of the upper and lower extremities. Though historically relegated to the operating room, the procedure is also readily applicable to outpatient use. Because of its reliability, safety, and ease of use, it is now commonly used in the emergency department (ED) and clinic. In the ED, the technique provides quick and complete anesthesia, muscle relaxation, and a bloodless operating field. The procedure is free from the troublesome side effects associated with other regional blocks, such as the axillary block. The procedure is easily mastered and has a very low failure rate; consistently good results can be expected. Although not a standard requirement of ED personnel, this technique can be safely used by trained ED clinicians, including physician's assistants and nurse practitioners, and does not have to be administered by an anesthesiologist. The first practical use of analgesia associated with the intravenous (IV) injection of a local anesthetic agent was described by August Gustav Bier in 1908. Colbern has since proposed the eponym Bier block (Video 32.1). Although the procedure has been in existence for many years, the need for special equipment and a safe anesthetic agent limited its use. However, the Bier block has now gained wide acceptance as a safe and effective procedure, and several papers extol its virtues. Even though complications do exist, no reported fatalities directly attributable to use of the Bier block with lidocaine have been reported. In this chapter, the techniques and complications are discussed according to their application in the ED.

INDICATIONS AND CONTRAINDICATIONS

Indications for IVRA include any procedure on the arm or leg that requires operating anesthesia, muscle relaxation, or a bloodless field, such as reduction of fractures and dislocations, repair of major lacerations, removal of foreign bodies, débridement of burns, and drainage of infection (Fig. 32.1). IVRA is commonly used for extremity surgery, such as carpal tunnel surgery or tendon repair. The procedure may be carried out on any patient of any age who is able to cooperate with the clinician.

The only absolute contraindications are allergy to the anesthetic agent and, possibly, uncontrolled hypertension. Relative contraindications include severe Raynaud’s disease, Buerger’s disease, or a crushed or already hypoxic extremity in which further transient ischemia would be detrimental. Homozygous sickle cell disease is a theoretical contraindication, but few data exist on the treatment of patients with this condition.

Intravenous Regional Anesthesia

**Indications**

Procedures of the arm or leg that require operating anesthesia, muscle relaxation, or a bloodless field:
- Reduction of fractures and dislocations
- Repair of major lacerations
- Removal of foreign bodies
- Débridement of burns
- Drainage of infection

**Contraindications**

**Absolute**
- Allergy to anesthetic agent
- Uncontrolled hypertension

**Relative**
- Severe Raynaud’s disease
- Buerger’s disease
- Crushed or already hypoxic extremity
- Procedures taking >90 minutes

**Complications**

- Lidocaine allergy (rare)
- Lidocaine toxicity
- Thrombophlebitis
- Tissue extravasation of anesthetic agent

**Equipment**

- 22-gauge IV catheter or 21-gauge butterfly needle
- 60-mL syringe
- Esmarch (elastic) bandage
- 1% lidocaine without epinephrine
- Sterile saline diluent
- Pneumatic tourniquet

**Review Box 32.1** Intravenous regional anesthesia: indications, contraindications, complications, and equipment. A double cuff tourniquet is shown. Not shown is a device required to provide continuous cuff pressure (see Fig. 32.2). Do not use a standard blood pressure cuff.
IVRA is best used for procedures requiring no more than 60 to 90 minutes of tourniquet time. Continuous cardiac or blood pressure monitoring is not standard and not required unless extenuating circumstances prognosticate potential cardiovascular problems. An uncooperative patient may delay the procedure rather than contraindicate it. The judicious use of standard ED sedation may facilitate the use of IVRA in an anxious or transiently uncooperative patient.

**EQUIPMENT**

The equipment required for IVRA consists of the following:

- 1% lidocaine (Xylocaine), without epinephrine, to be diluted to a 0.5% solution (Note: 1% lidocaine = 10 mg/mL; hence 1 mL of 1% lidocaine = 10 mg.)
- Clonidine, a parenteral opioid such as fentanyl, or ketorolac if used as additives
- Sterile saline solution as a diluent
- 50-mL syringe/18-gauge needle
- Pneumatic tourniquet (single or double cuff) such as the Zimmer A.T.S. 2000 Automatic Tourniquet System (Zimmer Inc., Bloomfield, CN) (Note: Do not use a standard blood pressure cuff.)
- IV catheters (20- or 22-gauge) or a 21-gauge butterfly needle
- Elastic bandage/Webril padding
- 500 mL of 5% dextrose in water (D₅W) and IV extension tubing (optional)

**PROCEDURE**

The procedure should be explained to the patient in advance. Premedication with midazolam (Versed), diazepam (Valium), or an opioid (e.g., morphine, fentanyl) may be helpful but need not be used routinely. The only painful portions of the procedure are establishment of the infusion catheter and exsanguination. The procedure should not be done on patients who are persistently uncooperative, intoxicated, or obtunded or who have had a previous reaction to a local anesthetic.

The patient need not be free of oral intake for a specific period before the procedure, but it is prudent to delay the procedure if the patient has just eaten a large meal. As a precaution, a suggested option is a large-bore catheter and an IV line with D₅W in the unaffected extremity. General resuscitation equipment, including suction, anticonvulsant drugs, bag-valve-mask (BVM) apparatus, and oxygen, should be available. Continuous cardiac and blood pressure monitoring is not routine but may be used as an option depending on the clinician’s assessment of the potential for cardiovascular events.

While the patient is being prepared, keep the lidocaine solution ready, but withhold it until the injured extremity is exsanguinated and the cuff is in place and inflated, as discussed later. The standard dose of lidocaine for the arm is 3 mg/kg. Inject it as a 0.5% solution (1% lidocaine mixed with equal parts sterile saline in a 50-mL syringe). Hence, for a 70-kg patient, infuse 210 mg of lidocaine (21 mL of 1% lidocaine) mixed with 21 mL of saline for a total volume in the infusing syringe of 42 mL of 0.5% lidocaine. Farrell and coworkers described a procedure termed the *minidose Bier block* in which 1.5 mg/kg of lidocaine is used and reported a 95% success rate. This lower dose may decrease the incidence of central nervous system side effects and is more desirable in the ED setting. Additional lidocaine may be infused if the initial dose is inadequate. Do not use lidocaine with epinephrine. Plain lidocaine is also available as a 0.5% solution and can therefore be used directly to avoid diluting the stronger solution. Some commercial preparations with preservatives are commonly used. Lidocaine 0.5% is also available.
prefer preservative-free lidocaine, but most clinicians use standard lidocaine with preservatives.

Apply a pneumatic tourniquet with cotton padding (to prevent ecchymosis) under the cuff proximal to the pathology. It is strongly advised that one not use a regular blood pressure cuff because these cuffs often leak or rupture and are not designed to withstand high pressure for any length of time. A specially designed portable double-cuff pneumatic system, such as that marketed by Zimmer Corporation, is ideal and preferred by the author.

Premix the anesthetic and saline solution in the syringe. Inflate the tourniquet and place a plastic catheter or a metal butterfly needle in a superficial vein as close as possible to the pathologic site as possible, and securely tape it in place (Fig. 32.3, step 1). It is usually desirable to use a vein on the dorsum of the hand, but importantly, the injection site should be at least 10 cm distal to the tourniquet to avoid injection of anesthetic proximal to or under the tourniquet. Keep the hub on the catheter to avoid backbleeding, or attach the syringe to the butterfly tubing. This catheter will be the route of injection of the anesthetic agent. Anesthesia from a fingertip-to-elbow direction seems to occur irrespective of the site of infusion of the anesthetic, but selecting an injection location near the site of pathology may provide more rapid anesthesia at a lower dosage.

Deflate the tourniquet used to obtain IV access, and exsanguinate the extremity so that when the anesthetic agent is injected, it will fill the drained vascular system. Exsanguination may be accomplished by either of two methods. Simple elevation of the extremity for a few minutes may be adequate, but wrapping the extremity in a distal-to-proximal direction with an elastic or Esmarch bandage, while being careful to not dislodge the infusion needle, significantly enhances exsanguination (see Fig. 32.3, step 2). Wrapping may be painful, so this step can be eliminated if it causes too much anxiety for the patient. If the wrapping procedure is not done, the extremity should be elevated for at least 3 minutes. During the wrapping procedure, take care to not dislodge or infiltrate the infusion catheter.

With the extremity still elevated or wrapped, inflate the tourniquet to 250 mm Hg (or 100 mm Hg above systolic pressure). Place the arm by the patient’s side, and remove the elastic exsanguination bandage (see Fig. 32.3, steps 3 to 5). In a child, inflate the tourniquet to 50 mm Hg above systolic pressure. In elderly obese patients with calcified peripheral vessels, arterial occlusion may not be achieved safely.13 In the leg, cuff pressure of 300 mm Hg or approximately twice the systolic pressure measured in the arms is suggested.

With the tourniquet now inflated, slowly inject the 0.5% lidocaine solution into the infusion catheter at the calculated dose (see Fig. 32.3, step 6). Note that the solution is placed in the arm in which the circulation is blocked, not in the precautionary keep-open IV line on the unaffected side. At this point, blotchy areas of erythema may appear on the skin. This is not an adverse reaction to the anesthetic agent but merely the result of residual blood being displaced from the vascular compartment, and it heralds success of the procedure.

In 3 to 5 minutes, the patient will experience paresthesia or warmth beginning in the fingertips and traveling proximally, with final anesthesia occurring above the elbow, to the level of the tourniquet. Complete anesthesia ensues in 10 to 20 minutes, followed by muscle relaxation. Note that adequate analgesia may exist even if the patient can still sense touch and position and has some motor function. If the “minidose” technique (initial dose of 1.5 mg/kg of lidocaine) does not provide adequate anesthesia, infuse an additional 0.5 to 1 mg/kg of diluted lidocaine at this time. As an example, for a 70-kg patient, an additional 0.5 mg/kg of lidocaine equals 35 mg lidocaine (3.5 mL of 1% lidocaine), and when the 1% lidocaine is diluted equally with saline, the final volume of the additional 0.5% lidocaine is 7 mL. Additional lidocaine was required in 7% of cases in one series in which the minidose regimen was used.14 The clinician should be patient, however, and wait a full 15 minutes before infusing additional lidocaine. Alternatively, if analgesia is slow or inadequate, an extra 10 to 20 mL of saline solution may be injected to supplement the total volume of solution to enhance the effect. Do not exceed a 3-mg/kg total dose of lidocaine. For obese patients, a maximum of 300 mg of lidocaine is suggested for the arm and no more than 400 mg for the leg. Data for very obese patients do not exist. Next, withdraw the infusing needle and tightly tape the puncture site to prevent extravasation of the anesthetic agent. Perform the surgical procedure or manipulation, including postreduction x-ray films and casting or bandaging (see Fig. 32.3, step 7).

On completion of the procedure, deflation of the tourniquet may be cycled to prevent a bolus effect of any lidocaine that may remain in the intravascular compartment. Deflate the cuff for 5 seconds and reinflate it for 1 to 2 minutes. Repeat this action two or three times. This is probably required only if the cuff has been inflated for less than 30 minutes. A single deflation is often performed if the cuff has been inflated for longer than this time.

If the tourniquet has been in place for less than 30 minutes, an increase in transient lidocaine-related side effects may be seen if cycled deflation has not been used because adequate tissue fixation of the lidocaine has probably not occurred. This may result in a higher peak plasma lidocaine level, with increased side effects. If the surgical procedure is completed rapidly and the 3-mg/kg limit of lidocaine has been infused, the tourniquet should remain inflated until 20 to 30 minutes has elapsed, and only then should it be deflated via the cycling technique. It is reasonable to use a 20-minute cutoff if the minidose technique is used or a total of 200 mg or less of lidocaine has been administered because this dose is equal to a commonly administered antiarrhythmic IV bolus.

Sensation returns quickly when the tourniquet is removed, and in 5 to 10 minutes the extremity returns to its pre-anesthetic level of sensation and function. Many patients describe a transient intense tingling sensation after cuff deflation. If the procedure takes longer than 20 or 30 minutes, many patients complain of pain from the tourniquet because it is not inflated over an anesthetized area. Use of a double-cuff tourniquet may alleviate the problem of pain under the cuff. A wide tourniquet cuff (14 cm) is less painful than a narrow tourniquet (7 cm) when the cuff is inflated 10 mm Hg above loss of the arterial pulse.15 The reason for pain under the tourniquet is unknown, but this can be a limiting factor because most patients begin to feel significant discomfort after 30 minutes if only a single cuff is used. Tourniquet pain can be significantly reduced and tourniquet time extended by adding ketorolac to the lidocaine anesthetic (see later).

In the preferred double-cuff system, two separate tourniquets are placed side by side on the extremity. One is termed the proximal cuff and the other the distal cuff. The proximal cuff is inflated at the beginning of the procedure, and anesthesia is obtained under the deflated distal cuff. When the patient begins to feel pain under the proximal cuff, the distal cuff is...
Inflate the tourniquet to 250 mm Hg or 100 mm Hg above systolic pressure. In the leg, inflate the cuff to 300 mm Hg or twice the systolic pressure measured in the arm.

Exsanguinate the extremity by elevating and wrapping it in a distal-to-proximal fashion. Here, an Esmarch bandage is being used.

Inflate the tourniquet to 250 mm Hg or 100 mm Hg above systolic pressure. In the leg, inflate the cuff to 300 mm Hg or twice the systolic pressure measured in the arm.

Slowly inject the 0.5% lidocaine solution into the infusion catheter at the calculated dose. See text for details and dosing information.

Once the procedure is complete, deflate the tourniquet in a cycling fashion (deflate for 5 seconds, reinflate for 1 to 2 minutes) 2 or 3 times. Then remove the tourniquet.

Figure 32.3 Intravenous regional anesthesia. A double cuff tourniquet is depicted in the figures. The procedure depicted demonstrates the use of a single tourniquet; refer to the text for details regarding the use of a double tourniquet.
MECHANISM OF ACTION

Some of the anesthesia is undoubtedly related to the ischemia produced by the tourniquet, but most of the anesthesia is secondary to the anesthetic agent itself. Although the exact mechanism by which anesthesia is produced is unknown, the site of action of the anesthetic may be at sensory nerve endings, neuromuscular junctions, or major nerve trunks.13 Contrast-enhanced studies have demonstrated that the anesthetic agent does not diffuse throughout the entire arm, yet anesthesia of the entire limb is obtained. For example, when the anesthetic agent is injected into the elbow and kept in that region with the entire limb is obtained. For example, when the anesthetic does not diffuse throughout the entire arm, yet anesthesia of enhanced studies have demonstrated that the anesthetic agent reported.

The primary reason that lidocaine is used is due to its relatively short duration of action. Peak plasma lidocaine concentrations after release of the tourniquet decrease as the time of vascular occlusion (tourniquet time) increases. If the tourniquet is inflated for at least 30 minutes and the deflation-reinflation technique is used when the procedure is finished, the plasma concentration of lidocaine should be approximately 2 to 4 µg/mL, below the 5- to 10-µg/mL level at which dose-related lidocaine reactions occur.14 Tucker and Boas demonstrated a peak plasma lidocaine level of 10.3 µg/mL after a 10-minute period of vascular occlusion,15 as opposed to 2.3 µg/mL if the tourniquet was inflated for 45 minutes.

First dilute solutions of lidocaine are associated with lower peak lidocaine levels. When equal doses of lidocaine are used, peak arterial plasma levels are 40% lower if the 0.5% solution is used than if the 1% solution is used.16 For example, after 10
minutes of vascular occlusion, the peak plasma concentration of lidocaine has been demonstrated to reach 10.3 µg/mL with the 1% solution versus only 5.6 µg/mL when the drug was given under similar circumstances at a 0.5% concentration.22

**Exsanguination**

Many clinicians consider exsanguination of the extremity before injection of the anesthetic agent to be essential for success. Others do not believe that it is a critical factor. Exsanguination by simple elevation of the extremity should be done in all cases, but in certain instances one should consider avoiding the painful wrapping of the extremity with an elastic or Esmarch bandage. (Note that applying an Esmarch wrap over a fracture site is usually quite painful.) A pneumatic splint, such as the type used for prehospital immobilization, is also a reasonable alternative to painful wrapping. The process of exsanguination is believed to allow better vascular diffusion of the anesthetic.

**Site of Injection**

Anesthesia is usually achieved no matter where the local anesthetic is injected, but some evidence indicates that the procedure is more successful when the anesthetic is injected distally. Sorbie and Chacha noted the following failure rates associated with specific sites of anesthetic injection23: antecubital fossa, 23%; middle of the forearm or leg, 18%; and hand, wrist, or foot, 4%. In most cases a vein in the dorsum of the hand or foot is used. If local pathology precludes use of the hand, midforearm or antecubital fossa of the elbow is an acceptable, albeit less desirable, alternative as long as the infusion catheter is well below the tourniquet to avoid systemic injection.

Although most of the literature stresses use of this technique on the upper extremity, it may also be used successfully on the leg. It cannot, however, be used for procedures above the knee. Tourniquet pain appears to be a limiting factor when the procedure is used on the leg. One must be certain to avoid damage to the peroneal nerve by using the tourniquet in the midcalf area only.

**COMPLICATIONS**

Although IVRA is both safe and simple, one should not be lulled into complacency because complications do occur and are usually related to equipment failure or mistakes in technique.

**Anesthetic Agent**

Serious complications seldom occur if proper attention is paid to technique. True lidocaine allergy is very rare. Other reactions to lidocaine are rare and are usually systemic reactions from high blood levels.8,18,24 High levels may result from miscalculation of dosages, from too rapid release of the tourniquet before the anesthetic has become fixed in tissue ("bolus effect"), or rarely, from advancement of the infusion catheter proximal to the tourniquet, thereby resulting in direct IV infusion.25 To emphasize the safety of this procedure, note that the dose of lidocaine used in the minidose technique is similar to an IV bolus routinely given to patients with significant cardiovascular disease in the presence of ventricular dysrhythmias.

Generally, the central nervous system effects of lidocaine are minor and result in mild reactions such as dizziness, tinnitus, lethargy, headache, or blurred vision. This should not occur in more than 2% to 3% of patients and requires no treatment.8 Transient hypotension and bradycardia may develop but are generally self-limited. Convulsions may occur but are extremely rare.

The most common complication related to the anesthetic agent is rapid systemic vascular infusion, which occurs when a blood pressure cuff explodes or slowly leaks, with consequent loss of anesthesia and high blood levels.26 Similar complications may occur if the cuff is deflated earlier than 20 to 30 minutes after the induction of anesthesia. Both complications are the result of a bolus effect of the anesthetic, similar to an IV injection.

Van Neikerk and Tonkin reported three seizures in a series of 1400 patients.28 Auroy and colleagues reported 23 seizures in 11,229 cases,7 with no cardiac arrest or fatalities, and deemed IVRA safer than general anesthesia or peripheral nerve blocks. Seizures are generally not recurrent and are treated with oxygen and anticonvulsant drugs. Transient cardiovascular reactions, such as bradycardia and hypotension, are possible with large doses of lidocaine. Vasovagal reactions do occur. If resuscitation equipment is available and a precautionary IV line is started in the opposite arm, there should not be any serious sequelae.

One case of cardiac arrest 15 seconds after the use of 200 mg lidocaine has been reported, but the actual clinical scenario may have been a vasovagal reaction rather than a true cardiac arrest.28

**Additional Complications**

Thrombophlebitis can occur following the IV administration of anesthetics, and the formation of insignificant amounts of methemoglobin with the use of prilocaine hydrochloride (Citanest) has been reported.29 Methemoglobinemia can also theoretically occur with lidocaine but has not been reported. Bupivacaine offers no benefit over lidocaine, has been associated with deaths, and should be avoided.20

A particularly bothersome problem is infiltration or dislodgement of the infusion catheter during exsanguination and tissue extravasation of the anesthetic agent. In addition, some leakage of anesthetic can occur after the infusion needle has been removed. Both problems may result in poor anesthesia but may be minimized if a small, well-secured plastic infusion needle is used instead of a metal scalp vein ("butterfly") needle and if the puncture site is tightly taped after withdrawal of the catheter.

This procedure cannot be used for manipulations or operations in which the pulse must be monitored as a guide to reduction (e.g., supracondylar fractures of the humerus) because the tourniquet occludes arterial flow. Use of the Bier block in patients with sickle cell disease is not well documented. It should be used with caution until the ischemic effect of the tourniquet on the red blood cells of such patients has been clarified. In all patients, tourniquet time should not exceed 90 minutes. Ischemia for less than this amount of time is not associated with serious sequelae.

An excellent summary of a very favorable experience with IVRA of both the arm and leg in 1900 outpatients is available.21
REFERENCES

Systemic Analgesia and Sedation for Procedures

Kenneth Deitch

Procedural sedation and analgesia (PSA) refers to the use of analgesic, dissociative, and sedative agents to relieve the pain and anxiety associated with diagnostic and therapeutic procedures. PSA is an integral element of emergency medicine residency and pediatric emergency medicine fellowship training and curricula, and graduates of these programs are skilled in the practice of PSA. Moreover, by virtue of their training, emergency clinicians are proficient in resuscitation, vascular access, and advanced airway management, which permits them to effectively recognize and manage the potential complications associated with PSA.1

In a study of all practitioners, the most common clinical errors associated with PSA were delayed recognition of respiratory depression and arrest, inadequate monitoring, and inadequate resuscitation,2 mistakes that are unlikely to be made by emergency clinicians. The safety of PSA techniques by emergency clinicians has been well documented in numerous series in both children and adults.3-2 Safe and successful application of PSA requires careful patient selection, customization of therapy to the specific needs of the patient, and careful monitoring of patients for adverse events. Emergency clinicians must ensure that all patients receive pain relief and sedation commensurate with their individual needs during any procedure.

TERMINOLOGY

The progression from minimal sedation to general anesthesia is a nonlinear continuum that does not lend itself to division into arbitrary stages. Low doses of opioids or benzodiazepines induce mild analgesia or sedation, respectively, with little danger of adverse events. If, however, clinicians continue administering additional medication beyond this initial level, progressively altered consciousness ensues with a proportionately increased risk for respiratory and airway complications. If further medications are administered, the patient will advance along this continuum until protective airway reflexes are lost and general anesthesia is ultimately reached. This continuum of sedation is not drug specific in that varying states from mild sedation to general anesthesia can be achieved with virtually all nondissociative PSA agents (e.g., opioids, benzodiazepines, barbiturates, etomidate, propofol).3

In 1985, the American Academy of Pediatrics (AAP) and the National Institutes of Health issued guidelines for the management and monitoring of children receiving sedation for diagnostic and therapeutic procedures in response to the growing use of opioids and sedative-hypnotic agents in the outpatient setting and a number of sedation-related deaths.4,5 In these documents, three levels of sedation were defined (conscious sedation, deep sedation, and general anesthesia) to create a common language for describing drug-induced alterations in consciousness (Box 33.1).6,9 A key development in the field of PSA has been revision of the original terminology and adoption of clearer descriptions of varying types and degrees of sedation (see Box 33.1). Though historically popular, the widely misinterpreted and misused term “conscious sedation” has fallen into disfavor;7 it has been labeled as “confusing,”8 “imprecise,”9 and an “oxymoron”8,9 and has been replaced with the term “moderate sedation.”10

Despite improvements in PSA terminology, the system is imperfect and there is still no objective way to assess the depth of sedation. Levels of responsiveness remain at best crude surrogate markers of respiratory drive and retention of protective airway reflexes. This is especially true for all levels of sedation in young children (infants and toddlers) who do not understand or are unreliable in following verbal commands. Although respiratory depression and respiratory arrest can be detected quickly with standard interactive and mechanical monitoring, there is no safe and practical way to assess the status of protective airway reflexes. Data are currently insufficient to determine whether deep sedation is associated with impairment of protective reflexes or whether such danger is encountered only when “pushing” deep sedation to the point at which it approaches or reaches general anesthesia.

PSA GUIDELINES

Before the promulgation of PSA guidelines by specialty societies and governmental agencies, clinicians simply administered sedatives in varied clinical settings and used individual judgment to determine the need for specific monitoring devices and supporting personnel. Since 1985, at least 27 sets of PSA guidelines have been published,11 each crafted for the unique and differing settings in which PSA is practiced. Naturally, not all are in agreement.1 The intent of each of these guidelines is to better standardize the manner in which PSA is performed to enhance patient safety. Those most pertinent to emergency clinicians are from the American College of Emergency Physicians (ACEP),1 the AAP,12 and the American Society of Anesthesiologists (ASA).16,17

In the early 1990s, the Joint Commission on Accreditation of Healthcare Organizations, an independent, not-for-profit organization that evaluates and accredits hospitals in the United States, took a special interest in PSA, with the central theme being that the standard of sedation care provided should be comparable throughout a given hospital. Thus, patients sedated in the emergency department (ED) should not receive a significantly different level of attention or monitoring than those sedated for a similar-level procedure in the operating room or in the endoscopy suite. To ensure this, the Joint Commission requires specific PSA protocols to be applied consistently throughout each institution. These hospital-wide sedation policies will vary from site to site based on the specific needs and expertise available within each institution. In 2001, the Joint Commission released new standards for pain management, sedation, and anesthesia care.19,10

At each hospital accreditation survey the Joint Commission determines whether practitioners practice PSA consistent with their hospital-wide sedation policy and whether they provide sufficient documentation of such compliance. Clinicians must be familiar with their hospital’s sedation policies and should work with their medical staff to ensure that such policies are
that hospital policies governing the provision of anesthesia
must adhere to these guidelines and provide supporting
documentation when requested. In 2011, CMS issued revised
guidelines to help clarify various provisions of their anesthesia
services (including PSA) must be based on nationally recognized
guidelines; the source of these guidelines may include a number
of specialty organizations, including ACEP. CMS also notes
that “The ED is a unique environment where patients present
on an unscheduled basis with often very complex problems that
may require several emergent or urgent interventions to proceed
simultaneously to prevent further morbidity or mortality,” and
that, “…emergency medicine-trained physicians have very
specific skill sets to manage airways and provide the ventilation
necessary for patient rescue. Therefore these practitioners are
uniquely qualified to provide all levels of analgesia/sedation
and anesthesia (moderate to deep general).”

EVALUATION BEFORE PSA

The practice of PSA has three essential components performed
in sequence: the initial presedation evaluation, sedation during
the procedure, and postprocedure recovery and discharge from
the ED. In all but the most emergency situations, perform a
directed history and physical examination before PSA. If this
evaluation suggests additional risk, reconsider the advisability
of sedation. High-risk cases may be better managed in the
more controlled environment of the operating room.

Presedation assessment is a Joint Commission requirement,
and most hospitals have developed specific forms to facilitate
consistent documentation of the required items. In general,
however, all the appropriate parameters are already documented
in the general ED record or are obvious by simply evaluating
the patient’s complaint. Except in emergency situations, discuss
the risks, benefits, and limitations of any PSA with the patient
suitably detailed, yet reasonable and realistic. Unduly restrictive
policies do a disservice to patients by discouraging appropriate
levels of analgesia and sedation. Most hospitals pattern their
sedation policies after the Joint Commission standards and
definitions. It is important to note that the unique ketamine
dissociative state does not fit into the existing Joint Commis-
sion definitions of sedation and anesthesia. A ready solution
is to assign a distinct definition for “dissociative sedation”
(see Box 33.1).

The Joint Commission requires that PSA practitioners who
are permitted to administer deep sedation be qualified to rescue
patients from general anesthesia. Emergency clinicians typically
perform all levels of sedation except general anesthesia. Moder-
ate sedation suffices for the majority of procedures in adults
and cooperative children, although it will not be adequate for
extremely painful procedures (e.g., hip reduction, cardioversion).
Deep sedation can facilitate such procedures, but with greater
risk for cardiorespiratory depression than is the case with
moderate sedation. Moderate sedation is frequently insufficient
for effective anxiolysis and immobilization in younger, frightened
children, and deep or dissociative sedation is an appropriate
alternative.

The Centers for Medicare & Medicaid Services (CMS) has
published similar requirements for hospital anesthesia services
that also includes the provision of PSA throughout a given
institution. Health care institutions wishing to bill Medicare
must adhere to these guidelines and provide supporting
documentation when requested. In 2011, CMS issued revised
guidelines to help clarify various provisions of their anesthesia
services requirements. The new CMS guidelines emphasize
that hospital policies governing the provision of anesthesia

suitably detailed, yet reasonable and realistic. Unduly restrictive
policies do a disservice to patients by discouraging appropriate
levels of analgesia and sedation. Most hospitals pattern their
sedation policies after the Joint Commission standards and
definitions. It is important to note that the unique ketamine
dissociative state does not fit into the existing Joint Commis-
sion definitions of sedation and anesthesia. A ready solution
is to assign a distinct definition for “dissociative sedation”
(see Box 33.1).

The Joint Commission requires that PSA practitioners who
are permitted to administer deep sedation be qualified to rescue
patients from general anesthesia. Emergency clinicians typically
perform all levels of sedation except general anesthesia. Moder-
ate sedation suffices for the majority of procedures in adults
and cooperative children, although it will not be adequate for
extremely painful procedures (e.g., hip reduction, cardioversion).
Deep sedation can facilitate such procedures, but with greater
risk for cardiorespiratory depression than is the case with
moderate sedation. Moderate sedation is frequently insufficient
for effective anxiolysis and immobilization in younger, frightened
children, and deep or dissociative sedation is an appropriate
alternative.

The Centers for Medicare & Medicaid Services (CMS) has
published similar requirements for hospital anesthesia services
that also includes the provision of PSA throughout a given
institution. Health care institutions wishing to bill Medicare
must adhere to these guidelines and provide supporting
documentation when requested. In 2011, CMS issued revised
guidelines to help clarify various provisions of their anesthesia
services requirements. The new CMS guidelines emphasize
that hospital policies governing the provision of anesthesia

suitably detailed, yet reasonable and realistic. Unduly restrictive
policies do a disservice to patients by discouraging appropriate
levels of analgesia and sedation. Most hospitals pattern their
sedation policies after the Joint Commission standards and
definitions. It is important to note that the unique ketamine
dissociative state does not fit into the existing Joint Commis-
sion definitions of sedation and anesthesia. A ready solution
is to assign a distinct definition for “dissociative sedation”
(see Box 33.1).

The Joint Commission requires that PSA practitioners who
are permitted to administer deep sedation be qualified to rescue
patients from general anesthesia. Emergency clinicians typically
perform all levels of sedation except general anesthesia. Moder-
ate sedation suffices for the majority of procedures in adults
and cooperative children, although it will not be adequate for
extremely painful procedures (e.g., hip reduction, cardioversion).
Deep sedation can facilitate such procedures, but with greater
risk for cardiorespiratory depression than is the case with
moderate sedation. Moderate sedation is frequently insufficient
for effective anxiolysis and immobilization in younger, frightened
children, and deep or dissociative sedation is an appropriate
alternative.

The Centers for Medicare & Medicaid Services (CMS) has
published similar requirements for hospital anesthesia services
that also includes the provision of PSA throughout a given
institution. Health care institutions wishing to bill Medicare
must adhere to these guidelines and provide supporting
documentation when requested. In 2011, CMS issued revised
guidelines to help clarify various provisions of their anesthesia
services requirements. The new CMS guidelines emphasize
that hospital policies governing the provision of anesthesia

or parent or guardian in advance and obtain verbal agreement. Formal written informed consent is not required as a standard of care (but may be an institutional requirement), although documentation, as discussed earlier, is essential.

**General**

Assess the type and severity of any underlying medical problems. This is usually best documented by the standard ED medical record, history, physical examination, and nursing notes. Another tool used for this purpose is the ASA physical status classification for preoperative risk stratification (Table 33.1). Verify current medications and allergies and inquire about previous adverse experiences with PSA or anesthesia.

**Airway**

Inspect the airway to determine whether any abnormalities (e.g., severe obesity, short neck, small mandible, large tongue, trismus) are present that might impair airway management. Consider assessments such as Mallampati scoring or the distance between the chin and hyoid bone (see Chapter 4; Fig. 4.3).

**Cardiovascular**

Perform cardiac auscultation to assess for disturbances in rhythm or other abnormalities. In patients with known cardiovascular disease, evaluate their degree of reserve because most PSA agents can cause vasodilatation and hypotension.

**Respiratory**

Perform lung auscultation to assess for active pulmonary disease, especially obstructive lung disease and upper respiratory infections that may predispose the patient to airway reactivity.

**Gastrointestinal**

Assess the time and nature of the last oral intake because pulmonary aspiration of gastric contents may occur if the patient vomits when protective airway reflexes are impaired. Fig. 33.1 shows a four-step assessment tool to stratify the risk for aspiration before sedation and to identify prudent limits of targeted sedation, although this tool has not yet been validated.

More conservative guidelines from the ASA for elective surgery or procedures in healthy patients specify an age-stratified fasting requirement of 2 to 3 hours for clear liquids and 4 to 8 hours for solids and nonclear liquids. Nonetheless, they acknowledge that, regarding PSA, “the literature provides insufficient data to test the hypothesis that preprocedure fasting results in a decreased incidence of adverse outcomes.” Since these guidelines were published, multiple ED studies comparing fasting times in both pediatric and adult patients undergoing PSA failed to demonstrate a significant difference in the rates of emesis or aspiration and no other serious adverse events resulting from emesis or aspiration were identified.

The concept of preprocedure fasting is logistically difficult or impossible for emergency clinicians, who have no control over patients’ oral intake before arrival at the ED. In actual practice, emergency clinicians routinely perform PSA safely on patients who are noncompliant with the ASA elective-procedure fasting guidelines. Procedures can sometimes be delayed for several hours; however, this must be balanced against prolongation of pain and anxiety in the patient, inconvenience for the patient and family, and expenditure of room space and other finite ED resources. In addition, many ED procedures require urgent if not immediate attention (e.g., débridement and repair of animal bite wounds, acute burn management, arthrocentesis for suspected septic arthritis, reduction of joint dislocations, lumbar puncture in an uncooperative septic patient, hernia reduction, eye irrigation for ocular trauma or chemical burns, cardioversion in a hemodynamically unstable patient). Though uncommon, there may be occasions in which nonfasting patients require urgent procedures with a substantial depth of sedation that may be more safely managed in the operating room with endotracheal intubation to protect the airway.

Selecting agents that are less likely to produce vomiting, such as fentanyl instead of morphine or meperidine, may decrease the potential for aspiration. Concomitant antiemetic

---

**TABLE 33.1 ASA Physical Status Classification**

<table>
<thead>
<tr>
<th>ASA CLASS</th>
<th>DESCRIPTION</th>
<th>EXAMPLES</th>
<th>SUITABILITY FOR SEDATION</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Normal healthy patient</td>
<td>Unremarkable past medical history</td>
<td>Excellent</td>
</tr>
<tr>
<td>2</td>
<td>Patient with mild systemic disease—no functional limitation</td>
<td>Mild asthma, controlled seizure disorder, anemia, controlled diabetes mellitus</td>
<td>Generally good</td>
</tr>
<tr>
<td>3</td>
<td>Patient with severe systemic disease—definite functional limitation</td>
<td>Moderate to severe asthma, poorly controlled seizure disorder, pneumonia, poorly controlled diabetes mellitus, moderate obesity</td>
<td>Intermediate to poor; consider benefits relative to risks</td>
</tr>
<tr>
<td>4</td>
<td>Patient with severe systemic disease that is a constant threat to life</td>
<td>Severe bronchopulmonary dysplasia; sepsis; advanced degrees of pulmonary, cardiac, hepatic, renal, or endocrine insufficiency</td>
<td>Poor; benefits rarely outweigh risks</td>
</tr>
<tr>
<td>5</td>
<td>Moribund patient not expected to survive without the operation</td>
<td>Septic shock, severe trauma</td>
<td>Extremely poor</td>
</tr>
</tbody>
</table>

ASA, American Society of Anesthesiologists.

Higher-risk patient

<table>
<thead>
<tr>
<th>Oral intake in the prior 3 hours</th>
<th>Procedural urgency(^b)</th>
<th>Procedural sedation and analgesia targeted depth and duration</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Emergency procedure</td>
<td>Urgent procedure</td>
</tr>
<tr>
<td>Nothing</td>
<td>All levels of sedation</td>
<td>All levels of sedation</td>
</tr>
<tr>
<td>Clear liquids only</td>
<td>All levels of sedation</td>
<td>All levels of sedation</td>
</tr>
<tr>
<td>Light snack</td>
<td>All levels of sedation</td>
<td>Up to and including brief deep sedation</td>
</tr>
<tr>
<td>Heavier snack or meal</td>
<td>All levels of sedation</td>
<td>Up to and including extended moderate sedation</td>
</tr>
</tbody>
</table>

Higher-risk patients are those with one or more of the following present to a degree individually or cumulatively judged clinically important by the treating clinician:

- Potential for difficult or prolonged assisted ventilation should an airway complication develop (e.g., short neck, small mandible/micrognathia, large tongue, tracheomalacia, laryngomalacia, history of difficult intubation, congenital anomalies of the airway and neck, sleep apnea)
- Conditions predisposing to esophageal reflux (e.g., elevated intracranial pressure, esophageal disease, hiatal hernia, peptic ulcer disease, gastritis, bowel obstruction, ileus, tracheoesophageal fistula)
- Extremes of age (e.g., > 70 years or < 6 months)
- Severe systemic disease with definite limitation in function (i.e., American Society of Anesthesiologists physical status ≥ 3)
- Other clinical findings leading the emergency physician to judge the patient to be at higher than standard risk (e.g., altered level of consciousness, frail appearance)

B. Procedural urgency:

- Emergency (e.g., cardioversion for life-threatening dysrhythmia, reduction of a markedly angulated fracture or dislocation with soft tissue or vascular compromise, intractable pain or suffering)
- Urgent (e.g., care of dirty wounds and lacerations, animal and human bites, incision and drainage of abscesses, fracture reduction, hip reduction, lumbar puncture for suspected meningitis, arthrocentesis, neuroimaging for trauma)
- Semiurgent (e.g., care of clean wounds and lacerations, shoulder reduction, neuroimaging for new-onset seizures, removal of foreign bodies, examination for sexual assault)
administration is an unproven adjunct but a common consideration. In summary, common sense should apply and clinical judgment should prevail, but it is standard for PSA to be performed in the ED on patients in the nonfasting state.

Hepatic and Renal

The implications of delayed metabolism or excretion of PSA agents in infants younger than 6 months, in the elderly, and in patients with hepatic or renal abnormality should be considered.

PERSONNEL AND INTERACTIVE MONITORING

The most important element of PSA monitoring is close and continuous observation of the patient by an individual capable of recognizing complications of sedation (Fig. 33.2). This person must be able to continuously observe the patient's face, mouth, and chest wall motion. Equipment or sterile drapes must not interfere with such visualization. Such careful observation allows prompt detection of adverse events such as respiratory depression, apnea, partial airway obstruction, emesis, and hypersalivation.

PSA personnel should understand the pharmacology of analgesic and sedative agents and their respective reversal agents. They must be proficient in maintaining airway patency and assisting ventilation if needed. PSA requires a minimum of two experienced individuals, most frequently one clinician and one nurse or respiratory therapist. The clinician typically oversees drug administration and performs the procedure, whereas the nurse or respiratory therapist continuously monitors the patient for potential complications. The nurse or respiratory therapist should also document the medications administered and the response to sedation and measure vital signs periodically. The nurse or respiratory therapist may assist in minor, interruptible tasks but must remain focused on the patient's cardiopulmonary status, and this responsibility must not be impaired. An individual with advanced life support skills should also be immediately available, which is a requisite easy to fulfill in the ED setting.

During deep sedation, the individual dedicated to patient monitoring should have experience with this depth of sedation and no other responsibilities that would interfere with the advanced level of monitoring and documentation appropriate for this degree of sedation. Individual hospital-wide sedation policies may have additional requirements regarding how and when deep sedation is administered based on the patient's specific needs and the clinician's expertise.

It is not mandatory to have intravenous (IV) access in situations in which sedation is administered by the intramuscular (IM), oral, nasal, inhalational, or rectal routes, but it may be preferable based on the anticipated depth of sedation, comorbid conditions, or additional drug titration. When sedation is performed without IV access, an individual skilled in initiating such access should be immediately available.

EQUIPMENT AND MECHANICAL MONITORING

The routine use of mechanical monitoring has greatly enhanced the safety of PSA. With current technology, oxygenation (via pulse oximetry), ventilation (via capnography), and hemodynam-ics (via blood pressure and electrocardiogram [ECG]) can all be monitored noninvasively in nonintubated, spontaneously breathing patients.

Pulse Oximetry

Mechanical monitoring for PSA should include continuous pulse oximetry with an audible signal. Pulse oximetry measures the percentage of hemoglobin that is bound to oxygen. If the patient is breathing room air (21% oxygen), pulse oximetry will detect a decrease in alveolar ventilation rather quickly. Most patients undergoing procedural sedation in the ED are given supplemental oxygen by nasal cannula. With preoxygenation or the continued use of supplemental oxygenation during a procedure, the pulse oximetry will take significantly longer to drop despite the complete absence of ventilation. With preoxygenation it may take 4 to 5 minutes of apnea before the pulse oximetry will drop significantly. Hence, pulse oximetry is not a substitute for monitoring ventilation because of the variable lag time between the onset of hypventilation or apnea and a change in the oxygen saturation of hemoglobin molecules.

Capnography

Capnography is a very useful tool that provides a continuous, breath-by-breath measure of the respiratory rate and CO₂ exchange. Capnography readings are not affected by the presence or absence of additional oxygen. When sedated patients begin to hypventilate, the end-tidal CO₂ will rise. When apnea occurs, the end-tidal CO₂ drops to zero. Importantly, capnography can detect the common adverse airway and respiratory events associated with PSA. Capnography detects the earliest airway or respiratory compromise and will show abnormally high or low end-tidal CO₂ pressure well before pulse oximetry detects falling oxyhemoglobin saturation, especially in patients receiving supplemental oxygen. Early detection of respiratory compromise is especially important in infants and toddlers, who have smaller functional residual capacity and greater oxygen consumption than older children and adults. Capnography provides a non–impedance-based respiratory rate directly from the airway (via an oral-nasal cannula). This is more accurate than impedance-based respiratory monitoring, especially in patients with obstructive apnea or laryngospasm, in whom impedance-based monitoring will interpret chest wall movement without ventilation as a valid breath.

Multiple studies of patients undergoing PSA in a variety of clinical settings have demonstrated that capnography identifies respiratory depression earlier and more frequently than standard monitoring techniques (e.g., pulse oximetry, observation). Moreover, several additional studies have demonstrated that the use of capnography reduces the incidence of hypoxic events in both children and adults. However, there is no evidence that capnography reduces the incidence of serious adverse events such as neurologic injury (secondary to hypoxia), aspiration, or death. An additive role for capnography in cases of PSA in the ED has not been proven. As a result, capnography is not an agreed upon standard for procedural sedation in the ED. The American College of Emergency Physician's current clinical policy on PSA gives the use of capnography a level B recommendation (recommendations for patient care that may identify a particular strategy or range of strategies that reflect moderate clinical
After a thorough presedation assessment, attach the patient to the monitoring system. Routine use of pulse oximetry, capnography, ECG monitoring, and blood pressure monitoring greatly enhances the safety of PSA.

Apply supplemental oxygen, especially for patients undergoing deep sedation with agents such as propofol. In addition to pulse oximetry, observe the rise and fall of the chest, for the earliest indication of apnea. Ideally, capnography should be used in all situations in which high-flow oxygen is administered.

Ensure that all necessary age-appropriate resuscitation equipment is available, including oxygen, a bag-valve-mask device, suction, reversal agents, and a defibrillator (for patients with significant cardiovascular disease).

Administer the agent selected. The choice of appropriate medications is discussed in detail in the text. Here, propofol is being administered.

PSA requires a minimum of two experienced individuals: one to perform the procedure and one to continuously monitor the patient for potential complications. The person monitoring the patient must be focused on the patient’s cardiopulmonary status.

Perform the procedure. Here, the patient is being cardioverted for new-onset atrial fibrillation. After the procedure, monitor the patient until there is no further risk for cardiopulmonary depression. Before discharge, be sure that the patient is alert and oriented with stable vital signs.

Figure 33.2 Procedural sedation and analgesia (PSA). ECG, electrocardiographic.
certainty [i.e., based on evidence from one or more Class of Evidence II studies or strong consensus of Class of Evidence III studies]), stating, “Capnography may be used as an adjunct to pulse oximetry and clinical assessment to detect hypoventilation and apnea earlier than pulse oximetry and/or clinical assessment alone in patients undergoing procedural sedation and analgesia in the ED.”

**ECG Monitoring**

Although continuous ECG monitoring cannot be considered mandatory or standard of care in the absence of cardiovascular disease, such monitoring is simple, inexpensive, and readily available.

**BIS Monitoring**

The bispectral index (BIS) is a monitoring modality that uses a processed electroencephalogram signal to quantify the depth of anesthesia or sedation. A BIS value of 100 (unitless scale) is considered complete alertness, 0 represents no cortical activity at all, and the range of 40 to 60 is believed to be consistent with general anesthesia. Although this technology has been used widely to monitor the depth of sedation in the operating room, the ASA has judged that its clinical applicability for this purpose “has not been established.” Furthermore, a 2011 study found that patients in whom a modified minimum alveolar concentration protocol (i.e., the inhalational anesthetic concentration needed for 50% of patients to not move with the application of a noxious stimulus) was used had fewer awareness events than did those in whom a BIS protocol was used. Even though PSA research has demonstrated statistical associations between BIS and standard sedation scores, these studies have also noted unacceptably wide ranges of BIS values at various depths of sedation, Thus, although BIS is correlated with the depth of sedation in aggregate groups, it lacks sufficient capacity to reliably gauge such depth in individual patients and therefore cannot currently be recommended for ED PSA.

**Resuscitation Equipment and Supplies**

Gather all necessary age-appropriate equipment for airway management and resuscitation in the sedation area, including oxygen, a bag-valve-mask device, suction, and drug reversal agents. For subjects with significant cardiovascular disease, include a defibrillator as well.

**Vital Signs**

Measure vital signs periodically at individualized intervals, in most cases including measurements at baseline, after drug administration, on completion of the procedure, during early recovery, and at completion of recovery. During deep sedation it is reasonable to assess vital signs approximately every 5 minutes. Patients are at highest risk for complications 5 to 10 minutes after IV medications are administered and during the immediate postprocedure period when external stimuli are discontinued. Continuous monitoring of the ECG, blood pressure, pulse rate, and pulse oximetry via a standard monitor generally fulfills the monitoring requirements. Actual documentation in the medical record varies, and fewer entries on the record are necessary when continuous monitoring is used. There are no standards mandating the frequency of vital sign documentation in the medical record. Thus, in the absence of institutional requirements, frequency is generally guided by the specific patient scenario, medications used, and depth of sedation, with common sense prevailing in most cases.

**SUPPLEMENTAL OXYGEN**

Substantial variation in practice exists with regard to the use of supplemental oxygen during PSA. The premise is a logical one: increasing systemic oxygen reserves should naturally delay, or perhaps avert, hypoxemia should an airway or respiratory adverse event occur. However, the price paid for this well-intentioned safeguard is the loss of pulse oximetry as an early warning device. Hyperoxygenated patients will desaturate only after the apnea is prolonged. Indeed, the time required for preoxygenated, apneic, healthy adults and adolescents to desaturate to 90% averages more than 6 minutes.

Deitch and colleagues have shown in a series of randomized controlled trials that high-flow supplemental oxygen decreases the incidence of hypoxia during propofol sedation (number needed to benefit of 4), whereas lesser amounts of oxygen (3 L/min) do so only marginally with propofol and not at all with lighter levels of sedation. Thus, high-flow oxygen is strongly recommended with propofol or other deep sedation, assuming that interactive monitoring includes capnography to promptly identify respiratory depression. For lighter levels of sedation, supplemental oxygen has no established benefit and may impair detection of respiratory depression when using pulse oximetry without capnography.

**DISCHARGE CRITERIA**

Monitor all patients receiving PSA until they are no longer at risk for cardiorespiratory depression (Table 33.2). Before

<table>
<thead>
<tr>
<th>TABLE 33.2 Complications After Sedation</th>
<th>COMPLICATION</th>
<th>ETIOLOGY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delayed awakening</td>
<td>Prolonged drug action, hypoxemia, hypercapnia, hypovolemia</td>
<td></td>
</tr>
<tr>
<td>Agitation</td>
<td>Pain, hypoxemia, hypercapnia, full bladder, Paradoxical reactions, Emergence reactions</td>
<td></td>
</tr>
<tr>
<td>Nausea and vomiting</td>
<td>Sedative agents, Premature oral fluids</td>
<td></td>
</tr>
<tr>
<td>Cardiorespiratory events</td>
<td>Pain, hypovolemia, impaired ventilation, Vagal stimulation, opioids, hypoxia, Laryngospasm, airway obstruction, oversedation</td>
<td></td>
</tr>
</tbody>
</table>

discharge be sure that patients are alert and oriented (or have returned to an age-appropriate baseline) with stable vital signs. Many hospitals have chosen to use standardized recovery scoring systems similar to those used in their surgical postanesthesia recovery areas (Table 33.3). Although no generally accepted minimum durations for safe discharge have been established, one large ED study found that, in children with uneventful sedation, no serious adverse effects occurred more than 25 minutes after final medication administration. This suggests that in most cases, prolonged observation beyond an hour is unlikely to be necessary.

Make sure that all patients leave the hospital with a reliable adult who will observe them after discharge for postprocedural complications. Document the name of the individual in the hospital record. Give written instructions regarding appropriate diet, medications, and level of activity (Boxes 33.2 and 33.3). Even though patients may appear awake and able to comprehend instructions, they may not remember details once they leave the ED.

To be eligible for safe discharge, children are not required to walk unaided or demonstrate that they can tolerate an oral

### TABLE 33.3 Sample Recovery Scoring Systems

<table>
<thead>
<tr>
<th>Steward Recovery Score</th>
<th>Arousable, giddy, agitated</th>
<th>1</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Unresponsive</td>
<td>0</td>
</tr>
<tr>
<td>Consciousness</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Awake</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Responding to stimuli</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Not responding</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Airway</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coughing on command or crying</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Maintaining good airway</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Airway requiring maintenance</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Movement</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Moving limbs purposefully</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Nonpurposeful movements</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Not moving</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Modified Aldrete Score</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vital Signs</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stable</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Unstable</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Respirations</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>Shallow respirations or tachypnea</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Apnea</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Level of Consciousness</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alert, oriented or returned to preprocedural level</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

Oxygen Saturation

| 95%–100% or preprocedural level | 2 |
| 90%–94%                          | 1 |
| < 90%                            | 0 |

Color

| Pink or preprocedural color      | 2 |
| Pale or dusky                    | 1 |
| Cyanotic                         | 0 |

Activity

| Moves on command or preprocedural level | 2 |
| Moves extremities or uncoordinated walking | 1 |
| No spontaneous movement             | 0 |

Sedation Score Action

| > 8                           | Consider discharge if no score = 0 |
| 7–8                          | Vital signs q 20 min |
| 4–6                          | Vital signs q 10 min |
| 0–3                          | Vital signs q 5 min—consider further evaluation if prolonged |


---

**BOX 33.2 Sample Adult Disposition Instructions After PSA**

1. Do not drive or operate heavy machinery for 12 hours.
2. Eat a light diet for the next 12 hours.
3. Take only your prescribed medications as needed, including any pain medication you were discharged with. Avoid alcohol.
4. Do not make any important decisions or sign important documents for 12 hours. You may be forgetful because of the medications that were administered.
5. If you experience any difficulty breathing or persistent nausea and vomiting, return to the emergency department.
6. You should have a responsible person with you for the rest of the day and during the night.

PSA, Procedural sedation and analgesia.
Before drug administration, every effort should be made to minimize a patient’s anxiety and distress, particularly in children. The emotional state of a patient on induction strongly correlates with the degree of distress on emergence and in the days immediately after the procedure. Avoid being pressured by consultants to cut corners or rush PSA. Incorporating into the presedation preparation a discussion with the consultant about the sedation plan and the length of time required to safely prepare and sedate the patient can avoid the risks associated with hurried sedation.

For pediatric PSA, the clinician should appreciate the adult dose of the sedative being administered and consider this the maximum threshold. Understand that the initial dose of midazolam for PSA in a 100-kg patient on a milligram-per-kilogram basis is far less than the 0.1 mg/kg used in a child to avoid unexpected mishaps in drug dosing.

### ROUTES OF ADMINISTRATION

For nondissociative agents, titrate IV medications to the patient’s response for the best method of achieving rapid and safe analgesia and sedation. Wait the appropriate time for the medications to produce the intended effect before adding more doses. When using opioids, administer doses in 2- to 3-minute increments and observe for side effects such as miosis, somnolence, decreased responsiveness to verbal stimuli, impaired speech, and diminished pain on questioning as appropriate initial end points. For sedative-hypnotics, use similar incremental dosing and end points such as ptosis (rather than miosis), somnolence, slurred speech, and alterations in gaze. Repeated doses may be given in a titrated fashion based on the patient’s response during the procedure.

The oral, transmucosal (i.e., nasal, rectal), and IM routes are more convenient means of administration when IV access is not necessary, but they are much less reliable for timely dose titration to a desired response. New drug delivery systems, however, are expanding the effectiveness and ease of use of these routes of administration. The refinement of intranasal drug delivery has significantly increased the efficacy of this route of administration. Before the development of metered-dose atomizers, the degree of absorption and effectiveness of intranasal drug administration were operator dependent. Furthermore, new drug formulations with concentrations appropriate for intranasal administration are becoming available for study.

The main advantage of these other routes is for pediatric patients in whom IV access may be problematic or for procedures that may require only minimal sedation in conjunction with the use of local anesthetics. These routes are also advantageous for simple sedation during diagnostic imaging.

With the exception of ketamine, agents administered intramuscularly have erratic absorption and a variable onset of action. Accordingly, prolonged preprocedural and postprocedural observation may be necessary. When required, the IM route offers little advantage over oral or transmucosal administration.

Another PSA route is via inhalation of nitrous oxide. This gas can either be delivered by a demand-flow system using a handheld mask or be delivered to young children using a nose mask in a continuous-flow system under close clinician supervision.

Because individual needs may vary widely, the application of arbitrary ceiling doses of analgesic and sedative regimens...
is unwarranted. The true ceiling dose of an agent is the level that provides adequate pain relief or sedation without major cardiopulmonary side effects such as respiratory depression, apnea, bradycardia, hypotension, or allergic reactions.

There are two absolute contraindications to PSA: severe clinical instability requiring immediate attention and refusal by a competent patient. Relative contraindications include hemodynamic or respiratory compromise, altered sensorium, or an inability to monitor for adverse events (e.g., magnetic resonance imaging [MRI] without remote monitoring). However, even in many of these circumstances, appropriate agents can be given to provide analgesia and sedation while minimizing the chance for further deterioration. Although safely sedating patients at the extremes of age is challenging and requires additional care, as well as reductions in drug dosing (because of decreased drug metabolism and excretion), age is not a contraindication to PSA.

**DRUG SELECTION STRATEGIES**

The majority of nonpainful or minimally painful ED procedures in older children and adults can be performed without systemic sedation and analgesia. Skilled practitioners can frequently combine a calm, reassuring bedside manner with distraction techniques, careful local or regional anesthesia, or both. Many procedures, however, cannot be technically or humanely performed without PSA. These situations can be divided into three categories.

**Insufficient Analgesia.** Despite a cooperative patient, for some procedures it is impossible to achieve effective pain control with local or regional anesthesia. Examples of procedures requiring systemic PSA include fracture reductions, dislocation reductions, incision and drainage of large loculated abscesses, wounds that require scrubbing such as “road rash,” cardioversion, bone marrow aspiration/biopsy, and extensive burn débridement.

**Insufficient Anxiolysis.** Despite effective local or regional anesthesia, some patients will be so frightened that procedures cannot be technically or humanely performed without PSA. Young children requiring repair of lacerations are frequently in anticipation of such repairs in sensitive or personal regions (e.g., face, genitalia, perineum).

**Insufficient Immobilization.** Despite effective local or regional anesthesia and anxiolysis, PSA may be indicated to prevent excessive motion during procedures that require substantial immobilization (e.g., repair of complex facial lacerations, diagnostic imaging studies). Immobilization is most commonly an issue with young children and the mentally challenged.

**General Considerations.** Clinicians must therefore base customization of their selection of drugs (e.g., anxiolysis, sedation, analgesia, immobilization) on the unique needs of the patient and their individual level of experience with specific agents (Table 33.4). A risk-benefit analysis should be performed before every sedation (Box 33.4). The benefits of reducing anxiety and controlling pain should be carefully weighed against the risk for respiratory depression and airway compromise.

Factors influencing the extent of pharmacologic management are listed in Box 33.5. Some general drug selection strategies are discussed later and shown in Boxes 33.6, 33.7, and 33.8.

**MINOR PROCEDURES IN COOPERATIVE ADULTS AND OLDER CHILDREN.** Such procedures can usually be managed with topical, local, or regional anesthesia. Systemic PSA is typically unnecessary, although mild anxiolysis (e.g., nitrous oxide, oral midazolam) can make these patients more comfortable.

**MORE COMPLEX PROCEDURES OF LONGER DURATION IN COOPERATIVE ADULTS AND OLDER CHILDREN.** Supplemental of topical, local, or regional anesthesia with either nitrous oxide or IV midazolam and fentanyl permits customization of the depth of sedation and pain relief to the specific needs of each patient.

**PROCEDURES IN UNCOOPERATIVE ADULTS OR THE MENTALLY CHALLENGED.** Essentially all procedures in uncooperative adult-sized patients are difficult without systemic PSA. Depending on operator experience, IV midazolam, IV propofol, IV etomidate, or IM/IV ketamine or midazolam may be used in these situations. Given that the sedatives midazolam, propofol, and etomidate lack specific analgesic properties, many emergency physicians attempt to control pain with an opioid such as fentanyl before the procedure. Midazolam can be titrated intravenously to a relatively deep level of sedation, although as discussed previously, the risk for adverse effects increases with the depth of sedation. Ketamine (typically with coadministered midazolam when used in adults) can also provide the profound analgesia and immobilization necessary to perform painful procedures. However, in adults there is a risk for unpleasant hallucinatory recovery reactions. Ketamine should be used with extreme caution in older adults because its

---

**BOX 33.4 Risk-Benefit Analysis for PSA**

- Why is PSA needed in the first place? Is the procedure very painful, frightening, or requiring extreme cooperation?
- Are the risks of PSA appropriate for the procedure involved?
- If the patient is a child, do the parents or guardian consent to the use of PSA?
- How long will the procedure take? If it is a short procedure, is it worth the added risk and expense to the patient? If it is a longer procedure, is there an appropriate agent that can be titrated to allow adequate PSA throughout the entire length of the procedure?
- Are there significant side effects that limit a particular drug’s usefulness?
- Are enough nurses and support personnel present to safely allow the use of PSA?
- What is the recovery period for a given agent? Are there enough treatment areas and staff in the ED to allow adequate observation during recovery?
- When did the patient last eat? Is a delay in waiting for a sufficient fasting time worth the time lost in performing the procedure?

ED, Emergency department; PSA, procedural sedation and analgesia.

sympathomimetic properties may aggravate underlying coronary artery disease or hypertension. Occasionally, procedures in extremely uncooperative adults or the mentally challenged are best managed in the operating room with general anesthesia.

MINOR PROCEDURES IN UNCOOPERATIVE OLDER CHILDREN AND IN YOUNG CHILDREN. Minor procedures (e.g., small lacerations, IV cannulation, venipuncture, removal of superficial foreign bodies) in uncooperative children can frequently be managed by skilled practitioners with a combination of nonpharmacologic techniques (e.g., distraction, guided imagery, hypnosis, comforting, breathing techniques) in conjunction with topical anesthesia, careful local anesthesia, and when necessary, brief forcible immobilization (by personnel or a restraining device). In other cases, supplementing nonpharmacologic techniques with topical or local anesthesia and anxiolysis with oral midazolam may be sufficient to permit successful wound repair. Although oral administration is most popular and least invasive, the nasal or rectal routes can also be used depending on operator experience and preference.

### TABLE 33.4 Indications for PSA and Sedation Strategies

<table>
<thead>
<tr>
<th>CLINICAL SITUATION</th>
<th>INDICATION</th>
<th>PROCEDURAL REQUIREMENTS</th>
<th>SUGGESTED SEDATION STRATEGIES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Noninvasive procedures</td>
<td>CT</td>
<td>Motion control Anxiolysis</td>
<td>Comforting alone Chloral hydrate PO (in patients &lt; 3 yr old) Methohexital PR Pentobarbital PO, IM, or IV Midazolam IV Propofol or etomidate IV</td>
</tr>
<tr>
<td></td>
<td>Echocardiography</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Electroencephalography</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>MRI</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ultrasonography</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Procedure associated with low pain and high anxiety</td>
<td>Dental procedures</td>
<td>Sedation Anxiolysis Motion control</td>
<td>Comforting and topical or local anesthesis Midazolam PO/IN/PR/IV Nitrous oxide</td>
</tr>
<tr>
<td></td>
<td>Flexible fiberoptic laryngoscopy</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Foreign body removal, simple</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Intravenous cannulation</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Laceration repair, simple</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lumbar puncture</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Ocular irrigation</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Phlebotomy</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Slit-lamp examination</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Procedures associated with a high level of pain, high anxiety, or both</td>
<td>Abscess incision and drainage</td>
<td>Sedation Anxiolysis Analgesia Amnesia Motion control</td>
<td>Propofol or etomidate IV Propofol and fentanyl IV Propofol and ketamine IV Ketamine IM/IV Midazolam and fentanyl IV</td>
</tr>
<tr>
<td></td>
<td>Arthrocentesis</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Bone marrow aspiration and biopsy</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Burn débridement</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cardiac catheterization</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cardioversion</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Central line placement</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Endoscopy</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Foreign body removal, complicated</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fracture or dislocation reduction</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hernia reduction</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Interventional radiology procedures</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Laceration repair, complex</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Paracentesis</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Paraphimosis reduction</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sexual assault examination</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Thoracentesis</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Thoracostomy tube placement</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*There is no universally accepted or clinically correct dose, medication, or combination. Many regimens are acceptable. This table is intended as a general overview. Sedation strategies should be individualized. Although the pharmacopoeia is large, clinicians should familiarize themselves with a few agents that are flexible enough to be used for the majority of procedures. In all cases it is assumed that practitioners are fully trained in the technique, appropriate personnel and monitoring are used as detailed in this chapter, and specific drug contraindications are absent. CT, Computed tomography; IM, intramuscularly; IN, intranasally; IV, intravenously; MRI, magnetic resonance imaging; PO, orally; PR, per rectum; PSA, procedural sedation and analgesia.

**Factors Influencing the Extent of Pediatric Pharmacologic Management**

**AGE**
Selected drugs and routes of administration have age limitations and are not recommended above or below a certain age (e.g., demand-flow nitrous oxide in children < 5 years, nasal and rectal routes of administration in children > 6 years).

**TIME OF DAY**
A toddler seen at naptime or at 9 PM who is tired and sleepy will usually require smaller dosing and possibly a lower level of PSA than required at 9 AM. Young children with facial lacerations at night, after their normal bedtime, may require only topical anesthesia and a quiet room for 20 to 30 minutes to achieve a painless laceration repair while the child sleeps.

**FASTING STATUS**
Young children can be extremely difficult and uncooperative when hungry, tired, or both. In anticipation of PSA, many children are kept without oral intake from the time that they are triaged in the ED. This can further increase hunger and irritability, especially if the child waits 1 to 2 hours to be seen by a clinician.

**AVAILABILITY OF STAFFING AND EQUIPMENT**
Staffing availability can affect the use and timing of sedation and is especially important in busy EDs with multiple sedations occurring concurrently and in smaller units that are set up for only one sedation at a time.

**LOCATION OF THE INJURY**
Injuries located in areas of cosmetic concern (especially on the face) or near sensory organs (e.g., ears, eyes, mouth, nose) will often require a high degree of agitation control and a concomitant level of PSA.

**PREVIOUS MEDICATIONS**
An accurate history of previous medication administration is important in situations in which a child is referred from another facility because this can affect the type and timing of PSA agents that can be given. In particular, a child may have received opioids or sedative-hypnotics before transfer and may still be sedated on arrival, thus necessitating an adjustment in the PSA regimen.

**LEVEL OF ANXIETY**
The level of anxiety of both the child and the accompanying adult or adults must be accurately assessed. Children manifest anxiety in many different ways, and emergency clinicians must be facile at recognizing the varying expressions of anxiety, especially in young children. A child with a facial laceration quietly sitting on the stretcher during the initial examination will not necessarily be a calm and cooperative patient during repair of the laceration (infants and toddlers). The nursing assessment at triage of the state of the child and accompanying adult or adults can be very helpful in some cases in determining the need for PSA. A child who was frightened and uncooperative in triage may be calm and compliant during a procedure. Unfortunately, the reverse is also true. When confronted with an extremely anxious child, ED personnel should ascertain what the parents have told the child about the upcoming procedure. Many parents, in the hope of lessening their child’s anxiety, will tell the child that she or he will get a “shot” or a “needle” and that the procedure will “only hurt for a minute.” This type of parental preparation, especially in young children who do not have the cognitive ability to mediate their anxiety, often results in a significant increase in the child’s anxiety and a decrease in the child’s ability to cooperate, especially if the child has had a previous negative experience with a procedure in the ED. It is also important to assess the parents’ level of anxiety because this will determine the degree to which they can assist during the procedure. An extremely anxious parent or a parent who must take care of other siblings during the procedure will find it difficult to assist in distracting the child or otherwise helping the child cope with the procedure.

**PREVIOUS EXPERIENCE**
Children’s previous experience in hospitals can greatly affect their response to the current situation. Direct experience is not the only way to create anxious, frightened, and uncooperative patients, though. Images from television, stories from peers, or previous witness of a sibling being forcibly restrained for repair of a laceration can leave a powerful and lasting impression. This type of influence should be especially suspected in children whose anxiety seems out of proportion to the present situation. Eliciting from the parents a history of a previous difficult experience in the ED can be a decisive factor in determining the degree of sedation required. Children who have had a recent unpleasant laceration repair and who now have a new laceration may well require PSA as opposed to simple anxiolysis (either pharmacologic or nonpharmacologic) had there been no previous trauma.

**CHILD’S BEHAVIOR AT ROUTINE PRIMARY CARE VISITS**
Inquiring into how a child behaves during routine primary care visits can yield important information on how the child reacts to stressful situations, how cooperative the child will be with the anticipated procedure, and whether pharmacologic management is needed. Children who cry but hold still when vaccinated may be more compliant than children who are described by their parents as being “afraid of doctors” or “wild” during visits to the primary care physician.

ED, Emergency department; PSA, procedural sedation and analgesia.

**Systemic Analgesia and Sedation for Procedures**

**Factors Influencing the Extent of Pediatric Pharmacologic Management**

**AGE**
Selected drugs and routes of administration have age limitations and are not recommended above or below a certain age (e.g., demand-flow nitrous oxide in children < 5 years, nasal and rectal routes of administration in children > 6 years).

**TIME OF DAY**
A toddler seen at naptime or at 9 PM who is tired and sleepy will usually require smaller dosing and possibly a lower level of PSA than required at 9 AM. Young children with facial lacerations at night, after their normal bedtime, may require only topical anesthesia and a quiet room for 20 to 30 minutes to achieve a painless laceration repair while the child sleeps.

**FASTING STATUS**
Young children can be extremely difficult and uncooperative when hungry, tired, or both. In anticipation of PSA, many children are kept without oral intake from the time that they are triaged in the ED. This can further increase hunger and irritability, especially if the child waits 1 to 2 hours to be seen by a clinician.

**AVAILABILITY OF STAFFING AND EQUIPMENT**
Staffing availability can affect the use and timing of sedation and is especially important in busy EDs with multiple sedations occurring concurrently and in smaller units that are set up for only one sedation at a time.

**LOCATION OF THE INJURY**
Injuries located in areas of cosmetic concern (especially on the face) or near sensory organs (e.g., ears, eyes, mouth, nose) will often require a high degree of agitation control and a concomitant level of PSA.

**PREVIOUS MEDICATIONS**
An accurate history of previous medication administration is important in situations in which a child is referred from another facility because this can affect the type and timing of PSA agents that can be given. In particular, a child may have received opioids or sedative-hypnotics before transfer and may still be sedated on arrival, thus necessitating an adjustment in the PSA regimen.

**LEVEL OF ANXIETY**
The level of anxiety of both the child and the accompanying adult or adults must be accurately assessed. Children manifest anxiety in many different ways, and emergency clinicians must be facile at recognizing the varying expressions of anxiety, especially in young children. A child with a facial laceration quietly sitting on the stretcher during the initial examination will not necessarily be a calm and cooperative patient during repair of the laceration (infants and toddlers). The nursing assessment at triage of the state of the child and accompanying adult or adults can be very helpful in some cases in determining the need for PSA. A child who was frightened and uncooperative in triage may be calm and compliant during a procedure. Unfortunately, the reverse is also true. When confronted with an extremely anxious child, ED personnel should ascertain what the parents have told the child about the upcoming procedure. Many parents, in the hope of lessening their child’s anxiety, will tell the child that she or he will get a “shot” or a “needle” and that the procedure will “only hurt for a minute.” This type of parental preparation, especially in young children who do not have the cognitive ability to mediate their anxiety, often results in a significant increase in the child’s anxiety and a decrease in the child’s ability to cooperate, especially if the child has had a previous negative experience with a procedure in the ED. It is also important to assess the parents’ level of anxiety because this will determine the degree to which they can assist during the procedure. An extremely anxious parent or a parent who must take care of other siblings during the procedure will find it difficult to assist in distracting the child or otherwise helping the child cope with the procedure.

**PREVIOUS EXPERIENCE**
Children’s previous experience in hospitals can greatly affect their response to the current situation. Direct experience is not the only way to create anxious, frightened, and uncooperative patients, though. Images from television, stories from peers, or previous witness of a sibling being forcibly restrained for repair of a laceration can leave a powerful and lasting impression. This type of influence should be especially suspected in children whose anxiety seems out of proportion to the present situation. Eliciting from the parents a history of a previous difficult experience in the ED can be a decisive factor in determining the degree of sedation required. Children who have had a recent unpleasant laceration repair and who now have a new laceration may well require PSA as opposed to simple anxiolysis (either pharmacologic or nonpharmacologic) had there been no previous trauma.

**CHILD’S BEHAVIOR AT ROUTINE PRIMARY CARE VISITS**
Inquiring into how a child behaves during routine primary care visits can yield important information on how the child reacts to stressful situations, how cooperative the child will be with the anticipated procedure, and whether pharmacologic management is needed. Children who cry but hold still when vaccinated may be more compliant than children who are described by their parents as being “afraid of doctors” or “wild” during visits to the primary care physician.

ED, Emergency department; PSA, procedural sedation and analgesia.

**Pharmacopeia**

There is no universally correct or preferred medication or drug regimen. Many options are acceptable and successful.
with a recovery time of an additional 1 to 2 hours. Residual sedation may persist for several hours beyond this period. Rectal administration is erratically absorbed and therefore not recommended.

Adult Use. Use of chloral hydrate is limited to diagnostic imaging studies in children. It has no current use in adults.

Pediatric Use. Chloral hydrate is widely used as a sedative to facilitate nonpainful outpatient diagnostic procedures such as electroencephalography, computed tomography (CT) or MRI. IV pentobarbital appears to be more effective than chloral hydrate for the latter indication, although many centers prefer chloral hydrate in younger children (e.g., < 18 months) simply to avoid the need for IV access.

Adverse Effects. Despite a wide margin of safety, chloral hydrate can cause airway obstruction and respiratory depression, especially at higher doses (75 to 100 mg/kg). The incidence was 0.6% in one large series. There is no known dosage threshold of chloral hydrate below which this potential

### Sedative-Hypnotic Agents

**Chloral Hydrate**

**Pharmacology.** Chloral hydrate is a pure sedative-hypnotic agent without analgesic properties. When administered orally, the average time to peak sedation is approximately 30 minutes, with a recovery time of an additional 1 to 2 hours. Residual motor imbalance and agitation may persist for several hours.

**CONTRAINDICATIONS—ABSOLUTE (RISKS ESSENTIALLY ALWAYS OUTWEIGH BENEFITS)**

- Active hemodynamic instability
- Active respiratory distress or hypoxemia

**CONTRAINDICATIONS—RELATIVE (RISKS MAY OUTWEIGH BENEFITS)**

- Respiratory depression or altered level of consciousness
- Anticipated difficulty if ventilatory assistance should become necessary (e.g., facial deformity or trauma, small mandible, large tongue, trismus)

**PROTOCOL**

1. Establish intravenous access.
2. Preoxygenate the patient.
3. Connect appropriate monitoring equipment to the patient.
4. Administer supplemental oxygen if deep sedation is anticipated.
5. The pulse, respiratory rate, blood pressure, and level of consciousness should all be recorded initially and periodically throughout the procedure, depending on the depth of sedation.
6. Suction equipment, oxygen, a bag-valve-mask assembly, and reversal agents should be available immediately. An age-appropriate resuscitation cart with oral and nasal airways, endotracheal tubes, and a functioning laryngoscope must be nearby.
7. The order of drugs is one of personal preference. The ratio of analgesia to sedation is determined by the nature of the procedure. Some procedures require primary analgesia and secondary anxiolysis or sedation (e.g., incision and drainage of an abscess, bone marrow aspiration, arthrocentesis, burn débridement, central catheter placement). In this case, administer fentanyl first. Others require primary anxiolysis or sedation with secondary analgesia (e.g., lumbar puncture, simple foreign body removal); administer midazolam first.
8. Administer a local anesthetic if indicated after procedural sedation and analgesia are initiated (this often serves to help gauge the effectiveness of systemic analgesia).
9. Perform the procedure. Additional doses of fentanyl or midazolam may be required if further pain or anxiety is noted based on the response and length of the procedure.
10. If hypoxemia, oversedation, or slowed respirations are seen during or after the procedure, the patient should first be stimulated while oxygen is applied and the airway repositioned. If the patient’s response is insufficient, assist ventilations with a bag-valve-mask device. Reversal agents should be considered if there is not a prompt response to assisted ventilation.
11. Continue close observation until the patient is awake and alert, and release the patient with a friend, parent, or relative only after a sufficient discharge score has been attained.

**Box 33.6 Procedure for Moderate to Deep Sedation With Intravenous Midazolam and Fentanyl**

**CAVEATS**

- Do not consider this procedure if you lack experience with the drugs or do not have the time to perform procedural sedation and analgesia properly. Do not attempt this procedure if the pulse oximeter, suction, oxygen, or bag-mask device is not working, the intravenous line is not secured, or the room is too small or not set up for procedural sedation and analgesia.
- This is a two-person procedure, one to monitor the patient and one to perform the procedure.
- The individual response to the drugs is variable and dependent on the patient’s underlying physiologic state and the presence of concomitant drugs and medications.
- The maximum drug effect occurs 2 to 3 minutes after administration. Proceed slowly and patiently and allow the medication to take full effect before giving the next dose.
- Have naloxone and flumazenil immediately available for oversedation or respiratory depression.
- If the patient seems overly sedated, begin the procedure. The pain of the procedure often stimulates respiration and lessens sedation.

**CONTRAINDICATIONS—ABSOLUTE (RISKS ESSENTIALLY ALWAYS OUTWEIGH BENEFITS)**

- Active hemodynamic instability
- Active respiratory distress or hypoxemia

**CONTRAINDICATIONS—RELATIVE (RISKS MAY OUTWEIGH BENEFITS)**

- Respiratory depression or altered level of consciousness
- Anticipated difficulty if ventilatory assistance should become necessary (e.g., facial deformity or trauma, small mandible, large tongue, trismus)
complication can be consistently avoided,\textsuperscript{1,80} and accordingly, standard interactive and mechanical monitoring precautions apply to chloral hydrate as they do to other PSA agents.

Because it is a halogenated hydrocarbon, overdoses of chloral hydrate can be arrhythmogenic and produce ventricular arrhythmias. β-Blockers may be most effective in terminating the arrhythmia.\textsuperscript{83} The addition of end-tidal CO\textsubscript{2} monitoring (capnography) can provide the earliest possible warning of impending airway and respiratory complications before clinical examination or pulse oximetry and is particularly important when supplemental oxygen negates pulse oximetry as a warning device.\textsuperscript{84}

**Midazolam**

**Pharmacology.** Benzodiazepines are a group of highly lipophilic agents that possess anxiolytic, amnestic, sedative, hypnotic, muscle relaxant, and anticonvulsant properties. They lack direct analgesic properties and thus are commonly coadministered with opioids. Caution must be exercised when using benzodiazepines and opioids together because the risk for hyponxia and apnea is significantly greater than when either is used alone.\textsuperscript{84}

Midazolam is by far the most common benzodiazepine used for PSA and is preferred over the longer-acting lorazepam and diazepam. The time to peak effect for midazolam is approximately 2 to 3 minutes when given intravenously. Unlike diazepam, midazolam and lorazepam are water soluble, thus making parenteral administration less painful and mucosal absorption faster. Midazolam is readily reversed with flumazenil, and individuals undergoing PSA with midazolam are good candidates for this antidote should reversal be required.
**DEFINITION OF DISSOCIATIVE SEDATION**
- A trancelike cataleptic state induced by the dissociative agent ketamine and characterized by profound analgesia and amnesia with retention of protective airway reflexes, spontaneous respirations, and cardiopulmonary stability.

**CHARACTERISTICS OF THE KETAMINE “DISSOCIATIVE STATE”**
- Dissociation: After the administration of ketamine, the patient passes into a fugue state or trance. The eyes may remain open, but the patient does not respond.
- Catalepsy: Normal or slightly enhanced muscle tone is maintained. On occasion, the patient may move or be moved into a position that is self-maintaining. Occasional muscular clonus may be observed.
- Analgesia: Analgesia is typically substantial or complete. Ketamine delivers sufficient analgesia, so there will be no pain.
- Amnesia: Total amnesia is typical.
- Maintenance of airway reflexes: Upper airway reflexes remain intact and may be slightly exaggerated. Intubation is unnecessary, but occasional repositioning of the head may be needed for optimal airway patency. Suctioning of hypersalivation may occasionally be necessary.
- Cardiovascular stability: Blood pressure and pulse rate are not decreased and are typically mildly increased.
- Nystagmus: Nystagmus is typical.
- Hypersalivation may occasionally be necessary. Emphasize, especially to school-aged children and teenagers, that ketamine delivers sufficient analgesia, so there will be no pain.

**INDICATIONS**
- Short, painful procedures, especially those requiring immobilization (e.g., facial laceration, burn débridement, fracture reduction, abscess incision and drainage, central line placement, tube thoracostomy).
- Examinations judged likely to produce excessive emotional disturbance (e.g., pediatric sexual assault examination). Emphasize, especially to school-aged children and teenagers, that ketamine delivers sufficient analgesia, so there will be no pain.

**CONTRAINDICATIONS: ABSOLUTE (RISKS ESSENTIALLY ALWAYS OUTWEIGH BENEFITS)**
- Age younger than 3 months (higher risk for airway complications).
- Known or suspected schizophrenia, even if currently stable or controlled with medications (can exacerbate the condition).
- Porphyria, thyroid disorder, or thyroid medication (enhanced sympathomimetic effect).

**CONTRAINDICATIONS: RELATIVE (RISKS MAY OUTWEIGH BENEFITS)**
- Major procedures stimulating the posterior pharynx (e.g., endoscopy) increase the risk for laryngospasm, whereas typical minor ED oropharyngeal procedures do not.
- History of airway instability, tracheal surgery, or tracheal stenosis (presumed higher risk for airway complications).
- Active pulmonary infection or disease, including upper respiratory infection or asthma (higher risk for laryngospasm).
- Known or suspected cardiovascular disease, including angina, heart failure, or hypertension (exacerbation caused by the sympathomimetic properties of ketamine). Avoid ketamine in patients who are already hypertensive and in older adults with risk factors for coronary artery disease.
- Central nervous system masses, abnormalities, or hydrocephalus (increased intracranial pressure with ketamine).
- Glaucoma or acute globe injury (increased intraocular pressure with ketamine).

**PERSONNEL**
- Dissociative sedation requires two persons, one (a nurse) to monitor the patient and one (a physician) to perform the procedure. Both must know about ketamine’s unique characteristics.

**PRESEDATION**
- Perform a standard presedation assessment.
- Educate accompanying family members about the unique characteristics of the dissociative state if they will be present during the procedure or recovery.
- Frame the dissociative encounter as a positive experience.
- Consider encouraging adults and older children to “plan” specific, pleasant dream topics in advance of sedation (believed to decrease unpleasant recovery reactions). Emphasize, especially to school-aged children and teenagers, that ketamine delivers sufficient analgesia, so there will be no pain.

**KETAMINE ADMINISTRATION: GENERAL**
- Ketamine is not administered until the physician is ready to begin the procedure because the onset of dissociation typically occurs rapidly.
- Ketamine is initially administered as a single IV loading dose or IM injection. There is no apparent benefit from attempts to titrate to effect.
- In settings in which IV access can be obtained with minimal upset, the IV route is preferable because recovery is faster and there is less emesis. The IM route is especially useful when IV access cannot be consistently obtained with minimal upset and in patients who are uncooperative or combative (e.g., the mentally disabled).
- IV access is unnecessary for children receiving IM ketamine. Because unpleasant recovery reactions are more common in adults, IV access is desirable in these patients to permit rapid treatment of these reactions should they occur.

**KETAMINE ADMINISTRATION: IV ROUTE**
- Administer an IV loading dose of 1.5 to 2.0 mg/kg in children or 1.0 mg/kg in adults, with this dose being administered during a 30- to 60-second period. More rapid administration produces high central nervous system levels and has been associated with respiratory depression or apnea.
- Additional incremental doses of ketamine may be administered (0.5 to 1.0 mg/kg) if initial sedation is inadequate or repeated doses are necessary to accomplish a longer procedure.

**KETAMINE ADMINISTRATION: IM ROUTE**
- Administer IM ketamine, 4 to 5 mg/kg in children; the IV route is preferred for adults.
- Repeat the IM ketamine dose (full or half dose) if sedation is inadequate after 5 to 10 minutes (unusual) or if additional doses are required.
**COADMINISTERED MEDICATIONS**

- Prophylactic anticholinergics are no longer recommended.
- Prophylactic benzodiazepines are no longer recommended for children; however, they should be available to treat rare, unpleasant recovery reactions should they occur. Prophylactic IV midazolam, 0.03 mg/kg, may be considered for adults (number needed to benefit, 6).
- Prophylactic ondansetron can slightly reduce the rate of vomiting (number needed to benefit, 9 or higher).

<table>
<thead>
<tr>
<th>ROUTE OF ADMINISTRATION</th>
<th>IV</th>
<th>IM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advantages</td>
<td>Ease of repeated dosing, less vomiting, slightly faster recovery</td>
<td>No IV access necessary</td>
</tr>
<tr>
<td>Peak concentrations and clinical onset, minutes</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Typical duration of effective dissociation (min)</td>
<td>5–10</td>
<td>20–30</td>
</tr>
<tr>
<td>Typical time from dose to discharge (min)</td>
<td>50–110</td>
<td>60–140</td>
</tr>
</tbody>
</table>

**PROCEDURE**

- Adjunctive physical immobilization may occasionally be needed to control random motion.
- An adjunctive local anesthetic is usually unnecessary when a dissociative dose is used.
- Suction equipment, oxygen, a bag-valve-mask device, and age-appropriate equipment for advanced airway management should be immediately available. Supplemental oxygen is not mandatory but may be used when capnography is used to monitor ventilation.

**INTERACTIVE MONITORING**

- Close observation of the airway and respirations by an experienced health care professional is mandatory until recovery is well established.
- Drapes should be positioned so that airway and chest motion can be visualized at all times.
- Occasional repositioning of the head or suctioning of the anterior pharynx may be indicated for optimal airway patency.

**MECHANICAL MONITORING**

- Maintain continuous monitoring (e.g., pulse oximetry, cardiac monitoring, capnography) until recovery is well established.

- The pulse and respiratory rate should both be recorded periodically throughout the procedure. Blood pressure measurements after the initial value are generally unnecessary because ketamine stimulates catecholamine release and does not depress the cardiovascular system in healthy patients.

**POTENTIAL ADVERSE EFFECTS**

Percent estimates are for children; corresponding adult estimates are not yet reliable enough to report.

- Airway misalignment requiring repositioning of the head (occasional)
- Transient laryngospasm (0.3%)
- Transient apnea or respiratory depression (0.8%)
- Hypersalivation (rare)
- Emesis, usually well into recovery (8.4%)
- Recovery agitation (mild in 6.3%, clinically important in 1.4%)
- Muscular hypertonicity and random, purposeless movements (common)
- Clonus, hiccuping, or short-lived nonallergic rash on the face and neck

**RECOVERY**

- Maintain minimal physical contact or other sensory disturbance.
- Maintain a quiet area with dim lighting, if possible.
- Advise parents or caretakers not to stimulate the patient prematurely.

**DISCHARGE CRITERIA**

- Return to the pretreatment level of verbalization and awareness
- Return to the pretreatment level of purposeful neuromuscular activity
- A predischARGE requirement of tolerating oral fluids or being able to ambulate independently not required or recommended after dissociative sedation

**DISCHARGE INSTRUCTIONS**

- Nothing by mouth for approximately 2 hours
- Careful family observation and no independent ambulation for approximately 2 hours


**Adult Use.** Midazolam can be used effectively for moderate and deep sedation through careful IV titration to effect, typically together with fentanyl (see Box 33.6).

**Pediatric Use.** Advantages of midazolam over other benzodiazepines for pediatric PSA are its short duration of action, reversibility, and availability in multiple routes of administration. Midazolam may be used for the same indications and in the same manner as in adults. Some children require larger doses than would be typical for adults on a milligram-per-kilogram basis, and paradoxical responses (e.g., hyperexcitability) are not uncommon. Midazolam does not reliably render a child motionless, and therefore methohexital or pentobarbital is generally preferred for neuroimaging.

To avoid the need for IV access in frightened children, midazolam has been alternatively administered via the IM, oral, intranasal, and rectal routes. However, the inability to effectively titrate with these routes dictates that a reliable depth of sedation cannot be predictably or regularly achieved. Consequently, these non-IV routes are primarily...
### TABLE 33.5 Drug Dosing Recommendations for PSA

<table>
<thead>
<tr>
<th>DRUG</th>
<th>CLINICAL EFFECTS</th>
<th>INDICATIONS</th>
<th>ADULT DOSE</th>
<th>PEDIATRIC DOSE</th>
<th>ONSET (min)</th>
<th>DURATION (min)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sedative-Hypnotics</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Choral Hydrate (Noect, Apothecon Pharmaceuticals, District Vadodara, Gujarat, India)</td>
<td>Sedation, motion control, anxiolysis</td>
<td>Not recommended</td>
<td>PO: 25–100 mg/kg, after 30 min may repeat 25–50 mg/kg</td>
<td>PO: 60–120</td>
<td>PO: 15–30</td>
<td>Effects unreliable if age &gt; 3 yr</td>
<td>Avoid in patients with significant cardiac, hepatic, or renal disease Rectal absorption is erratic May produce paradoxical excitement Because drugs cannot be titrated with the PO route, monitor closely for oversedation</td>
</tr>
<tr>
<td>Etomidate (Anidate, Pfizer Inc., New York, NY)</td>
<td>Sedation, motion control, anxiolysis</td>
<td>Procedures requiring sedation and/or anxiolysis</td>
<td>Sedation: 0.1 mg/kg IV; repeat if inadequate response</td>
<td>Not FDA approved in children</td>
<td>IV: &lt; 1</td>
<td>IV: 5–15</td>
<td>Adverse effects include respiratory depression, myoclonus, nausea, and vomiting Adrenocortical suppression occurs but is of no clinical significance</td>
</tr>
<tr>
<td>Midazolam (Versed, Sandoz Inc., East Hanover, NJ)</td>
<td>Sedation, motion control, anxiolysis</td>
<td>Procedures requiring sedation and/or anxiolysis</td>
<td>IV: Initial dose 1 mg, then titrated to max of 5 mg IM: 5 mg or 0.07 mg/kg</td>
<td>IV (0.5–5 yr): Initial dose of 0.05–0.1 mg/kg, then titrated to max of 0.6 mg/kg IV (6–12 yr): Initial dose of 0.025–0.05 mg/kg, then titrated to max of 0.4 mg/kg IM: 0.1–0.15 mg/kg PO: 0.5–0.75 mg/kg IN: 0.2–0.5 mg/kg PR: 0.25–0.5 mg/kg</td>
<td>IV: 2–3</td>
<td>IM: 10–20</td>
<td>Reduce the dose when used in combination with opioids May produce paradoxical excitement Because drugs cannot be titrated with the PO/PR/IN routes, monitor closely for oversedation</td>
</tr>
<tr>
<td>Methohexital (Brevital, Par Pharmaceutical, Chestnut Ridge, NY)</td>
<td>Sedation, motion control, anxiolysis</td>
<td>Diagnostic imaging</td>
<td>PR: 25 mg/kg IV (caution, limited research): 0.5–1 mg/kg</td>
<td>PR: 10–15</td>
<td>PR: 60</td>
<td>Avoid in patients with temporal lobe epilepsy or porphyria Because drugs cannot be titrated with the PR route, monitor closely for oversedation</td>
<td></td>
</tr>
</tbody>
</table>
### TABLE 33.5
Drug Dosing Recommendations for PSA

<table>
<thead>
<tr>
<th>DRUG</th>
<th>CONTRAINICATIONS</th>
<th>INDICATIONS</th>
<th>ADULT DOSE</th>
<th>PEDIATRIC DOSE</th>
<th>ONSET (min)</th>
<th>DURATION (min)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Diagnostic imaging</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Sedative-Hypnotics</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Pentobarbital</strong> (Nembutal, Akorn Inc., Lake Forest, IL)</td>
<td>Sedation, motion control, anxiolysis</td>
<td>Diagnostic imaging</td>
<td>Not recommended</td>
<td>IV: 1–6 mg/kg, titrated in increments of 1–2 mg/kg to desired effect</td>
<td>IV: 3–5</td>
<td>IV: 15–45</td>
<td>May produce paradoxical excitement. Avoid in patients with porphyria. Because drugs cannot be titrated with the PO/PR routes, monitor closely for oversedation.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>IM: 2–6 mg/kg, max of 100 mg</td>
<td>IM: 10–15</td>
<td>IM: 60–120</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>PO/PR (&lt; 4 yr): 3–6 mg/kg, max of 100 mg</td>
<td>PO/PR: 15–60</td>
<td>PO/PR: 60–240</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>PO/PR (&gt; 4 yr): 1.5–3 mg/kg, max of 100 mg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Propofol</strong> (Diprivan, APP Pharmaceuticals, Inc., Schaumburg, IL)</td>
<td>Sedation, motion control, anxiolysis</td>
<td>Procedures requiring sedation and/or anxiolysis</td>
<td>Load 1 mg/kg IV; may administer additional 0.5-mg/kg doses as needed to enhance or prolong sedation</td>
<td>IV: &lt; 1</td>
<td>IV: 5–15</td>
<td>Frequent hypotension and respiratory depression. Avoid with egg or soy allergies.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>PR: 25 mg/kg</td>
<td>PR: 10–15</td>
<td>PR: 60–120</td>
<td></td>
</tr>
<tr>
<td><strong>Thiopental</strong> (Pentothal, Hospira Inc., Lake Forest, IL)</td>
<td>Sedation, motion control, anxiolysis</td>
<td>Diagnostic imaging</td>
<td>Not recommended</td>
<td>PR: 25 mg/kg</td>
<td>PR: 10–15</td>
<td>PR: 60–120</td>
<td>Avoid in patients with porphyria. Because drugs cannot be titrated with the PR route, monitor closely for oversedation.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>PO/PR (&lt; 4 yr): 3–6 mg/kg, max of 100 mg</td>
<td>PO/PR: 15–60</td>
<td>PO/PR: 60–240</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>PO/PR (&gt; 4 yr): 1.5–3 mg/kg, max of 100 mg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Analgesic</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Fentanyl</strong> (Sublimaze, Akorn Inc., Lake Forest, IL)</td>
<td>Analgesia</td>
<td>Procedures with moderate pain</td>
<td>IV: 50 µg, may repeat q 3 min, titrate to effect</td>
<td>IV: 1 µg/kg per dose, may repeat q 3 min, titrate to effect</td>
<td>IV: 3–5</td>
<td>IV: 30–60</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Reversible with naloxone</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Dissociative Agent</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Ketamine</strong> (Ketalar, JHP Pharmaceuticals, Rochester, MI)</td>
<td>Analgesia, dissociation, amnesia, motion control</td>
<td>Procedures with moderate pain or requiring immobilization</td>
<td>IV: 1–1.5 mg/kg slowly over 30–60 sec, may repeat ½ dose q 10 min prn</td>
<td>IV: 1.5 mg/kg slowly over 30–60 sec, may repeat ½ dose q 10 min prn</td>
<td>IV: 1</td>
<td>IV: dissociation, 15; recovery, 60</td>
<td>Multiple contraindications.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>IM: 4–5 mg/kg, may repeat after 10 min (½ dose)</td>
<td>IM: 3–5</td>
<td>IM: dissociation, 15–30; recovery, 90–150</td>
<td></td>
</tr>
</tbody>
</table>
**TABLE 33.5** Drug Dosing Recommendations for PSA—cont’d

<table>
<thead>
<tr>
<th>DRUG</th>
<th>CLINICAL EFFECTS</th>
<th>INDICATIONS</th>
<th>ADULT DOSE</th>
<th>PEDIATRIC DOSE</th>
<th>ONSET (min)</th>
<th>DURATION (min)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Inhalational Agent</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nitrous oxide (Nitronox, Medical Gas Solutions Ltd., Flint, United Kingdom)</td>
<td>Anxiolysis, analgesia, sedation, amnesia (all mild)</td>
<td>Procedures requiring mild analgesia or sedation (age &gt; 4 yr)</td>
<td>Preset mixture with a minimum of 40% O₂ self-administered by a demand-valve mask (requires a cooperative patient)</td>
<td>Preset mixture for a cooperative child; continuous-flow nasal mask in an uncooperative child with close monitoring</td>
<td>&lt; 5</td>
<td>&lt; 5 following discontinuation</td>
<td>Requires specialized apparatus and gas scavenger capability. Several contraindications. Synergistic effect with recent opioids or sedative-hypnotics—use with caution in this setting.</td>
</tr>
</tbody>
</table>

| **Reversal Agents (Antagonists)** | | | | | | | |
| Naloxone (Narcan, Adapt Pharma Inc., Radnor, PA) | Opioid reversal | Opioid toxicity | IV/IM: 0.4–2 mg | IV/IM: 0.1 mg/kg per dose up to max of 2 mg/dose, may repeat q 2 min pm | IV: 2 | IV: 20–40 IM: 60–90 | If shorter acting than the reversed drug, serial doses may be required. |

| Flumazenil (Romazicon, Genentech USA Inc., San Francisco, CA) | Benzodiazepine reversal | Benzodiazepine toxicity | IV: 0.2 mg, may repeat q 1 min up to 1 mg | IV: 0.02 mg/kg per dose, may repeat q 1 min up to 1 mg | IV: 1–2 | IV: 30–60 | If shorter acting than the reversed drug, serial doses may be required. Do not use in patients chronically taking benzodiazepines, cyclosporine, isoniazid, lithium, propoxyphene, theophylline, tricyclic antidepressants. |

---

1. Alterations in dosing may be indicated depending on the clinical situation and the practitioner's experience with these agents. Individual dosages may vary when used in combination with other agents, especially when benzodiazepines are combined with opioids.

2. Use lower doses in geriatric patients and those with significant cardiopulmonary disease.

3. Midazolam is preferred over other benzodiazepines (e.g., diazepam, lorazepam) for procedural sedation and analgesia because of its shorter duration of action and multiple routes of administration.

4. Fentanyl is preferred over other opioids (e.g., morphine, meperidine) for procedural sedation and analgesia because of its faster onset, shorter recovery, and lack of histamine release.

5. Generally accepted contraindications to ketamine include age younger than 3 months; history of airway instability, tracheal surgery, or tracheal stenosis; procedures involving stimulation of the posterior pharynx; active pulmonary infection or disease (including active upper respiratory infection); cardiovascular disease, including angina, heart failure, or hypertension; significant head injury, central nervous system mass lesions, or hydrocephalus; glaucoma or acute globe injury; psychosis; porphyria; and thyroid disorder or thyroid medication.

6. Generally accepted contraindications to nitrous oxide include pregnancy (patient or personnel), preexisting nausea or vomiting, trapped gas pockets (e.g., middle ear infection, pneumothorax, bowel obstruction).

ED, Emergency department; FDA, US Food and Drug Administration; IM, intramuscularly; IN, intranasally; IV, intravenously; PO, orally; PR, per rectum; prn, as needed; PSA, procedural sedation and analgesia.

respiratory depression. Such respiratory depression from benzodiazepines, one must maintain continuous vigilance for apnea have occurred, thus underscoring the critical role of effects are exaggerated in the elderly. Deaths from undetected ethanol or other depressive drugs, especially opioids. These is dose dependent and greatly enhanced in the presence of properties. When carefully titrated intravenously, sedation is evident within 5 minutes with a duration of approximately 30 to 40 minutes.

**Adverse Effects.** When administered by skilled practitioners using standard precautions (see Box 33.6), the safety profile for midazolam is excellent. However, when administering benzodiazepines, one must maintain continuous vigilance for respiratory depression. Such respiratory depression is dose dependent and greatly enhanced in the presence of ethanol or other depressive drugs, especially opioids. These effects are exaggerated in the elderly. Deaths from undetected apnea have occurred, thus underscoring the critical role of continuous interactive and mechanical monitoring.

Benzodiazepines induce minimal cardiovascular depression. Although hypotension can occur, it is rare when the agents are carefully titrated. One reason that midazolam is ideal for painful procedures is its significant amnesic effect. Even though patients appear to feel pain during the procedure, it is often not remembered.

**Pentobarbital**

**Pharmacology.** Pentobarbital is a barbiturate capable of profound sedation, hypnosis, amnesia, and anticonvulsant activity in a dose-dependent fashion. It has no inherent analgesic properties. When carefully titrated intravenously, sedation is evident within 5 minutes with a duration of approximately 30 to 40 minutes.

**Adult Use.** Pentobarbital has no advantage over midazolam for adult PSA and is rarely used for this purpose.

**Pediatric Use.** Pentobarbital is the IV sedative of choice in many centers for diagnostic imaging in children. It is regarded as superior to midazolam or chloral hydrate for this indication. Pentobarbital, like midazolam, is available in multiple routes of administration.

**Adverse Effects.** Like other barbiturates, pentobarbital can lead to respiratory depression and hypotension because it is a negative inotrope.

**Ultrashort-Acting Sedative-Hypnotic Agents**

Ultrashort-acting sedatives (i.e., propofol, etomidate, thiopental, methohexitol) can rapidly produce potent sedation when administered intravenously, and all exhibit rapid awakening (< 5 minutes) after discontinuation of the drug. ED use of these agents (propofol in particular) for a variety of common short, painful procedures has expanded dramatically in recent years because their brief yet profound obtundation creates superlative conditions.

Substantial controversy surrounded the early administration of these agents for ED PSA. Proponents have cited their extremely rapid onset and recovery as enormous advantages over other sedatives. Critics have cited the level of continuous vigilance required to achieve a desired effect while simultaneously avoiding significant cardiopulmonary depression because these agents can result in rapid swings in levels of consciousness. Research that includes thousands of ED patients for propofol and hundreds for etomidate has subsequently demonstrated that the safety profiles of these agents are the same or better than those of other agents in the PSA pharmacopoeia.

**Propofol**

**Pharmacology.** Propofol is becoming an agent of choice for PSA in the ED because of its efficacy and safety profile (Fig. 33.3). When administered by IV bolus, its onset of action is typically within 30 seconds. The half-life for blood-brain equilibration is approximately 1 to 3 minutes, and its clinical effects typically resolve within 5 to 7 minutes. Longer procedures can be facilitated by repeated dosing. Patients are typically awake and alert within 15 minutes after discontinuation. Propofol exhibits inherent antiemetic and perhaps euphoric properties, and patient satisfaction is typically high. Propofol should be avoided in patients with known or suspected allergy to eggs or soy products.

**Adverse Effects.** Transient apnea and respiratory depression can occur with propofol but typically resolve spontaneously before intervention is necessary. Reported rates of assisted ventilation range from 0% to 4.6%. Similarly, transient hypotension (via direct negative inotropy as well as arterial and venous dilation) is common but typically resolves spontaneously without treatment. Injection site pain is noted less frequently in the ED setting than in the operating room, where higher doses are generally administered. Propofol usually causes pain on injection in awake individuals. Injecting slowly through a large vein, not a dorsal hand vein, and administering

**Figure 33.3** Propofol is emerging as an agent of choice for painful emergency department procedures. It is safe, effective, and very short acting. It can be cautiously combined with the short-acting narcotic fentanyl or with ketamine. All the safety caveats for procedural sedation and analgesia apply. The pain of a propofol injection can be minimized by choosing a large vein not on the dorsum of the hand and slowly injecting 2 to 3 mL of 2% lidocaine into the vein before injection of propofol.

Given the rapid onset and offset of propofol, consider (when available) an additional dedicated clinician (separate from the individual performing the procedure) to oversee administration of medication.
2 to 3 mL of 2% lidocaine slowly into the vein before propofol infusion will lessen the pain of injection.

**Etomidate**

*Pharmacology.* When administered by IV bolus, the onset of action of etomidate is typically within 30 seconds, and patients are usually awake and alert within 30 minutes after discontinuation.

*Adult Use.* Deep sedation can be achieved reliably with single loading doses. Dosing can be repeated as needed to enhance patients with volume depletion or cardiovascular compromise. Barbiturates also frequently cause hypotension at typical IV doses, so their use should be avoided whenever possible in

*Pediatric Use.* The safety and efficacy profile of etomidate appears to be similar to that in adults. For short-duration painful ED procedures, etomidate 0.2 mg/kg intravenously administered after fentanyl was associated with effective sedation, successful procedural completion, and readily managed respiratory events in children (oxygen supplementation, stimulation, or airway repositioning). Etomidate has also been compared to ketamine for orthopedic procedures in children, with equal rates of success and favorable safety profiles.

*Adverse Effects.* The primary adverse effects of etomidate are respiratory depression, myoclonus, nausea, and vomiting. Respiratory depression has been reported to be less common with propofol PSA than with methohexital, fentanyl/midazolam, or etomidate. Myoclonus is common yet generally benign, but it may be disconcerting and can interfere with the procedure. It consists of transient jerking or twitching movements that can be mistaken for seizure activity. Transient suppres-sion occurs with etomidate in septic patients but appears to lack clinical significance for single doses when used for ED PSA.

**Thiopental and Methohexital**

*Pharmacology.* Because of their lipid solubility, barbiturates are rapidly absorbed rectally. When given by the IV route, both thiopental and methohexital produce sedation within 1 minute. Clinical recovery is rapid (≈15 minutes) as a result of rapid redistribution from the central nervous system to the periphery.

*Adult Use.* There is limited published experience using these IV barbiturates for ED PSA, and propofol or etomidate would appear to be a better choice.

*Pediatric Use.* Rectal thiopental and methohexital can reliably produce sedation suitable for CT or MRI and obviates the need for IV access. Respiratory depression is unusual when using typical doses (see Table 33.5) but can occur. There is limited published experience using these IV barbiturates for ED PSA, and propofol would appear to be a better choice.

*Adverse Effects.* Barbiturates cause potent respiratory depression; in one ED report, apnea occurred in 10% of patients. Barbiturates also frequently cause hypotension at typical IV doses, so their use should be avoided whenever possible in patients with volume depletion or cardiovascular compromise.

---

**Analgesic Agents**

**Fentanyl**

Fentanyl is the most common opioid used for PSA because of its rapid onset, brief duration, rapid reversibility with naloxone, and lack of histamine release. It is often combined with other agents such as propofol, etomidate, and midazolam to provide additional effect and pain relief. The effects of fentanyl can be reversed immediately with naloxone should excessive sedation or respiratory depression occur. The longer-duration opioids morphine and meperidine are preferred for nonprocedural or preprocedural pain control and are frequently given initially for acute analgesia followed by fentanyl to facilitate the needed procedure. Although longer-acting opioids can be readily used for analgesia during PSA, they will be associated with longer recovery times and a higher incidence of histamine-related effects (e.g., nausea and vomiting, hypotension, pruritus). Fentanyl lacks these effects and is therefore preferred.

*Pharmacology.* Fentanyl is 100 times more potent than morphine and has no intrinsic anxiolytic or amnestic properties. A single IV dose has a rapid onset (<30 seconds) with a peak at 2 to 3 minutes and brief clinical duration (20 to 40 minutes). This increase in potency and onset of action is in part related to its greater lipid solubility, which facilitates passage of the drug across the blood-brain barrier. The effects of fentanyl can be rapidly and completely reversed with opioid antagonists (e.g., naloxone, nalbuphine).

*Adult Use.* Because of its pharmacokinetics, IV fentanyl is an ideal agent when analgesia is required for painful procedures; it can be easily and rapidly titrated. Because anxiolysis and sedation do not occur at low doses (1 to 2 µg/kg), concurrent administration of a pure sedative, most commonly midazolam, is advisable, especially in children (see BOX 33.6).

*Pediatric Use.* The combination of fentanyl and midazolam remains a popular PSA sedation regimen in children, with a strong safety and efficacy profile when both drugs are carefully titrated to effect. Any necessary level of mild to deep sedation can be achieved with these agents.

Fentanyl is also available in an oral transmucosal preparation. Although this novel and noninvasive delivery route obviates the need for IV access, titration is difficult and its efficacy is variable. Furthermore, the incidence of emesis is high (31% to 45%), and consequently this formulation has never become popular for PSA.

*Adverse Effects.* Like all opioids, fentanyl can cause respiratory depression. When used for PSA, standard interactive and mechanical monitoring is required. Because the opioid effect is most pronounced on the central nervous system respiratory centers, apnea precedes loss of consciousness. If apnea should occur, verbal or tactile stimulation should be attempted before the administration of opioid antagonists. As discussed earlier, caution must be exercised when using benzodiazepines and other PSA agents and opioids together because the risk for hypoxia and apnea is significantly greater than when either is used alone.

In the absence of significant ethanol intoxication, hypovolemia, or concomitant drug ingestion, hypotension is rare, even with very large doses of fentanyl (doses of 50 µg/kg are...
common in adult and pediatric cardiac surgery). Because of its safe hemodynamic profile, fentanyl is an ideal analgesic agent for use in critically ill or injured patients. In addition, nausea and vomiting are rare in comparison to analgesia with morphine or meperidine. A commonly observed reaction to fentanyl is nasal pruritus, and patients frequently attempt to scratch their nose during the procedure.

A rare side effect of fentanyl with potential for respiratory compromise is chest wall rigidity. This complication has not been problematic in the ED and is related to higher doses (>5 μg/kg as a bolus dose) than those used for PSA and has not been reported in any ED series. If it should occur, chest wall rigidity can usually be reversed with opioid antagonists, positive pressure ventilation, or both. Equipment for urgent pharmacologic paralysis should be available if reversal and positive pressure ventilation are unsuccessful.

**Diamorphine**

Diamorphine is a nasal opioid that is currently available in the United Kingdom, Australia, New Zealand, and Canada, but not in the United States. Diamorphine has an onset and duration of action similar to that of morphine; however, its higher water solubility permits potent doses to be delivered in the small (0.1 mL) volumes necessary for comfortable intranasal administration. In two studies of children and teenagers with fractures, intranasal diamorphine, 0.1 mg/kg, provided a similar level of analgesia with faster onset as IM morphine, 0.2 mg/kg. Intranasal spray administration was better tolerated than the injection, and there were no adverse events. Diamorphine may prove to be a useful initial analgesic for children and teenagers with acute pain, although in practice, an IV line would most likely be established to permit titration to full pain relief and PSA for any procedures needed (e.g., fracture reduction). The role of diamorphine in adults remains to be determined.

**Other Short-Acting Opioids**

Sufentanil, alfentanil, and remifentanil are other short-acting opioids that have a potential role in PSA. Currently, however, there is insufficient published experience to warrant their routine use. Although intranasal sufentanil, 0.75 μg/kg, appeared promising in one small pediatric trial, in another, doses of 1.5 μg/kg resulted in oxygen desaturation in 8 of the 10 children studied. This low toxic-to-therapeutic ratio and inability to titrate would appear to limit the utility of intranasal sufentanil by itself. However, a 2014 proof of concept trial using a lower dose of intranasal sufentanil (0.5 mcg/kg) with intranasal ketamine (0.5 mg/kg) showed promise with high patient acceptance and few adverse events. In the one published report of IV remifentanil with midazolam for PSA, there was an unacceptably high incidence of hypoxemia. Currently, there does not appear to be a clinically important advantage with these drugs versus fentanyl.

**Ketamine**

**Pharmacology.** Ketamine produces a unique state of cortical dissociation that permits painful procedures to be performed more consistently and effectively than with other PSA agents (Video 33.1). This state of dissociative sedation is characterized by profound analgesia, sedation, amnesia, and immobilization (Fig. 33.4) and can be rapidly and reliably produced with IV or IM administration. Ketamine is widely used worldwide and has demonstrated a remarkable safety profile in a variety of settings. Clinicians administering ketamine must be especially knowledgeable about the unique actions of this drug and the numerous contraindications to its use (see Table 33.5).

Ketamine differs from other PSA agents in several important ways. First, it uniquely preserves cardiopulmonary stability. Upper airway muscular tone and protective airway reflexes are maintained. Spontaneous respiration is preserved, although when administered intravenously, ketamine must be given slowly (over a period of 30 to 60 seconds) to prevent respiratory depression. Second, it differs from other agents in that it lacks the characteristic dose-response continuum to progressive titration. At lower doses ketamine produces analgesia and disorientation. However, once a dosage threshold (≈1 to 1.5 mg/kg intravenously or 3 to 4 μg/kg intramuscularly) is achieved, the characteristic dissociative state appears. This dissociation has no observable levels of depth, and thus the only value of ketamine “titration” is to maintain the presence of the state over time. Finally, the dissociative state is not consistent with formal definitions of moderate sedation, deep sedation, or general anesthesia (see Box 33.1) and must therefore be
considered from a different perspective than agents that exhibit the classic sedation continuum.\textsuperscript{13,15}

Ketamine is most effective and reliable when given intravenously or intramuscularly. Ketamine has a rapid circulation time when given intravenously, with onset of dissociation noted within 1 minute and effective procedural conditions lasting for approximately 10 to 15 minutes. When given intramuscularly, the same effect is achieved within 5 minutes, with effective procedural conditions for approximately 15 to 30 minutes. The typical duration from dosing to dischargeable recovery is 50 to 110 minutes when given intravenously and 60 to 140 minutes when given intramuscularly.\textsuperscript{14}

Like the benzodiazepines, ketamine undergoes substantial first-pass hepatic metabolism. As a result, oral and rectal administration results in less predictable effectiveness and substantially higher doses are required. Clinical onset and recovery are considerably longer than when given parenterally, and thus these routes are rarely used in the ED.\textsuperscript{100,146}

Ketamine can occasionally induce excessive salivation, and to combat this, some have historically coadministered atropine or glycopyrrolate. However, there is no evidence that either of these anticholinergic agents diminishes the risk for airway or respiratory adverse events,\textsuperscript{147–149} and their routine prophylactic use is no longer recommended.\textsuperscript{147} Instead, they can be reserved for treatment should excessive secretions occur (Fig. 33.5).

**Figure 33.5** Ketamine can occasionally induce excessive salivation, generally easily handled with minimal suctioning. Routine use of atropine or glycopyrrolate is not recommended; however, they can be administered should excessive secretions be noted.

**Adverse Effects.** In a meta-analysis of ketamine administration in 8282 pediatric patients, the overall incidence of airway and respiratory adverse events was 3.9%—primarily airway malalignment but also including transient laryngospasm (0.3%) and transient apnea (0.8%). None of these patients were intubated or had adverse sequelae.\textsuperscript{147} The frequency of such events was slightly higher when unusually high doses of ketamine were administered; otherwise, there was no clinically important association with age, doses within the typically recommended range, or other clinical factors.

Vomiting was noted in 8.4% overall in this same meta-analysis, with early adolescence representing the greatest risk and a lower incidence of emesis occurring in younger and older children.\textsuperscript{158} It occurs more frequently with the IM route than with the IV route, and there is no evidence to support any dose relationship within the usual range of clinically administered doses.\textsuperscript{158} When emesis occurs, it is typically late during the recovery phase when the patient is alert and can clear the airway without assistance.\textsuperscript{147} Vomiting does occur in some patients after discharge, including some who do not vomit in the ED.\textsuperscript{137} Although ondansetron prophylaxis (0.15 mg/kg up to a maximum of 4 mg intravenously) does reduce vomiting with ketamine, the magnitude of this effect is modest (number needed to benefit of 13), and thus it cannot be considered mandatory.\textsuperscript{161}

In 40 years of regular use there have been no documented reports of clinically significant ketamine-associated aspiration in patients without established contraindications.\textsuperscript{157} Because of its unique preservation of protective airway reflexes, ketamine may be preferred over other agents for urgent or emergency procedures when fasting is not ensured.\textsuperscript{13,152}

Mild agitation (whimpering or crying) during recovery was noted in 7.6% of children in the large meta-analysis, with
more pronounced agitation occurring in 1.4%. Such recovery reactions are not related to age, dose, or other factors to any clinically important degree, except for a higher incidence with subdissociative (<3 mg/kg intramuscularly) dosing.\textsuperscript{158} In contrast to traditional thinking, adolescents are not at substantially higher risk.\textsuperscript{158}

Despite the modest impact of prophylactic midazolam in reducing adult recovery agitation, in children, two controlled trials and a large meta-analysis failed to note even a trend toward a measurable benefit.\textsuperscript{158-160} Children have far fewer recovery reactions than adults do, and their reactions are milder when they do occur. Midazolam is therefore not recommended for routine prophylaxis but is optimally reserved to treat unpleasant ketamine-associated recovery reactions when they do rarely occur.\textsuperscript{147}

Ketamine is relatively contraindicated in patients with central nervous system masses, abnormalities, or hydrocephalus; however, there is no compelling evidence that it needs to be avoided in the setting of acute head trauma.\textsuperscript{147,162,163} Repeated reports that ketamine can increase intracranial pressure have prompted traditional caution against use of this drug in the setting of real or potential neurologic compromise,\textsuperscript{1,5,14,27,64-66} and there are case reports of deterioration in patients with hydrocephalus.\textsuperscript{167,168} However, newer suggestive evidence indicates that in most patients the resulting increases in pressure are minimal, assuming normal ventilation.\textsuperscript{162,163,167,169} and that the corresponding cerebral vasodilatory effect of ketamine may actually improve overall cerebral perfusion.\textsuperscript{162,163}

Dissociative sedation may represent a risk in patients with acute globe injury or glaucoma given the inconclusive and conflicting evidence of increased intraocular pressure with ketamine.\textsuperscript{170-174}

### Ketofol—Propofol Combination (Ketofol) for PSA in the ED

The PSA combination of ketamine and propofol, referred to by the portmanteau “ketofol,” is both popular and controversial.\textsuperscript{175} There is no doubt that ketofol is safe and effective in both adults and children;\textsuperscript{176-183} instead, the dispute is whether the combination exhibits any clinically important advantage over use of either drug alone.\textsuperscript{173} Although this question has not been definitively answered, a recent meta-analysis evaluating 6 trials using ketofol suggests that the adverse event profile of ketofol may be superior to propofol alone.\textsuperscript{184}

An important allure to ketofol proponents is how these two completely different sedatives balance each other’s deficits. Propofol is a superb sedative but lacks the analgesia that ketamine can amply provide. Ketamine mitigates propofol-induced hypotension, and propofol mitigates ketamine-induced vomiting and recovery agitation. The drugs exhibit synergistic and perhaps smoother sedation,\textsuperscript{176} and the combination has the theoretical benefits of minimizing the propofol dose and obviating the need for co-administered opioids.\textsuperscript{173}

Critics note that the existing ketofol literature fails to demonstrate superior procedural conditions or less respiratory depression than occurs with each drug alone\textsuperscript{176} and question the added complexity of administering two drugs when one is sufficient. They argue that the purported advantages related to hemodynamics and the total propofol dose are not clinically important. Ketofol uses a subdissociative dose of ketamine, and thus ketamine may just be replacing fentanyl as an analgesic rather than as a sedative.\textsuperscript{175}

Ketofol dosing is based on weight for both adults and children. One regimen uses an initial IV loading dose of ketamine of 0.5 mg/kg, followed by a 0.5 to 1.0 mg/kg IV propofol bolus, with subsequent titration of propofol alone.\textsuperscript{177-179} A second strategy mixes propofol and ketamine together 1:1 in the same syringe (both at 10-mg/mL concentrations) and then titrates in increments of 0.25 mg/kg of each drug to a usual effective dose of approximately 0.75 mg/kg.\textsuperscript{171,180,182,183}

Careful observation for respiratory depression is required, and the precautions listed for both agents should be stringently observed.

### Nitrous Oxide

**Pharmacology.** Inhaled nitrous oxide provides anxiolyis and mild analgesia. It is commonly dispensed at concentrations between 30% and 50%, with oxygen composing the remainder of the mixture. Nitrous oxide quickly diffuses across biologic membranes and, accordingly, has a rapid onset of action (30 to 60 seconds). Its maximum effect occurs after approximately 5 minutes, and the clinical effect wears off quickly on discontinuation. At typical PSA concentrations, there is preservation of hemodynamic status, spontaneous respirations, and protective airway reflexes.\textsuperscript{185-188} Nitrous oxide is widely used in dentistry at higher concentrations.\textsuperscript{189}

Nitrous oxide has an excellent safety profile; however, as a sole agent, it cannot reliably produce adequate procedural conditions.\textsuperscript{185-188} Given its relatively weak analgesic properties, in many cases nitrous oxide needs to be supplemented with an IV opioid or local or regional anesthesia (or both).

**Adult and Cooperative Child Use.** The safest method of administering nitrous oxide is via a self-administered demand-valve mask (Fig. 33.6).\textsuperscript{186-188} Patients must generate a negative pressure of 3 to 5 cm H2O within the handheld mask or mouthpiece to activate the flow of gas. They can thus self-titrates the dose by inhaling at will through the mask. Naturally, this will be effective only when the patient is cooperative. This technique provides a built-in fail-safe measure in that if patients become somnolent, the mask will fall from their face and gas delivery will cease.

Nitrous oxide can be used as an adjunctive anxiolytic during mildly painful procedures or during the administration of local or regional anesthesia for other procedures. It may also be administered during difficult or high-anxiety procedures such as pelvic examination or during difficult IV attempts.

A double-tank system is commonly used to deliver the nitrous oxide and oxygen mixture. The system relies on a mixing valve preset to deliver a fixed ratio and will deliver gas only when oxygen is flowing. The double-tank system contains a fail-safe device that automatically stops the flow of nitrous oxide when the oxygen supply is depleted.

**Uncooperative Child Use.** The primary limitation of self-administration is that it is ineffective in uncooperative patients, including most frightened young children. Continuous-flow nitrous oxide via a mask strapped over the nose or over the nose and mouth has been used in this population (Fig. 33.7).\textsuperscript{190-193} Nitrous oxide can effectively produce moderate or deep sedation when administered in this manner; however, this technique necessitates an additional clinician dedicated to continuous
it has the potential to increase the size of the closed space. This should be unlikely for short-term use in typical PSA concentrations.

Make sure that a scavenging system is in place to collect exhaled nitrous oxide, and take care to ensure compliance with occupational safety regulations. Avoid exposure of pregnant ED staff members to nitrous oxide because it is a known teratogen and mutagen.

Although the potential for abuse by ED staff exists, such abuse is rare if simple steps are taken. As with other agents, maintain a strict protocol of accountability. Add a simple locking device to the cylinders of gas. In addition, lock the delivery valve or mouthpiece in the same location as controlled substances.

**Antagonists**

Do not routinely administer reversal agents after the administration of opioids or benzodiazepines for PSA, but rather reserve them for the rare situations of oversedation or significant respiratory depression. The downside of reversal agents is that one must wait for the antagonist to dissipate before analyzing any residual effect of the PSA agent, with the caveat that long-acting PSA agents may last longer than the administered antagonist. Serious resedation after antagonism is not usually an issue with fentanyl or midazolam, but caution is advised, especially when large doses of PSA agents and large doses of antagonists have been used. Obviously, the length of observation varies with the amount of antagonist that has been administered.

**Naloxone**

Naloxone is an antagonist that competitively displaces opioids from opiate receptors. It rapidly reverses the analgesic and respiratory depressant effects of opioids. It may be administered intravenously, intramuscularly, subcutaneously, or even sublingually if needed, and dosing has been standardized for infants and children. Naloxone will not induce systemic opioid withdrawal symptoms in a patient without preexisting physiologic dependence. However, some patients will experience nausea with opioid reversal, and patients with persistent pain after their procedure will be quite uncomfortable. Rapid reversal may also lead to return of anxiety and stimulation of the sympathetic nervous system. If the situation permits, careful titration of small amounts of naloxone (0.1-mg aliquots...
been shown to substantially decrease the time of observation in the ED required for a patient undergoing PSA. Flumazenil lowers the seizure threshold and may rarely lead to life-threatening seizures. It should be avoided in patients with known benzodiazepine dependence, seizure disorder, cyclic antidepressant overdose, and elevated intracranial pressure. Flumazenil-induced seizures are generally associated only with drug overdose. Rapid reversal may also lead to return of anxiety and sympathetic stimulation. If the situation permits, careful titration of small amounts of flumazenil (0.1- to 0.2-mg aliquots intravenously) will reduce the risk for adverse effects and may permit partial rather than complete reversal. Observation times vary. For example, 1 hour after reversal will allow accurate assessment of residual sedation if less than 1 mg of flumazenil has been administered intravenously.

**Nalmefene**

Nalmefene is a long-acting opioid antagonist with a duration of action significantly longer than that of naloxone. Nalmefene may be given intravenously, intramuscularly, or subcutaneously, although intravenously is the preferred route. Intravenously, it can be titrated in incremental doses of 0.25 µg/kg every 2 to 5 minutes until the desired effect is attained. Although either naloxone or nalmefene will reverse the analgesia of opioids, naloxone is the preferred agent. For ED PSA, short-acting opioids such as fentanyl are commonly used, and administration of a reversal agent with a duration of action as prolonged as that of nalmefene does not confer any additional benefit. Furthermore, nalmefene would interfere with postprocedure control of pain with opioids.

**Flumazenil**

Flumazenil is a benzodiazepine antagonist that can promptly reverse benzodiazepine-induced sedation and respiratory depression. In the setting of PSA, flumazenil is a safe and effective method of reversing oversedation caused by benzodiazepines. It is not routinely used to reverse PSA because of the potential for resedation, and many clinicians prefer to allow patients to recover on their own. Flumazenil has not been shown to substantially decrease the time of observation in the ED required for a patient undergoing PSA. Flumazenil lowers the seizure threshold and may rarely lead to life-threatening seizures. It should be avoided in patients with known benzodiazepine dependence, seizure disorder, cyclic antidepressant overdose, and elevated intracranial pressure. Flumazenil-induced seizures are generally associated only with drug overdose. Rapid reversal may also lead to return of anxiety and sympathetic stimulation. If the situation permits, careful titration of small amounts of flumazenil (0.1- to 0.2-mg aliquots intravenously) will reduce the risk for adverse effects and may permit partial rather than complete reversal. Observation times vary. For example, 1 hour after reversal will allow accurate assessment of residual sedation if less than 1 mg of flumazenil has been used to reverse conscious sedation with midazolam.

**ACKNOWLEDGEMENT**

This chapter was previously superbly authored by Steven Green and Baruch Krauss and they are responsible for the majority of the current content.
REFERENCES


Principles of Wound Management

Richard L. Lammers and Kim N. Aldy

Management of acute traumatic wounds is one of the most common procedures in emergency medicine. Although many aspects of traumatic wound management remain controversial, the clinician can follow some basic principles to maximize the chance for successful healing. The purpose of this chapter is to give the clinician a general approach to wound care and to describe appropriate indications and techniques for wound management.

Wound care involves much more than closure of divided skin. The primary goal of wound care is not technical repair of the wound, but to provide optimal conditions so that the natural reparative processes of the wound may proceed. The cornerstones of wound care are cleaning, débridement, closure (when appropriate), and protection. The primary objectives in wound care are to:

1. Preserve viable tissue and remove nonviable tissue
2. Restore tissue continuity and function
3. Optimize conditions for the development of wound strength
4. Prevent excessive or prolonged inflammation
5. Avoid infection and other impediments to healing
6. Minimize scar formation
7. Provide suitable anesthesia during wound management

Patients who seek care in the emergency department (ED) because of acute wounds report that their top priorities include prevention of infection, return to normal function, a good cosmetic outcome, and minimal pain during repair.1,2 This chapter reviews the current strategies for attaining these goals.

BACKGROUND: WOUND HEALING

For centuries victims of wounds have commonly experienced inflammation, infection, and extreme scarring; in fact, these processes were considered part of normal wound repair. Emergency clinicians should have a basic understanding of the process of wound healing. Highlights of this complex phenomenon as they relate to clinical decision-making are presented here.

Wounds extending beneath the epithelium heal by forming scar tissue. Among the various proposed overlapping phases from injury to repair, epithelialization, fibroplasia, contraction, and scar remodeling/maturation constitute the main phases of this natural repair process.3-6

Inflammation is a beneficial response orchestrated by polymorphonuclear and mononuclear leukocytes that concentrate at the site of injury and phagocytose dead and dying tissue, foreign material, and bacteria in the wound, essentially performing a biologic débridement.7 As white blood cells die, their intracellular contents are released into the wound. In excessive amounts, the contents form the purulence that is characteristic of infected wounds. Some exudate is expected even in the absence of bacterial invasion, but infection with accumulation of pus interferes with epithelialization and fibroplasia, and impairs wound healing. Wounds contaminated with significant numbers of bacteria or foreign material may undergo a prolonged or persistent inflammatory response and not heal. Granuloma formation surrounding retained sutures is an example of chronic inflammation.

Although white blood cells remove debris within the wound, keratinocytes at the surface of the wound begin to migrate across the tissue defect during epithelialization. In most sutured wounds, the surface of the wound develops an epithelial covering impermeable to water within 24 to 48 hours. Surface debris, dead tissue, and eschar formation can impair this process. The epithelium thickens and grows downward into the wound along the course of skin sutures. Although there is some “adhesiveness” at the wound edges during the first few days, this is eventually lost because of fibrinolysis.

By the fourth or fifth day, newly transformed fibroblasts in the wound begin synthesizing collagen and protein polysaccharides, thereby initiating the stage of scar formation known as fibroplasia. Collagen is the predominant component of scar tissue.8 Wound strength is a balance between the lysis of old collagen and the synthesis of new collagen that “welds” the edges of the wound together. The amount of scar tissue that forms is influenced by physical forces (e.g., the stresses imposed by movement) acting across the wound. When the wound edges are approximated, either naturally or by mechanical closure within the first 24 hours, the wound can heal by “first intention”. In contrast, a wound with extensive tissue loss and not closed by sutures or other means heals by “secondary intention,” a combination of processes that include contraction, collagen formation, and epithelialization (Fig. 34.1).9 Contraction consists of movement of the skin edges toward the center of the defect, primarily in the direction of underlying muscle.

Significant gains in tensile strength do not begin until approximately the fifth day after the injury. Strength increases rapidly for 6 to 17 days, more slowly for an additional 10 to
is a wound that is either fully re-epithelialized or filled with an avascular scar.6

Decisions regarding the optimal time for suture removal and the need for continued support of the wound with tape are influenced by the following: (1) the tensile strength of the wound, (2) the period of scar widening, and (3) the cosmetically unacceptable effect of epithelialization along the suture track. Scars are quite red and noticeable at 3 to 8 weeks after closure. The appearance of a scar should not be judged before the scar is well into its remodeling phase. The cosmetic appearance of wounds 6 to 9 months after injury cannot be predicted at the time of suture removal.8 Therefore scar revision, if necessary, should be postponed until 6 to 12 months after injury.

One of the most important factors in predicting the cosmetic result of a wound is its location.9 In general, wounds on concave surfaces heal with better cosmetic results than do wounds on convex surfaces. Other factors that affect cosmesis include wound size, wound depth, and skin color.10 Small, superficial wounds in lax, light-colored skin, especially areas where the skin is thin, result in less noticeable scars. Wounds on convex surfaces look better after primary closure than after secondary healing. Static and dynamic forces, along with the propensity toward keloid formation, may influence the long-term cosmetic appearance of wounds more than the surgical skills of the clinician who repaired the wound.9 Repigmentation occurs over a period of 3 to 5 years, even in large wounds that heal by secondary intention.10 Patients should be advised to wear the primary wound closure for 14 days, and almost imperceptibly for as long as 2 years (Fig. 34.2). The strength of scar tissue never quite reaches that of unwounded skin, approaching a maximum of 80%.14 Although the process of collagen formation is essentially completed within 21 to 28 days, the scar widens for another month, and collagen continues to remodel and strengthen the wound for up to 1 year.15,17 The normal desirable outcome of this complex process

Figure 34.1 Steps in wound healing by first intention (left) and second intention (right). Note the large amounts of granulation tissue and wound contraction in healing by secondary intention.

Figure 34.2 Graphic representation of the various phases of wound healing. Note that the tensile strength of scar tissue never reaches that of unwounded skin. The values of tensile strength displayed are approximate and demonstrate the general concept of wound healing.
sunscreen over the scar, especially during the first few years after injury, because unprotected exposure to the sun can alter the pigmentation as the wound heals and may result in noticeably darker pigmentation than the surrounding skin.

INITIAL EVALUATION

The approach to management of a particular wound and the decision to close a wound immediately or after a period of observation is based primarily on factors that affect the risk for infection, and secondarily on cosmesis and long-term functionality. The history and physical examination should be directed toward identifying these factors. Some wounds may appear benign but conceal extensive and devastating underlying tissue damage. The following findings should alert the provider of a more complicated injury including an extremity wound caused by a roller or wringer device, a high-pressure injection gun, high-voltage electricity, heavy and prolonged compressive force, or the bite of a human or a potentially rabid animal. These findings radically alter the overall management of the patient. The American College of Emergency Physicians’ Clinical Policy for the Initial Approach to Patients Presenting with Penetrating Extremity Trauma provides a useful approach to the evaluation of all wounds.

History

In the initial evaluation of a wound, identify all the extrinsic and intrinsic factors that jeopardize healing and promote infection, including: (1) the mechanism of injury, (2) the time of injury, (3) the environment in which the wound occurred, and (4) the patient’s immune status.

Wound Age

In general, the likelihood of a wound infection increases with time, from the injury to definitive wound care. Definitive wound care does not always mean closing a wound. Some wounds should never be closed, such as small, contaminated lacerations on the bottom of the foot, whereas others can be closed many hours after the injury without increasing infection rates. A delay in wound cleaning is one of the most important factors in wound infection because it may allow bacteria contaminating the wound to proliferate. A delay of only a few hours in the treatment of a heavily contaminated wound can increase the risk for infection. Although no scientific data exist to fully answer the question, and there is no definitively accepted standard, it appears reasonable that most wounds that are not grossly contaminated can probably be closed safely 6 to 8 hours after injury, if the wound can be adequately cleaned. In contrast, some evidence suggests that wounds in highly vascular regions, such as the face and scalp, can be closed without increased risk for as long as 24 hours after injury. The “golden period”—the maximum time after injury that a wound may be closed safely without significant risk for infection—is an outdated term and not a fixed number of hours.

Many factors affect risk for infection, and closure decisions should not be based solely on time considerations. All data accumulated in the initial evaluation, both historical and physical, must be considered when making the decision to close a wound in a particular patient. Delayed primary closure is a reasonable alternative when there is clinical concern regarding closure at initial assessment. However, the techniques of wound care may extend the period; with skillful cleaning and debridement, a clinician may be able to convert a contaminated wound to a clean wound that can be safely closed in the ED. In most instances the clinician cannot totally eliminate infection risks but can favorably affect the probability and severity of infection with adequate wound care. Simply prescribing antibiotics in the hope that infection will somehow be averted is an unrealistic expectation, and can lead to antibiotic-resistant microorganisms.

In summary, any time frames offered are suggestions. Clinical judgment regarding potential for wound infection should be incorporated into the final decision-making process.

Other Historical Factors

Other factors that affect wound healing or the risk for infection include the patient’s age and state of health. Patient age appears to be an important factor in host resistance to infection, and individuals who are at the extremes of age, such as young children and the elderly, have the greatest risk for infection. Infection rates are reported to be higher in patients with concurrent medical conditions such as diabetes mellitus, immunologic deficiencies, malnutrition, anemia, uremia, congestive heart failure, cirrhosis, malignancy, alcoholism, arteriosclerosis, arteritis, collagen vascular disease, chronic granulomatous disease, smoking, chronic hypoxia, renal failure, and liver failure. Morbid obesity, in addition to other conditions in which patients are taking steroids or immunosuppressive drugs, or undergoing radiation therapy, are also at higher risk for infection. Shock, remote trauma, distant infection, bacteremia, retained foreign bodies, denervation, and peripheral vascular disease also increase wound infection rates and can slow the healing process.

Additional information pertinent to decision making in wound management includes the following:

- **Associated symptoms** (severe pain, paresthesia, or anesthesia)
- **Current medications** (specifically, anticoagulants, steroids, and immunosuppressive drugs)
- **Allergies** (especially to local anesthetics, antiseptics, analgesics, antibiotics, and tape)
- **Tetanus immunization status**
- **Potential exposure to rabies** (in bite wounds and mucosal exposures)
- **Potential foreign bodies** (embedded in the wound, especially when the mechanism of injury is unknown or the injury was associated with breaking glass or vegetative matter)
- **Previous injuries and deformities** (especially with extremity and facial injuries)
- **Associated injuries** (underlying fracture, joint penetration, crush injury, deep penetrating injury, animal or human bite)
- **Other factors** (availability for follow-up, patient understanding of wound care, compliance)

Physical Examination

All wounds should be examined for the amount of tissue destruction, degree of contamination, potential foreign bodies, and damage to underlying structures. The examiner should wear clean or sterile powder-free, latex-free gloves and avoid droplet contamination from the mouth. Examine the wound under good lighting and after the bleeding is controlled. Create a bloodless field, if necessary, with a tourniquet or sphygmonanometer (see Video 34.1 for equipment). Assess distal perfusion, motor and sensory function (prior to the
use of anesthetics), nearby tendon function, and document findings.

**Mechanism of Injury and Classification of Wounds**

The magnitude and direction of the injuring force and the volume of tissue on which the force is distributed determine the type of wound that is sustained. Three types of mechanical forces—shear, tension, and compression—produce soft tissue injury. The resulting disruption or loss of tissue determines the configuration of the wound. Wounds may be classified into six categories:

1. **Abrasions.** Wounds caused by forces applied in opposite directions result in the loss of epidermis and possibly dermis (e.g., grinding of skin against a road surface).

2. **Lacerations.** Wounds caused by shear forces produce a tear in tissues. Little energy is required to produce a wound by shear forces (e.g., a knife cut). Consequently, little tissue damage occurs at the wound edge, the margins are sharp, and the wound appears “tidy.” Tensile and compressive forces also cause separation of the tissues. The energy required to disrupt tissue by tensile or compressive forces (e.g., forehead hitting a dashboard) is considerably greater than that required for tissue disruption by shear forces because the energy is distributed over a larger volume. These lacerations often have jagged, contused, “untidy” edges, which have a higher risk for infection.

3. **Crush wounds.** Wounds caused by the impact of an object against tissue, particularly over a bony surface, compress the tissue. These wounds may contain contused or partially devitalized tissue and have higher rates of infection if not appropriately débrided.

4. **Puncture wounds.** Wounds with a small opening, whose depth cannot be entirely visualized, are caused by a combination of forces. They are particularly prone to infection because they are difficult to thoroughly clean and may retain foreign bodies.

5. **Avulsions.** Avulsions are wounds in which a portion of tissue is completely separated from its base, which is lost, or left with a narrow base of attachment (a flap). Shear and tensile forces cause avulsions. Skin tear avulsions often result from low-force friction or shearing forces that separate the layers of the skin (epidermis and dermis) from the underlying tissue. These wounds often occur in older adults as a result of the combination of minimal impact and vulnerable thin skin.

6. **Combination wounds.** Wounds can also have a combination of configurations. For example, a stellate laceration caused by compression of soft tissue against underlying bone can create wounds with elements of both crush injury and tissue separation. Missile wounds involve a combination of shear, tensile, and compressive forces that puncture, crush, and sometimes avulse tissue.

**Contaminants (Bacteria and Foreign Material)**

Infection rates in studies of traumatic wounds range from 1% to 38%. Numerous factors affect the risk for wound infection, but the primary determinants of infection are the amount of bacteria and dead tissue remaining in the wound, the patient’s immune response, and local tissue perfusion.

Essentially all traumatic wounds are contaminated with bacteria to some extent. The number of bacteria remaining in the wound at the time of closure is directly related to the risk for infection. A critical number of bacteria must be present in a wound before a soft tissue infection develops. In experimental wounds, fewer bacteria are required to infect wounds caused by a compressive force (≥10⁴ bacteria/g of tissue) than by a shear force (≥10⁶).

The nature and amount of foreign material contaminating the wound often determines the type and quantity of bacteria implanted. In general, visible contamination of a wound increases the risk for infection. The presence of undetected, reactive foreign bodies in sutured wounds almost guarantees an infection. Although bullet or glass fragments by themselves rarely produce wound infection, these foreign bodies may carry particles of clothing, gun wadding, or soil into the wound. Minute amounts of organic or vegetative matter, feces, or saliva carry highly infective doses of bacteria. The bacterial inoculum from human bites often contains 1 billion or more organisms per milliliter of saliva. Inorganic particulate matter, such as sand or road surface grease, usually introduces few bacteria into a wound and has little chemical reactivity. These contaminants are relatively innocuous. Soil that contains a large proportion of clay particles or a high organic content (such as that found in swamps, bogs, and marshes), however, has a high risk for infection.

In industrialized countries most wounds encountered in the practice of emergency medicine have low initial bacterial counts. If wound cleaning and removal of devitalized tissue are instituted before bacteria within the wound enter their accelerated growth phase (3 to 12 hours after the injury), bacterial counts will generally remain below the threshold needed to initiate infection.

**Devitalized Tissue**

Devitalized and necrotic tissue in a traumatic wound should be identified and removed. If left in place, it will allow bacteria to proliferate, inhibit leukocyte phagocytosis, and create an anaerobic environment suitable for certain bacterial species.

**Wound Location**

The anatomic location of the wound has considerable importance in the risk for infection. Bacterial densities on the surface of the skin range from a few thousand to millions per square centimeter. Distal extremity wounds are more at risk for the development of wound infections than are injuries on most other parts of the body. Levels of endogenous bacteria on hairy parts of the scalp, the mouth, axilla, nails, foreskin, perineum, and vagina may be high enough to serve as a potential source of infection in wounds in these locations. The high vascularity in areas such as the scalp, face, or perineum appears to offset the risk posed by the large numbers of endogenous microflora, whereas wounds in ischemic tissue are notoriously susceptible to infection.

**Underlying Structures**

If injury to underlying structures such as nerves, vessels, tendons, joints, bones, or ducts is found, the clinician may choose to forego wound closure and consult a surgical specialist (Fig. 34.3). Procedures such as joint space irrigation, reduction and débridement of compound fractures, neurorhaphy, vascular anastomosis, and hand flexor tendon repair are best accomplished in the controlled aseptic setting of the operating room,
CHAPTER 34  Principles of Wound Management 625

the procedure and assuring the patient that everything possible will be done to minimize pain. In general, all wound care should be performed with the patient in a supine position because fainting is a common occurrence during wound preparation and repair (Fig. 34.4). Relatives and friends can be allowed to stay with the patient, but they should be cautioned to report any dizziness or nausea, and they should remain seated throughout the procedure.

Anyone cleaning, irrigating, or suturing wounds should wear protective eyewear and gloves because virtually any patient may be seropositive for human immunodeficiency virus (HIV). Although mucosal exposure to blood or tissue products that are contaminated by HIV is considered a relatively low risk for subsequent infection, universal precautions are currently recommended, and a mask should be considered. Minimal aseptic technique requires the use of gloves during the cleaning procedure.

Thorough cleansing of bacteria, soil, and other contaminants from a wound cannot be accomplished without the patient's cooperation. Scrubbing most open wounds is painful, and the patient's natural response is to withdraw the injured area away from the provider. Local or regional anesthesia should precede examination and cleansing of a wound (Video 34.2). Despite adequate anesthesia, the patient may be too apprehensive to cooperate. If reassurance does not alleviate the fears of young children, consider both sedation and physical restraining devices.

Mechanical Scrubbing

Scrubbing the internal surface of a wound is controversial. Although scrubbing a wound with an antiseptic-soaked sponge decreases the risk of infection by removing foreign particulates, bacteria, and tissue debris, an abrasive sponge may inflict more damage on the tissue. As the amount of damage caused by scrubbing correlates with the porosity of the sponge, a fine-pore sponge (e.g., 90 pores/inch) should be used to minimize tissue abrasion. Mechanical scrubbing should be reserved for wounds contaminated with significant amounts of bacteria or foreign material (Fig. 34.5A and B), or should be considered if irrigation alone is ineffective in removing visible contaminants from a wound.
Wound Cleansing: Mechanical Scrubbing and Irrigation

Grossly contaminated wounds such as this need to be thoroughly cleansed prior to repair, by mechanical scrubbing and/or irrigation. Reserve mechanical scrubbing for wounds contaminated with significant amounts of bacteria or foreign material. Use a fine-pore sponge (e.g., 90 pores/inch) to minimize tissue abrasion.

Top, ZeroWet splash shield attached to the end of a syringe. The shield is held near or against the skin and protects the user from splashes from the high-pressure laminar flow nozzle. Bottom, a makeshift splash protector can be made from a medication cup.

Tap water and nonsterile gloves are commonly used to copiously irrigate wounds of the extremities by taking advantage of the volume and force parameters from the faucet. Anesthetize lacerations before cleaning. Be mindful of the potential for the patient to faint.

Use of the ZeroWet device. Note that the clinician is holding the laceration open with forceps to allow irrigation of the deep structures.

For foot wounds, the patient is placed on a stretcher and wheeled to the sink.

Figure 34.5 Wound cleansing: mechanical scrubbing and irrigation. Wound irrigation with tap water is acceptable.
Initially scrub a wide area of skin surface surrounding the wound with an antiseptic solution to remove contaminants that might be transferred into the wound by instruments, suture material, dressings, or the clinician’s gloved hand during wound management. It is important to remove all nonabsorbable particulate matter; any such material left in the dermis may be retained in the healed tissue and result in a disfiguring “tattoo” effect. Detergents have an advantage over saline in that they minimize friction between the sponge and tissue, thereby limiting damage to tissue during scrubbing. Detergents also dissolve particles and thus help dislodge them from the surface of the wound. Unfortunately, many of the detergents available are toxic to tissues. Use caution when considering detergents.

**Antiseptics During Cleaning**

For many years, antiseptic solutions have been used for their antimicrobial properties in and around wounds (Table 34.1). Studies on the use of antiseptics in wounds demonstrate that there is a delicate balance between killing bacteria and injuring tissue. Intact skin can withstand strong microbicidal agents, whereas leukocytes and the exposed cells of skin and soft tissue can be damaged by these agents.

Povidone-iodine (Betadine [Purdue Products L.P., Stamford, CT]) is widely available as a 10% stock solution. Although the undiluted solution may be used to prepare the skin surrounding a wound, it may be harmful to some tissue; therefore it should not be placed within the interior of the wound. Diluted povidone-iodine solution in concentrations of less than 1% appears to be safe and effective for cleaning contaminated wounds, but the precise concentration that provides the most benefit is unclear. Even dilute povidone-iodine may be particularly irritating when used for scrubbing contaminated wounds. In contrast, povidone-iodine surgical scrub (Betadine scrub, 7.5% iodine) and hexachlorophene (pHisoHex [sanofi-aventis U.S. LLC, Bridgewater, NJ]) both contain anionic detergents that are harmful to tissues. In vitro studies have demonstrated that chlorhexidine gluconate–alcohol (Hibiclens [Mölndlycke Health Care US, LLC, Norcross, GA]) is toxic to both fibroblasts and keratinocytes. When diluted to decrease its cytotoxic effect on these cells, it is no longer able to sufficiently kill *Staphylococcus aureus*. Its effect on the actual wound infection rate is unknown. Hydrogen peroxide is hemolytic, and there is little reason to use it except to clean surrounding skin encrusted with blood and coagulum, or to soak off adherent blood-saturated dressings. Peroxide should not be used on granulation tissue because oxygen bubbles lift newly formed epithelium off the wound surface.

Nonantiseptic, nonionic surfactants are attractive alternatives to these toxic cleansing agents. In contrast to antiseptic solutions, these preparations cause no tissue or cellular damage, leukocyte inhibition, or impairment of wound healing. The solutions cause no corneal injury, have low risk of conjunctival irritation, and do not cause pain on contact with the wound. Poloxamer 188 (Shur-Clen [ConvaTec, Greensboro, NC] or Pluronic F-68 [Thermo Fisher Scientific, Grand Island, NY]), a group of pluronic polyols, is nontoxic but has no antibacterial activity. Scrubbing experimental wounds with Shur-Clen 1% solution has reduced infection rates (though not statistically better than normal saline or povidone-iodine), thus proving it has the ability to cleanse a wound effectively and atraumatically. Pluronic polyols may be a good choice if the wound is near mucous membranes. The use of antiseptic preparations in

### Table 34.1 Summary of Agents Used for Wound Care

<table>
<thead>
<tr>
<th>AGENT</th>
<th>BIOLOGIC ACTIVITY</th>
<th>TISSUE TOXICITY*</th>
<th>SYSTEMIC TOXICITY*</th>
<th>POTENTIAL USES</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Povidone-iodine surgical scrub (Betadine 7.5%)</td>
<td>Virucidal; strongly bactericidal</td>
<td>Detergent component toxic to wound tissue</td>
<td>Painful in open wounds</td>
<td>Hand cleanser</td>
<td>Iodine allergy possible</td>
</tr>
<tr>
<td>Povidone-iodine solution (Betadine 10%)</td>
<td>Virucidal; bactericidal</td>
<td>Potentially toxic at full strength; 1% solution has no significant tissue toxicity</td>
<td>Extremely rare</td>
<td>Wound periphery cleanser; dilute to &lt;1% for wound irrigation</td>
<td>Dilute 10:1 (saline to Betadine) if used to irrigate wounds</td>
</tr>
<tr>
<td>Chlorhexidine gluconate (Hibiclens)</td>
<td>Bactericidal</td>
<td>Toxic to tissues, including eyes</td>
<td>Extremely rare</td>
<td>Hand cleanser</td>
<td>Avoid in open wounds, eyes, or ears</td>
</tr>
<tr>
<td>Poloxamer 188 (Shur-Clen); Pluronic F-68</td>
<td>No antibacterial or antiviral activity</td>
<td>None known; does not inhibit wound healing</td>
<td>None known</td>
<td>Wound cleanser</td>
<td>Nontoxic in wounds and eyes</td>
</tr>
<tr>
<td>Hexachlorophene (pHisoHex)</td>
<td>Bacteriostatic against gram-positive bacteria</td>
<td>Potentially toxic to wound tissue</td>
<td>Possibly teratogenic with repeated use</td>
<td>Alternative hand cleanser</td>
<td>Systemic absorption causes neurotoxicity</td>
</tr>
<tr>
<td>Hydrogen peroxide</td>
<td>Very weak antibacterial agent</td>
<td>Toxic to tissue and red cells</td>
<td>Extremely rare</td>
<td>Wound periphery cleanser</td>
<td>Foaming activity removes surface debris and coagulated blood</td>
</tr>
</tbody>
</table>

*Based largely on in vitro studies and animal data.
wounds can be considered in high risk, contaminated wounds where the benefits outweigh the risks.

Irrigation

It is important to distinguish between skin antiseptics and irrigating solutions. As a general rule, commercially available antiseptics should be used only to clean intact skin and not exposed wound surfaces. Most open wounds can be irrigated effectively with copious amounts of saline or tap water. Concerns about introducing infection by the use of common tap water are unfounded.

Properly performed irrigation (Video 34.3) is effective in removing particulate matter, bacteria, and devitalized tissue that is loosely adherent to the edges of the wound and trapped within its depths. The effectiveness of irrigation is primarily determined by the hydraulic pressure at which the irrigation fluid is delivered.

Low-pressure irrigation is defined at 1 to 2 psi, whereas high-pressure irrigation is greater than 8 psi. Port devices spiked into plastic intravenous bags that are squeezed by hand to deliver a stream of fluid, bulb syringes, or gravity flow irrigation devices all deliver fluid at very low pressures.

Although irrigation at either low or high pressures will reduce levels of Staphylococcus aureus, higher pressures should be more effective in rinsing wounds of small particulate matter. There appears to be no significant differences between tap water and saline delivered at high pressures. Although irrigation may not be required for low-risk, highly vascular, uncontaminated facial and scalp wounds, randomized, prospective trials are needed to answer this question.

Wound irrigation is best achieved with a large-volume syringe, 35 mL or 65 mL, and an 18- or 19-gauge catheter or needle to deliver irrigation volumes of at least 250 mL. The pressure that can be delivered with a syringe varies with the force exerted on the plunger and the internal diameter of the attached needle. A simple irrigation assembly consisting of a 19-gauge plastic catheter or needle attached to a 35-mL syringe produces 25 to 40 psi when the barrel of the syringe is pushed with both hands.

High-pressure irrigation can also be achieved through IV tubing by placing a pressure cuff around a saline bag and inflating it to 400 mm Hg. A standard running faucet can also generate approximately 45 psi. Irrigating extremity lacerations by holding them under a faucet of tap water is a common and accepted practice for wound preparations, and fears of introducing infection with a nonsterile irrigant are unfounded. (Note: One must anticipate potential fainting if a patient stands near a sink for faucet water irrigation.) These high-pressure irrigation systems remove significant numbers of bacteria and a substantial amount of particulate matter from the wound surface. Care must be taken to ensure that any method of irrigation is not exerting too much pressure because tissue damage can occur at pressures of 70 psi. Irrigation should continue until all visible, loose particulate matter has been removed. Warmed irrigating solutions are more comfortable for patients, even after the wound is anesthetized.

A potential complication of wound irrigation is that infectious material can be splashed into the face of the clinician, even when the tip of the irrigation device is held below the surface of the wound. Several commercial devices are available to contain the splatter, including devices that fit on the end of a syringe (see Fig. 34.5C and D) and devices that fit on the screw top of saline bottles (for this particular device the company reported variable pressure from 4 to 15 psi). An alternative strategy is to pierce the base of a small medicine cup with a shorter, large-bore needle. The cup can be placed upside down to cover the area to be irrigated and the syringe with the 19-gauge needle can be inserted through the base of the cup (see Fig. 34.5C). The wound should be positioned to allow continuous drainage of fluid during irrigation by any method.

Antibiotic Solutions for Irrigation

A variety of antibiotic solutions have been instilled directly into wounds or used as irrigation solutions, including ampicillin, a neomycin-bacitracin-polymyxin combination, tetracycline, penicillin, kanamycin, and cephalothin. Although there have been no reports of topical sensitization or toxic tissue levels of the antibiotic, studies have found inconsistent effectiveness in reducing infection rates. The indications for using antibiotic solutions to clean wounds have not been defined, and this practice is not considered standard.

Recommendations for Cleaning the Wound

The prerequisites for any wound-cleaning technique are a calm or sedated patient, satisfactory anesthesia, and thorough cleaning of the skin surface adjacent to the wound. The primary goal of wound cleaning is to rid the wound of major contaminants and lower the infective dose of bacteria (Video 34.4). Two strategies are recommended. A contaminated or “dirty” wound can be irrigated, or both scrubbed and irrigated, with a 0.9% saline solution. As an alternative, the wound can be scrubbed with pluronic polyols and irrigated with a normal saline solution. Only pluronic polyols or saline should be used near the eyes. Perform scrubbing with a soft, fine-pore sponge, and use high-pressure techniques for irrigation. Avoid using hydrogen peroxide on open wounds. Either gentle scrubbing with pluronic polyols and normal saline high-pressure irrigation or irrigation alone appears to be a satisfactory method for cleaning minimally contaminated wounds.

A sterile bottle of saline solution can be used to irrigate a wound, but once the bottle is opened, bacterial contamination occurs quite rapidly and it should be discarded after use. Patients frequently irrigate their wounds with tap water before going to the ED. Many clinicians routinely irrigate wounds, especially extremity wounds, with tap water instead of sterile saline, and infection rates have been found to be comparable to that of saline irrigation (see Fig. 34.5E and F). The advantage of using tap water from a faucet is that large volumes of irrigant can be quickly applied to an open wound at the high pressures required to remove debris and bacteria. The disadvantages of this technique are that irrigation pressures are difficult to control, and the patient may faint if allowed to stand at a sink. If tap water is used to irrigate wounds, it may also be safely delivered through a syringe at the bedside.

Preparation for Wound Closure

Before debridement or wound closure, prepare and drape the wound. Avoid shaving body hair because preoperative wounds that were shaved demonstrated higher infection rates. For wounds in hair-bearing areas, hair can be removed by clipping if it interferes with the procedure. Stubborn hairs that repeatedly invade the wound during suturing can be coated with petrolatum jelly or a water-soluble ointment to keep them out of the field. Do not shave eyebrows because critical landmarks for exact approximation of the wound edges would...
be lost. Although shaved eyebrows will grow back eventually, shaving also produces an undesirable cosmetic effect. Clip hair rather than shaving it.

Disinfect the skin surface adjacent to the wound (not the wound itself) with a standard 10% povidone-iodine or chlorhexidine gluconate (Hibiclens) solution. Paint the solution widely on the skin surrounding the wound, but do not allow it to seep into the interior of the wound itself (Fig. 34.6A).

After hand washing, the clinician and any assistants involved in the procedure must wear sterile or nonsterile but clean gloves. Sterile gloves are not required, and their use does not decrease infection rates. However, sterile gloves fit better on hands and result in better control of instruments and sutures. Wear face masks, especially if you have an upper respiratory bacterial infection. Because droplets of saliva may leak even from around the edges of a face mask, avoid talking

**Figure 34.6** Wound preparation and exploration.

1. This patient punched out a window, a setup for retained glass.
2. Do not explore this laceration with a finger. Instead, use radiographs and instruments to search for foreign bodies (the arrow defines a large piece of glass that could cut the examining finger).
3. Many patients are unaware of the presence of large foreign bodies in a deep wound, although they should be questioned about the possibility or sensation of retained material.
in proximity to the wound. Use a single fenestrated drape or multiple folded drapes around the wound site. Place a sterile glove on a patient who has a hand wound to provide a sterile field in lieu of a fenestrated drape. The area to be sutured can then be exposed by cutting the glove, and the extremity can be placed on a sterile towel (see Fig. 34.6B). This technique provides a clean field without the need to continually adjust the drape or to operate through a small opening.

Explore the entire depth and the full extent of every wound to locate hidden foreign bodies, particulate matter, bone fragments, or any injuries to underlying structures that may require repair (see Fig. 34.6C and D). Avoid the common temptation to initially explore wounds and assess their characteristics with a finger in search of a foreign body (see Fig. 34.6E) because embedded glass, metal fragments, or sharp pieces of bone may cut the clinician and cause exposure to blood-borne infections. Visualize directly with good lighting in a bloodless field, explore the wound with a metal probe, and use radiographs as safer approaches to wound exploration. Despite these measures, lacerations through thick subcutaneous adipose tissue are treacherous because large amounts of particulate matter can be completely obscured in deep folds of soft tissue. Unless a careful search is undertaken, contaminants may be inadvertently left in the depths of a sutured wound, resulting in infection. Some clinicians are reluctant to extend lacerations to properly clean or explore them; however, opening the wound to permit adequate visualization is often necessary for successful wound exploration.

Débridement
Débridement of foreign material and devitalized tissue is of undisputed importance in the management of a contaminated wound. Use this technique to remove tissue embedded with foreign matter or bacteria. Devitalized tissue impairs the ability of the wound to resist infection and also prolongs the period of inflammation. Débridement also creates a tidy, sharp wound edge that is easier to repair and results in a more cosmetically acceptable scar.

If the wound is already clean and the edges are viable, sharp débridement may not improve the outcome. Irregular wounds have greater surface areas than linear lacerations. Because skin tension is distributed over a greater length, the width of the scar is usually less with jagged wounds than if the wound is converted to an elliptical defect with tidy edges. If the edges are devitalized or contaminated though, the wound edges must be débrided (Fig. 34.7A and B). To avoid a wide scar in this situation, undermine the wound, and consider judicious removal of soft tissue within the wound to decrease the tension between the wound edges.

Excision
Excision is the most effective type of débridement because it converts a contaminated traumatic wound into a clean wound. If significant contamination occurs in areas in which there is laxity of tissues, and if no important structures such as tendons or nerves lie within the wound, the entire wound may be excised (see Fig. 34.7C). Complete excision of grossly contaminated wounds such as animal bites allows the primary closure of such wounds with no greater risk of infection than occurs with relatively unimicated lacerations.

When a puncture wound is excised, make the axis of the excision parallel to a wrinkle, a skin line, or a line of dependency. Make the long axis of this lenticular excision three to four times as great as the short axis (see Fig. 34.7D). Premark the skin with a surgical marking pen or by making a superficial “scoring” mark (cutting only down to the epidermis) around the wound with a No. 15 scalpel blade. Place tension on the surrounding skin with a finger or a skin hook. While steadying your hand on the table or on the patient, use the No. 15 blade and cut through the skin at right angles or at slightly oblique angles to the surface of the skin. If complete excision of the entire depth of the wound is not necessary, use tissue scissors to cut the edge of the wound by following the path premarked in the epidermis with the scalpel blade. If complete excision is desired, incise each wound edge past the deepest part of the wound. Excise and remove the wedge of tissue carefully without contaminating the fresh wound surface.

Plan the excision carefully because excessive removal of tissue can create a defect that is too large to close. Wounds on the trunk, the gluteal region, or the thigh are amenable to excision. In contrast, simple excision of a wound on the palm or the dorsum of the nose will make approximation of the resulting surgical wound edges difficult. In hair-bearing areas of the face, particularly through the eyebrows, angle the incision parallel to the angle of the hair follicles to avoid linear alopecia (see Fig. 34.7E).

Selective Débridement
Complete excision is impossible for most wounds because of insufficient skin elasticity and area, so selective débridement must be used. Stellate wounds and wounds with an irregular, meandering course have greater surface area and less skin tension per unit length than do linear lacerations. In some cases, excision of an entire wound would result in the loss of too much tissue (i.e., produce a gaping defect and excessive tension on the edges of the wound when closed). Avoid this problem by selective débridement and approximataion of the irregular wound edges. This technique involves sharp débridement of devitalized or heavily contaminated tissue in the wound piece-by-piece and eventual matching of one edge of the wound to the other. Selective débridement is time-consuming but preserves more of the surrounding viable tissue.

Identifying devitalized tissue in a wound remains a challenging problem. Tissue with a narrow pedicle or base, especially distally based narrow flaps on extremities, is unlikely to survive and should be excised. Approximately 24 hours after the wound occurs, a sharp line of demarcation often occurs and can help distinguish devitalized skin and viable skin, but in most fresh wounds, usually only a subtle bluish or blanched discoloration is present. Try to predict tissue viability by comparing the capillary refill of injured tissue with that of the adjacent skin. If circulation is adequate, viable tissue also becomes hyperemic soon after the release of a proximal tourniquet.

In heavily contaminated wounds, especially those with abundant adipose tissue, remove all exposed fat and all fat impregnated with particulate matter. The subcutaneous adipose tissue attached to large flaps or to avulsed viable skin should be débrided before reapproximation of the wound; removal of this fatty layer allows better perfusion of the flap or the graft. Contaminated bone fragments, nerves, and tendons are almost never removed. Every effort should be made to clean these structures and return them to their place of origin because they may be functional later. Fascia and tendons perform important functions despite potential loss of viability. If they can be cleaned adequately, these tissues should not be débrided.
Control of Hemorrhage

Hemostasis is essential at any stage of wound care. Persistent bleeding obscures the wound and hampers exploration and closure of the wound. If bleeding is not a problem before wound débridement, it may become a complication during the other. After débridement or excision, irrigate the wound again to remove any remaining tissue debris.

Wound Débridement

They may be left in wounds as free grafts and be covered by viable flaps of tissue. Instruments generally required for débridement include two fine single- or double-pronged skin hooks, a scalpel with a No. 15 blade, tissue scissors, hemostats, and a small tissue forceps. Using gentle tissue pressure, stabilize the jagged wound edges with skin hooks or forceps, and use the scalpel or scissors to cut away devitalized tissue from one end of the wound to the other. After débridement or excision, irrigate the wound again to remove any remaining tissue debris.

Figure 34.7 Wound débridement.
cleaning or after the edges of the wound are excised. If it is significant, hematoma formation in a sutured wound can separate the wound edges, impair healing, and cause dehiscence or infection.

Several practical methods of achieving hemostasis are available. Provide direct pressure with gloved fingers, gauze sponges, or packing material, and elevate the wound. This technique is usually effective in immediately controlling a single bleeding site or a small number of sites until the cut ends of vessels constrict and coagulation occurs. In a patient with multiple injuries and several urgent problems, control hemorrhage temporarily with a compression dressing. Apply several absorbent sponges directly over the bleeding site and secure them in place with an elastic bandage (e.g., Ace wrap) or elastic adhesive tape (Elastoplast). Then apply pressure with the elasticity of the bandage and elevate the bleeding part. Wound care can then be deferred while the clinician attends to more pressing matters.

Although simply crushing and twisting the end of a small vessel with a hemostat avoids dissection further into the wound, this method provides unreliable hemostasis. Ligation of the vessel with fine absorbable suture material is preferred. Clamp the bleeding ends of vessels with fine-point hemostats to provide immediate hemostasis. Because nerves often course with these vessels, clamp them only under direct visualization. The tip of the hemostat should project beyond the vessel to hold a loop of a ligature in place (Fig. 34.8A). With an assistant lifting the handle of the hemostat, pass a synthetic 5-0 or 6-0 absorbable suture around the hemostat from one hand to the other. Tie the first knot beyond the tip of the hemostat. Once the suture is securely anchored on the vessel, release the hemostat. Three knots are sufficient to hold the ligature in place. Cut the ends of the suture close to the knot to minimize the amount of suture material left in the wound.

Ligate vessels with diameters greater than 2 mm. Vessels smaller than 2 mm that bleed despite direct pressure can be controlled by pinpoint, bipolar electrocautery. A dry field is required for an effective electrical current to pass through the tissues. If sponging does not dry the field, use a suction-tipped catheter. Minimize trauma by using fine-tipped electrodes to touch the vessel, or touch the active electrode of the electrocautery unit to a small hemostat or fine-tipped forceps while gripping the vessel. Keep the power of the unit to the minimum level required for thrombosis of the vessel. Self-contained, sterilizable, battery-powered coagulation units are alternatives to electrocautery. These devices cauterize vessels by the direct application of a heated wire filament. Although these units may damage more surrounding tissue than electrocautery units do, they are compact, simple, and well suited for use in the ED (see Fig. 34.8B).

A cut vessel that retracts into the wall of the wound may frustrate attempts at clamping, ligation, or cauterization. First, control bleeding by downward compression on the tissue. Pass a suture through the tissue twice via a figure-of-eight or horizontal mattress stitch, and then tie it. This stitch will constrict the tissue containing the cut vessel (see Fig. 34.8C).

Large superficial varicosities may bleed spontaneously or from minor trauma. They may bleed profusely, especially when the patient stands up and increases venous pressure. Use a simple figure-of-eight suture to halt the bleeding (see Fig. 34.8D).

Epinephrine is an excellent vasoconstrictor. Place topical epinephrine (1:100,000) on a moistened sponge and apply it to a wound to reduce the bleeding from small vessels. When combined with local anesthetics, such as lidocaine with epinephrine, concentrations of 1:100,000 and 1:200,000 prolong the effect of the anesthetic and provide some hemostasis in highly vascular areas. Use vasoconstrictors only in situations in which widespread small-vessel and capillary hemorrhage in a wound cannot be controlled by direct pressure or cauterization. Hemostasis of a specific vessel may be achieved by directly injecting the soft tissues around the base of the bleeder with a small amount of lidocaine with epinephrine solution, even though the wound has previously been anesthetized. The combination of pressure and vasoconstriction may halt the bleeding long enough for the vessel to be ligated or cauterized, or to allow the wound to be closed and a compression dressing applied.

Fibrin foam, gelatin foam, and microcrystalline collagen may be used as hemostatic agents. Their utility is limited in that vigorous bleeding will wash the agent away from the bleeding site. Their greatest value may be in packing small cavities from which there is constant oozing of blood. Another novel hemostasis technique uses 2-octyl-cyanoacrylate (Dermabond [Ethicon, Cincinnati, OH]) to achieve hemostasis of skin avulsions on the fingertips, which are known to bleed aggressively. A tourniquet should first be used to minimize bleeding, and moisture should be dabbed away from the wound. Then the Dermabond can be layered on at least twice. This method is an off-label, anecdotal use of this product, and the consequence of exposing dermal skin layers directly to it have not been evaluated to date.

Do not spend excessive time attempting to tie off several small bleeding vessels while the patient slowly exsanguinates (Fig. 34.9). In highly vascular areas such as the scalp, it is sometimes best to suture the laceration after exploration and irrigation of the wound despite active bleeding; the pressure exerted by the closure will usually stop the bleeding. If bleeding is too brisk to permit adequate wound evaluation and irrigation, control hemorrhage by clamping and everting the galea or dermis of each wound edge with hemostats (Fig. 34.10A and B). Raney clamps or a large hemostat is an excellent way to stop scalp bleeding; they are used during neurosurgical procedures (see Fig. 34.10C). If the edge of the entire scalp is compressed the bleeding will generally stop. In the majority of simple wounds with persistent but minor capillary bleeding apposition of the wound edges with sutures, followed by a compression dressing, provides adequate hemostasis.

**Tourniquets**

If bleeding from an extremity wound is refractory to direct pressure, electrocauterization, or ligation, or if the patient has exsanguinating hemorrhage from the wound, place a tourniquet proximal to the wound to control the bleeding temporarily. Tourniquets are also helpful in examining extremity lacerations by providing a bloodless field. However, tourniquets can cause injury in three ways:

1. They can produce ischemia in an extremity.
2. They can compress and damage underlying blood vessels and nerves.
3. They can jeopardize the survival of marginally viable tissue. Although problems rarely develop from the use of tourniquets in routine wound care, potential problems can be minimized if (1) a limit is placed on the total amount of time that a tourniquet is applied, and (2) excessive tourniquet pressure is avoided. It is also imperative that all tourniquets be removed.
Hemorrhage Control

A
When tying off a bleeding vessel, the tip of the hemostat should project beyond the clamped vessel. An assistant raises the handles of the hemostat as a ligature is passed under them.

B
Battery-powered cautery can be used to coagulate minor subcutaneous bleeders.

C
Ligation of a retracted, bleeding vessel. 1, Horizontal mattress technique. 2, Figure-of-eight technique.

D
This elderly patient experienced massive spontaneous bleeding from a ruptured varicose vein. Blood loss was substantial and recurrent when she walked.

Figure 34.8 Hemorrhage control.
A few millimeters of difference in total stretch makes a large difference. In digits, tourniquet pressures of only 150 mm Hg are needed for hemostasis.

A surgical glove placed over a patient’s cleaned hand can also serve as a finger tourniquet (see Fig. 34.11C). Remove the tip of the glove covering the injured digit, and then roll it proximally along the patient’s finger to form a constricting band at the base. Another advantage of this technique is that contamination of the wound during closure is less likely. Rolled surgical gloves produce pressures ranging from 113 to 363 mm Hg, depending on the thickness, the amount of glove finger removed, the number of rolls, and the size of the glove in relation to the size of the patient’s hand. Ring-shaped exsanguinating digit tourniquets are available commercially (Tourni-Cot [Mar-Med Company, Grand Rapids, MI]) (see Fig. 34.11D). There is a real danger of forgetting to remove such a small tourniquet and accidentally incorporating it into the dressing.

These techniques provide bloodless fields in which to examine, clean, and close extremity wounds. Débridement of questionably devitalized tissue in a wound is best accomplished without a tourniquet or pharmacologic vasoconstriction because bleeding from tissues is often an indication of their viability.

**CLOSURE**

The various techniques for wound closure are presented in Chapter 35. The remainder of this chapter addresses issues related to wound management (e.g., secondary closure, wound dressings, use of antibiotics, aftercare instructions, and suture removal).

**Open Versus Closed Wound Management**

Wounds that heal spontaneously (i.e., by secondary intention) undergo much more inflammation, fibroplasia, and contraction than do those whose edges are reapproximated by wound closure techniques. During wound healing contraction covers the defect, but it may result in a deformity (contracture) or loss of function. Left to itself, the healing process may be unable to close a defect completely in areas in which the surrounding skin is immobile, such as on the scalp or in the pretibial area (Fig. 34.12). Exposed tendons, bone, nerves, or vessels may desiccate in an open wound. If the patient is careless with an otherwise adequate dressing that covers an open wound, the wound may be further contaminated. Surgical closure of wounds minimizes inflammation, fibroplasia, contracture, scar width, and contamination.

However, surgical closure of wounds can cause complications.Closure of contaminated wounds increases the probability of wound infection, with possible complications of impaired healing, dehiscence, and sepsis. For instance, raised pretibial flap lacerations in elderly patients often necrose when sutured, but survive and heal well by secondary intention if placed back into position with tape or tissue glue. Sutures in themselves are potentially detrimental to healing and can increase the risk for infection. Each suture inflicts a small intradermal incision that damages the surface epithelium, dermis, subcutaneous fat, blood vessels, small nerves, lymphatics, and epithelial appendages such as hair follicles, sweat glands, and ducts. Once divided and separated by a stitch, these appendages usually undergo inflammation and resorption. Each suture is another piece

---

**Figure 34.9** Scalp lacerations can bleed profusely. This patient’s pressure dressing fell off while he was on a stretcher in the back hall, and he experienced significant blood loss from a large scalp laceration. His intoxication prevented him from supplying his history of hemophilia.
Scalp lacerations may bleed profusely, and arterial vessels may not retract if they are in tough fibrous tissue. This forceps was used to crush a small pumping artery. Placing large (3-0 nylon) sutures that incorporate all scalp layers, defined as all tissue between the thumb and index finger (arrows), is preferred to attempting to cauterize or ligate small bleeders.

A hemostat placed on the scalp edge can similarly be used on a single bleeder.

Note that the suture will incorporate all layers of the scalp. Both the galea and the periosteum are identified along with exposed skull (x). The periosteum is not sutured.

Figure 34.10 Hemorrhage control of scalp lacerations. Note that the periosteum is often misidentified as the galea. The galea is adherent to the subcutaneous tissues, not covering the bone. (C, From Custalow CB: Color Atlas of Emergency Department Procedures, Philadelphia, 2005, Saunders, pp 136–137.)
of foreign material that can provoke inflammation. When a suture is removed, bacteria that have settled on the exposed portion of the suture are pulled into the suture track and deposited there.

Therefore the clinician must estimate the risk for infection prior to closure. If the wound is judged to be clean or is rendered clean by scrubbing, irrigation, and débridement, it may be closed. If the wound remains contaminated despite the best of efforts, it must be left open to heal by secondary intention. If the status of the wound is uncertain, delayed primary closure is an option that should be considered.

**Delayed Primary or Secondary Closure**

There is a common misconception that all wounds must be either sutured within a few hours or left open and relegated to slow healing and an unsightly scar. If there is a substantial risk that closure of a particular wound might result in infection, the decision to close or to leave the wound open can be postponed. After cleaning, wounds left unsutured appear to have higher resistance to infection than do closed wounds. The condition of the wound after 3 to 5 days will then determine the best strategy (Fig. 34.13).

Although cleaning and débridement should be accomplished as rapidly as possible, there is no urgency in closing a wound. Edlich and associates\(^8\) pointed out that “the rationale for delayed primary closure is that the healing open wound will gain resistance to infection and permit an uncomplicated closure.” Despite its effectiveness, delayed primary closure is a technique that remains largely unappreciated and probably underused by many clinicians (Fig. 34.14). It is a highly effective means of managing wounds that have a sufficient risk for
Open wound management is usually an outpatient procedure. The technique consists of careful cleaning and débridement, followed by packing of the wound with sterile, saline-moistened, fine-mesh gauze. The packed wound is covered with a thick, absorbent, sterile dressing. Depending on the specifics of the wound and the ability of the patient to perform his or her own wound care, the packing may be changed daily at home or in the ED, or may be left undisturbed for several days. Sterile, saline-soaked packing is standard, and there is no need to impregnate wounds with antiseptics. Prophylactic antibiotics are occasionally prescribed, but their use is neither mandatory nor of proven benefit. On the fourth or fifth postoperative day, the wound is reevaluated for closure. If no evidence of infection is present, the wound margins can be approximated (delayed primary closure), or the wound can be excised and then sutured (secondary closure) with minimal risk for infection.

Because the wound is closed before the proliferative phase of healing, there is no delay in final healing, and the results are indistinguishable from those of primary healing.

Certain wounds should almost always be managed open or by delayed closure (Fig. 34.15). Such wounds include those that are already infected and those heavily contaminated by soil, organic matter, or feces. Wounds associated with extensive tissue damage, such as high-velocity missile injuries, explosion injuries involving the hand, complex crush injuries, and most bite wounds are also included in this category. Deep or contaminated lacerations on the bottom of the foot, such as those occurring when the patient steps on an unknown object while wading in a stream or running through a field, or wounds that are deep punctures, are ideal candidates for delayed closure. Some are never sutured but left open for primary healing. Human bite wounds (extending past the dermis) should never be closed primarily, even if they appear clean, and are often opened or extended further for cleaning. Clinicians disagree about which animal bite wounds may be closed initially. Most would suture cosmetically deforming injuries, including facial bites and bite wounds that can be excised completely. Others would primarily suture dog bites not involving an extremity. In severe soft tissue injuries, delayed closure allows time for nonviable tissue to become demarcated from uninjured tissue. Débridement can then be accomplished with maximal preservation of tissue.

**PROTECTION**

**Dressings**

At the conclusion of wound repair, wipe away dried blood on the surface of the skin with moistened gauze to minimize subsequent itching, and cover the wound with a nonadherent dressing. Depending on the specifics of the wound and the type of repair, a dressing can consist of a simple dry gauze pad, a complex multilayer dressing, or an occlusive dressing. Some wounds, such as sutured lacerations, do not routinely require any dressing. Although they are infrequently used on acute, traumatic lacerations managed in the ED, occlusive dressings have been shown to accelerate healing in a variety of wounds. Various specialized (and expensive) synthetic dressings are available, including vapor-permeable adhesive films, hydrogels, hydrocolloids, alginates, synthetic foam dressings, silicone meshes, tissue adhesives, barrier films, and collagen-containing dressings.

**Function of Dressings**

Dressings serve various functions. They protect the wound from contamination and trauma, absorb excess exudate from the wound, immobilize the wound and surrounding area, exert downward pressure on the wound, and improve the patient’s comfort. Occlusive dressings on burns or abrasions maintain a moist environment and prevent painful exposure of the wound to air and dehydration of the wound’s surface. Sutured wounds are particularly susceptible to infection from surface contamination during the first 2 days after wound repair. Dressings protect the wound from contamination during this vulnerable period.

One of the primary functions of gauze dressing is to absorb the serosanguineous drainage that exudes from all wounds. Absorbent dressings also reduce the development of stitch
Delayed Primary Closure

Figure 34.14 Delayed primary closure. Not all wounds have to be repaired on the initial visit, but may be left open to slowly heal. A delayed primary closure in 3 to 4 days is often the best choice for dirty or old lacerations.

Abscesses to some extent. Surface sutures produce small indentations at their points of entrance; tiny blood clots and debris overlie these indentations and allow bacterial growth at the site. Small “stitch abscesses” can develop; they are initially undetectable but are nevertheless destructive to epithelium. Stitch abscesses rarely infect the entire wound but can slightly increase the width of the scar and produce noticeable, punctate suture marks.

The most common type of dressing is constructed in three layers: a nonadherent contact layer, an absorbent layer, and
Principles of Wound Management

make good temporary coverings for wounds requiring delayed closure several days after wounding. Petrolatum gauze (e.g., Adaptic [Acelity, San Antonio, TX], Xeroform [Medtronic, Minneapolis, MN], Betadine [Purdue Pharma, Stamford, CT], Kendall [Medtronic, Minneapolis, MN]) is commonly applied next to the surface of the wound to prevent the wound from sticking to the dry gauze in the absorbent layer and to protect the regenerating epithelium (Box 34.1). Always use nonadherent material to cover skin grafts. Some clinicians use fine-mesh gauze (41 to 47 warp threads/inch²) rather than petrolatum gauze on abrasions, especially on wounds that are heavily contaminated, because removal of this type of dressing débrides only the small tufts of granulation tissue that become fixed in the mesh pores and leaves a clean, even surface. Once a healthy, granulating surface is present and reepithelialization is proceeding, nonporous dressings can be used. Fine-mesh gauze is also used next to exposed tissue in wounds being considered for delayed primary closure. Though petroleum gauze is beneficial for nonadherence of subsequent dressings to wounds, studies have shown that the petroleum may decrease the tensile strength of the knots in a stitch.

Figure 34.15 A, Human bites are never closed primarily because of risk for infection. They are often opened and extended to facilitate cleaning and a search for foreign bodies (FBs) such as fragments of teeth. B, Dog bites in a cosmetic area may be closed primarily. Large cosmetic defects are best closed in the operating room by a surgeon with adequate time to correctly prepare the wound. C, This wound on the bottom of the foot, sustained by stepping on an unknown object while running in a stream, should not be closed but rather left open to heal primarily. Always check for foreign bodies in these wounds. Ask an assistant to hold the wound open while you perform pressure irrigation. This was initially a much smaller puncture wound that was extended to search for a suspected FB and also to clean and initially pack the wound open. Sutures were never required and the wound healed in 7 days.

Contact Layer: Dry, Semiocclusive, and Occlusive Dressings

Wounds covered with permeable dressings such as plain gauze tend to dry out. Although this is acceptable for dressing sutured wounds, drying of the wound surface damages a shallow layer of exposed dermis, which impedes epidermal resurfacing of abrasions, burns, and incisions. Wound desiccation results in further epidermal necrosis, crust formation, and increased inflammation. Coarse weaves of gauze, usually available in the form of multilayered pads, absorb blood and exudate, but the dressing will adhere if the interstices of the fabric are relatively large. Capillaries, fibrin, and granulation tissue will penetrate and become enmeshed in the material. If the proteinaceous exudate from the wound dries by evaporation, the scab usually clings to the dressing. Some clinicians use this effect to “débride” the wound when the gauze is removed. However, it may also destroy healing tissue, particularly the new epithelium. Even though débridement of the wound with wet-to-dry dressings is quick, careful débridement with surgical instruments is more controlled and less traumatic.

Adherence to the wound can be prevented if the dressing is nonabsorbent, occlusive, or finely woven. If the wound is kept moist by covering it with an occlusive film or nonadherent covering soon after wound management, and if the film is left in place for at least 48 hours, the epidermis will migrate over the surface of the dermis faster than when a dry scab is allowed to form. Protection of wounds that are healing by secondary intention with occlusive or semiocclusive dressings has several advantages including more rapid healing, less pain from exposure to air, better cosmetic result, fewer dressing changes, and protection from bacteria. Occlusive dressings make good temporary coverings for wounds requiring delayed closure several days after wounding.

Petrolatum gauze (e.g., Adaptic [Acelity, San Antonio, TX], Xeroform [Medtronic, Minneapolis, MN], Betadine [Purdue Pharma, Stamford, CT], Kendall [Medtronic, Minneapolis, MN]) is commonly applied next to the surface of the wound to prevent the wound from sticking to the dry gauze in the absorbent layer and to protect the regenerating epithelium (Box 34.1). Always use nonadherent material to cover skin grafts. Some clinicians use fine-mesh gauze (41 to 47 warp threads/inch²) rather than petrolatum gauze on abrasions, especially on wounds that are heavily contaminated, because removal of this type of dressing débrides only the small tufts of granulation tissue that become fixed in the mesh pores and leaves a clean, even surface. Once a healthy, granulating surface is present and reepithelialization is proceeding, nonporous dressings can be used. Fine-mesh gauze is also used next to exposed tissue in wounds being considered for delayed primary closure. Though petroleum gauze is beneficial for nonadherence of subsequent dressings to wounds, studies have shown that the petroleum may decrease the tensile strength of the knots in a stitch.

Box 34.1 Advantages of Occlusive Dressings

- More rapid healing
- Less pain from exposure to air
- Better cosmetic results
- Fewer dressing changes
- Better protection from bacteria

Wound Dressing

A common three-layer dressing consisting of antibiotic ointment, Adaptic, and gauze.

1. The inner layer is nonadherent gauze. The middle layer is 2- × 2-inch gauze sponges wrapped circumferentially and held in place with tape.
2. Begin with No. 2 tube gauze at the base of the finger. Hold this end with one finger while the tube gauze applicator is pulled toward the fingertip. A twisting motion firms the wrap about the digit; about 90° is necessary. Excessive stretch or twisting can compromise the circulation.
3. When the fingertip is reached, make a 360° twist, but avoid placing a constricting twist around the finger itself.
4. Pass the applicator toward the base of the finger with an additional 90° twist. Repeat once more; thus, three layers are in place.
5. Cut enough gauze to reach the base of the finger, and tape it there. As an alternative, pull the final layer beyond the tip while leaving it long enough to reach to and around the wrist (about three times the finger length). Split this gauze into two strands; bring them dorsally to the wrist, knot, and loosely wrap around the wrist.

For a distal finger dressing, covering the gauze with a finger cut from a clean glove provides protection from dirt and wetness.

Figure 34.16 Wound dressings.
There are over a thousand types of occlusive dressings available today.\textsuperscript{32} Occlusive dressings are ideal for covering abrasions and burns, but they are also used on lacerations and chronic wounds. Some experts report that they should be avoided if a wound is highly exudative, infected, or at high risk for infection.

There are four main types of occlusive dressings designed for the surfaces of wounds: films, hydrocolloids, foams, and hydrogels\textsuperscript{52} (Table 34.2). Films are transparent polyurethane-derived membranes that provide an occlusive effect; they include Omniderm (Omikron Scientific, Ltd., Alpharetta, GA), OpSite (Smith &Nephew, Ltd., Andover, MA), Tegaderm (3M, St. Paul, MN), and Bioclusive (Systagenix, Airebank Mills, Skipton, United Kingdom). They are best for wounds with minimal drainage and those that require frequent monitoring. Hydrocolloids are semiocclusive, waterproof dressings with soluble cellulose or gelatin backing, such as DuoDERM (ConvaTec) and Tegasorb (3M). They can be used in wounds with a moderate amount of exudate. Hydrogels are products with starch polymers and water, such as Curagel (Kendall) and Telfa (Covidien, Minneapolis, MN). Because they provide moisture to wounds they are suitable for dry or painful wounds. Foams are moderately absorbent polyurethane or silicone products that are semiocclusive and provide a cushion-effect.\textsuperscript{52,53} Silicon dressings such as Mepitel (Mölnlycke Health Care) can be used for the treatment of fragile wounds, like skin tears.\textsuperscript{59}

One fear of using occlusive dressings is that microorganisms will proliferate in the moist environment beneath the occlusive film and increase wound infection rates. However, occlusive dressings such as DuoDERM actually serve more as a barrier to external pathogenic bacteria.\textsuperscript{53,60} Although skin bacteria under occlusive dressings can multiply,\textsuperscript{61} even chronic wounds contaminated with large numbers of bacteria are routinely treated with occlusive dressings successfully.\textsuperscript{52}

Adhesive-backed dressings (e.g., DuoDERM and OpSite) may adhere to an open wound and remove new epidermis; however, the adhesive usually only adheres to dry intact skin, which is located around the wound.\textsuperscript{52} These film dressings have low absorptive capabilities and do not allow exudate to drain out the edges of the dressing. Exudate pooling overlying

### Table 34.2 Summary of dressings used for acute wound care

<table>
<thead>
<tr>
<th>AGENT AND TYPE</th>
<th>PROPERTIES</th>
<th>ADVANTAGES</th>
<th>DISADVANTAGES</th>
<th>CLINICAL APPLICATIONS</th>
<th>EXAMPLES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gauze</td>
<td>Cotton or synthetic; woven; Plain or petrolatum-impregnated gauze</td>
<td>Cost effective</td>
<td>Poor absorbance; dehydrates the wound; impedes wound re-epithelialization; adherence causes pain with dressing changes</td>
<td>Primary dressing; must be changed every 12 to 24 hours</td>
<td>Curity (Covidien) Xeroform (Medtronic, Minneapolis, MN)</td>
</tr>
<tr>
<td>Film</td>
<td>Transparent; Self-adhesive; water-proof, but gas and water vapor permeable; flexible</td>
<td>Transparency allows for frequent wound inspection; adhesive does not stick to wound and thus low pain with dressing changes</td>
<td>Not absorbent; trapped excess exudate can cause wound maceration</td>
<td>Primary or secondary dressing; best for wounds with low exudate; ideal for flexible areas such as hands and joints; can be changed every 7 days</td>
<td>Tegaderm (3M) Transeal (DeRoyal) Bioclusive (Systagenix) Omniderm (Omidron Scientific)</td>
</tr>
<tr>
<td>Foam</td>
<td>Polyurethane or silicone center and semi-occlusive outer layer; adherent or non-adherent</td>
<td>Moderate absorption; cushions wounds</td>
<td>As wound exudate decreases, can dehydrate the wound; non-adherent foam may require a film covering</td>
<td>Good for wounds with high exudate; Ideal over bony areas; can be changed from daily to weekly</td>
<td>Optifoam (Medline) Mepilex (Mölnlycke)</td>
</tr>
<tr>
<td>Hydrogel</td>
<td>Water-based gel; sheet or amorphous form</td>
<td>Donates moisture; reduces wound pain</td>
<td>Non-adherent and requires a secondary dressing</td>
<td>Good for dry wounds or with low exudate; changed every 1 to 3 days</td>
<td>Curagel (Covidien) Tegagel (3M)</td>
</tr>
<tr>
<td>Hydrocolloid</td>
<td>Adhesive semi-occlusive water-proof</td>
<td>Moderate absorption; conforms to wound</td>
<td>Discharge can be malodorous and appears purulence-like</td>
<td>Changed every week</td>
<td>DuoDERM (ConvaTec) NuDerm (Systagenix)</td>
</tr>
<tr>
<td>Silicon</td>
<td>Porous soft silicone; nonadherent</td>
<td>Reduces wound pain</td>
<td>Expensive</td>
<td>Good for fragile wounds, like skin tears; changed every week</td>
<td>Mepitel (Mölnlycke)</td>
</tr>
</tbody>
</table>
the wound will macerate skin. Between dressing changes, coat the wound with petrolatum or an antibiotic ointment before applying these products. Wounds covered with certain occlusive dressings or with silver sulfadiazine (Silvadene, Marion Laboratories, Kansas City, MO) appear to be blanketed with pus; this exudate actually represents the beneficial proliferation of macrophages and polymorphonuclear leukocytes.

Absorbent Layer
When dressing wounds with considerable drainage, use sufficient gauze to cover the wound and absorb all the drainage. Change the dressing whenever it becomes soiled, wet, or saturated with exudate. Once a dressing becomes moist, pathogens can pass through the mesh-like structure to the underlying wound. Consequently, a dressing that is used to absorb exudate or débride the wound must be changed more frequently than one designed solely to occlude. Absorbent dressings on draining wounds can be changed daily to avoid bacterial overgrowth beneath the dressing. Aspirate fluid accumulating under an occlusive dressing and change the dressing every 1 to 2 days during the first week, or until the exudate no longer accumulates.

Outer Layer
Bleeding may persist despite attempts to provide good hemostasis. Compressive dressings help prevent hematoma formation and eliminate dead space within a wound. They are particularly useful for wounds that have been undermined extensively, and for facial wounds in which subcutaneous capillary bleeding and swelling can exert tension on fine skin sutures and jeopardize skin closure. Pressure dressings should be used to immobilize skin grafts. Surgical tape can serve as a pressure dressing in areas on which bandages cannot be applied easily, such as fingertips.

Apply pressure dressings to all ear lacerations to prevent hematoma formation and subsequent deformation and destruction of cartilage. Envelop the ear in the dressing to distribute pressure from the outer bandage evenly across the irregular surface of the pinna. Pack moistened cotton into the concavities of the pinna until the cotton is level with the most lateral surface of the pinna. Cut square pieces of gauze to fit the curvature of the ear and place them behind (medial to) the pinna. Place several more gauze squares on the lateral surface of the ear. Secure the packing in place with a circumferential head bandage, but do not encompass the opposite ear because it would just as easily cause pressure necrosis of that ear if left unprotected.

Bandage traumatic wounds to compress, immobilize, secure, and protect the wound and underlying dressing. Most bandaging is performed on extremities, where dressings are difficult to secure with tape alone. Rolls of cotton (Kerlix [Covidien], Kling [Johnson & Johnson] stretch gauze) are well suited for this purpose. Wind the bandage around the extremity and advance it proximally with circular, overlapping turns. Take care to avoid making wrinkles in the bandage, which may create pressure points, and also be careful not to make loose turns, which shorten the effective life of the dressing. When joint surfaces are crossed, anchor the cotton distally with several turns, then unravel it obliquely across the joint several times in a figure-of-eight pattern, and anchor it again proximally with two complete turns. Repeat this process until the bandage is securely in place. Fasten the ends of the bandage to the skin with strips of adhesive tape.

Bandages over the forearm and the lower extremities are particularly prone to slippage because of the constant motion of these parts and the marked changes in diameter of the extremity over a short distance. To help prevent this problem, rotate the roll of bandage 180 degrees after each circular turn to produce a reverse spiral and reduce the bandage’s mobility (see Fig. 34.16B). A simple dressing for a single digit is to use tube gauze or cover it with a finger cut from a surgical glove (see Fig. 34.16C and D). When bandaging digits, be cautious not to create a band-like dressing that can slip down the finger and cause constriction and ischemia. This can be especially problematic in children. Always include the hand and wrist in the dressing to create an anchor point and reduce circumferential bunching.

Certain chemically treated wide-mesh weaves have the properties of cling and stretch, which hold it snugly in place but expand if edema develops. An elastic cotton roll (Kerlix) allows the bandage to conform to body contours, provides some mobility to bandaged joints, and permits the wound to swell without the circumferential bandage constricting the extremity. An inelastic Kling bandage better immobilizes the part. Use rigid immobilization with plaster splints or braces to protect wounds in mobile areas, such as around large joints. Scalp wounds do well when left uncovered; bulky occlusive dressings of the scalp are discouraged. Encourage the patient to shower daily to remove debris from a sutured scalp laceration. If a dressing is thought necessary, hold it in place with a bandage. Change the outer layer of the dressing when it becomes externally contaminated, it is saturated with exudate, or when inspection and wound cleaning are required.

Dressings vary in their absorbency, adhesiveness, occlusiveness, opacity, and insulating properties. Further research may identify types of dressings that are best suited for different phases of the healing wound. Currently a two- or three-layer dressing is used for most traumatic wounds. Base the choice of material for the contact layer on the characteristics of the individual wound (see Table 34.2).

Splinting and Elevation
Wounds and sutured lacerations may be immobilized to enhance healing and to provide patient comfort (Fig. 34.17). Immobilization of an injured extremity promotes healing by protecting the closure and by limiting the spread of contamination and infection along lymphatic channels. Wounds overlying joints are subjected to repeated stretching and movement, which delays healing, widens the scar, and potentially disrupts the sutures. Short-term splints are almost always beneficial for lacerations that overlie joints and are frequently necessary for the protection of wounds involving the fingers, hands, wrists, volar aspect of the forearms, extensor surface of the elbows, posterior aspect of the legs, plantar surface of the feet, and the extremities when skin grafts have been applied. A plaster or aluminum splint may be incorporated into a bandage to reduce the mobility of the injured part.

Elevate injured extremities in all but the most trivial injuries. Elevation limits edema formation, allows more rapid healing, and reduces throbbing pain. Patients given this information are often more motivated to elevate the extremity as instructed. Use slings to elevate wounds involving the forearm or the hand. The patient can also wrap a pillow around an injured hand to promote elevation at home (see Fig. 34.17A). Another technique to suggest is to lie down on a sofa with the injured
antibacterial capabilities against *Streptococcus* and *Staphylococcus* species in crush wounds, but topical nitrofurazone can increase the proliferation of granulation tissue. Although there is a risk for allergic sensitization or contact dermatitis with preparations containing neomycin, allergic reactions are uncommon (roughly 1%) unless the ointment is used repeatedly (Fig. 34.18).

One obvious benefit of using topical antibiotics is that ointments prevent adherence of the wound surface to the dressing. Use ointments to reduce the formation of a crust that covers and separates the edges of the wound. Lacerations surrounded by abraded skin are especially predisposed to coagulum formation. In such cases, instruct the patient to cleanse the wound frequently and to follow the cleansing with an application of ointment during the first few days.

Strong topical corticosteroids have detrimental effects on healing. Application of 0.1% triamcinolone acetonide in an ointment retards healing in wounds by as much as 60%, whereas hydrocortisone probably does not interfere with epithelialization. Some clinicians believe that single and low doses of oral corticosteroids probably have no effect on wound healing but that repeated, large doses of steroids (≤40 mg of prednisone per day) inhibit healing, particularly if used before the injury or during the first 3 days of the healing phase. There is some evidence that topical vitamin A may reverse some of the antiinflammatory and immunosuppressive effects of corticosteroids.

The exact value of ointments in the treatment of lacerations has yet to be determined. However, routine use of ointments after wound cleaning does encourage inspection of the wound by patients. Ointments without antibiotics that are used solely for moisture to the wound have a higher rate of infection and should not be used. Antibiotic ointments can be placed over abrasions or sutured lacerations. Do not use ointments on wounds closed with tissue adhesive because the ointment will dissolve the adhesive.

**Wound Cultures**

Tissue taken for culture at the time of wound preparation and closure in the ED serves no useful purpose and is not
recommended. The results of such cultures cannot logically guide future antibiotic selection. It is not necessary to routinely culture all minimally infected wounds after closure. Cultures should be considered if there is a subsequent infection that is extensive, unusual, or otherwise concerning, if the patient is immunocompromised, or if there is suspicion of methicillin-resistant *Staphylococcus aureus*. All wounds that are grossly infected at the time of follow-up should also be assessed for the presence of a foreign body.

It is not uncommon to encounter a minor infection or inflammation of the suture tracks at the time of suture removal, as evidenced by a small drop of pus when the suture is removed. Such minor infections, so-called stitch abscesses, usually do well with warm soaks or topical antibiotic ointments and do not require wound cultures or systemic antibiotics.

**Systemic Antibiotics**

Most traumatic soft tissue injuries sustain a low level of bacterial contamination. Uncomplicated wound infection rates in ED patients range from 2% to 5%, regardless of clinician intervention. In a number of clinical studies of relatively uncontaminated and uncomplicated traumatic nonbite wounds (which represent the majority of wounds managed in the ED), prophylactic antibiotics administered in various routes and regimens did not reduce the incidence of infection.72–74 Studies of antibiotic prophylaxis for animal bite wounds have produced variable results, and no large study providing stratification of the many prognostic factors has been conducted.75

As multiple studies on the use of prophylactic antibiotic regimens for wounds treated in the ED have failed to demonstrate a benefit, there is no clear practice standard.76 In most soft tissue wounds in which the level of bacterial contamination after cleaning and débridement is low, antibiotics are not recommended. Heavily contaminated wounds (such as wounds in contact with saliva, pus, or feces) often become infected despite antibiotic treatment. Nevertheless, antibiotics may have marginal benefit when the level of contamination is overwhelming, or if the amount of questionably viable tissue left in the wound is considerable (e.g., with crush wounds). Antibiotics may be considered for extremity bite wounds, puncture-type bite wounds in any location, intraoral lacerations that are sutured, orocutaneous lip wounds, wounds that cannot be cleaned or débrided satisfactorily, and highly contaminated wounds (e.g., those contaminated with soil, organic matter, purulence, feces, saliva, or vaginal secretions). They may also be considered for wounds involving tendons, bones, or joints; for wounds requiring extensive débridement in the operating room; for wounds in lymphedematous tissue; for distal extremity wounds when treatment is delayed for 12 to 24 hours; for patients with orthopedic prostheses; and for patients at risk for the development of infective endocarditis.72,77

Although some consider prescribing prophylactic antibiotics for immunocompromised patients (diabetics and others), a true benefit has not consistently been demonstrated in any subset. Many resistant strains of bacteria now encountered in clinical practice (patients and in the community) have been linked to the excessive use of needless antibiotics. Other disadvantages of routine antibiotic use include unnecessary expense and potential side effects (e.g., rash, anaphylaxis, diarrhea, vomiting). It seems most prudent to avoid routine antibiotic prophylaxis and opt for meticulous wound care, close follow-up, and the selective use of antibiotics for proven infection. If antibiotics are being considered in a specific case, give them as soon as possible after wounding and continue them for only 2 to 3 days in the absence of development of an infection. If the risk for infection is high enough to warrant antibiotics, delayed primary closure may also be considered.

Many patients have difficulty determining whether their wounds are infected and mistake the normal healing process for infection; therefore in high-risk patients (a clinical judgment), mandatory follow-up is usually necessary.78,79

The choice of antibiotic, particularly for bite wound prophylaxis, is as controversial as the indications for use.78 Many species of bacteria cause animal bite wound infections, thus making complete coverage impossible.77,80 Antibiotic regimens vary with the species of the biter and with evolving bacterial resistance. The duration of antibiotic prophylaxis is also in question. It is common practice to provide antibiotics for 72 hours. (See additional comments on animal bites at the end of this chapter.)

**Tetanus Immunoprophylaxis**

Although tetanus is rare, it still occurs in the United States and is a preventable disease. Therefore any wound should be assessed for its potential to cause tetanus, and prophylaxis should be considered in the ED. Approximately 70% of Americans older than 6 years have protective levels of tetanus antibodies, but levels decline as age increases, with elderly women having the lowest levels of protection. New immigrants are most likely to have inadequate immunity. Hence, efforts at preventing tetanus should especially be addressed in immigrants and the elderly. The recommendations of the Centers for Disease Control and Prevention for tetanus prophylaxis are listed in Fig. 34.19 and Table 34.3.82,83

When questioning patients about their tetanus immunization status, determine if they ever completed the primary immunization series, and if not, how many doses were given. Patients who have not completed a full primary series of injections (three or four doses) may require both tetanus toxoid and passive immunization with tetanus immune globulin. Tetanus immune globulin will decrease, but not totally eliminate, the subsequent development of clinical tetanus. Tetanus and diphtheria immunizations are often given together (as DT or DTaP for pediatrics, and Td or Tdap for adults). The preferred active tetanus immunization in patients 7 years of age and older is Tdap or Td. For passive immunity, prepare 500 units of tetanus immune globulin. Give most of this dose intramuscularly, and infiltrate the remainder around the wound.83

**TABLE 34.3 Summary of Tetanus Vaccine Recommendations**

<table>
<thead>
<tr>
<th>VACCINE HISTORY</th>
<th>Clean, Minor Wounds</th>
<th>All Other Wounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unknown or fewer than 3 doses</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>3 or more doses</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

Mild local reactions consisting of erythema and induration are common (≈20%) after tetanus toxoid injections; occasionally, they are accompanied by fever and mild systemic symptoms. Reactions are approximately twice as common if diphtheria immunization is coupled with tetanus immunization. This is a hypersensitivity reaction, not an infection, and does not represent an absolute contraindication to further immunizations. A minor febrile illness, such as an upper respiratory infection, is not a reason to delay immunization. Although serious reactions are rare, a hypersensitivity reaction consisting of tenderness, erythema, and swelling or serum sickness develops in some patients with high antibody levels. Generalized urticarial reactions and anaphylaxis are the severe reactions associated with the tetanus vaccine. Although peripheral neuropathy and Guillain-Barré syndrome have also been reported, there is inadequate evidence to accept or reject a causal relationship. The only absolute contraindication to tetanus toxoid is a history of anaphylaxis or a neurologic event. In such cases, tetanus immune globulin can be given safely. Pregnancy is not a contraindication to either toxoid or immune globulin, although some suggest that the toxoid be used “with caution” during the first trimester. Given the excellent amnestic response to the toxoid, it is likely that the primary immunization series, coupled with intermittent boosters every 5 to 10 years, depending on the cleanliness of the wound, conveys immunity for most of one’s life. However, a significant percentage of elderly patients fail to develop protective antitoxin antibody titers after 14 days when given tetanus toxoid boosters.

Treatment decisions are based on the differentiation between clean and contaminated wounds. Even though any break in the skin can harbor *Clostridium tetani*, traditional definitions of tetanus-prone wounds include the following: injuries more than 6 hours old; wounds contaminated by feces, saliva, purulent exudate, or soil; wounds with retained foreign bodies or containing devitalized or avascular tissue; established wound infections; penetrating abdominal wounds involving bowel; deep puncture wounds; and wounds caused by crush, burns, or frostbite.

Tetanus can develop despite prior immunization, and it can result from chronic skin lesions in apparently minor or clean wounds. In 10% to 20% of cases, no previous wound can be identified. Patients’ recall of past immunizations is imperfect, and immunity may be, on rare occasions, inadequate after a complete series of tetanus toxoid.³¹ Tetanus boosters given more frequently than advised increase the incidence of adverse reactions to subsequent injections. However, the benefits of overtreatment seem to outweigh the risks.

**PATIENT INSTRUCTIONS**

Successful wound healing is partly dependent on the care given to the wound once the patient leaves the ED. Patient satisfaction depends not only on the cosmetic result but also on the expectation of that result.³⁹ Both are reasons why patients should receive thorough and clear instructions.

Inform the patient that no matter how skillful the repair, any wound of significance produces a scar. Most scars deepen in color and become more prominent before they mature and fade. The final appearance of the scar cannot be judged until 6 to 12 months after the repair.³⁹ Some wounds heal with wide, unattractive scars despite ideal management and closure. Wounds more likely to have significant scars are those that cross perpendicular to joints, wrinkle lines or lines of minimum tension (Kraissl lines), and those that retract more than 5 mm. Wounds that are likely to scar are those that are located over convexities or in certain anatomic locations (e.g., anterior upper part of the chest, back, shoulders) where hypertrophic scars are common. A wound crossing a concave surface may result in a bowstring deformity; one crossing a convexity may leave a scar depression. To avoid these complications, a Z-plasty
Some areas such as the face or the scalp, but vigorous scrubbing dried blood and exudate is probably beneficial, especially in from the wound. If possible, keep the injured part elevated.

If an injured extremity or finger is protected by a splint, it should be left undisturbed until the sutures are removed. Patients with intraoral lacerations can be instructed to use warm salt water mouth rinses at least three times a day. Patients with complicated or infection-prone wounds should be examined in 2 to 3 days by a clinician or nurse. Inform patients that a painful wound is often a sign of infection or suture reaction, and pain should prompt inspection of the wound. If no sign of infection is present after 48 to 72 hours, the patient can care for the wound until it is time for removal of the sutures.

Because the edges of a wound are sealed by coagulum and bridged by epithelial cells within 48 hours, the wound is essentially impermeable to bacteria after 2 days. Generally, a wound should be protected with a dressing itself. Generally, a wound should be protected with a dressing for removal of the sutures at an appropriate time. Examine the patient to protect the wound and keep the dressing clean and dry for 24 to 48 hours. In this initial period change the dressing only if it becomes externally soiled or soaked by exudate from the wound. If possible, keep the injured part elevated.

Daily gentle washing with mild soap and water to remove dried blood and exudate is probably beneficial, especially in areas such as the face or the scalp, but vigorous scrubbing of wounds should be discouraged. Patients may bathe with sutures in place but should not immerse the wound for a prolonged time. Although diluted hydrogen peroxide can be used to remove blood from the skin surface, it should not be repeatedly used as a cleaning agent on the healing wound itself. Generally, a wound should be protected with a dressing during the first week and the dressing changed daily. Sutured scalp lacerations are usually left open, and showering is encouraged.

If an injured extremity or finger is protected by a splint, it should be left undisturbed until the sutures are removed. Patients with intraoral lacerations can be instructed to use warm salt water mouth rinses at least three times a day.

Patients may ask about the efficacy of various creams and lotions (e.g., vitamin E, aloe vera, cocoa butter) in limiting scar formation, and epithelialization.

Wounds in which extensive dissection of subcutaneous tissue has been performed may develop an intense inflammation similar in appearance to low-grade, localized cellulitis. It is rarely necessary to open these wounds. Remove one or two stitches to relieve some of the tension caused by mild swelling, if necessary. Cleanse daily with water and a mild soap and apply warm compresses, and this type of wound reaction should subside within 24 to 48 hours.

If a wound becomes infected, evaluate for the presence of a retained foreign body as the nidus of the infection (Fig. 34.20). In most sutured wounds that become infected, remove the sutures to allow drainage. If a wound exhibits a minor infection, remove a few sutures or all of them. Pack grossly infected wounds open to allow further drainage. Treat infected wounds with daily cleansing, warm compresses, and antibiotics. Leave wounds that have been opened to heal by secondary intention, which involves wound contraction, granulation tissue formation, and epithelialization.

Most wound infections can be treated in the outpatient setting with oral antibiotics and follow-up. Lymphangitis does not mandate intravenous antibiotics or hospitalization.

**SECONDARY WOUND CARE**

**Reexamination**

Patients with simple closed wounds may be released with appropriate instructions for home care and be told to return for removal of the sutures at an appropriate time. Examine high-risk wounds, such as bite wounds and other infection-prone wounds, in 2 to 3 days for signs of infection. Inspect the wound if the patient experiences increasing discomfort, develops a fever, or believes that the wound is infected. Evaluate wounds being considered for delayed primary closure in 3 to 5 days.

Wounds in which extensive dissection of subcutaneous tissue has been performed may develop an intense inflammation similar in appearance to low-grade, localized cellulitis. It is rarely necessary to open these wounds. Remove one or two stitches to relieve some of the tension caused by mild swelling, if necessary. Cleanse daily with water and a mild soap and apply warm compresses, and this type of wound reaction should subside within 24 to 48 hours.

If a wound becomes infected, evaluate for the presence of a retained foreign body as the nidus of the infection (Fig. 34.20). In most sutured wounds that become infected, remove the sutures to allow drainage. If a wound exhibits a minor infection, remove a few sutures or all of them. Pack grossly infected wounds open to allow further drainage. Treat infected wounds with daily cleansing, warm compresses, and antibiotics. Leave wounds that have been opened to heal by secondary intention, which involves wound contraction, granulation tissue formation, and epithelialization.

Most wound infections can be treated in the outpatient setting with oral antibiotics and follow-up. Lymphangitis does not mandate intravenous antibiotics or hospitalization.

**Suture Removal**

The optimal time for suture removal varies with the location of the wound, the rate of wound healing, and the amount of tension on the wound. Certain areas of the body such as the back of the hand heal slowly, whereas facial or scalp wounds heal rapidly. The speed of wound healing is affected by systemic factors such as malnutrition, neoplasia, and immunosuppression. Therefore only general guidelines can be given for the timing of suture removal. At the time that suture removal is being considered, one or two sutures may be cut to determine whether
the edges of the skin are sufficiently adherent to allow removal of all the sutures. Removing sutures too early invites wound dehiscence and widening of the scar, whereas leaving sutures in longer than necessary may result in epithelial tracks, infection, and unsightly scarring.85

Percutaneous sutures stimulate an inflammatory reaction along the suture track. Infection around a suture can lead to the formation of a stitch mark.86 Other factors that determine the severity of stitch marks include the length of time that stitches are left in place, skin tension, the relationship of the suture to the edge of the wound, the region of the body, and the patient’s tendency for keloid formation.86,87 The skin of the eyelids, palms, and soles and the mucous membranes seldom show stitch marks. In contrast, oily skin, and the skin of the back, the sternal area, the upper part of the arms, the lower extremities, the dorsum of the nose, and the forehead are likely to exhibit the permanent imprints of suture material on the skin surface.86

If sutures are removed within 7 days, generally no discernible needle puncture or stitch marks will persist.87 However, at 6 days the wound is held together by a small amount of fibrin and cells, and has minimal strength.89 The tensile strength of most wounds at this time is adequate to hold the wound edges together, but only if there is no appreciable dynamic or static skin force pulling the wound apart.88 Minimal trauma to an unsupported wound at this point could cause dehiscence. The clinician should decide on the proper time to remove the sutures after weighing these various factors. If early suture removal is necessary (such as on the face), wound repair can be maintained with strips of surgical skin tape. The key to wound tensile strength after suture removal is an adequate deep tissue layered closure.

Some general guidelines exist for suture removal. Remove sutures on the face on the fifth day after the injury, or remove alternate sutures on the third day and the remainder on the fifth day. On the extremities and the anterior aspect of the trunk, leave sutures in place for approximately 7 days to prevent disruption of the wound. Leave sutures on the scalp, back, feet, hands, and joints in place for 10 to 14 days, even though permanent stitch marks may result.89 Some clinicians recommend removal of the sutures used to repair eyelid lacerations as early as 72 hours to avoid epithelialization along the suture track with subsequent cyst formation.89

Removing sutures is usually relatively simple. Cleanse the wound and any remaining crust overlying the surface of the wound or surrounding the sutures. Wipe the skin with an alcohol swab. Cut each stitch with scissors or the tip of a No. 11 scalpel blade at a point close to the surface of the skin on one side. Grasp the suture knot on the opposite side with forceps and pull it across the wound (Fig. 34.21). The amount of exposed suture dragged through the suture track is thereby minimized. It is difficult to remove sutures with very short ends. At the time of suture placement, cut the length of the suture ends so they generally equal the distance between sutures. This makes it easier to grasp the suture during subsequent removal while avoiding entanglement during the knotting of adjacent sutures.

Once the skin sutures are removed, the width of the scar increases gradually over the next 3 to 5 weeks unless it is supported. Support is provided by previously placed subcutaneous stitches that bring the edges of the skin into apposition or by the application of skin tape at the time of removal. A nonabsorbable, subcuticular suture can be left in place for 2 to 3 weeks to provide continued support for the wound. Although complications such as closed epithelial sinuses, cysts, or internal tracts can occur from prolonged use of this stitch, they are unusual and can be avoided by placement of a buried subcuticular stitch with an absorbable suture.

Small stitch abscesses may occur in wounds when sutures remain in place for more than 7 to 10 days. Localized stitch abscesses generally resolve after removal of the sutures and application of warm compresses, and without antibiotics in simple cases.

If time and effort have been invested in cosmetic closure of the face, protect the repair with skin tape after the skin sutures have been removed. Wound contraction and scar widening continue for 42 days after the injury.89 Because the desired result is a scar of minimal width, the tape can be used for as long as 5 weeks after removal of the sutures. With exposure to sunlight, scars in their first 4 months redden to a greater extent than the surrounding skin does. In exposed cosmetic areas and when prolonged exposure to the sun is anticipated, appropriate sun protection and avoidance strategies should be used (e.g., a hat and sunscreen). Sunscreen may have a role in protecting scars from the sun, but more studies are needed to better understand its impact.
COMPLICATIONS

There are several reasons why wounds fail to heal; some are related to decisions made at the time of wound closure, and others are consequences of later events. Some of the impediments to healing include ischemia or necrosis of tissue, hematoma formation, prolonged inflammation caused by foreign material, excessive tension on the edges of the skin, and immunocompromising systemic factors. A primary cause of delayed healing is wound infection. Wound cleaning and débridement, atraumatic and aseptic handling of tissue, and the use of protective dressings minimize this complication. Inversion of the edges of a wound during closure produces a more noticeable scar, whereas skillful technique can convert a jagged, contaminated wound into a fine, apparent scar. The patient’s actions also affect wound healing. Delay in seeking treatment of an injury may significantly affect the ultimate outcome of the wound. Furthermore, in the first few days after an injury, the patient must take responsibility for protecting the wound from contamination, further trauma, and swelling.

Infection is probably the most common cause of dehiscence. If the patient is careless or unlucky, reinjury can reopen a wound despite the protection of a thick dressing. If the size of the suture is too small, the stitch may break. A stitch that is too fine or tied too tightly may cut through friable tissue and pull out. Knots that have not been tied carefully may unravel. The suture material may be extruded or absorbed too rapidly. Finally, if a stitch is removed too early (i.e., before tissues regain adequate tensile strength), the wound loses necessary support and can open. If the wound edges show signs of separating at the time of suture removal, alternate stitches can be left in place and the entire length of the wound supported by strips of adhesive tape. The final appearance of a scar is determined by several factors. Infection, tissue necrosis, and keloid formation widen a scar. Wounds located in sebaceous skin or oriented 90 degrees to dynamic or static skin tension lines result in wide scars. Dynamic tension is primarily a result of nearby joint movement. Wounds located in areas of high static skin tension will gape initially and often heal with wide scars despite adequate closure, whereas wounds in areas of loose or lax skin often heal with fine, narrow scars.

Miscellaneous Aspects of Wound Care

Traumatic wounds are created by a wide variety of mechanisms, and clinicians must sometimes adjust wound management techniques to match special circumstances.

The ED Approach to Puncture Wounds

Puncture wounds are common, yet there are no prospective studies to identify the most effective way to manage puncture wounds in the ED. Aside from evaluating tetanus immunization status and considering the possibility of a foreign body, the clinician has few proven options to prevent infection in a puncture wound. Scrubbing the surface of the puncture, evaluating the opening for retained foreign matter, and trimming jagged skin and tissue edges may be helpful. The value or appropriateness of coring, probing, or irrigating the puncture track has not been established. Deep irrigation is usually ineffective unless the puncture site is converted into a larger laceration to make the full depth of the puncture accessible, but routine enlargement of a puncture track is not recommended. For through-and-through punctures, the track can often be débrided by pulling gauze through the wound (Fig. 34.22).

It is impossible to accurately predict the final outcome of a puncture wound, though it may be determined at the time of the injury. Gross contamination or deep foreign bodies may not be appreciated for days. No prospective randomized trials have evaluated the role of prophylactic antibiotic administration to prevent infection in puncture wounds. Hence there are no standards on the use, type, or duration of prophylactic antibiotic therapy, even in high-risk patients. Most clinicians forego routine antibiotics and opt for simple cleaning and appropriate follow-up. Puncture wounds of the bottom of the foot may be an exception and are discussed in more detail in Chapter 51.

Gunshot Wounds

A subset of gunshot wounds may be definitively handled in the ED with outpatient follow-up. Studies by Ordog and colleagues documented a very low infection rate in gunshot wounds treated with standard wound care on an outpatient basis, even when the missile was left in place and minor fractures were present. Because most gunshot wounds are puncture wounds, only minimal deep wound cleaning is possible. Superficial soft tissue wounds with entrance and exit wounds in proximity may be débrided by passing sterile gauze back and forth through the wound track (Fig. 34.23). Though prescribed frequently, no data support the routine use of antibiotics following gunshot wounds.

Animal Bites

Many aspects of the treatment of animal bites are controversial, and no universal standards exist. Most bites are caused by dogs or cats that are family pets. Numerous organisms can be cultured from an infected bite wound caused by a dog or cat, and cultures may guide antibiotic therapy in infected wounds. However, cultures that are performed during the initial treatment of an animal bite are not predictive of subsequent infection or infecting organisms. The predominant pathogens in animal bites are the oral flora of the biting animal and human skin flora. Approximately 85% of bites harbor potential pathogens, and the average wound yields five types of bacterial isolates; almost 60% have mixed aerobic and anaerobic bacteria. Skin flora such as staphylococci and streptococci are isolated in approximately 40% of bites. The gram-negative rod Pasteurella canis (multocida), S. aureus, and Streptococcus viridans are common culprits in bite wound infections. Pasteurella species are isolated from 50% of dog bite wounds and 75% of cat bite wounds. Cat bites are usually puncture wounds that cannot be completely cleaned because of the sharp, pointed shape of their teeth. When compared with dog bites, cat bites may become infected rather quickly after the bite (within 24 hours), thus suggesting Pasteurella infection (Fig. 34.24). Cat bite wounds tend to penetrate deeply, with a higher risk for osteomyelitis, tenosynovitis, and septic arthritis than with dog bites, which are associated with crush injury and wound trauma. Capnocytophaga animorsus, a fastidious gram-negative rod, can cause bacteremia and fatal sepsis after animal bites, especially in asplenic patients or those with underlying hepatic disease. Anaerobes isolated from dog and cat bite wounds include Bacteroides, fusobacteria, Porphyromonas, Prevotella, propionibacteria, and peptostreptococci. Puncture wounds from a dog can be problematic because
amoxicillin-clavulanate (875/125 mg twice daily) given for 3 to 5 days may reduce infection rates after cat or dog bites, especially for a puncture wound, when the patient is seen more than 8 hours after the bite, or when wound cleaning has been inadequate. Some clinicians administer prophylactic antibiotics for all cat bites, for a dog bite that has been sutured, for hand wounds, for wounds close to a bone or joint, or for bites associated with deep tissue or crush injury. Although this is a reasonable and common approach, there are no data strongly supporting this practice. Alternative antibiotic combinations include clindamycin (300 mg four times daily) plus doxycycline.

Infected animal bites should be treated with antibiotics, but the use of prophylactic antibiotics for animal bites is controversial. Depending on location, dog bites may have an infection rate of 3% to 5%, whereas cat bites may become infected 50% of the time. Antibiotics for either prophylaxis or infection should have in vitro activity against *P. canis.*

they are difficult to clean unless they are through-and-through wounds (Fig. 34.25). Some clinicians have advocated primary closure of large dog bite lacerations that are centrally located on the body; however, markedly contused lacerations are good candidates for delayed primary closure (see Fig. 34.15B).

Infected animal bites should be treated with antibiotics, but the use of prophylactic antibiotics for animal bites is controversial. Depending on location, dog bites may have an infection rate of 3% to 5%, whereas cat bites may become infected 50% of the time. Antibiotics for either prophylaxis or infection should have in vitro activity against *P. canis.* Prophylactic

Figure 34.22 Puncture wounds cannot be cleaned meticulously and they may contain debris. The ultimate course of a puncture wound is likely set at the time of injury and they often do well. If an infection is seen at follow-up, a foreign body (FB) is highly likely. A–C, This nail gun puncture of the distal finger joint was able to be debrided by D, passing a small hemostat through the puncture wound and pulling gauze through the wound. E, A small piece of metal, used to attach the nails together (arrows), was removed with this maneuver. F, Note that nails from a nail gun clip are held together with a piece of metal or glue that may hamper removal or be left in the wound. A splint and 3 days of cephalexin prophylaxis (because of the discovered foreign body) were provided, and this wound healed well.
The best way to approach bite wounds is simply to adhere to the basic principles of wound care. Search for underlying fractures or tooth fragments with deep animal bites. When a wound results from the bite or scratch of either a wild or a domestic animal, give rabies prophylaxis if indicated (Tables 34.4 and 34.5).94

Human Bites

Human bite wounds are problematic for several reasons: these wounds are contaminated with human saliva; they may hide foreign material; deep structural damage may complicate the injury; and wound care may be delayed. Pathogens include aerobic bacteria (such as streptococci and S. aureus) and anaerobic bacteria (such as Eikenella, Fusobacterium, Peptostreptococcus, Prevotella, and Porphyromonas species). When cultured, most infected human bites harbor three to four pathogens, including both aerobes and anaerobes. Limited case reports suggest that viral pathogens including hepatitis, HIV, and herpes simplex virus, may be transmissible by human bites; however, data are lacking to guide the clinician on the best approach.

(100 mg twice daily), or trimethoprim-sulfamethoxazole (1 double-strength tablet twice daily). Antibiotics that do not have adequate activity against P. canis include first-generation cephalosporins (such as cephalaxin), penicillinase-resistant penicillins (such as dicloxacillin), macrolides (such as erythromycin), and clindamycin alone.
benign wound conditions. Pack them with gauze until follow-up. Do not use deep sutures. On reevaluation within a few days, perform delayed primary closure if desired. Facial bite wounds are an exception; primary closure may be considered for these wounds. Consider splinting the hand in a position of function with a short-arm volar splint for a few days to minimize joint movement. Advise the patient to elevate the injured area.

Most clinicians administer prophylactic antibiotics for 3 to 5 days for human bites, although there are no studies that

in the ED for these potential infections. Generally, prophylaxis is not suggested if the wound does not penetrate the dermis, is not over a joint, and does not involve the hands, feet, or cartilaginous structures.91

Clenched fist injuries caused by contact with another person’s teeth during a fight, or “fight bites,” are among the most serious human bite wounds. Lacerations typically occur over the third and fourth metacarpophalangeal or proximal interphalangeal joints of the dominant hand. Even small lacerations or punctures are highly prone to infection. Joint penetration is relatively common. Relaxation of the fist may disseminate organisms into the deep compartments and the deep tendon spaces of the hand, predisposing the patient to deep soft tissue infection, septic arthritis, and osteomyelitis. Many patients ignore these wounds until the onset of pain, swelling, or purulent discharge; as a result, these injuries are often complicated by established infection at the initial ED visit. In addition, some patients are not always forthcoming about the origin of the wound, especially if interpersonal violence is involved, thereby leading to incorrect ED treatment of a seemingly minor injury (Fig. 34.26).

Irrigate bite wounds copiously with tap water or sterile saline, and remove grossly visible debris. Many clinicians will extend a small laceration to allow visualization of the underlying structures and better cleaning. Look for tooth chips, tendon injury, and joint penetration. Wounds involving tendons or joint spaces are more serious and require close follow-up.

In general, leave all human bite wounds open (unsutured), even when treated soon after the injury and with seemingly

Figure 34.25 Dog bite puncture wounds are difficult to clean if not through and through. See text for discussion of the appropriate use of antibiotics.

Figure 34.26 Human bite. This patient was seen on a Monday morning with a small puncture wound on the dorsal surface of his hand (arrow), and he claimed that it was an injury sustained at work. Hence, minimal wound care was given. In reality, it was a human bite from a bar fight the night before. In 3 days, a serious infection was obvious and hospitalization was required. Note that the wrist is flexed, a patient held position that is common and increases hand edema.
All bites and wounds should be immediately cleansed thoroughly with soap and water. If antirabies treatment is indicated, both rabies immune globulin (RIG) and human diploid cell rabies vaccine (HDCV) should be given as soon as possible, regardless of the interval after exposure. Local reactions to vaccines are common and do not contraindicate continuing treatment. Discontinue vaccine if fluorescent antibody tests of the animal are negative.

During the usual holding period of 10 days, begin treatment with RIG and HDCV at the first sign of rabies in a dog or cat that has bitten someone. The symptomatic animal should be killed immediately and tested.

If RIG is not available, use antirabies serum, equine. Do not use more than the recommended dosage.

The animal should be killed and tested as soon as possible. Holding for observation is not recommended. Vaccination may be discontinued if immunofluorescence tests of the animal are negative.


### TABLE 34.4 Rabies Postexposure Prophylaxis Guide

The following recommendations are only a guide. In applying them, take into account the animal species involved, the circumstances of the bite or other exposure, the vaccination status of the animal, and the presence of rabies in the region. Local or state public health officials should be consulted if questions arise about the need for rabies prophylaxis.

<table>
<thead>
<tr>
<th>ANIMAL SPECIES</th>
<th>CONDITION OF ANIMAL AT TIME OF ATTACK</th>
<th>TREATMENT OF EXPOSED PERSONa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Domestic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dog and cat</td>
<td>Healthy and available for 10 days of observation</td>
<td>None unless rabies develops in the animal&lt;sup&gt;b&lt;/sup&gt; Local wound healing should be included in the treatment of the exposed person for each category RIG&lt;sup&gt;c&lt;/sup&gt; and HDCV.</td>
</tr>
<tr>
<td></td>
<td>Rabid or suspected rabid</td>
<td>RIG&lt;sup&gt;c&lt;/sup&gt; and HDCV</td>
</tr>
<tr>
<td></td>
<td>Unknown (escaped)</td>
<td>RIG&lt;sup&gt;c&lt;/sup&gt; and HDCV</td>
</tr>
<tr>
<td>Wild</td>
<td>Regard as rabid unless proved negative by laboratory tests&lt;sup&gt;d&lt;/sup&gt;</td>
<td>RIG&lt;sup&gt;c&lt;/sup&gt; and HDCV</td>
</tr>
<tr>
<td>Livestock, rodents, and lagomorphs (rabbits and hares)</td>
<td>Consider individually. Local and state public health officials should be consulted on questions about the need for rabies prophylaxis. Bites of squirrels, hamsters, guinea pigs, gerbils, chipmunks, rats, mice, other rodents, rabbits, and hares almost never call for antirabies prophylaxis.</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup>All bites and wounds should be immediately cleansed thoroughly with soap and water. If antirabies treatment is indicated, both rabies immune globulin (RIG) and human diploid cell rabies vaccine (HDCV) should be given as soon as possible, regardless of the interval after exposure. Local reactions to vaccines are common and do not contraindicate continuing treatment. Discontinue vaccine if fluorescent antibody tests of the animal are negative.

<sup>b</sup>During the usual holding period of 10 days, begin treatment with RIG and HDCV at the first sign of rabies in a dog or cat that has bitten someone. The symptomatic animal should be killed immediately and tested.

<sup>c</sup>If RIG is not available, use antirabies serum, equine. Do not use more than the recommended dosage.

<sup>d</sup>The animal should be killed and tested as soon as possible. Holding for observation is not recommended. Vaccination may be discontinued if immunofluorescence tests of the animal are negative.


### TABLE 34.5 Rabies Postexposure Prophylaxis Schedule, United States

<table>
<thead>
<tr>
<th>VACCINATION STATUS</th>
<th>TREATMENT</th>
<th>REGIMEN&lt;sup&gt;e&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Not previously vaccinated</td>
<td>Local wound cleansing RIG Vaccine</td>
<td>All postexposure treatment should begin with immediate thorough cleansing of all wounds with soap and water. 20 IU/kg of body weight; if anatomically feasible, up to half the dose should be infiltrated around the wounds and the rest administered IM in the gluteal area. Note: RIG should not be administered in the same syringe or into the same anatomic site as the vaccine because RIG may partially suppress active production of antibody. No more than the recommended dose should be given. HDCV or RVA, 1 mL IM (deltoid area&lt;sup&gt;e&lt;/sup&gt;), one each on days 0, 3, 7, 14, and 28.</td>
</tr>
<tr>
<td>Previously vaccinated&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Local wound cleansing RIG Vaccine</td>
<td>All postexposure treatment should begin with immediate thorough cleansing of all wounds with soap and water. RIG should not be administered. HDCV or RVA, 1 mL IM (deltoid area&lt;sup&gt;e&lt;/sup&gt;), one each on days 0 and 3</td>
</tr>
</tbody>
</table>

<sup>e</sup>These regimens are applicable for all age groups, including children.

<sup>c</sup>The deltoid area is the only acceptable site of vaccination for adults and older children. For younger children, the outer aspect of the thigh may be used. The vaccine should never be administered in the gluteal area.

<sup>c</sup>Any person with a history of preexposure vaccination with HDCV or RVA, previous postexposure prophylaxis with HDCV or RVA, or prior vaccination with any other type of rabies vaccine and a documented history of antibody response to the previous vaccination.

HDCV, Human diploid cell rabies vaccine; IM, intramuscularly; IU; international units; RIG, rabies immune globulin; RVA, rabies vaccine, adsorbed. Four formulations of three inactivated rabies vaccines are currently licensed for preexposure and postexposure prophylaxis in the United States.

Digital Nerves

Numbness in the area of digital innervation, concomitant injury to a digital artery (flash, pulsating bleeding), or an electric shock sensation when exploring a laceration should alert the clinician to a possible digital nerve injury. If there is uncertainty about nerve injury, the diagnosis can be established at the time of a follow-up visit. However, lacerations of digital arteries that impair the distal circulation must be identified early during the initial evaluation.

Débridement of hand and finger lacerations should be minimal, and wound cleaning should be gentle yet thorough. Digital nerves that are transected distal to the metacarpophalangeal joint may be candidates for surgical repair. It is unclear at which point along the course of a digital nerve a transection can be repaired successfully, so referral to a hand specialist is essential. Frequently injuries proximal to the distal interphalangeal joint are not repaired, but many other factors will influence operative decisions. Repair of a digital nerve will often result in return of good sensory function, but it takes months, and repair can prevent painful neuromas from developing. Most hand surgeons will not repair digital nerves at the initial visit. Instead, they advise wound cleaning, skin closure, splinting, and outpatient follow-up in 24 to 36 hours, followed by delayed nerve repair.

Accidental Soft Tissue Injection With an EpiPen

An EpiPen provides self-injected subcutaneous epinephrine (0.3 or 0.15 mg) for emergency treatment of anaphylaxis (Fig. 34.28).

Occasionally the device is inadvertently discharged, usually into a finger, producing intense distal vasospasm (Fig. 34.29). The patient has a blanched finger and minor sensory disturbances. The natural history of untreated vasospasm is unknown; although ischemia may last for an hour or two, it may be self-limited. Digit amputation from this event has not been reported. Nonetheless, clinicians faced with an obviously ischemic finger may consider an intervention. Local heat and nitroglycerin ointment are of no proven benefit. Placing a...
Both a digital block and direct injection of the reversal agent into the site of epinephrine injection have been described. Injecting the actual site of penetration is intuitively the best option, but few case studies have been reported. 

Within 5 to 10 minutes, the ischemia is reversed, and no further action is required.

Figure 34.29 Accidental discharge of an EpiPen into a finger will usually cause intense distal vasoconstriction. A, This patient sustained such an injection into the volar tuft of the index finger (arrow) 2 hours before evaluation in the emergency department. B, Blanching of the distal end of the finger (arrow) caused by the vasoconstriction. C, A pulse oximetry probe was applied to the finger, and normal saturation (96%) (arrow) was found. She was simply observed for another hour, the blanching resolved, and she was discharged without intervention. D, This patient suffered from a similar injection on the volar surface of the middle finger. Blanching is obvious (arrow). E, Phentolamine and lidocaine were injected along the volar surface of the finger. F, Five minutes later the finger was hyperemic (arrow). No further intervention was required.

Pulse oximeter on the fingertip can indicate the degree of resultant hypoxia, an indirect quantification of ischemia, and can also be used to assess reversal therapy.

Injecting the digit with the α-adrenergic antagonist phentolamine (Regitine) can be safely performed in the ED to immediately and permanently reverse the ischemia (see Fig. 34.29D–F). Phentolamine is reconstituted and diluted in 2% plain lidocaine at a ratio of 1.5:1, or diluted with saline at a ratio of 1:1. Inject a small aliquot of the mixture with a 25- to 27-gauge needle. Both a digital block and direct injection of the reversal agent into the site of epinephrine injection have been described. Injecting the actual site of penetration is intuitively the best option, but few case studies have been reported. Within 5 to 10 minutes, the ischemia is reversed, and no further action is required.


82. Updated recommendations for the use of tetanus toxoid, reduced diphtheria toxoid and acellular Pertussis (Tdap) vaccine from the Advisory Committee on Immunization Practices. MMWR Morb Mortal Wkly Rep 60:13, 2011.


CHAPTER 35

Methods of Wound Closure

Richard L. Lammers and Lovita E. Scrimshaw

WOUND TAPE

Surgical tape strips are now routinely used to close simple wounds. Tape strips can be applied by health care personnel in many settings, including emergency departments (EDs), operating rooms, clinics, and first aid stations. Advantages include ease of application, reduced need for local anesthesia, evenly distributed wound tension, no need for suture removal, no residual suture marks, minimal skin reaction, superiority for some grafts and flaps, and suitability for use under plaster casts. One main advantage of wound tape over standard sutures and wound staples is its greater resistance to wound infection.1–3 Wound closure strips are the most cost-effective means for closure of low-tension lacerations and provide results comparable to tissue adhesive.4–6

Background and Tape Comparisons

Currently, several brands of tape with different porosity, flexibility, strength, and configuration are available. Steri-Strips (3M Corporation, St. Paul, MN) are nonwoven microporous tapes with ribbed backing (Fig. 35.1 and Video 35.1). They are porous to air and water, and the ribbed backing provides extra strength. Shur-Strip (Deknatel, Inc., Floral Park, NY) is another nonwoven microporous tape. Cover-Strips (Beiersdorf, South Norwalk, CT) are woven in texture and have a high degree of porosity. They allow not only air and water but also wound exudates to pass through the tape. Clearon (Ethicon, Inc., Somerville, NJ) is a synthetic plastic tape whose backing contains longitudinal parallel serrations to permit gas and fluid permeability. An iodoform-impregnated Steri-Strip (3M Corporation) is intended to further retard infection without sensitization to iodine.4 Other tape products include Curi-Strip (Kendall, Boston), Nichi-Strip (Nichiban Co., Ltd., Tokyo), Cicagraf (Smith & Nephew, London), and Suture Strip (Genetic Laboratories, St. Paul, MN). Steri-Strip S (formerly marketed as ClozeX) is a novel method of tape application which has shown benefit in surgical wounds.9–14

Scientific studies of wound closure tapes provide some comparisons of products. Koehn15 showed that Steri-Strip tape maintains adheriveness approximately 50% longer than Clearon tape does. Rodeheaver and coworkers16 compared Shur-Strip, Steri-Strip, and Clearon tape in terms of breaking strength, elongation, shear adhesion, and air porosity. The tapes were tested in both dry and wet conditions. Steri-Strip tape had approximately twice the breaking strength of the other two tapes in both conditions; there was minimal loss of strength in all tapes when wetted. Shur-Strip tape showed approximately two to three times the elongation of the other tapes at the breaking point, whether dry or wet. Shear adhesion (amount of force required to dislodge the tape when a load is applied at the place of contact) was slightly better for the Shur-Strip tape than for the Steri-Strip tape and approximately 50% better than for the Clearon tape. Of these three wound tapes, the investigators considered Shur-Strips to be superior for wound closure.

One comprehensive study of wound tapes compared Curi-Strip, Steri-Strip, Nichi-Strip, Cicagraf, Suture Strip, and Suture Strip Plus.17 All tapes were 12 mm wide except for Nichi-Strip, which was 15 mm. Each tape was compared for breaking strength, elongation under stress, air porosity, and adhesiveness. Curi-Strip, Cicagraf, and Steri-Strip exhibited equivalent dry breaking strength. However, when wet (a condition that can occur in the clinical setting), Cicagraf outperformed all tapes. All the tapes tested had similar elongation-under-stress profiles with the exception of Suture Strip Plus. This tape did not resist elongation under low or high force. Excessive elongation may allow wound dehiscence. Nichi-Strip was the most porous to air, and Cicagraf was almost vapor impermeable. Nichi-Strip and Curi-Strip had the best adherence to untreated skin. When the skin was treated with tincture of benzoin, however, Steri-Strip dramatically outperformed all other products. When all study parameters were considered, Nichi-Strip, Curi-Strip, and Steri-Strip achieved the highest overall performance rankings.

Indications

The primary indication for tape closure is a superficial straight laceration under little tension. If necessary, tension can be reduced by placing deep closures. Areas particularly suited for tape closure are the forehead, chin, malar eminence, thorax, and non–joint-related areas of the extremities. Tape may also be preferred for wounds in anxious children when suture placement is not essential. In young children who are likely to remove the tape, tape closure must be protected with an overlying gauze bandage.

In experimental wounds inoculated with Staphylococcus aureus, tape-closed wounds resisted infection better than wounds closed with nylon sutures.2 Tape closure works well under plaster casts when superficial suture removal would be delayed. Wound tape effectively holds flaps and grafts in place, particularly over the fingers, the flat areas of the extremities, and the trunk (Fig. 35.2).18 Tape closure can also be applied to wounds after early suture removal, particularly on the face, to maintain approximation of the wound edges while reducing the chance of permanent suture mark scarring. Finally, because of the minimal skin tension created by tape, it can be used on skin that has been compromised by vascular insufficiency, altered by prolonged use of steroids, and in the fragile skin of the aged. For example, wounds on the pretibial area are difficult to close, especially so in the elderly because of tissue atrophy. Wound tape provides an alternative or adjunct to suture closure in this situation.

Contraindications

Tape closure has disadvantages as well. Tape does not work well on wounds under significant tension or on wounds that are irregular, on concave surfaces, or in areas of marked tissue laxity. In many cases, tape does not provide satisfactory apposition of the wound edges without concurrent underlying deep closure. Naturally moist areas, such as the axilla, the palms of the hands, the soles of the feet, and the perineum prevent tape from sticking well. Tape also has difficulty adhering to wounds
that will have secretions, copious exudates, or persistent bleeding. It is of little value on lax and intertriginous skin, on the scalp, and on other areas with a high concentration of hair follicles. Tape strips are also at risk for premature removal by young children.

Do not place tape tightly and circumferentially around digits because it has insufficient ability to stretch or lengthen (Fig. 35.3). If placed circumferentially, the natural wound edema of an injured digit can make the tape act like a constricting band, which can lead to ischemia and possible necrosis of the digit. Use semicircular or spiral placement techniques if digits are to be taped.

**Equipment**

For simple tape closure, the equipment required includes forceps and tape of the proper size (see Fig. 35.1). Most taping can be performed using forceps, Steri-Strips, or cotton-tipped applicators.

![Figure 35.1 Equipment required for the application of wound tape. Wound tape (3M Steri-Strip brand depicted here) comes in a variety of widths. The use of benzoin helps Steri-Strips stay in place longer.](image)

**Figure 35.2** Wound tape in the care of avulsion injuries is preferred over sutures (see Fig. 35.5). A skin avulsion in the elderly following minor trauma is an ideal wound to repair with closure tape because such injuries cannot be closed with sutures. The goal is to provide approximation of the avulsed skin and apply pressure to avoid movement of the skin flap or accumulation of fluid under the avulsion. Tissue glue can augment this procedure. A, An elderly woman who was taking steroids had extremely thin skin and suffered a skin avulsion that could not be repaired with sutures. B, The skin edges are uncurled, stretched, and anatomically replaced. C, The wound should heal while closure tape keeps the skin in place. Tissue glue (Dermabond, Ethicon Inc.) was also dabbed on various parts of the edges to allow egress of fluid. D, Another elderly patient with a large avulsion injury on the forearm. E, Repair with Steri-Strips (3M) and skin glue. F, A compression dressing, such as an elastic bandage or a Dome paste (Unna boot) dressing, can be applied to minimize movement of the flap and decrease buildup of fluid under the flap.

![Figure 35.3 After suturing this proximally based flap, Steri-Strips (3M) are applied under a tourniquet to compress the flap, arrest motion of the flap, and lessen buildup of fluid. Tissue movement and fluid buildup are some reasons why flaps and avulsed skin fail to heal. Tape should be placed in a semicircular or spiral pattern on digits to avoid constriction.](image)
be done in the ED with 1/4-inch × 3-inch strips. For wounds larger than 4 cm, however, 1/2-inch–wide strips provide greater strength. Most companies manufacture strips up to 1 inch wide and 4 inches long.

Procedure

Proper wound preparation, irrigation, débridement, and hemostasis must precede tape closure. Fine hair may be cut short. Dry the area of tape application thoroughly to ensure proper adhesion. Do not attempt to apply tape to a wet area or over a wound that is slowly oozing blood because it will usually result in failure of the tape to stick to the skin. On fingers, tape can be applied to a wound that is kept dry by temporarily placing a tourniquet at the base of the finger (see Fig. 35.3).

The technique of applying tape is shown in Fig. 35.4. After the wound has been dried, apply a liquid adhesive such as tincture of benzoin or Mastisol (Eloquest Healthcare Inc, Ferndale, MI) to the skin adjacent to the wound to increase adhesion of the tape. All tape comes in presterilized packages. Open them directly onto the operating field. Handle tape with gloved hands. With the backing still attached, cut the tape to the desired length or long enough to allow approximately 2 to 3 cm of overlap on each side of the wound. After the end tab is removed, gently peel off the tape from its backing using forceps by pulling straight back. Do not pull to the side because the tape will curl and be difficult to apply to the wound. Place half of the tape securely on one side at the midportion of the wound. Gently but firmly appose the opposite wound edge to its counterpart. Apply the second half of the tape next. Hold the wound edges as close together as possible and at equal height to prevent the development of a linear, pitted scar. Apply additional tape by bisecting the remainder of the wound. Place a sufficient number of tape strips so that the wound is completely apposed without totally covering the entire length of the wound. An arrangement of tape strips parallel to each other and perpendicular to the wound provides good tape adherence. An arrangement of tape strips parallel to each other and perpendicular to the wound provides good tape adherence over time. Finally, if using a woven tape, place additional cross strips to add support and prevent blistering caused by unsupported tape ends. Cross strips are unnecessary when using nonwoven tape, such as Steri-Strip or Shur-Strip, because nonwoven tapes have sufficient ability to elongate, which also prevents blister formation. The use of cross strips with nonwoven tape actually decreases the longevity of tape adherence.

Do not cover taped wounds with occlusive dressings. Adhesive bandages (e.g., Band-Aids) and other impermeable dressings promote excessive moisture, which can lead to premature separation of tape strips from the wound. An adhesive bandage may also adhere to the tapes and pull them off the skin during dressing changes. Keep the tape in place for approximately 2 weeks or longer, if necessary. Instruct the patient to clean the taped laceration gently with a slightly moist, soft cloth after 24 to 48 hours. However, emphasize that if excessive wetting or mechanical force is used, premature tape separation may result. Instruct patients to gently trim the curled edges of the closure tape with fine scissors to avoid premature loss of the tape.

Pretibial lacerations, particularly in the aged and in those with thin skin, may be challenging to manage. Some studies of pretibial lacerations suggest using wound tape alone. However, wounds in this location are under great tension, and wound tape alone may be inadequate. Several studies have shown good results with a combination of tape and sutures. The approach by Silk showed improved healing time and decreased need for skin grafting (Fig. 35.5). Steri-Strips are placed parallel to and 1 cm from the wound edges, and then 2-0 silk sutures are used to approximate the wound margins. For flap-type lacerations, an immobilizing suture placed in the middle of the flap would act much like a button, lessening the tension along the wound margins. Bain and coworkers advocate using tissue glue rather than tissue tape to support suture placement.

Complications

Complications are uncommon with tape closure. The wound infection rate in clean wounds closed with tape compares favorably with rates for other standard closures. The cosmetic result of closure with wound tape is comparable to other closure methods. Premature tape separation occurs in approximately 3% of cases. Other complications include: (1) skin blistering, which occurs if the tape is stretched too tightly across the wound, and (2) wound hematoma, which can result if hemostasis is inadequate. Tape may loosen prematurely over shaved areas as hair grows back.

When tincture of benzoin is used, apply it carefully to the surrounding, uninjured skin. If spillage into the wound occurs, there is a higher risk for infection. Benzoin vapor can cause pain when applied near an open wound that has not been anesthetized. Benzoin can also injure the conjunctival and corneal membranes of the eye.

Summary

Modern tape products and techniques serve a valuable role in the management of minor wounds in the ED. In selected wounds, tape closure is as successful as suture closure. Closure tape should be considered for superficial wounds in cosmetically unimportant areas and for wounds on relatively flat surfaces that are too wide for simple dressings but do not require sutures.

Tissue Adhesive (Tissue Glue)

Tissue adhesive (also called tissue glue) provides a simple, rapid method of wound closure that is likely underused. Tissue adhesive has been approved for use in the United States since 1998. Two types of tissue adhesive are available: 2-octylcyanoacrylate (Dermabond, Ethicon, Inc; derma+flex QS, Chemence Medical Products, Alpharetta, GA; SurgiSeal, Adhezion Biomedical, Hudson, NC) and N-butyl-2-cyanoacrylate (Indermil, Convexicon Medical Ltd., Dublin, Ireland; Histoacryl, TissueSeal LLC, Ann Arbor, MI; LiquiBand (Advanced Medical Solutions, Winsford, United Kingdom) is a blend of an octylcyanoacrylate and butylcyanoacrylate. Animal studies have shown octylcyanoacrylate-based adhesive to have significantly greater strength and flexibility than butylcyanoacrylate-based adhesive, and Dermabond Advanced has better viscosity, as well as strength and flexibility, among the products in this category. Dermabond, like other tissue adhesives, is packaged in sterile, single-use ampules (Fig. 35.6 and Video 35.2). These bonding agents can be used on superficial wounds, even in hair-bearing areas. Tissue adhesives polymerize on contact with water. These
1. After wound preparation (and placement of deep closures, if needed), dry the skin thoroughly at least 2 inches around the wound. Failure to dry the skin and failure to obtain perfect hemostasis are common causes of failure of tape to stick to the skin.

2. Apply a thin coating of tincture of benzoin around the wound to enhance tape adhesiveness. Benzoin should not enter the wound because it increases the risk of infection. Do not allow benzoin to enter the eye.

3. Cut the tape to the desired length before removing the backing.

4. The tape is attached to a card with perforated tabs on both ends. Gently peel the end tab from the tape.

5. Use forceps to peel the tape off the card backing. Pull directly backward, not to the side.

6. Place half of the first tape at the midportion of the wound; secure firmly in place.

7. Gently but firmly appose the opposite side of the wound with the free hand or forceps. If an assistant is not available, the operator can approximate the wound edges. The tape should be applied by bisecting the wound until the wound is closed satisfactorily.

8. Wound margins are completely apposed without totally occluding the wound.

9. Only if using woven tape strips, additional supporting strips of tape are placed approximately 2.5 cm from the wound and parallel to the direction of the wound. Taping in this manner prevents the skin blistering that may occur at the ends of the tape.

**Figure 35.4** Proper technique for the application of wound tape.
repairing nail bed lacerations and securing the nail plate to the nail bed (see later section on Nail Bed Lacerations).

**Procedure**

Tissue adhesive can be used to approximate wounds that do not require deep-layer closure and do not have significant tension on the edges of the wound (Fig. 35.7). In preparation for closure, clean and anesthetize the wound. Débride the wound if necessary. It is especially important to control bleeding from the wound.

Hold the wound edges together with forceps, gauze pads, or fingers. Squeeze the small, cylindrical plastic container to expel droplets of tissue adhesive through the cotton-tipped applicator at the end of the container. Apply the adhesive in at least three to four thin layers along the length of the wound's surface, and extend it approximately 5 to 10 mm from each side of the wound. The purple color of the solution facilitates placement of the droplets. Support and hold the edges of the wound together for at least 1 minute while the adhesive dries. Low-viscosity tissue adhesives may seep into the wound or trickle off rounded surfaces during application. This tendency toward migration or “runoff” can be minimized by using high-viscosity adhesives (e.g., Dermabond HV Topical Skin...
Adhesive, Ethicon, Inc.), positioning the wound horizontally, or applying the adhesive slowly. Contain runoff with wet gauze or by creating a barrier of petrolatum.

Wound closures with tissue adhesive can be reinforced by pulling the edges of the wound into apposition with a few strips of porous surgical tape before application of the adhesive. Tissue adhesive can be placed on top of surgical tape, but tape should not be placed on top of dried tissue adhesive. Once the adhesive has dried completely, further protect the closure with a nonocclusive bandage.

Tissue adhesives can also be used to treat superficial skin tears that do not extend past the dermis. After the wound and skin flap are cleaned, lay the flap over the base of the wound and approximate the edges of the wound. Express any remaining blood and serum from under the flap, and ensure the entire area is dry. Apply a thin layer of adhesive over the wound margins and 1 to 2 cm beyond the margin. After the first layer dries, apply a second layer of adhesive. A dressing is not required unless there is a reason to protect the wound from repeated injury.

The primary advantage of tissue adhesive is the speed of closure. Wounds can be closed in as little as one-sixth the time required for repair with sutures. Application is rapid and painless. Use of tissue adhesive avoids suture marks adjacent to the wound and reduces the risk for needledstick injuries to health care personnel. Wounds closed with tissue adhesive have less tensile strength in the first 4 days than do sutured wounds, but 1 week after closure, the tensile strength and overall degree of inflammation in wounds closed with tissue adhesive are equivalent to those closed with sutures. Cosmetic results are similar to those obtained with suture repair. Tissue adhesive serves as its own wound dressing and has an antimicrobial effect against gram-positive organisms. The material sloughs off in 5 to 10 days, thereby saving the patient from another visit to a clinician. Do not apply ointments or occlusive bandages on wounds closed with tissue adhesive.

**Complications**

Although tissue adhesive is classified as nontoxic, some authors warn against placing it within the wound cavity. However, various investigators have applied specially formulated octyl-2-octylcyanoacrylate (liquid adhesive bandage) directly to open minor lacerations, abrasions, and partial-thickness wounds. Though rare, exudates, pain, tenderness, swelling, and foreign body reactions have been reported after tissue adhesive has entered the wound. When compared with adhesive bandages, wounds covered with tissue adhesive had equivalent rates of healing and complications.
If hemostasis is inadequate or an excessive amount of adhesive is applied too quickly, the patient can experience a burning sensation or sustain a local burn from the heat of polymerization. After polymerizing, tissue adhesive can fracture with excessive or repetitive movement. Although gentle rinsing is permitted, if the adhesive is washed or soaked, it will peel off in a few days, before the wound is healed. 35,36

If the clinician’s gloved fingers, gauze, or plastic instruments contact the tissue adhesive during application, the glove may adhere to the patient’s skin. Tissue adhesive can be removed with antibiotic ointment or petrolatum jelly or more rapidly with acetone. 46 Indermil (Connexicon Medical) must be stored under refrigeration.

One risk involving the use of tissue adhesive is its ease of use; clinicians may fail to adequately clean wounds before closure with tissue adhesive. 34 It should not be used to close infected wounds. There is a slightly higher risk for wound dehiscence in closures with tissue glue than with sutures. 35,36

If the edges of the wound cannot be held together without considerable tension, tissue adhesive should not be used. 46 Percutaneous sutures provide a more secure immediate closure than tissue adhesive. 41 It should also not be used over or near joints, on moist or mucosal surfaces, or on wounds under significant static or dynamic skin tension. See Fig. 35.7 for information on managing eyelids that are accidentally glued shut.

Wound staples

Background

Automatic stapling devices have become commonplace for closure of surgical incisions and traumatic wounds (Fig. 35.8). Clinical studies of patients with stapled surgical incisions have found no significant difference between stapling and suturing when infection rates, healing outcome, and patient acceptance are compared. 37-41 Wound stapling (Video 35.3) and nylon suture closure of skin compared favorably in wound tensile strength, complication rates, patient tolerance, efficiency of closure, scar width, color, general appearance, suture or staple marks, infection rates, and cost. In animal models, staples caused less wound inflammation and offered more resistance to infection with contaminated wounds. 62-65

The most significant advantage of wound stapling over suturing is speed of closure. The most significant downside is loss of the cosmetic effect that can be achieved with meticulous suture closure. On average, stapling is three to four times faster than suturing traumatic wounds. 66,67 When clinician time and cost of instruments are considered, the difference in cost between stapling and suturing is either minimal 67 or favors stapling. 66,69

Indications and Contraindications

The indications for stapling are limited to relatively linear lacerations with straight, sharp edges located on an extremity, the trunk, or the scalp. Staples may be especially useful for superficial scalp lacerations in an agitated or intoxicated patient. Because of their superficial placement in the adult scalp (usually above the galea), staples are not recommended for deep scalp lacerations. Staples may not provide the same hemostasis that is possible with deep sutures. They should not be placed in scalp wounds if computed tomography head scans are to be performed because staples produce scan artifacts. Similarly, staples should not be used if the patient is expected to undergo magnetic resonance imaging because the powerful magnetic fields may avulse the staples from the surface of the skin. Staples should not be used on the face, neck, hands, or feet.

Equipment

Standard wound care should precede wound closure. In many cases, when debridement and dermal (deep) closure are unnecessary, only tissue forceps are needed to assist in evertting wounds.

Many stapling devices are commercially available. The most versatile and least expensive is the Precise stapler (3M Corporation). Different units that hold between 5 and 25 staples can be purchased. The 10-staple unit will suffice for most lacerations. Other devices include the Proximate 11 (Ethicon, Inc.), Appose (Covidien, New Haven, CT), and Reflex One (ConMed, Utica, NY).

A novel device, the Insorb (Incisive Surgical, Inc., Plymouth, MN) absorbable stapler, uses absorbable staples (made of polylactic and polyglycolic acids) placed intradermally. The device has shown both time and cost savings in repair of abdominal and breast surgical wounds comparable to intradermal sutures. 70,71 The device is not widely available in the emergency setting.

Procedure

Before stapling, sometimes it is necessary to place deep, absorbable sutures to close deep fascia and reduce tension in the superficial fascia and dermal layers. To facilitate the stapling process, the edges of the wound should be everted, preferably by a second operator (Fig. 35.9, step 1). The assistant precedes the operator along the wound and everts the edges of the wound with forceps or pinches the skin with the thumb and forefinger. Stapling flattened wound edges may place the staple precisely but results in inversion of the wound. Once the edges are held in eversion, the staple points are gently placed across the wound. The center of the staple device should be placed over the center of the wound to ensure the best closure and

![Image](Image.png)
Wound Staples

Approximate and evert the skin edges by hand or with forceps before they are secured with staples. If possible, have an assistant perform this duty. Failure to evert the wound edges is a common error that may cause an unacceptable result.

Align the stapler to the center of the wound. Squeeze the stapler handle to advance one staple into the wound margins. Do not press too hard on the skin to prevent placing the staple too deeply.

As the handle is squeezed, an anvil automatically bends the staple to the proper configuration.

Allow a small space to remain between the skin and the crossbar of the staple. Excessive pressure created by placing the staple too deep causes wound edge ischemia, as well as pain on removal. Note that the staple bar is 2 to 3 mm above the skin line.

Supply the patient with a staple remover when being referred to an office for removal or for self-removal. To remove the staple, place the lower jaw of the remover under the crossbar of the staple.

Squeeze the handle gently, and the upper jaw will compress the staple and allow it to exit the skin.

Figure 35.9 Wound staples. Failure to align the center of the staple device directly over the center of the laceration is a common cause of a less than ideal staple closure. (Steps 1 and 2, From Custalow C: *Color atlas of emergency department procedures*, Philadelphia, 2005, Elsevier; steps 3–6, from Edlich RF: *A manual for wound closure*, St. Paul, MN, 1979, 3M Medical-Surgical Products. Reproduced by permission.)
Methods of Wound Closure

To avoid overriding or misaligned edges (which can be difficult in an uncooperative patient). When the stapler handle or trigger is squeezed, the staple is advanced automatically into the wound and bent to the proper configuration (see Fig. 35.9, steps 2 and 3). The operator should not press too hard on the skin surface to prevent placing the staple too deeply and causing ischemia within the staple loop. When placed properly, the crossbar of the staple is elevated a few millimeters above the surface of the skin (see Fig. 35.9, step 4). A sufficient number of staples should be placed to provide proper apposition of the edges of the wound along its entire length. (One suggested method is to place a staple in the midpoint of the length of the wound and then bisect either side of the wound with additional staples. Continue the process until the edges of the wound are well opposed.) After the wound is stapled, an antibiotic ointment may be applied. Stapled wounds may be left uncovered, or a sterile dressing may be applied. The patient can remove the dressing and gently clean the wound in 24 to 48 hours. Scalp lacerations can be cleansed by showering within a few hours.

Removal of staples requires a special instrument made available by each manufacturer of stapling devices. The lower jaw of the staple remover is placed under the crossbar, and the handle is squeezed (see Fig. 35.9, steps 5 and 6). This action compresses the crossbar and bends the staple outward, thereby releasing the points of the staple from the skin. Many primary care physicians do not routinely stock the instrument. If referred to a primary care physician's office for removal of staples, the patient can be given a disposable staple removal device to bring with them. If well-instructed, patients can remove their own staples with the removal device. The interval between staple application and removal is the same as that for standard suture placement and removal. Staples can cause significant scarring if left in place too long (Fig. 35.10).

Complications

Patient acceptance, comfort, and rates of wound infection and dehiscence are similar with staple-closed wounds and sutured wounds. However, removal of staples can be somewhat more uncomfortable than removal of sutures.

A common error during insertion of staples is failure to evert the edges of the skin before stapling (see Fig. 35.10). Eversion avoids the natural tendency of the device to invert the closure. Eversion may be accomplished with forceps or by pinching the skin with the thumb and index finger, a procedure that requires some practice. Another common error is to fail to align the middle of the staple exactly in the midline of the wound.

Staples will cause marks in the skin that are similar to sutures. However, in patients who tend to scar more easily, the scar

Figure 35.10 A, A very poor result occurred when staples (some marked with arrows) were used to close this deep scalp laceration. The edges of the wound were not everted, staples were not in the midline, sections of the skin overlapped significantly, poor hemostasis was obtained, and the galea could not be closed by the superficial staples. The patient did not wash his hair as instructed. B, Three days later during inspection of the wound, the staples were removed and C, the laceration was closed with 3-0 interrupted nylon sutures. The clinician should attempt to obtain a cosmetic closure on all scalp lacerations because as patients lose their hair, a previously hidden, unsightly scar emerges. In general, staples should not be used to close full-thickness scalp lacerations, especially wounds that are actively bleeding. D, Sloppy stapling on an extremity with inversion rather than eversion of the edges of the wound. Some staples are totally misaligned and barely include the opposite skin edge. E, The staples were left in too long, which caused a poor cosmetic result.
resulting from staples may be more pronounced than that produced by sutures, especially if the staples are left in place for prolonged periods.

Wound stapling achieves results that are generally comparable to those of sutures for the closure of traumatic, linear lacerations in noncosmetic areas, such as the scalp, trunk, and extremities. Stapling is much faster than suturing. Wound stapling does not differ significantly from suturing in terms of cost, infection rates, wound healing, and patient acceptance, but cosmesis may suffer, especially if the staples are left in too long (see Fig. 35.10D and E).

SUTURES

In the United States, most traumatic wounds are closed by suturing.

Equipment

Instruments

In addition to the instruments used for débridement, a needle holder and suture scissors are required for suturing (Fig. 35.11). The mechanical performance of disposable needle holders distributed by different surgical instrument companies varies considerably. The size of the needle holder should match the size of the needle selected for suturing; the needle holder should be large enough to hold the needle securely as it is passed through tissue, yet not so large that the needle is crushed or bent by the instrument.

Instruments used to débride a grossly contaminated wound should be discarded and replaced with fresh instruments for closure. Instruments covered with coagulated blood can be cleansed with hydrogen peroxide, rinsed with sterile saline or water, and then used for suturing.

In addition, 2.5 loupe magnification may improve inspection of wounds and facilitate the repair of facial and hand wounds.

Suture Material

A wide variety of suture material is available to satisfy clinician preferences (Fig. 35.12). There are no definitive standards on which to base the use of suture material for a particular wound or site. For most wounds that require closure of more than one layer of tissue, the clinician must choose sutures from two general categories: an absorbable suture for the deeper, subcutaneous (SQ) layer and a nonabsorbable suture for surface (percutaneous) closure.

Sutures can be described in terms of four characteristics:
1. Composition (i.e., chemical and physical properties).
2. Handling characteristics and mechanical performance.
3. Absorption and reactivity.
4. Size and retention of tensile strength.

Composition

Sutures are made from natural fibers (cotton, silk), from sheep submucosa or beef serosa (plain gut, chromic gut), or from synthetic material such as nylon (Dermalon [Medtronic, Minneapolis, MN], Ethilon [Ethicon Inc.], Nurolon [Ethicon Inc.], Surgilon [Medtronic]), Dacron (Ethiflex, Mersilene [both Ethicon Inc.]), polyester (Ti-Cron [Medtronic]), polyethylene (Ethibond [Ethicon Inc.]), polypropylene (Prolene [Ethicon Inc.]), polyglycolic acid (Dexon [Medtronic]), and polyglactin (Vicryl, coated Vicryl, Vicryl Rapide [all Ethicon Inc.]). Stainless steel sutures are rarely, if ever useful in wound closure in the ED setting because of handling difficulty and fragmentation. Some sutures are made of a single filament (monofilament); others consist of multiple fibers braided together (Table 35.1).

Handling and Performance

Desirable handling characteristics in a suture include smooth passage through tissue, ease in knot tying, and stability of the knot once tied (Table 35.2). Smooth sutures pull through tissue easily, but knots slip more readily. Conversely, sutures with a high coefficient of friction have better knot-holding capacity but are difficult to slide through tissue. Smooth sutures will loosen after the first throw of a knot is made, and thus a second throw is needed to secure the first in place. Rougher, multifilament types of suture make it more difficult to tighten a knot further after the first throw is made.

Multifilament sutures have the best handling characteristics of all sutures, whereas steel sutures have the worst. In terms of performance and handling, significant improvements have been made in the newer absorbable sutures. Gut sutures have

Figure 35.11 Basic suturing equipment includes a needle driver, forceps, and suture scissors.

Figure 35.12 A wide variety of both absorbable and nonabsorbable suture material is available. Other variables include suture size (e.g., 4-0, 5-0, 6-0), needle size, and type (e.g., PS-2, reverse cutting, conventional cutting). These variables are discussed in detail in the text.
many shortcomings, including relatively low and variable strength, a tendency to fray when handled, and stiffness despite being packaged in a softening fluid. Multifilament synthetic absorbable sutures are soft and easy to tie and have few problems with knot slippage. Polyglyactin 910 (coated Vicryl, Ethicon Inc.) sutures have an absorbable lubricant coating. The frictional drag of these coated sutures as they are pulled through tissue is less than that of uncoated multifilament materials, and resetting of knots after the initial throw is much easier. This characteristic allows retightening of a ligature without knotting or breakage and with smooth, even adjustment of suture line tension in running subcuticular stitches. Synthetic monofilament sutures have the troublesome property of “memory”: a tendency of the filament to spring back to its original shape, which causes the knot to slip and unravel. Some nonabsorbable monofilament sutures are coated with polytetrafluoroethylene (Teflon) or silicone to reduce their friction. This coating improves the handling characteristics of these monofilaments but results in poorer knot security.

Three square knots will secure a stitch made with silk or other braided, nonabsorbable material, and four knots are sufficient for synthetic, absorbable, and nonabsorbable monofilament sutures. Five knots are needed for the Teflon-coated synthetic Tefdek (Teleflex, Morrisville, NC). When coated synthetic suture material is used, attention to basic principles of knot tying is even more important. An excessive number of throws in a knot weakens the suture at the knot. If the

### TABLE 35.1 Commonly Available Suture Material

<table>
<thead>
<tr>
<th>ABSORBABLE SUTURE</th>
<th>NONABSORBABLE SUTURE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Monofilament</strong></td>
<td></td>
</tr>
<tr>
<td>Plain gut</td>
<td>Dermalon (nylon)</td>
</tr>
<tr>
<td>Chromic gut</td>
<td>Ethilon (nylon)</td>
</tr>
<tr>
<td>PDS (polydioxanone)</td>
<td>Novafil (polybutester)</td>
</tr>
<tr>
<td>Maxon (polyglyconate)</td>
<td>Prolene (polypropylene)</td>
</tr>
<tr>
<td>Monocryl (poliglecaprone 25)</td>
<td>Silk</td>
</tr>
<tr>
<td>Biosyn (glycomer 631)</td>
<td>Steel</td>
</tr>
<tr>
<td></td>
<td>Surgilene (polypropylene)</td>
</tr>
<tr>
<td></td>
<td>Tevdek (Teflon coated)</td>
</tr>
<tr>
<td><strong>Multifilament</strong></td>
<td></td>
</tr>
<tr>
<td>Dexon (polyglycolic acid)</td>
<td>Ethibond (polyethylene)</td>
</tr>
<tr>
<td>Coated Vicryl (polyglactin 910)</td>
<td>Mersilene (braided polyester)</td>
</tr>
<tr>
<td></td>
<td>Nurolon (nylon)</td>
</tr>
<tr>
<td></td>
<td>Surgilon (nylon)</td>
</tr>
<tr>
<td></td>
<td>Ti-Cron (polyester)</td>
</tr>
</tbody>
</table>

### TABLE 35.2 Characteristics of Suture Material

<table>
<thead>
<tr>
<th>SUTURE MATERIAL</th>
<th>KNOT SECURITY</th>
<th>TENSILE STRENGTH</th>
<th>TISSUE REACTIVITY</th>
<th>DURATION OF SUTURE INTEGRITY (DAYS)</th>
<th>TIE ABILITY (HANDLING)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surgical gut</td>
<td>Poor</td>
<td>Fair</td>
<td>Greatest</td>
<td>5–7</td>
<td>Poor</td>
</tr>
<tr>
<td>Chromic gut</td>
<td>Fair</td>
<td>Fair</td>
<td>Greatest</td>
<td>10–14</td>
<td>Poor</td>
</tr>
<tr>
<td>Coated Vicryl</td>
<td>Good</td>
<td>Good</td>
<td>Minimal</td>
<td>30</td>
<td>Best</td>
</tr>
<tr>
<td>Dexon</td>
<td>Best</td>
<td>Good</td>
<td>Minimal</td>
<td>30</td>
<td>Best</td>
</tr>
<tr>
<td>PDS</td>
<td>Fair</td>
<td>Best</td>
<td>Least</td>
<td>45–60</td>
<td>Good</td>
</tr>
<tr>
<td>Maxon</td>
<td>Fair</td>
<td>Best</td>
<td>Least</td>
<td>45–60</td>
<td>Good</td>
</tr>
<tr>
<td>Biosyn</td>
<td>Poor</td>
<td>Good</td>
<td>Minimal</td>
<td>14–21</td>
<td>Good</td>
</tr>
<tr>
<td>Monocryl</td>
<td>Good</td>
<td>Best</td>
<td>Minimal</td>
<td>7</td>
<td>Good</td>
</tr>
<tr>
<td>Vicryl Rapide</td>
<td>Good</td>
<td>Good</td>
<td>Minimal</td>
<td>5</td>
<td>Good</td>
</tr>
<tr>
<td><strong>Nonabsorbable</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethilon</td>
<td>Good</td>
<td>Good</td>
<td>Minimal</td>
<td></td>
<td>Good</td>
</tr>
<tr>
<td>Novafil</td>
<td>Good</td>
<td>Good</td>
<td>Minimal</td>
<td></td>
<td>Good</td>
</tr>
<tr>
<td>Prolene</td>
<td>Least</td>
<td>Best</td>
<td>Least</td>
<td></td>
<td>Fair</td>
</tr>
<tr>
<td>Silk</td>
<td>Best</td>
<td>Least</td>
<td>Greatest</td>
<td></td>
<td>Best</td>
</tr>
</tbody>
</table>

PDS, Polydioxanone.

Absorption and Reactivity

Sutures that are rapidly degraded in tissue are termed absorbable; those that maintain their tensile strength for longer than 60 days are considered nonabsorbable (see Table 35.1). Plain gut may be digested by white blood cell lysozymes in 10 to 40 days; chromic gut will last 15 to 60 days. Remnants of both types of suture, however, have been seen in wounds more than 2 years after placement.\(^{51,58,61}\) Ethicon catgut is rapidly absorbed within 10 to 14 days and is associated with less inflammation than chromic catgut.\(^{62}\) Polydioxanone (PDS; Ethicon) and Vicryl produce less tissue reaction and erythema than polypropylene.\(^{63}\) Vicryl is absorbed from the wound site within 60 to 90 days\(^{5,78}\) and Dexon (Medtronic), within 120 to 210 days.\(^{84,85}\) When placed in the oral cavity, plain gut disappears after 3 to 5 days, chromic gut after 7 to 10 days, and Dexon after 16 to 20 days.\(^{86}\) When used on the skin surface, fast absorbing catgut disappears in 9 to 13 days.\(^{87}\) In contrast, SQ silk may not be completely absorbed for as long as 2 years.\(^{78}\) The rate of absorption of synthetic absorbable sutures is independent of suture size.\(^{84}\)

Sutures may lose strength and function before they are completely absorbed in tissues. Braided synthetic absorbable sutures lose nearly all their strength after approximately 21 days. In contrast, monofilament absorbable sutures (modified polygylcolic acid [Maxon, Davis & Geck] and PDS) retain 60% of their strength after 28 days.\(^{88,89}\) Gut sutures treated with chromium salts (chromic gut) have prolonged tensile strength; however, all gut sutures retain tensile strength erratically.\(^{51,58,61}\) Of the absorbable types of suture, a wet and knotted polygylcolic acid suture is stronger than a plain or chromic gut suture subjected to the same conditions.\(^{76,90}\)

Polypropylene remains unchanged in tissue for longer than 2 years after implantation.\(^{91}\) In comparison testing, sutures made of natural fibers such as silk, cotton, and gut were the weakest; sutures made of Dacron, nylon, polyethylene, and polypropylene were intermediate in tensile strength; and metallic sutures were the strongest.\(^{78}\) Comparison of suture strength versus wound strength is a measure of the usefulness of a suture. Catgut is stronger than the soft tissue of a wound for no more than 7 days; chromic catgut, Dexon, and Vicryl are stronger for 10 to 21 days; and nylon, wire, and silk are stronger for 20 to 30 days.\(^ {92}\)

All sutures placed within tissue will damage host defenses and provoke inflammation. Even the least reactive suture impairs the ability of the wound to resist infection.\(^ {93}\) The magnitude of the reaction provoked by a suture is related to the quantity of suture material (diameter \(\times\) total length) placed in the tissue and to the chemical composition of the suture. Among absorbable sutures, polygylcolic acid and poliglactin sutures are the least reactive, followed by chromic gut. Nonabsorbable polypropylene is less reactive than nylon or Dacron.\(^ {76,93,94}\) Significant tissue reaction is associated with catgut, silk, and cotton sutures. Absorbable polygylcolic acid sutures are less reactive than those made of nonabsorbable silk.\(^ {94}\) High tensile strength should be avoided in contaminated wounds.

The chemical composition of sutures is a factor in early infection. The infection rate in experimentally contaminated wounds closed with polygylcolic acid sutures is lower than the rate when gut sutures are used.\(^ {91}\) However, other authors have compared plain gut and nonabsorbable nylon sutures for skin closure in children and found comparable cosmetic results and infection rates.\(^ {96}\)

Lubricant coatings on sutures do not alter suture reactivity, absorption characteristics, breaking strength, or the risk for infection.\(^ {77,94}\) Multifilament sutures provoke more inflammation and are more likely than monofilament sutures to produce infection if left in place for prolonged periods.\(^ {97,98}\) Monofilament sutures elicit less tissue reaction than do multifilament sutures, and multifilament materials tend to wick up fluid by capillary action. Bacteria that adhere to and colonize sutures can envelop themselves in a glycolcalyx that protects them from host defenses,\(^ {99}\) or they can “hide” in the interstices of a multifilament suture and, as a result, be inaccessible to leukocytes.\(^ {97}\) PDS provides the advantages of a monofilament suture in an absorbable form, thus making it a good choice as a subcuticular stitch. Vicryl Plus and Monocryl (Ethicon Inc.) have been found to have less bacterial adherence when compared to Vicryl, making these two materials reasonable choices for subcuticular closure in contaminated wounds.\(^ {100}\) Polypropylene sutures have a low coefficient of friction and subcuticular stitches with this material are easy to pull out.\(^ {101}\)

Size and Strength

The size of suture material (thread diameter) is related to the tensile strength of the suture; threads with greater diameter are stronger. The strength of the suture is proportional to the square of the diameter of the thread. Therefore a 4-0 suture of any type is larger and stronger than a 6-0 suture. The correct suture size for approximation of a layer of tissue depends on the tensile strength of that tissue. The tensile strength of the suture material should be only slightly greater than that of the tissue because the magnitude of damage to local tissue defenses is proportional to the amount of suture material placed in the wound.\(^ {76,92}\)

Synthetic absorbable sutures have made the older, natural suture material unnecessary for most wound closures. Polyglycolic acid (Dexon) and polyglactin 910 (coated Vicryl) have improved handling characteristics, knot security, and tensile strength. Their absorption rates are predictable, and tissue reactivity is minimal.\(^ {103,104}\) The distinct advantages of synthetic nonabsorbable sutures over silk sutures are their greater tensile strength, low coefficient of friction, and minimal tissue reactivity.\(^ {91,100}\) They are extensible and elongate without breaking as the edges of the wound swell in the early postoperative period.\(^ {102,103}\) In contrast to silk sutures, synthetics can easily and painlessly be removed once the wound has healed. The monofilament synthetic suture Novafil (Medtronic) has elasticity that allows a stitch to enlarge with wound edema and to return to its original length once the edema subsides. Stiffer materials lacerate the incircled tissue as the wound swells.\(^ {105}\)

The suture material most useful to emergency clinicians for wound closure is coated Vicryl for SQ layers and synthetic nonabsorbable suture (e.g., nylon or polypropylene) for skin closure.\(^ {106}\) Fascia can be sutured with either absorbable or nonabsorbable material. In most situations, 3-0 or 4-0 suture is used for the repair of fascia, 4-0 or 5-0 absorbable suture for SQ closure, and 4-0 or 5-0 nonabsorbable suture for skin closure. The lips, eyelids, and skin layer of facial wounds are repaired with 6-0 suture, whereas 3-0 or 4-0 suture is used when the edges of the skin are subjected to considerable dynamic stress (e.g., wounds overlying joint surfaces) or static stress (e.g., scalp).
Absorbable sutures have been used as an alternative to nonabsorbable sutures for simple skin closure with similar cosmetic results and infection rates in both adult and pediatric lacerations. Sutures that were evaluated include fast-absorbing gut, Vicryl Rapide, and Monocryl. Absorbable suture used for skin closure obviates the need for suture removal, and thus may be particularly advantageous in children. Still, some authors advise usual patient follow-up for skin closed with nonabsorbable sutures to remove any remaining suture material and to evaluate for complications.

**Needles**

The eyeless, or “swaged,” needle is used for wound closure in most emergency centers (Fig. 35.13). Selection of the appropriate needle size and curvature is based on the dimensions of the wound and the characteristics of the tissue to be sutured. The needle should be large enough to pass through tissue to the desired depth and then to exit the tissue or skin surface far enough that the needle holder can be repositioned on the distal end of the needle at a safe distance from the tip of the needle (Fig. 35.14). Although it is tempting to use the fingers to grasp the tip of the needle to pull it through the skin, this practice risks a needlestick. The clinician should either reposition the needle holder or use forceps to disengage the needle from the laceration.

In wound repair, needles must penetrate tough, fibrous tissue (skin, SQ tissue, and fascia) yet should slice through this tissue with minimal resistance or trauma and without bending. The type of needle best suited for closure of SQ tissue is a conventional cutting needle in a three-eighths or one-half circle (Fig. 35.15). Double-curvature needles (coated Vicryl with PS-4-C cutting needles, Ethicon) may be easier to maneuver in narrow, deep wounds. For surface closure, a conventional cutting-edge needle permits more precise needle placement and requires less penetration force (Fig. 35.16).

**Suturing Techniques (Figs. 35.17 and 35.18)**

**Skin Preparation**

Before suturing, the clinician should ensure adequate exposure and illumination of the wound, and placement of the patient at the appropriate height. The clinician should assume a comfortable standing or sitting position at one end of the long axis of the wound.

The skin surrounding the wound is prepared with a povidone-iodine solution and covered with sterile drapes. A clear plastic drape (Steri-Drape, 3M Corporation) can be used to provide a sterile field and a limited view of the area surrounding the wound. Some surgeons do not drape the face but prefer to leave the facial structures and landmarks adjacent to the wound uncovered and within view. If no drapes are used on the face, the skin surrounding the wound should be widely cleansed and prepared. Wrapping the hair in a sheet or placing the patient’s hair in an oversized scrub hat prevents stray hair from falling into the operating field. For a finger laceration, a sterile glove can be placed on the patient’s hand to avoid using a drape.

**Closure Principles**

There is a tendency to overuse sutures for minor lacerations that will heal nicely with no intervention. Therefore, before suturing, one must assess the need for the procedure (Fig. 35.19).
General Suturing Technique

1. Cleanse the skin surrounding the wound with an antiseptic such as chlorhexidine or povidone-iodine. Avoid introducing antiseptic into the wound because it may be toxic to tissue.

2. Anesthetize the wound prior to exploration and irrigation. Introduce the needle through the wound (as opposed to through the epidermis).

3. Explore the wound to exclude the presence of foreign bodies, gross contamination, or injuries to deep structures. Débride grossly contaminated or devitalized tissue.

4. Irrigate the wound thoroughly until it is visibly clean. Use of a large syringe with a splash guard is ideal. Retract the wound edges with an instrument to facilitate thorough irrigation.

5. Apply a sterile drape, gather the instruments, and ensure that the field is appropriately lit.

6. Place the first suture at the center of the wound so that it bisects the laceration into two equal segments.

7. Tie the knot. The first throw should be a double throw (i.e., surgeon’s knot) to prevent it from loosening. Place an additional three (single) throws and then cut the sutures while leaving 1- to 2-cm tails.

8. Continue to place additional sutures by further bisecting each segment of the laceration. After the last stitch has been placed, cleanse the area and apply an appropriate dressing.

Figure 35.17 General suturing technique.
Instrument Tie

1. Place the needle driver parallel to the wound and wrap the suture end twice over the needle driver. This forms the surgeon’s knot, which prevents the first throw from loosening.

2. Rotate the needle driver 90 degrees and grasp the short suture end on the opposite side of the laceration.

3. Gently pull the suture ends to the side of the laceration opposite their origin. Tighten only enough to approximate the skin edges; avoid overtightening, which may lead to tissue strangulation.

4. To begin the second throw, again place the needle driver parallel to the laceration. Wrap the long suture end over and around the needle driver once. (Only one wrap is used on throws 2 to 4.)

5. Rotate the needle driver 90 degrees and grasp the short suture end on the opposite side of the laceration.

6. Pull the suture ends to the side of the laceration opposite their origin. On the second and subsequent throws, you can tighten the knot down snugly.

7. Place an additional two throws (for a total of four), as depicted in steps 4 through 6. Remember to place the needle driver parallel to the wound and pull the long suture end over the driver; this will ensure that all knots tied are square knots.

8. After the last throw, cut the ends of the suture while leaving 1- to 2-cm tails. Avoid cutting the ends too short, which may lead to knot unraveling or difficulty during suture removal.

Figure 35.18 Instrument tie. (From Custalow C: Color atlas of emergency department procedures, Philadelphia, 2005, Elsevier.)
bones cause the edges of a new wound to gape, and they also
continuously pull on the edges of the wound once it has been
closed. Traumatic loss of tissue or wide excision of a wound
may have the same effect. The best cosmetic result occurs
when the long axis of a wound happens to be parallel to the
direction of maximal skin tension; such alignment brings the
edges of the wound together. Muscles pulling at right angles to the axis of the wound
impose dynamic stress. Swelling after an injury creates additional
tension within the circle of each suture. Skin suture marks
result not only from tying sutures too tightly but also from
failing to eliminate any underlying forces distorting the wound.
Tension can be reduced during wound closure in two ways:
undermining of the wound edges and layered closure.

Undermining. The force required to reapproximate the edges
of a wound correlates with the subsequent width of the scar. Wounds subject to significant static tension require the
undermining of at least one tissue plane on both sides of the
wound to achieve a tension-free closure. To undermine a wound, the clinician frees a flap of tissue from its base at a distance
from the edge of the wound approximately equal to the width
of the gap that the laceration presents at its widest point (Fig. 35.20).

The depth of the incision can be modified, depending on
the orientation of the laceration to skin tension lines and the
 laxity of skin in the area. A No. 15 scalpel blade held parallel
to the surface of the skin is used to incise the adipose layer or
dermal layer of the wound. The clinician can also accomplish
this technique by spreading scissors in the appropriate tissue
plane. Undermining allows the edges of the skin to be lifted
and brought together with gentle traction. Potential complica-
tions of this procedure include injury to cutaneous nerves and
creation of a hematoma under the flap. Because undermining
may harm the underlying blood supply, this technique should
be reserved for relatively uncontaminated wounds when no
other methods adequately relieve wound tension.

Layered Closure. The structure of skin and soft tissue varies
with the location on the body (Fig. 35.21). Most wounds
handled in an ED require approximation of no more than
three basic layers: fascia (and associated muscle), SQ tissue, and
skin surface (papillary layer of the dermis and epidermis).
Any “dead space” (or unapposed edges) within a wound may

Three principles apply to suturing lacerations in any location:
(1) minimize trauma to tissues, (2) relieve tension exerted on
the edges of the wound by undermining and layered wound
closure, and (3) accurately realign landmarks and skin edges
by layered closure and precise suture placement.

Minimizing Tissue Trauma
The importance of carefully handling tissue has been empha-
sized since the early days of surgery. Skin and SQ tissue that
has been stretched, twisted, or crushed by an instrument or
strangled by a suture that is tied too tightly may undergo
necrosis, and increased scarring and infection may result. When
the edges of a wound must be manipulated, the SQ tissues
should be lifted gently with toothed forceps or a skin hook
while avoiding the surface of the skin.

When choosing suture size, the clinician should select the
smallest size that will hold the tissues in place. Skin stitches
should incorporate no more tissue than is needed to coapt the
wound edges with little or no tension. Knots should be tied
securely enough to approximate the edges of the wound without
blanching or indenting the skin surface.

Relieving Tension
Many forces can produce tension on the suture line of a reap-
proximated wound. Static skin forces that stretch the skin over

Figure 35.19 Not all wounds require suturing. A, This child sustained
a superficial forehead laceration just through the epidermis. The
swelling accentuated the defect. Treatment was simply keeping the
laceration clean. B, Four months after the injury, an excellent result
was achieved. As the laceration was healing, it became red and more
noticeable (as do all scars), but eventually faded.

Figure 35.20 The technique of undermining can markedly improve
cosmetic results by relieving wound tension. The scalpel is used to
find an appropriate site; a natural plane often exists at the junction
of the epidermis and dermis.
If a deep, gaping wound is closed without approximation of underlying SQ tissue, a disfiguring depression may develop at the site of the wound. Finally, layered closure provides support to the wound and considerably reduces tension at the skin surface. Several exceptions exist to the general rule of multilayered closure. The adipose layer of soft tissue should not be closed separately. A “fat stitch” is not necessary because little support is provided by closure of the adipose layer and additional suture material may increase the possibility of infection.2,118

Scalp wounds are generally closed in a single layer. For lacerations penetrating the dermis in the fingers, hands, toes, and feet and the sebaceous skin of the nasal tip, the amount of SQ tissue is too small to warrant layered closure; in fact, SQ stitches may leave tender nodules in these sensitive locations. Layered closure is not recommended for wounds without tension, those with poor vascularity, and those with a moderate or high risk for infection. With single-layer closure, the surface stitch should be placed more deeply.92

Suture Placement

SQ Layer Closure

Once the wound has been assessed and appropriately managed for fascial reapproximation, the SQ layer should be addressed (Video 35.4). Although histologically the fatty and fibrous SQ tissue (hypodermis) is an extension of (and continuous with) the reticular layer of the dermis,119 suturing these layers is traditionally referred to as an SQ closure. One approach is to close the length of this layer in segments by placing the first stitch in the middle of the wound and bisecting each subsequent segment until closure of the layer has been completed. This technique is useful for the closure of wounds that are long or sinuous, and it is particularly effective for wounds with one elliptical and one linear side. The needle is grasped with the

---

**Figure 35.21** Variation in the structure of skin. A, Section of the skin of the scalp, ×15. B, Section of human sole perpendicular to the free surface, ×100 (after Maximow AA). C, Section through human thigh perpendicular to the surface of the skin (after Maximow AA.) Blood vessels are injected and appear black (low magnification). (A, Courtesy H. Mizoguchi; B and C, From Bloom W, Fawcett DW: A textbook of histology, ed 10, Philadelphia, 1975, Saunders. Reproduced by permission.)

**Figure 35.22** Note how a subcutaneous suture almost closes the wound with minimal tension on the edges of the skin while obliterating any subcutaneous space. Tie and bury the knot by pulling the sutures in the long axis of the wound as shown.

fill with blood or exudate and enhance the development of infection. Closure of individual layers obliterates this dead space (Fig. 35.22).

Approximation of a lacerated muscle hastens healing and return of function to the muscle. However, the fascia, not muscle, should be sutured. Muscle tissue itself is too friable to hold a suture. Layered closure is particularly important in the management of facial wounds; this technique prevents scarring of muscle to the SQ tissue and consequent deformation of the surface of the wound with contraction of the muscle.
Some clinicians can suture more rapidly if the fingers are placed on the midshaft of the needle holder rather than in the rings of the instrument (Fig. 35.24). The suture enters the SQ layer at the bottom of the wound (Fig. 35.25A) or, if the wound has been undermined, at the base of the flap (see Fig. 35.25B) and exits in the dermis. Once the suture has been placed on one side of the wound, it can be pulled across the wound to the opposite side (or the edges of the wound pushed together) to determine the matching point on the opposite side. The needle is then advanced into this point. The needle should enter the dermis at the same depth as it exited from the opposite side, pass through the tissue, and exit at the bottom of the wound (or the base of the flap). The edges of the wound can be closely apposed by pulling the two tails of the suture in the same direction along the axis of the wound (see Fig. 35.25C).

Some clinicians place their SQ stitch obliquely rather than vertically to facilitate knot tying. When the knot in this SQ stitch is tied, it will remain inverted, or “buried,” at the bottom of the wound. Burying the knot of the SQ stitch avoids a painful, palpable nodule beneath the epidermis and keeps the bulk of this foreign material away from the surface of the skin. Most emergency clinicians construct knots with the instrument tie technique (see Fig. 35.18). Hand and instrument knot-tying techniques are described and illustrated in wound care texts.121,122

Once the knot has been secured, pull the tails of the suture taut for cutting. Hold the scissors with the index finger on the junction of the two blades. Slide the blade of the scissors down the tail of the suture until the knot is reached. With the cutting edge of the blade tilted away from the knot, cut the tails. This technique prevents the scissors from cutting the knot itself and leaves a 3-mm tail, which protects the knot from unraveling.123 Suture the entire SQ layer in this manner.

After the SQ layer has been closed, the distance between the skin edges determines the approximate width of the scar in its final form. If this width is acceptable, insert surface sutures.124 Despite undermining and placement of a sufficient number of SQ sutures, on occasion a large gap between the edges of the wound may persist. In such cases, use a horizontal dermal stitch to bridge this gap (see Fig. 35.31).
Surfaced Closure

The epidermis and the superficial layer of dermis are sutured in a single layer with nonabsorbable synthetic suture. The choice of suture size, the number of sutures used, and the depth of suture placement depend on the amount of skin tension remaining after SQ closure.

If the edges of the wound are apposed after closure of the deeper layers, small 5-0 or 6-0 sutures can be used simply to match the epithelium on each side. Wounds with greater tension and separation should have skin stitches placed closer to each other and closer to the edge of the wound with consideration for an interrupted vertical mattress suture; layered closure is important in such wounds. If the edges of the wound remain separated, or if SQ stitches were not used, a larger-size suture placed deeply may be required.

The number of sutures used in closing any wound will vary with the wound's location, the amount of tension on the wound, and the degree of accuracy required by the clinician and patient. For example, sutures on the face are often placed between 1 and 3 mm apart. Unless the edges of the wound are uneven, sutures should be placed in a mirror-image fashion such that the depth and width are the same on both sides of the wound.

In general, the distance between each suture should be approximately equal to the distance from the exit of the stitch to the edge of the wound.

Skin closure may be accomplished by placing the appropriate number of sutures in segments (Fig. 35.26). When suturing the skin, right-handed operators should pass the needle from the right side of the wound to the left. The needle should be driven through tissue by flexing the wrist and supinating the forearm; the course taken by the needle should result in a curve identical to the curvature of the needle itself (see Fig. 35.26, steps 1 and 2, and Fig. 35.27). The angle of exit for the needle should be the same as its angle of entrance so that an identical volume of tissue is contained within the stitch on each side of the wound (see Fig. 35.26, steps 3 and 4).

Complications

Sutures act as foreign bodies in a wound, and any stitch may damage a blood vessel or strangulate tissue. Therefore, the clinician should use the smallest size and the least number of sutures that will adequately close the wound. However, if spaced too widely, surface stitches will leave a “crosshatch” pattern of marks.

Encompassing too much tissue with a small needle is a common error. Forcefully pushing or twisting the needle in an effort to bring the point out of the tissue may bend or break...
Simple Interrupted Sutures

1. Hold the needle pointing downward by excessively pronating the wrist so that the needle tip initially moves farther from the laceration as the needle penetrates deeper into the skin. Drive the needle tip downward and away from the cut edge into the subcutaneous layer.

2. Advance the needle into the laceration. The needle tip is directed toward the opposite side at the same level by rolling the needle holder. The arc of the needle pathway is controlled by retracting the skin edge. This method incorporates more tissue within the stitch in the deeper layers of the wound than at the surface. As an alternative, if a small needle is used in thick skin or the distance across the wound is great, the needle can be removed from the first side, remounted on the needle holder, and advanced to the opposite side.

3. Advance the needle upward toward the surface so that it exits at the same distance from the wound edge as on the contralateral side of the wound. Grasp the needle behind the tip and roll it out in the arc of the needle.

4. The final position, with more tissue in the depth than in the surface. The distance from each exit of the suture to the laceration is half the depth of the dermis.

5. Close the surface of the wound in segments rather than from one end. Place the first suture in the center of the wound for a straight suture line.

6. A, Too few stitches used. Note the gaping between the sutures. B, Too many stitches used. C, Correct number of stitches used for a wound under an average amount of tension.

Figure 35.26 Simple interrupted sutures. See also Figs. 35.17 and 35.18. (Steps 1–4, From Kaplan EN, Hentz VR: Emergency management of skin and soft tissue wounds: an illustrated guide, Boston, 1984, Little, Brown, p 86. Reproduced by permission.)
the body of the needle. Using a needle of improper size will defeat the best suturing technique.

If sutures are tied too tightly around the edges of the wound or if individual stitches are under excessive tension, blood supply to the wound may be compromised, thereby increasing the chance of infection. Suture marks may form even after 24 hours.78,125

If the techniques described are applied to most wounds, the edges will be matched precisely in all three dimensions, with the least number of sutures required to appose the edges and relieve tension, while avoiding excessive scarring.

Eversion Techniques

If the edges of a wound invert or if one edge rolls under the opposite side, a poorly formed, deep, noticeable scar will result. Excessive eversion that exposes the dermis on both sides will also result in a larger scar than if the edges of the skin are perfectly apposed, but eversion produces a more visible scar than eversion does. Because most scars undergo some flattening with contraction, optimal results are achieved when the epidermis is slightly everted without excessive suture tension (Fig. 35.28). Wounds over mobile surfaces, such as the extensor surfaces of joints, should be everted. In time, the scar will be flattened by the dynamic forces acting in the area.

Numerous techniques can be used to avoid inversion of the edges of the wound. If the clinician angles the needle obliquely away from the laceration, a surface stitch can be placed so that it is deeper than it is wide116 and the stitch encircles more tissue in the SQ layer than at the surface. In other words, the stitch is wider at its deepest part and narrower at the surface. If this “bottle-shaped stitch” is intended to produce some eversion of the wound edges, the stitch must include a sufficient amount of SQ tissue (see Fig. 35.26, step 4).

To accomplish eversion, lift and turn the edge of the wound outward with a skin hook or fine-toothed forceps before inserting the needle on each side (Fig. 35.29, plate 1). Eversion can also be achieved simply by pressing on the skin adjacent to the wound with closed forceps (see Fig. 35.29, plate 2).

Vertical mattress sutures are particularly effective in evert ing the edges of the wound, and they can be used exclusively or alternated with simple interrupted sutures (see Fig. 35.29, plate 3).126 In wounds that have been undermined, an SQ stitch placed at the base of the flap on each side can by itself evert the wound (see Fig. 35.29, plate 4).

Interrupted Stitch

The simple interrupted stitch (Video 35.5) is the most frequently used technique for closure of skin. It consists of separate loops of suture tied individually. Although tying plus cutting each stitch is time-consuming, the advantage of this method is that if one stitch in the closure fails, the remaining stitches continue to hold the wound together (see Fig. 35.26).

Continuous Stitch

A continuous stitch is an effective method for closing relatively clean, low-risk wounds that are under little or no tension and are on flat, immobile skin surfaces. In a continuous, or “running,” stitch, the loops are the exposed portions of a helical coil tied at each end of the wound. A continuous suture line can be placed more rapidly than a series of interrupted stitches. A continuous stitch has the additional advantages of strength (with tension being evenly distributed along its entire length), fewer knots (which are the weak points of stitches), and more effective hemostasis. This stitch will accommodate mild wound swelling. The continuous technique is useful as an epithelial or surface stitch in cosmetic closures; however, if the underlying SQ layer is not stabilized in a separate closure, a continuous surface stitch tends to invert the edges of the wound.

The continuous suture technique has some disadvantages. This technique should not be used to close wounds overlying joints. If a loop breaks at one point, the entire stitch may unravel. Likewise, if infection develops and the incision must be opened at one point, cutting a single loop may allow the entire wound to fall open. A simple continuous stitch has a tendency to produce suture marks if used for the closure of large wounds and if left in place for more than 5 days.114 However, if all tension on the wound can be removed with SQ sutures, stitch marks are seldom a problem.

Among the variations of the continuous technique, the simple continuous stitch is the most useful to emergency clinicians (Fig. 35.30a). Place an interrupted stitch at one end of the wound and cut only the free tail of the suture. Next, encircle the tissue...
Sutures. Some sutured wounds are covered by plaster casts. On occasion, the patient (child or adult) is likely to be as frightened by and uncooperative with suture removal as they were for suture placement. Surface sutures are more likely to produce stitch marks in children because the wounds are under greater tension than those in adults. The continuous subcuticular (or dermal) suture technique is ideal for these situations; the wound can be closed with an absorbable subcuticular stitch, thereby obviating the need for later suture removal. In patients prone to keloid formation, the subcuticular technique can be used in lieu of surface stitches to avoid disfiguring stitch marks. Buried, absorbable subcuticular stitches do not appear to provoke more inflammation than percutaneous running stitches with monofilament nylon.113 Because stitch marks are not a problem, a nonabsorbable subcuticular suture can be left in place for a longer period than a surface suture can.

Continuous Subcuticular Stitch

Nonabsorbable sutures used for surface closure outlast their usefulness and must be removed. On occasion, wounds require an extended period of support, longer than that provided by surface stitches. Some patients with wounds that require skin closure are unlikely or unwilling to return for removal of the sutures. Some sutured wounds are covered by plaster casts. On occasion, the patient (child or adult) is likely to be as frightened by and uncooperative with suture removal as they were for suture placement. Surface sutures are more likely to produce stitch marks in children because the wounds are under greater tension than those in adults. The continuous subcuticular (or dermal) suture technique is ideal for these situations; the wound can be closed with an absorbable subcuticular stitch, thereby obviating the need for later suture removal. In patients prone to keloid formation, the subcuticular technique can be used in lieu of surface stitches to avoid disfiguring stitch marks. Buried, absorbable subcuticular stitches do not appear to provoke more inflammation than percutaneous running stitches with monofilament nylon.113 Because stitch marks are not a problem, a nonabsorbable subcuticular suture can be left in place for a longer period than a surface suture can.
Although this technique is commonly used for cosmetic closures, closure of the subcuticular layer alone may not alter the width of the scar. This technique does not allow perfect approximation of the vertical height of the two edges of a wound, and in cosmetic closures it is often followed by a surface stitch. The subcuticular stitch requires a 4-0 or 5-0 suture made of either absorbable material or nonabsorbable synthetic monofilament. An absorbable suture can be buried within the wound, whereas a nonabsorbable suture is used for a “pullout” stitch. The absorbable synthetic monofilament suture PDS (Ethicon) is designed for subcuticular closure. It passes through tissues as easily as nonabsorbable monofilament sutures do and is absorbed if left in the wound.

Before beginning the subcuticular stitch, approximate the SQ layer with interrupted sutures to minimize tension on the wound. Start the pullout subcuticular stitch at the skin surface approximately 1 to 2 cm away from one end of the wound. Insert the needle and exit the dermis at the apices of the wound (see Fig. 35.30C, plate 1). Take bites through tissue in a horizontal direction, with the needle penetrating the dermis 1 to 2 mm from the skin surface. Make the intradermal bites small, of equal size, and at the same level on each side of the

---

**Continuous Sutures**

1. Place a suture at one end of the laceration in an analogous fashion to a simple interrupted stitch. However, cut only the distal end of the suture while leaving the needle end attached.

2. Cross over the wound at a 45-degree angle, and reenter the wound parallel to the first pass. Do not tie the suture or cut the ends.

3. Advance the needle back to the original side and exit the skin at an equal distance from the previous suture pass.

4. Continue in this fashion until the wound edges are closed and the end of the wound is approached.

5. On the last pass, leave a loop of suture. Use this loop as a free end to tie.

6. Tie the loop to the needle end of the suture with an instrument tie, and then cut the knot.

**Figure 35.30** A, Continuous sutures.  
Continued
of the suture to each other over a dressing. The subcuticular stitch will become lax as tissue swelling subsides 48 hours after wound closure. Tighten the stitch at this time.

Subcuticular closure can be accomplished with absorbable sutures that do not penetrate the skin. Begin closure with a dermal or SQ suture placed at one end of the wound and secure it with a knot. After placement of the continuous subcuticular stitch from apex to apex, pull the suture taut, and tie a knot with a tail and a loop of suture (see Fig. 35.30C, plate 3). Bury the final knot by inserting the needle into deeper tissue and exit with the needle several millimeters from the edge of the wound. The advantage of this technique is that there are no suture marks on the skin.

Nonabsorbable subcuticular sutures can be left in place for 2 to 3 weeks, thus providing a longer period of support than with surface sutures and without the problem of stitch marks. If skin sutures are used in conjunction with a subcuticular stitch, remove the surface stitches in 3 to 4 days. A subcuticular closure in itself is stronger than a tape closure. The primary disadvantage of the subcuticular stitch is that it is time-consuming, especially when supporting surface stitches are used. Another, faster method that avoids penetrating

---

**Continuous Locked Sutures**

1. Begin the repair as you would a regular continuous stitch (Fig. 35.30A, steps 1 and 2). As the needle exits the near side, “lock” the suture by passing through the circle of the previous suture loop.

2. As the suture is pulled through the circle (arrow), it will become locked.

3. Keep tension evenly distributed across the suture as it is pulled down to the skin. The lock should be parallel to the wound edge (arrow).

4. Continue in this fashion by locking each throw along the way. On the last pass, leave a loop of suture and use it to tie to the needle end of the suture.

---

Figure 35.30, cont’d B, Continuous locked sutures.
Methods of Wound Closure

The horizontal mattress stitch (Videos 35.7 and 35.8) is an SQ stitch that is oriented 90 degrees to the interrupted SQ stitch, described previously (see Fig. 35.32B). The horizontal mattress stitch apposes the skin edges closely while providing may be used to take a deep bite of skin, thereby eliminating the need for layered closure in areas where excessive tension does not result. If the superficial loop is placed first, the tails can be pulled upward while the deep loop is placed; this technique ensures eversion of the wound in less time than needed with the traditional technique.

The skin is the interrupted subcuticular stitch (Fig. 35.31). Wounds with strong static skin tension may benefit from a few interrupted dermal stitches placed horizontal to the skin surface instead of a continuous subcuticular stitch.

Mattress Stitch

The various types of mattress stitches of use to the emergency clinician are interrupted stitches. The vertical mattress stitch (Video 35.6) is an effective method of everting skin edges (Fig. 35.32A; see also Fig. 35.29, plate 3). A vertical mattress stitch
some degree of eversion.\textsuperscript{114} A horizontal mattress suture may be ideal for areas where eversion is desirable but there is little SQ tissue.

The \textit{half-buried horizontal mattress stitch} is particularly useful in suturing the easily damaged apex of a V-shaped flap (see Fig. 35.32\textit{B}). In performing the “corner stitch” (Video 35.9), penetrate the skin at a point beyond the apex of the wound with the suture needle and exit through the dermis. Elevate the corner of the flap, and pass the suture through the dermis of the flap. Then, place the needle in the dermis at the base of the wound and return it to the surface of the skin. Place all dermal bites at the same level. Tie the suture with sufficient tension to pull the flap snugly into the corner without blanching the flap.\textsuperscript{114,130} The tip of a large flap with questionable

---

**Figure 35.31** Interrupted subcuticular stitch (also called a horizontal dermal stitch). Absorbable sutures are used. (A vertical suture also closes the deep tissue.)

---

**Vertical Mattress Suture**

1. Pass the needle through both sides of the wound in the same manner as the first pass in a simple interrupted suture. This “inner” pass should be very close to the wound edge.

2. Reinsert the needle vertical to the previous exit site at a greater distance from the wound edge.

3. Pass the needle deeper than the previous pass, and exit about 0.5 cm vertical to the previous entrance site. Pull the suture through so that 2 cm of the short end remains outside the skin.

4. Tie the ends of the suture with an instrument tie, while everting the edges of the wound. Place subsequent sutures until the wound edges are apposed.

---

**Steps in the vertical mattress stitch.** The key to close apposition and exact alignment of edges is to place the inner sutures very close to the suture line (wound edge). Note: in steps 1–4 at the top of this figure the inner pass was placed first, whereas in steps a–e at the bottom the outer pass was placed first. Either method is acceptable.

---

**Figure 35.32** A, Vertical mattress sutures.
viability may be further jeopardized by postoperative swelling, so a cotton stent can be placed underneath the knot of the corner stitch. The cotton absorbs the tension produced by swelling.

The only disadvantage of the horizontal and vertical mattress stitches is that they cause more ischemia and necrosis inside their loops than do either simple or continuous stitches. With mattress sutures, alternating the side of the wound margin on which the knot is tied has been found to reduce wound tension and may reduce the risk of edge necrosis.

The invertible horizontal mattress stitch is a variation of the horizontal mattress stitch (Fig. 35.33). This stitch produces wound inversion and should only be used when wound inversion is desired, such as when re-creating the alar crease of the nose or the rolled helical rim of the ear. In most other wounds, inversion will create a depressed and more noticeable scar.
Figure 35.33 Inverting horizontal mattress sutures. 

A. Traditional everting horizontal mattress stitch and inverting horizontal mattress stitch. B. Application of the inverting horizontal mattress suture to the helix of the ear and final cosmetic result with re-creation of the helical rim. 

Methods of Wound Closure

CHAPTER 35

Figure-of-Eight Sutures

Figure-of-eight stitch, two methods. These stitches are useful in wounds with friable tissue or in areas where buried sutures are undesirable. This stitch reduces the amount of tension placed on the tissue by the suture.

Vertical figure-of-eight suture technique. It can be used to close parallel lacerations.

Figure 35.34 Figure-of-eight sutures. (Plate 1, Modified from Dushoff IM: About face, Emerg Med 6:11, 1974. Reproduced by permission; Plate 2, from Mitchell GC: Repair of parallel lacerations (letter), Ann Emerg Med 16:924, 1987.)

Figure 35.35 Technique for closure of parallel lacerations in which the central tissue island has an intact base. (Redrawn from Samo DG: A technique for parallel lacerations, Ann Emerg Med 17:297, 1988.)

Figure-of-Eight Stitch

A figure-of-eight stitch is useful for wounds with friable tissue, on the eyelids where the skin is too thin for buried sutures, or in areas in which buried sutures are undesirable (Fig. 35.34, plate 1). This stitch reduces the amount of tension placed on the tissue by the suture, which allows the stitch to stay in place when a simple stitch would tear through the tissue. The disadvantage of this technique is that more suture material is left in the wound. A vertical variation of the figure-of-eight stitch is sometimes used to approximate close, parallel lacerations (see Fig. 35.34, plate 2). Another technique involves a vertical mattress stitch (Fig. 35.35).

Correction of Dog-Ears

When the edges of a wound are not precisely aligned horizontally, there will be excess tissue on one or both ends. This small flap of excess skin that bunches up at the end of a sutured wound is commonly called a dog-ear. This effect also occurs when one side of the wound is more elliptical than the opposite side or when excision of a wound is not sufficiently elliptical because it is either too straight or too nearly circular.

If a dog-ear is present, it can be eliminated on one side of the wound in the following manner. Elevate the flap of excess skin with forceps or a skin hook. Make an incision at an oblique angle from the apex of the wound toward the side with the excess skin. Undermine the flap and lay it flat. Trim the resulting triangle of skin, and complete the closure (Fig. 35.36, plate 1). An alternative method consists of carrying the incision directly from the apex in line with the wound. Pull the flap of excess tissue over the incision while using skin hooks to retract the extended apex of the wound. Excise excess tissue, and suture the remainder of the wound. If dog-ears are present on both sides of one end of the wound, excise the bulge of excess tissue in an elliptical fashion and close the wound (see Fig. 35.36, plate 2).

Stellate Lacerations

Repair of a stellate laceration is a challenging problem. Usually a result of compression and shear forces, these injuries contain large amounts of partially devitalized tissue. The surrounding
Correction of Dog-Ears

Correction of a dog-ear. Excision of bilateral dog-ears.

Management of Stellate Lacerations

View from above a stellate laceration showing closure with half-buried mattress stitches. For some stellate lacerations it is best to cover with Steri-Strips and revise the scar later or, if small, excise the laceration and convert it to a linear repair.

Left, significant soft tissue contusion with stellate lacerations. Right, the result at 1 year. No tissue was débrided; instead, meticulous attention was paid to accurate soft tissue realignment with fine suture (7-0).

Correction of Dog-Ears

1. a) Excess tissue; b) Line to be incised; c) Incision extended; d) Pull tight; e) Triangle to be excised.

2. a) Ellipse to be excised; b) Cut along edge; c) New wound edge; d) Forceps; e) Final suture line.

Management of Stellate Lacerations

1. View from above a stellate laceration showing closure with half-buried mattress stitches. For some stellate lacerations it is best to cover with Steri-Strips and revise the scar later or, if small, excise the laceration and convert it to a linear repair.

Figure 35.36 Correction of a dog-ear. The critical maneuver is shown, where the skin is pulled tight to align the starred areas and identify the triangular piece of excess tissue to be excised (1d). Note that a straight laceration becomes somewhat curved with this technique. (From Dushoff IM: A stitch in time, Emerg Med 5:1, 1973. Reproduced by permission.)

Figure 35.37 Management of stellate lacerations.

soft tissue is often swollen and contused. Much of this contused tissue cannot be débrided without creating a large tissue defect. Sometimes tissue is lost, yet the amount is not apparent until key sutures are placed. In repairing what often resembles a jigsaw puzzle, the clinician can remove small flaps of necrotic tissue with iris scissors; large, viable flaps can be repositioned in their beds and carefully secured with half-buried mattress stitches. If interrupted stitches are used to approximate a thin flap, take small bites in the flap and larger, deeper bites in the base of the wound. A modification of the corner stitch can be used to approximate multiple flaps to a base (Fig. 35.37).

Thin flaps of tissue in a stellate laceration with beveled edges
may be more easily repositioned and stabilized with a firm dressing.\textsuperscript{114}

Closure of stellate lacerations cannot always be accomplished immediately, especially if considerable soft tissue swelling is present. It may be best in some instances to consider delayed closure or revision of the scar at a later date. In complicated lacerations, inexact tissue approximation may be all that is possible initially. For small stellate lacerations, it may be possible to excise the lesion totally and turn it into a linear repair.

**Repair of Special Structures**

**Facial Wounds (General Features)**

The ideal result in the repair of a facial laceration is an extremely narrow, flat, and inapparent scar. Facial and forehead lacerations that follow natural skin creases or lines will heal with a less noticeable scar than those that are oblique or perpendicular to the natural wrinkles of the skin (Fig. 35.38). In addition to basic wound management, a few additional techniques can be used to achieve satisfactory cosmetic results.

Although necrosis of partially devitalized wound edges contributes to wide scars, facial skin with apparently marginal circulation may survive because of excellent vascularity. SQ fat, which in other locations may be débrided thoroughly, should be preserved if possible in facial wounds to prevent eventual sinking of the scar and to preserve normal facial contours. Therefore débridement of most facial wounds should be conservative\textsuperscript{117} (Fig. 35.39). Carefully avoid underlying structures when débriding facial wounds. For example, the cheek contains both the seventh cranial nerve and the parotid gland. Clear drainage of the wound, and that side is higher than the opposite side, take

A layered closure has long been considered essential in the cosmetic repair of many facial wounds. However, the importance of layered closure in facial wounds was called into question by Singer and associates.\textsuperscript{138} These investigators found similar cosmetic outcomes and scar widths in small facial wounds less than 3 cm in length and 10 mm in width that were repaired with and without deep dermal sutures. Further confirmation of these results is needed.

If a layered closure is undertaken, approximate the dermis with an SQ stitch or with a combination of SQ and subcuticular stitches to bring the edges of the wound together or to within 1 to 2 mm of apposition. This may be close enough that the use of additional sutures seems almost unnecessary.\textsuperscript{124} If an SQ stitch is the only stitch used to close the deeper layers, pass the needle through the dermal-epidermal junction, or within 1 to 2 mm of the surface of the skin, without causing a dimpling effect. Tie the stitch snugly by pulling the two ends of the suture in the same direction (see Fig. 35.25C).

In cosmetic areas, a surface stitch should not be used to relieve a wound of significant tension. A surface stitch on the face is most appropriately used to match the epidermal surfaces precisely along the length of the wound. If the edges of the wound are separated more than approximately 2 to 4 mm after closure of the SQ layer, use a 5-0 or 6-0 subcuticular suture to eliminate the tension produced by this separation and to provide prolonged stability. Alternatively, use a few guide stitches to hold sections of the wound together before definitive closure with surface stitches. Guide stitches allow the surface stitches to be placed with little tension on each individual stitch; they match irregular edges; and they protect the SQ stitches from disruption. Place the first guide stitch at the midpoint of the wound, and subsequent guide stitches to bisect the intervening spaces. Once the definitive surface stitches have been placed, the guide stitches, if slack, can be removed. Because a needle damages tissue with each passage through the skin, use guide stitches only when necessary.

In a straight laceration, achieve better apposition during surface closure by stretching it lengthwise with finger traction or with skin hooks. When the needle is placed on one side of the wound, and that side is higher than the opposite side, take a shallow bite. Use the needle to depress the edges of the wound to the proper height, and allow the needle to “follow through” to the other side and pin the two sides together. If the first side entered is lower, elevate the needle when entering the second side to match the epithelial edges.

If the skin edges are apposed closely by the SQ stitch or a subcuticular stitch, a small, shallow epithelial stitch can be used in lieu of the standard, deeper, surface stitch to correct discrepancies in vertical alignment (Fig. 35.40).\textsuperscript{136} Achieve precise alignment of wound edges by inserting the needle as close to the edge as possible without tearing through the tissue. A 6-0 synthetic nonabsorbable suture is an excellent material for this stitch. A continuous stitch is preferable because it can be placed quickly, but interrupted stitches are acceptable. Space epithelial stitches no more than 2 to 3 mm apart and encompass no more than 2 to 4 mm of tissue.\textsuperscript{136} Once skin closure is complete, make final adjustments in the tension on a continuous suture line before tying the end of the stitch. If any discrepancies in level persist, use interrupted sutures or tape to flatten these few irregularities. The disadvantages of epithelial stitches are that they are time-consuming and add more suture material to the wound. Discrepancies in level can often be corrected with surgical tape.

Surgical tape is useful as a secondary support to protect the surface stitch from the stress produced by normal skin movement (Fig. 35.41). Facial wounds have a tendency to swell and place excessive stretch on a surface stitch. Minimize this by applying a pressure dressing and cold compresses to the wound after closure. Surgical tape can serve to a limited extent as a

![Figure 35.38](image-url) Lacerations following natural skin lines (shown here) heal with a less noticeable scar than those that are oblique or perpendicular to natural lines (or wrinkles).
bone, adhesions might develop that in time may limit the movement of skin during facial expressions. Close the frontalis muscle fascia and adjacent fibrous tissue as a distinct layer; if left unsutured, the retracted ends of this muscle may bulge beneath the skin. If the gap in a muscle belly is later filled with scar tissue, movement of the muscle may pull on the entire scar and make it more apparent.117

A U-shaped flap laceration with a superiorly oriented base poses a difficult problem. Immediate vascular congestion and later scar contraction within the flap produce the “trapdoor” effect, with the flap becoming prominently elevated (Fig. 35.43, plates 1 and 2). To minimize this effect, approximate the bulk of the SQ tissue of the flap to a deeper level on the base side of the wound. This way the skin surfaces of the two sides are apposed at the same level (see Fig. 35.43, plate 3). Apply a firm compression dressing to eliminate dead space and hematoma formation within the wound. Despite these efforts, secondary revision is sometimes necessary.118 Frequently, swelling of the

**Forehead**

Although the forehead is actually a part of the scalp, lacerations in this region are treated as facial wounds. Vertical lacerations across the forehead are oriented 90 degrees to skin tension lines, and the resulting scars are more noticeable than those from horizontal lacerations. Midline vertical forehead lacerations may result in cosmetically acceptable scars with standard closure techniques; uncentered lacerations may benefit from S-plasty or Z-plasty techniques during the initial repair or during later revision of the scar.

Superficial lacerations may be closed with skin stitches alone, but deep forehead lacerations must be closed in layers. First, approximate significant periosteal defects before closing more superficial layers (Fig. 35.42). If skin is directly exposed to small pressure dressing. In simple, low-tension facial wounds, wound closure with surgical tape provides results that are equivalent to closure with tissue adhesive.119

**Figure 35.39** A, This woman was punched in the face, suffered a laceration of her cheek, and went to the emergency department 35 hours later. The wound was not infected, but it had contracted and was beginning to heal by granulation. Under local anesthesia, the wound was opened, irrigated, and minimally debrided, and the skin edges were trimmed. B, Using a No. 15 blade, a 1-mm-deep incision was made in the skin along the edges of the wound border. C, The incised edges were then cut away with tissue scissors. D, The wound was undermined to relieve tension on the skin. E, The wound is clean, undermined, and ready to close. F, The wound was closed with 6-0 interrupted sutures, which were removed in 5 days. No antibiotics were used, and only a small linear scar resulted.
Epithelial stitches. If the skin edges are well apposed by a subcutaneous or subcuticular stitch, epithelial stitches can be used to correct discrepancies in vertical alignment. Achieve precise alignment by inserting the needle as close to the edge as possible without tearing the tissue, no more than 2 to 3 mm apart and encompassing no more than 2 to 4 mm of tissue. This wound should be carefully evaluated for an occult globe injury.

Wound closure tape can be used to provide additional support while sutures are in place and after they are removed. This may be especially useful in cosmetic areas such as the face.

This patient suffered a full-thickness flap (partial avulsion) type of forehead laceration. The surface of the scalp is visible in the upper portion of the wound. Injuries such as these must be closed in layers. The galea aponeurosis is repaired with absorbable suture. Additional subcuticular and subcutaneous stitches were placed meticulously, which resulted in excellent apposition of the wound edges. The surface is repaired with a combination of simple interrupted and continuous epithelial sutures.

Flap resolves over a 6- to 12-month period. Because flap elevation can be quite disconcerting, forewarn the patient and family about a possible trapdoor effect.

When a forehead laceration borders the scalp and the thick scalp tissue must be sutured to thinner forehead skin, use a horizontal or vertical mattress stitch with an intradermal component (see Fig. 35.43, plates 3E and 3F). Note that even a minor forehead contusion or laceration may bleed subcutaneously and, in a few days, produce blackness around the eyes (Fig. 35.44). Forewarn patients about this.
Repair of "Trapdoor" Injuries

Elevation of a forehead flap. The trapdoor effect is a natural healing process of elliptical or round lacerations. Patients should be advised of this phenomenon.

Repair of a U-shaped flap laceration with a superiorly oriented base to minimize the trapdoor effect. A, Excision of edges. B, Undermining. C, Approximation of subcutaneous tissue on the flap to subcutaneous tissue at a deeper level on the base. B and C, When a laceration in the thin skin of the forehead borders the thicker skin of the scalp, a horizontal mattress suture with an intradermal component can enhance healing by bringing tissues to the same plane. These figures show eversion of thinner skin to obtain adequate approximation with thicker scalp tissue. D-F, Skin closure.


Figure 35.44 A, Patients should be informed that minor forehead or nasal bridge trauma can produce benign blackness around the eyes in a few days. B, “Raccoon eyes” are most often benign; however, this phenomenon can be impressive. C, Under bilateral supraorbital nerve blocks, multiple small lacerations from this windshield injury are explored with a metal instrument and good lighting to remove tiny pieces of glass. Most superficial cuts can be left alone and others sutured with 6-0 nylon suture.
So-called raccoon eyes were once thought to represent a fracture, and although often associated with fractures, this is usually a common, benign finding, although occasionally a striking one.

Windshield injuries involving the forehead can be challenging to manage (see Fig. 35.44C) because multiple superficial cuts can harbor small glass particles and the injuries do not readily lend themselves to closure with suture. Supraorbital blocks can be used to anesthetize the forehead while the clinician meticulously looks for glass in each skin defect, often feeling pieces only with forceps or a small hemostat. Some pieces of glass are best felt; others are appreciated as shining objects under a good light source.

**Eyebrow and Eyelid Lacerations**

Manage jagged lacerations through eyebrows with little, if any, débridement of untidy but viable edges. The hair shafts of the eyebrow grow at an oblique angle and vertical excision may produce a linear alopecia in the eyebrow, whereas with simple closure, the scar remains hidden within the hair.

If partial excision is unavoidable, angle the scalpel blade in a direction parallel to the axis of the hair shaft to minimize damage to the hair follicles. Align points on each side of the lacerated eyebrow precisely. Precede SQ closure with a single percutaneous stitch on each margin of the eyebrow. Use the edges of the eyebrow as landmarks for reapproximation; therefore do not shave the eyebrow because these landmarks will be lost. Shaved eyebrows grow back slowly and sometimes incompletely, and shaving them often results in more deformity than caused by the injury itself. Take care not to invert hair-bearing skin into the wound.128

The thin, flexible skin of the upper eyelid is relatively easy to suture. A soft 6-0 suture (or smaller) is recommended for closure of simple lacerations. Traumatized eyelids are susceptible to massive swelling; compression dressings and cool compresses can be used to minimize this problem.

The emergency clinician must recognize that eyelid lacerations, especially complicated ones, require the expertise of an ophthalmologist with experience in ocular plastic surgery. Lacerations that traverse the lid margin require exact realignment to avoid entropion or ectropion. Injuries penetrating the tarsal plate frequently cause damage to the globe (Fig. 35.45). A deep horizontal laceration through the upper lid that divides the thin levator palpebrae muscle or cuts its tendinous attachment to the tarsal plate can result in ptosis. In most cases an ophthalmologist should repair the injury primarily. A laceration through the portion of the upper or lower lid medial to the punctum frequently damages the lacrimal duct or the medial canthal ligament and requires specialized techniques for repair (see Fig. 35.45B). Adipose tissue seen within any periorbital laceration may be retrobulbar fat herniating through the wound, and further evaluation is required (see Fig. 35.45C). Leave the repair of lid avulsions, extensive lid lacerations with loss of tissue, and complex types of lid lacerations to ophthalmologists.

**Ear Lacerations**

The primary goals in the management of lacerations of the pinna are expedient coverage of exposed cartilage and prevention of wound hematoma (Fig. 35.46). Cartilage is an avascular tissue, and when ear cartilage is denuded of its protective, nutrient-providing skin, progressive erosive chondritis ensues. The first step in the repair of an ear injury is to trim away jagged or devitalized cartilage and skin. If the skin cannot be stretched to cover the defect, remove additional cartilage along the wound margin. Depending on the location, as much as 5 mm of cartilage can be removed without significant deformity. Approximate the cartilage with 4-0 or 5-0 absorbable sutures placed at folds or ridges in the pinna, representing major landmarks. Sutures tear through cartilage; therefore include the anterior and posterior perichondrium in the stitch. Do not apply more tension than is needed to touch the edges together.

Next, in through-and-through ear lacerations, approximate the posterior skin surface with 5-0 nonabsorbable synthetic suture. Once closure of the posterior surface is completed, approximate the convoluted anterior surface of the ear with 5-0 or 6-0 nonabsorbable synthetic suture, joining landmarks point by point (see Fig. 35.46C and D). In repair of the helical fold, use the inverting horizontal mattress stitch (see prior section in this chapter on mattress suture). On the free rim, evert the skin to avoid notching later. Take care to cover all exposed cartilage. In heavily contaminated wounds of the ear (e.g., bite wounds) that already show evidence of inflammation, debride the necrotic tissue, cover the cartilage with a loose approximation of skin, and give the patient antibiotics.134,140 Once the lacerated ear has been sutured, enclose it in a compression dressing (see Chapter 63). Complex or complicated ear lacerations generally require consultation and close follow up.

**Nose Lacerations**

Lacerations involving the margin of the nostril are complicated and should be repaired accurately to ensure that unsightly notching does not occur (Fig. 35.47). In the medial portion of the nostril and superior columella, the lower lateral cartilages are quite close to the margin and relatively superficial. If the extent of the laceration is not recognized or repaired, wound healing may cause superior retraction of the margin of the nostril. Carefully palpate nasal wounds for underlying bone injury. If bony deformity is noted, consider consulting a plastic surgeon, because the fragments may require surgical wiring. Secondary repair of complications is difficult.

In repairing superficial lacerations of the nose, reapproximation of the edges of the wound is difficult because the skin is inflexible. Even deeply placed stitches will slice through the epidermis and pull out. When the edges of the wound cannot be coapted easily before skin closure, place 6-0 absorbable sutures in the fibrofatty junction with an SQ stitch. Because it is difficult to approximate gaping wounds in this location, keep débridement to a minimum. Fortunately, the nose has a rich supply of blood and often will heal without débridement. Nasal cartilage is frequently involved in wounds of the nose, but it is seldom necessary to suture the cartilage itself.

Align the free rim of the nostril precisely to avoid unsightly notching. The inverting horizontal mattress stitch may be useful in the alar crease of the nostril to help with inversion (see prior section in this chapter on mattress suture). Many clinicians recommend early removal of stitches to avoid stitch marks, yet the oily nature of skin in this area makes it difficult to keep the wound closed with tape. A running subcuticular stitch may be preferable when repairing nasal lacerations, but simple interrupted stitches are also acceptable.141 If the wound is gaping before closure, a subcuticular stitch is recommended to provide support for a prolonged period.142 If there is significant tissue injury, consultation with a facial plastic surgeon may be warranted.
Figure 35.45 A. A laceration of the eyelid margin is a complicated repair usually performed by an ophthalmologist or plastic surgeon. The principles of repair are demonstrated here. 1, The suture is placed precisely in the plane of the meibomian glands at the eyelid margin, approximately 2 mm from the edges of the wound and 2 mm deep. This placement should provide adequate eversion of the margins. 2, Partial-thickness lamellar sutures are placed across the tarsus and tied anteriorly. 3, The anterior skin and muscle lamella are closed with fine sutures, and these are tied over the long marginal sutures to prevent them from touching the cornea. B, A method of identifying and repairing the canalicus. This repair is best left to the ophthalmologist, but the emergency clinician must recognize the potential for a canalicus injury. C, Deep laceration of the left upper lid with herniation of orbital fat. For fat to prolapse, the orbital septum (and potentially the globe itself) must have been perforated. This is a wound requiring operating room exploration and repair.

Lip and Intraoral Lacerations

Lip lacerations are cosmetically deforming injuries, but if the clinician follows a few guidelines, these lacerations usually heal satisfactorily.

The contamination of all intraoral and lip wounds is considerable, and they must be thoroughly irrigated. Regional nerve blocks are preferred over local anesthetic injection because the latter method distends tissue, distorts the anatomy of the lip, and obscures the vermilion border. Loss of less than 25% of the lip permits primary closure with little deformity. The following types of wounds require initial surgical consultation or later reconstructive surgery: loss of more than 25% of the lip, extensive lacerations directly through the commissure of the mouth, and deep scars in the vermilion of the upper lip (which can later result in a redundancy of tissue). Small puncture-type lacerations heal well only if the skin is closed and the small intraoral laceration is left open (Fig. 35.48). Such injuries commonly occur when the victim’s tooth
Methods of Wound Closure

Figure 35.46 A, Lacerations of the ear require a special repair aimed at covering cartilage and preventing hematoma formation. With this through-and-through laceration of the margin of the pinna, the cartilage is trimmed just enough to allow the skin to be approximated to cover all exposed cartilage. Sutures are not used in the cartilage itself for this laceration, but the edges are approximated by skin sutures that incorporate the perichondrium. B, The repair is easiest if the posterior pinna is sutured first. C and D, Lacerations of the helical rim that traverse the two skin surfaces and the cartilage require a three-layer repair with accurate reapproximation of the auricular cartilage, as is done for the nose, to avoid notching. The cartilage is repaired by placing 5-0 or 6-0 clear absorbable suture through the perichondrium and cartilage. The skin is repaired with 6-0 or 7-0 nonabsorbable suture. E, An ear compression dressing should be used to prevent hematoma (see Chapter 63 for discussion of anesthesia and dressing for this injury).

Figure 35.47 A, Initially benign-appearing laceration of the left nostril of a 2-year-old patient. B, However, further investigation shows a full-thickness injury with a laceration of the lower lateral cartilage. A three-layer closure with reapproximation of the cartilage was performed.
Check all lip lacerations for tooth fragments embedded in the wound. **A**, This superficial mucosal laceration produced by the teeth can be cleaned, minimally trimmed, and left open to heal. Note the broken upper teeth (arrow); fragments may be embedded in the laceration. Healing produces a whitish tissue that can be mistaken for infection. **B**, This extensive laceration of the mucosa requires a layered suture closure. **C**, Small through-and-through lacerations made by the teeth can be irrigated and closed with skin and mucosal sutures in two layers. Small defects in the mucosa may be left open. Large through-and-through injuries and lacerations of the tongue margins require sutures to achieve anatomic healing. The muscle layer should be closed separately (with absorbable sutures) to prevent hematoma formation. In general, buried sutures are better tolerated by the patient.

A poorly aligned vermilion border (arrow) distorts the lip contour. To prevent this complication, place the first stitch at the vermilion-cutaneous border to obtain proper alignment.

Small through-and-through lacerations can be thoroughly cleaned and closed in two layers (skin and mucosa). It is acceptable to leave the mucosal side open if the defect is small (see Fig. 35.48C). Close large through-and-through lacerations of the lip in three layers. With a multilayer closure, approximate the muscle layer with a 4-0 or 5-0 absorbable suture securely anchored in the fibrous tissue located anterior and posterior to the muscle. The vermilion-cutaneous junction of the lip is a critical landmark that, if divided, must be repositioned with precision. Even a 1-mm step-off is apparent and cosmetically unacceptable.

Approximate the vermilion border with a 5-0 or 6-0 nonabsorbable guide stitch before any further closure to ensure proper alignment throughout the remainder of the repair (Fig. 35.49). Close the vermilion surface of the lip and the buccal mucosa with interrupted stitches of 4-0 or 5-0 absorbable suture. Finally, close the skin with 6-0 nonabsorbable suture (Fig. 35.50). Evaluate all lacerations that penetrate the oral mucosa for the presence of tooth fragments, especially if teeth are missing.
CHAPTER 35 Methods of Wound Closure

Figure 35.50 A, This patient sustained a large through-and-through lip laceration that will require a layered closure. A submental nerve block may be used, or a local anesthetic (lidocaine with epinephrine) can be injected if it does not distort the tissue landmarks. The muscle layer should be repaired with 4-0 or 5-0 absorbable suture. B, The vermilion border (arrow) must be repositioned precisely to prevent permanent disfigurement. A 5-0 or 6-0 guide suture should be placed at the border before any other closure. Nonabsorbable monofilament sutures are then used to close the skin. C, Absorbable sutures are used to close the buccal mucosa.

Figure 35.51 A, A lacerated tongue usually heals well without sutures, but they were placed in this patient because of a large rent in the middle of the tongue. Dexon (Medtronic), Vicryl (Ethicon), or silk sutures (avoid nylon) are ideal for suturing the surface of the tongue. Bleeding is usually controlled with direct pressure and local infiltration of lidocaine with epinephrine; others require deep sutures for hemostasis. Many seemingly large central tongue lacerations (such as occur during a seizure) heal well with no suturing if the margins of the tongue are intact. B, When a forked tongue is possible or flaps are pronounced, the tongue requires anatomic repair. C, This laceration will heal well without sutures.

or chipped. Intensify the search if the patient returns with an infection of a sutured wound. Probe the wound with forceps to identify fragments not seen directly in the wound. In the setting of marked facial swelling, take a radiograph of the soft tissue to help identify embedded tooth fragments.

Tongue Lacerations
Some controversy exists regarding when to suture tongue lacerations. Small, simple, linear lacerations, especially those in the central portion of the tongue, heal quickly with minimal risk for infection. Many small tongue lacerations that occur in children or from falls or seizures do not require sutures. In general, lacerations that involve the edge or pass completely through the tongue, flap lacerations, and all lacerations bisecting the tongue need to be sutured. Small flaps on the edge of the tongue may be excised, but large flaps should be sutured. If dilute peroxide mouth rinses and a soft diet are used for a few days, healing can be rapid. Persistent bleeding from minor lacerations brings most patients to the ED, and closure with deep sutures may be necessary to prevent further bleeding.

Repair of a tongue laceration in any patient is difficult, but in an uncooperative child, the procedure may prove impossible without general anesthesia. A Denhardt-Dingman side mouth gag aids in keeping the patient’s mouth open. Anesthetize a localized area of the tongue topically by covering the area with 4% lidocaine-soaked gauze for 5 minutes. Determine the maximum safe dose of local anesthetic and avoid exposure to greater doses. Large lacerations require infiltration anesthesia (1% lidocaine with buffered epinephrine) or a lingual nerve block. If the tip of the tongue has been anesthetized, a towel clip or suture can be used to maintain protrusion of the tongue in an uncooperative patient. Further anesthesia and subsequent wound cleansing and closure are possible while an assistant applies gentle traction on the tongue.

Use size 4-0 absorbable sutures to close all three layers (inferior mucosa, muscle, and superior mucosa) in a single stitch; alternatively the stitch should include half the thickness
of the tongue, with sutures placed on the superior and inferior surfaces, as well as on the edge of the tongue. Sutures on the tongue frequently become untied. This problem can be prevented if the stitches are buried. Avoid nylon sutures because the sharp edges are uncomfortable. Closure of the lingual muscle layer with a deep absorbable suture alone may be sufficient to control bleeding and return motor function to the lacerated tongue; mucosal healing is rapid.

**Scalp Lacerations**

The scalp extends from the supraorbital ridges anteriorly to the external occipital protuberances posteriorly and blends with the temporals fascia laterally. The scalp has five anatomic layers: skin, superficial fascia, galea aponeurotica, subaponeurotic areolar connective tissue, and periosteum (Fig. 35.52). Clinically, however, the scalp may be divided into three distinct layers. The outer layer consists of the skin, superficial fascia, and galea (the aponeurosis of the frontalis and occipitalis muscles), which are firmly adherent and surgically considered as one layer. The subaponeurotic layer and the periosteum form the second and third layers. The integrity of the outer layer is maintained by inelastic, tough, fibrous septa, which keep wounds from gaping open unless all three portions have been traversed. Wounds that gape open signify a laceration extending beneath the galea layer (Fig. 35.53). The galea itself is loosely adherent to the periosteum by means of the slack areolar tissue of the subaponeurotic layer. The galea is firmly attached to the underside of the SQ fascia and is rarely identified as a distinct layer in the depths of a wound. The periosteum covers the skull. The tissue-thin periosteum is often mistakenly identified as the galea, but the periosteum is flimsy and adherent to the skull and cannot be sutured.

Several unique problems are associated with wounds of the scalp. Multiple scalp wounds that are hidden by a mat of hair are easily overlooked. Stellate lacerations are common in this region, not only because the scalp is vulnerable to blunt trauma but also because its superficial fascial layer is inelastic and firmly adherent to the skin. Stellate lacerations pose additional technical problems in closure and have a greater propensity for infection. Shear-type injuries can cause extensive separation

---

**Figure 35.52** Anatomy of the scalp. The skin, superficial fascia, and galea constitute the outer layer. Blood vessels in the fascia are the major source of the bleeding in scalp lacerations.

**Figure 35.53** A, The galea has been transected in wounds that gape open like this one, and to achieve hemostasis and obtain the best closure, the galea should be sutured. B, This nongapping wound does not include the galea and can be closed with superficial sutures.
of the superficial layers from the galeal layer. Debris and other contaminants can be deposited several centimeters from the visible laceration; therefore, careful exploration plus cleaning of scalp wounds is important. When scalp wounds are debrided, obviously devitalized tissue should be removed, but débridement should be conservative because closure of large defects on the scalp is difficult.

The presence of a rich vascular network in the superficial fascia results in profuse bleeding from scalp wounds. Severed scalp vessels tend to remain patent because the fibrous SQ fascia hinders the normal retraction of blood vessels that have been cut and allows persistent or massive hemorrhage with simple lacerations. The subgaleal layer of loose connective tissue contains emissary veins that drain through diploic vessels of the skull into the venous sinuses of the cranial hemispheres. In scalp wounds that penetrate this layer, bacteria may be carried by these vessels to the meninges and the intracranial sinuses. Thus, a scalp wound infection can result in osteomyelitis, meningitis, or a brain abscess. Closure of galeal lacerations not only ensures control of bleeding but also protects against the spread of infection.

Profuse bleeding, especially from extensive scalp lacerations, is best controlled by suturing (see also Chapter 34). Unless the vessels are large or few, ligation of multiple scalp vessels seldom provides effective hemostasis, and considerable blood loss can occur during the attempt. Ask an assistant to maintain compression around the wound while you complete closure of the wound. A simple procedure that often provides hemostasis of scalp wounds is to place a wide, tight rubber band or Penrose drain around the scalp from the forehead to the occiput. You can also control bleeding temporarily in some cases by grasping the galea and the dermis with a hemostat and everting the instrument over the edge of the skin (Fig. 35.54). The disadvantage of this technique is that tissue grasped by the hemostat may be crushed and devitalized. If the SQ tissue is also everted for a prolonged period, necrosis can occur.

If an assistant is not available to apply direct pressure, use local anesthetics containing epinephrine because this is sometimes effective in controlling persistent bleeding from small vessels in a scalp wound. If bleeding from the edge of the scalp wound is vigorous and definitive repair must be postponed while the patient is resuscitated, Raney scalp clips or a hemostat can be applied quickly to the edge of the scalp wound to control the hemorrhage.

Before wound closure, visually examine the underlying skull and palpate it for skull fractures (Fig. 35.55). In wounds that can be visualized or explored, more small skull fractures are detected with the clinician’s eyes and gloved finger than with radiographs. However, a common error is to mistake a tear in the galea or the periosteum for a fracture during palpation inside the wound. Direct visualization of the area should resolve the issue. Of particular importance are stab or puncture wounds in the scalp and forehead, such as from a nail, screwdriver, knife, or ice pick. Without a laceration to explore, such wounds may seem benign, and the patient can initially appear relatively asymptomatic, yet the skull or brain has been penetrated. When evaluating a puncture or stab wound to the head, a computed tomography scan may provide unexpected findings of a skull fracture, linear or depressed, or an underlying brain injury or early hemorrhage. In wounds that expose bone but do not penetrate the skull, prolonged exposure may leave a nidus of dead bone in which osteomyelitis could develop. If exposed bone is visibly necrosed, consider appropriate consultation with neurosurgery for partial removal of the bone with rongeurs.

Figure 35.54 To temporarily control bleeding from vessels in the fascia, the galea can be everted to compress the fascia.

Figure 35.55 A, A finger and direct vision can be used to identify skull fractures. Do not mistake a rent in the soft tissues as a fracture. B, Puncture wounds in the forehead and scalp from such objects as a knife, nail, screwdriver, or ice pick can penetrate the skull and brain, and initially the patient appears well and the wound looks benign. This patient was in a bar fight and was “stabbed” in the forehead. He had no complaints. The wound could not be fully explored but appeared to be benign. C, Computed tomography scan demonstrating a depressed skull fracture (arrows).
until active bleeding appears. Clip hair surrounding the scalp wound far enough from the edge of the wound so that suturing can proceed without entangling the hair or burying it in the wound. As stated in other sections, avoid shaving, and instead clip hair to avoid further injury. If hairs along the wound edges become embedded in the wound, they will stimulate excessive granulation tissue and delay healing. Place Vaseline or tape on stubborn hairs that persistently fall into the wound to help hold them back. Although clipping scalp hair is not popular with some patients, failure to adequately expose an area is a common cause of improper cleaning and closure of scalp wounds. Because of the extensive collateral blood supply of the scalp, most lacerations in this area heal without problem, even after delayed treatment. Nonetheless, wound cleaning must be sufficient to avoid the devastating complication of scalp infection.

With microvascular techniques, large sections of skin avulsed from the scalp can be reimplanted. Some of these same techniques are used to salvage avulsed scalp, similar to those used for amputated extremities.

Unlike most wounds involving multiple layers of tissue, scalp wounds can usually be closed with a single layer of sutures that incorporate skin, SQ fascia, and the galea. The periosteum does not need to be sutured; in fact, sutures generally do not hold in this friable, thin tissue. Separate closure of the galea introduces additional suture material into the wound and may increase the risk for infection. However, in extremely large wounds, separate closure may be necessary to provide a more secure approximation of the galea than can be obtained with a large-needle, single-layer closure. In this situation, an inverted stitch (with an absorbable 3-0 or 4-0 suture) will bury the knot beneath the galea.

In superficial wounds, approximate the skin and SQ tissue with simple interrupted or vertical mattress stitches using nonabsorbable 3-0 nylon or polypropylene suture on a large needle. Avoid smaller suture material because it tends to break when firm knots are being tied. Leave the ends of the tied scalp sutures at least 2 cm long to facilitate subsequent removal. The use of blue nylon rather than black may make suture removal easier. Absorbable sutures may provide similar cosmetic and functional outcomes in scalp laceration repair.

Sutured scalp lacerations need not be bandaged. Instruct patients that they can wash their hair in 2 hours. If bleeding persists, use an elastic bandage as a compression dressing. Place gauze sponges over the laceration to provide direct local pressure beneath the elastic bandage.

An easy method of scalp laceration repair is the hair apposition technique (HAT). HAT should not be used in deep wounds as it only apposes the superficial layers of the scalp. Advantages of this technique include cosmetic results similar to suturing, increased speed of repair, no need for clipping of hair, and no need for return visit for suture or staple removal. The HAT technique should not be used with lacerations greater than 10 cm in length, irregular wound margins, hair less than 3 cm in length, and continued bleeding. Wounds between 1 and 3 cm in length may be repaired with the modified HAT technique. Care must be taken to place only a small drop of glue onto the twist of hair (not the scalp) to prevent tissue glue from entering the wound. In the HAT technique, the hair is twisted and no

---

**Figure 35.56** Closure of scalp lacerations.
actual knot is tied in the hair. The patient should be instructed to wait two days prior to hair washing. The patient should be informed that the glue will gradually fall off within two weeks. If the drop of glue fails to fall off, it may be simply removed with tweezers or comb.

**Nail Bed Lacerations**

Injuries to the nail and nail bed (also called the nail matrix) are common problems in emergency medicine, yet controversy exists over proper management (Fig. 35.59). As a general rule, repair nail bed lacerations unless they are well approximated. An exception is a nail bed laceration that causes a simple subungual hematoma.

**Subungual Hematomas**

Subungual hematomas develop from a nail bed laceration. Some nail bed lacerations are minor and of no consequence, whereas others portend a poor outcome unless the bed is repaired. A simple subungual hematoma (even in the presence of a tuft fracture) in which the nail is firmly adherent and disruption of the surrounding tissue is minimal, is not an indication to remove the nail to search for nail bed lacerations (Fig. 35.60). Despite the presence of a nail bed laceration, a good result can be expected as long as the tissue is held in anatomic approximation by the intact fingernail. Nail trephination is discussed in Chapter 37.

Suture the nail bed if a large subungual hematoma is associated with an unstable or avulsed nail. Remove the nail completely and repair the nail bed under a bloodless field (Fig. 35.61). Always use absorbable sutures in the nail bed so they do not have to be removed. A good outcome depends on maintaining the space under the eponychium (cuticle) as the laceration heals and the new nail grows out, which is a slow process.

**Partial Nail Avulsions**

If the nail is partly avulsed (especially at the base) or loose, lift the nail to assess and potentially repair the nail bed. If the nail is intact, it is best to leave it in place if the nail bed laceration

![Figure 35.57](image)

*Figure 35.57* Once the scalp is sutured, the patient is encouraged to wash the hair at home that day to remove blood and debris. Prohibiting bathing with sutures in place in any part of the body has no scientific merit.

**Figure 35.58**

A, Hair apposition technique (HAT) for superficial scalp lacerations in which hair is greater than 3 cm in length. 1, Select 4 to 5 hair strands on each side of the laceration; separately twist each group of hair strands to form bundles. 2, Cross the bundles using artery forceps. 3, Twist the bundles around each other once; there is no need to form an actual knot. 4, Place one small drop of glue on the twist of hair (not on the scalp) to secure. Repeat, every 1–2 cm along the wound, bisecting the wound just as if using sutures. B, Modified HAT, which is useful for hair lengths of 1 cm to 3 cm. 1, On each side of the wound, a bundle of 5 to 15 hairs is grasped by a clamp and the clamps are twisted around one another. 2, Tighten the twist of hairs to appose the wound margins. 3, One small drop of tissue glue is placed on the twist of hairs. 4, Repeat this every 1 to 1.5 cm to secure the entire laceration. (A, from Ong M, Ooi S, Saw, et al: A randomized controlled trial comparing the hair apposition technique with tissue glue to standard suturing in scalp lacerations (HAT study), *Ann Emerg Med* 40:21, 2002. B, from Karaduman S, Yurukrumen A, Guryay S, et al: Modified hair apposition technique as the primary closure method for scalp lacerations, *Am J Emerg Med* 27:1050, 2009.)
can be repaired (Fig. 35.62). The method for atraumatically removing a nail is demonstrated in Fig. 35.67.

Nail Bed Repair
When the integrity of the nail bed is significantly disrupted, a rippled nail may develop (see Fig. 35.62F). Anatomic repair of the nail bed may minimize subsequent nail deformity. Approximate a simple nail bed laceration with 6-0 or 7-0 absorbable suture (to obviate the need for suture removal); either Vicryl Rapide or chromic gut may be used for nail bed repair. If available, use loupe magnification and a finger tourniquet to maintain a bloodless field (Fig. 35.63). Repair

**Figure 35.59** Anatomy of the fingernail. The fingernail rests on the nail bed, also termed the matrix. The distal end of the nail covers the sterile matrix; the proximal end arises from and covers the germinal matrix. The tissue adherent to the proximal dorsal surface of the nail is the eponychium (also termed the cuticle), and the potential space between the nail and the eponychium is the nail fold.

**Figure 35.60** A, This subungual hematoma occupies most of the nail and should be drained by simple nail trephination. The injury does not require nail removal or repair of the nail bed because the nail is still firmly attached to the matrix. Even though a nail matrix laceration (the source of the bleeding) is present, the cosmetic result will be excellent. The presence of an underlying digital tuft fracture does not change management. This hole is too small and did not evacuate all the blood. B, There are two issues with this injury. First, an inadequately sized hole was placed (too small) and did not drain the subungual hematoma. All blood should be drained in the emergency department before discharge to ensure proper treatment. Note the subtle ecchymosis under the eponychium (arrow). This can occur only if the nail base was avulsed. The base of the nail is lying just under the skin. Compare it with A, where no blood is seen in the paronychial area as the nail base remains intact. C, After the hole is enlarged and the blood is totally drained, and without removing the nail, a hemostat is used to pull the nail forward and manipulate the base back into its proper position under the eponychium.
such as Silastic sheet, sterile polypropylene foil (obtained from the sterile infusion reservoir used on IV tubing), or gauze packing for approximately 2 to 3 weeks. Reinsertion of the nail occasionally results in infection, so clean the nail carefully. After cleaning, suture the avulsed nail in place or secure it with wound closure tape or tissue glue. If only the distal portion of the nail has been avulsed, it can still be used as a temporary splint or “dressing” to protect and maintain the integrity of the underlying nail bed (see Fig. 35.63).

The figure-of-eight technique can be used to secure the nail, which minimizes additional trauma to the nail bed and

Figure 35.61 A. Classic “finger slammed in a door” with forced flexion and avulsion of the base of the nail. This large subungual hematoma can be misleading, but it is associated with blood under the cuticle (arrow) proximal to the nail, a clue that this is not a simple injury. B, When the blood is drained, the extent of the injury is more obvious. C, Careful removal of the nail exposes a laceration of the nail bed, which is sutured with absorbable suture. D, A drainage hole is placed in the nail (arrow) because it will be used as a temporary splint for the nail bed and to keep the cuticle space open to prevent scarring. E, The avulsed nail is placed under the eponychium to its base and sutured in place for 3 to 4 weeks while a new nail grows. This replaced nail may be removed, or it is pushed out by the new nail.
the eponychial fold. A tourniquet is often necessary to maintain a bloodless field during tissue adhesive application.

Complete Nail Avulsions
If the entire nail is avulsed but intact, it should be replaced after repairing the nail bed laceration for three reasons: (1) it acts as a splint or mold to maintain the normal anatomy of the nail bed, (2) it covers a sensitive area and facilitates dressing changes, and (3) it maintains the fold for new nail

---

**Figure 35.62** A, This fingernail was avulsed at the base, a common result of having a door slam on the digit. The nail is tightly adherent to the nail bed, so it is not removed but simply replaced under the eponychium to its former position. B, Sutures are placed in the nail to keep it stable. This nail may simply start to grow on its own. C, A saw-induced laceration of the fingertip with an open fracture, nail bed laceration, and skin laceration. D, Only part of the nail was removed and the nail bed repaired through the window. The skin is closely approximated. Such injuries usually heal well with attention to detail, and infection is unusual despite the open fracture. Oral antibiotic therapy for 5 to 7 days is reasonable. E, Because this avulsed nail is unstable and subungual bleeding is present, the nail can be removed and the nail bed inspected. Any large nail bed laceration should be repaired meticulously with absorbable suture (6-0). After repair of the nail bed, a drainage hole is placed in the nail, and the nail is replaced under the eponychium (cuticle) and fixed in place with sutures that incorporate the nail edge and the skin bilaterally. In 2 to 3 weeks, the new nail will begin to push out the old nail (usually growing under it while maintaining the eponychium), and the old nail is removed. The exposed nail bed will be sensitive for a while. See Fig. 35.67 for a simple technique to remove the fingernail. F, This nail is permanently deformed with ridges. Although crush injury to the nail bed is probably responsible for this deformity, repair of the nail bed might have minimized the resultant deformity.

---

eponychium (Fig. 35.64). This technique has also been used to secure sterile polypropylene foil in the event that the nail is too injured to be replaced (Fig. 35.65). Tissue adhesive is a time-saving option for repairing nail bed lacerations in both adult and pediatric patients (Fig. 35.66). Tissue glue obviates the need for suture removal and produces similar cosmetic and functional results. Tissue glue is applied on nail bed lacerations after meticulous alignment. The glue can also be used to secure the nail plate under
Approximately 6 months is required for a new nail to reach to the fingertip.

Complicated Nail Bed Injuries

If the germinal matrix of the nail bed is avulsed but intact, reimplant the nail with a 5-0 or 6-0 absorbable suture using a mattress stitch. If an open fracture exists, do not allow any part of the matrix to remain trapped in the fracture line.

If the nail bed is found to be extensively lacerated, it may be prudent to refer the patient to a hand surgeon, who can raise a flap of tissue extending from the proximal nail fold, explore the wound for foreign bodies, and clean under the nail bed. A fingertip avulsion that involves the germinal matrix or an isolated nail bed avulsion should not be allowed to heal on its own (i.e., by secondary intention). If the exposed germinal matrix is left open to granulate, it will form scar tissue that could produce a distorted and sensitive digit. If part of the

Growth. If the proximal portion of the nail is not replaced, either of two complications may result: (1) longitudinal scar bands may form between the proximal nail fold and the germinal matrix and cause a permanent split or deformity of the nail, or (2) the space between the proximal nail fold and the germinal matrix of the nail bed may be obliterated within a few days. Consequently, splints over proximal nail beds should be maintained for 2 to 3 weeks. The proximal portion of the traumatized nail often needs to be trimmed so that it will fit more easily into the nail fold. It is usually necessary to suture or glue the nail in place. A replaced nail may grow normally and act as a free graft, but it is often dislodged by a new nail. If the nail was lost or irreparably destroyed, insert a piece of nonadherent, petrolatum-impregnated gauze (such as Adaptic [Acelity, San Antonio, TX] or Xeroform [DeRoyal, Powell, TN]) between the proximal nail fold and the germinal matrix. Nails grow at a rate of 0.1 mm/day, and approximately 6 months is required for a new nail to reach to the fingertip.

Complicated Nail Bed Injuries

If the germinal matrix of the nail bed is avulsed but intact, reimplant the nail with a 5-0 or 6-0 absorbable suture using a mattress stitch. If an open fracture exists, do not allow any part of the matrix to remain trapped in the fracture line.

If the nail bed is found to be extensively lacerated, it may be prudent to refer the patient to a hand surgeon, who can raise a flap of tissue extending from the proximal nail fold, explore the wound for foreign bodies, and clean under the nail bed. A fingertip avulsion that involves the germinal matrix or an isolated nail bed avulsion should not be allowed to heal on its own (i.e., by secondary intention). If the exposed germinal matrix is left open to granulate, it will form scar tissue that could produce a distorted and sensitive digit. If part of the

This nail was removed and the nail bed repaired with absorbable suture.

Because the nail was macerated and unable to be replaced, gauze is used to maintain the eponychial space for 2 to 3 weeks. A small piece of gauze is placed with forceps to gently pack open the space to prevent scar formation.

Figure 35.63 Nail bed repair. (Plate 2, from Dushoff IM: Handling the hand, Emerg Med 8:111, 1976. Reproduced by permission.)
Drains in Sutured Wounds

Drains are used primarily to keep wounds open for drainage of existing purulence or blood that may otherwise collect in the wound. Drains do not prevent infection. When no infection exists, the use of drains in soft tissue wounds “prophylactically” is rarely required in the ED setting. Drains in uninfected wounds may wick surface bacteria into the wound and impair resistance of the wound to infection. Drains placed in experimental wounds contaminated with subinfective doses of bacteria behave as foreign bodies by enhancing the rate of infection, regardless of whether the drain is placed entirely within the wound or brought out through the wound.

If the wound is considered to be at high risk for infection, instead of suturing the wound with a drain in place (in anticipation of infection), it may be more prudent to leave the wound open and consider delayed primary closure later when the risk for infection subsides. Drains should not serve as substitutes for other methods of achieving hemostasis in traumatic wounds.

Lacerations Over Joints

Lacerations over joints may enter the joint itself or injure adjacent tendons or muscle groups. It may be difficult to determine whether the joint has been violated. A plain radiograph of a large joint such as the knee may demonstrate air within the joint, which is evidence of joint penetration. If a wound penetrates the joint cavity, open débridement may be required, but the approach varies with the joint involved.

Fingertip Amputations

Treatment of fingertip amputations has undergone evolution from complicated grafts and flaps to nonsurgical follow-up and primary healing. If bone is not...
Figure 35.65 A. Sterile infusion set reservoir used as fingernail substitute. B. Fingernail avulsion secondary to trauma. C. Fingernail substitute trimmed to fit in the nail bed and to provide the nail bed protection while healing. D. Secured fingernail substitute. E. Clinical outcome at one year. (From Tos S, Artiaco S, Coppolino S, et al: A simple sterile polypropylene fingernail substitute, Chirurgie de la main 28:144, 2009.)
Figure 35.66 Dermabond (Ethicon) use for securing the nail to the nail bed. (Courtesy Michelle Lin, MD. From Rezaie S: Trick of the trade: nail bed repair with tissue adhesive glue, *Acad Life Emerg Med* 2014. Available at: http://www.aliem.com/trick-trade-nail-bed-repair-tissue-adhesive-glue/. Accessed 2016.)

**Nail Removal**

1. To remove a fingernail or toenail atraumatically, the blades of iris scissors are held parallel to the nail bed to avoid lacerating the matrix. A digital block is necessary. The closed blades are slowly advanced in the plane between the nail and the nail bed and then gently spread to loosen the nail. The scissors are advanced and spread in stages until the base of the nail is reached and the entire nail is loose. The nail is grasped with a hemostat and pulled from the base.

2. Once the nail is removed, repair any nail bed lacerations with absorbable suture.

3. Replace the nail into the eponychium.

4. Replace the nail into the eponychial fold.

**Figure 35.68** A, This laceration looks benign but may involve the knee joint or quadriceps tendon. A radiograph to look for air in the joint or a saline arthrogram (see Chapter 53) should be performed. B, Air in the joint space (arrows) on a plain radiograph proved joint space violation.

**Figure 35.67** Nail removal.
Figure 35.69 A and B, Guillotine amputation of the fingertip through the distal end of the nail (arrow). The distal phalanx is not involved. Approaches vary widely, and referral can be made in a few days if not immediately available. C, Although it is tempting to replace the amputated tip, many would allow this wound to heal spontaneously, albeit slowly (8 to 12 weeks), and not perform skin grafts or flaps. D, The amputated nail is placed under the eponychium and sutured into place. A rounded but shortened tip with sensation can be expected. Periodic débridement and use of the nail as a splint resulted in a good final appearance. E and F, In only 4 weeks, this amputation has nearly healed to the original length and contour with only dressing changes and minimal débridement. Note that a new nail is growing (arrow) and the replaced nail has been removed.

Figure 35.70 A and B, This macerated tip has islands of skin left and good tissue volume and will do quite well with sutures to restore the basic anatomy. Healing will take 4 to 5 weeks.
involved, a good result can be expected with attentive wound care, occasionally minimal debridement, and protective dressing changes. It may take 6 to 12 weeks for healing to occur, but acceptable length, function, and sensation can be expected. A motivated patient and follow-up with a knowledgeable specialist are required. There are no data or standard that mandates long-term antibiotics for such injuries, but it is reasonable to provide gram-positive antibiotic coverage for 7 to 10 days.

**DISTALLY AND PROXIMALLY BASED FLAP LACERATIONS**

Minor flap lacerations can be managed primarily in the ED. Large flaps, such as scalping lacerations, are best handled by a consultant. Both proximally and distally based flaps are not always simple lacerations. The major problem is vascularity: venous drainage at the end of a distally-based flap and the potential for ischemic necrosis in any flap configuration. The body has the ability to augment the circulation in some flaps (Fig. 35.71A). The length-to-width ratio of the flap is not the main variable in survival of the flap (see Fig. 35.71B). Perfusion pressure is the most critical factor. Intravascular perfusion pressure decreases along the length of the flap. If intravascular perfusion pressure is less than interstitial pressure at the distal end of the flap, capillaries will collapse (the critical closing pressure). As edema is generated by ischemia and inflammation, interstitial pressure increases, which decreases chances for tissue survival.

Many proximally and distally based flaps (see Fig. 35.71C) may be closed in the ED and monitored on an outpatient basis. Healing of a distally based flap is hampered by loss of venous and lymphatic drainage and by subsequent edema of the flap, which decreases capillary flow. Warn the patient about the possibility of flap necrosis and the need for revision or even skin grafting at a later date. Partial flap survival is better than total loss of the flap, so flaps with questionable viability should not be removed. Flaps are similar to free skin grafts, and the keys to a more successful outcome include undermining the flap to relieve tension, limiting buildup of fluid under the
flap with a compression dressing, and decreasing movement of the flap as it heals. Cautious minor defatting may be performed on the underside of the flap. Tissue adhesive instead of multiple sutures can be helpful.

Self-inflicted wounds with a vague or inaccurate history may be encountered in the ED. Characteristic self-injury patterns are depicted in Fig. 35.72. These patterns of injury should be recognized and appropriate psychiatric follow-up provided.
REFERENCES


Foreign Body Removal

Daniel B. Stone and David J. Scordino

CHAPTER 36

EVALUATION AND DIAGNOSIS

A foreign body (FB) is any substance that is not naturally part of the body. These cases are common in the clinical setting. An FB should be suspected whenever the skin is broken. A thorough history and physical examination are essential to assess the risk for an FB. During assessment, FBs may not be obvious especially when the wound may appear closed, but they should be considered whenever the history is particularly concerning. For example, a patient with an apparent “sprained foot” who was walking without shoes and experienced a sharp, sudden pain in the foot may have a needle, toothpick, or any other similar type of FB (Fig. 36.1). Certain mechanisms of injury, such as puncturing or kicking out a window or stepping on an unknown object while walking in a field or stream, are often associated with a retained FB. Mechanisms that make FBs less likely include lacerations from metal objects; however, if considerable force was applied during the injury and the object is not available for inspection, radiographic imaging may be warranted because bone may have been encountered and a small section might have splintered off the offending object. The history should also include whether the patient perceives or suspects an FB. Steele and associates found that the negative predictive value of patient perception was 89% but that the positive predictive value was just 31%. Importantly, the patient’s past medical history should be explored for allergies to local anesthetics, bleeding diatheses, diabetes mellitus, vascular disease, uremia, immunocompromised state, or other diseases that would affect wound healing or management.

Before the physical examination, be sure to have enough examination time, as well as proper space and equipment. It is important to confirm that the patient is willing to undergo the procedure because cooperation is essential to optimize success. Attempts at removing an FB in an overly uncooperative patient, such as one who is intoxicated, drugged, cognitively limited, or confused, are potentially injurious to both the examiner and the patient. In most noncritical situations, if a patient is uncooperative, perform the examination again when the patient is able to cooperate. In benign cases, follow-up can occur a few days later.

During the physical examination, carefully palpate the periphery of all wounds to elicit tenderness because retained FBs often produce pain during palpation. In addition, some FBs can migrate away from the wound or perceived entry point. Some superficial FBs may be palpated through the skin, but surprisingly large FBs may be found in seemingly minor wounds without much external evidence. In some cases, the external characteristics of the wound do not yield firm evidence regarding the presence or absence of an FB. Deeper FBs may not be palpable and must be localized by other techniques.

Exploration is an important part of the bedside evaluation, whether done initially or after further imaging studies (Fig. 36.2). This requires adequate pain control, good lighting, proper equipment, a bloodless field, a cooperative patient, and appropriate positioning to visualize as much of the wound as possible (Figs. 36.3 and 36.4). A metal probe may help identify the FB by feel or sound. Glass, as an example, is difficult to identify by sight in soft tissue, but touching it with metal causes a characteristic grating sound. Probing a wound with a gloved finger to locate or identify an FB is strongly discouraged because of the risk of the FB penetrating the glove and exposing the clinician to infection such as human immunodeficiency virus (HIV) and hepatitis (Fig. 36.5). Alternatively, some authors have suggested injecting the entrance wound with methylene blue to outline the track of the FB. The blue line of injected dye is followed into the deeper tissues. This technique is of limited value because the track of the FB often closes tightly and does not allow passage of the methylene blue.

Augmenting the Physical Examination: Imaging Techniques

Many emergency clinicians mistakenly believe that in the absence of adipose tissue, if the base of the wound can be clearly visualized and explored, an FB can always be ruled out. Orlinsky and Bright found the reliability of exploration to be related to the depth of the wound. In their study, only 2 of 133 superficial wounds, deemed adequately explored, had an FB on plain films, but 10 of 130 wounds had FBs beyond the subcutaneous fat despite negative explorations. Anderson and coworkers reported that 37.5% of the foreign bodies were initially missed by the treating physicians. In many of these cases, a radiograph of the injured area was not taken. Avner and Baker detected glass with routine radiographs in 11 of 160 wounds (6.9%) that were inspected and believed by the clinician to be free of glass. Clinicians evaluating for FBs should maintain a low threshold for ordering or performing imaging studies. Options for imaging include plain radiographs, ultrasound (US), computed tomography (CT), magnetic resonance imaging (MRI), and fluoroscopy. See Table 36.1 for a summary.

Plain Radiography

Plain radiographs are readily available, are easily interpreted, and cost significantly less than CT, formal US, or MRI. The ability of plain films to detect FBs in soft tissue depends on the object’s composition (relative density), configuration, size, and orientation (Fig. 36.6). Detection of FBs on plain films can be enhanced by requesting an underpenetrated soft tissue technique and by obtaining multiple views to prevent FBs from being obscured by superimposed bone or soft tissue folds. Plain films are often sufficient; however, digitized radiographs may be manipulated to enhance identification of a suspected FB. Indirect evidence of an FB may include trapped or surrounding air, a radiolucent filling defect, or secondary bony changes such as periosteal elevation, osteolytic or osteoblastic alterations, or pseudotumors of bone.

Metallic objects are readily visualized on radiographs. Despite a common misconception that glass must contain lead to be visualized on a plain radiograph, almost all glass objects in soft tissue (bottles, windshield glass, lightbulbs, microscope cover slips, laboratory capillary tubes) are at least somewhat radiopaque and can be detected by plain radiographs unless they are obscured by bone or are very small (< 1 mm) (Fig. 36.7). The absence of a glass FB on multiple projections is strong, though not absolute evidence that glass is not contained in a
CHAPTER 36
Foreign Body Removal

Entrance wound (Fig. 36.8). This also helps identify the path that leads to the FB and the relative distance from the surface to the FB. Needles at two angles may also be used to aid in localization (Fig. 36.9).

Liberal use of plain film radiography makes sound clinically relevant medicolegal practice. A review of 54 wound FB claims against 32 physicians from 22 institutions found glass, a radiopaque substance, to be the most common material. However, in only 35% of the cases involving glass were plain films taken. Cases with a glass FB without a radiograph ordered were associated with unsuccessful defense (60%) and higher indemnity payments.

US has become the modality of choice for imaging radiolucent FBs such as wood and thorns because most soft tissue FBs are hyperechoic on US. In addition, if in place for more than 24 hours, most FBs will be surrounded by a hypoechoic area corresponding to granulation tissue, edema, or hemorrhage, which may aid in making the diagnosis. Metal will leave a linear trail of echoes deep to the FB, referred to as the comet...
Figure 36.2 Some wound mechanisms are classic for the presence of a foreign body (FB). Note that even small pieces of all types of glass are readily seen on a plain radiograph. A, This intoxicated patient kicked out a window and sustained seemingly minor puncture wounds. He did not believe that glass was in the wound, there was little pain, and no FB could be palpated externally. B, A radiograph revealed a large shard of glass (arrow) deeply embedded in the wound. C, The shard of glass was removed, but only after 20 minutes of exploration. D, Another classic scenario for a retained glass FB is putting the arm through a window. An FB was not suspected or sensed by the patient or clinician. E, When a radiograph was taken, a large shard of glass (arrow) was readily detected. Despite its size, removal was difficult and time-consuming.

Figure 36.3 A, This patient has an obvious wooden foreign body (FB) on the dorsal aspect of the proximal part of the forearm. B, Manual removal of the object was not difficult. C, However, thorough exploration of the wound cavity via an incision over the entire length of the FB tract revealed multiple small wooden fragments. Wood fragments retained in a wound will always cause an infection. Proper exploration requires adequate analgesia, good lighting, hemostasis, and a cooperative patient. Use of a metal probe may help identify the foreign object by feel or sound (see Fig. 36.17).
A wooden object leaves an acoustic shadow without artifact.\textsuperscript{17,18} US may be performed at the bedside. To do so, use a high-frequency transducer (at least 7.5 MHz, such as a high-frequency linear vascular probe) because most FBs are small and superficial. A lower frequency may also be necessary if a deeper FB is suspected, but it may miss small FBs.\textsuperscript{19} A spacer may be needed to adjust the “focal zone” (i.e., where the beam is the narrowest and signal intensity is the highest). US may be particularly difficult in the hand or foot, which have many echogenic structures and web spaces that may limit visualization when the FB is adjacent to bone.\textsuperscript{16,18,20} Keep in mind that the presence of air, scars, calcification, sutures, and sesamoid bones in surrounding tissue may lead to false-positive results.\textsuperscript{20,21}

Advantages of US include low cost, no ionizing radiation, the ability to define the object in three dimensions, and real-time bedside imaging, which may be used during removal.\textsuperscript{9} Many studies report that US is highly sensitive for the detection of FBs by adequately trained personnel, either radiologists, technicians, or emergency physicians.\textsuperscript{1} It is difficult to cite a precise sensitivity or specificity because of the wide variation in these studies with regard to FB size, material, and location; operator experience; and the models used in these studies.\textsuperscript{21-23}

Figure 36.4 A windshield forehead injury usually harbors multiple retained pieces of glass that are difficult to find. Probing with a needle or forceps in each wound to feel or hear contact with the fragments may help find them. A supraorbital nerve block is ideal to facilitate probing.

Figure 36.5 What’s wrong here? Though historically suggested as a useful technique to find foreign bodies, probing the depths of a wound with a gloved finger may result in a puncture wound in the operator. The practice is strongly discouraged because of the prevalence of hepatitis and human immunodeficiency virus infection.

tail artifact. A wooden object leaves an acoustic shadow without artifact.\textsuperscript{17,18} US may be performed at the bedside. To do so, use a high-frequency transducer (at least 7.5 MHz, such as a high-frequency linear vascular probe) because most FBs are small and superficial. A lower frequency may also be necessary if a deeper FB is suspected, but it may miss small FBs.\textsuperscript{19} A spacer may be needed to adjust the “focal zone” (i.e., where the beam is the narrowest and signal intensity is the highest). US may be particularly difficult in the hand or foot, which have many echogenic structures and web spaces that may limit visualization when the FB is adjacent to bone.\textsuperscript{16,18,20} Keep in mind that the presence of air, scars, calcification, sutures, and sesamoid bones in surrounding tissue may lead to false-positive results.\textsuperscript{20,21}

Advantages of US include low cost, no ionizing radiation, the ability to define the object in three dimensions, and real-time bedside imaging, which may be used during removal.\textsuperscript{9} Many studies report that US is highly sensitive for the detection of FBs by adequately trained personnel, either radiologists, technicians, or emergency physicians.\textsuperscript{1} It is difficult to cite a precise sensitivity or specificity because of the wide variation in these studies with regard to FB size, material, and location; operator experience; and the models used in these studies.\textsuperscript{21-23}

Figure 36.6 Radiographic appearance of various foreign bodies. Arrows point to A, glass B, pencil lead (graphite, but not wood, is radiopaque), and C, a metallic pin. Plain radiographs should be taken in multiple projections. Do not ignore subtle soft tissue irregularities. Generally absent on radiographs are aluminum, plastic, wood, thorns and other organic objects, such as fish spines, and small pieces of gravel.

Figure 36.7 Almost all glass is visible on a plain radiograph, regardless of its lead content. Do not ignore small white specs on a radiograph. If glass is superimposed on bone, it may be missed, so multiple projections are recommended. Plain film radiography shows sharp corners characteristic of glass. Proximity of the foreign body to the bone may be deceiving.

CT

CT depends on x-ray absorption; thus, it generally visualizes the same material detected on plain films, but subtle differences in soft tissue densities may help identify FBs not seen on radiographs.\textsuperscript{7,11} In addition, CT produces a better
A reasonable initial approach for localizing nonvisualized FBs in the ED is to obtain multiple-projection plain radiographs with a soft tissue technique. This technique will visualize the majority of FBs, especially metal and glass. US should be considered for objects known to be radiolucent, such as wood or thorns. The role of CT and MRI for evaluation of FBs in the ED is limited, but they are the definitive imaging tests in confusing cases. For suspected intraorbital or intracranial FBs, CT is recommended. CT or MRI is also warranted when a previously negative explored wound exhibits recurrent infection, poor healing, or persistent pain. CT or MRI may be the appropriate initial test for patients with nonspecific swelling to define FBs and possible alternative diagnoses such as abscesses, masses, or other inflammatory processes. Bedside US and fluoroscopy (for radiopaque FBs) may be used to guide difficult FB extractions if initial attempts at removal are unsuccessful and the proper equipment and experienced personnel are available.

### TABLE 36.1 Imaging Summary

<table>
<thead>
<tr>
<th></th>
<th>Radiography</th>
<th>US</th>
<th>CT</th>
<th>MRI</th>
<th>Fluoroscopy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Materials visualized</strong></td>
<td>Metal, Glass, Graphite, Plastic (variable)</td>
<td>Wood, Thorns, Organic matter, Metal, Glass</td>
<td>Metal, Glass, Wood (late), Plastic (variable), Organic material (variable)</td>
<td>Glass, Plastic, Organic matter</td>
<td>Metal, Glass, Plastic (variable)</td>
</tr>
<tr>
<td><strong>Pros</strong></td>
<td>Cheap, Available, Easy to interpret</td>
<td>Bedside use for real-time extraction, Reliably expensive</td>
<td>3D images with some improved visualization of FBs vs. plain film</td>
<td>Improved detection of radiolucent objects vs. CT</td>
<td>Useful for real-time bedside extraction</td>
</tr>
<tr>
<td><strong>Cons</strong></td>
<td>Unable to visualize all material, May miss small FBs in areas adjacent to bone</td>
<td>Results vary based on operator experience, Difficult visualization in the hands and potential for false positives</td>
<td>Expensive, Increased exposure to ionizing radiation</td>
<td>Expensive, Not readily available in all EDs, Dangerous for metallic FBs</td>
<td>Unfamiliar technique to most ED clinicians, Not widely available</td>
</tr>
<tr>
<td><strong>Conclusions</strong></td>
<td>Reasonable initial approach, particularly for metal and glass, Obtain multiple views via a soft tissue technique</td>
<td>Recommended for radiolucent structures such as vegetative material (wood, thorns, etc.), Allows direct visualization, Can assist with operative removal, Potentially a bedside procedure depending on the clinician’s skill level</td>
<td>Recommended approach for intracranial, intraorbital FBs, Has a role for repeated visits and concerns for retained FBs, Limited role during initial evaluation</td>
<td>Role limited on initial evaluation, Has a role with repeated visits and concern for retained FBs</td>
<td>Use limited to individual clinicians with experience and availability</td>
</tr>
</tbody>
</table>

3D, Three dimensional; CT, computed tomography; ED, emergency department; FB, foreign body; MRI, magnetic resonance imaging; US, ultrasonography.

A reasonable initial approach for localizing nonvisualized FBs in the ED is to obtain multiple-projection plain radiographs with a soft tissue technique. This technique will visualize the majority of FBs, especially metal and glass. US should be considered for objects known to be radiolucent, such as wood or thorns. The role of CT and MRI for evaluation of FBs in the ED is limited, but they are the definitive imaging tests in confusing cases. For suspected intraorbital or intracranial FBs, CT is recommended. CT or MRI is also warranted when a previously negative explored wound exhibits recurrent infection, poor healing, or persistent pain. CT or MRI may be the appropriate initial test for patients with nonspecific swelling to define FBs and possible alternative diagnoses such as abscesses, masses, or other inflammatory processes. Bedside US and fluoroscopy (for radiopaque FBs) may be used to guide difficult FB extractions if initial attempts at removal are unsuccessful and the proper equipment and experienced personnel are available.

### MRI

Although MRI is expensive and not as readily available to emergency clinicians as plain films and CT are, it may be superior to CT in detecting small, nonmetallic, radiolucent FBs such as plastic, particulary in the orbit. MRI may not visualize wood, which may appear as a linear signal with associated inflammation and looks hypointense with respect to muscle on T1- and T2-weighted sequences, on which it appears as a signal void. Plastic is more easily visualized with MRI than with CT. MRI cannot be used for metallic objects and gravel, which contain various ferromagnetic particles that
Figure 36.8  A, This patient had an obvious diabetic foot abscess. B, A radiograph was taken to rule out osteomyelitis or soft tissue gas. To everyone's surprise, including the patient's, a sewing needle (arrow), origin and timing unknown, was the source of the infection. C and D, A paper clip was taped to the skin to localize the depth and position of the foreign body (FB; arrow) on the radiograph. An anteroposterior radiograph (not shown) also localized the FB. E, For orientation, the outline of the paper clip was traced on the foot. A dot marks a possible entrance site. F, A generous incision was made over the projected site of the FB. G, The needle (arrow) was felt and extracted with a hemostat. H, The wound was left open and packed with a cotton wick.
Figure 36.9 When a small entrance wound (A) is noted but the foreign body (FB) is not seen, noninvasive localization is preferable to blind probing. Metal markers taped to the skin or needles inserted close to the FB under local anesthesia (B) and radiographed at different angles provide a guide to localization and extraction of the FB. (Reproduced from Hospital Medicine, January 1981, with permission of Cahner’s Publishing Co.)

Many foreign bodies are radiopaque and can easily be seen with traditional radiography. Such foreign bodies include glass, metal, and some plastics. Other foreign materials, such as splinters, spines, and thinner plastics, are radiopaque and easily missed on radiographs. Ultrasound is an optimal modality for both identifying retained foreign bodies and aiding in their removal. A number of studies have evaluated the sensitivity and specificity of ultrasound in identifying foreign bodies in soft tissue. The findings have been variable, depending on the type and size of foreign body.1-3

General Considerations
A preliminary radiograph may be helpful in narrowing down the area to be evaluated with ultrasound, particularly when there is a large area where the foreign body may be found. Asking the patient to identify the point of maximal tenderness may also be helpful in narrowing down the overall area to be examined. Once the area has been clarified, a high-frequency (10 to 12.5 MHz) transducer should be selected. Higher frequencies will convey sufficient resolution to distinguish foreign material from normal soft tissue structures.

Each type of foreign body has specific identifiable characteristics; however, certain general findings suggest the presence of foreign material. The finding of soft tissue edema, represented by anechoic (black) or hypoechoic (dark gray) areas within the normal soft tissue, is highly suggestive of recent tissue disruption (Fig. 36.US1). The area in question should be evaluated from a number of different angles to find the object in its long axis. A small foreign body may easily be overlooked if only a small portion of it is seen.

Additionally, the use of a “stand-off” pad may be helpful, especially when dealing with superficial structures such as the hand or foot. A slim, fluid-filled structure, such as a 100-mL bag of saline or a glove filled with water, is placed over the area of interest. The transducer is then placed on top of this pad. This extra layer creates an acoustic window to allow greater resolution and eliminate some superficial artifacts that may impede the examination.

Metallic foreign bodies are strongly echogenic and are very straightforward to locate. They will appear bright white and often give off strong reverberation artifacts (Fig. 36.US2).

Glass foreign bodies appear very similar to metallic objects in that they are highly echogenic (white). Glass may cause a reverberation artifact but more typically will cause acoustic shadowing to extend deep to the object (Fig. 36.US3).

Fluoroscopy
More recently, portable, low-power, C-arm fluoroscopy has become available in some emergency departments (EDs), particularly for orthopedic reductions. Its use has also been reported for the removal of BB pellets, metal, glass, and coins from patients.26 Like radiography, fluoroscopy can visualize objects that are radiopaque but not radiolucent such as wood.
Ultrasound Box 36.1: Foreign Body Removal—cont’d

Figure 36.US3 Glass foreign body (arrow) identified as a hyperechoic (white) object causing a strong shadow (arrowhead) within the otherwise normal-appearing soft tissue.

Figure 36.US4 Sonographic appearance of a retained wooden foreign body (arrow).

Wooden objects (such as splinters) are more challenging to locate, particularly with very small foreign bodies. Wooden foreign bodies do not create as strong an echogenic focus as other types of material do and may appear only slightly brighter than normal tissue (Fig. 36.US4). Very subtle shadowing may be seen extending deep to the object. The presence of surrounding edema is often key to locating the FB.

Removal

Once the object has been localized, it can be removed either blindly or under direct sonographic guidance.

Another technique that may be helpful, particularly with smaller objects, is to insert two 25-gauge needles under sonographic guidance. These needles should be inserted at right angles to each other so that the tips of each of them rest at the foreign body. The clinician can then cut the skin and overlying soft tissue and dissect down to the intersection of these two needles.

References


FB REMOVAL

Removal Decisions

One should judiciously evaluate and manage each FB scenario individually. The composition and location of an FB, as well as the patient’s medical status and vocational and avocational activities, greatly affect decision making related to FB removal, including the best time and place for removal. Reactive material, such as wood, should be removed immediately when accessible because retained wood will invariably lead to inflammation and infection. Other inert material, such as glass or plastic, may often be removed on an elective basis. Some innocuously located glass and metallic FBs may frequently be left permanently embedded in soft tissue. With deeply embedded, small, inert material (BB, bullets, etc.) that is not located near any vital structures, the time, effort, and trauma involved in removal may be excessive when compared with the possible adverse effects of the foreign material remaining in place. An ill-conceived extended search for an elusive but otherwise harmless FB often results in frustration for the clinician and discomfort, dissatisfaction, and possible injury for the patient.

If localization is certain and removal can be accomplished within a manageable period without worsening of the injury, an attempt at removal is generally indicated on the initial visit (given the availability of clinicians and support staff).

When reviewing the decision regarding when and how to remove the FB, the possibility of the FB migrating to involve vital structures, though quite remote, should be discussed with
the patient. Cases of reported missile embolization in the vascular system are influenced by missile caliber, impact velocity, physical wound characteristics, point of vessel entrance, body position and movement, and velocity of blood flow. Retained bullets usually remain in soft tissue but can rarely make their way into the vascular system. Schurr and colleagues reported a paradoxical bullet embolization from the left external iliac vein to the left iliac artery via a patent foramen ovale. When clinicians first examined the patient, a bullet was noted on the chest radiograph, and an isolated chest wound was suspected. However, the bullet had apparently entered the chest, traversed the abdomen to the iliac vein, and then embolized back to the chest and arterial system.

After the initial history, examination, and preoperative and preanesthetic documentation of the neurovascular status of the patient, a decision must be made regarding the time and place of removal. Thirty minutes is a reasonable self-imposed limit for the provider when attempting removal of an FB in the ED. More difficult procedures should be referred. Many FBs appear superficial on radiographs, thus suggesting that removal will be quite easy. However, surprisingly large or presumed superficial FBs still often prove quite elusive.

If an FB is left in place, inform the patient why it was not removed. If the patient is referred for delayed removal, this should also be carefully explained and documented. Regardless of whether the FB is removed, clean all wounds appropriately and update tetanus prophylaxis if indicated.

**Equipment and Preparation**

Good space and lighting, a comfortable patient and operator position, a standard suture tray, and a scalpel are usually adequate equipment for the removal of most simple FBs (Fig. 36.10). Tissue retractors, special pickups, and loupes may be added if needed.

Local soft tissue injection or selected nerve blocks with buffered bupivacaine or lidocaine (or both) are the recommended anesthesia for the removal of most soft tissue FBs. Judicious use of sedation (parenteral, rectal, or oral) is advised if the clinician senses undue apprehension or anxiety in the patient. Sedation may be especially helpful in children, with ketamine often being an excellent choice. If the patient is totally uncooperative, postpone exploration to a more appropriate time and setting (e.g., under regional or general anesthesia in the operating room).

To successfully remove an FB in the soft tissue of an extremity, a time-limited arterial tourniquet can help provide a bloodless field. Inflate a blood pressure cuff or portable self-contained pneumatic cuff above arterial pressure on the upper part of the arm, forearm, leg, or thigh. To limit bothersome backbleeding, elevate the extremity and wrap it with an elastic bandage to exsanguinate the extremity before inflating the tourniquet. A Penrose drain or specialized tourniquet may be used as a tourniquet at the base of a finger or toe. Alternatively, use a sterile glove as a finger tourniquet. Cut the fingertip of the glove on the involved finger and roll the glove down to the base of the finger. Most patients can tolerate an ischemic tourniquet for 15 to 30 minutes, and it is safe to stop the circulation to an extremity for this length of time.

**Operative Technique**

The specific technique for removal of an FB is tailored to each clinical situation (Videos 36.1 to 36.4). In general, FBs should be removed only under direct vision. Grasping blindly into a wound with a hemostat to remove an FB should be avoided. This technique is especially dangerous in the hand, foot, neck, or face, where sensitive or vital structures may easily be damaged.

After obtaining appropriate informed consent and following sterile preparation, consider enlarging the entrance wound with an adequate skin incision because it can be advantageous. Numerous techniques are used, depending on the clinical scenario (Fig. 36.11). Attempting to remove an FB through a puncture wound or an inadequate skin incision is a common error that is both frustrating and self-defeating. After a proper skin incision, explore the wound carefully by spreading the soft tissue with a hemostat. Frequently, the FB can be felt with an instrument before it can be seen (see Fig. 36.11, plate 1). After placing a tourniquet on an extremity, follow the track of the FB, although it often cannot be identified when surrounded by muscle or fat. If the FB, such as one that is made of fiberglass or plastic, is difficult to visualize, is located in the superficial soft tissue, or has contaminated the surrounding soft tissue, excising a small block of tissue rather than removing the FB alone may be necessary. Excise the block of tissue only under direct vision and after nerves, tendons, and vessels have been identified and excluded from the excision area.

If an FB such as a thorn or needle enters the skin perpendicularly, a linear incision may pass to one side or the other of the FB, and it may be difficult to determine where the FB lies in relation to this incision (see Fig. 36.11, plate 2). For this reason, the search must then be extended into the walls of the incision rather than simply through the skin. In such cases, excise a small ellipse of skin and undermine the skin for 0.5 to 1.0 cm in all directions. Next, compress the tissue from the sides in the hope that the FB will extrude and can then be grasped with a hemostat.

After removal of the FB, it is important to irrigate and cleanse the wounds. If a small incision has been made in a noncosmetic area (such as the bottom of the foot), leave the incision open and bandaged. The area may be periodically soaked in hot water for a few days. A return visit is necessary.

![Figure 36.10](image-url) The contents of a standard suture tray, with the addition of a scalpel, are usually adequate for the removal of most foreign bodies.
Foreign Body Removal Techniques

A. A sewing needle completely embedded below the surface is easily located with a radiograph. B. After the application of local anesthesia, a small incision over the superficial end permits removal with a hemostat. C. The hemostat is introduced through an adequate incision, spread to open the tissue, and used to "feel" the FB as the hemostat is advanced.

If a linear skin incision is used to locate a mobile FB that is perpendicular to the skin in subcutaneous fat (A), the FB may be displaced (B). C. A modified elliptical incision is made and the skin edges are undermined to displace the FB into the middle of the wound. Pressure with the thumbs may be applied to the skin to force the FB into view.

A. FBs in deep fat (A) may be approached by making a small elliptical incision around the entry point (B). C. The incision is then laterally undercut and grasped (without pulling) with forceps. D. The ellipse is then further undercut until contact with the FB is made. E. The FB may be grasped and removed along with the entry tract and the soiled local fat.

Figure 36.11 Foreign body (FB) removal techniques. (1 and 3, Reproduced from Hospital Medicine, January 1981, with permission of Cahner’s Publishing Co; 2, From Rees CE. The removal of FBs: a modified incision. JAMA. 113:35, 1939. Copyright 1939, American Medical Association. Reproduced with permission.)

Special Scenarios and Techniques

Puncture Wounds in the Sole of the Foot

Puncture wounds in the feet from unknown objects and under unknown circumstances present problems to the clinician. In general, puncture wounds in the feet are at risk for retained FBs and infection. It is impossible to adequately explore a puncture wound. In noncosmetic areas, therefore, a puncture

only if signs of infection develop. If a large incision has been made, the skin may be sutured primarily as long as no other contraindications are present. In cases in which gross contamination has occurred or there is significant tissue injury, do not close the wound on the initial visit. Leave the wound open but packed. Suture the skin after 3 to 5 days if the wound is free of inflammation or infection (known as delayed primary closure; see Chapters 34 and 35 for details).
One technique is to bend the tip of a sterile hypodermic needle and slide it under the nail. The needle is used to hook the FB and then remove it. Alternatively, slide a 19-gauge hypodermic needle under the nail to surround a small splinter. Bring the needle tip against the underside of the nail and secure the splinter. Remove the needle and splinter as a unit.35

Another possible technique is to shave the overlying nail plate with a No. 15 scalpel blade via light strokes in a proximal-to-distal direction. This creates a U-shaped defect in the nail and exposes the entire length of the sliver.36

Wooden splinters are commonly embedded under the fingernail. Such FBs must be removed completely because of the high risk of subsequent infection. If complete removal

---

Figure 36.12 This patient stepped on “something” while walking through a polluted urban creek, a setup for a foreign body and infection. Mandatory radiographs were negative. A, Under proper lighting (and a calf blood pressure cuff for a tourniquet), the patient is placed prone. Local lidocaine with epinephrine is infiltrated through the cut skin edges, not via the intact sensitive skin. (Alternatively, a posterior tibial nerve block could be used.) B, Inspection reveals a dirty skin flap that is cut off. C, Because a puncture wound cannot be explored or irrigated, this wound should be lengthened with a scalpel to allow further care. D, The depths of the wound are explored. E, An assistant holds the wound open with a hemostat so that copious irrigation can be accomplished. This wound should be packed open and checked in 24 to 48 hours, not closed primarily with sutures. Delayed closure can be performed in 4 to 5 days if necessary, but this injury healed well without sutures. Prophylactic antibiotics are of no proven value but may be given.

---

wound can be converted to a laceration to adequately explore and clean it (Fig. 36.12). Under most circumstances, delayed primary closure is recommended (see Chapter 34). This topic, including stepping on a nail and puncture wounds in the sole of the foot, is discussed extensively in Chapter 51.

Subungual FBs

Once a subungual FB has been identified, perform a digital block before manipulation of the nail or nail bed. Pay special attention to deeply embedded subungual FBs.35 Some cases may require removing a small portion of the nail with double-pointed heavy scissors to expose the FB and grasp the foreign material with splinter forceps (Fig. 36.13).
is gross contamination or significant tissue devitalization. Entrance or exit wounds should not be closed. Antibiotics are usually unnecessary for minor uncomplicated gunshot wounds in otherwise healthy patients, but may be beneficial in patients with multiple injuries, multiple comorbidities, gross wound contamination, significant tissue devitalization, large wounds, or delays in treatment.

There is a concern about lead toxicity from bullet fragments. If a bullet is bathed in synovial, pleural, peritoneal, or cerebrospinal fluid, the lead may leach out over time and produce a significant elevation in blood lead levels. Though rare, lead fragments in contact with synovium are a reason for concern and potentially a reason for removal (see the previous discussion). In one study, Farrell and associates found that patients with retained lead FBs had statistically significant elevated blood lead levels. The study did not differentiate the location of the FB and whether patients were, in fact, symptomatic from plumbism. Nevertheless, it may be prudent for the patient’s primary care provider to monitor blood lead levels in patients with known lead FBs to prevent the future development of lead toxicity. The value of routine prophylactic antibiotics for metallic FBs left in soft tissue has not been proved and antibiotics are generally not used.

For superficial metallic FBs, a sterile magnet can be used to facilitate the removal of small metallic fragments. Introduce the magnet into the entry site, with a small scalpel and hemostat used to extend and open the wound as needed. When the magnet comes in contact with the metal, a click is heard and the FB is removed while attached to the magnet. If resistance to extraction occurs, exploration can be done with the FB attached to the magnet. Sometimes, instead of introducing the magnet directly into the wound, it can be placed on the overlying skin to guide superficial small FBs out of the wound. Try not to grasp or touch the bullet because it can interfere with ballistics evidence during any subsequent legal investigation.

**FBs in Fatty Tissue**

FBs in fatty tissue can be removed by making an elliptical incision around the entrance wound. Grasp the incised skin loosely with Allis forceps. Undercut the incision until the FB is contacted. Remove the FB, skin, and entrance track in one
block. Remove a small portion of subcutaneous fat along with the FB to minimize infection. FBs in fat are very mobile, and probing may displace them even further. FBs that are embedded in fat and are perpendicular to the skin can also be removed, as shown in Fig. 36.11. As always, the removal of soft tissue, including adipose, should be done in as conservative a manner as possible, as any tissue removal may have implications in scar formation.

**Pencil Lead/Graphite**

Use careful judgment in removing pencil graphite when it is lodged in the skin. Graphite invariably leaves a pigmented tattoo in the soft tissue, and it is preferable to excise the material en bloc (see Fig. 36.11) when pencil lead is found in a cosmetic area. The graphite specks cannot be irrigated or scrubbed off, and tattooing results if they are not removed. Furthermore, a pencil lead FB may resemble a malignant melanoma over time. US has been shown to be useful in differentiating between pencil lead FBs and melanoma.40

**Fishhooks**

Traditionally, four fishhook removal techniques (see Video 36.1) have been described: advance and cut, string-yank, needle cover, and retrograde.41 The preferred method depends on the type of fishhook, location, depth of penetration, and conditions under which the removal is to take place (Fig. 36.15).42–44 Initially, note whether the fishhook is single or multiple and determine the number and location of barbs. Remove or cover any remaining exposed hooks to prevent subsequent injury. As with most injuries, document the patient’s vascular and neurologic status before and after removal. As with all wounds, administer tetanus prophylaxis if indicated. Prophylactic antibiotics are not generally necessary.

**Advance and Cut Technique**

To perform the advance and cut technique for removal, advance the fishhook and cut proximal to the barb (Fig. 36.16, plate 1). This method is particularly well suited for superficially embedded fishhooks. Generally, infiltrate local anesthetic (1% lidocaine) into the tissue overlying the barb. Force the barb through the surface of the anesthetized skin and clip it off. Move the rest of the hook retrogradely, along the direction of entry. Because this technique is almost always successful, it can be considered a first-line option.

**String-Yank Technique**

In the field or stream, removal of a fishhook may be accomplished without local anesthesia by using the string-yank technique. This technique may be used in the ED as well. Some clinicians prefer to use local 1% lidocaine to facilitate removal. For the “stream” technique (see Fig. 36.16, plate 2), pass a looped string or fishing line around the belly of the hook at the point where it enters the skin. Wrap approximately 1 ft of string around the dominant hand to provide strong traction. Hold the shank of the hook parallel to and in approximation to the skin with the index finger of the opposite hand. Use the thumb and middle finger of the opposite hand to stabilize and depress the barb, which helps the index finger disengage the barb from the subcutaneous tissue. When the barb has been disengaged, give a sharp pull (i.e., a quick tug with a snapping motion) with the dominant hand to remove the hook. Take care to keep bystanders out of the expected path of the hook because it often flies out of the patient.

**Needle Cover Technique**

For this technique use an 18-gauge needle to cover the barb (see Fig. 36.16, plate 3). After adequate local anesthesia has been achieved, pass the needle through the entrance wound of the hook parallel to the shank of the hook to sheath the barb and allow the hook to be backed out while the barb is covered. An alternative to this procedure is to insert a No. 11 blade parallel to the shank of the hook down to the barb and use the point of the blade to free the subcutaneous tissue that is engaged on the barb. Cover the barb with the point of the
Removal of wooden splinters is a common FB removal procedure. By simply grasping the end of a superficial, protruding splinter, it may be adequately removed, but care should be taken to not leave small pieces of material in the wound. Some splinters cannot be visualized at the point of entry but can easily and readily be palpated beneath the skin. When a wood FB is in subcutaneous tissue, it is advisable to cut down on the long axis of the FB to remove it via a skin incision rather than pulling it out through the entrance wound (Figs. 36.17 and 36.18). Although an incision may seem extensive and creates...
a laceration where only a puncture wound existed, opening the track allows thorough cleaning and removal of all the small pieces of the splinter that might otherwise remain. Any remaining piece of wood can lead to infection. The deeper the splinter, the more likely that small pieces of the splinter will remain in the deep part of the wound. If the incision is linear, it may be sutured. Occasionally, the fastest method of removing small wooden splinters is to completely excise the entrance track and the FB en bloc, followed by linear closure. Particular mention should be made of certain wood splinters that are pliable and reactive, such as California redwood and northwest cedar. Any wood that is easily fragmented requires meticulous care to ensure removal of all material.

**Traumatic Tattooing**

Ground-in foreign material or tattooing of the skin is a difficult problem because permanent disfigurement may occur (Fig. 36.19). These injuries occur most often from falls on blacktop surfaces or asphalt or falls from bicycles or motorcycles on a variety of surfaces. Many cases may be managed

![Figure 36.17](image)

**Figure 36.17** A, The course and depth of penetration of this large wooden splinter in the leg are uncertain, but it is axiomatic that all pieces of wood must be removed to prevent infection. B–D, To ensure complete removal of foreign bodies, an incision is made over the entire course of the splinter. E–G, All pieces of wood are carefully removed under direct vision. H, The laceration is sutured primarily. Although it may be tempting to simply pull the splinter out and irrigate the puncture track, such actions often lead to retained particles and complications.
with adequate local anesthesia and meticulous débridement with a sponge, scrub brush, or toothbrush. If all foreign material cannot be removed with these methods, give careful consideration to secondary excision of the tattooed area and primary closure with subsequent plastic surgery to repair the defect. However, it is usually impossible to completely remove traumatic tattooing in the ED. Referral for more extensive surgical treatment after local wound care is quite acceptable. Dermabrasion may be an acceptable delayed treatment when the tattooing is superficial. An alternative is to refer patients to a dermatologist for laser removal of traumatic tattoos. Certain lasers, such as a yttrium-aluminum-garnet (YAG) laser, have proved to be an excellent alternative. The process is more specific in removing embedded material without harming the surrounding tissue. The type of material will determine the number of treatments required. Not all tattoos can be removed completely.

**Marine FBs**

Although most marine FBs, such as shell fragments, may be treated like other FBs, a number of marine animals carry toxins and may leave FBs that require special consideration. Saltwater marine FBs may be contaminated with *Vibrio* species, which are usually sensitive to tetracyclines, aminoglycosides, or third-generation cephalosporins. Even if all foreign material has been removed, stings from marine animals may initiate a prolonged local irritation that simulates cellulitis. However, the presence of a wound infection on a subsequent visit strongly raises concern for an occult retained FB rather than the simple conclusion that the wound is merely infected by bacteria.

**Figure 36.18** A, A splinter was removed from the foot of this 3-year-old 7 days earlier, and a few days after removal it began to swell. A retained foreign body (FB) was suspected. Plain radiographs were negative. A small pyogenic granuloma is seen. After ketamine sedation, a 5-mm core of tissue was removed, but no FB was found on extensive exploration. This was an abscess from the puncture. The wound was packed open and cephalexin was given. B, It healed well (1 week later), but if it recurs, computed tomography or magnetic resonance imaging would be indicated to search for an occult FB.

**Figure 36.19** A, This patient was seen approximately 1 week after a blast injury. Traumatic tattoos are evident. The patient subsequently underwent successful laser removal of most of the pigment. B, Traumatic tattoo of the chin. Bluish discoloration and slight erythema, predominantly caused by silica, are apparent. (A, From Flint PW, Haughey BH, Lund VJ, et al, editors: *Cummings Otolaryngology: Head & Neck Surgery*, ed 5, St. Louis, 2010, Mosby; B, From Bolognia JL, Jorizzo JL, Shaffer JV, editors: *Dermatology*, ed 3, St. Louis, 2012, Saunders.)
edema, and cardiorespiratory failure.\textsuperscript{48,49} Dysrhythmias, hypotension, syncope, bronchospasm, laryngeal sequelae ranging from malaise and headache to paralysis and muscle spasm, but severe reactions may result in neurologic response. Systemic reactions usually consist of fever, chills, proteinaceous substances are implicated in the systemic reactions. Vinegar (5\% acetic acid) is the initial decontaminating agent of choice because it will inactivate the unfired nematocysts of most (but not all) species of jellyfish, Portuguese man-of-war, and sea anemones.\textsuperscript{48,50,51} Apply it continuously for 30 minutes or until the pain is gone.\textsuperscript{49} Do not apply fresh water to the area because the osmotic shock will activate any remaining unfired nematocysts and cause them to discharge toxin and increase pain and toxicity. Other suggested, but unproven, remedies include meat tenderizer, ammonia, baking soda, urine, olive oil, sugar, and papaya latex.

After the wound has been decontaminated, remove any remaining tentacle fragments. Extract large fragments with forceps; however, individual nematocysts are very small ($<1$ mm in length) and are not easily visible. To remove the remaining nematocysts, scrape the skin with a hard edge, such as a credit card held perpendicular to the skin. An alternative is to apply shaving cream and shave the area gently to eliminate the remaining fragments.\textsuperscript{49} Do not use sand to remove nematocysts because sand can increase the discharge of venom. If the pain persists, it should be assumed that organelles still remain and further cleansing is required.

After decontamination, use topical anesthetics or steroids, but prophylactic antibiotics are not necessary. Follow routine wound care. Treat allergic and systemic reactions appropriately. The application of warm or cold packs to the area has not been shown to reduce pain.\textsuperscript{50} Antivenin is available only for box jellyfish envenomation, but its use remains incompletely understood and controversial.

In summary, a 2013 Cochrane review found inconclusive evidence defining the most effective treatment for jellyfish envenomation, but a 2016 article suggested several interventions may be effective for nematocyst inhibition. These authors wrote that topical vinegar for at least 30 seconds for some species while seawater, baking soda slurry and water warmed to 42°C to 45°C can help, along with oral analgesics.\textsuperscript{52,53} Ultimately species-specific interventions are the most effective, but there is inadequate research to make definitive recommendations.

Coral
Coral is composed of a calcium carbonate core and thousands of small marine animals. Fire coral, a specific type of coral, is another type of coelenterate that produces toxicity with stinging nematocysts. After contact, a burning and intense pruritus may occur along with a series of skin eruptions. Within minutes of contact, pruritus, erythema, and urticaria-like lesions may appear, and blister formation may result within hours (Fig. 36.21). Eventually, the lesions will become lichenoid, but complete resolution may not take place for 15 weeks after contact. Ultimately, hyperpigmented areas will form at the point of initial contact. Immediate care with oral antihistamines and topical steroids tends to reduce, but not prevent, the symptoms.

“Coral cuts,” which can be deep lacerations, occur in divers and snorkelers exploring coral reefs. With these wounds, delayed healing may take place with the secondary development of cellulitis or ulceration, perhaps as a result of contamination of the wound with bacteria or microparticles of coral.\textsuperscript{48} Treatment consists of copious saline irrigation. Hydrogen peroxide may be used to help remove small coral particles from the wound.\textsuperscript{48} Do not close the wound; wet-to-dry dressings are advised instead.

Sponges
Sponges produce both an irritant and a contact dermatitis. The irritant dermatitis occurs as a result of sponge spicules embedded in the victim’s skin. Remove the spicules with adhesive tape (applied to the skin and then peeled back), and then bathe...
handled or inadvertently stepped on, these spines may become embedded in the patient and a severe local reaction may result from venom in the spines. Systemic symptoms occur and include muscle weakness; paralysis of the lips, tongue, and face; hypotension; abdominal pain; and respiratory distress. Local pain responds quite well to immersion in hot water (43.3°C to 46.1°C [110°F to 115°F]) for 30 to 90 minutes. Retained spines may become infected or cause delayed (∼1 to 2 months) FB granulomas. This reaction is not adequately understood, but it may be due to an intense and persistent inflammatory reaction. Spines that penetrate joints may induce synovitis. Echinoderm spines may discharge a purple dye that may be mistaken for a retained spine (Fig. 36.22).49 Spines are usually visible on radiographs and should be removed if possible, although they are brittle and can break off in the skin. A YAG laser may be an effective alternative to remove sea urchin spines. If spines are located in a joint or near a nerve, surgical extraction using an operative microscope may be necessary.48 Otherwise, if removal is difficult, leave the spine in place until it is resorbed or a local reaction takes place. If appropriate, open the wound and drain it to allow it to close by secondary intention.

**Catfish**

Several species of catfish in North America contain toxic venom, and a sting from the dorsal or pectoral spines can embed an FB. The spine secretes venom from an epidermal gland at its base.54 The pain is usually ephemeral, and because no specific antitoxin exists, treatment consists of local care and analgesics. Immerse the affected part in hot water (∼43.3°C [110°F]) for at least 30 minutes if the pain is severe. This is believed to provide relief of symptoms by decreasing vascular and muscle spasm.55 Local injection of the wound with alkalized bupivacaine provides local analgesia and may also neutralize the toxin.54 Inspect the wound and remove any remaining spines (Fig. 36.23). A radiograph may be taken to confirm the absence of FBs, but catfish spines and cartilage may be radiolucent. Bedside

---

**Figure 36.21** Fire coral stings. (From James WD, Berger TG, Elston DM, editors: *Andrews’ diseases of the skin: clinical dermatology*, ed 11, St. Louis, 2011, Saunders.)

**Figure 36.22** A, Multiple sea urchin punctures in the hand soon after injury and following a soak in hot water. B, The same hand after 6 days without intervening therapy other than soaking. Lack of discoloration indicates absorption of dye from the sea urchin spines and probable absence of retained fragments. (From Auerbach PS, editor: *Wilderness Medicine*, ed 6, St. Louis, 2011, Mosby.)
true infection, and surgical exploration is often required in persistent cases. Administer tetanus prophylaxis as per routine recommendations.

Cactus Spines
The size of cactus spines fluctuates considerably. The difficulty of removal is generally inversely proportional to the size of the FB. Larger embedded cactus spines are managed like
wood splinters and sea urchin spine FBs. More advanced imaging techniques (US, CT, or MRI) may be required for localization of deeply embedded spines.

Deeply embedded cactus spines generally produce granulomatous reactions, but infections are rare.\(^5\) Dermatitis from embedded cactus spines is a well-described phenomenon. Hence, make an effort to remove deeply embedded spines after carefully weighing the benefit and potential harm related to deep exploration, especially in a sensitive location.\(^5\) Using forceps, remove superficially embedded, medium to large cactus spines by direct axial traction on each spine. Smaller spines (glochids) may be difficult and tedious to remove individually. Apply an adherent facial mask gel to remove the spines en masse with the gel (Fig. 36.25). Depilatory wax melted in a microwave oven and applied warm, commercial facial gels, and household glue (Elmer’s Glue-All, Borden, Inc., Columbus, OH) have all been recommended for this purpose.\(^5\) Over-the-counter “home use” facial mask gels are not adherent enough to be effective without multiple (eight or more) applications.

**String-Wrap Method**

An occasional patient can remain calm during this procedure, but if the swelling is significant or the digit has been traumatized, anesthesia is necessary. Perform a proximal digital or metacarpal block to provide sufficient anesthesia and to minimize tissue disention at the ring site. Before removal of the ring, wrap a wide Penrose drain circumferentially in a distal-to-proximal direction to reduce the soft tissue swelling (Fig. 36.26). Leave the wrap in place for a few minutes to reach the maximum effect. Some nonanesthetized patients panic during the procedure because of increasing pain from compression and unwinding.\(^6\)

First, pass a 20- to 25-inch piece of string, umbilical tape, or thick silk suture between the ring and the finger. Shorter lengths are discouraged because one may need to repeat the wrapping procedure midway. If marked soft tissue swelling is present, pass the tip of a hemostat under the ring to grasp the string and pull it through. Wrap the distal string clockwise around the swollen finger (proximal to distal) to include the proximal interphalangeal (PIP) joint and the entire swollen finger. Start the wrapping next to the ring. Wrap it snugly enough to compress the swollen tissue. Place successive loops of wrap next to each other to keep any swollen tissue from bulging between the strands. When the wrapping is complete, carefully unwind the proximal end of the string in the same clockwise direction to force the ring over that portion of the finger that has been compressed by the wrap. The PIP joint is the area that is most difficult to maneuver over and causes the most pain.

Occasionally, the finger must be rewrapped if it was not done carefully the first time. It is not uncommon to produce abrasions or other kinds of trauma in the skin during this procedure. If the finger with the ring is lacerated or there are underlying fractures, it is prudent to cut the ring off instead of attempting this technique.

Certain rings are made of extremely hard material such as tungsten carbide or ceramic. In these cases, cracking the material with standard locking pliers can break the ring. Place the pliers on the ring and adjust the jaws to fit tightly, and then remove and readjust them while increasing tension with each subsequent adjustment. Continue until the material cracks and falls apart. Some rings may be lined with a metal band. Use a standard ring cutter to remove the band.

**Ring Cutter**

A ring cutter should be used when the swelling is excessive or other methods fail (Fig. 36.27). A ring cutter has a small hook that fits under the ring and serves as a guide for the saw-toothed wheel that cuts the metal. The cut ends of the ring are spread with large hemostats (e.g., Kelly clamps), and the ring is removed. If the tension is too great to spread the ring, another cut 180 degrees apart from the original ring cut can be performed. This will allow the ring to fall off in two pieces. A jeweler can subsequently repair cut rings.

Certain hardened metal rings, such as tungsten carbide, may not be amenable to the use of a ring cutter. Case studies have demonstrated that a dental-tipped drill or dental volvere can be used successfully. However, because of the nature of these instruments and the possibility of injury from them, such as lacerations from the blade, thermal burns, or ocular FBs, take precautions to limit further damage to the finger, as well as the use of eye protection to shield the patient’s and operator’s eyes.
**Ring Removal: String-wrap Method**

1. Lessen the edema by compressing the finger with a Penrose drain left tightly wrapped for 3 to 5 minutes.
2. Slide a small hemostat under the ring, grab a long piece of umbilical tape, and pull it under the ring.
3. Begin to carefully wrap the tape around the finger in a proximal-to-distal fashion.
4. Continue winding the distal strand to compress the skin distal to the ring. Take time to place successive loops next to each other, and keep tissue from bulging between the strands.
5. Remove the ring by unwrapping the proximal strand in the same direction that it was wrapped. The most difficult area to negotiate is the proximal interphalangeal joint. You may have to repeat the wrapping procedure to totally remove the ring.

**Figure 36.26** Ring removal: string-wrap method.

**Body Piercing and Removal**

The art of body piercing predates most history books. Over the past decade an enormous increase in the practice of body piercing has occurred. Accordingly, so have the complications associated with the practice. For centuries the ears were the most common place. Today, the lips, tongue, eyebrow, nose, navel, nipples, and genital areas have become sites of body piercing. To date, there are only a limited number of studies on infection after piercings in areas other than the ears. Three major types of jewelry are used: (1) barbell studs, which are
straight bars with a ball threaded onto both ends; (2) labret studs, which are straight bars with a ball threaded on one end and a disk permanently fixed on the other end (more commonly used on the lips); and (3) a captive bead ring, which consists of a bead with small dimples on opposite sides and an incomplete ring with rounded ends to fit into the dimples (Fig. 36.28). The bead is held “captive” by tension from both sides of this incomplete ring. The bead ring is a variation of this: one bead is permanently fixed to one end, and an opening is made by removing the free end of the ring.

The most common reason for removal is infection (Fig. 36.29). Other symptoms such as bleeding, edema, allergic reaction, and keloid formation may also prompt removal. Occasionally, tongue piercings must be removed to permit intubation. To remove barbell- and labret-type studs, hold the bar with forceps and unscrew the bead on the other end. To remove a captive bead ring, hold the ring on both sides of the captive bead to release tension on the bead. This will dislodge the bead from the ring, which is holding it in place. If the jewelry is near the mouth or nose, take care to prevent aspiration.

Figure 36.27 Ring removal: ring cutter method.

Figure 36.28 Various types of piercing jewelry. To remove a barbell stud or labret stud, unscrew the ball that is threaded onto the bar. To remove a captive bead ring, snap the ball out of the incomplete ring (it is held in place simply by tension and is not screwed in). You may need to insert needle-nose pliers into the center of the ring and spread it to pry the ring open. To remove a bead ring, unscrew the ball from one end of the ring.
Tick Removal

It is important to remove ticks early because of tick-borne disease, including the hard tick of the Ixodid family. Rocky Mountain spotted fever, Lyme disease, tularemia, and ascending paralysis are among the many infections identified as tick-borne diseases. It is important to note that the rate of disease transmission before 48 hours of attachment is exceedingly low. Removal of Ixodid ticks is difficult because the mouthparts become cemented within 5 to 30 minutes of contact with the host’s skin. Removal will become more difficult the longer the tick is attached. Inadequate or partial removal of the tick may cause infection or chronic granuloma formation. Removal by mechanical means is recommended. Nonmechanical, traditional, and
lacerates the skin and increases the amount of tissue caught in the mechanism. Although the clinician may anesthetize the skin and excise the entrapped tissue, a less invasive method can be considered.

Cutting the median bar between the faceplates of the zipper mechanism remains the most common method. The interlocking teeth of the zipper then fall apart when the median bar (diamond or bridge) of the zipper is cut in half (Fig. 36.32), and the skin is subsequently freed. A bone cutter or wire clippers and a folk methods of forcing the tick to disengage (e.g., the use of petroleum jelly, fingernail polish, a hot match, or alcohol) are not advised and can cause the tick to regurgitate, thereby increasing the possibility of transmission of infection.

The use of straight or curved forceps or tweezers is the recommended method of removal. If these instruments are not available, use a gloved hand. Grasp the tick as close to the patient’s skin surface as possible and gently apply steady axial traction (Fig. 36.30). Take care to not squeeze, crush, twist, or jerk the tick’s body because this may expel infective agents or leave mouthparts in the skin. If mouthparts are left behind after removal of the body, they may be removed with tweezers. If one is still unable to remove the mouthparts, consider excision under local anesthesia to prevent local infection.

Many patients have great anxiety about the development of tick-borne diseases after tick removal. Some studies have demonstrated that single-dose doxycycline (200 mg) may prevent the development of Lyme disease. Children younger than 8 years may be given a single dose of doxycycline (4 mg/kg). However, prophylactic antibiotic treatment of all tick bites is not recommended. Amoxicillin prophylaxis is not effective. When a patient is in an area where the incidence of Lyme disease is high or when a partially engorged deer tick in the nymphal stage is discovered on the body, the patient is more likely to benefit from prophylaxis. Regardless of whether prophylaxis is given, instruct patients about the symptoms and signs of Lyme disease and encourage them to return or seek medical evaluation if these symptoms develop.

**Zipper Entrapment**

The skin of the penis may become painfully entangled in a zipper mechanism (Fig. 36.31). Unzipping the zipper frequently...
moderate amount of force may be required to break the bar. The addition of mineral oil followed by traction has been demonstrated to achieve some success. Patients with penile lacerations warrant urologic follow-up to assess for urethral injury.

**Infiltration of Radiographic Contrast Material**

The infiltration of high-osmolality intravenous contrast material has the potential to cause skin necrosis, but the use of low-osmolality dye, which is well tolerated in soft tissues, has essentially eliminated this problem (Fig. 36.33). The use of high-pressure power injectors, with the technologist out of the room, calls for careful inspection of the intravenous site before injection. However, because of the high-pressure rapid auto-injection of contrast material and the absence of a technician in the room, extravasation of contrast material is not a rare event and occurs even with meticulous technique. Infiltration occurs in approximately 0.5% to 1% of auto-injects. Once contrast material has infiltrated, no intervention has been demonstrated to ameliorate the local reactions, which are usually mild. Swelling, erythema, and mild discomfort at the infiltration site may occur. It is best to resist the temptation to use excessive heat or cold. Elevation of an extremity is appropriate but of unproven value. Injection of steroids and other agents has no known benefit. Low-osmolality dye is usually totally resorbed in a few days, with no serious consequences in the vast majority of cases. Progress of the dye’s egress may be followed with radiographs. The majority of contrast material extravasations resolve with no long-term consequences with conservative management. If large volumes of contrast material are injected (>75 to 100 mL), especially in confined areas such as the dorsum of the hand, it is possible that skin necrosis from the pressure or a compartment syndrome may develop, but this is rare. The clinician should resist attempts to routinely remove even large volumes of extravasated contrast material by incision or aspiration. Although some contrast material may be removed with surgical techniques during fasciotomy, the actual benefit is unknown. Surgical intervention is rarely required and is based on measurement of compartment pressure and clinical evaluation. Patients with small-volume and minimally symptomatic extravasations may be discharged. Large-volume extravasations warrant consultation or further observation.

**TASER Darts**

The Thomas A. Swift Electric Rifle, or TASER, is a conducted electrical weapon used in many areas by police to subdue violent patients (see Chapter 70). The TASER fires two barbed electrodes on long copper wires (Fig. 36.34). The barbs attach to skin or clothing and create an arc that delivers an electrical jolt that causes overwhelming pain and involuntary muscle contraction and incapacitates the subject. The electricity is of such high frequency that it is believed to stay near the surface and not penetrate to the depth of internal organs. The barbs are designed to not penetrate deeper than 4 mm, and police are taught to remove them by stretching the surrounding skin and tugging sharply. If this fails, cutting down on the dart after local anesthesia should facilitate removal. Patients may need medical evaluation after ED removal of the darts for the underlying state of agitation that required the use of a TASER, complications of electrical injury, injury from the fall after incapacitation, and injury from the barb, especially if struck in the mouth, eye, neck, or groin. Most patients do well with minimal intervention and proper wound management if the patient is not unduly agitated and the TASER did not involve the critical areas of the body just mentioned.
CHAPTER 36  Foreign Body Removal  733

Figure 36.33  The entire volume (100 mL) of nonionic low-osmolality contrast material (Isovue) for a computed tomography scan was pressure-injected into the antecubital soft tissue when the intravenous line infiltrated. High-volume injections are automatically administered by a programmed auto-injector; the technician is not in the room to stop it. A, There was only mild pain but considerable soft tissue swelling, and most of the redness and blistering occurred when the technician taped an unprotected ice pack directly to the skin and nearly caused frostbite. B, X-ray evidence of the infiltration. C, Within 36 hours, the dye was absorbed, without further treatment. No skin necrosis occurred, as has been seen when older ionic high-osmolality agents infiltrate. There is no known way to ameliorate the potential soft tissue injury. The antecubital fossa is the preferred site for an intravenous line for injection of contrast material, and such an extravasation on the dorsum of the hand may be more serious. Do not attempt to remove the contrast material by aspiration or surgical incisions.

Human and Animal Bite FBs
The most common FB after a human bite is a piece of tooth. These FBs can be difficult to find and may not be appreciated if the patient does not admit or disclose the bite wound. Usually, the bite puncture is small and not easily cleaned or visualized. The puncture can be widened with a formal incision to aid in cleaning and evaluation for tendon integrity, injury to the joint capsule, fracture, or an FB (Fig. 36.35). Occasionally, small pieces of teeth can become embedded in a wound, such as with dog, cat, or snake bites (Fig. 36.36). Radiographic detection is variable, and exploration is often the only alternative.

Pyogenic Granuloma (Lobar Capillary Hemangioma)
A pyogenic granuloma is a benign acquired polypoid, friable vascular lesion of the skin (hand, neck, foot, fingers, and trunk) and mucous membranes (Fig. 36.37). They are common in children; in pregnancy, the lesion is termed epulis gravidarum (pregnancy tumor). The cause is unknown, but there is some association with topical retinoids and the protease inhibitor indinavir; they are not due to infection, and they are not granulomas. They grow rather rapidly over a period of a few weeks, are occasionally associated with minor trauma, have a glistening dark red appearance, and may bleed. Histologically, a pyogenic granuloma is a hemangioma. A variety of topical therapies (silver nitrate, cryotherapy), laser, or cautery are available, but removal by sharp dissection with primary suturing

Figure 36.34  A and B, Barbed TASER darts can be removed by a quick pull or through a small incision made over the barb.

Figure 36.35  This human bite puncture wound was enlarged with an incision to aid in cleaning and search for a foreign body, usually a piece of tooth. Even though this wound appears clean, it should not be closed with sutures.
A and B. The bite of a boa constrictor, though not poisonous, may contain small tooth fragments. As with cat and dog bites, radiographic evaluation is variable and exploration may be required.

Figure 36.37 Pyogenic granuloma. Despite their name, these lesions are neither pyogenic (i.e., infectious) nor granulomas. Rather, they are a type of hemangioma. Pyogenic granulomas can develop rapidly over a period of weeks and are common in pregnant women.

is usually curative (Fig. 36.38). The recurrence rate may be as high as 40% with nonsurgical intervention. 

Hair-Thread Tourniquet
Hair or thread fibers adherent to infants’ clothing occasionally become tightly wrapped around a child’s digits or genitals (Fig. 36.39). If they are left in place, amputation may eventually occur. The offending fibers may be difficult to visualize, and the child is often brought for evaluation only after signs of distal ischemia appear. Occasionally, the fiber can be grasped with toothless forceps or a small hemostat and then unwrapped. More commonly, fibers cannot be identified because they are deeply embedded in swollen tissue. Removal not requiring minor surgery includes use of a chemical depilatory, such as the over-the-counter product Nair (calcium thioglycolate), to dissolve or weaken the hair. A generous amount of depilatory cream is worked into the involved area and allowed to dissolve or weaken the hair, which is then removed mechanically. If a depilatory is not successful in 30 to 60 minutes, surgical removal is indicated.

A No. 11 blade can be used to cut the constricting bands under a regional nerve block. It may be difficult to identify individual hairs that are deeply embedded in a swollen digit and even more difficult to assess the success of the intervention. Frequently, multiple hairs are involved. Because the bands may be quite deep, the incision should avoid known neurovascular tracts. Barton and coworkers recommend a dorsal, rather than a lateral, incision on the digits. If the soft tissue on the distal end of the digit has been rotated after a circumferential dermal laceration from the tourniquet, the distal tissue can be realigned with the proximal tissue and two dorsolateral sutures placed or tissue adhesive glue applied to maintain the digit in alignment.

Generally, conservative wound care is sufficient once the band has been removed. Application of an antibiotic ointment may enhance healing and allow easier removal of serous drainage from the circumferential laceration. Clinical reassessment in 24 hours will indicate whether any constricting bands remain.

DISPOSITION MANAGEMENT

Tetanus
Wounds with FBs should be considered contaminated wounds, and tetanus status should be updated according to the recommendations for patients with contaminated wounds.

Antibiotics
There is no consensus or standard for the use of antibiotics after removal of wound FBs. Antibiotics may be indicated for immunocompromised patients, but there are no data to support the routine use of antibiotics in patients with wounds that have been thoroughly cleaned and from which all foreign material has been extracted. Prophylaxis may be considered if there was excessive time between injury and removal or obvious contamination or when it is difficult to adequately clean the
of community-acquired methicillin-resistant *S. aureus* in many areas, clindamycin, trimethoprim-sulfamethoxazole, or tetracycline may provide alternative coverage. However, infections associated with FBs are not likely to be from methicillin-resistant *S. aureus*. Under certain circumstances, alternative antibiotics may be indicated. For infected plantar punctures through a shoe, a fluoroquinolone to cover *P. aeruginosa* is appropriate coverage, although most *Pseudomonas* infections are complex and may need extensive débridement and intravenous antibiotics. Saltwater marine FBs may be contaminated with *Vibrio* species, which are usually sensitive to tetracyclines, aminoglycosides, or third-generation cephalosporins. Freshwater FBs, on the other hand, are more likely to harbor *A. hydrophila*, which can be treated with tetracyclines.

**FB Reactions**

If the FB is not removed or cannot be removed, an FB reaction may occur. Some FBs produce an inflammatory reaction or infection a few days after introduction into the body. Other objects may not cause problems for weeks, months, or even years until they flare up for no apparent reason. The primary factors that affect the extent of tissue reactions are contamination and whether the material is inert or reactive with human tissue. Reactive FBs, such as wood, will generally produce inflammation eventually, whereas inert FBs, such as bullets, rarely do. Some inert FBs carry dirt particles, pieces of clothing, or other sources of bacterial contamination. Expeditious removal may be necessary, even if the FB itself is relatively small and unlikely to cause a reaction.
A purulent bacterial infection may develop in the presence of any FB; therefore, any abscess or cellulitis that recurs or wounds that do not heal as expected should always be investigated for retained FBs. Karpman and coworkers found a 15% rate of infection (S. aureus and Enterobacteriaceae) in a series of 25 patients treated for cactus thorn injuries on the extremities. Certain thorns (black thorns, rose thorns), redwood and northwest cedar splinters, toothpicks, hair, and stingray or sea urchin spines are noted for their ability to initiate chronic FB reactions. Sea urchin spines and other marine FBs are covered with slime, calcareous material, and other debris that commonly initiate an FB granuloma. The inflammatory reaction seen with cactus thorns may be an allergic reaction to fungus found on the cactus plant.

Many FB reactions are thought to result from an inflammatory response to organic material, or they may represent infection from bacteria introduced at the time of the wound. Clinically evident reactions may be delayed for weeks or even years after injury (Fig. 36.40). The chronic infection or inflammatory reaction may not be accompanied by the production of pus, but it may be quite painful or result in loss of function. FBs may also be associated with the formation or development of a chronic pseudotumor, a sinus tract, or an osteomyelitis-like lesion of bone and soft tissue. In addition, organic material has been noted to induce chronic tenosynovitis, chronic monarticular synovitis, and chronic bursitis.

Rapidly traveling projectiles with considerable inherent heat (e.g., bullets) are less likely to cause infection but are more apt to cause other difficulties. Damage to surrounding areas can occur during passage through tissue. Rarely do retained lead FBs, such as bullets or shotgun pellets, leach out lead into the general circulation and produce systemic lead poisoning unless they are in contact with synovium (Fig. 36.41). If this process does occur, it may take years to develop and can result in vague or nondescript symptoms (e.g., fatigue, arthralgia, headache, or abdominal pain) many years after the initial injury.

Figure 36.40 This foreign body (FB) granuloma developed after the FB was stable for 6 months. There was no gross infection and it was dissected en masse.

Figure 36.41 There is no need to routinely remove retained bullets. Most lead foreign bodies are well tolerated, but if a bullet is bathed in synovial, pleural, peritoneal, or cerebrospinal fluid (CSF), the lead may leach out over time and produce a significant elevation in blood lead levels. The symptoms are often vague, and the relationship between the retained lead and the patient’s clinical scenario may be difficult to sort out. A, This patient had chronic neurologic findings, including wristdrop, and was wheelchair bound. B, The symptoms were due to chronic lead poisoning from a 20-year-old retained bullet (arrows) in or near the spinal canal, possibly being bathed by CSF. C, Note the lead lines in the gingiva, indicative of lead poisoning.
Elevated blood lead levels are more likely to occur if body fluids such as joint, pleural, peritoneal, or cerebrospinal fluid bathe the lead. Bullets retained in muscle or other soft tissue are not likely to produce any sequelae related to their lead content. However, Farrell and colleagues reported unsuspected elevated lead levels in patients with retained lead fragments who were seen in the ED with a variety of complaints. Lead levels of up to 50 µg/dL were reported. Levels greater than 45 µg/dL are generally considered an indication for chelation therapy. The relationship between the retained lead and the symptoms was unclear, but this report verifies the observations of others that retained lead FBs in selected areas can significantly elevate blood lead levels and may produce symptomatic plumbism.

**Discharge Instructions**

The patient should be informed that despite every effort there can be no absolute guarantee that all foreign material has been identified or extracted, regardless of whether some or any FB was removed during the initial exploration. Prudent clinicians should always suggest close follow-up and should leave open the possibility that an occult FB may still remain in the wound. Discharge instructions should also include the signs and symptoms of problems related to retained material. Some centers routinely add this caveat on all discharge instructions to patients treated for lacerations or soft tissue defects. Written instructions should inform patients that additional steps may be undertaken if foreign material is subsequently suspected. Discharge instructions should indicate that close, continued and careful follow up is important with any evaluation for FB, FB removal, and post-FB removal assessment.

**ACKNOWLEDGMENT**

The authors and editors wish to thank Ted Koutouzis and Matthew Levine for contributions to this chapter in previous editions.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Incision and Drainage

Gina Ambrose and Donald Berlin

Incision and drainage (I&D) procedures (Videos 37.1-37.5) in the emergency department (ED) are most commonly performed for soft tissue abscesses (Fig. 37.1). The total number of ED visits increased from 90 million to 115 million over a 10-year period, with visits for abscess-related complaints increasing faster than overall ED visits. Of an estimated 34.8 million ambulance visits for skin and soft tissue infections, 33% are seen in EDs.

The emergence and predominance of methicillin-resistant Staphylococcus aureus (MRSA) as the cause of cutaneous abscesses during the past several decades have necessitated major revisions in long-standing guidelines for antibiotic administration. In light of this significant etiologic change, MRSA should be considered as a probable cause in most skin and soft tissue infections.

ABSCESS ETIOLOGY AND PATHOGENESIS

Localized pyogenic infections may develop in any region of the body. Abscesses generally begin as a localized superficial cellulitis. Some organisms cause necrosis and liquefaction, as well as the accumulation of leukocytes and cellular debris. This is followed by loculation and subsequent walling off of these products leads to abscess formation. The cause of an abscess depends on its anatomic location and flora indigenous to that area. Different organisms cause disease based on environmental exposure. For example, direct inoculation of extraneous organisms may occur during a mammalian bite (e.g., Eikenella, Pasteurella), exposure to saltwater (e.g., various Vibrio strains) or freshwater (e.g., Aeromonas), or meat or fish exposure (e.g., Erysipelothrix rhodanines). Pseudomonas folliculitis has been associated with the use of a hot-tub, as this organism thrives in a warm, wet environment.

Staphylococcal strains, which are normally found on the skin, produce rapid necrosis, early suppuration, and localized infections with large amounts of creamy yellow pus, the typical manifestation of an abscess. Conversely, group A β-hemolytic streptococcal infections tend to spread through tissues and cause a more generalized infection characterized by erythema, edema, a serous exudate, and little or no necrosis, typical manifestations of cellulitis. Anaerobic bacteria, which proliferate in the oral and perineal regions, produce necrosis with profuse brownish, malodorous pus and may cause both abscesses and cellulitis.

Normal skin is extremely resistant to bacterial invasion, and few organisms are capable of penetrating intact epidermis. In a normal healthy host with intact skin, the topical application of even very high concentrations of pathogenic bacteria does not result in infection. The requirements for infection usually include a high concentration of pathogenic organisms, such as in hair follicles in the adnexa; occlusion of glands or other structures that prevent desquamation and normal drainage; a moist environment; adequate nutrients; and trauma to the corneal layer, which allows organisms to penetrate into deeper tissues. Tissue perfusion may also play a role in the ability to prevent infection. Trauma may be the result of abrasions, shaving, insect bites, hemotoma, injection of chemical irritants, incision, or occlusive dressings that macerate the skin. The presence of a foreign body can potentiate skin infections by enabling a lower number of bacteria to establish an infection. For example, abscesses occasionally develop at suture sites in otherwise clean wounds. In addition, abscesses can develop at any site used for body piercing. Ear piercings through the cartilage of the pinna seem to be at particular risk for infection because of the avascularity of auricular cartilage.

When favorable factors are present, the normal flora that colonize cutaneous areas flourish and infect the skin and deeper structures. In persons performing manual labor, the arms and the hands are infected most frequently. In women, the axilla and submammary regions are frequently infected because of minor trauma from shaving, contact with undergarments, a moist environment, and an abundance of bacteria in these areas. Infections may develop anywhere on the body in intravenous (IV) drug users, although the upper extremities are most commonly affected. Deep soft tissue abscesses can be caused by an addict’s attempts to access deep venous structures when peripheral venous access sites are exhausted. In addition, areas with compromised blood supply are more prone to infection because normal host defenses, including cell-mediated immunity, are less available.

Bacteriology of Cutaneous Abscesses

Although most abscesses contain bacteria, 5% of abscesses are sterile, especially those associated with IVDU. Clinically, sterile
abscesses cannot be differentiated from those caused by bacteria. They are caused by injected irritants that are not fully absorbed, causing local inflammation at the site of injection. Somewhat atypical abscesses develop in parenteral drug users. Injection of a cocaine-heroin mixture (“speedball”) may predispose users to abscesses by inducing soft tissue ischemia. Jenkins and colleagues compared the microbiology of abscesses in IVDU-patients versus patients without a history of IVDU and found that streptococcal species and anaerobes were identified more commonly than *Staphylococcus* in IVDU-patients.

The microbiology and the underlying cause of skin and soft tissue abscesses are related to their location. Abscesses involving the extremities are generally the result of a breach in the skin’s integrity from trauma such as cuts, abrasions, or needle punctures. Abscesses involving the head, neck, and perineal region are usually associated with obstruction of the apocrine sweat glands. These types of abscesses increase in frequency after puberty because of the increased apocrine and sebaceous gland activity. Perirectal abscesses are typically the result of bacterial spread from adjacent anal glands. Vulvovaginal abscesses usually result from obstruction of a Bartholin gland, which then causes duct and gland edema and subsequent infection. Pilonidal abscesses are hypothesized to be caused by sacrococcygeal infections from ingrown hairs in the intergluteal cleft.

In 2002, Brook compiled the findings from more than 15 bacteriologic studies of 676 polymicrobial abscesses. *S. aureus* and group A β-hemolytic streptococci were the most prevalent aerobes in skin and soft tissue abscesses and were isolated in specimens from all body sites. Gastrointestinal and cervical flora (enteric gram-negative bacilli and *Bacteroides fragilis*) were found most often in intraabdominal, buttock, and leg lesions. Group A β-hemolytic streptococci, pigmented *Prevotella*, *Porphyromonas* species, and *Fusobacterium* species—all normal residents of the oral cavity—were most commonly found in lesions of the mouth, head, neck, and fingers.

In a study of the bacteriology of cutaneous abscesses in children, Brook and Finegold found aerobes (staphylococci and group A β-hemolytic streptococci) to be the most common isolates from abscesses of the head, neck, extremities, and trunk, with anaerobes predominating in abscesses of the buttocks and perirectal sites. Mixed aerobic and anaerobic flora was found in the perirectal area, head, fingers, and nail bed. This study noted an unexpectedly high incidence of anaerobes in non-perineal abscesses. Anaerobes were found primarily in areas adjacent to mucosal membranes (e.g., the mouth), where these organisms tend to thrive, and in areas that are easily contaminated (e.g., by sucking fingers, which causes nail bed and finger infections or bite injuries). If an unexpected or atypical organism is found in an abscess culture, the clinician should consider an underlying process that is not readily apparent from the history or physical examination. For example, tuberculosis or fungal isolates are sometimes found in immunocompromised patients (e.g., those with diabetes or acquired immunodeficiency syndrome). Finding *Escherichia coli* suggests an enteric fistula or even self-inoculation of feces in some patients with a psychiatric illness such as Munchausen’s syndrome. Recurrent abscesses without an obvious underlying cause could indicate clandestine drug use. What appears to be a typical recurrent abscess may be a manifestation of an underlying septic joint, osteomyelitis, or rarely, metastatic or primary cancer (see Fig. 37.2).

**Special Considerations**

Parenteral drug users, insulin-dependent diabetics, hemodialysis patients, cancer patients, transplant recipients, and individuals with acute leukemia have an increased frequency of abscess formation when compared with the general population. At initial evaluation, the patient may emphasize an exacerbation of the underlying disease process or an unexplained fever, with symptoms of an abscess being a secondary complaint. In these situations, abscesses tend to have exotic or uncommon bacteriologic or fungal causes and typically respond poorly to therapy. Patients with diabetes-induced ketoacidosis (DKA) should be evaluated extensively for an infectious process; a rectal examination should be included with the physical examination to rule out a perirectal abscess as the infectious trigger of DKA. This is also true for patients who are immunocompromised. There are several reasons why patients with diabetes and parenteral drug users are at increased risk for abscess formation: intrinsic immune deficiency, an increased incidence of staphylococcal carriage, potentially compromised tissue perfusion, and frequent needle punctures, which allow a mode of entry for pathogenic bacteria.

Patients who use IV drugs frequently use veins in the neck and in the femoral areas, which can produce abscesses and other infectious complications at these sites. Any abscess near a vein of the antecubital fossa or dorsal of the hand should alert the clinician to possible IV drug use; however, substance users may also inject directly into the skin (“skin popping”), which can cause cutaneous abscesses distant from veins (Fig. 37.3).

A foreign body may serve as a nidus for abscess formation. Patients with a history of IVDU frequently break needles in skin that has been toughened by multiple injections, so the clinician should maintain a high index of suspicion for retained needle fragments. If an abscess is recurrent or if the patient is a known or suspected IV drug user, consider radiographs or other
Virulent community-acquired MRSA (CA-MRSA) causes rapid and destructive soft tissue infection because of the presence of two bacterial toxins elaborated by the omnipresent USA-300 and USA-400 strains. Panton-Valentine leukocidin enhances tissue necrosis, and phenol-soluble modulin is toxic to neutrophils. Methicillin resistance is mediated by PBP-2a, a penicillin-binding protein encoded by the \textit{mecA} gene, which permits the organism to grow and divide in the presence of methicillin and other \textit{β}-lactam antibiotics. \textit{S. aureus} acquires methicillin resistance through a mobile staphylococcal cassette chromosome (SCC) that contains the \textit{mecA} gene complex (SCC\textit{mec}). MRSA probably arose as a result of antibiotic selective pressure.\textsuperscript{25,26}

A single clone probably accounted for most MRSA isolates discovered during the 1960s; by 2004, six major MRSA clones

**MRSA**

First acknowledged in the 1960s as a cause of infection in patients in health care settings, MRSA has now become the most common identifiable cause of community-acquired skin and soft tissue infections in many metropolitan areas in the United States. The spread of this organism is considered an epidemic and it is very virulent and aggressive.\textsuperscript{22,23} One study found an MRSA prevalence as high as 75% to 80% in some parts of the country.\textsuperscript{24} Ultrasound is also a useful adjunct to evaluate for foreign bodies that are not radiopaque.

**Figure 37.2** An abscess that appears in an atypical place or recurs after successful initial treatment should raise the possibility of rare or underlying conditions. A, This patient had a large “abscess” on the lateral chest wall that initially drained unusual gelatinous material, not frank pus. B, At follow-up 3 days later, the abscess was much improved. The contents of the abscess had been sent for pathologic analysis because it had an unusual consistency, and a highly undifferentiated soft tissue malignancy was demonstrated. The fluid was sterile. Normally, analyzing or culturing the contents of an abscess will not yield helpful information, but in this case the unusual consistency of the collection prompted further analysis. C, This intravenous drug user had an “abscess” of the chest wall drained in various emergency departments several times over a 2-month period, and it seemed to initially respond to drainage and antibiotics. He still had an area of cellulitis, minor fluctuance, and continued drainage near the center of the chest. This is an atypical place for a simple cutaneous abscess. Magnetic resonance imaging demonstrated osteomyelitis and an abscess of the sternoclavicular joint that was draining to the skin and simulating a recurrent cutaneous abscess. He required extensive surgical débridement and prolonged antibiotics. The etiologic organism was never ascertained, but \textit{Pseudomonas} is often present. D, This patient underwent a sternotomy for bypass surgery a few months previously. She had been treated sporadically for a minor wound infection, but then a draining fluctuant mass developed at the inferior border of the sternum. This is the external manifestation of extensive sternal osteomyelitis.
known risk factors. A small pustule can become a large abscess in 24 to 48 hours (Fig. 37.4). Such lesions are often mistaken for a spider bite or drug use because of their rapid progression and seemingly spontaneous onset in an otherwise healthy person. These observations have led to the identification of some risk factors for CA-MRSA, including skin trauma (e.g., lacerations, had emerged. The spread of resistance is thought to be mediated by horizontal transfer of the mecA gene and related regulatory sequences thereon.

In 1980, the spread of MRSA from hospitals into communities became evident. More recently, community-acquired infections have occurred more frequently, even in people without

**Figure 37.3** This patient had a large abscess in the deltoid area (arrow) and could offer no explanation for it. This is a typical scenario for a drug user who injects directly into the skin. B, The characteristic circular skin lesion from “skin popping” found on the arms (arrows) confirmed the clinical suspicion. Even though a drug screen was positive for opioids, the patient denied drug use and attributed the leg lesions to frequent trauma on the job. Drug users with abscesses are at risk for numerous infections, including brain abscess, endocarditis, and occult osteomyelitis.

**Figure 37.4** A–C, Examples of community-acquired methicillin-resistant *Staphylococcus aureus* (CA-MRSA) infections. These aggressive infections can spread rapidly. The patient frequently describes a small pustule that becomes an abscess in 24 to 48 hours. Patients often believe that it is a spider bite because of its rapid onset in an otherwise healthy person with no other reason for the lesion. A CA-MRSA abscess, though clinically aggressive, is usually treated like any other cutaneous abscess.
tattoos, IV and intraderal drug use, shaving), incarceration, shared razors or towels, and close contact with others colonized or infected with MRSA. Animals can also carry MRSA and can function as a source of transmission. Importantly, many patients with CA-MRSA have no identifiable risk factors for acquisition of the disease. CA-MRSA tends to be more virulent than health care–associated MRSA (HA-MRSA) and is associated with more frequent serious complications such as osteomyelitis, joint infections, sepsis, and death. However, these organisms fortunately tend to be susceptible to a broader array of antibiotics.

The prevalence of MRSA has increased in both health care and community settings. For example, the prevalence of methicillin resistance among *S. aureus* isolates in intensive care units in the United States has been reported at 60%, and more than 90,000 invasive infections by MRSA occurred in the United States in 2005. HA-MRSA and CA-MRSA differ with respect to their clinical epidemiology and molecular structures. HA-MRSA is defined as MRSA infection that occurs following hospitalization (hospital onset, formerly “nosocomial”) or MRSA infection that occurs outside the hospital within 12 months of exposure to a health care setting (e.g., history of surgery, hospitalization, dialysis, or residence in a long-term care facility—community onset instead of community acquired).

HA-MRSA is usually associated with severe, invasive disease, including skin and soft tissue infection, bloodstream infection, and pneumonia. In fact, *S. aureus* continues to be a significant cause of surgical site infections. HA-MRSA strains tend to be resistant to multiple drugs. MRSA is one of the few pathogens routinely implicated in nearly every type of hospital–acquired infection. This is probably related in part to the organism’s capacity for biofilm formation on indwelling lines and tubes in hospital settings. Biofilm facilitates survival and multiplication of MRSA on these surfaces, thereby prolonging the duration of exposure of the organism to antibiotics, as well as promoting the potential for the development of genetic resistance.

CA-MRSA is defined as MRSA infection that occurs in the absence of health care exposure. It is often associated with skin and soft tissue infections in young, otherwise healthy individuals. Most CA-MRSA strains are sensitive to non–β-lactam antibiotics, although a multidrug-resistant isolate has been described in men who have sex with men. This strain contains the pUSA03 plasmid and carries resistance genes for β-lactams, fluoroquinolones, tetracycline, macrolides, clindamycin, and mupirocin.

The CA-MRSA and HA-MRSA classifications are no longer distinct as MRSA colonization can develop in one realm and manifestations of infection in another. In the mid-2000s in San Francisco, the annual incidence of CA-MRSA surpassed that of HA-MRSA.

Furthermore, community-onset HA-MRSA infections have been observed with increasing frequency. This was illustrated in a study of 209 patients discharged from hospitalized care; within 18 months following hospital discharge, 49% of new MRSA infections began outside the hospital. In another series of 102 patients with CA-MRSA infections, 29% had molecular typing consistent with HA-MRSA.

CA-MRSA was initially reported in injecting drug users in the early 1980s and has since become the most frequent cause of skin and soft tissue infections seen in US EDs and ambulatory clinics. In an assessment of the prevalence of MRSA across the United States, Moran and colleagues compiled data from adults who sought treatment of acute skin and soft tissue infections in EDs in 11 American cities in August 2004. *S. aureus* was isolated from three fourths of the 422 patients who met the study criteria. Seventy-eight percent of the *S. aureus* isolates were resistant to methicillin. MRSA was isolated from 59% of patients in the study. The prevalence of MRSA ranged from 15% to 74% in the participating EDs. MRSA was the most common identifiable cause of skin and soft tissue infections in all but one of the EDs.

Frazee and associates, reporting from an ED in northern California, found that half of the 137 patients in their study were either infected with or colonized by MRSA. Three fourths of all *S. aureus* isolates were MRSA. In addition, 76% of cases met a strict clinical definition of CA-MRSA. The incidence of CA-MRSA, genetically unrelated to nosocomial isolates, increased steadily from 1990 to 2001 and then dramatically in 2002 and each year thereafter.

MRSA has also emerged as a potential sexually transmitted disease. Roberts and colleagues described their treatment of two patients who came to their urban ED with abscesses, probably transmitted by heterosexual oral–genital contact. Both tested positive for MRSA. A 2010 case report drew a similar conclusion. It reported orogenital transmission of MRSA to an immunocompetent 22-year-old man who tested orally positive for MRSA and group B (genital) *Streptococcus* after oral contact with a female partner in whom MRSA-positive gluteal lesions had previously been diagnosed. MRSA abscesses have been described following skin-to-skin contact during a lap dance.

In a retrospective chart review, Roberts and colleagues found that 18% of the 524 subcutaneous abscesses treated in their urban ED in 2006 were confined to the genital area. Almost three fourths of the 272 outpatient wound cultures performed on that year’s patient population were positive for MRSA.

**MANIFESTATIONS OF CUTANEOUS ABDOMESSES**

The diagnosis of cutaneous abscess formation is usually straightforward. The presence of a fluctuant mass in an area of induration, erythema, and tenderness is clinical evidence that an abscess exists (see Fig. 37.1). An abscess may appear initially as a definite, tender, soft tissue mass, but in some cases a distinct abscess may not be readily evident. If the abscess is deep, as is true of many perirectal, pilonidal, and breast abscesses, the clinician may be misled by the presence of a firm, tender, indurated area without a definite mass. If the findings on physical examination are equivocal, needle aspiration or ultrasound examination may be performed to assist in the diagnosis. This approach may also identify a mycotic aneurysm or an inflamed lymph node simulating an abscess. A specific entity commonly mistaken for a discrete abscess is the sublingual cellulitis of Ludwig’s angina (see Chapter 65).

Parenteral injection of illicit drugs can produce simple cutaneous abscesses that unpredictably advance to extensive necrotizing soft tissue infections. The emergency clinician must maintain a high index of suspicion to avoid missing this potentially life-threatening condition. Cellulitis and abscess formation can lead to bacteremia and sepsis, especially in immunocompromised patients.

The pain of an abscess often brings the patient to the hospital before it spontaneously ruptures, or the patient can have a draining abscess that appears to have undergone spontaneous...
rupture and is self-resolving. The patient may have even punctured the abscess in an attempt to drain it. In most cases, a formal I&D procedure will be necessary to effectively manage the condition, even though copious drainage may not be encountered. Although no formal drainage may be required after the spontaneous rupture of a simple cutaneous abscess, conditions such as a perirectal abscess, Bartholin gland abscess, and breast abscess are usually best managed with further appropriate drainage and packing.

**IMAGING**

**Ultrasound-Guided Needle Aspiration**

Radiologists have been performing ultrasound-guided needle aspiration of abscesses for some time, and emergency clinicians are now becoming more comfortable with the procedure. High-resolution ultrasound technology is being used to obviate “blind” procedures done in the ED (e.g., joint aspiration and central line placement).

For ultrasound-guided drainage of a cutaneous abscess, use a high-resolution probe (7.5–10 MHz) and maintain sterility throughout the procedure. Place the sterile transducer over the main body of the abscess and insert the needle through the skin adjacent to the transducer. Adjust their relative relationships in keeping with the depth and location of the abscess cavity. Guide the needle, seen as a bright artifact, directly into the abscess. Watch the abscess cavity collapse as pus drains out. Scan the entire area of the suspected abscess and beyond to capture unexpected extensions of the abscess. Be sure to drain all pockets.

**LABORATORY FINDINGS**

A complete blood count (CBC), blood cultures, and Gram stain are not standard or required for the treatment of straightforward cutaneous abscesses in the ED. Recommendations for culturing abscesses encountered in the ED are confusing and clinical practice varies. Firm recommendations for the emergency clinician are difficult to standardize, partly because of insufficient data but also because the recommendations promulgated are not confined to ED abscess treatment. In addition, “complicated” and “uncomplicated” criteria are somewhat arbitrary. Traditionally, culturing the contents of a readily drainable cutaneous abscess was not indicated, nor standard. It simply provided no additional useful information to the clinician under most circumstances. Many clinicians still forgo routine culturing, even in the CA-MRSA milieu. Currently, there is no agreement concerning routine culturing, and reasonable arguments can be made for a culture or no-culture approach to most abscesses treated in the ED. The authors support selective, not routine culturing but acknowledge that some now consider cultures to be indicated for all abscesses drained in the ED. A culture will potentially identify an unusual or resistant organism, especially if I&D is not curative. Culture will also permit identification of antibiotic susceptibility and assist in customization of antibiotic therapy. Cohort results also provide a framework for local epidemiology and resistance patterns. Culturing the abscess contents will distinguish between MRSA and nonresistant abscesses and will provide useful sensitivity information when managing complicated cases.

Culturing should be performed for recurrent, unusual, or atypical abscesses. This information could be useful if the patient responds poorly to initial surgical drainage, if secondary spread of the infection occurs, or if bacteremia develops. It also appears prudent to obtain cultures from abscesses and other purulent skin and soft tissue infections in patients already taking antibiotics, in immunosuppressed patients, in those with signs of systemic illness, in patients who have not responded adequately to initial treatment, if there is concern for a cluster or outbreak of infection, or in patients with severe local infection. Severe local infection can be defined as an abscess larger than 5 cm in diameter, multiple lesions, or extensive surrounding cellulitis. However, the degree of surrounding cellulitis qualifying as “extensive” is ill defined.

When obtaining a specimen for culture, the most accurate and complete culture results will be obtained if one aspirates pus with a needle and syringe before I&D. The material should be cultured for aerobic and anaerobic bacteria. Most clinicians, however, still culture free-flowing purulent material obtained with a cotton swab during I&D. For uncomplicated ED abscesses, this culture technique is standard and adequate to isolate aerobic organisms, including MRSA. A “sterile” culture from a specimen collected with a standard cotton swab after incision is frequently the result of improper anaerobic culture technique. In selected patients, such as immunocompromised hosts or IV drug users, isolation of possible anaerobic organisms by needle aspiration with a syringe through appropriately cleaned (e.g., with chlorhexidine) skin before I&D will enhance the results of culture and can add information that may be clinically relevant to subsequent therapy. There is a general misconception that foul-smelling pus is a result of *E. coli*. This foul odor is actually caused by the presence of anaerobes; the pus associated with *E. coli* is odorless.

The discovery of solid or suspicious material in an abscess should prompt histologic evaluation because a malignancy may mimic cutaneous abscesses (see Fig. 37.2A and B).

The majority of patients with an uncomplicated cutaneous abscess will have a normal CBC and will not experience fever, chills, or malaise. Therefore, in the absence of extenuating circumstances, it is not standard to analyze blood from patients with typical cutaneous abscesses because laboratory test results do not lead to a specific therapeutic path. An abscess may produce leukocytosis, depending on the severity and duration of the purulent process; however, the presence or absence of leukocytosis has virtually no diagnostic or therapeutic implications. Bacteremia may occasionally be manifested as a peripheral abscess resulting from septic emboli, and it usually produces clinical characteristics dissimilar to those associated with simple cutaneous abscesses. A cutaneous abscess itself rarely produces bacteremia.

Gram stain is neither indicated nor standard in the care of uncomplicated simple abscesses. However, patients who appear “toxic” or immunocompromised and those who require prophylactic antibiotics (see the section on Prophylactic Antibiotics later in this chapter) may benefit from Gram stain in addition to cultures. Gram stain results have been shown to correlate well with subsequent culture results, so in compromised hosts the test can be used to direct the choice of antibiotic therapy. Anaerobic infections should be suspected when multiple organisms are noted on Gram stain, when a foul odor is associated with the purulence, when free air is noted on radiographs of the soft tissue, and when no growth is reported on cultures.
Ultrasound offers a distinct advantage when evaluating a patient with suspected soft tissue infection and may change management. A recent study from *Academic Emergency Medicine* by Tayal and colleagues evaluated the effect of soft tissue ultrasound on the management of cellulitis in the emergency department. The authors found that in patients with a low suspicion for abscess, ultrasound changed management in 56% of cases. Peritonsillar abscesses are difficult to diagnose from the physical examination alone, and some clinicians may feel hesitant to attempt blind drainage. Ultrasound of suspected peritonsillar abscesses has been found to be reliable in making the diagnosis. The overall size of the abscess, as well as its proximity to the carotid artery, can be evaluated with ultrasound, which will perhaps improve the confidence of the clinician in attempting drainage.

**General Considerations**

Typically, a high-frequency (7.5 to 10 MHz) transducer should be used to evaluate the superficial soft tissues. The higher frequency will allow the clinician sufficient resolution to identify changes consistent with soft tissue infection. The entire area should be scanned in detail, in multiple planes, to identify fluid pockets. Surrounding structures in the area should also be evaluated, especially when incision and drainage are planned. When evaluating the posterior pharynx for a potential peritonsillar abscess, an intracavitary transducer should be used.

**Normal Soft Tissue**

Normal soft tissue is characterized by well-defined layers, with clear demarcation between these layers (*Fig. 37.US1*). The top of the screen corresponds to the most superficial soft tissue, including the epidermis and dermis. It should appear hyperechoic (light gray to white), thin, and clearly separate from the underlying layers. Subcutaneous tissue is found beneath the dermis and is of varying thickness. However, as with the most superficial layers, this layer should appear thin and well demarcated from the surrounding layers. Underneath the subcutaneous tissue, muscle will typically be seen as layers of striated tissue separated by bright layers of fascia.

**Cellulitis**

Cellulitis is recognized on ultrasound by thickening of the skin and subcutaneous layers (*Fig. 37.US2*). The tissue may also appear more hyperechoic than normal soft tissue. When a significant amount of edema is present within the tissue, bands of hypoechoic (dark gray) or anechoic (black) fluid may be seen within the area of thickened tissue. This is known as “cobblestoning” (*Fig. 37.US3*). Cobblestoning appears as thin bands of fluid throughout the tissue and can be distinguished from an abscess by the lack of a discrete fluid collection.

**Abscess**

An abscess is seen as a focal, discrete fluid collection within an area of cellulitis (*Fig. 37.US4*). The presence of surrounding cellulitis is the key to distinguishing an abscess from other fluid collections such as cysts. The character of the fluid may be variable, depending on the...
Incision and Drainage

CHAPTER 37 Incision and Drainage

745

drained cutaneous abscesses in immunocompetent hosts. For simple abscesses, I&D alone is likely to be quite adequate and curative, even if the causative organism is MRSA. Routine administration of antibiotics after I&D is not currently standard, although the topic is subject to ongoing investigations. Simply stated, drainage alone for uncomplicated abscesses is usually curative. Furthermore, the use of antibiotics may in fact be harmful. Antibiotic misuse has also been shown to complicate resistance patterns, both in general and in specific patients.

Even though no randomized controlled trial data definitively demonstrate the need for antibiotic therapy in conjunction with I&D of uncomplicated cutaneous abscesses in healthy, immunocompetent patients (without the specific types of valvular heart disease discussed later), there is strong consensus for antibiotic treatment of abscesses associated with the following conditions: severe or extensive disease (e.g., involving multiple sites of infection) or rapid progression in the presence of associated cellulitis, signs and symptoms of systemic illness, associated comorbid conditions, immunosuppression, extremes of age, abscess in an area difficult to drain (e.g., face, hand, and genitalia), associated septic phlebitis, and lesions that are unresponsive to I&D alone.13

Antibiotic use is indicated for abscesses with associated severe cellulitis (this term is not well defined and generally is a clinical judgement) and those with purulent cellulitis. Purulent

ULTRASOUND BOX 37.1: Cellulitis and Abscesses—cont’d

Figure 37.US4 Ultrasound appearance of an abscess. A large, well-contained hypoechoic (dark gray) fluid collection (arrow) can be seen surrounded by thickened subcutaneous tissue.

Figure 37.US5 Ultrasound image of a peritonsillar abscess. A rounded area of mixed density (both anechoic and hypoechoic areas) is seen in the center of the image.

INDICATIONS FOR AND CONTRAINDICATIONS TO I&D

Surgical I&D is the definitive treatment of a soft tissue abscess,2; antibiotics alone are often inadequate. Drainage of a suppurative focus generally results in marked resolution of the symptoms in most uncomplicated cases. In the initial stages, only induration and inflammation may be found in an area destined to produce an abscess. Premature incision, before localization of pus, will not be curative and may theoretically be deleterious because extension of the infectious process and, rarely, bacteremia can result from manipulation. In some cases, the application of heat to an area of inflammation may ease the pain, speed resolution of the cellulitis, and facilitate the localization and accumulation of pus. Nonsurgical methods are not a substitute for surgical drainage and should not be continued for more than 24 to 36 hours before the patient is reevaluated.

PROPHYLACTIC AND THERAPEUTIC ANTIBIOTIC THERAPY

The utility of antibiotics remains unproven for prophylaxis against and for treatment of uncomplicated and adequately

References:
cellulitis is defined as cellulitis associated with purulent drainage or exudate in the absence of a drainable abscess. Purulent cellulitis is usually caused by MRSA. Nonpurulent cellulitis, defined as cellulitis with no purulent drainage or exudate and no associated abscess, is usually due to β-hemolytic streptococci. Empirical therapy for MRSA, pending culture results, is recommended for patients who have failed non-MRSA treatment, for those with a previous history of or risk factors for MRSA, and for those with severe infection or systemic signs and symptoms. Empirical therapy for infection with β-hemolytic streptococci is likely to be unnecessary under these circumstances.

The duration of therapy for skin and soft tissue infections has not been well defined, although no differences in outcome were observed in adult patients with uncomplicated cellulitis receiving 5 versus 10 days of therapy in a randomized, controlled trial.\(^5\) In the Food and Drug Administration licensing trials for complicated skin and soft tissue infections, patients were typically treated for 7 to 14 days. However, in the outpatient setting of uncomplicated infections, 3 to 5 days of antibiotic therapy is reasonable but should be individualized on the basis of the patient’s clinical condition and response to treatment. Accordingly, a return wound inspection or primary care follow-up is an important component of the care plan.

Patients with a history of IVDU who have an abscess and fever require parenteral antibiotic therapy after blood has been drawn for culture, until bacterial endocarditis can be ruled out.\(^2\) Additionally, patients who have extensive cellulitis or are clinically septic require immediate IV antibiotics, as well as aggressive surgical drainage of any significant abscess. By administering IV ampicillin/sulbactam (2 g/1 g) every 6 hours, Talan and colleagues\(^5\) achieved 100% eradication of pathogens from major abscesses in hospitalized IV drug users and non-drug users.

**Therapeutic Antibiotics**

In contrast to prophylaxis before surgery, the routine use of therapeutic oral antibiotics after I&D of simple cutaneous abscesses in otherwise healthy patients who are not immunocompromised appears to have no value, and their empirical use cannot be scientifically supported. Llera and Levy\(^3\) performed a randomized, double-blind study to compare the outcomes of patients treated with a first-generation cephalosporin after the drainage of cutaneous abscesses in the ED with those who received a placebo. They found no significant difference in clinical outcome between the two groups and concluded that antibiotics are unnecessary for abscesses in individuals with normal host defenses. This is in agreement with several previous studies.\(^37\)–\(^39\) One study found that, although the use of trimethoprim-sulfamethoxazole (TMP-SMX) did not decrease treatment failure rates, it may help to decrease recurrence of subsequent lesions.\(^60\) It should be noted that high-risk patients were often excluded from these studies. Immunocompromised patients have not been adequately studied in this situation and are therefore often given antibiotics empirically; but this practice, though common, has not been supported by rigorous prospective studies.

For empirical coverage of CA-MRSA in outpatients with skin and soft tissue infection, oral antibiotic options include the following: clindamycin, TMP-SMX, a tetracycline (doxycycline or minocycline), and linezolid. If coverage of both β-hemolytic streptococci and CA-MRSA is desired, options include the following: clindamycin alone or the use of TMP-SMX or a tetracycline in combination with a β-lactam (e.g., amoxicillin or cephalaxin), or linezolid alone. The use of rifampin as a single agent or as adjunctive therapy for the treatment of skin and soft tissue infections is not recommended.\(^19\)

Facial abscesses should be handled cautiously and checked frequently. Any abscess above the upper lip and below the brow may drain into the cavernous sinus, so manipulation may predispose to septic thrombophlebitis of this system. Treatment with antistaphylococcal antibiotics and warm soaks after I&D has been recommended pending resolution of the process. Areas not in this zone of the face can be treated in a manner similar to that used for other cutaneous abscesses.

A relatively unstudied but a common and currently accepted strategy for patients with soft tissue infections (especially CA-MRSA infections) that are borderline, by clinical judgment, for hospital admission and therapeutic IV antibiotics, is to administer a single dose of an IV antibiotic in the ED, followed by oral antibiotics and close outpatient follow-up. When a CA-MRSA infection is likely, IV vancomycin or linezolid is a reasonable option. Oral linezolid may be as effective as the IV form.

**Prophylactic Antibiotics**

**Prophylaxis for Endocarditis**

The precise risk for endocarditis after I&D of a cutaneous abscess remains unknown, and it is difficult to predict in which patients an infection will develop and which particular therapeutic procedures subject the patient to the highest risk for infection. However, bacteremia clearly occurs with manipulation of infected skin, and infected musculoskeletal tissue only in patients with cardiac conditions that carry the highest risk for an adverse outcome from infective endocarditis.\(^61\) These conditions are listed in Box 37.1. Most skin infections are polymicrobial, but only staphylococcal and β-hemolytic streptococci are likely to cause infective endocarditis. Therefore the therapeutic regimen should include an agent active against these organisms, such as an antistaphylococcal penicillin or a cephalosporin. For patients who cannot tolerate a β-lactam or if MRSA is suspected, vancomycin or clindamycin can be substituted.

Two clinical situations deserve special mention. First, because of the frequent incidence of valvular damage in IV drug users, prophylactic antibiotics may be indicated before I&D of abscesses in these patients; however, no standards exist. It is prudent to inquire about previous endocarditis or auscultate for a heart murmur if IV drug abuse is suspected or known and administer antibiotics under these circumstances. Second, any patient with a documented history of endocarditis must receive prophylactic antibiotics before the I&D procedure.

Cutaneous abscesses may result from active endocarditis and prophylactic antibiotics may obscure subsequent attempts to identify the causative organism. With this in mind, two or three blood cultures (aerobic and anaerobic) should be
considered before antibiotic therapy for those at risk for endocarditis. Patients with a diagnosis of mitral valve prolapse have traditionally been included for treatment with prophylactic antibiotics, but the indication for this is unclear and antibiotic prophylaxis for uncomplicated mitral valve prolapse is no longer part of the AHA guidelines.66

Kaye67 suggested prophylaxis only for patients who have a holosystolic murmur secondary to mitral valve prolapse.

Prophylaxis for Bacteremia in Other Conditions

Conflicting results have been reported from the few studies investigating the relationship between I&D of cutaneous abscesses and bacteremia. For example, in 1985 Fine and associates68 concluded that I&D of cutaneous abscesses is often accompanied by transient bacteremia. They compared blood culture results from specimens obtained before and at 1, 5, and 20 minutes after I&D procedures in 10 patients with soft tissue infections. None of the cultures of blood obtained before I&D were positive; however, six patients had at least one positive culture after the procedure. Eleven of the 30 postprocedure cultures yielded growth. In contrast, in 1997 Bobrow and coworkers69 concluded that I&D of a localized cutaneous abscess is unlikely to result in transient bacteremia in afebrile adults. Their study included 50 patients with localized cutaneous abscesses. Blood samples were collected before and at 2 and 10 minutes after I&D. In addition, specimens from the wound were collected after drainage. None of the blood cultures were positive, even though 64% of the wound cultures were positive, primarily for S. aureus. Bobrow and coworkers69 noted that prophylactic antibiotics should be given to patients at high risk for bacterial endocarditis. In a discussion of the differences between these findings and those reported by Fine and associates,68 Bobrow and coworkers69 noted that Fine and associates’ cultures were obtained from indwelling, heparinized IV catheters, a practice that allows ample opportunity for contamination. Furthermore, half the patients in Fine’s group had perirectal abscesses, and if these abscesses involved mucosal surfaces, the risk for bacteremia was potentially increased.68

Immunocompromised patients may benefit from the prophylactic administration of antibiotics in preparation for I&D of cutaneous lesions. In contrast to patients with risks for endocarditis, immunocompromised patients are at risk for septicemia secondary to brief bacteremia. IV drug users have a high incidence of diseases associated with human immunodeficiency virus (HIV),65,67,71 and the treating clinician must anticipate various degrees of immunodeficiency among them. Because no specific standard of care exists, clinical judgment must guide the use of antibiotics in these situations.

No specific guidelines have been offered for the antibiotic regimen to be used before I&D of infected cutaneous tissue in patients at risk for conditions other than endocarditis. The choice of antimicrobial agent is based on the organism anticipated to cause the bacteremia. Although the location of the abscess will give some clue to the organism involved, most abscesses contain multiple strains of bacteria. Not all bacteria are potent pathogens, so their mere presence does not predict their role in subsequent morbidity. Because Staphylococcus continues to be the most common cause of cutaneous abscesses, a broad-spectrum antistaphylococcal drug is indicated. Prophylaxis can consist of a single IV dose given half an hour before I&D. A first-generation cephalosporin or penicillinase-resistant penicillin is a good initial choice. Vancomycin may also be considered. Others may prefer cefazolin (Ancef [GlaxoSmithKline, Research Triangle Park, NC], Kefzol [Eli Lilly and Co Indianapolis, IN]), 1 g intravenously, for adults. This regimen covers staphylococcal and streptococcal species, many gram-negative organisms, and many anaerobes.

Although it has not been studied, it is reasonable to also give antibiotics before I&D to all patients who will subsequently be administered therapeutic antibiotics. The ideal duration of therapeutic antibiotics is unknown. As a general guideline, immunocompromised patients should receive antibiotics for 5 to 7 days and immunocompetent patients for 3 to 5 days after the procedure, depending on the severity of the condition and their clinical response at follow-up.

**RECURRENT INFECTIONS**

The first-line approach to the prevention of recurrent soft tissue infections is education regarding personal hygiene and appropriate wound care. Patients should be directed to keep wounds covered with clean, dry bandages. Maintain good personal hygiene with regular bathing and frequent hand washing with soap and water or an alcohol-based hand gel, especially after touching the wound or items that came in contact with the wound. Avoid reusing or sharing personal items that have contacted the infected area.

Environmental hygiene measures should be considered in patients with recurrent skin and soft tissue infections in the household or community setting. Focus cleaning efforts on high-traffic surfaces (i.e., surfaces that come in frequent contact with skin each day, such as counters, doorknobs, and toilet seats) that may contact bare skin or uncovered infections.

An approach to individual treatment of recurrent CA-MRSA soft tissue infections is outlined in Box 37.2.

Oral antimicrobial therapy is recommended for the treatment of active infection only and is not routinely recommended for
Recurrent CA-MRSA soft tissue infections have been linked to a carrier state in affected individuals, with the nose and skin areas being colonized. It may be difficult to totally or permanently eradicate colonization, and there are no proven methods to accomplish this. The following strategies have been used in an attempt to eradicate the carrier state in individuals with recurrent CA-MRSA soft tissue infections. The appropriate time to implement these procedures is not known, but it would be reasonable to institute first-line techniques if more than two to three episodes of proven CA-MRSA infection are documented.

**FIRST-LINE**
Apply 1 cm of mupirocin (Bactroban [GlaxoSmithKline]) ointment (an intranasal form is available) to the anterior nares three times a day for 7 days, plus
Daily total body wash with (4%) chlorhexidine (Hibiclens [Mölnlycke Health Care US, LLC, Norcross GA]) for 5-7 days.
Note: For multiple recurrent infections, consider consultation with an infectious disease expert.

**SECOND-LINE**
Obtain specimens for culture to ascertain sensitivity to various antibiotics.
Repeat mupirocin and chlorhexidine as above.
Rifampin, 300 mg twice a day, plus trimethoprim-sulfamethoxazole DS or doxycycline (100 mg) twice a day for 1-2 weeks.
Note: For multiple recurrent infections, consider consultation with an infectious disease expert.

*a* May cause burning, pruritus, dry membranes, and erythema. Avoid long-term use. Do not substitute bacitracin.

*b* May cause skin and eye irritation. Apply for 5 minutes and then rinse thoroughly.

*c* Do not use rifampin alone because resistance may develop. It can cause an orange color of urine and tears and stain contact lenses. Adjust the dose for patients with renal impairment. Do not use in those with impaired hepatic function.

CA-MRSA, Community-acquired methicillin-resistant Staphylococcus aureus.

Box 37.2 Strategies to Eradicate CA-MRSA Carrier States in Patients With Recurrent CA-MRSA Soft Tissue Infections

decolonization. In the case of multiple recurrent infections, consider consulting an infectious disease expert.

**I&D PROCEDURE**

**Procedure Setting**
Definitive I&D of soft tissue abscesses can be performed in either the ED or the operating room (OR). When abscesses are drained in the ED, some centers prefer to use a special area to avoid contamination of general treatment rooms, but protocols vary greatly (Box 37.3).

The choice of the locale for the procedure depends on a number of important factors. The location of the abscess may dictate management in the OR. Large abscesses or abscesses located deep in soft tissues require a procedure involving a great degree of patient cooperation, which might be achieved only under general or regional anesthesia. Proximity to major neurovascular structures, such as in the axillae or antecubital fossa, may necessitate specific management. Infections of the hand (with the exception of distal finger infections) have traditionally been managed in the OR because of the many important structures involved and the propensity for limb-threatening complications.

Lack of adequate anesthesia is the most common factor limiting I&D in the ED. The current increased use of ED procedural sedation (see Chapter 33) has changed previous OR cases to ones that can be managed well in the ED. If the clinician believes that the abscess cannot be fully incised and drained because of inadequate anesthesia, the patient should be taken to the OR for management under general anesthesia. In addition to limiting proper drainage, it is inhumane and unethical to subject a patient to extreme pain when alternatives are available.

**Equipment and Anesthesia**

A standard suture tray provides adequate instruments if a scalpel and packing material are added (Fig. 37.5). Although sterility is impossible during the procedure, one should avoid contamination of surrounding tissue. Some clinicians prefer to use an obligatory skin scrub with an antiseptic solution, but the value of this step is dubious. Most clinicians use nonsterile gloves while draining pus, but practices vary.

It is often quite difficult to achieve local anesthesia by direct infiltration because of the poor function of local anesthetic agents in the low pH of infected tissue. Furthermore, distention of sensitive structures by a local injection is quite painful and hence poorly tolerated by most patients. Skin anesthesia is usually possible, but total anesthesia of the abscess cavity itself cannot generally be achieved. If a regional block can be performed (see Chapters 29-32), this type of anesthesia is preferred. Alternatively, a field block may be used. It should be noted that infected tissue is very vascular and local anesthetics are therefore absorbed quickly. Strict adherence to maximum safe doses of local anesthetics is required.

The skin over the dome of an abscess is often quite thin, thus making skin anesthesia difficult. If a 25-gauge needle is used carefully, one can frequently inject the dome of the abscess with a local anesthetic.
Historically, drainage of an abscess has involved the creation of a large skin incision to ensure adequate initial drainage of purulent material, followed by filling the abscess cavity with packing to promote continued drainage. The loop drain technique, an old technique now undergoing renewed interest, confers several advantages over the classic I&D. There are no packing changes, smaller incisions result in better cosmesis, and incisions drain as long as the loop is in place. This technique, though perhaps counterintuitive to classic teaching, provides excellent results for abscesses that previously would have been subjected to traditional incision, drainage, packing, and repacking protocols.

**WHAT TYPE OF DRAIN TO USE**

The original article describing this technique used a silicone vessel loop, a small Penrose drain, or a sterile rubber band. A readily available product is a vessel loop used by surgeons (Fig. 37.13, step 1).

**TECHNIQUE**

**Step 1:** Make a small (5–10 mm) incision with a No. 11 scalpel blade at the periphery of the abscess but still within its borders (see Fig. 37.13, step 2). Make the first incision where the abscess is pointing or already draining. Then probe the cavity with a hemostat and break up loculations (see Fig. 37.13, step 4). Pus should flow out of this puncture incision.

**Step 2:** With the hemostat inside the abscess cavity, find the opposite edge of the abscess cavity and position the tip of a hemostat underneath the area where you will make the second incision (see Fig. 37.13, step 5). Tent the skin from underneath. Make another stab with a No. 11 scalpel blade over the hemostat tip. Slide the tip of the hemostat through the new incision. At this stage, you should have two small incisions with a hemostat going into one, tunneling through the abscess cavity, and coming out of the other incision. There should be a maximum separation between the two incisions of up to 4 cm. If the abscess is larger than this, you can place more loops, depending on the shape of the abscess cavity.

**Step 3:** Using the hemostat that has traversed the cavity, grab the end of the loop drain and pull it back through the wound (see Fig. 37.13, steps 7 and 8). There should be a long end of the loop drain exiting each incision.

**Step 4:** Tie the two ends of the drain together without tension. Too much tension is painful and can necrose the skin between the incisions. One way to avoid excessive skin tension is to place a syringe between the skin and the drain while you are tying it (see Fig. 37.13, step 9). Tie the knots flush with the syringe; when the syringe is pulled away there will be a loosely placed loop that will exert no tension on the incision sites. A finger should be able to be passed between the skin and loop. The drain is going to be slippery, so make about five knots. Tie the last two very tightly by stretching the loop almost to the breaking point. Trim the ends of the drain (see Fig. 37.13, step 10). An option at this point is to irrigate the abscess cavity through the stab incision. Cover the operative site with gauze as drainage will continue.

**DISCHARGE INSTRUCTIONS**

With the dressing removed and the loop still in place, the patient should bathe or shower twice a day for the first 3 days to promote continued drainage. Change the dressing at least twice daily or whenever saturated. Gently pull the loop back and forth once or twice a day to help keep the wound open.

**WHEN TO REMOVE THE DRAIN**

Remove the drain when the discharge stops and cellulitis improves. Simply cut the loop and pull it out. The loop is generally removed in approximately 5–10 days but may be removed sooner with a small abscesses or rapid healing. The patient now has only two small puncture sites that will heal over rather than a large scar that must heal by secondary intention.


**Incision**

One should make all incisions conform with skin creases or natural folds to minimize visible scar formation (Fig. 37.7). Care should be taken in areas such as the groin, the posterior aspect of the knee, the antecubital fossa, and the neck so that vascular and neural structures are not damaged.

A No. 11 or 15 scalpel blade, held perpendicular to the skin, is used to nick the skin over the fluctuant area, and then a simple linear incision is carried the total length of the abscess cavity (see Fig. 37.6, step 3). This will afford more complete drainage and facilitate subsequent breakup of loculations. Attempting to drain an abscess with an inadequate incision is counterproductive and makes packing changes more difficult.

**subcutaneously. Without moving the tip of the needle, the anesthetic solution spreads over the dome through the subcutaneous layers into the surrounding skin and provides excellent skin anesthesia. If the needle is in the proper plane (best accomplished by holding the syringe parallel rather than perpendicular to the skin), the surrounding skin blanches symmetrically during infiltration without having to reposition the needle (Fig. 37.6, step 2). In an extremely anxious or uncomfortable patient, judicious use of preoperative sedation (see Chapter 33) with IV opioids and sedatives or with nitrous oxide makes the procedure easier for both the patient and clinician. Ketamine, propofol, or a combination of these drugs is a popular option in the ED setting.

Some clinicians recommend the use of topical ethyl chloride or Fluori-Methane spray (Gebauer Company, Cleveland, OH) for the initial skin incision, but although this is an attractive concept to patients, the pain relief offered by these agents is variable and fleeting. Ethyl chloride is also highly flammable. These vapocoolant sprays may be useful to provide momentary anesthesia for injection of a local anesthetic or for the initial skin incision if the injection or incision is made immediately after blotching of the skin. In general, however, these agents are of minimal benefit as a stand-alone anesthetic agent for all but the smallest of superficial abscesses (e.g., purulent folliculitis).**
Identify and confirm the location of the abscess. This patient has a pilonidal abscess, which is found at the superior portion of the gluteal cleft.

Anesthetize the region. Use a 25-gauge needle to inject the dome of the abscess. Hold the needle parallel to the skin during injection. Blanching will occur as the anesthetic spreads.

Use an 11-blade scalpel to make a linear incision over the total length of the abscess cavity. If possible, the incision should conform to skin creases or natural folds.

Culture the purulent drainage.

Probe the depth of the abscess and use a hemostat to break open loculations. This can be painful; inject additional anesthetic if needed through the cut skin edges and into the deeper tissues.

Irrigate the abscess cavity to assist with removal of residual debris.

Loosely pack the abscess cavity with gauze wick. The purpose of the packing is to keep the incision open, which allows continued drainage of the cavity. Avoid overpacking the abscess.

Place an absorbent gauze dressing over the packed abscess. Arrange for follow-up care, which usually entails a repeat exam in 1 to 3 days.

Figure 37.6 Incision and drainage. Packing the cavity with gauze is a common intervention but this procedure has not been proven to be of significant value. Routine culturing and irrigation are optional and left to the discretion of the clinician.
Exceptions regarding aggressive incision are abscesses in cosmetic areas, in areas under significant skin tension (e.g., extensor surfaces), and in areas with extensive scar tissue (e.g., sites of multiple previous drainage procedures). In these special circumstances, a stab incision or simple aspiration alone may be attempted initially, with the goal of limiting tissue injury and resultant scar formation. Use of this less aggressive approach requires that the patient be counseled that multiple decompressions (e.g., via needle aspiration) or delayed aggressive I&D may be required. The abscess will need to be reassessed in 24 to 48 hours to determine whether additional intervention is needed.

**Wound Dissection**

Following a standard incision, the operator should probe the depth of an abscess to assess its extent and ensure proper drainage by breaking open loculations (see Fig. 37.6, step 5). An ideal instrument for this procedure is a hemostat, optionally wrapped in gauze (or a cotton swab for small abscesses), which is placed into the abscess cavity and spread and manipulated throughout the cavity (Fig. 37.9). Traditionally, the operator's gloved finger has been suggested as an ideal way to assess the depth of the abscess cavity and to break up loculations, but this is a potentially dangerous practice that should be avoided unless it is certain that the abscess contains no sharp foreign body. Of particular concern is an abscess caused by skin-popping of IV drugs. These abscesses occasionally harbor broken needle fragments (see Fig. 37.9D and E). In addition, patients who engage in this practice have a high incidence of hepatitis and HIV infection. Clinicians are often surprised at the depth or extent of abscesses discovered during probing. Sharp curettage of the abscess cavity is not usually required and may produce bacteremia. Although tissue probing is generally the most painful aspect of the technique and total local anesthesia is difficult to attain, this portion of the procedure should not be abbreviated. If pain persists, additional local anesthetic can be administered through the cut skin edges and into deeper tissues to provide additional anesthesia (Fig. 37.10A). If the procedure is limited because of pain, use of appropriate analgesia or anesthesia is mandated. Failure to adequately pack the abscess on the first visit makes follow-up packing changes more problematic.

A blunt-ended suction device can be used to extract copious pus from large or deep abscesses while also assisting in breakup of loculations (see Fig. 37.10B).

**Wound Irrigation**

Following the breakup of loculations, some clinicians advocate copious irrigation of the abscess cavity with normal saline to ensure adequate removal of debris from the wound cavity (see Fig. 37.6, step 6). Though seemingly beneficial, irrigation of the abscess cavity has not been experimentally demonstrated to significantly augment healing or affect outcome, and hence may be eschewed. Hyperemic tissue may bleed profusely, but the bleeding usually stops in a few minutes if packing is used. Abscesses of the extremities can be drained with the use of a tourniquet to provide a bloodless field.

**Packing and Dressing**

Multiple studies, most from countries outside the United States, suggest that primary suture closure of incised and drained
When used, a loose packing of gauze or other material is placed gently into the abscess cavity to prevent the wound margins from closing and to afford continued drainage of any exudative material (see Fig. 37.6, step 7). The packing material should make contact with the cavity wall so that on removal, gentle débridement of necrotic tissue will occur spontaneously. A common error is to attempt to pack an abscess too tightly with excessive packing material. In essence, the pack merely keeps the incision open, and its main purpose is not to absorb all drainage as a dressing accomplishes this goal. Care must be exercised to ensure that the packing does not exert significant pressure against the exposed tissue and lead to further tissue necrosis. Some prefer to use plain gauze, some use gauze soaked in saline or povidone-iodine, and some use gauze impregnated with iodine (iodoform). For large abscess cavities, gauze pads (without cotton backing) are ideal packing material (Fig. 37.11). If gauze pads are used, the number of pads placed in the wound should be counted and charted. Ideally, the corner of each pad should exit from the wound. The clinician must ensure that all gauze pads will be removed when the packing is changed or discontinued. More commonly, thin (0.6 to 1.2 cm) packing strip gauze, either plain or iodoform, is used. The iodoform gauze may sting the patient for a few minutes after it is inserted. Packing, especially packing strips containing iodine, will be radiopaque on a plain radiograph. If a foreign body is considered, a radiograph should be obtained before packing.

When used, a loose packing of gauze or other material is placed gently into the abscess cavity to prevent the wound margins from closing and to afford continued drainage of any exudative material (see Fig. 37.6, step 7). The packing material should make contact with the cavity wall so that on removal, gentle débridement of necrotic tissue will occur spontaneously. A common error is to attempt to pack an abscess too tightly with excessive packing material. In essence, the pack merely keeps the incision open, and its main purpose is not to absorb all drainage as a dressing accomplishes this goal. Care must be exercised to ensure that the packing does not exert significant pressure against the exposed tissue and lead to further tissue necrosis. Some prefer to use plain gauze, some use gauze soaked in saline or povidone-iodine, and some use gauze impregnated with iodine (iodoform). For large abscess cavities, gauze pads (without cotton backing) are ideal packing material (Fig. 37.11).

Abscesses results in faster healing than after packing and secondary closure, and results in similarly low abscess recurrence rates. Many of the reported cases involved abscesses in the anogenital region, a site teeming with bacteria. Acute superficial abscesses have been managed with incision, curettage, and primary suture closure without antibiotics or packing, which is safe and cost-effective. This clinical approach is not currently in wide practice in the United States.

Although the specific technique and the clinical value of routine packing of abscesses have not been well studied, packing is a traditional intervention and is often performed in the ED. Some, however, advocate that packing is neither needed nor beneficial for easily drained abscesses, and the intervention adds to cost and patient discomfort. Although most studies evaluating the necessity of wound packing have been small, there does not appear to be evidence of increased recurrence of abscesses when not packed. There is no proven benefit to routine packing of drained cutaneous abscesses as it does not appear to prevent recurrence of abscesses or to decrease further intervention at follow-up.

As noted earlier, some advocate curettage and irrigation of the abscess cavity and then primary suture closure, without packing. Overall, it appears reasonable to avoid packing of small, easily drained abscesses. The loop drainage technique described hereafter is evidence that formal packing is not a prerequisite for abscess healing. Hence, clinical judgment, common sense, and individualized treatment based on the particular scenario should prevail. Simply stated, no packing recommendation is universally accepted, but the procedure is likely overused.

When used, a loose packing of gauze or other material is placed gently into the abscess cavity to prevent the wound margins from closing and to afford continued drainage of any exudative material (see Fig. 37.6, step 7). The packing material should make contact with the cavity wall so that on removal, gentle débridement of necrotic tissue will occur spontaneously. A common error is to attempt to pack an abscess too tightly with excessive packing material. In essence, the pack merely keeps the incision open, and its main purpose is not to absorb all drainage as a dressing accomplishes this goal. Care must be exercised to ensure that the packing does not exert significant pressure against the exposed tissue and lead to further tissue necrosis. Some prefer to use plain gauze, some use gauze soaked in saline or povidone-iodine, and some use gauze impregnated with iodine (iodoform). For large abscess cavities, gauze pads (without cotton backing) are ideal packing material (Fig. 37.11). If gauze pads are used, the number of pads placed in the wound should be counted and charted. Ideally, the corner of each pad should exit from the wound. The clinician must ensure that all gauze pads will be removed when the packing is changed or discontinued. More commonly, thin (0.6 to 1.2 cm) packing strip gauze, either plain or iodoform, is used. The iodoform gauze may sting the patient for a few minutes after it is inserted. Packing, especially packing strips containing iodine, will be radiopaque on a plain radiograph. If a foreign body is considered, a radiograph should be obtained before packing. The value of antibiotic-impregnated gauze is uncertain.

An absorbent gauze dressing should be placed over the packed abscess, or if an extremity is involved, a lightly wrapped circumferential dressing should be used (see Fig. 37.6, step 8).
hours after the procedure, with the first but possibly the only packing change occurring at this time. Some wounds warrant closer monitoring. Diabetic patients or other patients with impaired healing capacity, mental impairment, or physical disabilities may require a home care nurse or hospital admission for more frequent wound care and packing changes. Wounds that are at high risk for complications, such as those about the face or hands or those with significant cellulitis, require close follow-up depending on the specific scenario. The patient should be encouraged to play an active role in wound care. During the first follow-up visit, compliant and able patients should be taught to change the packing and dressing. If this is anatomically impossible, a friend or family member can be instructed in the technique. If long-term packing or complicated wound care is required, referral to wound care centers rather than multiple repeated ED visits may be more practical.

The technique for changing packing material is usually one of personal preference. It should be emphasized that patients often fear a repeated visit and expect significant pain with subsequent wound care, especially if the initial I&D was difficult. Therefore the specifics of packing change should be addressed

The technique for changing packing material is usually one of personal preference. It should be emphasized that patients often fear a repeated visit and expect significant pain with subsequent wound care, especially if the initial I&D was difficult. Therefore the specifics of packing change should be addressed

FOLLOW-UP EXAMINATION

A drained abscess should be reevaluated in 1 to 3 days, depending on a number of factors. Most lesions are reevaluated 48 hours after the procedure, with the first but possibly the only packing change occurring at this time. Some wounds warrant closer monitoring. Diabetic patients or other patients with impaired healing capacity, mental impairment, or physical disabilities may require a home care nurse or hospital admission for more frequent wound care and packing changes. Wounds that are at high risk for complications, such as those about the face or hands or those with significant cellulitis, require close follow-up depending on the specific scenario. The patient should be encouraged to play an active role in wound care. During the first follow-up visit, compliant and able patients should be taught to change the packing and dressing. If this is anatomically impossible, a friend or family member can be instructed in the technique. If long-term packing or complicated wound care is required, referral to wound care centers rather than multiple repeated ED visits may be more practical.

The technique for changing packing material is usually one of personal preference. It should be emphasized that patients often fear a repeated visit and expect significant pain with subsequent wound care, especially if the initial I&D was difficult. Therefore the specifics of packing change should be addressed

FOLLOW-UP EXAMINATION

A drained abscess should be reevaluated in 1 to 3 days, depending on a number of factors. Most lesions are reevaluated 48 hours after the procedure, with the first but possibly the only packing change occurring at this time. Some wounds warrant closer monitoring. Diabetic patients or other patients with impaired healing capacity, mental impairment, or physical disabilities may require a home care nurse or hospital admission for more frequent wound care and packing changes. Wounds that are at high risk for complications, such as those about the face or hands or those with significant cellulitis, require close follow-up depending on the specific scenario. The patient should be encouraged to play an active role in wound care. During the first follow-up visit, compliant and able patients should be taught to change the packing and dressing. If this is anatomically impossible, a friend or family member can be instructed in the technique. If long-term packing or complicated wound care is required, referral to wound care centers rather than multiple repeated ED visits may be more practical.

The technique for changing packing material is usually one of personal preference. It should be emphasized that patients often fear a repeated visit and expect significant pain with subsequent wound care, especially if the initial I&D was difficult. Therefore the specifics of packing change should be addressed

FOLLOW-UP EXAMINATION

A drained abscess should be reevaluated in 1 to 3 days, depending on a number of factors. Most lesions are reevaluated 48 hours after the procedure, with the first but possibly the only packing change occurring at this time. Some wounds warrant closer monitoring. Diabetic patients or other patients with impaired healing capacity, mental impairment, or physical disabilities may require a home care nurse or hospital admission for more frequent wound care and packing changes. Wounds that are at high risk for complications, such as those about the face or hands or those with significant cellulitis, require close follow-up depending on the specific scenario. The patient should be encouraged to play an active role in wound care. During the first follow-up visit, compliant and able patients should be taught to change the packing and dressing. If this is anatomically impossible, a friend or family member can be instructed in the technique. If long-term packing or complicated wound care is required, referral to wound care centers rather than multiple repeated ED visits may be more practical.

The technique for changing packing material is usually one of personal preference. It should be emphasized that patients often fear a repeated visit and expect significant pain with subsequent wound care, especially if the initial I&D was difficult. Therefore the specifics of packing change should be addressed
is present, packing may be discontinued. The patient should then be instructed to begin warm soaks of the wound. Gentle hydrostatic débridement may be performed by the patient in the shower at home: the patient holds the skin incision open and directs the shower or faucet spray into the abscess cavity. Appropriate dressing changes should then follow until healing is completed. When all signs of infection (e.g., erythema, drainage, pain, and induration) have resolved and healthy granulation tissue is present, the patient may be discharged from medical care.

**SPECIFIC ABSCESS THERAPY**

**Folliculitis, Furuncles, and Carbuncles**

Folliculitis, furuncles, and carbuncles are common ED presentations in healthy patients with no specific predisposing risk factors, due to chronic colonization with a *Staphylococcus* bacterium. Those in close contact with infected individuals are also at increased risk for similar infections. The pathogenesis of staphylococcal disease is a complex host-bacterium interaction. *S. aureus* can invade the skin by way of a hair follicle or any open wound. This can lead to destruction of the local tissue followed by vasodilation of the blood vessels, increasing the blood flow to the affected area, which is called hyperemia. Subsequently, an exudative reaction occurs, with polymorphonuclear cell invasion. The process will extend, following the path of least resistance, and form an abscess. The abscess may then extend and form sinus tracts. Eventually dissemination may occur through invasion of blood vessels, leading to infection of other organ systems including osteomyelitis, meningitis, and endocarditis.7,74

Inflammation of a hair follicle caused by bacterial invasion is known as folliculitis. Folliculitis is a common inflammatory skin syndrome of the young and middle-aged, but can be observed in all age groups. The underlying pathogenic mechanisms are unclear, but microbial involvement has been suggested.7 Staphylococci, especially *S. aureus*, are mainly cultured from inflamed lesions and pustules76–78 but fungal involvement (e.g., Malassezia species) has also been documented.79–81

*Pseudomonas aeruginosa* is a rare cause of folliculitis. *Pseudomonas* folliculitis can develop after contact with contaminated water from swimming pools, hot tubs, and spa baths.82

Fungal folliculitis may be seen solely or in conjunction with *Staphylococcus* in individuals who use antibacterial antibiotics, oral corticosteroids, immunosuppressive treatments, topical steroids, have hyperhidrosis, or are obese.83 Treatment generally consists of warm compresses to the affected area and antimicrobial soaps. If multiple sites are involved systemic antibiotics may be required. Recurrence may warrant repeat treatment along with chlorhexidine baths and bacitracin to the nares.

Furuncles, or “boils,” are acute circumscribed abscesses of the skin and subcutaneous tissue that usually involve the hair follicles (Fig. 37.14). They are most frequently seen on the face, neck, buttocks, thigh, perineum, breast, and axilla. Carbuncles are groups of interconnected furuncles, most frequently seen at the base of the neck where the skin is thickest. The thickness of the skin leads to lateral extension. *S. aureus* is the most common organism seen. Large carbuncles can cause systemic symptoms and complications. A common risk factor for carbuncles is diabetes. These large carbuncles often

---

Figure 37.12 Wound check. A, Packing gauze is removed from the abscess on the first wound inspection. If drainage is present, the abscess may be repacked. B, Small abscesses can be cleaned with a cotton swab swirled in the cavity, which is left open. This care can be continued at home.
Vessel Loop Method of Incision and Drainage

Instead of traditional gauze packing, a sterile silicone loop is used. This is commonly referred to simply as a “vessel loop” and is readily available from the operating room (this device is used by vascular surgeons).

After sterile preparation and anesthetic infiltration, make a small (5–10 mm) incision at the periphery of the abscess. If possible, make the incision where the abscess is already pointing, noted by the white arrow.

Express as much pus as possible from the incision. Obtain cultures if clinically indicated.

Use a hemostat to probe the abscess cavity and break up loculations.

Optionally irrigate the cavity with a syringe and plastic intravenous catheter.

Probe with the hemostat to find the opposite edge of the abscess cavity. Position the tip of the hemostat underneath the area of the second incision, tent the skin, and make another stab incision over the hemostat tip.

Place the hemostat through the cavity so that it exits the second incision. Grab the end of the vessel loop (arrow).

Pull the vessel loop through the abscess cavity so that it exits through the first incision.

Tie the two ends of the vessel loop together. To avoid excessive skin tension, tie the loops over a syringe. Tie at least five knots, as the loop material is slippery. Tie the last two knots tightly, by stretching the loop almost to the breaking point.

Remove the syringe, and then trim the ends of the vessel loop. Note that it is not tied tightly against the skin. Refer to the text for information on discharge instructions and loop removal.

Hidradenitis Suppurativa

HS is a chronic, relapsing, inflammatory follicular occlusive disease process that affects the apocrine sweat glands in the axilla, the inguinal region, and the perineum. Its prevalence is 1% to 4% in industrialized countries; most affected individuals are young women. The condition results from follicular inflammation and subsequent occlusion of the apocrine ducts by keratinous debris, which leads to ductal dilation, inflammation, and rupture into the subcutaneous area. The clinical manifestations vary, ranging from recurrent inflamed nodules and abscesses to draining sinus tracts and bands of severe scar formation. The associated pain, malodor, drainage, and disfigurement that accompany HS contribute to a profound psychosocial impact of the disease on many patients. Dysregulation of the immune response is thought to play a role, with some studies showing a response to immunosuppressive agents. Occlusion of the hair folicle predisposes to secondary bacterial infection and subsequent abscess formation. In its early stages, it is indistinguishable from a simple furuncle. Progression and recurrence, however, lead to the distinctive appearance of multiple foci coupled with areas of induration and inflammation that are in various stages of healing. This chronic process can create draining fistulous tracts, often involving large areas that are not amenable to simple I&D procedures (see Fig. 37.15). Genetic factors appear to be an important contributor in HS. A hormonal impact has been suggested by observations requiring widespread surgical excision rather than simple I&D. Systemic therapy may be indicated in patients with widespread lesions, systemic symptoms, or in immunosuppressed patients. Pedicure-associated nontuberculous mycobacteria has been reported. This is believed to be associated with the foot baths used and suboptimal cleaning methods.

Most cases of recurrent staphylococcal skin infections are caused by autoinfection from skin lesions or nasal reservoirs. Treatment is directed at prevention, through eliminating the organisms. This can be accomplished by the application of bacitracin to the nares, chlorhexidine baths, and good hygiene. If unsuccessful, then systemic oral antistaphylococcal treatment is required, usually for 2 to 3 weeks. Close contacts often require treatment as well.

S. aureus can cause suture abscesses. A suture abscess is often misdiagnosed as a wound infection, but in fact it is a local nidus of inflammation, infection, or both caused and potentiated by suture material. Such an abscess usually appears after sutures have been in place for 3 to 5 days, with single or multiple discrete areas of erythema and tenderness being noted at the site where the suture penetrates the skin. Simply removing the suture (a drop of pus may be expressed) and providing warm compresses and topical antibiotic ointment is generally all that is required. Wide opening of the wound and systemic antibiotics are seldom required. When the suture is buried, a small incision should be followed by probing the wound with a small hook or bent needle (see Chapter 35) tosnare the suture for removal.

Figure 37.14 A, Staphylococcus aureus is a common cause of folliculitis, although other bacteria such as Pseudomonas may be responsible. Treatment consists primarily of local measures such as warm compresses and antibacterial soaps and ointments. B, Furuncles are circumscribed abscesses of the skin and subcutaneous tissue that usually involve a hair follicle. C, A carbuncle is a complicated abscess on the nape of the neck. It is very common in diabetics. Carbuncles are usually caused by S. aureus, including community-acquired methicillin-resistant S. aureus. Because of its many crypts, loculations, and intercommunicating small abscesses, treatment by simple incision and drainage is often not readily curative. When fluctuance is appreciated, incision is indicated. One should avoid multiple small incisions because tissue circulation may be compromised. Antibiotics may augment healing of this abscess, but wide surgical excision may be required. (A, from Weston WL, Lane AT, Morelli JG, editors: Color Textbook of Pediatric Dermatology, ed 4, St. Louis, 2007, Mosby; B, from Long SS, Pickering LK, Prober CG, et al: Principles and Practice of Pediatric Infectious Diseases Revised Reprint, ed 3, Philadelphia, 2009, Churchill Livingstone.)
Bacterial cultures of specimens from hidradenitis suppurativa lesions are frequently sterile or reflect organisms seen in other soft tissue abscesses. *Staphylococcus* is the most commonly isolated organism, with *E. coli* and β-hemolytic streptococci being other important pathogens. In the perineal region, enteric flora are often found. Many of these abscesses have multiple isolates, and anaerobic bacteria are frequently cultured. CA-MRSA is an increasing cause of such abscesses.

Clinical manifestations may be influenced by gender. Primary sites of involvement in women are the groin, upper inner thigh, axilla, chest, and the buttocks or gluteal clefts. In men, primary sites of involvement are the groin or thigh, axilla, perineal or perianal regions, and buttocks or gluteal cleft. Both sexes regularly show involvement in nonintertriginous skin, particularly at sites of skin compression and friction. Beltlines, waistbands, and brassiere straps or bands are also common locations.

Multiple disorders have been associated with HS including diabetes, obesity, dyslipidemia, hypertension, cardiovascular disease, Crohn’s disease, and acne vulgaris. Long-standing, poorly controlled HS may lead to significant physical and emotional consequences such as strictures and contractures, lymphatic obstruction, lymphedema of limbs and genitalia, weakness, depression, suicide, and long-term effects of chronic suppuration including anemia, hypoproteinemia, amyloidosis, infectious complications (lumbosacral epidural abscess, sacral bacterial osteomyelitis), arthritis, squamous cell carcinoma, and anemia.

Multiple therapeutic approaches exist for the management of HS. Any fluctuant area requires drainage, as described in the section on I&D Procedure. In patients with extensive cellulitis, a broad-spectrum, antistaphylococcal antibiotic should be used. Unfortunately, HS cannot be cured with localized I&D and/or antibiotics. If I&D is required for relief of a secondary abscess, it should be made clear to the patient that this procedure does not cure the underlying disease. The chronic nature of the disease produces multiple areas of inflammation and subcutaneous fistulous tracts that induce routine recurrences. The patient must be informed of this unfavorable prognosis and should be referred to a dermatologist or surgeon for long-term care.

Milder forms of the disease are initially treated with conservative measures. Many different approaches have been tried, with only limited degrees of success. Few controlled studies of treatment strategies have been performed. Patients are often counseled to lose weight, stop smoking, avoid skin trauma, wear light clothing, refrain from shaving, stop using scented deodorants, and improve personal hygiene. The benefits of these efforts are unknown. Topical clindamycin is often utilized as a first-line therapy for mild HS. A randomized 3-month trial supported its efficacy for mild inflammatory lesions and demonstrated the relative safety and tolerability of this regimen. Oral antistaphylococcal antibiotics are most commonly used, with varying results. Retrospective reviews looking at the efficacy of combination therapy (clindamycin and rifampin) have been promising, with some patients experiencing long-term remission, although diarrhea is a common side effect. There have been reports of success with topical clindamycin, hormone therapy in females, and laser therapy. Finally, immunosuppressive therapy with tumor necrosis factor-α inhibitors (infliximab, adalimumab, and etanercept) has been successful in treating a few intractable cases.

**Figure 37.15** A, Axillary abscesses are common and recurrent. Community-acquired methicillin-resistant *Staphylococcus aureus* is an increasing cause. B, Vessel loop drainage of an axillary abscess. C, Hidradenitis suppurativa of the groin or axilla is a complicated series of abscesses that may not be amenable to simple incision and drainage. In this patient with involvement of the groin, extensive surgery was required to excise recurrent infection.
Advanced stages of the disease are managed by wide or local excision. Recurrence is more likely with smaller excisions than wider excisions. Delayed closure may offer better results, but this has not been directly compared with primary closure. Wide excision may warrant skin grafting.

**Breast Abscess**

Breast abscesses are localized collections of pus in the breast tissue. They can occur in lactating (14%) and nonlactating breasts (86%). They occur more commonly in African Americans, obese individuals, and smokers. They are caused by normal skin bacteria that enter through a break or crack in the skin, usually the nipple (Fig. 37.16). The infection becomes established in parenchymal tissue and causes pain, swelling, and localized hyperthermia. In its early stages, when cellulitis predominates, a breast abscess can be difficult to diagnose. In equivocal cases, antibiotics may be curative. Cellulitis may progress to frank abscess formation. Women with a significant breast abscess can be quite ill and appear toxic.

The estimated incidence of inflammatory processes of the breast (mastitis) in lactating women ranges from 2% to 33%. In lactating women the infection is usually precipitated by milk stasis after weaning or missed feedings and usually occurs in the first 6 weeks of breastfeeding or during the weaning phase. The cause is usually bacterial invasion through a cracked or abraded nipple by *S. aureus*, including MRSA, or streptococci originating from the mouth of the nursing child. Manifestations include redness, heat, pain, fever, and chills. Treatment consists of antistaphylococcal antibiotics, continued breast emptying with a breast pump, and application of heat. During treatment it is important to encourage continued breast emptying to promote drainage. Nursing can be continued with the noninfected breast.

Breast abscesses in nonlactating women are more common. People at risk are smokers and diabetics. These are also risk factors for recurrence. In addition, abscesses may be a complication of nipple piercing or breast implants inserted under appropriate medical care, as well as via illicit cosmetic procedures. Most primary breast abscesses are caused by *S. aureus*, with MRSA becoming increasingly prominent.

Less prominent pathogens include strains of *Staphylococcus pyogenes*, *E. coli*, *Bacteroides*, *Corynebacterium* species, coagulase-negative staphylococci (e.g., *Staphylococcus lugdunensis*), *P. aeruginosa*, *Proteus mirabilis*, and anaerobes. Mixed flora are more commonly found in recurrent abscesses.

In the setting of nonsevere infection, in the absence of risk factors for MRSA, outpatient therapy may be initiated with dicloxacillin or cephalaxin, pending culture results. In the setting of β-lactam hypersensitivity, clindamycin may be used. If the community prevalence of MRSA is high, treatment with clindamycin or TMP-SMX should be considered. In the setting of severe infection, empiric inpatient therapy with vancomycin should be initiated.

Ultrasound-guided needle aspiration, as opposed to formal I&D, is becoming the standard of care for most breast abscesses. Multiple aspirations may be required. When repeat aspirations are required, hospitalization may be the best way to facilitate treatment. When compared with I&D, aspiration causes less scarring, does not interfere with breastfeeding, and does not require procedural sedation. Christensen and associates recommended that ultrasound-guided needle drainage replace surgery as first-line treatment of uncomplicated puerperal and nonpuerperal breast abscesses. Emerging treatment parameters are ultrasound-guided needle aspiration for abscesses less than 3 cm in diameter and ultrasound-guided catheter drainage for abscesses 3 cm in diameter or larger. Ultrasound images of breast abscesses appear as inhomogeneous, hyperchoic masses (Fig. 37.17). Repeat aspiration daily or every other day helps to achieve complete resolution, with a mean of 3.5 aspirations being required. Surgical drainage is warranted in the setting of compromise to the overlying skin and in cases in which the abscess is not responding to needle aspiration and antibiotics.

Recurrent abscesses are a common, troublesome complication after traditional treatment with I&D and antibiotics. Fortunately, the reported recurrence rate associated with ultrasound-guided aspiration or drainage procedures is quite low. Patients with persistent recurrences should be managed by a surgeon for total excision of the involved area.

Although a breast abscess is rarely a harbinger of malignancy, it could be the initial manifestation of a metastatic process. After needle aspiration or catheter drainage, send the aspirate for culture and cytologic examination, and perform a postdrainage mammogram and ultrasound to evaluate for underlying or coexistent malignancy.

A breast abscess in a male is an unusual occurrence, so consider malignancy and underlying bone or joint infection in these cases (see Fig. 37.2A and B).

**Pilonidal Abscess**

Pilonidal sinuses are common malformations that occur in the sacrococcygeal area (Fig. 37.18). The cause of the sinus formation is unclear, and the malformation may occur during embryogenesis. Although pilonidal sinuses are present from birth, they are not usually manifested clinically until adolescence or the early adult years. Pilonidal cyst formation is thought to be secondary to blockage of a pilonidal sinus. The blockage is most commonly the result of hair in the region, and the lesion may in part be a foreign body (hair) granuloma. The result of this obstruction is repeated soft tissue infection,
As a person sits or bends, the natal cleft stretches, traumatizing hair follicles and thus creating pores. The pores collect debris and serve as a fertile environment for roots of hairs to lodge and become embedded. As the skin is drawn taut over the natal cleft with movement, negative pressure is created in the subcutaneous space drawing hair deeper into the pore, and the friction causes the hairs to form a sinus. Pilonidal disease typically develops in people with deep natal clefts. Subsequently, hairs become ingrown, bacteria invade, and an abscess develops in the sacrococcygeal region. At the current time, CA-MRSA infections are not common.

Patient presentation is highly variable. They may be completely asymptomatic with just a pilonidal cavity and sinus, or present with acute infection and drainage, or even chronic inflammation with drainage. There may even be multiple areas of drainage. Patients with a pilonidal abscess will seek care for back pain and local tenderness. On examination the area is found to be indurated. Frank abscess formation may not be appreciated as sometimes the abscess is deep. One will usually see barely perceptible dimples or tiny openings at the rostral end of the gluteal crease (Fig. 37.19). A hair or slight discharge may be noticed at the opening. The discharge will usually be purulent or bloody. One may find a more caudal cyst or abscess, possibly with a palpable sinus tract connecting the two. The sinus and cyst may be draining chronically, or they may become infected as the size increases and blockage occurs.147

Treatment of an acutely infected cyst is the same as previously discussed for any fluctuant abscess: all hair and pus should be removed, and the lesion should be packed (see Figs. 37.6 and 37.13). The abscess cavity may be quite large and thus necessitate a lengthy incision to ensure complete drainage. The incision should be made laterally, off-midline, or over the area of maximal fluctuance. A retrospective study looked at 48 patients who had undergone lateral incisions and matched them with 48 patients who had undergone midline incisions. Those who had undergone midline incisions took approximately 3 weeks longer to heal than those drained with a lateral incision.148 In general, reported median healing times after I&D vary between 12 and 63 days.149,150 The area may be repacked followed by drainage and partial resolution, with eventual reaccumulation. Pilonidal abscess formation most frequently affects young adults with a 2:1 male to female ratio. The sinuses and cysts are lined with stratified squamous epithelium and, after excision, may be found to contain wads of hair and debris. When cultured, pilonidal abscesses generally yield mixed fecal flora with a preponderance of anaerobes.143 The specific mechanism for the development of pilonidal disease is unclear, although hair and inflammation are contributing factors.144–146

Figure 37.17 Longitudinal images of ultrasound-guided drainage of an abscess in the lower inner quadrant of a 27-year-old woman’s right breast 8 weeks after delivery. A, Well-defined, slightly loculated, inhomogeneous hypoechoic abscess measuring 6 cm x 4 cm. Dots extend across the abscess, with the cursor at one margin of the abscess. B, Pigtail catheter (7 Fr) containing a trocar (arrows) in the abscess cavity. C, After removal of the trocar and aspiration of 70 mL of pus. The parallel lines represent the walls of the catheter (arrows). (From Ulitzsch D, Nyman MKG, Carlson RA: Breast abscess in lactating women: US-guided treatment, Radiology 232:904, 2004.)

Figure 37.18 Pilonidal sinus. A, Sinuses occur in the midline, some 5 cm above the anus in the natal cleft. B, Longitudinal section showing sinuses and pits. (From Hill GJ II, editor: Outpatient Surgery, ed 3, Philadelphia, 1988, Saunders. Reproduced with permission.)
infection (e.g., tuberculosis, lymphogranuloma venereum), malignancy, or trauma.\textsuperscript{154}

The prevalence of anal abscesses in the general population is probably much higher than seen in clinical practice as the majority of patients with symptoms referable to the anorectum do not seek medical attention. It is estimated that there are approximately 100,000 cases per year in the United States.\textsuperscript{155} The mean age for presentation of anal abscess and fistula disease is 40 years (range 20 to 60 years).\textsuperscript{156-159} Adult males are twice as likely to develop an abscess and/or fistula compared with women.\textsuperscript{155,159}

Perirectal infections can range from minor irritation to fatal illness. Surprisingly, clandestine infections often occur in diabetics. Successful management depends on early recognition of the disease process and adequate surgical therapy. Small abscesses can initially be managed on an outpatient basis with simple I&D, described previously. Because of the morbidity and mortality associated with inadequate treatment of these conditions, patients with large and deep abscesses should be promptly admitted to the hospital for evaluation and treatment under general or spinal anesthesia (Fig. 37.20).

It is important to understand the anatomy of the anal canal and the rectum to appreciate the pathophysiology of these abscesses and their treatment (Fig. 37.21). The mucosa of the anal canal is loosely attached to the muscle wall. At the dentate line, where columnar epithelium gives way to squamous epithelium, there are vertical folds of tissue called the rectal columns of Morgagni that are connected at their lower ends by small semilunar folds called anal valves. Under these valves are invaginations termed anal crypts. Within these crypts are collections of ducts from the anal glands. These glands are believed to be responsible for the genesis of most, if not all perirectal abscesses. These glands often pass through the internal sphincter but do not penetrate the external sphincter.

The muscular anatomy divides the perirectal area into compartments that may house an abscess, depending on the direction of spread of the foci of the infection (Fig. 37.22).\textsuperscript{160,161} The circular fibers of the intestinal coat thicken at the rectum-anus junction to become the internal anal sphincter. The muscle fibers of the levator ani fuse with those of the outer longitudinal fibers of the intestinal coat as it passes through the pelvic floor. These conjoined fibers are connected by fibrous tissue to the external sphincter system, which consists of three circular muscle groups.

**Pathophysiology**

As described previously, the anal glands are mucous-secreting structures that terminate in the area between the internal and external sphincters. It is believed that most perirectal infections begin in the intersphincteric space secondary to blockage and subsequent infection of the anal glands. Normal host defense mechanisms are overwhelmed, and this results in invasion and overgrowth by bowel flora.\textsuperscript{162}

If the infection spreads across the external sphincter laterally, an ischiorectal abscess is formed. If the infection dissects rostrally, it may continue between the internal and external sphincters and give rise to a high intramuscular abscess. The infection may also dissect through the external sphincter over the levator ani to form a pelvirectal abscess.\textsuperscript{164}

When infection of an anal crypt extends by way of the perianal lymphatics and continues between the mucous membrane and the anal muscles, a perianal abscess forms at the anal orifice. A perianal abscess is the most common variety of perirectal abscess.
infection. A perianal abscess lies immediately beneath the skin in the perianal region at the lowermost part of the anal canal. It is separated from the ischiorectal space by a fascial septum that extends from the external sphincter and is continuous with the subcutaneous tissues of the buttocks. The infection may be small and localized or it may be very large, with a wall of necrotic tissue and a surrounding zone of cellulitis. Perianal abscesses may be associated with a fistula-in-ano. A fistula-in-ano is an inflammatory tract with an external opening in the skin of the perianal area and an internal opening in the mucosa of the anal canal. A fistula-in-ano is usually formed after partial resolution of a perianal abscess, and its presence is suggested by recurrence of these abscesses with intermittent drainage. The external opening of the fissure is usually a red, elevated piece of granulation tissue that may exhibit purulent or serosanguineous drainage on compression. In many cases the tract can be palpated as a cord. Patients with anal fistulas should be referred for definitive surgical excision.

Ischiorectal abscesses are fairly common. They are bounded superiorly by the levator ani, inferiorly by the fascia over the perianal space, medially by the anal sphincter muscles, and laterally by the obturator internus muscle. These abscesses may commonly be bilateral, and if so, the two cavities communicate by way of a deep postanal space to form a “horseshoe” abscess. Intersphincteric abscesses are less common. They are bounded by the internal and external sphincters and may extend rostrally into the rectum, thereby separating the circular and longitudinal muscle layers.

Causes of perirectal abscesses other than the cryptoglandular process have been documented but are fairly rare. It is believed that hemorrhoids, anorectal surgery, episiotomy, or local trauma may cause abscess formation by altering the local anatomy and thus destroying natural tissue barriers to infection.

Perirectal abscesses may serve as a portal of entry for organisms responsible for necrotic soft tissue infections such as Fournier’s gangrene.

**Physical and Laboratory Findings**

A perianal abscess is not generally difficult to diagnose. The severe throbbing pain in the perianal region is acute and
agrivated by sitting, coughing, sneezing, and straining. There is swelling, induration, tenderness, and a small area of cellulitis in proximity to the anus. Rectal examination of a patient with a perianal abscess reveals that most of the tenderness and induration are located below the level of the anal ring. Deeper abscesses may be difficult to localize and identify. Computed tomography is often the first imaging study, given its ready availability. The sensitivity of computed tomography for anorectal abscesses, however, is only 77%. Therefore, when clinical suspicion is high, magnetic resonance imaging should also be considered.

Patients with ischiorectal abscesses have fever, chills, and malaise, but at first there is less pain than with a perianal abscess. Initially on physical examination, one will see an asymmetry of the perianal tissue; later, erythema and induration are apparent. Digital examination reveals a large, tense, tender swelling along the anal canal that extends above the anorectal ring. If both ischiorectal spaces are involved, the findings are bilateral.

Patients with intersphincteric abscesses usually have dull, aching pain in the rectum rather than in the perianal region. No external aberrations of the perianal tissues are noted, but tenderness may be present. On digital examination one frequently palpates a soft, tender, sausage-shaped mass above the anorectal ring; if the mass has already ruptured, the patient may give a history of passage of purulent material during defecation.

Diagnosis of pelvic rectal abscesses may be very difficult. Usually, fever, chills, and malaise are present, but because the abscess is so deeply seated, few or no signs or symptoms are present in the perianal region. Rectal or vaginal examination may reveal a tender swelling that is adherent to the rectal mucosa above the anorectal ring.

Laboratory findings do not always aid in the diagnosis. Kovalcik and colleagues found that less than 50% of their patients had a white blood cell count greater than 10.0 x 10^9/L. Cultures of perirectal abscesses usually show mixed infections involving anaerobic bacteria, most commonly B. fragilis and gram-negative enteric bacilli. MRSA, though less common, should also be considered in endemic areas.

**Treatment**

Successful management of perirectal abscesses depends on adequate surgical drainage. Complications from these infections may necessitate multiple surgical procedures, prolong hospital stays, and result in sepsis and death. Bevais and associates retrospectively studied the charts of 184 patients who were treated surgically over a 10-year period. These patients were evaluated primarily to identify factors that contributed to morbidity and mortality. Initial drainage was performed under local anesthesia in 38% of the patients and under spinal or general anesthesia in 62%. The authors identified three key factors in those with excessive morbidity and mortality: (1) delay in diagnosis and treatment, (2) inadequate initial examination or treatment, and (3) associated systemic disease. They believed that the only way to effectively examine and adequately drain all but superficial well-localized perirectal abscesses was under spinal or general anesthesia. This assessment was supported by evidence of an increased incidence of recurrence, sepsis, and death in patients treated under local anesthesia.

Drainage of deep abscesses under local anesthesia generally does not allow drainage of all hidden loculations. In addition, local anesthesia is not adequate for the treatment of associated pathologic conditions.

Small, well-defined perianal abscesses without deeper perirectal involvement are the only perirectal infections that lend themselves to outpatient therapy. All other perirectal abscesses should be drained in the OR. The result of I&D is almost immediate relief of pain and rapid resolution of infection. Indications for inpatient drainage are failure to obtain adequate anesthesia, systemic toxicity, extension of the abscess beyond a localized area, or recurrence of a perianal abscess. Recurrence may be caused by the presence of a fistula-in-ano.

Drain a perianal abscess through a cruciate incision because if a simple linear incision is used, the abscess cavity has a propensity to close prematurely without adequate drainage. With either technique, make an incision over the area that is most fluctuant. If a simple linear incision is used, lightly pack the abscess cavity for at least 24 hours to ensure adequate drainage. It is extremely painful to probe a perianal abscess and to break up loculations, so liberal analgesia is advised and conscious sedation should be considered. Advise the patient to begin sitz baths at home 24 hours after surgery. Replace the packing at 48-hour intervals until the infection has cleared and granulation tissue has appeared. This usually occurs within 4 to 6 days. Cultures are generally not required in the treatment of an anal abscess as antibiotics are not generally needed. Cultures and antibiotics may be useful to distinguish between a cryptoglandular abscess (typically due to colonic flora) and an abscess of the perianal skin (typically due to staphylococcal species). Cultures may also be useful in patients who would typically be treated with antibiotics, including those with valvular heart disease, immunosuppression, extensive cellulitis, and diabetes; patients who have been on multiple courses of antibiotics; patients with pain out of proportion to findings, who may have an unusual pathogen (e.g., immunocompromised patients); and patients with leukemia or lymphoma who may have unusual or resistant bacteria.

Use of de Pezzer catheters in anorectal abscesses has been described as an alternative to traditional incision and packing. In a series of 91 patients treated in this manner, Kyle and Isbister found equivalent rates of subsequent fistula surgery, less need for general anesthesia, and a shorter postoperative hospital stay than in patients treated with traditional incision and packing. Beck and coworkers reported successful use of catheter drainage in 55 patients with an ischiorectal abscess. Because of the complexity of ischiorectal abscesses, this technique is probably best left to the surgeon who is providing ongoing care.

Perirectal abscesses are currently recognized as a fairly common cause of fever in granulocytopenic patients. These abscesses have a different bacteriologic profile: *P. aeruginosa* organisms are isolated most frequently. These patients are initially seen later because pain develops later in the course, and fever may be the first manifestation. Therefore any patients who are granulocytopenic with vague anorectal complaints, especially those with fever, should be examined carefully for perirectal abscesses. Any abscess that is found should be drained immediately under appropriate anesthesia, and extensive IV antibiotic coverage should be initiated.

**Infected Sebaceous Cyst**

A common entity that appears as a cutaneous abscess is an infected sebaceous cyst. Such infections are increasingly being caused by MRSA. Sebaceous cysts, caused by obstruction of sebaceous gland ducts, can occur anywhere on the body. The
Sebaceous Cyst Excision

This patient has an infected sebaceous cyst on the posterior of the earlobe (arrow). These lesions are typically tender, fluctuant, subcutaneous masses, often with overlying erythema.

Drain the cyst as you would a typical abscess. The thick sebaceous material must be expressed manually, because it is too thick to drain spontaneously.

Sebaceous cysts differ from simple abscesses in that they have a distinct capsule with a pearly white appearance (arrow). This capsule must be removed to prevent recurrence. This may be done initially or on a return visit once the inflammation has subsided.

Appearance of the cavity after excision of the capsule. Treat the lesion as you would any other abscess, with wound packing and timely follow-up care.

Figure 37.23 Excision of an infected sebaceous cyst. Loop drainage is not applicable to this abscess due to the need for capsule removal.

cyst becomes filled with a thick, cheesy, sebaceous material, and the contents frequently become infected. Sebaceous cysts can be quite large and may persist for many years before they become infected. When infected, they appear clinically as tender, fluctuant subcutaneous masses, often with overlying erythema.

The initial treatment of an infected sebaceous cyst is simple I&D (Fig. 37.23). The loop drainage technique may not be suited for this abscess. The thick sebaceous material must be expressed because it is too thick to drain spontaneously. An important difference exists between infected sebaceous cysts and other abscesses. A sebaceous cyst has a definite pearly white capsule that must be excised to prevent recurrence. Traditionally, in the presence of significant inflammation it is preferable to drain the infection initially and remove the shiny capsule on the first follow-up visit or at a later visit, when it can be more easily identified. Alternatively, the entire cyst can be removed during the initial incision. At the time of capsule removal, the edges are grasped with clamps or hemostats, and the entire capsule is removed by sharp dissection with a scalpel or scissors. After excision of the capsule, the area is treated in the same manner as a healing abscess cavity. Simple drainage without excision of the capsule often leads to recurrence.

Kitamura and colleagues reported a randomized study of 71 patients treated by either traditional I&D or primary resection of the cyst, followed by irrigation and wound closure. In this study, the patients treated by primary resection had faster healing, fewer days of pain, and less scarring.

Paronychia

Paronychia is an infection localized to the area around the nail root (Fig. 37.24). It is a common infection, probably caused by frequent trauma to the delicate skin around the fingernail and the cuticle. These type of infections have also been linked to “insults” induced by nail cosmetics (mechanical trauma, irritant reactions, and allergic reactions) and to a number of occupations (e.g., hair cutting and meat handling).
a minor infection begins, the nail itself may act like a foreign body. Usually, the infectious process is limited to the area above the nail base and underneath the eponychium (cuticle), but occasionally, it may spread to include tissue under the nail as well and form a subungual abscess. *Staphylococcus* is the most common cultured organism from these lesions; however, anaerobes and numerous gram-negative organisms may be isolated. Paronychia in children is often caused by anaerobes, and is believed to be the result of fingernail and nail biting. Occasionally, a group A β-hemolytic streptococcal infection will develop in a paronychia in a child with streptococcal pharyngitis who engages in thumb sucking. Paronychia involving CA-MRSA has been reported. Paronychia has also been reported in association with antiretroviral therapy for HIV infection and with use of epidermal growth factor inhibitors.

A paronychia appears as a swelling and tenderness of the soft tissue along the base or the side of a fingernail (Fig. 37.25). Pain, often around a hangnail, usually prompts a visit to the ED. The infection begins as a cellulitis and may form a frank abscess. If the nail bed is mobile, the infectious process has extended under the nail, and a more extensive drainage procedure should be performed. Although no comparative trials of surgical management versus oral antibiotics have been performed, some general guidelines for the management of paronychias have become established in clinical practice. If soft tissue swelling is present without fluctuance, treatment consists of frequent warm soaks (three times a day) and a short course of oral antibiotics. Topical antibiotics with or without topical corticosteroids have also been used. Incision will be of little value at the early cellulitic phase. Antibiotic treatment of localized infection is summarized in Table 37.1. If significant cellulitis is present, a broad-spectrum antistaphylococcal antibiotic (cephalosporin or semisynthetic penicillin) may be tried.

When a definite abscess has formed, drainage is usually quickly curative. A number of invasive operative approaches have been suggested. Actual skin incision or removal of the nail is rarely required, and neither procedure should be the initial form of treatment. One can invariably obtain adequate drainage by simply lifting the eponychial fold away from the nail matrix to allow the pus to drain. This is usually curative because a paronychia is not a cutaneous abscess per se, but rather a collection of pus in the potential space between the cuticle and the proximal end of the fingernail. Drainage may be accomplished after a digital nerve block is performed. After softening the eponychium by soaking, advance a No. 11 scalpel blade, scissors, or a 21- to 23-gauge needle parallel to the nail and under the eponychium at the site of maximal swelling (Fig. 37.26). When using the needle approach, some patients

---

**Figure 37.24** Paronychia. 1, Abscess at the side of the nail. 2, The infection has extended around the base of the nail. It has raised the eponychium but has not penetrated under the nail. 3, End stage of paronychia, with a subeponychial and subungual abscess. (From Wolcott MW, editor: Ferguson’s Surgery of the Ambulatory Patient, ed 5, Philadelphia, 1974, Lippincott. Reproduced with permission.)

**Figure 37.25** Paronychia in various degrees of severity. A, Paronychia of the index finger. In this early stage of abscess formation, the collection is limited to the lateral nail fold (arrow). B, More advanced paronychia of the great toe. The abscess has spread around the base of the nail and lifted the eponychium (arrows) but has not yet spread under the nail. There is associated cellulitis on the distal portion of the toe. C, Even more advanced paronychia, with subungual extension (black arrow) and cellulitis extending up the finger and onto the dorsum of the hand (white arrow). This patient required removal of the nail and intravenous antibiotics.
may tolerate the procedure without the need for digital blockade. Pus escapes rapidly, with immediate relief of pain. A tourniquet placed at the base of the finger may limit bleeding and aid the clinician in determining the exact extent of the infection during the drainage procedure.

If more than a tiny pocket of pus is present, fan the knife tip or needle or spread the scissors under the eponychium while keeping the instrument parallel to the plane of the fingernail (see Fig. 37.26, step 5 and 6). When a large amount of pus is drained, slip a small piece of packing gauze under the eponychium for 24 hours to provide continual drainage. Given the prevalence of MRSA, cultures are generally indicated. Antibiotics are frequently prescribed, although they are not essential if drainage is complete or the surrounding area of cellulitis is minimal. An alternative to systemic antibiotics is to keep the operative site bathed in antibiotic ointment. After the anesthesia has worn off, start the patient on frequent soaks in warm tap water at home. In most cases the patient may easily remove the packing. At 24 to 36 hours, soak the finger in warm water and pull the gauze out; a repeated visit to a clinician is not required if healing is progressing, but follow-up examinations should always be encouraged. Once the packing is removed, cover the area with a dry, absorbent dressing. Apply an antibiotic ointment periodically to the site for several days.

An alternative technique, called the Swiss roll technique, was described by Pabari and colleagues for severe, acute paronychia with a run-around or contiguous infection of both nail folds. With a No. 15 scalpel blade pointing away from the nail bed, make an incision on both sides of the nail fold. The nail fold, after thorough irrigation with saline, is then rolled proximally over nonadherent dressing, like a Swiss roll, and sutured to the skin with a nonabsorbable dressing. Apply a finger dressing. Remove the sutures in 48 hours and unfold the nail fold back to its original position to allow it to heal by secondary intention. The authors argue that this technique spares the nail plate and allows rapid healing, although no controlled studies have compared this technique with simple incision followed by packing gauze described previously.

If the infection has produced purulence beneath the nail (subungual abscess), remove a portion of the nail or trephinate the nail to ensure complete drainage. As an alternative to nail removal, place a hole in the proximal part of the nail with a hot paper clip or a portable electrocautery device. Make a large opening or multiple holes to ensure continued drainage.

The proximal portion of the nail is involved in most cases. After a digital block has been performed under sterile conditions, proximal subungual collections can be managed by bluntly elevating the eponychium to expose the proximal edge of the nail. Elevate the proximal third of the nail from the nail bed and resect it with scissors. Leave the distal two thirds of the nail in place to act as a physiologic dressing and to decrease postoperative pain (Fig. 37.27). If purulence is found below the lateral edge of the nail, gently elevate the affected part and excise it longitudinally.

Exercise care during this procedure to avoid damage to the nail matrix. Place a wick of gauze beneath the eponychium for 48 hours to ensure continued drainage.

Most paronychias resolve in a few days, and one to two postoperative visits should be scheduled to evaluate healing and reinforce home care. For compliant patients with a small paronychia, home care alone may suffice after the initial drainage. Clinical infection lasting longer than a few weeks should prompt evaluation for osteomyelitis of the distal phalanx, a known but rare complication of even a properly drained paronychia.

Patients occasionally come to the ED complaining of a chronic, indolent paronychial infection. These seldom respond to ED intervention. Frank purulence is rarely present, and

<table>
<thead>
<tr>
<th>CONDITION</th>
<th>MOST COMMON OFFENDING ORGANISMS</th>
<th>RECOMMENDED ANTIMICROBIAL AGENTS</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Paronychia</td>
<td>Usually <em>Staphylococcus aureus</em> or streptococci; <em>Pseudomonas</em>, gram-negative bacilli, and anaerobes may be present, especially in patients with exposure to oral flora</td>
<td>First-generation cephalosporin or antistaphylococcal penicillin; if anaerobes or <em>Escherichia coli</em> are suspected, oral clindamycin (<em>Cleocin</em>) or a β-lactamase inhibitor such as amoxicillin-clavulanate potassium (<em>Augmentin</em>)</td>
<td>I&amp;D should be performed if infection is well established If infection is chronic, suspect <em>Candida albicans</em> Early infections without cellulitis may respond to conservative therapy</td>
</tr>
<tr>
<td>Felon</td>
<td><em>S. aureus</em>, streptococci</td>
<td>First-generation cephalosporin or antistaphylococcal penicillin</td>
<td>I&amp;D should be performed if infection is well established Oral antibiotic therapy is usually adequate</td>
</tr>
<tr>
<td>Herpetic whitlow</td>
<td>Herpes simplex virus</td>
<td>Supportive therapy</td>
<td>Antivirals may be prescribed if infection has been present for &lt;48 hr For recurrent herpetic whitlow, suppressive therapy with an antiviral agent may be helpful Consider antibiotics if secondarily infected I&amp;D is contraindicated</td>
</tr>
</tbody>
</table>

*I&D, Infection and drainage.* 
Paronychia Drainage

In selected patients the procedure can be completed without anesthesia. After softening the eponychium by soaking, advance a No. 11 scalpel blade under the eponychium parallel to the nail. Pus escapes rapidly, with immediate relief of pain. Antibiotics are not required for minor cases. Instruct the patient to frequently soak the digit in warm water at home.

Many patients, especially those with a large paronychia, will require a digital block before the procedure. (See Chapter 31 for details on digital blocks.) Lift the eponychial fold away from the nail. Here, the blunt side of a No. 15 blade is being used. No sharp incision is required; simply lifting the eponychium away from the nail is sufficient.

Pus will readily escape as the fold is lifted. Keep the scalpel parallel to the nail and continue to lift the eponychium to release all the pus. Place a small piece of packing gauze under the eponychium for 24 hours, encourage warm soaks, and prescribe antibiotics if cellulitis is present.

Figure 37.26 Paronychia drainage.
Incision and Drainage

Incision and Drainage typically involve the digits but can also involve other areas of the hand. The lesions are usually quite painful but are self-limited and resolve in 2 to 3 weeks. Unlike bacterial infections the pulp space is soft and not tense. The treatment of herpetic whitlow is conservative. Drainage of herpetic whitlow is contraindicated as it may induce a secondary bacterial infection and delay healing. Treatment is symptomatic and consists of splinting, elevation, and analgesia as needed. Oral antiviral agents effective against herpes infections (acyclovir, famciclovir, or valacyclovir) can shorten the course of the disease if given early. Infection recurs in 30% to 50% of cases, but the initial infection is usually the most severe. Acyclovir, 200 mg taken orally three to four times a day, can decrease recurrence rates.

Consideration must be given to preventing spread of the infection. An occlusive dressing decreases the chance of viral transmission, but health care providers with herpetic whitlow should limit and perhaps even refrain from patient contact, especially until all lesions have crusted over and viral shedding has stopped. Compliance with universal precautions will decrease the likelihood of patient-to-provider transmission.

Felon

A felon is an infection of the pulp of the distal end of the finger (Fig. 37.29). The usual cause is trauma with secondary invasion by bacteria. A felon may develop in the presence of a foreign body, such as a thorn or a splinter, but often a precipitating trauma cannot be identified. Finger-stick felons are common in diabetic patients. An important anatomic characteristic of this area is that many fibrous septa extend from the volar skin of the fat pad to the periosteum of the phalanx; these septa subdivide and compartmentalize the pulp area. When an infection occurs in the pulp, these structures make it a closed-space infection. The septa limit swelling, delay pointing of the abscess, and inhibit drainage after incomplete surgical decompression. Pressure may increase in the closed space and initiate an ischemic process that compounds conservative treatments are often unsatisfactory. Many causes of this frustrating condition have been described, including fungal, bacterial, viral, and psoriatic conditions. Screen patients for malignancy if they have chronic paronychia that is unresponsive to therapy. Meticulous hand care, oral and topical antimicrobial medications, topical corticosteroids, and, occasionally, aggressive surgical intervention have been suggested. Refer these patients to a dermatologist or hand surgeon because of the prolonged treatment required.

Herpetic Whitlow

Herpetic whitlow is an extremely contagious infection of the distal phalanx caused by herpes simplex virus (type 1 or 2) (Fig. 37.28). Herpes infection of the hand predominantly presents in a bimodal age distribution: children younger than 10 years of age and young adults between 20 and 30 years of age. In children, herpetic whitlow tends to be associated with gingivostomatitis caused by herpes simplex 1, whereas adults most commonly harbor herpes simplex 2. Inoculation occurs through a discontinuity in the skin. Health care providers exposed to oral secretions (e.g., dental hygienists and respiratory therapists) and patients with other herpes infections are most commonly infected. After a 2-day to 2-week incubation period, an infected individual may experience prodromal signs and symptoms such as fever, malaise, lymphadenitis, and axillary lymphadenopathy. In the affected finger the infection is characterized by tenderness, followed by throbbing or burning pain (out of proportion to the findings on physical examination), edema, and erythema. Vesicles containing clear, bloody, or cloudy fluid form and mark the most infectious stage of the process. Vesicles may coalesce to form a bulla. Viral vesicles typically involve the digits but can also involve other areas of the hand. The lesions are usually quite painful but are self-limited and resolve in 2 to 3 weeks. Unlike bacterial infections the pulp space is soft and not tense. The treatment of herpetic whitlow is conservative. Drainage of herpetic whitlow is contraindicated as it may induce a secondary bacterial infection and delay healing. Treatment is symptomatic and consists of splinting, elevation, and analgesia as needed. Oral antiviral agents effective against herpes infections (acyclovir, famciclovir, or valacyclovir) can shorten the course of the disease if given early. Infection recurs in 30% to 50% of cases, but the initial infection is usually the most severe. Acyclovir, 200 mg taken orally three to four times a day, can decrease recurrence rates.

Consideration must be given to preventing spread of the infection. An occlusive dressing decreases the chance of viral transmission, but health care providers with herpetic whitlow should limit and perhaps even refrain from patient contact, especially until all lesions have crusted over and viral shedding has stopped. Compliance with universal precautions will decrease the likelihood of patient-to-provider transmission.

Felon

A felon is an infection of the pulp of the distal end of the finger (Fig. 37.29). The usual cause is trauma with secondary invasion by bacteria. A felon may develop in the presence of a foreign body, such as a thorn or a splinter, but often a precipitating trauma cannot be identified. Finger-stick felons are common in diabetic patients. An important anatomic characteristic of this area is that many fibrous septa extend from the volar skin of the fat pad to the periosteum of the phalanx; these septa subdivide and compartmentalize the pulp area. When an infection occurs in the pulp, these structures make it a closed-space infection. The septa limit swelling, delay pointing of the abscess, and inhibit drainage after incomplete surgical decompression. Pressure may increase in the closed space and initiate an ischemic process that compounds
the infection. The infection can progress to osteomyelitis of the distal phalanx, septic arthritis, and flexor tenosynovitis. Although the septa may facilitate an infection in the pulp, they also provide a barrier that protects the joint space and tendon sheath by limiting proximal spread of the infection.

In most cases the offending organism is *S. aureus*, but mixed infections and gram-negative infection may also occur. A study looking at the bacteriology of hand infections over an 11-year period found that of 159 hand infections, 48 (30%) harbored CA-MRSA, the incidence of CA-MRSA increased by 41% per year, and IV drug use and felon infections in particular were risk factors for CA-MRSA. These changing patterns, along with the often protracted course of felons and concern for osteomyelitis, make this one of the soft tissue infections in which culture may be helpful.

A patient in whom a felon is developing will describe a gradual onset of pain and tenderness of the fingertip. In a few days, the pain becomes constant and throbbing and gradually becomes severe. In the initial stages, physical examination may be quite unimpressive because the fibrous septa limit swelling in the closed pulp space. As the infection progresses, swelling and redness become obvious. Occasionally, one may elicit point tenderness, but frequently, the entire pulp space is extremely tender. The patient characteristically arrives with the hand elevated over the head because the pain is so intense in the dependent position. Cessation of pain indicates necrosis and nerve degeneration.

During the early stages of cellulitis, a felon may be controlled by treatment consisting of elevation, oral antibiotics (see Box 37.2), and warm water or saline soaks. For more developed felons, proper treatment consists of early and complete I&D. Antibiotics alone are not curative once suppuration has occurred. Delaying surgery may result in permanent disability and deformity.

A minor felon can usually be drained on an outpatient basis. The distal phalanx was almost completely resorbed because of pressure, inflammation, and chronic osteomyelitis. This infection is extensive and warrants consultation with a hand specialist.

Figure 37.29 A well-developed felon. In this advanced case the patient had little pain at initial evaluation. The distal phalanx was almost completely resorbed because of pressure, inflammation, and chronic osteomyelitis. This infection is extensive and warrants consultation with a hand specialist.
Felon Drainage

The preferred initial incision for draining a felon is made directly into the area of most fluctuance (1). More aggressive incisions should be reserved for complicated cases because they have greater morbidity and require more complicated wound care. The unilateral longitudinal approach is a good first choice. Some prefer a similarly located through-and-through incision (see below).

A fat pad incision is generally avoided but can be acceptable for localized infections. They may be associated with a painful scar in an area that is often traumatized. The transverse fat pad incision should avoid the digital nerves (2), and the longitudinal fat pad incision should avoid the flexor tendon (3).

The infection has spread volarly and progressed to a felon. Drainage of both areas is required. The eponychium is lifted, which resulted in immediate drainage of large quantities of pus.

A hemostat is placed under the eponychium to facilitate complete evacuation of the pus.

To drain the felon, a through-and-through incision is made on the volar side. A hemostat is used to break up loculations in the fat pad and then to grab gauze for a pull-through pack.

The gauze pack is pulled through the incision and left in place for 24 to 48 hours to ensure continued drainage.

Figure 37.30 Felon drainage. For a felon the authors prefer the through-and-through drainage procedure shown in B. Antibiotics should be used postoperatively. Culture can aid in the selection of long-term antibiotics, but most initially cover for community-acquired methicillin-resistant Staphylococcus aureus.
examined this possible association, with fracture subsets totaling 26, no infections occurred. These studies suggest that the presence of an underlying fracture does not contraindicate nail trephination for fear of causing infection. There are no data supporting the routine use of prophylactic antibiotics following trephination if an underlying fracture is present.

Subungual hematomas should be trephinated if they are acute (less than 24 to 48 hours old), are not spontaneously draining, are associated with intact nail folds, or are painful. After 48 hours, most subungual hematomas have clotted and trephination is typically not effective.

Nail removal is unnecessary, even if there is an underlying distal phalanx fracture, as long as the nail margins and nail are intact. If the nail is split, avulsed or disrupted, or a nail bed laceration extends to the skin, the nail should be removed and the underlying nail bed repaired (see further discussion in Chapter 35). Simple trephination of an uncomplicated subungual hematoma, even if it involves the entire subungual area, results in a good outcome functionally and cosmetically. Prophylactic antibiotics are not necessary for patients with uncomplicated subungual hematomas that have been trephinated. An underlying tuft fracture does not mandate antibiotic prophylaxis. For more serious injuries, a slight risk for infection exists, but in most cases rigorous wound cleaning and careful soft tissue repair constitute adequate treatment.

Methods of Trephination

All methods of trephination require aseptic technique. Clean the nail thoroughly and place the digit in a sterile field. Follow universal precautions because the fluid under the nail is under pressure and can spurt. Ensure that adequate analgesia has been achieved. Anesthesia is not always required, but a digital block can be used to calm an anxious patient and is commonly performed.

An 18- or 21-gauge needle can be rotated between the thumb and the index finger while gentle pressure is applied so that the sharp end of the needle corkscrews through the nail. When using a needle, the “give” is harder to feel, so...
the clinician should proceed slowly until blood is drawn. Because
the hole made with a needle is small, a second hole may be
required.

An alternative needle-based approach is being used by a
group of Turkish dermatologists. Kaya and associates\textsuperscript{208} use
extra fine, 29-gauge insulin syringes for evacuation of hema-
tomas. After the nail is trimmed, the needle is inserted parallel
to the nail plate and advanced to the distal edge of the
hematoma. This technique is especially effective for small
hematomas of the second, third, and fourth toenails, which
are hard to trephine.

A No. 11 scalpel blade can be used to score and then cut
through the affected nail. This approach is painful for the
patient because the hole is usually larger than required.

Hot cautery is the most common form of trephination and
is usually done with a paper clip which has been straightened
and heated in a flame (Figs. 37.32 and 37.33). Place the hot
end on the nail above the center of the hematoma and apply
gentle pressure until the nail is breached and the hematoma
expressed. A give is felt as the instrument passes through the
nail. Stop the pressure at this point to avoid damage to the
nail bed. Blood exits rapidly, and the blackened nail regains
its normal color (see Fig. 37.33, step 3). The blood usually
remains fluid for 24 to 36 hours and can easily be expressed
with slight pressure. Multiple holes may be needed for continued
drainage. Do not use hot cautery trephination on artificial
nails because they are flammable.\textsuperscript{202} Hot cautery with a paper
clip is easy to perform, but hematomas treated in this way
tend to reform. It has also fallen out of favor secondary to the
need of using an open flame in front of the patient. In addition,
most paper clips are made of aluminum, which is difficult to
sufficiently heat to penetrate the nail. A portable hot-wire
electrocautery unit can be used, but it is difficult to obtain
an adequate drainage hole without adapting the instrument
(see Fig. 37.33). It can be modified to burn a larger hole by
“fattening” the end of the wire loop and rotating the device
slowly as the nail is penetrated or by removing a small rectangle
of nail. In addition to being convenient, the cautery device is
desirable because the wire stays hotter longer, thereby enhancing
nail penetration.

The PathFormer (Path Scientific, Carlisle, MA) is a relatively
new device that allows controlled nail trephination (“mesoscis-
sion”) and therefore minimizes patient discomfort during the
procedure.\textsuperscript{207} It uses electrical resistance in the nail bed as
feedback to stop and retract the drill when it penetrates the
nail plate. The nail bed, with its blood supply and nerve endings,
is not disturbed.

Figure 37.32 A straightened paper clip, held with a hemostat and
heated with a flame, can be used for hot cautery trephination.
laceration. A common mechanism is slamming a finger in a car door, which causes sudden flexion of the distal phalanx in conjunction with a crush injury of the nail bed. The nail itself is usually stable, so generally no repair of the nail bed appears to be required. However, if the subungual hematoma extends past the confines of the nail bed, such as in the paronychial space (under the skin of the cuticle), there must be a communication between the nail bed and this space. This produces a paronychial hematoma, with blood occupying the space where pus would be located in an infectious paronychia (Fig. 37.34).

When this condition is present, the avulsed proximal portion of the fingernail overlies the nail fold of the cuticle but is appreciated only after the overlying skin is opened. Open reduction (replacement) of the nail must be performed. The replaced nail often grows normally, but a lost or deformed nail is possible. Repair of the nail bed laceration is optional at this juncture but may not be required if the nail is stable. Once the injury is anatomically aligned, splinting and soaking

**Outcome**

It is difficult to predict the fate of the fingernail after drainage of a subungual hematoma. Obviously, there must be a nail bed laceration if bleeding occurs, but with a stable nail bed, repair is unnecessary. Even if a small tuft fracture is present, most do well with simple drainage. Some patients will lose the nail, but if the nail root or nail bed is not significantly disrupted and the nail remains implanted, a normal-appearing nail is the usual final result.

After a hematoma has been drained, the nail should be cleaned thoroughly and a dry dressing applied. Patients should be advised to keep the digit dry for 2 days. Any fracture should be splinted and given appropriate follow-up at a surgical clinic.

**Conditions With a Similar Appearance**

One condition that may be mistaken for a simple subungual hematoma is closed avulsion of the base of a fingernail occurring in conjunction with a subungual hematoma from a nail bed laceration. A common mechanism is slamming a finger in a car door, which causes sudden flexion of the distal phalanx in conjunction with a crush injury of the nail bed. The nail itself is usually stable, so generally no repair of the nail bed appears to be required. However, if the subungual hematoma extends past the confines of the nail bed, such as in the paronychial space (under the skin of the cuticle), there must be a communication between the nail bed and this space. This produces a paronychial hematoma, with blood occupying the space where pus would be located in an infectious paronychia (Fig. 37.34). When this condition is present, the avulsed proximal portion of the fingernail overlies the nail fold of the cuticle but is appreciated only after the overlying skin is opened. Open reduction (replacement) of the nail must be performed. The replaced nail often grows normally, but a lost or deformed nail is possible. Repair of the nail bed laceration is optional at this juncture but may not be required if the nail is stable. Once the injury is anatomically aligned, splinting and soaking

---

**Figure 37.34** Complicated subungual hematoma. **A,** This patient slammed his finger in a car door and sustained acute flexion of the distal phalanx and a crush injury. It appears to be a simple subungual hematoma, but note the blood in the paronychial space (arrow). There is communication between the nail bed laceration and the eponychium. **B,** All the blood did not drain when the nail was trephinated. **C,** Blood accumulated in this area because the base of the fingernail has been avulsed from its origin and now lies between the eponychium (cuticle) and skin. This is appreciated when the skin is débrided. Note the white avulsed base of the fingernail just under the skin. The closed nature of the injury causes the confusion. **D,** With the nail removed, the nail bed laceration (arrow) can be seen and repaired. **E,** The old trephined nail; **F,** this can now be replaced and sutured in its original position to keep the eponychial space open, or the space can be packed with gauze for a few weeks to discourage scar formation and subsequent nail deformity. Note the drainage hole in the original nail.
follow as for a simple subungual hematoma. These injuries rarely become infected, and there is no evidence that the prophylactic use of antibiotics is necessary.

Subungual hematomas can also be a manifestation of diseases such as Kaposi's sarcoma and melanoma. These conditions should be considered when no trauma has occurred and the findings on physical examination are not consistent with a simple subungual hematoma.

ACKNOWLEDGMENTS

The authors acknowledge the significant contributions of Liam Holtzman, Eveline Hitti, and Jeffrey Harrow to this chapter in previous editions.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


151. Deleted in review.


CHAPTER 38

Burn Care Procedures

Anthony S. Mazzeo

Two million people suffer a burn-related injury every year in the United States. The American Burn Association (ABA) estimates that almost 500,000 of these patients received medical evaluation and treatment in 2014 and approximately 40,000 required hospitalization.1 According to the 2014 National Burn Repository,2 patients who suffer burn injuries are predominately male (69%), and their mean age is 32 years old. Children younger than 5 years account for 19% of burns, and patients older than 60 years account for an additional 13%. Seventy-four percent of all burns involve less than 10% of the total body surface area (TBSA). Nearly 80% of all burns are caused by flame or fire or by scalds, with scald injury occurring most in children younger than 5 years.

Overall, the incidence of burn injury has declined in recent decades and advances in medical care have improved the morbidity and mortality of burn patients.2,3 Enhancements in resuscitation, surgical and anesthetic techniques, intensive care, infection control, nutrition, and metabolic support have all contributed to dramatic improvements in the survival, preservation of body function, physical appearance, and emotional outcomes of patients with this injury. The initial care provided to burn patients by emergency medical providers can improve outcomes by preventing the conversion of superficial burns to deep burns requiring surgery and by improving the long-term functional and cosmetic outcomes of the affected tissues.

The classification of burns is based on three criteria: depth of skin injury, percentage of TBSA involved, and source of injury (thermal, chemical, electrical, or radiation). The seriousness of a burn injury is determined by the characteristics and temperature of the burning agent, the duration of exposure, the location of the injury, the presence of associated injuries, and the age and general health of the victim (Table 38.1).

The ABA defines minor burns as uncomplicated partial-thickness burns involving less than 5% TBSA in children (<10 years old) or the elderly (>50 years old), less than 10% TBSA in adults, or full-thickness burns less than 2% TBSA.4 Moderate or major burns include injuries that involve a greater TBSA, as well as burns in areas of specialized function, such as the face, hands, feet, and perineum. More serious burns also include those caused by a high-voltage electrical injury or those with associated inhalation injuries or other major trauma.

Throughout the course of history, clinicians have experimented with burn therapies to relieve pain and promote healing. Many treatment regimens and home remedies have been successful, largely because minor burns generally do well with a modicum of intervention and commonsense wound care. Although little has changed in the care of minor ambulatory burns over the past 3 decades, treatment of major burns has significantly improved, including the development of sophisticated burn centers, increased knowledge of burn wound physiology, and prevention of infection.

WOUND EVALUATION

Emergency clinicians should be aware that the depth of a burn wound cannot always be determined accurately on clinical grounds alone at initial evaluation and that burn injury is a dynamic process that may change over time, particularly during the 24 to 48 hours after the burning process has been arrested. It is common, for example, for a seemingly minor or superficial burn to appear deeper on the second or third return visit (Fig. 38.1). This phenomenon is not a continuation of the burning process that can be altered by clinician intervention but is considered to be a pathophysiologic event related to tissue edema, dermal ischemia, or desiccation.5

Estimating Burn Depth

The depth of a burn has historically been classified by degree.7 First degree involves the epidermis only, second-degree (or partial-thickness) burns extend into the dermis, and third-degree (or full-thickness) burns destroy the entire skin. An additional fourth degree is sometimes used to describe injuries to the underlying muscle, tendon, or bone (Fig. 38.2).

First-degree burns involve the epidermis only (Fig. 38.3A). The skin is reddened but is intact and not blistered. This injury ranges from mildly irritating or even pruritic to exquisitely painful. Minor edema may be noted. Causes include ultraviolet light (as in sunburn) and brief thermal “flash” burns. First-degree burns may blister within 24 to 36 hours, and the patient should be warned about this possibility. Frequently, the epidermis may flake or peel within 5 to 10 days. Healing occurs spontaneously, usually without scarring.

Second-degree burns involve the epidermis and extend into the dermis to include the sweat glands and hair follicles. Superficial second-degree burns involve only the papillary dermis (see Fig. 38.3B). These burns are pink, moist, and extremely painful. Blisters are common and the skin may slough. The burn blanches with pressure, and mild to moderate edema is common. Hair follicles often remain intact. Superficial second-degree burns are the most common burns seen in the emergency department (ED). The usual causes are scalds, contact with hot objects, or exposure to chemicals. Barring infection or repeated trauma, these burns heal spontaneously and without scarring in approximately 2 weeks. These areas may be sensitive to sunburn, windburn, and skin irritation for months after the original injury has healed.

Deep second-degree burns involve the reticular dermis and appear mottled white or pink (see Fig. 38.3C). There is obvious edema and sloughing of the skin, and any blisters are usually ruptured. Blanching is absent. These burns are not generally painful initially and may have decreased sensation, but pressure may be perceived. Within a few days, however, these burns can become exquisitely painful. This type of burn may be converted to a full-thickness injury by further trauma or infection.

Third-degree burns result from complete loss of the dermis and may extend into subcutaneous (SQ) tissue (see Fig. 38.3D). These burns usually appear dry, pearly white, or charred. They are initially painless, with a leathery texture. Circumferential third-degree burns on an extremity or the torso cause a loss of elasticity that may impair the circulation or ventilation and necessitate an escharotomy.

Fourth-degree burns extend deeply into SQ tissue, muscle, fascia, or bone (see Fig. 38.3E). These burns are
**TABLE 38.1 Characteristics of Burns, by Depth**

<table>
<thead>
<tr>
<th>CLASSIFICATION OF BURN</th>
<th>ETIOLOGY</th>
<th>APPEARANCE</th>
<th>SENSATION</th>
<th>TIME TO COMPLETE HEALING</th>
<th>SCARRING</th>
</tr>
</thead>
<tbody>
<tr>
<td>First Degree</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Superficial epidermal layers</td>
<td>Sunburn, other UV exposure</td>
<td>Dry, red Blanches with pressure</td>
<td>Present May be quite painful</td>
<td>3–7 days</td>
<td>No</td>
</tr>
<tr>
<td>Second Degree</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Varying depth, blisters, or bullae formation</td>
<td>Dermal appendages spared (e.g., sweat glands, hair follicles)</td>
<td>Includes entire epidermis and some portion of the dermis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Supersicial partial thickness</td>
<td>Water scald Longer flash burn</td>
<td>Blisters, peeling skin Blanches with pressure Skin red and moist under blisters</td>
<td>Painful Exposure to air and temperature painful</td>
<td>7–21 days</td>
<td>Unusual if no infection and proper follow-up Pigment change may be seen Burned area may be sensitive to frostbite, windburn, and sunburn for many months Itching may be problematic for weeks after healing</td>
</tr>
<tr>
<td>Deep partial thickness</td>
<td>Flame Water immersion Oil, grease, hot foods (e.g., soup)</td>
<td>Variable color Wet or waxy dry, does not blanch Blisters easily removed, skin peeling off</td>
<td>Pressure only</td>
<td>&gt;21 days</td>
<td>Severe; risk for contracture</td>
</tr>
<tr>
<td>Third Degree</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Loss of all skin elements, thrombosis and coagulation of vessels</td>
<td>Flame, steam, oil grease Immersion, scald Caustic chemical, high voltage</td>
<td>Leathery appearance, white or charred, dry, inelastic; blanching with pressure May be present under blisters</td>
<td>Deep pressure only</td>
<td>Never heals Requires grafting</td>
<td>Very severe, high risk for contracture</td>
</tr>
</tbody>
</table>

*U*V, Ultraviolet.


---

**Figure 38.1** It may be difficult to accurately assess the depth or severity of a burn on the first visit. **A,** This full-thickness burn will not heal without a skin graft. **B,** This blistered hot water burn is probably second degree, but full-thickness burns can develop under blisters. **C,** At 2 weeks, a second-degree burn and a small area of third-degree burn (arrows).
Figure 38.2 Depths of a burn. First-degree burns are confined to the epidermis. Second-degree burns extend into the dermis (dermal burns). Third-degree burns are full thickness through the epidermis and dermis. Fourth-degree burns involve injury to underlying tissue structures such as muscle, tendons, and bone. (From Townsend CM, Beauchamp RD, Evers BM, et al., editors: Sabiston textbook of surgery, ed 19, St. Louis, 2012, Saunders.)

Figure 38.3 Depth of thermal injury. A, Patient with sunburn on the lower extremity (a superficial or first-degree burn with associated blisters on the anterior tibial surface). B, Partial-thickness injury of the hand (superficial second-degree burn). C, Partial-thickness injury extending beyond the subcutaneous layers (deep second-degree burn). D, Full-thickness (third-degree) burn. E, Full-thickness injury with extensive tissue loss (fourth-degree burn). (From Davis PJ, Cladis FP, Motoyama EK, editors: Smith’s anesthesia for infants and children, ed 8, St. Louis, 2011, Mosby.)
Calculating burn size is necessary to determine treatment plan, fluid requirements, and aids in prognosis. It is important for the emergency provider to accurately estimate the burn size. Note that first-degree burns do not count in the calculation of total burn surface area when utilizing fluid resuscitation formulas. Overestimating the burn size is a common error, especially in children. Several formulas are available to estimate TBSA in burn patients. In 1944, Lund and Browder published the now famous Lund-Browder chart (Fig. 38.4). In their landmark paper, they used direct measurements and body surface area formulas to produce a chart that clinicians can use to estimate %TBSA. Burn centers typically utilize the Lund-Browder chart for estimating burn size. The initial Lund-Browder chart was developed from human anatomic studies derived from 11 adults (3 women and 8 men) and produced a unisex chart. A 2004 study involving 60 volunteers determined that the Lund-Browder chart significantly underestimates the size of chest burns in large-breasted women. The investigators developed a table that incorporates a correction using brassiere cup size.

The simplest method for estimating TBSA in adults is the “rule of nines.” This formula was developed in the late 1940s by Pulaski and Tennon, who observed that the percentage of each body segment was approximately a multiple of nine (Fig. 38.5). Similar formulas for children adjust estimates for their disproportionately large head surface area. However, in a study of obese patients it was determined that this formula underestimates %TBSA of the legs and torso and overestimates %TBSA of the arms and head. The authors suggested replacing the “rule of nines” with a “rule of fives” for obese patients heavier than 80 kg.

The size of a burn can also be estimated by using the patient’s hand as representing approximately 1% TBSA. With this method, the hand is a rectangle. However, two studies using planimetry have determined that the hand actually represents from 0.5% to 0.78% of a patient’s TBSA.

Regardless of the method used, it is important for the emergency provider to take time to quantify the %TBSA as accurately as possible. It should be reemphasized that the emergency provider to accurately estimate the burn size. Note that first-degree burns do not count in the calculation of total burn surface area when utilizing fluid resuscitation formulas. Overestimating the burn size is a common error, especially in children. Several formulas are available to estimate TBSA in burn patients. In 1944, Lund and Browder published the now famous Lund-Browder chart (Fig. 38.4). In their landmark paper, they used direct measurements and body surface area formulas to produce a chart that clinicians can use to estimate %TBSA. Burn centers typically utilize the Lund-Browder chart for estimating burn size. The initial Lund-Browder chart was developed from human anatomic studies derived from 11 adults (3 women and 8 men) and produced a unisex chart. A 2004 study involving 60 volunteers determined that the Lund-Browder chart significantly underestimates the size of chest burns in large-breasted women. The investigators developed a table that incorporates a correction using brassiere cup size.

The simplest method for estimating TBSA in adults is the “rule of nines.” This formula was developed in the late 1940s by Pulaski and Tennon, who observed that the percentage of each body segment was approximately a multiple of nine (Fig. 38.5). Similar formulas for children adjust estimates for their disproportionately large head surface area. However, in a study of obese patients it was determined that this formula underestimates %TBSA of the legs and torso and overestimates %TBSA of the arms and head. The authors suggested replacing the “rule of nines” with a “rule of fives” for obese patients heavier than 80 kg.

The size of a burn can also be estimated by using the patient’s hand as representing approximately 1% TBSA. With this method, the hand is a rectangle. However, two studies using planimetry have determined that the hand actually represents from 0.5% to 0.78% of a patient’s TBSA.

Regardless of the method used, it is important for the emergency provider to take time to quantify the %TBSA as accurately as possible. It should be reemphasized that

**Estimating Burn Size**

Calculating burn size is necessary to determine treatment plan, fluid requirements, and aids in prognosis. It is important for

**Table 38.2 American Burn Association’s Grading System for Burn Severity and Disposition of Patients**

<table>
<thead>
<tr>
<th>Type of Burn</th>
<th>Minor</th>
<th>Moderate</th>
<th>Major</th>
</tr>
</thead>
<tbody>
<tr>
<td>Criteria</td>
<td>&lt;10% TBSA burn in adult</td>
<td>10%–20% TBSA burn in adult</td>
<td>&gt;20% TBSA burn in adult</td>
</tr>
<tr>
<td></td>
<td>&lt;5% TBSA burn in young or old</td>
<td>5%–10% TBSA burn in young or old</td>
<td>&gt;10% TBSA burn in young or old</td>
</tr>
<tr>
<td></td>
<td>&lt;2% full-thickness burn</td>
<td>2%–5% full-thickness burn</td>
<td>&gt;5% full-thickness burn</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High-voltage injury</td>
<td>High-voltage burn</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Suspected inhalation injury</td>
<td>Known inhalation injury</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Circumferential burn</td>
<td>Any significant burn involving the</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Concomitant medical problem</td>
<td>face, eyes, ears, hands, feet,</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(e.g., diabetes, sickle cell disease)</td>
<td>genitalia, or joints</td>
</tr>
<tr>
<td>Disposition</td>
<td>Outpatient management</td>
<td>Hospital admission vs. higher-level, structured outpatient care</td>
<td>Consider referral to a burn center</td>
</tr>
</tbody>
</table>

*Criteria: Burn, partial-thickness or full-thickness burn, unless specified; young, patient younger than 10 years; adult, patient 10 to 50 years of age; old, patient older than 50 years.*

fluid recommendations based on %TBSA DO NOT include first-degree burns in the %TBSA, rather they are based on %TBSA of second- and third- (and fourth-) degree burns. Including first-degree burns in the calculation of %TBSA results in an overestimate of the total burned areas, and is a common error.

**HISTOPATHOLOGY OF BURNS**

One thermal wound theory describes three zones of injury in burns (Fig. 38.6):

1. Zone of coagulation: dead, avascular tissue that must be débrided.
2. Zone of stasis: injured tissue in which blood flow is impaired. Desiccation, infection, or mechanical trauma may lead to cell death.
3. Zone of hyperemia: minimally injured, inflamed tissue that forms the border of the wound. The hyperemia usually resolves within 7 to 10 days but may be mistaken for cellulitis. Histologically, full-thickness burns are characterized by confluent vascular thrombosis involving arterioles, venules, and capillaries. Edema secondary to loss of microvascular integrity results not only from the effects of direct thermal injury but also from the release of vasoactive mediators. The increase in vascular permeability is linked to activation of complement and release of histamine. Histamine increases catalytic activity...
of the enzyme xanthine oxidase, with resultant production of hydrogen peroxide and hydroxyl radicals. These by-products increase the damage to dermal vascular endothelial cells and result in progressive vascular permeability.\(^6\)

The cellular debris and denatured proteins of the eschar provide a milieu that supports the proliferation of microorganisms. The devitalized tissue (eschar) sloughs spontaneously, usually as a result of the proteolytic effect of bacterial enzymes. The greater the degree of wound bacteriostasis, the greater the delay in sloughing.

Partial-thickness burns result in incomplete vascular thrombosis, usually limited to the upper dermis. The dermal circulation is restored gradually, generally over a period of several days, thus resulting in a significant interval of relative ischemia. The eschar in deep partial-thickness burns is thinner than in a full-thickness burn and sloughs as a result of reepithelialization rather than bacterial proteolysis.

**Figure 38.5** The “rule of nines.” The rule of nines is a rough estimate of the total body surface area (TBSA) burned. Note that adults and children are different. This formula frequently overestimates the extent of a burn in clinical practice. As a rough guide, the area covered by the individual’s palm is approximately 1.25% TBSA. See Fig. 38.4 for a more accurate method of determining TBSA burned in children.

![Figure 38.5](image.png)

**Figure 38.6** Zones of injury after a burn. The zone of coagulation is the portion that is irreversibly injured. The zones of stasis and hyperemia are defined in response to the injury. (From Townsend CM, Beauchamp RD, Evers BM, et al., editors: Sabiston textbook of surgery, ed 19, St. Louis, 2012, Saunders.)

**OUTPATIENT VERSUS INPATIENT CARE**

One of the first steps in minor burn care is to select patients for whom outpatient care is appropriate. For patients determined to require inpatient care, the decision to admit or transfer the patient depends on the burn care capabilities of the initial treating facility. Guidelines set forth by the ABA\(^7\) regarding criteria for referral to a burn center are listed hereafter. Burn injuries that should be considered for referral to a burn center include the following:

1. Partial thickness burns greater than 10% TBSA
2. Burns involving the face, hands, feet, genitalia, perineum, or major joints
3. Third-degree burns in any age group
4. Electrical burns, including lightning injury
5. Chemical burns
6. Inhalation injury
7. Burn injury in patients with preexisting medical disorders that could complicate management, prolong recovery, or affect mortality
8. Any patient with burns and concomitant trauma in which the burn injury poses the greatest risk for morbidity or mortality after emergency or surgical stabilization of the traumatic injuries. In such cases if the trauma poses the greatest immediate risk, the patient’s condition may be stabilized initially in a trauma center before transfer to a burn center. Physician judgment will be necessary in such situations and should be in concert with the regional medical control plan and triage protocols.
9. Burned children in hospitals without qualified personnel or equipment for the care of children
10. Burn injury in patients who will require special social, emotional, or rehabilitative intervention

Note, the previously-mentioned guidelines are not absolute clinical mandates and clinical judgment and patient characteristics may allow certain patients listed in these categories to be treated appropriately without transfer to a burn center.\(^8\)

If in doubt, the decision to admit a patient with an acute burn injury is rarely inappropriate. Candidates who can be considered for outpatient treatment are generally adults and children meeting the ABA criteria for minor burn criteria. Burns usually better managed initially on an inpatient basis are large or deep burns involving the hands, face, feet, neck, or perineum; burns resulting from abuse or attempted suicide; burns occurring in association with other trauma or inhalation injuries; and chemical or electrical burns.

Patients who are at risk for poor outcomes with even minor burns include patients with concomitant medical problems such as diabetes mellitus, peripheral vascular disease, congestive heart failure, and end-stage renal disease; patients taking steroids or other immunosuppressive agents; patients who are very young or very old; those who are mentally impaired or have drug and alcohol dependency; homeless persons; those who are malnourished; and patients without a sufficient home support system. Whereas very minor burns in these patients may still be treated appropriately in the outpatient setting, inpatient treatment might be necessary in these circumstances even though the burn might be considered “minor” by ABA criteria. Other admission considerations include pain control, the ability to return for follow-up care, the degree of incapacity, the ability to receive wound care at home, and the overall social situation; all should influence the final decision of whether admission or transfer is warranted.\(^9\)
Additional guidelines that can guide emergency providers in determining the need for admission following an acute burn injury include the following:

1. Patients requiring intravenous (IV) access. Following a burn there is an immediate capillary leak of plasma-like fluid that can last for 18 to 24 hours. In burns involving greater than 20% TBSA, the leak occurs in both burned and nonburned tissues. If not replaced, this fluid loss can lead to hypovolemic shock and renal failure. IV fluid resuscitation is indicated for all patients with second- and third-degree burns greater than 10% TBSA, in patients younger than 10 or older than 50 years, and for burns greater than 20% TBSA in all other age groups.

2. Anticipated surgery. Deep burns are best treated by early surgical excision and skin grafting. This permits faster wound healing, provides more stable skin, and reduces contractures. Hospital admission facilitates wound care and preparation for surgery.

3. Respiratory problems. Patients with respiratory distress requiring oxygen therapy and those suspected of inhaling toxic fumes or vapors should be admitted for observation or intubation and for mechanical ventilation (Box 38.1). Direct bronchoscopic evaluation of the airway may assist in the evaluation and diagnosis of tracheobronchitis or pneumonitis from a smoke inhalation injury.

4. Need for special nursing care. Specialized wound care, dressings, and nursing care are often required for burns involving the face, hands, feet, perineum, and genitalia and may best be treated on an inpatient basis if outpatient care is not feasible or ideal based on the case specifics. Patients unable to care for themselves or those lacking family and friends able to assist them may also require admission.

5. Special burn injuries. Chemical injuries are often more severe than the initial examination would suggest. Unlike thermal burns, tissue destruction may continue many hours after injury. Patients with chemical injuries should be admitted whenever the injuries are of indeterminate depth, affect a large area, or are deep and require surgical excision or if there are systemic manifestations of chemical toxicity or when the chemical responsible for injury requires a specific antidote. Swelling from deep circumferential burns may constrict the chest or limbs and result in compartment syndrome. Such burns should be monitored frequently to determine the potential need for escharotomy or fasciotomy (Fig. 38.7). Whenever the patient’s condition prevents a reliable clinical examination, direct measurement of compartment pressures can provide an objective measurement of intracompartmental pressure and assist in the decision to perform these surgical procedures (see Chapter 54). Adult and pediatric patients with extensive burn injuries that require fluid resuscitation with large volumes of crystalloid should be monitored closely for the development of abdominal compartment syndrome. Patients with mechanical burns involving large areas of skin loss or with significant frostbite injuries are often admitted for specialized wound care and parenteral pain management. Patients with certain electrical injuries may require admission for cardiac monitoring, specialized wound care, or IV fluids.

**PROCEDURE**

**Emergency Treatment**

Home or field treatment and ED care overlap. Initial treatment of a thermal injury begins immediately following the burn. If safe to do so, patients should be rapidly removed from the source of injury. Flames are extinguished by smothering the fire with a blanket, jacket, or equivalent item; by dousing the fire with water; or by using a chemical fire extinguisher. Most chemical injuries are best treated by irrigating the affected area with copious quantities of clean water. Patients with electrical injuries are removed from contact with the electrical source as soon as it is safe to do so.

Cooling is most beneficial for small burns if started within 3 minutes of injury and possibly of additional benefit if continued for the first few hours after the burn. Doing so has been shown to reduce pain significantly and can limit tissue damage by decreasing thromboxane production. When cooling a burn...
wound, it is important to avoid hypothermia or freezing of tissue because this may deepen the injury. In the field, room-temperature or cold tap water irrigation, immersion, or compresses (20°C to 25°C) will provide some pain relief without the risk of further injuring burned tissues and inducing hypothermia, which can occur with iced solutions (Fig. 38.8). Placing ice on a burn should not be done. Sterile dressings are not required for field treatment; a moist towel or nonadherent sheet may be used. Nonmentholated shaving cream makes an excellent temporary covering for out-of-hospital use if a sheet may be used. Nonmentholated shaving cream makes an excellent temporary covering for out-of-hospital use if a sheet may be used. Towels are not required for field treatment; a moist towel or nonadherent sheet may be used. Nonmentholated shaving cream makes an excellent temporary covering for out-of-hospital use if a sheet may be used. Towels are best avoided but are probably benign. Remove jewelry and gross debris in the field if possible.

Initial Care of Major Burns

Major burns require the specialized resources of a burn center. In such cases, emergency providers should initiate the resuscitation, consult the burn center for referral, and transfer the patient as soon as practically possible. Initial resuscitation should follow standardized trauma protocols, including a primary and secondary survey, and provide immediate interventions directed at airway management, breathing, and cardiovascular support, as needed. IV catheters can be inserted initially through burned skin when unburned sites are unavailable. Early IV access permits the administration of resuscitative fluids, medications, and parenteral narcotics to relieve the pain. Patients with burns exceeding 20% TBSA should receive IV fluid resuscitation with lactated Ringer’s (LR) solution based on various available formulas. LR solution is preferred because large saline infusions will produce hyperchloremic acidosis. One need not use bolus fluid resuscitation as fluids are lost via capillary leak and via the raw skin, and significant fluid loss does not occur within the first hour.

Historically, the Parkland or Brooke formulas have been used to estimate fluid needs. It should be noted that the 2011 Advanced Burn Life Support (ABLS) Manual specifies that research has shown that the Parkland formula’s 4 mL/kg × % TBSA formula commonly results in excessive edema formation and over-resuscitation. Whether this is because first-degree burns may have been erroneously included in the %TBSA or other reasons is unclear. For adults the ABLS advises a total of 2 mL LR × body weight in kilograms × % second- and third-degree burns in the first 24 hours. For pediatric patients (under age 14 and weight less than 40 kg), the ABLS advises a total of 3 mL LR × body weight in kilograms × % second- and third-degree burns in the first 24 hours. It should be noted that 24 hours starts at the time of the burn and not the time medical care begins. After the total amount is calculated, fluids should be administered as a constant infusion rather than by bolus administration. Half of the calculated fluid requirement should be given in the first 8 hours and the remaining half provided over the following 16 hours (Box 38.2). Lastly, these fluid quantities are guidelines and urine output monitoring will assist in adjusting the rate or volume to meet the target outputs of 0.5 mL/kg per hr for adults and 1 mL/kg per hr for children weighing under 40 kg. According to the ABA, Foley catheter placement is advised in major burns to accurately monitor urine output.

More recently, the US Army Institute of Surgical Research has advocated a simpler formula for estimating hourly fluid requirements in burn patients. This simpler formula may be more useful for prehospital providers or ED resuscitation (Box 38.3). The formulas estimate hourly fluid requirements and must be adjusted up or down to achieve a urine output of 0.5 to 1.0 mL/hr. Insertion of a Foley catheter is usually necessary to accurately measure hourly urine output.

Patients exposed to carbon monoxide should have carboxyhemoglobin levels measured and empirically receive 100% oxygen. The duration of oxygen administration will depend on the level and symptoms of the individual patient, but often

**Figure 38.8** To cool a burn that cannot easily be immersed in water, cover the area with unfolded gauze pads that have been soaked in room-temperature saline. Continue to frequently soak the gauze with cool saline or tap water drawn up in a syringe. Adding a few ice chips to the liquid is helpful, but do not cover the burn with ice. Towels are generally too bulky for this procedure. Narcotics are the best way to control pain in any burn, and should be used liberally.

**Box 38.2 ABA ABLS Initial Fluid Recommendations for Burn Patients in First 24 Hours**

1. Lactated Ringer’s (LR) is the fluid of choice.
2. Estimate %TBSA (count second and third degree only)
4. Utilize appropriate formula as detailed below:
   - **Adults with Thermal and Chemical Burns:**
     - 2 mL LR × patient’s body weight in kg × %TBSA (second and third degree only)
   - **Children 14 years or younger and body weight of less than 40 kg:**
     - 3 mL LR × child’s weight in kg × %TBSA (second and third degree only)-add normal maintenance fluid to the fluid requirements, consider including dextrose in maintenance fluids for infants and young children who may rapidly deplete glycogen stores, particularly for those 2 years and younger.
   - **Adult patients with High Voltage Electrical Injuries**
     - If there is evidence of deep tissue injury or red pigments in urine use:
       - 4 mL LR × patient’s body weight in kg × %TBSA (second and third degree only)
     - 5. Infuse half of the calculated 24-hour fluid needs in the first 8 hours following the burn. Infuse the second half of the 24-hour fluid needs in the next 16 hours (hours 9–24)
     - 6. Increase or decrease the fluid rate based on the patient’s urine output (and pigment). Adjust fluids to maintain urine output at 0.5 mL/kg per hr in adults and 1 mL/kg per hr in children.

ABA, American Burn Association; ABLS, advanced burn life support; TBSA, total body surface area.
Simplified ED Burn Fluid Resuscitation: the Rule of 10

1. Estimate burn size (%TBSA) to the nearest 10.
2. %TBSA × 10 = initial fluid rate in mL/hr (for adult patients weighing 40 to 80 kg).
3. For every 10 kg above 80 kg, increase the rate by 100 mL/hr.

BOX 38.4 Baux Score and Modified Baux Score to Predict Mortality in Burn Injury

Original Baux Score: % Mortality = Age + %TBSA
Modified Baux Score: % Mortality = Age + %TBSA + 17*(Inhalation Injury, Yes=1, No=0)

Plot of the observed mortality versus the group midpoint computed within 10 groups of the revised Baux score.


require 6 hours or more. Once considered a traditional empirical treatment, there is no evidence-based proven benefit from hyperbaric oxygen therapy for carbon monoxide poisoning. A Cochrane review (http://www.summaries.cochrane.org/CD002042) concluded that there is insufficient evidence to support the use of hyperbaric oxygen for the treatment of patients with carbon monoxide poisoning. The Cochrane review of published trials found conflicting, potentially biased, and generally weak evidence regarding the usefulness of hyperbaric oxygen for the prevention of neurologic injury. Per an evidence-based analysis, existing randomized trials do not establish whether the administration of hyperbaric oxygen to patients with carbon monoxide poisoning reduces the incidence of adverse neurologic outcomes. Because there may still be advocates of hyperbaric oxygen therapy, consultation with a local hyperbaric center is reasonable in certain cases, but it is not standard that this intervention be routinely implemented. Critically ill and pregnant patients are still often offered hyperbaric treatment, but controversy over the efficacy and safety persists even for these subgroups.

Cyanide is released from mattress fires and burning of upholstery. Patients suspected of having been exposed to significant levels of cyanide and manifesting symptoms should receive hydroxocobalamin (Cyanokit [Meridian Medical Technologies, Inc., Columbia, MD]). If not available, the Cyanide Antidote Package may be used despite lack of proven benefit of this traditional cyanide therapy. It is reasonable to empirically administer hydroxocobalamin or the sodium thiosulfate portion of the cyanide kit to burn victims in coma or to those exhibiting metabolic (lactic) acidosis after smoke exposure.

In addition to the logical increased mortality with larger areas of TBSA involved, the mortality of patients with burns is also increased when concomitant inhalation is present. The Baux Score was developed a half-century ago by Professor Serge Baux to estimate the mortality of burns. This score was updated in 2010 using data from the National Burn Registry to include the effect of inhalation injury to mortality (Box 38.4). In short, the original Baux score became outdated because advances in care of the burn victim which has improved the mortality of patients. The modified Baux score is now more accurate for modern burn survival rates.

Burn patients have an impaired ability to regulate their core body temperature and will quickly become hypothermic if untreated. Core temperature should be measured frequently, and active and passive warming strategies should be implemented to prevent hypothermia from developing. This can include minimizing exposure by covering patients with sheets and blankets, warming IV fluids, warming the room, or applying radiant or convective warming systems. In anticipation of transfer to a burn center or before surgical consultation, remove any wet cooling dressings that may have been applied initially and cover the wounds with dry gauze dressings.

Initial Care of Minor Burns

Prompt cooling of the burned part is an almost instinctive response and is one of the oldest recorded burn treatments, having been recommended by Galen (AD 129–199) and Rhazes (AD 852–923). In the ED, room-temperature or cold tap water irrigation, immersion, or compresses (20°C to 25°C) are optimal in obtaining pain relief and providing some measure of protection for burned tissues without the problems of hypothermia that iced solutions can cause. If not done before ED treatment, immediate cold water immersion may still have some ability to limit the extent of a burn and will provide significant pain relief. It is acceptable to add a few ice chips to the water, but packing the wound in ice must be avoided.

All involved clothing and jewelry (such as rings), along with any gross debris, should be removed from the burned area. Chemical burns involving the skin or eyes require prolonged tap water irrigation. The burn should otherwise be covered with a moist, sterile dressing. In the ED and prehospital phase, appropriate analgesics, usually narcotics, are the best way to control pain and should not be forgotten in the initial phase of burn care. The burned area may be immersed immediately in room-temperature water or covered with gauze pads soaked in room-temperature water or saline (see Fig. 38.8). The gauze may be kept cool and moist to provide continued pain relief; the patient will let the clinician know when additional cooling is desired. Many clinicians use sterile saline for cooling, but it has no proven benefit over tap water, even when the skin is broken. It is acceptable to add ice chips to water or saline to lower the temperature. As stated previously, immersion of burned tissue in ice or ice water should be avoided because ice immersion increases pain and risks frostbite injury or systemic hypothermia.
to remove only grossly devitalized tissue. Additional débridement of the wound can take place, if needed, during subsequent follow-up visits when the wound has matured.
3. Consider applying a layer of antibiotic cream or ointment such as 1% silver sulfadiazine (Silvadene [Pfizer Inc., New York, NY]) or bacitracin directly to the wound (see Fig. 38.10D).
4. Apply fine-mesh gauze or commercial nonadherent gauze such as Adaptic or petrolatum gauze impregnated with 3% bismuth tribromophenate (Xeroform [DeRoyal, Powell, TN]) to the burn wound.
5. Cover and pad the wound with loose gauze fluffs. If fingers and toes are involved, pad the web spaces and the digits individually and separate them with strips of gauze (see Figs. 38.9D and 38.10E and F). Wrap the entire dressing snugly (but not tightly) with an absorbent, slightly elastic material such as Kerlix (Medtronic, Minneapolis, MN).
6. Instruct the patient to elevate the affected limb to prevent swelling, which may cause delayed burn conversion or wound infection.

**Open Burn Care**
Following cleansing of the wound with chlorhexidine soap and débridement of blisters and any loose skin, wounds that are not amenable to a dressing, such as those on the face, can be managed initially by the application of a bland topical antibiotic such as bacitracin. The wound can be washed two or three times per day, followed by reapplication of the topical agent. This is the preferred method for managing burns on the face and neck.

**Outpatient Care of Minor Burns**
Minor burns are generally those that will heal spontaneously and do not require surgery or specialized wound care. These wounds are not associated with immunosuppression or hypermetabolism, nor are they highly susceptible to infection, a quality associated with larger burns.2829 Treated conservatively, most minor burns will heal without significant scarring. Many complications seen with minor burns are thought to result from overtreatment rather than undertreatment. Examples include the use of dry dressings that can adhere to newly forming skin and secondary infections from the overzealous use of topical or systemic antibiotics. The most important characteristic of a dressing is that it controls fluids within the wound. Burn dressings that keep the surface of the wound moist and avoid pooling of fluids will speed healing.32 The best material for this purpose is a generous use of topical or systemic antibiotics.

Limit extent of thermal damage
Reduction or cessation of pain
Elimination of local hyperthermia
Inhibition of postburn tissue destruction
Decreased edema
Reduced metabolism and toxin production

The potential benefits of burn cooling are listed in Box 38.5. Because most patients with minor burns seek medical attention after initial self-instituted prehospital cooling, it is unlikely that the clinician can favorably affect the burned tissue with any intervention in the ED. With the exception of pain relief and removal of debris, the primary benefits of burn cooling are probably experienced only if the burn is cooled promptly, within the first 3 minutes after injury, thus making home care important.28,29 Minor burns are considered tetanus prone, and tetanus toxoid should be administered if patients are unsure of their tetanus immunization status or when it has been more than 10 years since the last immunization. Nonimmunized patients should receive tetanus toxoid and immunization with subsequent boosters in accordance with current guidelines.

**Biologic Dressings**
Biologic dressings are natural tissues, including skin that consists of collagen sheets containing elastin and lipid. They are not routinely used in the emergency care of minor wounds and are primarily treatment options in burn centers. The benefits of biologic dressings include a reduction in surface bacterial colonization, diminished fluid and heat loss, avoidance of further wound contamination, and prevention of damage to newly developed granulation tissue. Examples of biologic dressings include cadaveric human skin and commercially available porcine xenograft or collagen sheets.

**Synthetic Dressings**
Synthetic dressings are manufactured in various forms. Film-type dressings have a homogeneous structure and are usually polymers. Because these dressings are nonpermeable, problems with retention of wound exudates have occurred. Some second-generation dressings have been developed to address these problems. These products include Tegaderm (3M Medical, St. Paul, MN), Vigilon (Bard Medical, Covington, GA), DuoDERM (Convatec, Bridgewater, NJ), Biobrane (Smith & Nephew, Op-Site (Smith & Nephew, Inc., Andover, MA), Acticoat (Smith & Nephew), Aquacel products (Convatec, Greensboro, NC), and TransCyte (Advanced Tissue Sciences, La Jolla, CA).3233 These preparations have theoretical benefits under certain circumstances, but are not proven to have superior performance over simple gauze dressings for minor outpatient burns. These products are most often used by burn centers and have little applicability for minor burns in patients discharged from the ED. For patients admitted or transferred to a burn center, simple gauze dressings are appropriate. Some

**BOX 38.5 Advantages of Prompt Burn Cooling**

Limit extent of thermal damage
Reduction or cessation of pain
Elimination of local hyperthermia
Inhibition of postburn tissue destruction
Decreased edema
Reduced metabolism and toxin production

The potential benefits of burn cooling are listed in Box 38.5. Because most patients with minor burns seek medical attention after initial self-instituted prehospital cooling, it is unlikely that the clinician can favorably affect the burned tissue with any intervention in the ED. With the exception of pain relief and removal of debris, the primary benefits of burn cooling are probably experienced only if the burn is cooled promptly, within the first 3 minutes after injury, thus making home care important.28,29 Minor burns are considered tetanus prone, and tetanus toxoid should be administered if patients are unsure of their tetanus immunization status or when it has been more than 10 years since the last immunization. Nonimmunized patients should receive tetanus toxoid and immunization with subsequent boosters in accordance with current guidelines.

**Burn Dressings**
Biologic dressings are natural tissues, including skin that consists of collagen sheets containing elastin and lipid. They are not routinely used in the emergency care of minor wounds and are primarily treatment options in burn centers. The benefits of biologic dressings include a reduction in surface bacterial colonization, diminished fluid and heat loss, avoidance of further wound contamination, and prevention of damage to newly developed granulation tissue. Examples of biologic dressings include cadaveric human skin and commercially available porcine xenograft or collagen sheets.

**Synthetic Dressings**
Synthetic dressings are manufactured in various forms. Film-type dressings have a homogeneous structure and are usually polymers. Because these dressings are nonpermeable, problems with retention of wound exudates have occurred. Some second-generation dressings have been developed to address these problems. These products include Tegaderm (3M Medical, St. Paul, MN), Vigilon (Bard Medical, Covington, GA), DuoDERM (Convatec, Bridgewater, NJ), Biobrane (Smith & Nephew, Op-Site (Smith & Nephew, Inc., Andover, MA), Acticoat (Smith & Nephew), Aquacel products (Convatec, Greensboro, NC), and TransCyte (Advanced Tissue Sciences, La Jolla, CA).3233 These preparations have theoretical benefits under certain circumstances, but are not proven to have superior performance over simple gauze dressings for minor outpatient burns. These products are most often used by burn centers and have little applicability for minor burns in patients discharged from the ED. For patients admitted or transferred to a burn center, simple gauze dressings are appropriate. Some
procedures such as débridement and dressing changes are planned. We prefer to use IV opioids (occasionally supplemented with a short-acting benzodiazepine such as midazolam) for all painful procedures. For complicated débridement or dressing changes adequate analgesia is a minimum requirement with some patients requiring procedural sedation (see Chapter 33). Regional or nerve block anesthesia is an excellent alternative when practical, and if feasible, nitrous oxide analgesia may be used. Ketamine may also be a reasonable alternative. Oral opioids may be inadequate for the initial treatment of significant pain but can be used for continued outpatient analgesia. Local anesthetics may be injected in small quantities when appropriate, such as for the débridement of a deep ulcer or other small burn. Topical analgesics have no role in burn care. A properly designed dressing will do much toward preventing further discomfort after release home; however, home burn care and dressing changes may be quite painful. For this reason, an adequate supply of an oral opioid analgesic should be provided, and responsibility in analgesic use should be encouraged.

burn centers prefer that topical agents not be applied before transfer so that the full extent of the burn can be assessed immediately.

**Specific Clinical Issues in Minor Burn Care**

**Analgesia**

Pain is a critical feature of any burn injury. Relief of pain by the appropriate and judicious use of narcotic analgesics is of the utmost importance in the initial care of all burn patients. Prehospital narcotics are very appropriate when standard contraindications do not exist. Analgesia should be provided before extensive examination or débridement is performed. Inadequate analgesia is probably the most common ED error in the treatment of burn injuries, especially when burns occur in children. Parenteral narcotic analgesics have been erroneously relegated to pain control only for major burns, but it is suggested that narcotics be generously administered in the initial treatment of even minor painful burns.

Parenteral opioids (e.g., fentanyl, 1 to 2 µg/kg, or morphine, 0.1 to 0.2 mg/kg) are usually required, especially if painful procedures such as débridement and dressing changes are planned. We prefer to use IV opioids (occasionally supplemented with a short-acting benzodiazepine such as midazolam) for all painful procedures. For complicated débridement or dressing changes adequate analgesia is a minimum requirement with some patients requiring procedural sedation (see Chapter 33).

Regional or nerve block anesthesia is an excellent alternative when practical, and if feasible, nitrous oxide analgesia may be used. Ketamine may also be a reasonable alternative. Oral opioids may be inadequate for the initial treatment of significant pain but can be used for continued outpatient analgesia. Local anesthetics may be injected in small quantities when appropriate, such as for the débridement of a deep ulcer or other small burn. Topical analgesics have no role in burn care. A properly designed dressing will do much toward preventing further discomfort after release home; however, home burn care and dressing changes may be quite painful. For this reason, an adequate supply of an oral opioid analgesic should be provided, and responsibility in analgesic use should be encouraged.
Pruritus

Postburn pruritus is one of the most common and distressing complications of burn injury and is estimated to affect 87% of burns. It typically develops in the subacute phase and is therefore not a common issue for the emergency provider in the acute treatment phase. Despite the limited literature on the treatment of postburn pruritus, available therapies include oral antihistamines, topical antihistamines, and topical moisturizers. The use of topical therapies should be withheld until sufficient wound healing has occurred.
Edema

Minor burns lead to immediate inflammation mediated by the release of histamine and bradykinins, which cause localized derangements in vascular permeability with resultant burn wound edema. This edema may be harmful in several ways. First, the increase in interstitial fluid increases the diffusion distance of oxygen from capillaries to cells and thereby increases hypoxia in an already ischemic wound. Second, the edema may produce untoward hemodynamic effects by a purely mechanical mechanism: compression of vessels in muscular compartments. Third, edema has been associated with the inactivation of streptococcidial skin fatty acids, thus predisposing the patient to burn cellulitis. 35,36

Successful management of burn edema hinges on immobilization and elevation. Most patients are unfamiliar with the medical definition of elevation and are not aware or convinced of its value. Patient education in this regard is critical; however, certain burns (e.g., burns in dependent body areas) are prone to edema despite everyone’s best intentions. It is for this reason that lower extremity burns in general and foot burns in particular are prone to problems. Major burns of the hand should be elevated while the patient is still in the ED. This is most readily accomplished by hanging the injured hand from an IV pole with a stockinette to support the bandaged hand (Fig. 38.11).

Use of Topical Preparations and Antimicrobials

Minor burns result in insignificant impairment of normal host immunologic defenses, and burn wound infection is not usually a significant problem. Topical antimicrobials are often used; however, some believe that these agents may actually impair wound healing. 16 Although the procedure is of unproven value, many clinicians routinely use antibiotic creams or ointments on even the most minor burns. Most patients expect some type of topical concoction, so a discussion of their use—or nonuse—is prudent.

Topical antimicrobials were designed for the prevention and care of burn wound sepsis or wound infection, primarily in hospitalized patients with major burns, and there is no convincing evidence that their use alters the course of first-degree burns and superficial partial-thickness injuries. As noted, the burn dressing is the key factor in minimizing complications in all burns. Nonetheless, topical antimicrobials are often soothing to minor burns, and their daily use prompts the patient to look at the wound, assess healing, perform prescribed dressing changes, or otherwise become personally involved in the care. Keep in mind that if a topical antimicrobial is used, its effectiveness is decreased in the presence of proteinaceous exudate, thus necessitating regular dressing changes if the antimicrobial benefit of topical therapy is to be realized. In reality, once-daily dressing changes are more practical and are commonly prescribed, and no data indicate that this regimen is inferior to more frequent dressing changes.

All full-thickness burns should receive topical antimicrobial therapy because the eschar and burn exudate are potentially good bacterial culture media and deep escharotic or subescharotic infections may not be easily detected until further damage is done. All deep partial-thickness injuries likewise benefit from the application of a topical antimicrobial. As stated, this intervention can await definitive therapy in a burn unit.

Criteria for choosing a specific topical agent include its’ in vitro and clinical efficacy, toxicity (absorption), superinfection rate, ease and flexibility of use, cost, patient acceptance, and side effects. It is important to note that no firm scientific data convincingly support the use of any specific topical antimicrobial for minor outpatient burns.

Specific Topical Agents

**Silver Sulfadiazine (Silvadene, Pfizer).** This poorly soluble compound is synthesized by reacting silver nitrate with sodium sulfadiazine. It is the most commonly used topical agent for outpatients, and it is well tolerated by most patients (Fig. 38.12A; also see Fig. 38.10D). It has virtually no systemic effects and moderate eschar penetration, and it is painless on application. Although silver sulfadiazine is commonly used, its popularity is waning. It must be applied daily and the thick white cream can be difficult to remove. There are no well-designed studies confirming improved burn healing or a reduced rate of infection with silver sulfadiazine. It may impede reepithelialization and should be stopped when this occurs. Hence, many burn specialists prefer plain bacitracin ointment as the topical of choice because of its cost, equal efficacy, and good patient acceptance.

Silver sulfadiazine is available as a “micronized” mixture with a water-soluble white cream base in a 1% concentration that provides 30 mEq/L of elemental silver. It does not stain clothes, is not irritating to mucous membranes, and washes off with water. It may be used on the face, but such use may be cosmetically undesirable for open treatment. It should not be used near the eyes. Its broad gram-positive and gram-negative antimicrobial spectrum includes β-hemolytic streptococci, *Staphylococcus aureus* and *Staphylococcus epidermidis*, *Pseudomonas spp.*, *Proteus spp.*, *Klebsiella spp.*, Enterobacteriaceae, *Escherichia coli*, *Candida albicans*, and possibly herpesvirus hominis.

![Figure 38.11](image-url) Begin elevation of a burned hand in the emergency department. After a hand dressing is applied, suspend the arm from an intravenous pole with stockinette. A plaster or fiberglass splint may also be incorporated into the dressing.
BROAD-SPECTRUM ANTIBIOTIC OINTMENTS. Many nonprescription topical antimicrobials are used for minor burn therapy despite a paucity of data attesting to specific benefits. Included are bacitracin zinc ointment, polymyxin B–bacitracin (Polysporin, Johnson & Johnson, New Brunswick, NJ). These are all soothing, cosmetically acceptable for open treatment (such as on the face), and effective antiseptics under burn dressings. Some researchers caution against agents containing neomycin because of a potential for sensitization (see Fig. 38.12B). Though commonly applied by patients without adverse effects, we advise against the use of topicals that contain neomycin (Neosporin, Johnson & Johnson) because of the potential for contact dermatitis. The authors suggest plain bacitracin or Polysporin ointment as the routine topical agents for most burns, although Silvadene is a very acceptable, albeit more expensive alternative.

ALOE VERA CREAM. Aloe vera cream is commercially available in a 50% or higher concentration with a preservative. It exhibits antibacterial activity against at least four common burn wound pathogens: *Pseudomonas aeruginosa*, *Enterobacter aerogenes*, *S. aureus*, and *Klebsiella pneumoniae*. Heck and coworkers and others37,38 compared a commercial aloe vera cream with silver sulfadiazine in 18 patients with minor burns. Healing times were found to be similar, and there was no increase in wound colonization in the aloe vera group in comparison to patients treated with silver sulfadiazine. Other authors have promulgated the use of aloe gel preparations for minor burns.38 Aloe vera cream is an acceptable, inexpensive option for open or dressed outpatient care of minor burns.

HONEY. Honey has long been advocated as an inexpensive and effective topical treatment for minor outpatient burns. The physicochemical properties of honey (osmotic effect, pH) give this substance the antibacterial and antiinflammatory properties that support its use. It may be superior to silver sulfadiazine with regard to minor burn wound healing. Honey is not widely used, but it has been promulgated as a safe, effective, and inexpensive dressing for the outpatient management of burn wounds.39-41

CORTICOSTEROIDS. High-potency topical steroid preparations have no beneficial effects on the rate of healing or limitation of scarring of thermal burns. Though probably not harmful in most cases, their use is not supported.12

FOLLOW-UP CARE OF MINOR BURNS

The specifics of outpatient follow-up of minor burns are controversial and often based on clinician preference and personal bias rather than on firm scientific data. Follow-up should be individualized for each patient and should be based on the reliability of the patient, the extent of the injury, the frequency and complexity of dressing changes, and the amount of discomfort anticipated during a dressing change. Depending on the characteristics of the patient and the resources available to them as an outpatient follow-up plans may include primary care physician rechecks, follow-up at wound care center, home health visits by wound care team, or even return to an ED or “fast-track” setting. Outpatient physical therapy departments or wound care centers often have excellent facilities to monitor outpatient burns with clinician oversight.
If a topical antibiotic agent is used, the dressing should be changed daily with removal and reap苹plication of the topical preparation. The wound should be rechecked by a clinician after 2 to 3 days and periodically thereafter, depending on wound size, compliance, healing, and other social issues. If a dry dressing is chosen, follow-up every 3 to 5 days is usually adequate. The purpose of any burn dressing changes or home care regimen is defeated if the patient cannot afford the material or is not instructed in the specifics of burn care. Many EDs supply burn dressing material on patient release. A complete pack includes antibiotic ointment or cream, gauze pads (fluffs), an absorbent gauze roll, a sterile tongue blade to apply the cream, and tape (Fig. 38.13). Providing limited supplies of the items necessary for dressing changes may enhance compliance with follow-up if the patient has to return for additional supplies. Writing a prescription and merely stating that the dressing should be changed daily may not be sufficient.

Daily home care can be performed by the patient with help from a family member or visiting nurse (Box 38.6). The dressing may be removed each day and the burn area gently washed with a clean cloth or a gauze pad, tap water, and a bland soap. Sterile saline and expensive prescription soaps are not required. A tub or shower is an ideal place to gently wash off burn cream. The area can be gently washed with mild soap and a clean cloth or gauze pads.

Outpatient Physical Therapy for Burn Care
When the hospital's outpatient physical therapy department or wound care center is equipped to treat minor burns, it is prudent to consider this option as a means of longitudinal follow-up. Many centers make available daily or periodic burn treatment consisting of dressing changes, whirlpool débridement, and range-of-motion exercises. An additional advantage is that medically trained personnel evaluate the burn daily, thereby decreasing clinician visits and enabling identification of problems before serious complications develop. Similar services are available in many areas where providers visit the patient in their home to perform wound/burn care.

Burn Wound Healing
Burn wound healing differs from healing of other soft tissue wounds. The duration is variable but is often proportional to burn depth. Within 1 to 3 weeks and following the initial inflammatory response, neovascularization of the burn occurs and is accompanied by fibroblast migration. Collagen production begins but it is often deposited randomly, thereby leading to scar formation. Reepithelialization follows collagen deposition. The persistence of necrotic tissue and eschar in the wound will impede all aspects of healing. The extent of scar formation is related directly to healing time. Wound healing that occurs

1. Take pain medicine 1/2 hour before dressing change if you find dressing changes to be painful.
2. If the burn is on the hand, foot, or other areas that are difficult to reach, have someone help you.
3. Have all materials available. Gloves may be worn.
4. Remove the dressing and rinse off all burn cream or ointment with tap water, under a shower, or in the bathtub. The area can be gently washed with mild soap and a clean cloth or gauze pads.
5. Look at the burn and assess the healing, blistering, and amount of swelling. Note any signs of infection.
6. Gently exercise the area through range of motion.
7. Apply the burn ointment with a sterile tongue blade.
8. Cover the cream with fluffed-up gauze.
9. Wrap the area in bulky gauze.
10. Repeat this dress change daily.

BOX 38.6 How to Change a Burn Dressing at Home: Patient Instructions

Figure 38.13 Providing patients with burn dressing material on discharge encourages proper home care. Dispensing only limited supplies of the items may enhance compliance with follow-up visits.
in fewer than 16 days often results in decreased scar formation. Proper wound débridement is also associated with faster wound healing and minimizes scar formation.

Healing of superficial burns occurs by reepithelialization from the wound edge and from residual dermal elements containing epithelial cells. This process generally requires 10 to 14 days. After healing, the initial epithelial layer is often fragile and is easily reinjured. The application of bland, lanolin-containing creams for 4 to 8 weeks after initial healing may reduce dryness and cracking of the healing wound.

Deep burns lack residual dermal elements within the wound and heal by reepithelialization from the wound edge. Healing is slow and often unsatisfactory; it frequently takes longer than 3 weeks, and an unstable epithelium is produced that is prone to hypertrophic scarring and contractures. This is a particular problem in burns that extend across joints and limit motion. Burns that take longer than 2 to 3 weeks to heal are also prone to infection, which may be reduced by using topical antimicrobial agents. Deep wounds should be referred for surgical consultation and generally require early excision, grafting, and physical therapy.

**SPECIAL MINOR BURN CARE CIRCUMSTANCES**

**Blisters**

Management of blisters in minor burns is controversial. In reality, there is little one can do wrong when it comes to a clinical approach to blisters in minor burns (Fig. 38.14). Management arguments are generally theoretical or based on local tradition; the ultimate outcome of a minor burn is rarely determined by how one treats blisters. Intact blisters do offer some pain control and a physiologic dressing that rarely becomes infected; however, most large blisters spontaneously rupture after 3 to 5 days and eventually require débridement. When the integrity of the blister is breached, the fluid becomes a potential culture medium. Clinical choices for blister management include débridement (immediate and delayed) or simply leaving the blister intact. In general, ruptured blisters should be débrided. Needle aspiration of blisters should NOT be performed as this may increase the risk of infection.

Some studies suggest that intact burn blisters may allow reversal of capillary stasis and less tissue necrosis. Madden and colleagues showed that burn exudate (as contained within intact blisters) is beneficial in stimulating epidermal cell proliferation.

Swain and associates demonstrated that the density of burn colonization with microorganisms was much lower in minor burns with blisters left intact. They also found that 37% of patients with aspirated blisters (not recommended) experienced a reduction in pain versus none of those whose blisters were unroofed. Other investigators believe that undressed wounds with débrided blisters are subject to additional necrosis secondary to desiccation, which can convert a partial-thickness burn to a full-thickness injury. Finally, intact blisters clearly provide some pain relief, as evidenced by the sudden increase in pain immediately after débridement. Increased pain should be anticipated and analgesia offered as appropriate when débridement is necessary. We suggest the steps listed in Fig. 38.10 as a general approach to burn blisters.

**Figure 38.14** It is difficult to do anything wrong with minor burn blisters, and many regimens are acceptable. Blister aspiration is not supported. Eventually, however, any unroofed blisters will have to be débrided. A, An expeditious and relatively painless way to débride a burn is to use a dry gauze pad to grasp the dead skin and B, peel it off. Meticulous instrument débridement is often time-consuming and stressful to the patient. Be aware that pain occurs when air comes in contact with the débrided skin, and prophylactic analgesia should therefore be provided. Large burns can be débrided under procedural sedation.

**Minor Burn Infections**

Prophylactic systemic antibiotics are not warranted in the routine treatment of outpatient burns. It may be difficult to separate the erythema of the injury or healing process from cellulitis, but minor burns rarely become infected, with infection rates being well under 5%. There are bacteria on the skin at all times; normal skin usually harbors nonvirulent pathogens such as *S. epidermidis* and diphtheroids. Therefore, all burns are contaminated but not necessarily infected. Because thermal trauma results in coagulative necrosis, burn wounds contain a variable amount of necrotic tissue which, if infected, acts much as an undrained abscess and prevents access of antibiotics and host defense factors.

The microbial flora of outpatient burns varies with time after the burn. Shortly after injury, the burn becomes colonized with gram-positive bacteria such as *S. aureus* and *S. epidermidis*. After this period there is a gradual shift toward inclusion of gram-negative organisms, 80% of which originate from the patient’s own gastrointestinal tract. Common organisms seen on days 1 to 3 include *S. epidermidis*, β-hemolytic streptococci, *Bacillus subtilis*, *S. aureus*, enterococci, *Mima polymorpha*, *Enterobacter* spp., *Acinetobacter* spp., and *C. albicans*. One week after the burn these organisms may be seen along with *E. coli*, *P. aeruginosa*, *Serratia marcescens*, *K. pneumoniae*, and *Proteus vulgaris*. 
Anaerobic colonization of burn wounds is rare unless there is excessive devitalized tissue, as occurs with a high-voltage electrical injury. For this reason, routine anaerobic cultures are generally unnecessary in an assessment of infective organisms that produce minor infections.

A healing burn may produce leukocytosis and a mild fever in the absence of infection, especially in children. Early (days 1 to 5) burn infections are generally caused by gram-positive cocci, especially β-hemolytic streptococci. Streptococcal cellulitis is characterized by marked spreading erythema extending outward from the wound margins. Despite the plethora of organisms and the presence of some gram-negative pathogens in superficial burn cultures, standard gram-positive cellulitis coverage is appropriate initial therapy in most cases.

Effective topical treatment at the time of initial burn care and subsequent dressing changes is meant to delay bacterial colonization, maintain the bacterial density of wounds at low levels, and produce a less diverse wound flora. Because outpatient management of burns should be attempted only when the risk for infection is minimal, the use of systemic antibiotics is unnecessary for minor burns, even in the setting of delayed treatment, diabetes, and steroid use. Unnecessary antibiotic use may select for resistant organisms. Antibiotics in the management of minor burns have been recommended for patients undergoing an autograft procedure. There are no data on the use of antibiotics as prophylaxis for patients with burns in the setting of valvular heart disease.

In minor burn care, wound cultures are not required or recommended. It is useless, for example, to culture blister fluid in a patient who arrives for emergency care immediately after a thermal injury. Cultures are necessary only when overt infection develops, especially when it occurs when a topical or systemic antibiotic is being used. Cultures may also be of benefit when the infected wound is old, when hygiene is poor, or when there are preexisting abrasions nearby. Although they may adequately reflect wound flora, falsely sterile cultures are relatively frequent. In general, superficial cultures do not reflect deep burn flora and provide no quantitative information.

Sterile wound biopsy for culture is most satisfactory for the assessment of intraescharotic, subescharotic, or invasive infections and allows quantification of bacterial flora.

**Foot Burns**

Despite their relatively small surface area, foot burns tend to heal poorly, usually because of excessive edema; therefore, they are generally considered major burns. Foot burns are the most common burn category to fail outpatient therapy, and subsequently require admission and inpatient care (Fig. 38.15). Zachary and coworkers reported on a series of 104 patients with foot burns. In no patient admitted on the day of injury did burn cellulitis develop; in contrast, 27% of delayed-admission patients had cellulitis. Their study also noted a higher incidence of hypertrophic scarring with the need for skin grafting in the delayed-admission group. Overall, fewer days of hospitalization were required for the initially admitted group.

Specific problems in the care of foot burns include pain, wound drainage, difficulty changing dressings without help, inability of even motivated patients to comply with the requirements for elevation, and prolonged convalescence. The benefits of hospital admission include splinting, intensive local burn care, physical therapy, and bed rest with elevation, which minimizes edema. For these reasons, initial admission should be considered for all but the most minor of foot burns. Close outpatient follow-up wound checks are required for those foot burns that are most appropriate for outpatient care.

**Hand Burns**

Because of its functional significance, burns involving the hand can result in significant functional loss even when the TBSA burned is small. Losing the use of one or both hands can become seriously disabling and affect the patient’s activities of daily living regardless of whether the cause of the loss is a burn dressing, the late onset of scar contractures, or loss as a result of amputation.
As with other burns, the depth and extent of the burn determine the severity of the injury. The entire surface of one hand represents only 2.5% TBSA, yet even small burns can cause a disproportionate loss of function. Deep partial- or full-thickness hand burns, even if quite small, often warrant referral for early excision and grafting to limit scarring and maintain function. The skin on the dorsum of the hand is thinner than that on the palm and is more susceptible to burn injury so it must remain flexible to allow finger motion. Any exposed tendon or bone, such as may be seen with an electrical burn, constitutes a true fourth-degree injury, and either flap closure or amputation is required to heal the wound.

Many of the issues complicating outpatient management of foot burns are relevant to the care of hand burns. After initial burn cooling, the wound should be gently cleansed with mild soap. Any loose skin or ruptured blisters should be gently débrided, rinsed, patted dry, and covered with a topical antimicrobial agent and a nonadherent, bulky gauze dressing. The fingers should be carefully separated and bandaged individually. Small, intact blisters that do not interfere with hand function are prudent for all significant hand burns, particularly full-thickness burns on the dorsum of the hand should be splinted after bandaging to avoid the development of contractures or a boutonnière deformity.

Hospital admission or burn center referral should be considered for all significant hand burns, particularly full-thickness injuries and circumferential burns involving the digits (Fig. 38.16). If outpatient treatment is appropriate, the patient should have appropriate follow-up, must be given comprehensive instructions, and should have the resources available to perform daily dressing changes and range-of-motion exercises of the fingers and wrist during these dressing changes. An initial follow-up visit should occur in 48 to 72 hours, but the patient should be encouraged to return if burn cellulitis, worsening pain, fever, or lymphangitis develops. Ideally, the patient should be seen twice in the first week after injury and then, if clinically appropriate may be reduced to once a week until the burn is healed.

**Facial Burns**

Facial burns commonly result from unexpected ignition flash burns (e.g., from a stove, oven, or charcoal grill) or from car radiator accidents (Fig. 38.17A and B). Facial burns from these sources usually do well, but singeing of facial hair, significant edema, and pain often result. However, facial burns from these causes may produce airway problems and might require skin grafting. Singed nasal hairs or any sign of significant heat exposure to the face should prompt an evaluation of airway injury, which may result in airway compromise at a time later than the initial incident (see Fig. 38.17C).

Concurrent globe or corneal injury is quite rare because of protective blinking reflexes. If the eye is burned, it is usually in the setting of a life-threatening concomitant burn injury. Burns involving the eyelids can cause significant scarring. Fluorescein staining and slit-lamp examination should be performed to confirm or exclude the diagnosis of corneal injury (Fig. 38.18). Treatment of a corneal injury can involve irrigation, topical ophthalmic antibiotics, and consideration of eye patching versus protective soft contact lens (see Chapter 62). Referral to an ophthalmologist is usually prudent. Facial burns are otherwise treated in the usual fashion and with an open (no dressing) technique. Patients are instructed to wash their face two or three times a day with a mild soap and then apply a thin layer of antibiotic ointment, such as bacitracin zinc. Car radiator burns result from the combination of a hot liquid and steam burn (see Fig. 38.17B). Antifreeze exposure to skin does not produce a caustic injury, nor is it systemically absorbed. Neck burns are treated similarly.

A flash burn in a patient smoking cigarettes while using nasal oxygen causes a facial burn that is not uncommonly seen in the ED (see Fig. 38.17D). These burns may involve the nose and lips, may have melted plastic particles on the skin or in the nose, and can be quite painful. Although such patients generally do well, facial burns can make the continued use of nasal cannula oxygen problematic until healing takes place. Though not generally an inhalation burn issue, careful evaluation of the upper airways and assessment of lung function are prudent. Many patients using oxygen are relatively immunocompromised. They may not be able to tolerate even minor physical insults and have fragile conditions that require careful evaluation, short-term observation in some cases, and close follow-up for delayed healing and infection. Hospitalization for burn care and general supportive measures may be prudent for all but minor burns in this patient population. Minor burns can be treated on an outpatient basis in an open fashion with topical ointments, such as bacitracin.

All patients with head or neck burns should be evaluated carefully for a concomitant inhalation injury. Such patients may have direct evidence of injury, such as oral burns, blisters, soot, or hyperemia and a history of being in an enclosed space, or have indirect evidence, such as dyspnea, wheezing, arterial hypoxemia, or an elevated carboxyhemoglobin level. The definitive diagnostic test for inhalation injury is fiberoptic bronchoscopy. Flash ignition burns involving the face do not pose a problem with carbon monoxide poisoning, and although inhalation injuries generally do not occur with minor flash ignition burns, airway management should remain a consideration.

Inpatient care should be considered for all patients with significant facial burns. Outpatient pain control may be difficult in those with facial burns, the degree of edema may be difficult

---

**Figure 38.16** This badly burned hand requires referral to a surgeon or burn center and should not be definitively handled in the emergency department. Note the very tight ring (arrow).
to predict, and home care can be problematic. There are no universally agreed standards for admission versus outpatient treatment of facial burns.

Corneal contact burns, as from accidental contact with a curling iron, are often manifested rather dramatically as opacified, “heaped-up” corneal epithelium (see Fig. 38.18). Despite their appearance, the end result is usually excellent. Treatment is the same as for a corneal abrasion.59

Abuse of Children and Elderly Individuals

Recognition of the possibility of deliberate abuse by burning in the pediatric and geriatric populations is essential. In addition, children younger than 2 years have a thinner dermis and a less well-developed immune system than adults do. Elderly patients (>65 years) likewise tolerate burns poorly. These two populations are the most prone to abuse, often by family members (Fig. 38.19). For these reasons, both these groups of patients often require inpatient care.23
Figure 38.19 Burns can be a manifestation of child abuse, spouse abuse, or abuse of the elderly. 
A, Abuse burns from contact with a hot metal grate as a result of a child allegedly falling. B, This burn was the result of spouse abuse caused by throwing hot soup during an argument. Domestic abuse is often denied initially, but the delayed arrival at the hospital was a clue. C, Burns of the face and neck are common when a toddler pulls hot liquid from a stove. This case was never proved to be child abuse, but burns in young children are often due to abuse, especially if they are in atypical places. Although the body surface area of this burn is relatively small, the patient’s age and the burn’s location, coupled with the possibility of child abuse, require that this child be hospitalized. D, This infant received a severe blistering sunburn at the beach despite being in the shade most of the day. Reflection of sunlight from the sand and water can injure the delicate skin of an infant, who should have sunscreen applied. E, Self-inflicted cigarette burns in a psychiatric patient.
The majority of abused children are 18 to 36 months old, and for unknown reasons, the majority are male. Immersion burns are a common type of abuse and are characterized by circumferential, sharply demarcated burns on the hands, feet, buttocks, and perineum. Cigarette burns and burns from hot objects such as irons should be obvious. Contact burns on “nonexploring” parts of the child also warrant suspicion. A delay in seeking treatment may be a tip-off that a burn resulted from abuse. In older populations, the presence of confirmed self-inflicted burns such as cigarette burns suggests psychiatric disease (see Fig. 38.19E).

**Burns in Pregnancy**

There is little information in the literature concerning the special problems of pregnant burn victims. Ying-bei and Ying-jie reported on 24 pregnant burn patients representing a wide range of burn severity. Complications of the burn injuries included abortion and premature labor, although all patients in this series with burns covering less than 20% TBSA did well and delivered living full-term babies.

Because the resistance of pregnant women to infection is lower than that of nonpregnant women, control of burn wound infection is paramount. Gestational age appears to have no direct bearing on prognosis. Silver sulfadiazine cream should be avoided near term because of the potential for kernicterus.

**SPECIFIC BURNING AGENTS**

**Hot Tar Burns**

Asphalt is a product of the residues of coal tar and is commonly used in roofing and road repair. It is kept heated to approximately 450°F. When spilled onto the skin, the tar cools relatively rapidly, but the retained heat is sufficient to produce a partial-thickness burn. Fortunately, full-thickness burns are unusual. Cooled tar is nonirritating and does not promote infection. When cooled tar is physically removed, the adherent skin is usually avulsed (Fig. 38.20). Careless removal of the tar may inflict further damage on burned tissues. Agents such as alcohol, acetone, kerosene, or gasoline have been used to remove the tar, but these are flammable and may cause additional skin damage or a toxic response secondary to absorption.

There is no great need to meticulously remove all tar at the first visit. Obviously devitalized skin can be débrided, but adherent tar should be emulsified or dissolved rather than manually removed (Fig. 38.21). Polyoxyethylene sorbitan (Tween 80 [Sigma-Aldrich Corp., St. Louis, MO] or polysorbate 80) is the water-soluble, nontoxic, emulsifying agent found in Neosporin and several other topical antibiotic creams. Note that the cream formulations, not the ointments, contain the most useful tar dissolvers. The creams contain a complex mixture of ethers, esters, and sorbitol anhydrides that possess excellent hydrophilic and lyophilic characteristics when used as nonionic, surface-active emulsifying agents. With persistence, most tar can be removed (emulsified) on the initial visit, although this may be unnecessary as previously detailed.

Another household product (De-Solv-It [Orange-Sol Blending and Packaging, Inc., Chandler, AZ] multipurpose solvent) also appears logical for topical ED use. De-Solv-It has a surface-active moiety that wets the chemical’s surface and emulsifies tar and asphalt. Because De-Solv-It is itself a petroleum-based solvent, it should be applied only briefly, and the operator should wear gloves and protective eyewear during application. It should be used only for external exposure to tar or asphalt. After removal, the skin should be cleansed gently and dressed appropriately.

Many clinicians instead prefer to emulsify the majority of tar on an outpatient basis. A generous layer of polysorbate-based ointment can be applied under a bulky absorbent gauze dressing. The patient is then released home, and the residual tar is easily washed off after 24 to 36 hours (see Fig. 38.20B). Several dressing changes may be required. Once the residual tar is removed, the wound is treated as any other burn (see Fig. 38.21).

Shur-Clens (Convatec), a nontoxic, nonionic detergent, also works well for tar burn wound cleansing, as do mineral oil, petrolatum, and Medi-Sol (Orange-Sol), a petroleum-citrus
Immediate flushing with water is recommended for all chemical burns, with the exception of those caused by alkali metals. Flushing serves to cleanse the wound of unreacted surface chemical, dilute the chemical already in contact with tissue, and restore lost tissue water. Leonard and colleagues clearly demonstrated that patients receiving immediate copious water irrigation for chemical burns had less full-thickness burn injury and a 50% or greater reduction in hospital stay. Flushing should be thorough and may require at least 30 minutes (or as much as 2 hours) for maximal benefit, depending on the nature of the chemical.

Acid and Alkali Burns

Chemical burns cause progressive tissue damage until the chemical is inactivated or removed. Acids damage tissue by coagulation necrosis, a process that limits the depth of penetration into tissue. Alkalis react with lipids in skin and result in liquefactive necrosis. This process permits penetration of the chemical into tissues until neutralized. Thus, exposure to alkali is more likely to produce deep tissue wounds. Skin exposed to caustic substances should be decontaminated aggressively until neutralized and the resulting wounds considered deep until demonstrated otherwise.

Desiccant acids, such as sulfuric acid, create an exothermic reaction with tissue water and can cause both chemical and thermal injury. With extensive immersion injuries, acids may be absorbed systemically, thereby leading to systemic acidosis and coagulation abnormalities.

Chemical burns may be excruciatingly painful for long periods. Discomfort can be out of proportion to what one might expect from the perceived depth or extent of the burn. When caring for a chemical burn, the emergency care team should remove all potentially contaminated clothing. Any dry (anhydrous) chemical should first be brushed off the patient’s skin. The involved skin should then be irrigated with large amounts of water under low pressure. Any remaining particulate matter should be carefully débrided during irrigation.

<table>
<thead>
<tr>
<th>ACIDS</th>
<th>ALKALIS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Picric</td>
<td>Sodium hydroxide</td>
</tr>
<tr>
<td>Tungstic</td>
<td>Ammonium hydroxide</td>
</tr>
<tr>
<td>Sulfosalicylic</td>
<td>Lithium hydroxide</td>
</tr>
<tr>
<td>Tannic</td>
<td>Barium hydroxide</td>
</tr>
<tr>
<td>Trichloroacetic</td>
<td>Calcium hydroxide</td>
</tr>
<tr>
<td>Cresylic</td>
<td>Sodium hypochlorite</td>
</tr>
<tr>
<td>Acetic</td>
<td>Formic</td>
</tr>
<tr>
<td>Formic</td>
<td>Sulfuric</td>
</tr>
<tr>
<td>Hydrochloric</td>
<td>Hydrofluoric</td>
</tr>
<tr>
<td>Chromic</td>
<td>Chrome</td>
</tr>
</tbody>
</table>

Immediate flushing with water is recommended for all chemical burns, with the exception of those caused by alkali metals. Flushing serves to cleanse the wound of unreacted surface chemical, dilute the chemical already in contact with tissue, and restore lost tissue water. Leonard and colleagues clearly demonstrated that patients receiving immediate copious water irrigation for chemical burns had less full-thickness burn injury and a 50% or greater reduction in hospital stay. Flushing should be thorough and may require at least 30 minutes (or as much as 2 hours) for maximal benefit, depending on the nature of the chemical.

Acid and Alkali Burns

Chemical burns cause progressive tissue damage until the chemical is inactivated or removed. Acids damage tissue by coagulation necrosis, a process that limits the depth of penetration into tissue. Alkalis react with lipids in skin and result in liquefactive necrosis. This process permits penetration of the chemical into tissues until neutralized. Thus, exposure to alkali is more likely to produce deep tissue wounds. Skin exposed to caustic substances should be decontaminated aggressively until neutralized and the resulting wounds considered deep until demonstrated otherwise.

Desiccant acids, such as sulfuric acid, create an exothermic reaction with tissue water and can cause both chemical and thermal injury. With extensive immersion injuries, acids may be absorbed systemically, thereby leading to systemic acidosis and coagulation abnormalities.

Chemical burns may be excruciatingly painful for long periods. Discomfort can be out of proportion to what one might expect from the perceived depth or extent of the burn. When caring for a chemical burn, the emergency care team should remove all potentially contaminated clothing. Any dry (anhydrous) chemical should first be brushed off the patient’s skin. The involved skin should then be irrigated with large amounts of water under low pressure. Any remaining particulate matter should be carefully débrided during irrigation.
Strong alkali burns may require irrigation for 1 to 2 hours before tissue pH returns to normal. Some recommend that if the burn continues to feel “slippery” or tissue pH has not returned to normal after extensive irrigation, chemical neutralization may be helpful.\textsuperscript{64,65} Given that any heat of neutralization will be carried away with the irrigation solution,\textsuperscript{66} prompt irrigation with a dilute acid (e.g., vinegar, or 2% acetic acid) may hasten neutralization and patient comfort.

**Contact Burns From Wet Cement**

The major constituent of Portland cement, an alkaline substance, is calcium oxide (64%), combined with oxides of silicon, aluminum, magnesium, sulfur, iron, and potassium. There is considerable variability in the calcium oxide content of different grades of cement, with concrete having less and fine-textured masonry cement having more.\textsuperscript{63} The addition of water exothermically converts the calcium oxide to calcium hydroxide (Ca\(_2\)OH\(_2\)), a strongly corrosive alkali with a pH of 11 to 13. As the cement hardens, the calcium hydroxide reacts with ambient carbon dioxide and becomes inactive.

Both the heat and the Ca(OH)\(_2\) produced in this exothermic reaction can result in significant burns. Because of its low solubility and consequent low ionic strength, long exposure to Ca(OH)\(_2\) is required to produce injury. This usually occurs when workers spill concrete into their boots or kneel in it for a prolonged period. The burn wound and the resultant protein denaturation of tissues produce a thick, tenacious, ulcerated eschar. Concrete burns are insidious and progressive. What may appear initially as a patchy, superficial burn might in several days become a full-thickness injury requiring excision and skin grafting.\textsuperscript{67} The pain associated with these burns is often severe and more intense than the appearance of the wound might suggest (Fig. 38.22). Interestingly, many workers are not warned of the dangers of prolonged contact with cement, and because the initial contact with cement is usually painless, exposure may not be realized until the damage is done.

Treatment is as follows. Remove any loose particulate cement or lime, usually by brushing off, remove contaminated clothing, and irrigate the wound copiously with tap water (the pH of the effluent is tested and irrigation continued if the effluent is still alkaline). Apply compresses of dilute acetic acid (vinegar) to neutralize the remaining alkali and provide relief of pain after irrigation. Apply antibiotic ointment to the eschar during the early postburn period.

Sutilains ointment (Travase, Flint Pharmaceuticals, Deerfield, IL) is often recommended because it contains proteolytic enzymes that help speed eschar separation, but any common topical burn preparation is acceptable. The depth of burns from wet cement can be difficult to assess in the first several days. If it becomes apparent that the burns are full-thickness burns, early excision and skin grafting are recommended.

Cement burns should be differentiated from cement dermatitis, which is far more common. The latter is a contact sensitivity reaction, probably from the chromates present in cement. The contact dermatitis can initially be treated as a superficial partial-thickness burn.

**Air Bag Keratitis and Thermal Burns**

Safety legislation has mandated increased use of air bags to protect automobile occupants in the event of collision. Burns from air bags can be thermal, friction, or chemical (Fig. 38.23). The automobile air bag is a rubberized nylon bag that inflates on spark ignition of sodium azide to yield nitrogen gas, ash,
and a small amount of sodium hydroxide. Within seconds, the superheated air is vented, and this can produce a thermal burn if it contacts an extremity, the face, or the upper part of the torso. If the air bag ruptures, the alkaline contents of the bag are dispersed as a fine, black powder that usually causes no problems unless the eyes are exposed. Patients with eye exposure exhibit clinical evidence of a chemical keratoconjunctivitis, including photophobia, tearing, redness, and decreased visual acuity. Tear pH is usually elevated, and there may be particulate matter in the fornices.

The severity of an ocular alkaline burn is related to the duration of exposure and the concentration and pH of the chemical. For this reason, prompt, copious irrigation of the eyes with frequent assessment of tear pH is essential to prevent or minimize the injury (see Chapter 62). A rising pH suggests that trapped particulate matter is releasing additional chemical. Corneal edema and conjunctival blanching are signs of serious injury and can necessitate immediate ophthalmologic consultation.

**Hydrocarbon Burns**

Hydrocarbons are capable of causing severe contact injury by virtue of their irritant, fat-dissolving, and dehydrating properties. Cutaneous absorption may cause even more dangerous systemic effects. Gasoline, the usual agent involved, is a complex mixture of C₄ to C₁₁ alkane hydrocarbons and benzene; the hydrocarbons appear to be the major toxic agent. Lead poisoning caused by absorption through intact skin or burns from exposure to leaded gasoline have been reported previously but are currently quite rare because unleaded gasoline has virtually replaced the leaded version for most purposes.

The depth of injury is related to the duration of exposure and the concentration of the chemical agent. Gasoline immersion injuries resemble scald burns and are usually partial thickness. Occasionally, gasoline-injured skin exhibits a pinkish brown discoloration, possibly related to dye additives. A common source of gasoline exposure is motor vehicle collisions in which a comatose patient has been lying in a pool of gasoline.

The lungs are the usual site of systemic absorption and are often the only major route of excretion. The resultant high pulmonary concentrations may lead to pulmonary hemorrhage, atelectasis, and acute respiratory distress syndrome. To treat hydrocarbon burns, remove contaminated clothing, administer prolonged irrigation or soaking of the contaminated skin, débride significant burns caused by lead-containing gasoline (to reduce systemic lead absorption), and apply topical antibiotic ointments.

**Phenol Injury**

Phenol is a highly reactive aromatic acid alcohol that acts as a corrosive. Carbolic acid, an earlier term for phenol, was noted to have antiseptic properties and was used as such by Joseph Lister in performing the first antiseptic surgery. Hexylresorcinol, a phenol derivative, is in current use as a bactericidal agent. Phenols, in strong concentrations, cause considerable eschar formation, but skin absorption also occurs and can result in systemic effects such as central nervous system depression, hypotension, hemolysis, pulmonary edema, and death. Interestingly, phenol acts differently from other acids in that it penetrates deeper in a dilute solution than in a more concentrated form. Therefore, irrigation with water is suboptimal for phenol burns, but, because water is commonly and readily available, it is frequently used for irrigation.

Full-strength polyethylene glycol (PG 300 or 400) is more effective than water alone in removing phenolic compounds and should be obtained and used after water irrigation has begun. Polyethylene glycol is nontoxic and nonirritating and may be used anywhere on the body. When immediately available, polyethylene glycol can be used to remove the surface chemical before water irrigation (and chemical dilution) is begun.

**Hydrofluoric Acid Injury**

Hydrofluoric acid (HFA) is one of the strongest inorganic acids known; it has been widely used since its ability to dissolve silica was discovered in the late 17th century. Currently, HFA is used in masonry restoration, glass etching, and semiconductor manufacturing; for control of fermentation in breweries; and in the production of plastics and fluorocarbons. It is also used as a catalyst in petroleum alkylating units. It is available in industry as a liquid in varying concentrations up to 70%. It is also readily sold in home improvement and hardware stores. Significant concentrations of HFA are present in many home rust removal products, aluminum brighteners, automobile wheel cleaners, and heavy-duty cleaners in concentrations of less than 10%. Despite its ability to cause serious burns, unregulated and poorly labeled HFA products are recklessly used on a regular basis in the home and in small businesses. The public and many clinicians are generally unaware of the potential problems with this acid (Fig. 38.24).

Although HFA is quite corrosive, the hydrogen ion plays a relatively insignificant role in the pathophysiology of the burn injury. The accompanying fluoride ion is a protoplasmic poison that causes liquefaction necrosis and is notorious for its ability to penetrate tissues and cause delayed pain and deep tissue injury. This acid can penetrate through fingernails and cause nail bed injury. With home products, the unwary user does not realize that the substance is caustic until the skin (usually the hands and fingers) is exposed for a few minutes to hours, at which time the burning begins and becomes progressively worse. At this point the damage is done and the absorbed HFA cannot be washed off. With higher-strength industrial products, symptoms are almost immediate.

The initial corrosive burn is due to free hydrogen ions; secondary chemical burning is due to tissue penetration of fluoride ions. Fluoride is capable of binding cellular calcium, which results in cell death and liquefaction necrosis. The ionic shifts that result, particularly shifts of potassium, are believed to be responsible for the severe pain associated with HFA burns.

In high concentrations, the fluoride ions may penetrate to bone and produce deminerlization. Exposure of skin to concentrated HFA involving as little as 2.5% TBSA can lead to systemic hypocalcemia and death from intractable cardiac arrhythmias; it has been calculated that exposure to 7 mL of anhydrous HFA (HFA gas) is capable of binding all the free calcium in a 70-kg adult. Hyperkalemia and hypomagnesemia can also develop. If the hands are exposed, the acid characteristically penetrates the fingernails and injures the nail bed and cuticle area. As with most caustics, the pain is generally out of proportion to the apparent external physical injury. HFA burns produce variable areas of blanching and erythema, but blisters or skin sloughing are rarely seen initially. Skin necrosis and cutaneous hemorrhage may be noted in a few days.

Immediate treatment should begin with copious irrigation with water. Another approach is to wash the area with a solution.
of iced magnesium sulfate (Epsom salts) or a 1:500 solution of a quaternary ammonium compound such as benzalkonium chloride (Zephiran, Sanofi-Synthelabo Inc., New York, NY) or benzethonium chloride (Hyamine 1622, Sigma-Aldrich). Magnesium and calcium salts form an insoluble complex with fluoride ions, thus preventing further tissue diffusion. Though frequently recommended, topical preparations are often ineffective in limiting injury or controlling pain.

If there is no or only minimal visible evidence of skin injury and minimal pain, the burn may be dressed with topical calcium gluconate paste. This is not commercially available in the United States but is easily compounded in the pharmacy by mixing 3.5 to 7 g of pulverized calcium gluconate with 5 oz of a water-soluble lubricant such as K-Y jelly. This will form a thick paste with a calcium gluconate concentration of 2.5% to 5.0%. Some have suggested dimethyl sulfoxide as a vehicle to aid in skin penetration of the calcium. Plastic wrap (e.g., Saran Wrap, SC Johnson, Racine, WI) is used over a standard dry burn dressing to cover the calcium paste on the limbs. A vinyl or rubber glove filled with the paste can be used to cover the fingers and hands. Completely re-dress the wound and reapply the paste every 6 hours for the first 24 hours. As with most topical treatments of HFA burns, calcium gluconate is only minimally effective in relieving pain, and its value is probably overestimated in the literature.

A digital or regional nerve block with long-acting bupivacaine is an excellent way to provide prolonged pain relief if the hands are involved, but this does nothing to ameliorate the injury. In most cases, oral opioids are required. If bullae or vesicles have formed, they should be débrided to decrease the amount of fluoride present, and the wound should then be treated as any partial-thickness burn. Burns with HFA of less than 10% strength will heal spontaneously, usually without significant tissue loss, but pain and sensitivity of the fingertips may persist for 7 to 10 days. In addition, the fingernails may become loose.

The presence of significant skin injury or intense pain implies penetration of the skin by fluoride ions. This scenario is particularly common with exposure to HFA solutions in concentrations of 20% or greater, but tissue injury can occur with prolonged exposure to less concentrated products.

Initial treatment of a more concentrated exposure begins as described earlier and includes immediate débridement of necrotic tissue to remove as much fluoride ion as possible. After débridement, inject a 10% solution of calcium gluconate (note: avoid calcium chloride for tissue injections) intradermally and subcuticularly with a 30-gauge needle about the exposed area, using approximately 0.5 mL per square centimeter of burn. Pain relief should be almost immediate if this therapy is adequate. Because the degree of pain is a measure of the effectiveness of treatment, the use of anesthetics, especially by local infiltration, may be deleted if the burn is on the arm or leg. HFA can penetrate fingernails without damaging them. Soft tissue can be injected without prior anesthesia, but if the fingertips or nail beds are involved, they may be injected after a digital nerve block has been performed (Fig. 38.25). Before anesthesia and injection of calcium, the patient can outline the affected areas with a pen to ensure accurate injection of the antidote (see Fig. 38.25C). Although some investigators recommend that the fingernails be removed routinely, we strongly advise against this unless the nails are very loose or there is obvious necrosis of the nail bed. Fingers are best injected with a 25- or 27-gauge needle (a tuberculin syringe works well). Nails frequently become loose in a few days, but they often return to normal and do not require removal, particularly when lower-concentration nonindustrial products are involved.

Although infiltration of calcium gluconate is somewhat effective, the technique has certain limitations. Injections are painful, and the calcium gluconate solution itself causes a burning sensation. Because of the volume restrictions, not enough calcium may be delivered to bind all the free fluoride ions present. For example, 0.5 mL of 10% calcium gluconate contains 4.2 mg (0.235 mEq) of elemental calcium, which will neutralize only 0.025 mL of 20% HFA.

Several authorities have advocated intraarterial calcium infusions in the treatment of serious HFA burns of the extremities. Though very effective, this technique is not recommended for burns secondary to dilute HFA (i.e., concentrations <10%) because the morbidity is usually quite mild. When using this technique, 10 mL of 10% calcium gluconate is diluted in 50 mL of a 5% dextrose and water solution. The dilute solution is administered by slow infusion into an arterial catheter. It is unclear which artery best delivers the calcium...
to injured tissues. If only the radial three digits are involved, probably only the radial artery needs to be cannulated. Otherwise, a percutaneous catheter is inserted into the brachial artery. However, some investigators have advocated use of the radial artery in all cases, and because the arterial supply of the hand is interconnected, this may be a reasonable recommendation.79 The radial artery is usually more easily cannulated than the brachial artery. When arterial access has been accomplished, the solution is infused slowly over a 4-hour period. At this point the catheter is left in place and the patient is observed. If pain returns at any time over the next 4 hours, the infusion is repeated. If the patient is pain free over the 4-hour observation period, the burn is dressed and the patient is released home. This technique may be initiated in the ED, but many clinicians are reluctant to cannulate an artery and infuse calcium in the ED. Such patients require hospitalization or burn center referral for further evaluation and observation.

Advantages of the intraarterial method are elimination of the need for painful SQ injections and avoidance of the volume limitations of the SQ route while providing substantially more calcium to neutralize the fluoride. Disadvantages of intraarterial calcium therapy include the possibility of local arterial spasm (which can be treated with vasodilators such as phentolamine or removal of the catheter), local arterial injury or thrombus, and the long duration of treatment required.

Infusing calcium into the general venous circulation is of no benefit for HFA burns. Some authors have advocated the use of regional IV calcium gluconate, similar to the method used with the Bier block for regional anesthesia (see Chapter 32).30 Case reports have noted variable success, but this technique...
has neither been well studied nor rigorously compared with other options. This method would be useful only for upper extremity burns. To perform regional calcium therapy, place an IV catheter in the dorsum of the hand on the involved extremity. Partially exsanguinate the arm by elevation, wrapping with an elastic bandage, or both. Apply a Bier block tourniquet or a heavy-duty blood pressure cuff to the burn and inflate it to 20 to 30 mm Hg above the systolic pressure to stop blood flow to and from the arm. Because slow deflation of a regular blood pressure cuff may thwart success of the procedure, using this specialized tourniquet is recommended. Then, dilute 10 mL of 10% calcium gluconate with 30 to 40 mL of saline, and infuse the solution into the venous catheter. Keep the solution in the arm by keeping the tourniquet on for 20 to 30 minutes. Some patients cannot tolerate arm ischemia for this period, thus limiting the effectiveness of this procedure. Theoretically, the calcium diffuses out of the venous system and into the injured tissues. After 20 to 30 minutes, deflate the cuff to restore normal circulation to the extremity. It may require 10 to 20 minutes after deflation of the tourniquet before the patient experiences relief of pain. This procedure is safe, but its efficacy is variable.

HFA burns involving the eye are potentially devastating injuries that deserve special mention. Ophthalmologist referral is mandatory. Ocular exposure to liquid or gaseous HFA will result in severe pain, tearing, conjunctival inflammation, and cornal opacification or erosion. Complications include decreased visual acuity, globe perforation, uveitis, glaucoma, conjunctival scarring, lid deformities, and keratitis sicca. Optimal therapy for ocular HFA burns, other than initial irrigation, is unknown. Irrigation may be performed with water, isotonic saline, or magnesium chloride. We advise copious saline irrigation. Topical antibiotics and cycloplegics, along with light pressure patching, are also recommended. The use of topical steroids has been advocated by some to lessen corneal fibroblast pressure patching, are also recommended. The use of topical antibiotics and cycloplegics, along with light

**Chromic Acid Injury**

Chromic acid compounds are used extensively in industry, mainly in metallic electroplating. Chromic acid is commonly used in concentrated solutions containing up to 25% sulfuric acid. It causes sufficient skin damage to allow absorption of the toxic chromium ion if intensive irrigation is not undertaken immediately. Heated (60°C to 80°C) chromic acid makes the problem of chromium absorption much worse.

Dichromate salts containing hexavalent chromium are the most readily absorbed and the most toxic because they can cross cell membranes. The mortality rate from these burns is very high if the burn exceeds 10% TBSA. Chromium absorption leads to diarrhea, gastrointestinal bleeding, hemolysis, hepatic and renal damage, coma, encephalopathy, seizures, and disseminated intravascular coagulation.

To treat, immediately excise the burned tissues to lessen the total body dichromate burden. Wash wounds with a 1% sodium phosphate or sulfate solution and dress them with bandages soaked in 5% sodium thiosulfate solution. These actions reduce the hexavalent chromium ion to the less well absorbed trivalent form.

Institute chelation therapy with ethylenediaminetetraacetic acid (EDTA) and give IV sodium thiosulfate and ascorbic acid. Hemodialysis, peritoneal dialysis, or exchange transfusion may be indicated.

**Phosphorus Burns**

White phosphorus is a translucent, waxy substance that ignites spontaneously on contact with air. For this reason, it is usually stored under water. It is used primarily in fireworks, insecticides, rodenticides, and military weapons.

Phosphorus causes both thermal burns from the flaming pieces and acid burns from the oxidation of phosphorus to phosphoric acid. The burns classically emit a white vapor with a characteristic garlic odor.

Treat these burns first by immersion in water. Débride any gross debris. Wash the wound with a 1% copper sulfate solution, which reacts with the residual phosphorus to form copper phosphate. Copper phosphate appears as black granules, which allow for easy débridement. After débridement, remove the residual copper with a thorough water rinse, dress the wound, and treat it as any other burn.

**Elemental Alkali Metal Burns**

The commonly encountered alkali metals (sodium, lithium, and potassium) are highly reactive with water and with water vapor in the air and produce their respective hydroxides with liberation of hydrogen gas. Therefore, water should never be used for extinguishing or débriding the metal. A class D fire extinguisher or plain sand may be used to smother the fire, followed by the application of mineral oil or cooking oil to isolate the metal from water and allow safe débridement. Then, treat the burn as an alkali burn.

Magnesium burns in a less intense fashion but otherwise acts as other alkali metals do. These burns may be particularly injurious, however, because if all the metallic debris is not removed, the small ulcers that form will slowly enlarge until they become quite extensive.

The initial topical treatment of unusual chemical burns is outlined in Table 38.3.

**Electrical Burns**

Electrical burn wounds occur when energy traveling through the body across a potential difference is converted to heat. This energy has the ability to destroy deep tissues, including muscles, tendons, and nerves (Fig. 38.26). In addition, electrical injuries can arise from the arc produced when electricity passes through the air and from flames caused by the ignition of clothing. Electrical injuries from high-voltage or high-current sources (> 1000 V and > 5000 mA) are more likely to result in deep soft tissue damage, whereas low voltage or low current (< 1000 V and 5 to 60 mA) causes less soft tissue damage but is more likely to result in cardiac arrhythmias.

**TEN and SJS**

Toxic epidermal necrolysis (TEN) and Stevens-Johnson syndrome (SJS) are severe blistering diseases. They are primarily associated with the intake of medications that cause apoptosis of keratinocytes, which results in the separation of large areas of skin at the dermal-epidermal junction and produces the appearance of a scald. More than 200 medications have been associated with this condition, although infections and immunizations have also been associated. A major factor in improving outcomes has been high-quality intensive support and trained
incidence of 0.4 to 1.2 and 1.2 to 6.0 per million persons, respectively, and are more common in females and the elderly. Patients at risk are those who are severely immunocompromised (e.g., human immunodeficiency virus [HIV] infection, lymphoma). Death occurs on average in every third patient with TEN. More than 200 drugs, including antibiotics (particularly sulfonamides), nonsteroidal antiinflammatory drugs, and anticonvulsants, have been implicated.

Various theories exist to explain the precise sequence of molecular and cellular events responsible for the development of TEN. The 1- to 3-week interval between the onset of TEN and the commencement of drug therapy favors an immune etiology. Cytotoxic T cells are seen in cutaneous lesions, and it is hypothesized that necrolysis is due to their recognition of complexes between drug metabolites and major histocompatibility complex class I molecules on the surface of keratinocytes. Exfoliation is due to the death of keratinocytes via apoptosis, and data suggest that the latter is mediated by interaction of the death receptors, transmembrane proteins, Fas, and its ligand FasL. This activates the proteolytic cascade (caspases), which leads to cellular disintegration. Evidence has shown upregulation of FasL in patients with TEN.

**Clinical Features**

TEN and SJS are usually characterized by fever, corneal irritation, and painful swelling (representing oral mucocutaneous involvement). These symptoms can precede the rash by 1 to 3 days. In more than 99% of patients, erythema and erosions of the buccal, ocular, and genital mucosa develop and are painful. Diagnosis may require skin biopsy as the condition might be difficult to clinically differentiate from other conditions such as staphylococcal scalded skin syndrome or other dermatologic emergencies. The epithelium of the respiratory tract is involved in 25% of cases of TEN, and gastrointestinal lesions can occur. The skin lesions first appear as erythematous, dusky red, or purpuric macules, irregular in size and shape, that tend to coalesce. Nikolsky’s sign (blistering following pressure with the finger) may be evident. A gray appearance of the macule heralds necrosis of the epidermis, which soon separates from the dermis and leaves a raw painful area. Wound infections

---

**TABLE 38.3 Chemical Burn Treatment**

<table>
<thead>
<tr>
<th>WATER LAVAGE</th>
<th>CHROMIC ACID</th>
<th>TANNIC ACID</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potassium permanganate</td>
<td>Tannic acid</td>
<td>Sulfsalicylic acid</td>
</tr>
<tr>
<td>Cantharides</td>
<td></td>
<td>Trichloroacetic acid</td>
</tr>
<tr>
<td>Lyes (hydroxide salts)</td>
<td></td>
<td>Cresylic acid</td>
</tr>
<tr>
<td>Clorox</td>
<td></td>
<td>Acetic acid</td>
</tr>
<tr>
<td>Dichromate salts</td>
<td></td>
<td>Formic acid</td>
</tr>
<tr>
<td>Picric acid</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calcium salt injection</td>
<td>Oxalic acid</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hydrofluoric acid</td>
<td></td>
</tr>
<tr>
<td>Oil immersion</td>
<td>Sodium metal</td>
<td></td>
</tr>
<tr>
<td></td>
<td>White phosphorus</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mustard gas</td>
<td></td>
</tr>
<tr>
<td>Avoid water lavage</td>
<td>Sodium metal</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Potassium metal</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lithium metal</td>
<td></td>
</tr>
<tr>
<td>Specific approaches</td>
<td>Sodium metal</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lyes (hydroxide salts)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hydrofluoric acid</td>
<td></td>
</tr>
<tr>
<td></td>
<td>White phosphorus</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Excision</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Weak acid lavage (vinegar)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Calcium gluconate injection</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Copper sulfate solution</td>
<td></td>
</tr>
</tbody>
</table>

---

*Figure 38.26* Electrical burn. The patient experienced a contact burn across the dorsa of the toes from an exposed electrical wire. (From Davis PJ, Cladis FP, Motoyama EK, editors: *Smith’s anesthesia for infants and children*, ed 8, St Louis, 2011, Mosby.)

*Figure 38.27* Stevens-Johnson syndrome (SJS). Purpuric macules became bullous. Note the inflammation of the conjunctivea and lips. By definition, in SJS the lesions occupy less than 10% of total body surface area (TBSA). (From Paller AS, Mancini AJ, editors: *Hurwitz clinical pediatric dermatology*, ed 4, St. Louis, 2011, Saunders.)

---

nursing care with expertise in wounds. Thus these disorders are ideally suited to treatment at burn centers.

The distinction between TEN and SJS is one of extent, with lesions occupying less than 10% TBSA qualifying as SJS (Fig. 38.27) and lesions involving greater than 30% TBSA being called TEN (Fig. 38.28); when the extent of involvement lies between 10% and 30%, an intermediary term is coined, SJS-TEN overlap. These disorders are rare with an annual
against Fas that can block the binding of Fas with FasL, but recently granulysin has been identified as the most important of known mediators for SJS & TEN. Importantly, subsequent research has not demonstrated a significant mortality advantage for patients using either high- or low-dose IVIG compared with patients treated with supportive care only, so IVIG is not currently established as standard of care therapy.

Mortality of TEN approaches 30% on average, depending on stage of diagnosis at presentation, supportive measures, age, and comorbidities. A modified scoring system for patients (S. aureus and P. aeruginosa) and fluid and electrolyte loss often follow. Death is mainly due to sepsis, acute respiratory distress syndrome, and multiorgan failure. Reepithelialization and healing of wounds can occur within 3 weeks but may be prolonged. There are, however, late sequelae such as scarring and pigmentary abnormalities, as well as serious ophthalmic complications (e.g., symblepharon, conjunctival synechiae, entropion), which can range from sicca syndrome to blindness.

**Management**

Initial management of TEN requires immediate discontinuation of all medications, including antibiotics, if there are no signs of infection. Supportive care is similar to that required in the treatment of thermal burns. Depending on the body surface area involved patients may require the resources of a Burn Center that has dermatology consultants available. Medical attention is paid to correction of fluid and electrolyte abnormalities, renal insufficiency, nutrition, and sepsis. Involvement of respiratory mucosa may warrant intubation and ventilation. The wounds are carefully débrided and a biopsy specimen is taken to confirm the diagnosis. Regular hydrotherapy and topical antimicrobials are used to decrease infection. Silvadene and mafenide are best avoided because they may be implicated in causing the disorder, but alternative dressings include the various silver products (e.g., Acticoat, Smith & Nephew), as well as bacitracin and Xeroform (DeRoyal). Synthetic dressings such as Biobrane (Smith & Nephew) and biologic materials, including allograft, have all been used with various success.

Regular examination by an ophthalmologist is also recommended. The eyelids should be cleansed daily and followed with a daily application of antibiotic ointment. Attention to oral hygiene is also mandatory because oral lesions are common. Several case reports and uncontrolled series suggest the utility of specific therapies for the treatment of TEN/SJS; however, thus far there are no strong evidence-based standards to promote any particular therapy. Studies have included cyclosporine, cyclophosphamide, IVIG, plasmapheresis, and N-acetylcysteine, but these therapies are not established as standard of care. The use of systemic corticosteroids remains controversial, and they may even increase mortality. Promising results were shown with the use of IV immunoglobulins that contain antibodies against Fas that can block the binding of Fas with FasL, but recently granulysin has been identified as the most important of known mediators for SJS & TEN. Importantly, subsequent research has not demonstrated a significant mortality advantage for patients using either high- or low-dose IVIG compared with patients treated with supportive care only, so IVIG is not currently established as standard of care therapy.

Mortality of TEN approaches 30% on average, depending on stage of diagnosis at presentation, supportive measures, age, and comorbidities. A modified scoring system for patients

---

**TABLE 38.4 Scorten Score for Predicted Mortality in TEN**

<table>
<thead>
<tr>
<th>RISK FACTOR</th>
<th>0</th>
<th>1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>&lt;40 years</td>
<td>≥40 years</td>
</tr>
<tr>
<td>Malignancy</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Heart rate</td>
<td>&lt;120</td>
<td>≥120</td>
</tr>
<tr>
<td>Serum BUN (mg/dL)</td>
<td>&lt;27</td>
<td>&gt;27</td>
</tr>
<tr>
<td>%BSA involved</td>
<td>&lt;10%</td>
<td>&gt;10%</td>
</tr>
<tr>
<td>Serum bicarbonate (mEq/L)</td>
<td>≤20</td>
<td>&lt;20</td>
</tr>
<tr>
<td>Serum glucose (mg/dL)</td>
<td>&lt;250</td>
<td>&gt;250</td>
</tr>
</tbody>
</table>

**NUMBER OF RISK FACTORS** | **MORTALITY RATE**
--- | ---
0–1 | 3.2% |
2 | 12.1% |
3 | 35.3% |
4 | 58.3% |
5 or more | 90% |

exposure of more than 100 rad causes acute radiation syndrome within hours of exposure. This is characterized acutely by nausea, vomiting, diarrhea, fever, fatigue, and headache. The symptoms may resolve transiently during a latent period only to recur as hematopoietic, gastrointestinal, or vascular complications.93,94

**EMERGENCY ESCHAROTOMY**

Full-thickness burns result in an eschar that is inelastic and may become restrictive and result in compartment syndrome. Intracutaneous and interstitial edema can develop and progress, both because of fluid resuscitation and as a direct result of transcapillary extravasation of fluid from the thermal injury. As the soft tissues become edematous and pressure rises under the unyielding eschar, first venous and then lymphatic, capillary, and ultimately arterial flow to the underlying and distal unburned tissue may be compromised. Full-thickness and extensive partial-thickness circumferential extremity burns are most likely to impede peripheral blood flow. Circumferential chest burns may restrict chest wall movement and impair ventilation, and circumferential neck burns may result in tracheal obstruction. In such cases, immediate escharotomy may be indicated. On occasion, because of high-volume fluid resuscitation, noncircumferential and deep partial-thickness burns require surgical decompression to prevent the complications of nerve or muscle damage. Once signs and symptoms of vascular impairment are present, the clinician must act quickly to prevent tissue hypoxia and cellular death. This pathophysiology may be manifested within a time frame that requires an emergency clinician to intervene. Clinical assessment of tight compartments may be aided by measurements such as capillary refill, Doppler signals, pulse oximetry, and direct measurement of compartment pressures. Escharotomy, when required, is usually performed within the first 2 to 6 hours of a burn injury. The need for non–burn specialists to identify the need for and perform an adequate escharotomy is illustrated by the report of Brown and associates,95 who found that 44% of pediatric burn cases were inadequately decompressed before arrival at a referral burn unit.

**Frostbite**

Frostbite is the result of exposure to low environmental temperatures (Fig. 38.29). The formation of ice crystals within extracellular fluid causes direct cellular injury and cellular dehydrothrombosis through transmembrane osmotic shifts. In addition, a secondary vascular effect of cooling leads to endothelial damage and progressive dermal ischemia.92

Initial management of acute frostbite should entail determination of the core temperature and a full physical examination. The frostbitten part should be rapidly rewarmed in a water bath (temperature of 40°C to 42°C) with adequate analgesia for 15 to 30 minutes. Treatment of deep injury consists of elevation of the injured part to control edema, adequate analgesia, splinting, and the application of topical antibiotics. Traditionally, white blisters are débrided. They generally represent superficial injury, and débridement is thought to be beneficial because it helps in the removal of thromboxane A2 and prostaglandin F2α. Hemorrhagic blisters, however, are said to represent deeper injury and are best left intact to protect tissues against desiccation. The use of antithromboxane drugs such as aspirin or ibuprofen has been shown to be useful, as has the application of aloe vera. The avoidance of refreezing is critical to reduce complications. Patients should be educated on this important precaution. Premature amputation should be avoided but may be necessary for definitive closure. Long-term consequences include a predilection for future frostbite, vasospastic syndromes, and cold hypersensitivity.

**Radiation Burns**

Accidents involving ionizing radiation are not common, but when they occur, the clinical findings may range from erythema to charring of the superficial layers of skin. Whole-body exposure of more than 100 rad causes acute radiation syndrome. With TEN has been published by Bastuji-Garin and colleagues that predicts the mortality of these patients91 (Table 38.4). Although rarely useful for the emergency provider, this scoring system and predicted mortality may aid in the collaborative decision to hospitalize at a non-burn center or arrange for transfer to a burn center.

**Figure 38.29 Frostbite.** Initial management includes rapid rewarming in a water bath (temperature of 40°C to 42°C) for 15 to 30 minutes. Hemorrhagic blisters should be left intact; white or clear blisters may be debrided. Avoidance of refreezing is imperative.
It is not standard of care that emergency clinicians be skilled in emergency escharotomy, nor can it be expected that this procedure will be done in the ED. The technique is described here for circumstances when escharotomy must be performed by a non–burn specialist.

**Indications**

The indications for escharotomy are based on clinical examination, compartment pressure, or both. A high index of suspicion and a low threshold for intervention are essential for a successful outcome. Skin temperature and palpation of pulses are unreliable and imprecise indicators of the adequacy of circulation because of peripheral vasoconstriction and local edema. A patient with circulatory embarrassment significant enough to warrant escharotomy may complain of deep aching pain, progressive loss of sensation, or paresthesias, but these parameters are difficult to quantify in a severely burned, sedated, or mechanically ventilated patient. However, motor activity and peripheral pulses may remain intact despite severe underlying muscle ischemia. In the series by Brown and associates,95 peripheral pulses were present in 74% of the limbs that required decompression. Muscle compartments with pressures in excess of 30 mm Hg should be decompressed. Measurements should be taken before and after escharotomy to ensure adequate decompression.

In a patient with absent distal arterial flow (as determined with a Doppler ultrasonic flow meter) but otherwise adequate blood pressure, immediate escharotomy is indicated. Bardakjian and coworkers96 suggested that an oxygen saturation below 95% in the distal end of the extremity as demonstrated by pulse oximetry (in the absence of systemic hypoxia) is also a reliable indicator of the need for emergency escharotomy.

**Technique of Escharotomy**

Because full-thickness burns are insensate to pain and involve coagulation of superficial vessels, no anesthesia is needed. Patients with deep partial-thickness burns may still possess pain sensation, and escharotomy may be performed with local anesthesia or systemic analgesia. A properly executed escharotomy releases the eschar to the depth of SQ fat only. This results in minimal bleeding, which can be controlled with local pressure or electrocautery. These incisions, even though life or limb saving, represent potential sources of infection for the burn patient and should be treated as part of the burn wound. The wounds should be loosely packed with sterile gauze impregnated with an appropriate topical antimicrobial such as silver sulfadiazine cream. Fasciotomy, which involves a deeper incision, may be needed for thermal or electrical burns.

**Limbs**

Under sterile conditions, incise the lateral and medial aspects of the involved extremity with a scalpel or electrocautery 1 cm proximal to the burned area and 1 cm distal to the involved area of constricting burn (Fig. 38.30). Carry the incision through the full thickness of skin only and this should result in immediate separation of the constricting eschar to expose SQ fat. Because joints are areas of tight skin adherence and potential vascular impingement, incisions should cross these structures (Fig. 38.31). Take care to avoid vital structures, such as the ulnar nerve at the elbow, the radial nerve at the wrist, the superficial peroneal nerve near the fibular head, and the posterior tibial
artery at the ankle. In circumferential burns of the feet, if escharotomy is indicated, extend the incision to the great toe medially and the little toe laterally. In circumferential burns of the hands in which escharotomy is indicated, extend the incisions to the thenar and hypothenar aspects of the hands (see Figs. 38.30 and 38.31). Softening of the compartment, improved distal tissue perfusion, return of sensation, Doppler flow signal strength, and oximetry values indicate adequate release.96

**Chest**

Full-thickness circumferential chest or upper abdominal burns may impair respiration. Nearly all these patients are expected to be intubated and mechanically ventilated. Evidence of the need to release the eschar is increased airway pressure or an inability to ventilate. Escharotomy of the chest wall should extend from the clavicle to the costal margin in the anterior axillary line bilaterally, while avoiding breast tissue in females. This may be joined by transverse incisions to result in a chevron-shaped subcostal incision (see Fig. 38.30).

**Neck**

Neck escharotomy should be performed laterally and posteriorly to avoid the carotid and jugular vessels.

**Penis**

Penile escharotomy is performed midlaterally to avoid the dorsal vein.

**Complications**

Complications of escharotomy include bleeding, infection, and damage to underlying structures. Complications of inadequate decompression include muscle necrosis, nerve injury (such as footdrop), and even amputation of the limb. Systemic complications of inadequate decompression include myoglobinuria and renal failure, hyperkalemia, and metabolic acidosis.

**CONCLUSION**

Patients with circumferential or nearly circumferential burns should be evaluated for the risk of developing compartment syndrome and deep tissue ischemia. Emergency clinicians should not hesitate to perform an escharotomy before transfer of the patient to a burn center if there is evidence of reduced limb perfusion or impaired ventilation.

**ACKNOWLEDGMENTS**

The significant contributions of Courtney A. Bethel, MD, Leigh Ann Price, MD, and Kevin B. Gerold, MD, to earlier editions remain appreciated.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


805.e2  SECTION VI  SOFT TISSUE PROCEDURES


89. Deleted in review.


Patients with foreign bodies (FBs) lodged in the esophagus commonly present to the emergency department (ED) for evaluation and treatment. Though most commonly accidental, FBs may sometimes be swallowed purposefully. Patients may have a sensation of a recently passed FB, minor irritation, life-threatening airway obstruction, or other significant complications. Because of the anatomic and physiologic features of the esophagus, FBs in this area of the gastrointestinal (GI) tract present unique clinical issues to the clinician.

GENERAL FEATURES

Anatomy

The esophagus is a muscular tube 20 to 25 cm in length. There are three anatomic areas of narrowing in which FBs are most commonly entrapped: in the upper esophageal sphincter, which consists of the cricopharyngeus muscle; in the midesophagus at the crossover of the aortic arch; and in the lower esophageal sphincter (LES) (Fig. 39.1). The LES is the narrowest point of the esophagus and the entire GI tract.

Epidemiology

There are almost 100,000 cases of FB ingestions reported annually in the United States, and they are the eighth most common cause of calls to poison control centers. Patients with retained esophageal FBs generally fall into one of the following categories: pediatric patients, psychiatric patients, prisoners, and adults who either are edentulous or have underlying esophageal pathology. Children account for 75% to 85% of esophageal FBs seen in the ED, with the peak incidence occurring at the age of 18 to 48 months. The incidence is equal in boys and girls. Inquisitive children frequently place objects in their mouth and unintentionally swallow them. As a result, children most commonly ingest coins, but they also swallow buttons, marbles, beads, screws, and pins. Unlike adults, children who have entrapped, accidentally swallowed FBs do not normally have underlying esophageal disorders. However, this is not the case in children with esophageal meat or food impaction, and these patients will need further evaluation for underlying esophageal disease, the most common of which is eosinophilic esophagitis (55%), or stricture (27%).

Patients with an anatomic abnormality of the esophagus or a motor disturbance are more prone to FB entrapment. Anatomic abnormalities include strictures, webs, rings, diverticula, and malignancies. Motor disturbances include achalasia, scleroderma, and esophageal spasm. Adults who have dentures or underlying esophageal anatomic or motor abnormalities may accidentally ingest food boluses, chicken bones, fish bones, glass, toothpicks, fruit pits, or pills while in the act of eating. Prisoners and psychiatric patients ingest a wide variety of objects, some of which may be quite unusual: spoons, razor blades, pins, nails, or practically any other object.

Complications

Impacted FBs of the esophagus must be removed or dislodged. The time frame under which this mandate must be carried out varies widely and depends on many circumstances. In general, however, the esophagus does not tolerate FBs well or for prolonged periods because it is prone to pressure, edema, necrosis, infection, and eventually perforation. FBs can transit the esophagus in a matter of seconds or minutes or may adhere to the mucosa. Retained objects may become less symptomatic after time, and the clinician must resist the urge to allow esophageal FBs to “pass by themselves” or “dissolve.” Once FBs become stuck in the mucosa, they may become less symptomatic, but they rarely pass on their own. The one exception may be children with coins or other smooth round FBs such as small marbles, especially those lodged at the LES. Approximately one third of these objects may pass spontaneously within 24 hours, and some authorities have advocated an observational approach, although this tends to be more poorly accepted by parents. Button batteries must be immediately removed because of the risk of rapidly occurring esophageal erosion and perforation. The literature is replete every year with a wide array of case reports and small reports of series of complications that can arise from retained esophageal FBs (Box 39.1), including benign mucosal abrasions, lacerations, esophageal stricture, and necrosis from corrosive agents such as button batteries. Esophageal perforation can lead to life-threatening conditions such as retropharyngeal abscess, mediastinitis, pericarditis, pericardial tamponade, pneumothorax, pneumomediastinum,
Dysphagia is a common initial complaint with esophageal FBs. Drooling is suggestive of high-grade obstruction, and complete inability to handle oral secretions is a sign of total obstruction. Infants with a clandestine esophageal FB can exhibit wheezing or a chronic cough. They may appear to have bronchospasm and may be treated for asthma by a number of clinicians before an FB is suspected. Stridor from an FB can mimic epiglottitis.

The esophagus is well innervated proximally, and patients can typically accurately localize FBs in the oropharynx or upper third of the esophagus. However, scratches or abrasions of the esophagus can create a persistent FB sensation. Upper esophageal FBs often cause gagging or vomiting. In rare cases, an upper esophageal FB can impinge on the trachea, especially in children, and mimic infection by inducing wheezing, stridor, or frank respiratory distress. The lower two thirds of the esophagus is not as well innervated, and FBs in this location typically cause vague symptoms of discomfort, fullness, or nonlocalizing pain. Swallowed coins that lodge in the lower part of the esophagus in children may cause no overt symptoms until feeding is attempted.

The location of retained esophageal FBs is related to age and areas of physiologic narrowing of the esophagus (Table 39.1). Children more typically have objects entrapped at the cricopharyngeus muscle, whereas adults more commonly have entrapment at the LES.

**Clinical Findings**

Esophageal FB impaction is usually an acute condition, particularly in adults who have a clear history of ingestion. Children also commonly remember an ingestion, but some will have a vague history or symptoms. As many as one third of children with proven esophageal FBs are asymptomatic on initial evaluation, therefore a high index of suspicion is indicated, especially in children who were seen with an object in their mouth that subsequently disappeared. This is particularly true if transient coughing or gagging occurred, even though the actual ingestion was not witnessed. Poor feeding, irritability, fever, stridor, cough, wheezing, and aspiration can all be caused by an underlying esophageal FB in a child, especially a young infant.

Dysphagia is a common initial complaint with esophageal FBs. Drooling is suggestive of high-grade obstruction, and complete inability to handle oral secretions is a sign of total obstruction. Infants with a clandestine esophageal FB can exhibit wheezing or a chronic cough. They may appear to have bronchospasm and may be treated for asthma by a number of clinicians before an FB is suspected. Stridor from an FB can mimic epiglottitis.

The esophagus is well innervated proximally, and patients can typically accurately localize FBs in the oropharynx or upper third of the esophagus. However, scratches or abrasions of the esophagus can create a persistent FB sensation. Upper esophageal FBs often cause gagging or vomiting. In rare cases, an upper esophageal FB can impinge on the trachea, especially in children, and mimic infection by inducing wheezing, stridor, or frank respiratory distress. The lower two thirds of the esophagus is not as well innervated, and FBs in this location typically cause vague symptoms of discomfort, fullness, or nonlocalizing pain. Swallowed coins that lodge in the lower part of the esophagus in children may cause no overt symptoms until feeding is attempted.

The location of retained esophageal FBs is related to age and areas of physiologic narrowing of the esophagus (Table 39.1). Children more typically have objects entrapped in the upper part of the esophagus at the level of the cricopharyngeus muscle, whereas adults more commonly have entrapment at the LES.

**Evaluation**

The most useful aspect of the evaluation is the history. The time of the ingestion, size and shape of the ingested object, and any current symptoms should be ascertained. Findings on physical examination are frequently normal in patients with esophageal FBs, unless complete obstruction is present. In this case they will be drooling, spitting, and unable to handle oral secretions. Even though a patient may be asymptomatic at initial encounter, transient coughing or gagging should raise the index of suspicion for an esophageal FB. Examination of the oropharynx, neck, respiratory system, cardiac system, and abdomen is essential in the evaluation of potential complications.

After attending to life-threatening conditions such as airway compromise, the goal of ED evaluation is to localize the FB to determine what, if any, interventions need to be undertaken to remove it or assist its transit into the stomach. Once an FB passes into the stomach, it has a greater than 90% likelihood of passing through the entire GI tract without any further problems. Even irregular, and seemingly dangerous FBs will often transit the entire GI tract with relative ease.
RADIOLOGY OF ESOPHAGEAL FOREIGN BODIES

Background
Radiographic imaging of a patient with a suspected esophageal FB is a common practice and is particularly useful for detecting radiopaque FBs. Traditionally, an inability to quickly identify the object by physical examination encouraged the use of plain radiography in an attempt to verify and localize the retained FB. However, the limitations of plain radiography require that other diagnostic approaches also be considered.

Indications
Essentially every patient with a suspected esophageal FB warrants radiographic evaluation with some exceptions, including those patients who have an obvious complete obstruction with an inability to manage their secretions. These patients need endoscopy to relieve the obstruction. If there is any question of aspiration, a chest x-ray may be indicated. Patients with a history of sharp, nonradiopaque FBs such as toothpicks will also need urgent endoscopy. If there has been a delay in presentation in the case of a sharp FB and there are any signs of infection such as fever or chest pain, computed tomography (CT) may be indicated to evaluate for perforation or mediastinitis. Finally, those patients with an FB sensation that is visualized in the upper pharynx on exam and removed may forego radiographs. In nonverbal patients, including preschool children and those who are demented or debilitated, maintain a low threshold for screening radiography in cases with a suspicious history. Examples include a child seen with an object in the mouth that “disappeared” or a patient with symptomatology suggestive of an esophageal FB, such as drooling, gagging, or unexplained respiratory symptoms.

Plain Radiographs
Plain radiographs reliably verify and localize radiopaque FBs such as glass and metal of sufficient size, and are indicated as the main method of radiologic evaluation for these objects.

When used, a complete oropharyngeal radiographic series includes the nasopharynx to the lower cervical vertebra in both lateral and anteroposterior views. Optimum-quality radiographs are mandatory. Patients should be positioned upright with the neck extended and the shoulders held low. Use of a soft tissue technique enhances the discrimination of weak radiopaque FBs. Phonation of “eeeee” during radiography prevents motion artifact from swallowing, distends the hypopharynx, and enhances soft tissue landmarks. As previously mentioned, FBs are most frequently entrapped at one of three locations of physiologic narrowing in the esophagus: the cricopharyngeus muscle (Fig. 39.2), the aortic crossover (Fig. 39.3), and the LES (Fig. 39.4). Objects that become lodged in the middle portion of the esophagus most likely represent esophageal pathology, such as stricture related to tumor or eosinophilic esophagitis.

Posteroanterior (PA) and lateral views of the chest are used to evaluate the remainder of the esophagus. Both projections are indicated to identify multiple objects and FBs visible in only one plane. Esophageal FBs typically lie in the vertical plane and are differentiated from airway bodies or calcifications by their location posterior to the tracheal air column on lateral radiographs. As a rule, flat objects such as coins perch in the

Figure 39.2 Posteroanterior radiograph of an esophageal foreign body (coin) lodged at the level of the cricopharyngeus muscle. This is the most common area of the esophagus to harbor a coin in children. Coins remaining in the upper tract are usually removed unless there is a steady progression with observation. This coin would probably be symptomatic in an infant and cause respiratory distress, drooling, wheezing, and possibly stridor. Note: The chance of spontaneous passage is approximately 20% to 25%.

Figure 39.3 Posteroanterior radiograph of an esophageal foreign body (coin) lodged at the level of the aortic crossover. Note: The chance of spontaneous passage is approximately 20% to 25%.
varies with the fish species and between different samples of them radiolucent on plain films. The degree of bony calcification is incomplete, but cooking alters the structure of bones and makes passage is approximately 25% to 60%; chances increase with prolonged exposure can be minimized if adult-sized radiograph cassettes in case the FB has passed into the stomach or beyond. Radiation is sometimes obtained for reassurance of the presence of the FB in the stomach.

In adults, if neck or chest films are negative, abdominal films are used. Swallowed coins or other FBs may become lodged in the nasopharynx, usually after gagging or vomiting, and may aid in the verification of radiopaque FBs. Soft tissue swelling, indirect evidence of esophageal obstruction, and may aid in the setting of suspected FB entrapment. CT in cases in which radiographic evaluation is required. CT of the neck and mediastinum is an easy, rapid, cost-effective, and accurate noninvasive means of detecting or excluding esophageal FBs, and has garnered support in the setting of suspected FB entrapment. CT further excels at localization and characterization of the impacted FB and identification of associated complications, such as perforation. CT has a sensitivity and accuracy of 97% and 98%, respectively, for the diagnosis of esophageal FBs compared to plain films. CT provides improved diagnostic utility for fish bone FBs over plain radiography when obtained either with or without barium enhancement. The sensitivity of CT can be improved with 3D reformations. These allow better visualization of the FB and any esophageal injury. Use of CT in patients in whom clinical suspicion for a retained FB is high has the potential to reduce the number of unnecessary endoscopies.

The use of intravenous (IV) contrast with CT is not indicated, unless there is concern for an inflammatory process (abscess, peritonitis, fistula formation), or vascular injury (aortic or caval perforation). Use of oral contrast with CT in the setting of suspected FB remains controversial. If used, it can impair visualization on endoscopy. Use of oral contrast is generally reserved for those rare cases where either esophageal perforation is suspected (water-soluble media should be used), or a nonradiopaque FB is suspected and the radiologist wants a small amount of contrast to “outline” the potential FB.

**Computed Tomography**

CT of the neck and mediastinum is an easy, rapid, cost-effective, and accurate noninvasive means of detecting or excluding esophageal FBs, and has garnered support in the setting of suspected FB entrapment. CT further excels at localization and characterization of the impacted FB and identification of associated complications, such as perforation.

**Contrast-Enhanced Esophagograms**

**Background**

A contrast-enhanced esophagogram is a test that is almost never indicated in the ED as a routine intervention to evaluate...
Contrast material (barium) may interfere with the detection and extraction of FBs at endoscopy, and may increase the risk for aspiration pneumonitis. Therefore routine, serial contrast-enhanced esophagograms after negative plain radiography in patients with known or suspected FBs are unnecessary for diagnostic purposes in most cases. Selective use is reasonable, but CT or endoscopy are interventions with a better and more cost-effective yield. The American Society for Gastrointestinal Endoscopy guidelines suggest “avoiding contrast radiographic examinations” before endoscopic removal. The rare indication in the ED would be a questionable FB or perforation where the radiologist or consultant requests the procedure to “outline” a possible FB to better visualize it, or to evaluate for leakage of contrast material.

**Procedure**

Esophagograms couple voluntary ingestion of an enteric contrast agent (Gastrografin [Bracco Diagnostics Inc., Monroe Township, NJ] or barium) and plain radiography. Immediately after ingestion, erect and horizontal radiographs are performed at right-angle projections (PA and lateral, or right and left anterior oblique). In addition to anatomic abnormalities, radiolucent FBs may be identified by contrast delineation or filling defects within the contrast column (Fig. 39.7). The initial choice of contrast agent is debated and should be individualized according to the threat of aspiration and perforation. Water-soluble Gastrografin is indicated first in most cases of suspected perforation because it causes less mediastinal inflammation when extravasated; however, it can give rise to severe chemical pneumonitis if aspirated, and is relatively contraindicated in patients with complete esophageal obstruction. Patients without evidence of complete esophageal obstruction are instructed to swallow progressively larger aliquots of contrast agent up to approximately 50 mL. If these films are normal, the procedure is repeated with half-strength and then full-strength barium to delineate small esophageal injuries. Note that water-soluble contrast material (Gastrografin) causes more pulmonary reaction than barium does when inadvertently aspirated, and should be used in small aliquots if aspiration or complete esophageal obstruction is a concern.
Conclusions

Diagnostic radiography for esophageal FBs is indicated in almost all cases with the exception of those with a complete obstruction where there is no concern for aspiration. Plain radiographs clearly assist the clinician in several situations: (1) screening of children, adults with dementia, and nonverbal patients with a history or symptoms suspicious for purposeful or inadvertent FB ingestion that can be assumed to be radiopaque; and (2) localization of known radiopaque FBs to clarify underlying esophageal pathology.

Follow-up Radiographs

Often FBs are visualized in the stomach, or procedures performed in the ED (described later) result in the FB moving into the stomach. A variety of organizations have proposed recommendations for how to manage FBs in the stomach and intestines (Table 39.2).

Barium interferes with endoscopy and should not be used when endoscopy is anticipated.

<table>
<thead>
<tr>
<th>FOREIGN BODY TYPE</th>
<th>LOCATED IN THE ESOPHAGUS</th>
<th>LOCATED IN STOMACH OR BOWEL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coin</td>
<td>Remove within 24 hours if still present</td>
<td>Stomach—x-ray weekly</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Remove if no progression within 3–4 weeks</td>
</tr>
<tr>
<td>Button battery</td>
<td>Remove emergently</td>
<td>Stomach—repeat x-ray in 2 days</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Past pylorus—repeat x-ray in 3–4 days</td>
</tr>
<tr>
<td>Magnet</td>
<td>Remove all magnets within endoscopic reach (proximal duodenum or stomach)</td>
<td>Remove all magnets within endoscopic reach (proximal duodenum or stomach).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Daily x-ray to assess progression</td>
</tr>
<tr>
<td>Sharp object</td>
<td>Remove emergently endoscopically</td>
<td>Remove all objects &gt;2.5 cm wide or 6 cm long that are within endoscopic reach (proximal duodenum or stomach) urgently.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Daily x-ray to assess progression</td>
</tr>
<tr>
<td>Blunt object</td>
<td>Remove endoscopically within 24 hours</td>
<td>Remove all objects &gt;2.5 cm wide or 6 cm long that are within endoscopic reach (proximal duodenum or stomach).</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Weekly x-ray</td>
</tr>
</tbody>
</table>

Barium interferes with endoscopy and should not be used when endoscopy is anticipated.

Table 39.2 Summary of Recommendations for Esophageal Foreign Bodies

Conclusions

Diagnostic radiography for esophageal FBs is indicated in almost all cases with the exception of those with a complete obstruction where there is no concern for aspiration. Plain radiographs clearly assist the clinician in several situations: (1) screening of children, adults with dementia, and nonverbal patients with a history or symptoms suspicious for purposeful or inadvertent FB ingestion that can be assumed to be radiopaque; and (2) localization of known radiopaque FBs to clarify
the necessity for and means of FB extraction. Conversely, attempts to verify radiolucent FBs, including bones, by plain radiography are often misleading. CT is more accurate and sensitive than plain films in diagnosing FBs, and in most cases, contrast is not needed.

**VISUALIZATION OF ESOPHAGEAL AND PHARYNGEAL FBs**

Patients with an FB sensation in their oropharynx typified by a “fish bone” or “chicken bone” sensation need to have some form of visualization of their oropharynx performed as part of the physical examination. The three procedures are direct pharyngoscopy, which is simply direct visualization or examination using a tongue blade with a light source that may be a pen light, wall light, or head light; indirect laryngoscopy, which involves using a handheld mirror reflecting a light to allow visualization of the epiglottis, vallecula, arytenoids, arytenoid folds, and vocal cords (a procedure that requires experience and a cooperative patient); or nasopharyngoscopy, a procedure using a flexible nasopharyngoscope. If an FB is visualized (Fig. 39.8), it should be removed with forceps and the oropharynx carefully reexamined for any injury or additional FB. All three of these procedures are discussed in detail in Chapter 63.

**ESOPHAGOSCOPY**

Esophagoscopy is the definitive diagnostic and therapeutic procedure for impacted esophageal FBs. Although esophagoscopy is not a procedure performed by the emergency provider, its proper role in the ED evaluation of FBs must be understood. With esophagoscopy, the provider can document the presence and location of the FB along with any underlying lesion. The clinician can then remove the object and reevaluate the esophagus after removal of the FB to rule out perforation or underlying pathology. Esophagoscopy may be necessary even if a radiologic contrast-enhanced study does not reveal complete obstruction because plain radiographs are not always conclusive. Esophagoscopy may be necessary to exclude predisposing pathology or resultant perforation, even when symptoms presumed to be caused by an esophageal FB have resolved.

Esophagoscopy is the preferred method for removal of sharp or pointed objects such as bones, open safety pins, and razors. In the case of sharp objects prone to causing esophageal perforation, intravenous antibiotics should be administered before the procedure. Endoscopy is the preferred way to remove an impacted meat bolus and to evaluate for possible esophageal pathology at the same time. Esophagoscopy is also indicated for an FB retained for more than 24 to 48 hours, both to remove it and to examine for esophageal wall erosion or perforation. Esophagoscopy is the only appropriate removal technique for multiple or large esophageal FBs. This technique is also indicated for patients with an FB proved to have passed into the stomach and for those who have persistent symptoms possibly caused by esophageal wall injury. Flexible endoscopic procedures can usually be performed without general anesthesia, even in most children. The success rate of flexible endoscopy in patients with retained esophageal FBs exceeds 96%.

Traditionally, esophagoscopy is up to 10 times more expensive than other maneuvers, such as Foley catheter removal or esophageal bougienage (described later), largely because of charges for the surgical suite, but it has a higher success rate than the other two techniques. ED removal of esophageal FBs in children by experienced endoscopists, while the child is under conscious or deep sedation administered by the emergency provider, has been described. In selected cases this approach can shorten the interval to completion of the procedure and reduce expense.

**ESOPHAGEAL PHARMACOLOGIC MANEUVERS**

**Background**

As the LES is the narrowest portion of the entire GI tract, most FBs that reach the stomach eventually move through the GI tract without further problems. Because a large number of entrapped esophageal FBs are lodged at the LES, especially in adults, several therapeutic maneuvers have been proposed to assist transit into the stomach, including pharmacologic relaxation of the LES. In theory, agents that promote smooth muscle relaxation should improve mobility through the LES. Although many clinicians use pharmacologic adjuncts for all esophageal FBs, objects lodged at the LES will probably benefit most from such interventions. Nonspecific pain relief, anxiolysis, vomiting, and spontaneous passage over time may account for the success attributed to many pharmacologic manipulations of esophageal FBs as there is little scientific evidence that any of these agents offer any benefit over placebo.

**Indications and Contraindications**

The indication for pharmacologic relaxation of the LES is the presence of a smooth or blunt FB such as a coin or food bolus. Angulated, abrasive, or sharp FBs should not be treated with pharmacologic modalities but instead should be removed by esophagoscopy. Analgesics and sedatives are routinely indicated if pain is present or the patient is excessively anxious.
Glucagon

**Pharmacology**

Glucagon was historically a prototype for the spasmolytic agents. Glucagon theoretically relaxes esophageal smooth muscle and decreases LES resting pressure. Glucagon has no effect on the upper third of the esophagus, a common site of coin impaction in children, where striated muscle is present and some voluntary control is operative. It only minimally affects the middle third of the esophagus. Peristalsis is not affected by glucagon. Results with glucagon have been mixed, and the only randomized study, done in children, showed no better results than those achieved with placebo. Nonrandomized studies in adults showed that glucagon is approximately equivalent to placebo, with a 33% success rate. A meta-analysis has also shown no benefit over placebo. However, its use is still advocated by some authorities such as the American Society for Gastrointestinal Endoscopy as an acceptable option as long as it does not “delay definitive endoscopic removal of a food impaction.” Likewise, the North American Society for Pediatric Gastroenterology, Hepatology and Nutrition has stated that glucagon “may be considered in patients with distal esophageal impactions” or in “facilities where endoscopic care is not readily available.” As glucagon has little downside and emergency providers may be asked to administer it by their consultants for esophageal food impactions, it is reasonable to understand its use despite lack of scientific evidence for efficacy.

**Indications and Contraindications**

Glucagon is used for smooth FBs or food impactions at the LES that are suspected because of a patient’s complaint of pain or “something stuck” in the lower part of the chest or epigastrium. The clinical diagnosis is usually straightforward, with a 33% success rate. In children, where striated muscle is present and voluntary control is operative, it is reasonable to understand its use despite lack of scientific evidence for efficacy.

**Complications**

Glucagon is associated with a few minor side effects. If administered too rapidly, it causes nausea and vomiting. Therefore adult patients must be alert and mobile enough to avoid aspiration. Occasionally vomiting dislodges the impacted food bolus. Theoretically, there is a risk for rupture of the obstructed esophagus during induced emesis, so slow injection is preferred to minimize this side effect.

**Administration of Glucagon**

The therapeutic dose is 0.25 to 2 mg administered intravenously (Table 39.3) over a period of 1 to 2 minutes in a seated patient, although one study found that in normal subjects 1 mg of glucagon provides no significant additive benefit over 0.5 mg. The patient is given water orally within 1 minute after the injection of glucagon to stimulate normal esophageal peristalsis; this helps push the food through the relaxed LES into the stomach. Glucagon has a rapid onset and short duration of action: GI smooth muscle relaxes within 45 seconds, and its duration of action is approximately 25 minutes. If no results are seen within 10 to 20 minutes, a second administration of 0.25 to 2 mg may be tried. Success rates are higher when glucagon is combined with gas-forming agents or even carbonated beverages. It is recommended that a small volume of some oral fluid be routinely given to enhance the activity of glucagon.

**TABLE 39.3** Recommended Pharmacologic Therapies for Esophageal FBs

<table>
<thead>
<tr>
<th>CLASS AND AGENTS</th>
<th>SITE OF ACTION</th>
<th>DOSE AND ROUTE</th>
<th>ADVERSE EFFECTS</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Spasmolytics</strong></td>
<td>Glucagon</td>
<td>1–2 mg IV&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Nausea, vomiting, hyperglycemia, hypersensitivity</td>
</tr>
<tr>
<td>Nitroglycerin</td>
<td>Body and LES</td>
<td>0.4–0.8 mg SL&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Hypotension, tachycardia, bradycardia</td>
</tr>
<tr>
<td>Nifedipine</td>
<td>LES</td>
<td>5–10 mg SL&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Hypotension, tachycardia Use with caution</td>
</tr>
<tr>
<td><strong>Gas-Forming Agents</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tartaric acid</td>
<td>Distal and proximal</td>
<td>15 mL tartaric acid (18–20 g/100 mL)&lt;sup&gt;d&lt;/sup&gt;</td>
<td>Vomiting, increased intraesophageal pressure</td>
</tr>
<tr>
<td>Sodium bicarbonate</td>
<td>Distal and proximal</td>
<td>15 mL sodium bicarbonate (10 g/100 mL)&lt;sup&gt;d&lt;/sup&gt;</td>
<td>Vomiting</td>
</tr>
<tr>
<td>Carbonated beverage</td>
<td>Distal and proximal</td>
<td>100 mL PO</td>
<td>Increased intraesophageal pressure</td>
</tr>
</tbody>
</table>

<sup>a</sup>May be repeated once or used in conjunction with nitroglycerin.

<sup>b</sup>One to 2 inches of nitroglycerin paste applied under an occlusive dressing may be an alternative.

<sup>c</sup>A capsule is punctured, chewed, held in the mouth for 3 minutes, and then swallowed. Because of hypotension, a 5-mg dose may be used in the elderly. Do not use if the patient has cardiovascular disease, is hypotensive, or has also recently been given nitroglycerin.

<sup>d</sup>Alternatively, dissolve 2 to 3 g tartaric acid and 2 to 3 g sodium bicarbonate in 30 mL water.

FB, Foreign body; IV, intravenously; LES, lower esophageal sphincter; PO, per os (orally); SL, sublingually.
of clinical concern, and blood glucose levels do not need to be monitored. No fatalities have been reported. Although theoretically glucagon can stimulate release of catecholamine in patients with pheochromocytoma and can induce hypoglycemia from reflex release of insulin with an insulinoma, these endocrine tumors are rare. Nonetheless, precipitation of either profound catecholamine or insulin reactions with the use of glucagon should direct a workup for these underlying tumors.

**Further Evaluation and Therapy**

If the patient experiences relief of symptoms after the administration of glucagon, a postprocedure radiograph or contrast-enhanced study may be obtained to confirm passage of a radiopaque object, but this is not mandatory (Fig. 39.9). Adult patients with successful FB passage into the stomach may be discharged home, but careful follow-up should be obtained to rule out coexistent esophageal pathology because a significant number of patients (65% to 80%) will have underlying esophageal disorders. Pediatric patients with food boluses lodged in the esophagus also have a high rate of underlying esophageal disease and need referral for further evaluation; this is not the case for accidentally swallowed FBs in children.

If glucagon fails to produce relief of the symptoms or resolve the radiograph findings, its use does not preclude other methods from being used.

**Nitroglycerin and Nifedipine Pharmacology**

Both sublingual nitroglycerin and nifedipine have been used in a manner similar to glucagon to relieve LES tone and allow the passage of a distal esophageal FB. Although these two agents have been used less than glucagon for the treatment of esophageal FBs, both are useful for the relief of chest pain associated with esophageal smooth muscle spasm and may be administered concurrently with glucagon. Nifedipine, conversely, significantly reduces LES pressure without changing contraction amplitudes in the body of the esophagus. Thus nitroglycerin may relieve partial or complete obstruction of the middle or lower part of the esophagus secondary either to intrinsic esophageal disease or to simple FB impaction; and nifedipine, like glucagon, is most likely to succeed when the bolus is lodged at the gastroesophageal junction.

**Indications and Contraindications**

Similar to clinical indications for the use of glucagon, any patient with an impacted smooth esophageal FB, especially a food bolus, may be a candidate for nitroglycerin or nifedipine (or both). In addition, similar to the mode of action of glucagon, neither of these agents is expected to relax a fixed fibrotic stricture or ring at the gastroesophageal junction. Nevertheless, because both agents have a relatively benign side effect profile, if the patient has no contraindication to their use, they may be tried with or without previous documentation of the distal esophageal obstruction with a contrast-enhanced study. Contraindications to their use include a history of allergic reactions, a sharp esophageal FB, hypovolemia, and hypotension.

**Use and Complications**

Doses of 1 or 2 (0.4 mg) sublingual nitroglycerin tablets, 1 to 2 inches of nitroglycerin paste, or 5 to 10 mg of nifedipine have been reported. Some patients with esophageal FBs may have some degree of dehydration because of the inability to swallow liquids or their own saliva. These patients may be prone to hypotension from the vasodilation associated with the use of either agent. Ideally, rehydration should precede therapy with these agents. Do not use both agents simultaneously. The smaller dose of nifedipine is suggested in the elderly or those with cardiovascular disease.

**Further Evaluation and Therapy**

As with the use of glucagon, if nitrate therapy fails to produce relief of the symptoms or resolve the radiographic findings, its use does not preclude trying another method. If a patient experiences symptomatic relief, a postprocedure radiograph may be obtained to confirm passage of a radiopaque object, but this is not mandatory. Adult patients may be discharged home, but careful follow-up should be obtained to rule out coexistent esophageal pathology because a significant number of patients will have underlying esophageal disorders.

**Gas-Forming Agents**

**Pharmacology**

The use of gas-forming agents for the treatment of distal esophageal food impaction, especially meat boluses, was first described in 1983. The combination of tartaric acid solution
followed immediately by a solution of sodium bicarbonate or even carbonated beverages has been reported. In theory, use of this acid-base mixture or a carbonated beverage may produce sufficient carbon dioxide to distend the esophagus, relax the LES, and push impacted food through the gastrosophageal junction into the stomach.  

**Indications and Contraindications**

Gas-forming agents are indicated for the relief of smooth distal esophageal FB impaction, with or without prior FB confirmation by a radiographic study (see Fig. 39.9). They are often given to patients with food impaction or retained coins. Although gas-forming agents are more likely to succeed with distal esophageal impactions, they have also been successful in relieving obstructions in the proximal part of the esophagus. Concurrent administration of spasmolytic agents may improve the effectiveness of gas-forming agents.

**Use and Complications**

A solution of 15 mL of tartaric acid (18.7 g/100 mL), followed by 15 mL of a sodium bicarbonate solution (10 g/100 mL), or 1.5 to 3 g of tartaric acid and 2 to 3 g of sodium bicarbonate dissolved in 15 mL of water can be used. Carbonated beverages (100 mL) have also been successful in the transit of FBs into the stomach and are more readily available in the ED. Many patients with esophageal FB impaction have been noted to retch after receiving gas-forming agents, which theoretically puts patients at risk for esophageal trauma, including rupture. Gas-forming agents should not be given to patients with impactions of greater than 6 hours’ duration, or to patients with chest pain that might be indicative of an esophageal injury.

**Further Evaluation and Therapy**

As with the use of glucagon, nitroglycerin, or nifedipine, even if administration of the gas-forming agent is successful, as judged by relief of symptoms, follow-up evaluation is necessary to determine the underlying esophageal abnormality that potentially led to the FB impaction.

**Papain**

The use of papain, a proteolytic enzyme used commercially as a meat tenderizer, is NOT recommended for treatment of an esophageal food FB. The North American Society for Pediatric Gastroenterology, Hepatology and Nutrition states that it “is contraindicated.” Papain is not currently recommended because of the unacceptable risk for complications and the availability of safer, more effective interventions. It has been associated with esophageal injuries, pulmonary edema, and hemorrhages. There is one published report of a retrospective series with no complications from using this agent, but the preponderance of data and expert opinion would weigh against its use in the ED.

**REMOVAL OF ESOPHAGEAL FBs IN THE ED**

In the United States the most prevalent method for removing FBs lodged in the esophagus is referral for removal under direct visualization via endoscopy, normally done in either an endoscopy suite or operating room under sedation. Endoscopy is costly and time-consuming, and if the patient is being seen after hours, the procedure requires additional personnel to come into the hospital to care for the patient. With procedural sedation becoming more common in EDs, procedures to remove lodged esophageal FBs, when appropriate, are becoming more common. Currently three procedures are used in the ED to remove FBs lodged in the esophagus, and these are somewhat dependent on the type and location of the FB. These procedures include Magill forceps removal, Foley catheter removal, and esophageal bougienage. Each will be described separately. Another strategy, “watchful waiting,” is reserved for single, smooth, asymptomatic FBs (usually coins) typically at the LES in children and will also be discussed.

**Magill Forceps Removal of Esophageal FBs**

**Background**

In children, the most common accidentally ingested FB is a coin, and the most common location for the FB to be lodged is at the cricopharyngeus muscle (Fig. 39.10). Given these facts, when children have a single coin at the level of the cricopharyngeus muscle confirmed by radiographs, they are candidates for Magill forceps removal. The procedure requires sedation (see Chapter 33), and thus all airway equipment must be available, in addition to monitoring equipment and expertise in its use. In centers that are using this procedure, success rates have ranged from 95% to 100%. All centers reporting this procedure used sedation and visualization with a laryngoscope or video-assisted system. The procedure was performed rapidly, usually in less than a minute. Complications were rare and typically consisted of minor bleeding or vomiting. In the one center that intubated patients before the procedure, complications were higher and associated with the intubation. Magill forceps removal seems to be ideally indicated for stable children with a coin at the cricopharyngeus muscle in a facility that is well equipped and staffed and experienced in managing procedural sedation and airways in children.
**Foley Catheter Removal of Esophageal FBs**

**Background**

Foley catheter removal of esophageal FBs was first described in the thoracic surgery literature in 1966 and in the emergency medicine literature in 1981. The technique has essentially been unchanged since the first reports and is now used by emergency providers, radiologists, otolaryngologists, and general surgeons. The classic patient for this technique is a small child who is brought to the hospital shortly after swallowing a coin that is documented by radiography, but the procedure may be used for a wide variety of smooth FBs in patients of all ages. Success rates for Foley catheter removal of FBs have been cited from 85% to 100%, with complication rates of 0% to 2%. Many of the reported complications were caused by nasal insertion of the catheter, and complication rates are lower when the catheter is inserted orally and at centers that perform the procedure frequently. Foley catheter extraction of FBs costs significantly less than endoscopy. Fluoroscopic assistance may be preferable, but it is not essential. Whether the procedure is performed in the ED or the radiology department, equipment and personnel

---

**Procedure (Fig. 39.11)**

1. Ensure that adequate procedural sedation and appropriate monitoring are both implemented (refer to Chapter 33). Place the patient supine on the stretcher with the head slightly extended in the “sniffing position.”
2. Insert the laryngoscope or video-assisted laryngoscope. Suction as necessary.
3. Visualize the upper part of the esophagus, where the FB is normally lodged.
4. Grasp the FB with the Magill forceps and then slowly remove it.
5. Visualize the esophagus after removal for any injuries such as erosion or bleeding.
6. Recover the patient from procedural sedation.

**Aftercare**

No specific aftercare is necessary if no complications have occurred from the procedure. If there is any evidence of esophageal injury, the patient needs to be referred to gastroenterology immediately. Children who ingest FBs are at risk for future ingestions, with the “repeat offender” rate being as high as 18% to 20%.

---

**Figure 39.11** Removal of an esophageal foreign body with Magill forceps.
capable of emergency pediatric airway management must be present.

**Indications**
Recently ingested smooth, blunt objects that are radiographically opaque are most suitable for balloon catheter extraction. Recently ingested FBs carry little likelihood of causing pressure necrosis, perforation, or other significant injury; however, 24 to 48 hours’ duration of impaction should be the upper limit for consideration of this technique.\(^3^2,^6^7\) Coins are particularly amenable to Foley manipulation, but food boluses and button batteries have also been extracted successfully.\(^7^3\) Radiographically opaque objects are most easily located with plain radiographs. Radiopaque objects can be manipulated, but uncertainty about location mandates contrast-enhanced esophagography.

**Contraindications**
Contraindications to catheter removal of esophageal FBs include total esophageal obstruction, as manifested by an air-fluid level on a plain radiograph or contrast-enhanced esophagogram or when patients are unable to handle oral secretions. The presence of a total obstruction prevents passage of the tip of the catheter distal to the FB. Esophageal perforation and airway distress, as recognized by the typical symptoms and signs, require immediate surgical consultation and preclude blind esophageal manipulation. The presence of multiple esophageal FBs also precludes use of the Foley catheter. Sharp, irregularly shaped FBs should not be removed with this technique because esophageal perforation or laceration can result, and the balloon may burst during the procedure. Finally, lack of expertise or equipment to handle an airway problem arising during the procedure is a contraindication.

**Procedure (Fig. 39.12)**
1. Coach the patient on the procedure. Topical oropharyngeal anesthesia may be used (although this increases the risk for aspiration). Light sedation may be used.
2. Place the patient in the Trendelenburg, lateral decubitus, or prone position.
3. Insert an uninflated catheter (10 to 16 Fr in children) so that the tip is past the FB (as visualized on fluoroscopy or as estimated on radiographs).
4. Fill the catheter balloon slowly with 3 to 5 mL of saline or contrast material (if fluoroscopy is used). Stop inflating the balloon if the patient complains of increased pain.

**Figure 39.12** Removal of an esophageal foreign body with a Foley catheter.
5. Use steady, gentle traction to withdraw the catheter.
6. Once the FB and the tip of the catheter reach the hypo-
  pharynx, grasp the object with forceps, or instruct the patient
to spit it out.
7. Obtain another radiograph to ensure that multiple FBs
were not present.
8. If the catheter slips by the FB, reinsert the catheter, inflate
it with an additional 2 to 3 mL of fluid, and make one
additional attempt at withdrawal.
9. If fluoroscopy is not used and no FB is retrieved, another
radiograph should be obtained because 10% to 20% of the
time the FB will pass distally into the stomach.

Complications
Complication rates of 0% to 2% have been reported.75,77–81
Many complications (nosebleeds or displacement of the FB
into the nose) have been related to nasal insertion of the catheter.
Complication rates are lower when the catheter is inserted
orally and generally lower at centers that perform the procedure
frequently. No deaths have been reported. Laryngospasm and
aspiration are rare complications. Failure to either remove the
object or displace it into the stomach occurs in approximately
2% to 10% of carefully selected patients;75,79,80 but success
rates are lower in adults or patients with underlying esophageal
disorders.80

Aftercare
Children who have an FB removed successfully with a Foley
catheter need no further follow-up if they remain asymptomatic.
If the FB was moved into the stomach, clinical follow-up should
be adequate to verify movement of gastric FBs through the
alimentary tract. Discharge instructions should include warnings
about the potential symptoms of GI obstruction, perforation,
and hemorrhage. Parents of children who swallow coins can be
instructed to watch for coins in their stool. Adults with esopha-
geal FBs that have been removed successfully must be referred
for evaluation of possible esophageal pathology, as should
children with food impactions. Should an FB remain lodged in
the esophagus, immediate referral for endoscopy is necessary.

Esophageal Bougienage

Background
Displacement of esophageal FBs into the stomach can be done
with nasogastric or orogastric tubes or via esophageal bougie-

age. Esophageal bougienage is a technique for dislodging
impacted esophageal coins by blind mechanical advancement
of the coin into the stomach, a procedure first described in
1965.82 The technique has greater than a 95% success rate
with essentially no reported complications when used for the
proper indications.50–53,81,83,84 Rates as high as those with
endoscopy have been reported.67,81 Furthermore, bougienage
is unrivaled in overall cost-effectiveness because it is approxi-
mately 10% of the cost of endoscopic removal.50–53

This technique does not allow visualization of the esophagus
or retrieval of the object. There have traditionally been warnings
against forceful advancement of esophageal FBs, but growing
evidence verifies the efficacy and safety of blind esophageal
bougienage as first-line therapy for coin ingestions in properly
selected patients. Although early articles suggested that
esophageal bougienage should be performed exclusively by
pediatric surgeons, the technique is easily mastered and used
by emergency clinicians.50,51,53,83,84

Indications and Contraindications
Strict patient selection is paramount for successful and uncom-
plicated bougienage. The criteria have changed little since
initially proposed and define a group in whom a round, smooth
object can be forcibly passed into the stomach with little risk.50,51
Although many swallowed objects meet this description, only
coins hold clear supportive evidence in the literature. Because
many patients with impacted food or meat have underlying
esophageal pathology, we do not suggest that it be used in the
ED for this condition. Selection criteria are the following: a
single, smooth FB lodged less than 24 hours in a patient with
no respiratory distress or history of esophageal disease, including
previous FBs, or surgery. The FB should be likely to pass
beyond the stomach without complications. The procedure is
contraindicated in patients who do not satisfy all the criteria.
It is important to ascertain the duration of esophageal impaction
to avoid performing the procedure when there may be underly-
ing esophageal injury. Plain radiographs are indicated to verify
coin location and the absence of multiple esophageal bodies.
Preprocedure esophagograms are not required.

Procedure (Fig. 39.13)
1. Coach the patient (and parents) on the procedure. Place
the patient in an upright position.
2. Achieve topical anesthesia with gurgled 2% to 4% viscous
lidocaine, atomized 2% lidocaine, or topical benzocaine
spray.
3. Flex the patient’s head, open the mouth, and ask the patient
to protrude the tongue. Use a tongue blade if necessary.
In some patients a bite block may be needed; several tongue
blades can be taped together for this purpose.
4. Pass a well-lubricated, appropriately sized bougie (Table
39.4) posteriorly along the roof of the mouth and caudally
to the hypopharynx.
5. The patient may gag momentarily; ask the patient to
swallow and gently pass the dilator past the cricopharyngeus
muscle.
6. Ask the patient to phonate while passing the bougie at
this point.
7. Once past the cricopharyngeus, extend the head to enable
the bougie to pass distally to the stomach with little
resistance.
8. Withdraw the bougie after a single pass.
9. Terminate the procedure immediately if pain or resistance
to advancement is encountered.
10. Obtain a postprocedure x-ray to ascertain passage of the
coin into the stomach.

Complications
Gagging and self-limited nonbloody vomiting are not uncom-
mon after the procedure and may reveal the dislodged coin.
Patients may experience a residual FB sensation for several
hours. Pulmonary aspiration and inadvertent passage into the
airway are potential complications. Likewise, traumatic pha-
ryngeal and esophageal injury, ranging from mild self-limited
bleeding to frank esophageal perforation with concomitant
infection, are possible but rare.

Aftercare
Plain radiographs of the chest and abdomen are indicated to
verify passage of the FB into the stomach and to check for
any potential complications. Discharge asymptomatic patients
with appropriate precautions, including the need to return if
Esophageal Bougienage

1. Place the patient in the upright position. Achieve topical anesthesia with gargled 2% to 4% viscous lidocaine, atomized 2% lidocaine, or topical benzocaine spray.

2. Coach the patient (and parents) on the procedure.

3. Pass the lubricated bougie posteriorly along the roof of the mouth and caudally to the hypopharynx.

4. The patient may gag momentarily; ask him to swallow, and then gently pass the dilator past the cricopharyngeus muscle.

Once past the cricopharyngeus, extend the head to enable the bougie to pass distally to the stomach with little resistance.

Withdraw the bougie after a single pass.

Terminate the procedure immediately for pain or resistance to advancement.

Coat the patient (and parents) on the procedure.

Pass the lubricated bougie posteriorly along the roof of the mouth and caudally to the hypopharynx.

The patient may gag momentarily; ask him to swallow, and then gently pass the dilator past the cricopharyngeus muscle.

Withdraw the bougie after a single pass.

Terminate the procedure immediately for pain or resistance to advancement.

Figure 39.13 Esophageal bougienage.

### TABLE 39.4 Hurst Dilator Size by Age for Esophageal Bougienage

<table>
<thead>
<tr>
<th>PATIENT AGE</th>
<th>HURST DILATOR SIZE</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 to 2 years</td>
<td>28 F</td>
</tr>
<tr>
<td>2 to 3 years</td>
<td>32 F</td>
</tr>
<tr>
<td>3 to 4 years</td>
<td>36 F</td>
</tr>
<tr>
<td>4 to 5 years</td>
<td>38 F</td>
</tr>
<tr>
<td>5 years or older</td>
<td>40 F</td>
</tr>
</tbody>
</table>

SPECIAL SITUATIONS

**Childhood Coin Ingestion**

Coins are the most commonly ingested objects by preschool-age children. In most cases the ingestion is quickly realized by a caretaker, and in the majority of cases the coins pass uneventfully. Rarely, an esophageal coin can be clandestine for many weeks or months and produce a variety of vague respiratory or GI symptoms. In addition, many coin ingestions are not witnessed, so maintain a high index of suspicion for children with dysphagia, drooling, or crying, who may have esophageal FBs, most likely coins.

Most coins pass from the esophagus to the stomach with only transient symptoms. The child may be in pain for a few minutes as the coin migrates, but on arrival at the ED, the child is often asymptomatic. Coins that remain in the esophagus are likely to, but do not always, produce continued symptoms (e.g., drooling, pain, dysphagia, and refusal to eat or drink). Rarely, esophageal coins can cause airway distress by external compression of the trachea and simulate an asthmatic attack. Coins below the diaphragm are asymptomatic, and the presence of pain or symptoms requires further evaluation. Coins in the trachea produce immediate and obvious respiratory distress.

Although up to 44% of children with esophageal coins may be asymptomatic, and up to two thirds of coins will be in the stomach at the time of ED evaluation, it is prudent to obtain plain radiographs in all children with a suggestive history. In most cases a single film that includes the pharynx, esophagus, and stomach will suffice to prove or exclude an ingested coin (Fig. 39.14). Another advantage of obtaining radiographs is to rule out multiple FB ingestions, which are not uncommon in children. Only a single PA chest film is needed to prove signs of respiratory compromise, chest or abdominal pain, dysphagia, hematemesis, persistent vomiting, or other concerns are present. Follow-up abdominal radiographs may be performed to document passage of the coin if it is not identified in feces within 1 week.
the presence of a coin, but a lateral projection is also suggested. If the flat surface of the coin is seen (see Figs. 39.2 to 39.4), this orientation ensures an esophageal position. If the edge of the coin is seen, this orientation suggests that it has traversed the vocal cords, but a coin in the airway is not subtle and produces obvious distress. It is advisable to also routinely obtain a lateral radiograph to determine whether multiple coins are stacked on top of each other (Fig. 39.15).

Once a coin’s presence has been documented, a decision concerning removal must be made. The approach varies and there is no agreed standard. Overall, approximately 25% of coins will pass spontaneously, even if the coin is located proximally. Observation for 8 to 16 hours is a reasonable approach for asymptomatic children if the ingestion has taken place within 24 hours. Coins in the upper and middle third of the esophagus are less likely to pass spontaneously, and some prefer to remove them as soon as the diagnosis is made.\(^\text{19}\)

Coins in the distal end of the esophagus will pass spontaneously in one third to one half of patients within 24 hours.\(^{34,88,89}\)

The decision regarding management of these patients depends on various factors: clinician comfort and experience with removal techniques, local protocols and procedures developed by the medical staff of each institution, and comfort level of the caretakers with various therapeutic options. Regardless of the approach, a radiograph should be taken just before removal to ensure that spontaneous passage has not occurred.

One suggested protocol (Fig. 39.16) involves radiologically localizing the coin, and if the child is symptomatic, immediately removing the coin. If the patient is asymptomatic, the coin may be removed immediately, or the patient may be observed either as an inpatient or at home. If the child is asymptomatic, one common practice is to allow the child to drink a carbonated beverage and eat a small amount of soft food in the ED, wait approximately 1 to 2 hours, and obtain another radiograph. If sent home with an asymptomatic retained FB, the patient is allowed to eat or drink but should be rechecked in 12 to 24 hours with the knowledge that up to 50% of asymptomatic coin FBs will pass into the stomach spontaneously.\(^{34,88,89}\) Both the American Society for Gastrointestinal Endoscopy and the North American Society for Pediatric Gastroenterology, Hepatology, and Nutrition recommendations for impacted esophageal coins in children state that asymptomatic coins in the esophagus may be observed up to 24 hours before removal.\(^{10,20}\)

The techniques of Magill forceps removal, esophageal bougienage, and Foley catheter removal have been described earlier. All are options for single coins present in the esophagus for less than 24 to 48 hours. Another option for coins impacted at the LES is pharmacologic relaxation of the sphincter to aid passage into the stomach, although scientific evidence for efficacy is lacking. The most common method to remove esophageal coins in use today is esophagoscopy.

Approximately two thirds of ingested coins are in the stomach at the time of first investigation, and such patients can be released home safely to allow almost certain spontaneous passage with a normal diet. Spontaneous passage of a coin from the stomach to the anus usually requires 3 to 7 days. There is no need for routine cathartic use. Parents should be advised to check the stool for the coin and return for repeated radiographs if the coin is not found in 1 to 2 weeks. Most coins are passed unknowingly by the patient. Any abdominal discomfort or distention warrants reevaluation in the ED. If a follow-up radiograph demonstrates a persistent coin in the intestines for more than 3 to 4 weeks, an obstructive lesion may be present, and further evaluation is warranted.

Finally, there are theoretical concerns about US pennies, which contain 97.5% zinc. Theoretically zinc can lead to mucosal ulceration from its caustic nature; however, the evidence to date suggests no increased risk from ingested pennies.\(^{92}\)

### Fish or Chicken Bones in the Throat

Patients who complain of a “bone” in their throat usually arrive at the ED within several hours of the onset of symptoms, and have generally tried a home remedy such as swallowing a piece of bread. These patients are typically able to pinpoint the location of their discomfort and have an FB sensation that is exacerbated by swallowing. Patients who are markedly symptomatic, vomiting, or unable to swallow require definitive therapy. Those with minor complaints may be evaluated safely over a period of a few days, often as outpatients.

In cooperative patients, careful examination of the oropharynx should be done by either direct or indirect laryngoscopy, or both. If the bone is seen, it should be removed with forceps. If the patient feels pain in the upper part of the throat, special attention is directed to the tonsils because bones often lodge in this area (Fig. 39.17). Strands of saliva may mimic a bone, and small bones may be difficult to see. More commonly, the area of complaint is below the oropharynx. In these patients, indirect laryngoscopy or nasopharyngoscopy should be the
If no bone is seen on physical examination, the bone may have passed after causing local irritation that persists, or the bone is present and not visualized because of location or consistency. Minor symptoms in the upper part of the throat probably represent persistent local irritation. Minimally symptomatic patients can be discharged and reevaluated in 24 hours. Those with complaints of an FB below the visualized pharynx, or with very bothersome persistent symptoms, should be evaluated with a CT scan of the neck or possibly the chest if the symptoms are distal. Positive scans are an indication for endoscopic removal of the bone. If the CT scan reveals no FB or postbone complication and the patient is stable, the patient is discharged home with follow-up within 24 hours.

Patients with oropharyngeal abrasions will usually be asymptomatic at that time. If still symptomatic on follow-up, endoscopy is advocated.

A small bone lodged in the esophagus for a few days is annoying and painful, but it is not generally an emergency. However, impacted bones can cause serious sequelae, often weeks later, and continued complaints cannot be ignored. Importantly, a lodged bone will not dissolve and rarely passes spontaneously once lodged in the mucosa. Referral and possible endoscopy are necessary if complaints persist for more than 2 to 3 days, even if the examination and CT scan are negative.

**Figure 39.15** A, Lateral radiograph of a child showing four stacked coins at the same location. A posteranterior (PA) view suggested a single coin. Multiple swallowed coins are common in children. It is important to obtain both PA and lateral films to ascertain the exact number and location of swallowed coins. B, A single coin was seen on the PA chest film, but this lateral film suggests three coins. However, they do not seem to be stacked directly on top of each other. This digital radiograph was accidentally exposed three times; in fact, only one coin was swallowed and x-rayed three times during minimal movement.
**Sharp Objects in the Esophagus**

Sharp objects cause the majority of complications seen in patients with esophageal FBs. Such objects include tacks, pins, open paper clips, bobby pins, toothpicks, and razor blades (Fig. 39.18). They will not usually pass spontaneously and should be removed urgently. The only appropriate removal technique is under direct visualization with endoscopy. Swallowed dentures or partial plates are a particular hazard in elderly, demented, or mentally challenged patients. Frequently there is no history of such, and patients have a variety of complaints, such as a sore throat or persistent vomiting (Fig. 39.19). Prisoners and psychotic patients are well known to clandestinely swallow multiple bizarre and sharp objects.

Attempts at radiographic localization are appropriate for metallic or radiopaque FBs. Most objects in the stomach, even those considered problematic, will transit the remainder of the GI tract if less than 6 cm in length or 2.5 cm in diameter. If larger than this, consult a gastroenterologist (see Table 39.2). If radiographs show the FB in the esophagus, endoscopic removal is indicated, and attempts to remove such objects in the ED by other methods are not indicated.

**Nonradiopaque Objects in the Esophagus**

Objects such as toothpicks, aluminum pull tabs from beverage cans, plastic, and food boluses cannot be visualized on plain radiographs and will normally not pass spontaneously. Toothpicks cause a higher percentage of complications than any other type of esophageal FB. As with fish bones, toothpicks often lodge in the tonsils or posterior pharynx and can be seen on direct vision. The imaging modality of choice is CT. Removal is mandatory and time-sensitive for sharp objects, especially toothpicks.

**Impacted Food Bolus**

A large bolus of food may become impacted in the esophagus, usually at the LES. This occurs most frequently in the elderly, those intoxicated while eating, or those with dentures. Frequently, underlying esophageal pathology is present such as a stricture or web, even in the young (see Fig. 39.6). Children with an impacted food bolus often have underlying esophageal pathology, and it may occur following prior surgery for congenital esophageal malformations. The diagnosis is usually straightforward, and patients may be in significant distress, gagging, and unable to swallow. A barium swallow may be used to confirm the diagnosis, but this is rarely necessary and is discouraged as it impairs visualization on endoscopy, and in cases of complete obstruction, risks pulmonary aspiration. The American Society for Gastrointestinal Endoscopy suggests avoiding contrast radiographic exams before endoscopic removal of esophageal FBs. Proceeding directly to endoscopy appears to be the most reasonable approach. Food boluses may be amenable to pharmacologic relaxation of the LES, but the definitive intervention is endoscopy to both remove the bolus and evaluate the esophagus for pathology.

The most logical ED approach is initial aggressive relief of symptoms (judicious narcotics, sedatives, antiemetics), followed by endoscopic removal. Papain is contraindicated.
Removal of impacted food is an urgent issue but need not be done immediately on arrival at the ED or in the middle of the night with inadequate resources; endoscopy can wait several hours. Frequently, pain relief and a few hours of relaxation will allow the bolus to slowly pass; if so, follow up with gastroenterology for evaluation of any esophageal pathology is mandatory (this is true for food boluses in patients of any age). Vomiting occasionally dislodges the impaction.

**Figure 39.17** Many fish bones become impaled in the soft tissues of the upper digestive tract. A, This woman felt a bone catch in her throat while eating fish. As is often the case, she was able to consistently localize the foreign body to the right submandibular area, thus suggesting that it could be seen by direct visualization. B, With only a tongue blade, local anesthetic spray, and good lighting, a fish bone was found embedded in the tonsil and was easily removed with forceps. Removal provided immediate and total relief, as is usually the case. Strands of saliva can mimic a fish bone, so be careful when probing and grasping. C, Fishbone visualized (arrow) embedded in a patient's right tonsil. D and E, This patient felt a fish bone in her left pharynx, and a small bone (arrow) was removed from her left tonsil, a common place to find a bone with such symptoms. (C, Courtesy Geoff deLaurier, MD, and Jerahme Posner, MD.)

**Button Battery Ingestion**

Button batteries lodged in the esophagus should be considered an emergency because of the potential for serious morbidity and mortality. They must be emergently removed. These batteries range in size from 7 to 25 mm and are radiopaque (Fig. 39.20). Batteries appear as round densities, similar to an impacted coin, but some demonstrate a “double-contour” configuration. It is important to distinguish between a coin and a button battery because button batteries require immediate removal. Batteries consist of two metal plates joined by a plastic seal. Internally, they contain an electrolyte solution (usually concentrated sodium or potassium hydroxide) and a heavy metal such as mercuric oxide, silver oxide, zinc, or lithium.

If ingested, these batteries often lodge in the esophagus. Mechanisms of injury include electrolyte leakage, injury from electrical current, heavy metal toxicity, and pressure necrosis. Of particular concern is the development of corrosive esophagitis or perforation as a result of caustic injury and prolonged mucosal pressure. Though essentially harmless in the stomach and intestines, batteries lodged in the esophagus should be considered an emergency because even new batteries are subject to corrosion and leakage, which can result in mucosal necrosis within a few hours of contact with the esophagus.

Esophageal impaction of batteries mandates immediate removal. Options include Magill forceps removal, Foley catheter removal, esophageal bougienage, or esophagoscopy. Esophagoscopy allows direct esophageal evaluation and a more controlled extraction. In addition, the “invasive” nature of
Swallowed magnets within endoscopic reach much be urgently removed.\textsuperscript{10,13,20,103} Some children with complications from multiple magnet ingestion have underlying conditions such as developmental delay or autism, but older children can inadvertently swallow magnets when using them to imitate a pierced tongue. Between 2002 and 2011 there were approximately 1600 magnet ingestions annually in the United States,\textsuperscript{104} almost exclusively in children. In 2011 the US Consumer Safety Product Commission issued an alert describing the safety risks from swallowed magnets (https://www.cpsc.gov/content/cpsc-warns-high-powered-magnets-and-children-make-a-deadly-mix). These toy magnets are small round strong magnets made from rare earth elements and are much stronger than traditional magnets used to make toys of various shapes, and if multiple magnets are swallowed, they will become adherent inside the bowel (Fig. 39.21). Multiple magnet ingestion is common, occurring up to 57\% of the time.\textsuperscript{105,106} Multiple magnets, especially if ingested at different times, may attract each other across layers of bowel and lead to pressure necrosis, fistula, volvulus, perforation, infection, or obstruction.\textsuperscript{103–107} A piece of bowel may be pinched by two magnets that are swallowed simultaneously and are in different loops of bowel. Hence, suspected magnet ingestion requires urgent evaluation. Radiographs of the neck and abdomen should be obtained to prove magnet ingestion,\textsuperscript{46,107} but radiographs cannot determine whether bowel wall is compressed between the magnets. Identification of magnets that appear to be stacked but slightly separated raises concern for bowel entrapment.

Management of swallowed magnets depends on the timing, location, type, and number of magnets. Because even single magnets have some risk, endoscopic removal is mandated if the magnets are within endoscopic reach. Magnets in the esophagus or stomach should be promptly removed via endoscopy.\textsuperscript{10,13,20} Single magnets passed beyond the stomach can generally be managed conservatively initially, but serial outpatient radiographs should be obtained to confirm that the

\textbf{Magnets}

Swallowed small magnets from toys and household items have become a serious health hazard in children. All such batteries may lead to rapid edema, thus making the catheter technique more difficult. In some localities rapid transfer of button battery ingestions to trauma centers or referral centers has resulted in much more rapid removal of the batteries.\textsuperscript{100,101} Once in the stomach, button batteries do not require removal. They may be monitored radiographically to demonstrate passage, with little risk for GI injury or heavy metal poisoning, even if the battery opens\textsuperscript{10,10,46,102} (see \textit{Table 39.3}).
The Patient in Distress

Pharyngeal or upper esophageal FBs can cause respiratory distress or respiratory arrest, usually in infants and the elderly. The Heimlich maneuver can be attempted in the prehospital setting or the ED when the situation is appropriate. The first intervention is to ensure an adequate airway, which can be obvious by the situation or may require laryngoscopy or other means of direct visualization. Forceps may be required to remove obstructions under direct vision. Because FBs may mimic multiple other pathologies, the approach to an acutely choking patient is challenging and every situation individual. Upper esophageal FBs can compress the trachea and cause stridor and respiratory compromise; these are indications for immediate removal.

ED Evaluation of FB Sensation in the Throat

FB impactions in the esophagus are usually straightforward, but patients may seek treatment in the ED because of a lump in the throat or difficulty swallowing with no apparent reason or history of FB ingestion. Such complaints require an examination and an investigation based on the clinical encounter and individual circumstances. Complete evaluation of these complaints is beyond the scope of this chapter, but initial modalities available to the clinician to evaluate these complaints are barium swallow, CT, and pharyngoscopy or laryngoscopy. The need for consultation is based on the clinical scenario.

If no cause is suspected by the history or examination, globus pharyngeus may be the causative factor. This may be associated with anxiety or a panic attack. The sensation of a painless
lump in the throat is called globus pharyngeus or globus hystericus. It has many causes other than FBs. Palpate, visualize, or review the anatomic structures in the area: the chin, laryngeal cartilage, cricothyroid cartilage, tracheal rings, sternum, and cricopharyngeal muscle.

FB sensation may be caused by infection, acid reflux, esophageal spasm, esophageal strictures, pill esophagitis, benign and malignant tumors, hiatal hernia, scleroderma, and many other causes. Globus sensations may also persist after an FB has been completely removed because of mucosal injury. Neurologic causes include botulism, myasthenia gravis, cerebrovascular accident, and amyotrophic lateral sclerosis. If the patient otherwise appears well and is able to drink liquids and keep hydrated, referral to a gastroenterologist as an outpatient is standard.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


16.RELATED


Nasogastric (NG) intubation (Videos 40.1–40.5) is commonly used to evaluate or treat bowel obstruction, ileus, or gastric hemorrhage, preoperatively or postoperatively, or to administer food or medication into the gastrointestinal (GI) tract. Patients with long-term feeding tube complications and those requiring replacement or other manipulation of tubes are frequently seen and treated in the emergency department (ED).

PROPERTIES OF NG AND FEEDING TUBES

Polypropylene is the material most commonly used for Levin and Salem sump NG tubes (Fig. 40.1), but it is too rigid for long-term use as a feeding tube. Polypropylene tubes are less likely to kink than others but are more capable of creating a false passage during placement. Latex (rubber) tubes are moderately firm, require greater lubrication for passage, are relatively thick walled, and induce a greater foreign body reaction than tubes made of other common materials. Latex, especially in latex balloons, deteriorates more rapidly than other materials.1

Foley catheters are primarily latex, although silicone Foley catheters are available for patients with latex allergies. Silicone tubes are thin walled, pliable, and nonreactive; however, the walls of silicone tubes are weaker and may rupture if fluid is introduced into a kinked tube. Polyurethane tubes are nonreactive and relatively durable. Rigidity varies from manufacturer to manufacturer, depending on the thickness of the tube. A stylet may aid in the passage of polyurethane and silicone tubes, but it increases rigidity and the potential for tissue dissection, especially with tubes that have a small distal end-bulb.1 Some feeding tubes have weights, which are usually made of tungsten and are nontoxic if released into the GI tract.

NG TUBE PLACEMENT

Indications and Contraindications

The simplest NG tube is the Levin tube, which has a single lumen and multiple distal “eyes.” The advantage of the Levin tube is its relatively large internal diameter (ID) in proportion to its external diameter. The theoretical disadvantage is that a Levin tube should not be left hooked up to suction after the initial contents of the stomach have been drained because the suction will cause the stomach to invaginate into the eyes of the tube, blocking future tube function and potentially causing injury to the stomach lining. Levin tubes are therefore rarely used in the ED. The Salem sump tube is preferred over the Levin tube for chronic use as a drainage device because it has a separate (blue-colored) channel that vents the distal main lumen to the atmosphere (Fig. 40.2). This vent helps prevent an excessive vacuum from forming at the tip of the tube. Note that both intermittent suction and wall unit vacuum can exceed the venting capacity of the second lumen, so the vacuum setting should be less than 120 mm Hg. The most commonly used size is 16 Fr, although larger and smaller sizes are available.

Review Box 40.1 Nasogastric tube placement: indications, contraindications, complications, and equipment.
The major indication for NG tube placement and suction is to aspirate the stomach contents in patients with gastric bleeding, gastric outlet or intestinal obstruction, gastric or bowel distention, prolonged ileus, or gastric, esophageal, or bowel perforation. Draining the dilated stomach of excessive contents lessens the chance of vomiting and possibly aspiration and provides marked relief of symptoms in patients with intestinal obstruction. The presence or anticipation of depressed mentation potentially justifies NG tube placement to protect the airway, especially if the stomach is full or vomiting is uncontrolled. Bag-valve-mask ventilation often distends the stomach with air, and a postintubation NG tube can improve ventilation, prevent vomiting, and increase patient comfort. Trauma patients may need an NG tube as part of the evaluation for GI injury or to decompress the stomach before surgery or peritoneal lavage. A radiopaque NG tube may help delineate transdiaphragmatic hernia of the stomach after trauma. A deviated NG tube is a nonspecific sign of traumatic aortic rupture. An NG tube may be used to instill air into the stomach for documentation of a suspected gastric perforation by enhancing visualization of free air under the diaphragm on an upright chest film. In patients unable to swallow, an NG tube may be passed to administer medications or oral contrast material for a computed tomography (CT) scan.

For the evaluation and treatment of upper GI bleeding, indications for NG tubes differ among clinicians, and practices vary widely. Clinical judgement prevails as the best arbitrator, but the true clinical value of an NG tube in patients with GI bleeding is probably less than traditionally promulgated. Although insertion of an NG tube may prompt earlier endoscopy, the procedure has not been demonstrated to improve clinical outcomes. NG aspiration can help localize bleeding in only a minority of patients with GI hemorrhage. Patients who vomit a significant amount of proven bloody material have had upper GI bleeding; they do not need an NG tube for diagnosis. Such patients are best evaluated by endoscopy, although when a history of bleeding is suspected, stomach aspiration may have a role in diagnosis to confirm the presence of blood. In patients without hematemesis, NG aspiration uncovers less than half the patients with upper GI hemorrhage and thus cannot be used to unequivocally rule out upper GI bleeding. Even a rapidly bleeding duodenal ulcer may not produce blood in the stomach. Some contend that emptying a markedly dilated stomach of blood and stomach contents may reduce the rate of bleeding, but data are lacking. Patients with presumptive upper GI bleeding, which includes those with melena, those younger than 50 years old, and those with a hematocrit lower than 30, need upper endoscopy.5,6 Patients with a bleeding pattern indicative of a Mallory-Weiss tear may need neither an NG tube nor endoscopy.

NG tubes can provide the earliest indication of high-volume esophageal or gastric bleeding. Although variceal rupture may potentially be caused by insertion of instruments into the esophagus, several studies suggest that passage of an NG tube is generally safe, even in patients with esophageal varices. Use of NG tubes for the evaluation and monitoring of upper GI bleeding should be judicious rather than universal.

Increasingly, the literature suggests that most other causes of vomiting are best controlled with medication. Postoperative ileus ends sooner and patients recover faster without an NG tube after various studied abdominal surgeries.7 NG tubes prolong the hospital stay, pain, and hyperamylasemia in those with mild to moderate pancreatitis.7,8 Many patients are better off without an NG tube from the point of view of safety, comfort, and speed of recovery.

Figure 40.1 Salem sump tube. This tube contains a second lumen that allows venting during continuous suction. 1, Pigtail extension (blue) of the air vent lumen. 2, Connector for attachment of the suction lumen to the vacuum line. 3, Gastric end with suction eyes. 4, Insertion depth markers.

Figure 40.2 Diagram of the Salem sump tube. A, General design. B, Diagram of the double-lumen principle for suction. (A and B, Courtesy the Argyle Division of Sherwood Medical, St. Louis, MO.)
In an awake and alert patient with a preserved gag reflex, passage of an NG tube has not been demonstrated to result in significant pulmonary aspiration, even when vomiting occurs during passage. NG tubes are, however, contraindicated in patients with a special predisposition to injury from placement of the tube. Patients with facial fractures who have sustained an injury to the cribriform plate may suffer intracranial penetration with a blindly placed nasal tube.\(^9\) Severe coagulopathy is a relative contraindication to passage of an NG tube. In patients with a coagulopathy or significant facial or head trauma, passing the desired tube through the mouth may be a better alternative (Fig. 40.3). Patients who have esophageal strictures or a history of alkali ingestion may suffer esophageal perforation. Gagging will decrease venous return and increase cervical and intracranial venous pressure. Comatose patients may vomit during or after NG tube placement. Indwelling NG tubes predispose patients to pulmonary aspiration because of tube-induced hypersalivation, depressed cough reflex, or mechanical or physiologic impairment of the glottis.\(^7\) Aspiration is also quite common with nasoenteral feedings in debilitated patients, hence the use of a gastrostomy feeding tube for this condition. An NG tube should be avoided when possible in patients who have undergone gastric bypass surgery or lap banding procedures.

Extended irrigation of the stomach with water in a patient with upper GI hemorrhage can lower serum potassium levels.\(^10\) and animal studies suggest that cold water lavage can cause rather than control bleeding.\(^11,12\) No study has shown irrigation to be effective in the control of bleeding.\(^13,14\) and vigorous lavage with cold water may lower the body temperature. Erythromycin has been demonstrated to be effective in clearing the stomach for endoscopy,\(^15,16\) and its success is better substantiated than irrigation.

If the bib area is covered with a towel and a supply of tissues or washcloths is available. For standard NG tube placement, a piston or bulb syringe (with a catheter slip-tip) should be available. NG feeding tubes should have a compatible 50-mL or 60-mL syringe (some are Luer compatible and others are slip-tip compatible).

Tape torn into 4-inch strips or a commercial NG tube holder (e.g., Suction Tube Attachment Device, Hollister, Libertyville, IL) should be handy for securing the tube after placement. Cotton-tipped applicators and tincture of benzoin may be helpful in securing the tube to the nose if the skin is greasy. Make sure that the feeding tube is designed for duodenal passage if this is desired; such tubes are usually longer than regular feeding tubes.

### Procedure

Most NG tubes are placed at the bedside in an awake patient. Explain the procedure to the patient. Written informed consent is not standard. If the patient is alert, raise the head of the bed so that the patient is upright. Place a towel over the patient’s chest to protect the gown, and place an emesis basin on the patient’s lap.\(^17\) Position the tube (typically a 16- or 18-Fr sump) so that the insertion distance can be estimated, and mark the distance with tape or by noting the markers printed on the proximal end of the tube. A simple method is to measure the tube from the xiphoid to the earlobe and then to the tip of the nose. Then add 15 cm (6 inches) to this number (Fig. 40.4, step 5).\(^18\) It is a common error to fail to estimate the proper length of tube before passage, which can result in the tip of the tube positioned in the esophagus or coiled excessively in the stomach. Check the nares for obstruction. Assess patency by direct visualization, by gentle digital nasal examination, and by having the patient sniff whilst first one and then the other nostril is occluded. Pass the tube down the more patent naris.

### Relief of Discomfort

Ameliorate the pain and gagging associated with tube placement by using vasoconstrictors, topical anesthetics, and antiemetics. Because patients rate NG tube placement as very painful, one of the most painful procedures performed in the ED, use these adjuncts whenever time and the clinical situation permit (Fig. 40.5). Spray topical vasoconstrictors, such as phenylephrine (0.5% Neo-Synephrine [Hospira Inc., Lake Forest, IL]) or oxymetazoline (0.05% Afrin [Bayer Healthcare LLC, Whippany, NJ]), into both nares at first in case one side proves to be problematic (see Fig. 40.4, step 2). The nares, nasopharynx, and oropharynx should all be anesthetized at least 5 minutes before the procedure. Gagging is reduced if the pharynx is anesthetized as well as the nose. Combinations of tetracaine, butyl aminobenzoate, and benzocaine (Cetacaine [Cetylene Inc., Pennsauken, NJ]), nebulized or atomized (spray cans or bottles) lidocaine (4% or 10%), and lidocaine gels (2%) are most commonly used. Lidocaine preparations of 10% are most useful. Lidocaine may be nebulized and delivered by face mask with the equipment used to administer bronchodilators to asthmatics (see Fig. 40.4, step 3). This method has been found to be superior to lidocaine spray for reducing gagging and vomiting and increasing the chance of successful passage.\(^19,20\) Cullen and coworkers concluded that nebulized nasal and pharyngeal lidocaine (4 mL of a 10% solution) reduces the discomfort associated with passage of an NG tube better than placebo and without any lidocaine-related toxicity.\(^21\)
**Nasogastric Tube Placement**

1. Choose the most patent naris by visual inspection and the sniff test. Alternatively, insert a gloved finger into each nostril to assess patency.

2. Apply a topical vasoconstrictor, such as phenylephrine or oxymetazoline.

3. Anesthetize the naris, nasopharynx, and oropharynx at least 5 minutes before the procedure. Nebulized lidocaine is ideal because it reduces both nasal and pharyngeal discomfort.

4. Apply 2% viscous lidocaine along the floor of the nasal cavity, and allow it to drip into the nasopharynx and be swallowed.

5. Slowly insert the nasogastric tube along the floor of the nostril under direct vision until it passes into the oropharynx.

6. Measure the distance from the tip of the nose to the earlobe to the xiphoid process. Add another 15 cm to this distance. Note the total distance on the nasogastric tube markers.

7. While the patient sips water from a straw, rapidly pass the tube to the predetermined depth. Coordinate tube advancement with the swallowing mechanism to promote easy passage.

8. Assess tube placement with air insufflation and aspiration.

9. Attach the tube to intermittent wall suction.

10. Secure the tube to the nose with tape.

*Figure 40.4* Nasogastric tube placement.
After a topical vasoconstrictor and anesthetic are administered, lubricate the tube with viscous lidocaine or lidocaine jelly. Lubrication and anesthesia of the nares can be facilitated by using a syringe (without needle) filled with 5 mL of an anesthetic lubricant, such as 2% lidocaine gel (see Fig. 40.4, step 4). Simply putting anesthetic jelly on the tube just before insertion will not provide any anesthesia. Topical anesthetics are generally quite safe, but pay attention to the total dose of anesthetic administered to avoid toxicity. Note that each milliliter of a 10% lidocaine solution contains 100 mg of lidocaine and can be absorbed systemically. In rare cases topical benzocaine has caused methemoglobinemia, even with the relatively small amounts used for endoscopy.

Although no standard exists and supportive data are sparse, some clinicians administer ondansetron (Zofran [GlaxoSmithKline, Research Triangle Park, NC], 4 mg) or metoclopramide (Reglan [Baxter Healthcare Corp., Deerfield, IL], 10 mg) intravenously 5 minutes before passage of an NG tube to potentially reduce nausea and gagging and, secondarily, to improve the pain and prolongation of the procedure that gagging engenders. Metoclopramide may have additional benefits on the discomfort associated with NG tube insertion, unrelated to its antinausea effects. Ondansetron is preferred for nausea because metoclopramide can cause agitation or facial and tongue spasm, but these effects can be reversed rapidly with the administration of 25 mg diphenhydramine intravenously.

Under direct vision, not blind forcing, insert the tube gently into the naris along the floor of the nose under the inferior turbinate and NOT upward toward the nasal bridge (Fig. 40.6; also see Fig. 40.4, step 6). If mild resistance is felt in the posterior nasopharynx, apply gentle pressure to overcome this resistance. If significant resistance is encountered, it is better to try the other nostril because bleeding or dissection into retropharyngeal tissue may occur if force is used. Once the tube passes into the oropharynx, pause to help the patient regain composure and enhance the chance for cooperation with the rest of the procedure.

If the patient is alert and cooperative, a common option is to ask the patient to sip water from a straw and swallow while you advance the tube into and down the esophagus (see Fig. 40.4, step 7). This may facilitate passage of the tube. Once the tube is in the nasopharynx, flex the patient’s neck to direct the tube into the esophagus rather than the trachea. In an intubated, anesthetized, or paralyzed patient, two stacked positive pressure breaths lasting 1 to 2 seconds each via a face mask provide similar relaxation of the upper esophageal sphincter for a brief 4- to 5-second window, and this markedly increases successful tube placement.

Some gagging during the procedure is common and is not an indication to halt attempts at passage. Withdraw the tube promptly into the oropharynx if the patient has excessive choking, gagging, coughing, a change in voice, or condensation appears on the inner surface of the tube because this indicates the possibility of passage of the tube into the trachea. With the mouth open, inspect the tube in the posterior pharynx to detect coiling or respiratory passage. If the tube is lateral to the midline, this suggests correct position in the esophagus. Once the tube is in the esophagus, advance it rapidly to the previously determined depth. Passing the tube slowly prolongs discomfort and may precipitate more gagging.

**Confirmation of Tube Placement**

Before the NG tube is secured, confirm successful placement by nonradiographic means or by auscultation. Use more than
one method when in doubt because all methods of confirmation have some possibility of error. Radiographic evaluation is the most definitive way to confirm the position of an NG tube, but it is not standard to routinely obtain radiographic confirmation. Radiographs may be more useful in the obtunded patient. Alert and cooperative patients will feel when a tube is entering the trachea or lungs and will be very symptomatic and therefore obviously alert the clinician. Patients who are struggling or have altered sensorium may not give any sign when the tube passes into the lungs, and therefore careful confirmation of placement, usually with radiography, is critical for these individuals. Ultrasound, when successful, is also highly reliable (see Ultrasound Box 40.1).27

A quick, simple, but unreliable method of confirmation is to insufflate air into the NG tube and auscultate for a rush of air over the stomach (see Fig. 40.4, step 8). If increased pressure is required to instill the air or if no sounds are heard, the tube may be malpositioned or kinked. Suspect an esophageal location if the patient immediately burps on insufflation. Left pulmonary tubes and tubes which have advanced past the stomach into the small bowel often sound so similar to properly placed tubes that misplacements are not detected.3

Aspiration of stomach contents, especially if pH-tested, is more reliable and can be performed if the tube position is in question. If the pH is less than 4, there is an approximately 95% chance that the tube is in the stomach and nonrespiratory placement is almost guaranteed.28 Although aspirated fluid can occasionally be obtained from the lung or pleural space, the pH should be 5.5 or higher.29 Approximately 4% of correctly placed tubes have aspirates with a pH higher than 5.5. Causes include duodenal reflux, antacids, H2 blockers, or recent instillation of formula or medications.29

If awake and cooperative, ask the patient to talk. If the patient cannot speak, suspect respiratory placement. Note that with small-bore tubes, patients may still be able to speak despite tracheal placement.19

Once correct tube position is tentatively confirmed, secure the tube. If the patient requires abdominal or chest radiographs for other diagnostic purposes, place the NG tube before obtaining the films. An NG tube deviated to the right may occasionally be seen in patients with traumatic rupture of the aorta, but this is not a reliable indicator.

Securing the Tube

The NG tube is generally secured to the patient with tape attached to both the tube and nose (see Fig. 40.4, step 10). Excellent purpose-made tape NG holders exist. A butterfly bandage (or tape on each side of the nose) that coils around the NG tube is a typical approach. Clean and prepare the nose and tube with tincture of benzoin if possible. If the tube lets go or requires repositioning, replace both the tape and the tincture of benzoin. It is wise to also secure the tube to the patient’s gown so that a tug on the tube will encounter this resistance before pulling on the tape securing the tube to the patient’s nose. A rubber band tied around the tube with a slipknot (Fig. 40.7) and pinned to the gown near the patient’s shoulder is effective. It is critical to ensure that the tube is secured in such a way that it does not press on the medial or lateral aspect of the nostril. Necrosis or bleeding can result if a tube is not secured correctly.

When a Salem sump is used, keep the blue pigtail above the level of the fluid in the patient’s stomach or the stomach contents may leak back through the vent lumen. If a patient needs to ambulate with a sump tube in place, fit the blue pigtail into the plastic connector at the end of the suction lumen. This creates a closed loop that should not leak.

Placement Issues

If the patient is intubated, deflate the balloon of the endotracheal (ET) tube briefly to allow passage of the NG tube. In a nonintubated unconscious patient, the NG tube is easily misplaced into the pulmonary tree. This complication may be missed during the procedure in cases in which the gag and cough reflexes are suppressed and the patient cannot talk. In addition, the absence of swallowing may prevent successful passage of the tube. In comatose or ventilated patients, several techniques have been studied to aid passage of the NG tube. Using Magill forceps to grip the tube or stenting the end of the tube with a gum bougie passed into the most distal tube eye can help stiffen and control the tube. The combination and sequence of techniques used will depend on the clinician’s expertise, the equipment available, and the level of sedation.

If less traumatic strategies fail, an ET tube can be prepared with an ID that is slightly larger than the external diameter of the NG tube. Slit it along its lesser curvature from the proximal end to a point 3 cm from its distal end. Pass the NG tube through a naso into the oropharynx. Visualize the tip of the tube with a laryngoscope, grasp it with Magill forceps, and pull it out of the mouth. Pass the slit ET tube (generally 8 mm in ID) through the mouth into the esophagus.31 Alternatively, pass a 7-mm-ID slit ET tube directly through the nose into the esophagus.32 Passage into the esophagus is facilitated by the stiffness of the larger ET tube and does not require active swallowing. Thread the tip of the NG tube into the ET tube and advance it into the stomach (Fig. 40.8). Remove the slit ET tube from the esophagus. When the distal part of the tube is visible, slit the unslit 3-cm distal part with scissors. Remove the ET tube and the NG tube will remain in place.33 Advance any slack tubing with forceps or pull it back nasally, depending on the final depth required for the NG tube. The technique can also be performed by passing the slit ET tube nasally, which saves the trouble of orally advancing or nasally retracting any slack in the tubing.33 Make sure that the NG tube can pass into the 7-mm-ID tube and that the ET tube is well lubricated inside and out.
Ultrasound has emerged as a new modality to assist in placement verification of nasogastric tubes, both during the procedure and as a confirmatory test after the procedure.

Place the high-frequency transducer on the patient’s neck transversely between the cricoid cartilage and the suprasternal notch, slightly off to the side with the orientation button to the patients’ left. (Fig. 40.US1). Obtaining a clear view of the esophagus sometimes involves slight angling of the probe and occasionally even benefits from some minor tracheal manipulation, usually leftward displacement. With these maneuvers, it is probably possible to visualize the esophagus over 90% of the time. (Fig. 40.US2)

When the nasogastric tube passes the probe image level, the esophagus acquires a bright echogenic density with posterior shadowing (Fig. 40.US3). The nasogastric tube’s strong interaction with the ultrasound may obscure clear visualization of the esophageal wall. A brighter echogenic density is obtained if there is a metal guidewire in the tube, as is common in feeding tubes but not in Salem sump tubes.

If 20 cm of tube has been passed but the esophageal appearance hasn’t changed, the tube is likely in the trachea or coiled in the mouth and failing to pass. Sometimes the empty esophagus isn’t easily seen, and then it becomes even more important to distinguish the course of the tube and ensure that the tube is in an extratracheal soft tissue

---

**ULTRASOUND BOX 40.1: Assistance in Nasogastric Tube Placement**

by Christine Butts, MD

---

**Figure 40.US1** Proper positioning of the high-frequency transducer between the cricoid and the suprasternal notch.

**Figure 40.US2** Angling of the probe and gentle tracheal (*) manipulation facilitates visualization of the collapsed esophagus (arrow).

**Figure 40.US3** When the nasogastric tube passes the probe, the esophagus becomes brightly echogenic and creates posterior shadowing (arrow). In this image, an endotracheal tube (asterisk) can be visualized in the trachea.

**Figure 40.US4** Lateral cross-sectional views may help confirm that the nasogastric tube is clearly outside of the trachea.
times easier to pass nasally than an NG tube, and then the lubricated NG tube can be passed through it. In addition, it affords some protection to the nasal mucosa if multiple attempts at passing the NG tube are necessary or if it is particularly important to minimize bleeding or trauma. Cooling an NG tube increases its rigidity, and coiling it can increase the curvature of the tube, both of which may help pass the tube.

Alternatively, the NG tube and larynx can be visualized with a laryngoscope, endoscope, or the GlideScope (Verathon, Bothell, WA) (see Fig. 40.9). Under laryngeal visualization, manipulation and lifting of the jaw and neck flexion can help align the tube for passage under direct vision into the esophagus.

Ultimately, if all other methods fail, place a flexible fiberoptic bronchoscope or esophagoscope into and through the esophagus under direct visualization. Thread a guidewire into the stomach. Place the NG tube over the guidewire into the stomach and then remove the guidewire.

Complications

Complications of standard NG tube placement are similar to the problems noted with placement of an NG feeding tube. The complications related to tube misplacement are discussed in that section. In addition, clinicians placing an NG tube in a patient with neck injuries should be cautious of potentiating cervical spine injuries with excessive motion during passage (especially in association with coughing and gagging in an awake patient). Furthermore, passage of an NG tube in an awake patient with a penetrating neck wound may exacerbate hemorrhage should coughing or gagging result. Particularly serious forms of tube misplacement are pulmonary placement (Fig. 40.10) and intracranial placement (Fig. 40.11). In confusing cases, a CT scan will clarify most misplacement issues (Figs. 40.12 and 40.13). NG tubes, when in place for prolonged periods, are a common cause of innocuous gastric bleeding and gastric erosions.
Tension gastrothorax can develop in patients with an intrathoracic stomach. It can occupy much of the left hemithorax and thus displace the heart and lungs and cause a clinical syndrome identical to tension pneumothorax. Even though successful passage of an NG tube will relieve tension gastrothorax, the high pressures of a tension gastrothorax often develop because torsion of the stomach in the chest prevents egress of air; such torsion may also prevent ingress of the therapeutic NG tube. The condition is rare enough that further emergency therapy is based on case reports rather than substantial series. Relief of tension gastrothorax has been accomplished by transthoracic puncture of the stomach with a 16-gauge catheter-over-the-needle device inserted into the second intercostal space in the midclavicular line and then removing the needle. The catheter is left in place attached to intravenous tubing with the distal end under a water seal [38]. A single 16-gauge puncture of the stomach is unlikely to leak and cause pleuritis; such punctures have long been used for placement of percutaneous endoscopic gastrostomy (PEG) tubes. Inserting a chest tube into the stomach is not advisable because gastric fluid may leak into the pleural space. Once the tension on the stomach is relieved, it may be possible to pass the NG tube to prevent reoccurrence of the problem. The stomach, no longer tense and wedged in the chest, can twist to allow the tube to pass. Surgical correction of the condition permitting, intrathoracic herniation of the stomach is the definitive treatment to prevent recurrence of tension gastrothorax.
Feeding Tube Site

Three major classes of enteral feeding tubes are in common use and are classified according to the site of insertion. Tubes can enter through the nares (a cervical ostomy) or the abdomen (an abdominal ostomy). Enteral tubes are often categorized by the location of the tip of the tube. Tubes may terminate primarily in the stomach, such as a gastrostomy (G) or PEG tube (Fig. 40.14). They may terminate in the jejunum (J tube) (Fig. 40.15) or in both the stomach and the small intestine (a PEG-J tube). To confuse the issue, some tubes enter the stomach and terminate in the stomach (G tube) or in the proximal part of the small bowel (J tube), whereas others enter the GI tract directly through the wall of the small bowel (J tube). Practically speaking, almost all gastric tubes are PEG tubes. They are placed endoscopically with local anesthesia and without a surgical incision. J tubes that enter the small bowel directly are inserted surgically under general anesthesia, require a surgical incision, and result in a surgical scar at the insertion site. Gastric feeding results in better digestion than intestinal feeding does. Jejunal feeding reduces reflux and aspiration. Normally, approximately 20% of the gastric antral contents pass into the duodenum, with 80% refluxing back into the body of the stomach for further mixing, so proximal duodenal feeding is of limited benefit in preventing aspiration. If the feeding tube is placed in the antrum of the stomach or in the small bowel, enteral feeding solution passing into the small bowel may not be tolerated and can result in diarrhea and paradoxical decreased nutrition.

Procedure

The clinician should explain the procedure to the patient before passage of the tube. Replacement of a nasoenteric feeding tube requires greater time and effort if the patient is uncooperative or has a physically obstructing lesion. It is generally advisable to restrain the hands of demented, impaired, or otherwise uncooperative patients. Prepare the nares before passage of the tube using the procedure for primary NG tube placement. If a feeding tube stylet is used, lubricate and insert it into the feeding tube before introducing it into the naris. Tube stylets can be lubricated with water-soluble jelly. If using the Dobhoff, Entri-Flex (Biosearch [Biosearch, Branchburg, NJ]), or another tube with preapplied lubricant, you may need to activate the lubricant with a 5-mL flush of water. Never allow the stylet to protrude beyond the end of the feeding tube because these stiff, small-diameter wires have the capacity to scratch the esophagus and allow the creation of a false passage. The stylet may lock into position on the tube at the proximal end and should be properly secured.

When the patient is uncooperative or cannot drink, introduce 5 to 15 mL of water into the mouth or into the proximal end of the feeding tube with a syringe; this may induce swallowing and facilitate passage of the tube. Although the patient may not swallow for several minutes, wait for swallowing because this may mean the difference between a coiled or pulmonary tube placement and successful passage.

Confirmation of Placement

Auscultatory confirmation of tube placement can be misleading, so confirm proper placement of the tube with a radiograph before feeding. However, radiographic confirmation of tube placement should always be obtained. A nasoenteric tube that is placed correctly may exit through the nose, mouth, or both. The correct placement is determined by the following criteria:

1. Auscultation
2. Radiographic examination
3. Proper advancement
4. Incorrect placement

REPLACEMENT OF NASOENTERIC FEEDING TUBES

Indications and Contraindications

The most common indication for replacement of a feeding tube in the ED is unintentional removal of a preexisting feeding tube. In one prospective study, 38% of tubes were removed unintentionally. Although some of these tubes had fallen out or had been coughed out, more than half were pulled out by the patient. Tube rupture, deterioration, or clogging may also necessitate replacement. Management of a clogged or nonirrigating feeding tube is discussed in the later section on clogged feeding tubes.

Figure 40.11 Anteroposterior and lateral skull radiographs demonstrate intracranial insertion of a nasogastric tube in a patient with multiple skull fractures. (From Johnson JC: Letter to the editor: back to basics for morbidity-free nasogastric intubation, JACEP 8:289, 1979.)
tip in the duodenum, or it may occupy a position overlying the right side of the abdomen yet not have entered the duodenum. A contrast-enhanced study is necessary to ascertain duodenal position when pulmonary placement has been ruled out.44,45

Examine the radiograph also for the presence of mediastinal air or pneumothorax, which may suggest pulmonary or esophageal puncture. An esophageal puncture should be evaluated with endoscopy and may require surgery, depending on the size of the rent.

The end-bulb of many nasoduodenal tubes will pass into the duodenum after positioning the patient in the right decubitus position for an hour. Some researchers recommend pretreatment with metoclopramide to enhance gastric emptying.46–48 One investigator found that metoclopramide enhances duodenal passage of nasogastrically placed feeding tubes in diabetic, but not nondiabetic, patients.44 Gastric antral motility in diabetic patients is often impaired; metoclopramide helps restore normal synchronized activity in these patients but has little effect on emptying in those with normal antral function. The usual dose of metoclopramide is 10 mg, administered intravenously. In addition, 3 mg/kg of erythromycin lactobionate given intravenously over a 1-hour period works similarly and may be effective even if metoclopramide fails.49 Endoscopy or fluoroscopy may be necessary if positioning and metoclopramide are not successful.

Complications

Pulmonary intubation is an uncommon but well-known and potentially fatal complication of insertion of nasal feeding tubes (see Fig. 40.10). Coughing and respiratory distress are the most common symptoms of respiratory passage of an NG tube, but there may be relatively few apparent symptoms in a demented or comatose patient.50 Decreased mentation and an absent cough reflex are predisposing factors for unrecognized
Misplacement is far more likely in patients with ET or tracheostomy tubes, conscious or not, perhaps because tracheal stimulation by a misplaced tube is less likely to be appreciated or communicated. A small end-bulb (e.g., 2.7 mm in diameter) can slip past a tracheal high-volume, low-pressure cuff and easily pass to the periphery of the lung. Pneumothorax may result when an NG tube dissects into or is withdrawn from the pulmonary parenchyma. Bloody aspirate from a tube should heighten awareness of possible tissue damage.

A clogged or nonfunctional NG tube may be difficult to remove. Fluoroscopy may allow careful insertion of a guidewire or stylet into an in situ tube to facilitate removal. Fluoroscopy may also identify the mechanical problem interfering with removal of the tube. Segments bent double are probably the most common cause. Knots, though uncommon, do occur. Do not use excessive force to remove an NG tube because serious injury to the patient may result.

Premature removal of an NG tube is the most frequent complication of the use of feeding tubes. For long-term use

---

**Figure 40.14** Placement of a percutaneous endoscopic gastrostomy tube. A, Under conscious sedation, pass a lighted endoscope into the stomach. Indent the skin with a finger to determine the optimal puncture site where the stomach and abdominal wall are closest, with no bowel between. B, Fill a syringe with saline and advance it percutaneously to the selected entry point until the tip of the needle is seen entering from the gastric lumen through the endoscope. If air is aspirated and no needle tip is seen, the needle is in the bowel, not the stomach. C, Push and pull the scope/snare/feeding tube combination into position (arrows show direction of travel). D, Pull the head of the feeding tube into contact with the gastric mucosa. E, Use an external bolster or crossbar to keep the tube snug against the skin and gastric wall, but not so tight that it causes ischemia of the intervening tissue.

**Figure 40.15** Formation of a Witzel tunnel and final permanent placement of a jejunal catheter. (From Wiedeman JE, Smith VC: Use of the Hickman catheter for jejunal feedings in children, *Surg Gynecol Obstet* 162:69, 1986.)
Gastrointestinal Procedures

SECTION VII

Gastrointestinal Procedures

840

SECTION VII

Gastrointestinal Procedures

or frequent removal of an NG tube by a patient, a gastric feeding tube will probably be inserted. To help prevent removal by an uncooperative patient, secure the NG tube to a loop anchor passed in the same naris. The anchor works by aversive stimulation of the soft palate and nose, with distraction from the NG tube rather than by mechanical stabilization of the tube.

Sax and Bower recommend a technique for creating a separate NG-tube anchor. Cut a soft, weighted nasoenteric tube approximately 12 inches from the top. Pass a heavy (2-0) silk suture through the tube to exit the side hole. Insert the guidewire with care because it must not protrude from the inserted end. Sedate the patient if uncooperative. Insert the tube through the anesthetized naris into the nasopharynx, grasp it with Magill forceps, and pull to remove it through the mouth (Fig. 40.16A). Trim excess tubing without cutting the silk suture. Make a closed loop by tying the silk suture in front of the nose. Leave the loop long enough so that it does not apply continuous pressure to the nose or palate while at rest. Pass the nasal feeding tube through the same nostril and secure it to the loop (see Fig. 40.16B). This anchor is simpler to construct and more comfortable than anchors that pass through the opposite naris.

Complications of properly placed nasoenteric tubes include nasopharyngeal erosion, esophageal reflux, tracheoesophageal fistula, gagging, rupture of esophageal varices, and otitis media. One survey of nasogastrically fed patients found that the most distressing features of having an NG tube for feeding were deprivation of tasting, drinking, and chewing of food; soreness of the nose; rhinitis; esophagitis; mouth breathing; and the sight of other patients who were eating.

Checking feeding tolerance is difficult with small-gauge feeding tubes. Aspiration of tubes to check for residuals is not recommended with tubes 9 Fr in size or smaller. Aspiration is likely to clog the tubes because they collapse under pressure and relatively small particles can occlude the tube. For the same reasons, the residual is likely to be inaccurate.

**Patient or Nursing Instructions**

To maintain patency of the catheter, small tubes should be flushed with 20 to 30 mL of tap water at least two to three times daily and after the administration of medication. Water is a more effective irrigant than cranberry juice. Medications should be in liquid form or completely dissolved or they may clog the tube. Methods of dealing with a clogged tube are discussed subsequently. The tube should be anchored to the nose and face in such a way that it is not in contact with the skin at the nasal opening. This reduces tube discomfort and prevents necrosis of the alae, nares, and distal septum. Patients who exhibit a tendency to pull on their tubes need adequate restraints. Patients receiving tube feedings should have their head elevated to at least 30 degrees above the horizontal.

**PHARYNGOSTOMY AND ESOPHAGOSTOMY FEEDING TUBES**

Cervical pharyngostomy and cervical esophagostomy have both been developed relatively recently. Cervical esophagostomies are generally performed at the time of cervical or maxillofacial operations. Malignant growths in the proximal part of the esophagus, head, or neck are the primary indications for esophagostomy. Cervical esophagostomies may eventually evolve into a permanent sinus, thereby allowing the feeding tube to be removed between meals. Such tubes are unlikely to be replaced in the ED, but the concept of these feeding tubes is illustrated in Fig. 40.17. Complications of pharyngostomy and esophagostomy include local soft tissue irritation, accidental extubation because of excessive length of the external tube, pulmonary aspiration from vomiting, arterial erosion with exsanguination, and esophagitis or stricture of the esophagus from reflux.

**GASTROENTEROSTOMY AND JEJUNOSTOMY TUBES**

A nursing home patient with a nonfunctioning or displaced feeding tube represents a common ED scenario. The clinician cannot always determine the location of the original feeding tube by simply looking at a patient who arrives in the ED for replacement of the tube. Nevertheless, the emergency clinician should attempt to ensure that the terminal end of a replaced...
Some type of tube must be placed to stent the stoma, or the stoma will quickly close (in a matter of hours) and the patient might require a more complicated procedure to regain access. The only real concern of placing a gastric tube into the jejunum is that the balloon will produce intestinal obstruction if it is fully inflated.

If the tube is nonfunctioning yet still in place, the clinician must make a judgment regarding the risk versus benefit of removal and replacement, versus an attempt at unclogging the tube (see the subsequent discussion on unclogging). The major concern is that a new tube may be misplaced (i.e., into the peritoneal cavity). If it appears that a skin incision was used to place the tube, it is unlikely that the patient has an easily removable tube. If the patient has signs of a complication (e.g., infection, ileus, intestinal obstruction), surgical consultation is warranted. Note that a migrated tube, with the balloon or tube obstructing the gastric outlet, is a common cause of gastric distention, persistent vomiting, or signs of intestinal obstruction. This is easily remedied (see procedural description in section on Complications and Fig. 40.22).

Most PEG tubes do not have sutures joining the stomach with the abdominal wall, so there is potential for a replaced tube to end up in the peritoneal cavity. Adhesions, however, usually keep the stomach appropriately positioned, but only after the tract has matured. Nonoperative tube replacement techniques are safe only through an established tract between the skin and the bowel. Catheter replacement should not be attempted in the immediate postoperative period. A simple gastrostomy takes approximately a week to form a tract.61 A stable tract may take 2 or 3 weeks longer to form if healing is compromised.

Poor nutrition is the most common element compromising wound healing in patients requiring a feeding tube. A Witzel tunnel may take up to 3 weeks after the operation to mature sufficiently for safe nonoperative tube replacement (see Fig. 40.15).
in use to prevent leakage. Some tubes are placed with the aid of accompanying guidewires or stents.

The easiest tube to replace is one that has been removed in the ED or dislodged for only a few hours (Fig. 40.23). The stoma closes quite quickly, so replacement is best done as soon as possible. Gently probe the stoma site with a cotton-tipped applicator to determine patency and direction of the tract. In selected cases, a hemostat can gently dilate the opening to accept a replacement tube. Do all such attempts carefully to

**Figure 40.18** A, Without old records, the exact type and positioning of this nonfunctioning feeding tube are unknown. The operative scar on the abdominal wall suggests an implanted tube, not a simple gastric tube. B, Injection of contrast material before removal of the tube demonstrates the tip of the tube ending in the small bowel, not the stomach. C, If the tube has been removed and questions remain about the type of tube and circumstances of placement, a new tube is best placed under fluoroscopy with guidewire assistance.
avoid creating a false tract. Use local anesthetic around the stoma if exploration causes pain. Bleeding is common. After the tube is passed, restrain the hands to avoid removal of the tube by uncooperative patients.

**Removal of a Transabdominal Feeding Tube**

A feeding tube may need to be removed because it is irreversibly clogged, leaking, or broken; persistently develops kinks; is too large or too small; causes a hypersensitivity reaction; is associated with an abscess; or is not the appropriate length for feeding into the desired viscus. Before a new transabdominal feeding tube is inserted, remove the old tube. Most, but not all tubes, can be removed without endoscopy. Before attempting to remove the tube in place, it is imperative to know whether it is safe to remove. Standard de Pezzer or mushroom catheters that have been modified with bolsters or rings at the time of endoscopic or surgical insertion may no longer be safe to remove with traction. Tubes are occasionally secured with sutures or rigid internal bumpers or stays. It is rare, however, to encounter a tube that cannot be removed with traction/countertraction. Modest force may be required. Use a hand for countertraction, and be prepared for a pop and splattering of gastric contents (Fig. 40.24). This causes the tube and end mushroom to narrow, and the tube should come out easily. The inner crossbar, if present, may remain in the stomach when the rest of the feeding tube complex is removed by traction. The crossbar will pass in stool, and obstruction from it has yet to be reported in adults. In small children, obstruction is a possibility, and the crossbar should be removed by endoscopy.63,64

Recently placed feeding tubes may need to be left in until a tract has formed (1 to 3 weeks, depending on the procedure), even if the tube is nonfunctional. A simple Foley catheter G tube is the easiest to remove. Deflate the Foley balloon and the tube should slide right out. If the Foley balloon cannot be deflated, cut the tube to allow the balloon to deflate. Do not cut the catheter so close to the abdomen that it will be impossible to maintain a grip on it for removal by traction if the balloon still does not deflate. The balloon may also be punctured to cause it to deflate. To puncture a Foley balloon, apply traction on the catheter to draw the balloon up against the ostomy (Fig. 40.25). Using the taut feeding tube as a guide, pass a small-gauge needle along the tube to puncture the balloon. It may be necessary to try again on the other side of the catheter because the balloon may be inflated asymmetrically and contact with the needle may be established on one side and not the other. Be careful not to
track away from the ostomy into the patient’s abdominal wall or to cause separate punctures of the stomach. Allow a minute for the balloon to deflate before making another attempt at removal by traction. Large, nondeflating balloons should probably be punctured, whereas small balloons may be removed with traction.

If it is not possible to pull the inner bolster or mushroom out through the ostomy, cut the tube at the skin, push the remaining short stump into the stomach, and rely on later rectal passage. Although obstruction or impaction is infrequent, it can occur, and this alternative has the potential to be problematic in children or patients who have had previous impaction, potential for bowel obstruction, or stool-passing problems. Rigid or large internal mushrooms and bolsters, the very kind that cause the most difficulty with percutaneous removal, are also more likely to cause difficulty with rectal passage. In no case should a device be released into the gut with a long length of tubing attached. Remember that double-part tubes may have an additional length of tubing for duodenal or jejunal feeding that extends far past the inner bolster. Korula and Harma reported successful intestinal elimination of 63 of 64 G tubes that were cut at the skin entrance and advanced
G-Tube Replacement (With Foley Catheter)

1. To make a bolster, cut a 3-cm segment of tubing from the proximal end of another Foley catheter. (A red rubber catheter is used in this case). A bolster will prevent migration of the tube, which can cause gastric obstruction.

2. Cut a small hole (on each side) in the middle of the 3-cm segment.

3. Insert a hemostat through the holes in the new bolster and grasp the feeding tube.

4. Pull the feeding tube through the bolster. It should be placed 1 cm above the external abdominal skin.

5. Identify and inspect the stoma. Surrounding scars may suggest that the tube is in fact an implanted jejunal tube, not a simple G tube.

6. Lubricate the tube and gently pass it into the stoma.

7. Inflate the balloon with saline once it is in proper position.

8. Advance the bolster so that it is 1 cm above the skin.

9. Confirm proper tube placement with auscultation and aspiration. A contrast-enhanced radiograph may be also be obtained.

10. Remember that Foley catheters are not ideal feeding tubes but are useful in temporarily maintaining stoma patency. Disintegration of the catheter latex may occur in a few months.

Figure 40.21 G-tube replacement (with a Foley catheter). Foley catheters are not ideal feeding tubes but can be useful temporarily to maintain stoma patency.
the tape places on the skin as the tube pulls on the tape can lead to skin damage. Strong adhesive tape can also damage the skin on removal. Tape that is not sufficiently adhesive can let go, particularly if it gets wet. Because tape is less durable, home or nursing home caregivers who are less skilled may retape the tube (Fig. 40.26). If the tape is replaced at home, it may be placed under too much tension and cause thinning of the abdominal wall at the stoma. If too much slack is allowed, the tube may be pulled in. Strong tape can also damage the tube during tape changes if it adheres too strongly to the tube. Special adhesive devices or bolsters are preferred.

**Verification of Placement**

There is no universally agreed standard with regard to performing a confirmatory contrast-enhanced study for all easily replaced feeding tubes. The editors of this text, and most clinicians, verify position routinely with a contrast-enhanced radiograph, whereas others use the clinical criteria outlined earlier. Radiographic verification is easily performed and interpreted in the ED. Routine use of postplacement contrast-enhanced radiography to confirm proper placement should be mandatory when the tube tract is immature (i.e., < 1-month duration), when passage of the replacement catheter has been difficult, when the clinician is unable to aspirate intestinal contents, or when the patient is unable to communicate any symptoms that might occur with tube misplacement. Peritoneal infusion of feeding solution can be fatal.

The position of the G tube may be checked by air insufflation and aspiration of gastric fluid, as is done with nasoenteric tubes. Air should enter the stomach without resistance and should produce immediate borborygmi. Gastric fluid should return with aspiration. It may be necessary to insert a small volume of water to obtain good return. Water pooling in soft tissue may be aspirated back through a misplaced catheter. Good tube placement is indicated when more fluid returns with aspiration than was originally placed into the catheter.

Correct tube placement is easy and readily verified radiographically with a small amount of contrast material passed into the tube (Fig. 40.27). Use a catheter-tipped syringe to introduce water-soluble contrast solution (e.g., diatrizoate meglumine–diatrizoate sodium [Gastrografin, Bracco Diagnostics Inc., Monroe Township, NJ]) into the tube. Barium is contraindicated because of the potential for peritoneal contamination. Inject 20 to 30 mL of water-soluble solution to document the intraluminal tube position. Take a supine abdominal film 1 to 2 minutes after instillation of dye to optimize visualization of the gut. Because the film must be obtained quickly, it is easiest to perform the injection in the radiology suite, followed by a radiograph in only a few minutes. If the contrast material does not flow freely into the tube, the procedure should be terminated immediately and the position of the tube questioned. With proper positioning, contrast material will outline the part of the gut containing the tube (e.g., stomach with a G tube). An irregular or rounded blotch with wispy edges or streamers suggests peritoneal leakage. In questionable cases, injection of dye can be performed under fluoroscopy.

**Complications**

Viscus puncture, viscus–abdominal wall separation, and creation of a false tract with subsequent misplacement of the tube are...
Figure 40.23 A, The feeding tube was dislodged at 11 PM and by 9 AM the stoma was too tight for easy replacement of the tube. It was accomplished under fluoroscopic guidance, always the best option in questionable cases. B, The stoma opening and direction of the tract can be investigated by gently probing the site and tract with a Q-tip; in this case, it easily entered the stomach. C, This tight stoma was carefully dilated with a hemostat under local anesthesia. A false passage can easily be created, and this area usually bleeds readily. D, To give a Foley catheter sufficient rigidity to aid in passage, the end of a Q-tip was inserted in the side port of the distal end of the catheter, and traction was applied to the catheter. E, If a de Pezzer catheter is used, an endotracheal tube stylet distends the flange for passage, and the tip reforms once in the stomach. F, This patient removed her recently replaced feeding tube, with the balloon inflated, while still in the emergency department awaiting transfer. This could have been avoided if her hands were restrained.
of fistulas and small bowel obstruction from adhesions or volvulus around the jejunostomy site. The most common complications of gastrostomy and gastroenterostomy are local skin erosion from leakage, wound infection, hemorrhage, and dislodgment of the tube. Peritonitis and aspiration are the most critical complications of gastrostomy feedings. Jejunalostomies are less prone to stomal leakage and cause less nausea, vomiting, bloating, and aspiration than do gastrostomies. Dislodgment of G and J tubes is most common in the 2 weeks after creation of the ostomy. Extrusion of the G tube is usually caused by excessive tension applied to the tube. Only gentle contact of the gastric and abdominal walls is desirable.

Figure 40.24 Gentle, firm traction using the flat part of the opposite hand for countertraction will remove most percutaneous endoscopic gastrostomy tubes, even those with internal mushroom bumpers. Modest force may be required; be prepared for a sudden pop and splattering of gastric contents.

Figure 40.25 A, If a Foley balloon will not deflate, use traction to bring the balloon against the abdominal wall. Gently pass a small needle along the course of the catheter while puncturing the balloon as many times as necessary. Wait a few minutes for the fluid to egress. B, Once the balloon is deflated, it can be withdrawn. Note the encrusted condition of this long-standing Foley catheter used as a percutaneous endoscopic gastrostomy tube. C, Occasionally, the wire from a central line kit can clear the inflation lumen and allow deflation. D, If the valve mechanism malfunctions, cut the catheter and attempt to drain the balloon by placing a needle in the inflation channel and flushing and withdrawing fluid.

Figure 40.26 Tape is best avoided to minimize skin maceration. If needed, this is a simple but less secure technique for securing a gastrostomy tube to the skin.
Uncooperative patients should be restrained and mittens are often particularly helpful. Sutures and large mushrooms or balloons do not prevent purposeful removal of the G tube by an uncooperative patient.

A small amount of drainage is to be expected at the tube entry site. Local leakage of gastric juices may macerate and irritate the skin, which can predispose the site to local infections and abscesses and encourage the development of small granulomas. Granulomas are particularly common in children. They can be treated with silver nitrate at the time of dressing changes. Any dressing used around the entry site of an enteral nutrition tube should absorb fluid and not encourage persistent moisture. An unusually large stoma may promote a leak. Although insertion of a larger tube or firmer traction on the tube might be transiently effective, these measures often result in further enlargement of the stoma. Rigid gastrostomies promote leakage by widening the stoma as they pivot. Insertion of a soft, pliant feeding tube through the widened stoma is often easy and allows later contraction of the stoma. If these techniques are ineffective, temporary removal of the feeding tube may allow the stoma to shrink. Large amounts of drainage around the stoma site may occur with high residual volumes. The residual should be checked and feedings withheld until residuals are less than 100 mL. Feeding residuals should be checked every 4 hours when a patient is receiving continuous-drip feeding.

Pneumoperitoneum after percutaneous gastrostomy is neither unusual nor dangerous. Benign pneumoperitoneum may be...
present as long as 5 weeks after PEG. Pneumatosis cystoides intestinals can occur through the defect in the bowel wall created for the enterostomy tube. Though often clinically insignificant, its occurrence suggests air under pressure in the small bowel. NG suction and diet change generally permit resolution of the problem. Catheter or feeding tube removal is not usually required.

Clinically significant pulmonary aspiration can occur with G-tube feeding. Methods of checking for silent pulmonary aspiration include assessing tracheal aspirates with a glucose oxidant reagent strip or placing methylene blue in the formula and monitoring tracheal aspirates for pigmentation.

A Foley balloon accidentally inflated in the small bowel or esophagus can lead to perforation or obstruction. Carefully inflate the balloon soon after it has entered the stomach to prevent perforation of the vescu. A G tube may migrate in the stomach and obstruct the gastric outlet. This complication is manifested clinically by vomiting and high residuals of feeding solution. Volvulus and jaundice may also occur as a result of balloon migration. To correct inward migration, first deflate the balloon. Pull the tube back just deep to the desired final position. Reinflate the balloon and pull it the rest of the way back into position. The external bolster should be repositioned or one made and positioned, or inward migration will probably reoccur. If the balloon is not deflated before repositioning, the bowel in which it is wedged may be drawn back with it, thereby preventing relief of the obstruction and possibly precipitating intussusception.

A gastrocolic fistula is usually manifested as copious diarrhea. Once it is confirmed, treatment consists of removal of the G tube. Later creation of a gastrostomy in a different location may be possible. The patient may require hospital admission for nutritional support and monitoring of fluid and electrolyte status.

An external bolster that is snugged down too tightly might result in a short stoma and embedding of the internal bolster in the abdominal wall. An abscess may result. Overly tight external bolster's should be loosened. The correct position is 1 cm from the external surface of the abdomen.

CLOGGED FEEDING TUBES

Clogging, leaking, fracture, and kinking are problems common to all feeding tubes. Clearing a clogged tube may be a temporary benefit but is rarely a long-term solution. Hence, whenever feasible, replace nonfunctioning or leaking tubes with new ones. Although it may be only a temporary solution, if the tube has a complex placement or the clinician is unsure how the tube is secured internally, it is prudent to attempt to unplug the tube rather than replace it. Large G tubes are the least likely to clog. G tubes at least 28 Fr in size can tolerate home blenderized foods and viscous feeding solutions. Isosmotic feeding solutions are tolerated by fairly narrow tubes and cost one-sixth of what elemental feedings cost. Isosmotic feedings will clog needle catheters. When tube lumens are 14 Fr or smaller, all pills and the contents of all capsules should be dissolved in water to prevent obstruction of the tube.

Kinking is a frequent cause of tube blockage during the immediate period after reinsertion. Withdrawing the tube a few centimeters usually relieves the kink and obstruction. A persistently recurring kink requires removal of the tube and insertion of a fresh tube.

Accumulated feeding solution or medication precipitates are very difficult to clean or remove. Milking a pliable tube backward may remove some of the cheesy precipitates. Guidewires or stylets may clear the proximal portion of a clogged tube lumen but are unsafe to use in subcutaneous areas of the lumen because they can puncture the tube and injure the patient or create a leak in the tube. Bionix Medical Technologies (Toledo, OH) supplies tube decloggers in two lengths and several diameters for clearing clogs in G and J tubes (Fig. 40.28). After selecting a declogger shorter than the feeding tube, insert it gently into the tube until the end of the screw hits the clog and then rotate it clockwise to bury the end in the clog. Next, slide the declogger up and down to break up and dislodge the clog. Insertion, rotation, and sliding might have to be repeated several times until the dislodged material can be flushed into the patient with saline or water. Gravity alone should be sufficient to allow fluid to pass through the feeding tube into the patient and is a better test of successful declogging than is passage of fluid with a syringe. Theoretically, the pointed screw end of the declogger can puncture the tube, or if a declogger that is too long is put in, it may puncture the bowel or stomach directly. According to the company, no cases have been reported.

Fogarty arterial embolectomy catheters can be used to unclog J and G tubes. Insert the soft tip of the Fogarty catheter into the feeding tube and advance it while monitoring the insertion...
Irrigation with carbonated beverages and high-pressure irrigation with small-volume syringes have also been recommended as techniques for unclogging feeding tubes. Although irrigation seems like a straightforward and simple solution, these techniques are generally ineffective; furthermore, the possibility exists for dangerous tube rupture with internal leakage. Broviac catheters are especially prone to tube aneurysms, which can rupture under pressure.71 Tubes unclogged by forceful irrigation or by deep luminal probing should be radiographed after injection of contrast material to check the integrity of the tube.

References are available at www.expertconsult.com
P H CUTOFF LEVEL FOR RELIABLE CONFIRMATION OF NASOGASTRIC TUBE PLACEMENT.


INTRODUCTION
Managing patients with acute gastrointestinal bleeding from gastrointestinal varices can be one of the most challenging scenarios in emergency medicine. These patients often have advanced liver disease and can arrive at the emergency department (ED) with massive hematemesis, airway compromise, hemodynamic instability, critical anemia, thrombocytopenia, and coagulopathy. Gastroesophageal varices are the fourth most common cause of upper gastrointestinal bleeding (UGIB) and account for almost 12% of cases (Fig. 41.1). In patients with cirrhosis, varices account for up to 80% of cases of UGIB. In patients with established gastric or esophageal varices, the annual incidence of acute hemorrhage ranges from 4% to 15%. Over the past 3 decades, advances in resuscitation, critical care, pharmacology, and endoscopy have significantly reduced the mortality rate associated with acute variceal bleeding. In fact, mortality rates in patients with acute variceal bleeding currently range from 15% to 20%. Despite advances in management, up to 20% of patients with acute variceal bleeding fail standard therapy. Rescue therapies for this group of patients are limited and include surgery, placement of a transjugular intrahepatic portosystemic shunt or a covered esophageal metal stent, or balloon tamponade. This chapter details the indications and contraindications for balloon tamponade in patients with acute variceal bleeding, the techniques for placement of the various devices, and the potential complications of this intervention. Although this procedure is rarely needed and placement in the ED is not considered a standard intervention, emergency physicians with knowledge of the technique can attempt to place these critical and potentially lifesaving devices.

BACKGROUND
In 1950, Sengstaken and Blakemore developed and described the use of a double-balloon device to control variceal hemorrhage. Since that time, the Sengstaken-Blakemore tube (Video 41.1) has become the most widely known balloon tamponade device. The Sengstaken-Blakemore tube has an esophageal and a gastric balloon, along with a gastric aspiration port that allows continuous suction of stomach contents (Fig. 41.2). In 1968, Edlich and colleagues, from the University of Minnesota, modified the Sengstaken-Blakemore tube by adding an esophageal aspiration port and increasing the capacity of the gastric balloon.

Currently, three balloon tamponade devices are commercially available: the Linton-Nachlas, the Sengstaken-Blakemore, and the Minnesota tubes. In contrast to the Sengstaken-Blakemore and Minnesota tubes, the Linton-Nachlas tube is a single-balloon device that consists of a gastric balloon and two ports (esophageal and gastric) for aspiration and lavage. Because placement of these tubes remains a relatively rare procedure, most hospitals stock only one type of device. Regardless of the type of device, success rates for the control of hemorrhage with balloon tamponade tubes range from 60% to 90%.

Balloon Tamponade of GE Varices

**Indications**
Unstable patients with massive variceal bleeding in the following scenarios:
- Endoscopy is not available
- Endoscopy is unsuccessful at controlling bleeding
- Consultant physicians are unavailable and vasoactive agents have failed to stop bleeding

**Contraindications**
- History of esophageal stricture
- Recent esophageal or gastric surgery

**Complications**
- Airway obstruction
- Esophageal rupture
- Aspiration pneumonitis
- Pain
- Ulceration of lips, mouth, tongue, or nares
- Esophageal and gastric mucosal erosions
- Arrhythmia
- Dislodgment of previous variceal bands

**Equipment**
- 60-mL syringe (catheter-tip)
- Nasal spray
- Viscous lidocaine
- Manometer
- Bulb inflator
- “Y” tube
- Tube clamps
- Scissors
Endoscopy by a gastroenterologist remains the “gold standard” for the diagnosis and treatment of acute variceal hemorrhage. Sclerotherapy and band ligation are the two endoscopic techniques used to control bleeding esophageal or gastric varices. Endoscopic band ligation has been shown to be superior to sclerotherapy in initially controlling hemorrhage and improving survival. In fact, endoscopic band ligation is considered the treatment of choice for esophageal varices.

Balloon tamponade is indicated in unstable patients with massive hemorrhage in whom endoscopy either cannot be performed or is unsuccessful in controlling the bleeding. Balloon tamponade can be used to stabilize a patient for transfer to another institution with the resources to continue care. It is important to recognize that balloon tamponade is only a temporizing measure. Even though success rates in controlling the initial hemorrhage are high, up to 50% of patients rebleed when the device is deflated. Although rebleeding rates can be reduced with the concomitant use of vasoactive agents, arrangements must be made for more definitive control of varices in patients with a balloon tamponade device in place.

**INDICATIONS**

The general management of unstable patients with acute variceal bleeding is described in detail elsewhere. In brief, initial resuscitation should focus on early endotracheal intubation; circulatory resuscitation, including blood transfusion and administration of vasoactive agents and antibiotics; and, most importantly, early endoscopy. Vasoactive agents should be given as soon as possible in cases of confirmed or suspected variceal hemorrhage. Vasoactive medications reduce portal pressure and have been shown to decrease or stop variceal bleeding. Somatostatin and its synthetic analogue octreotide decrease release of the vasodilator hormone glucagon, thereby indirectly resulting in splanchic vasoconstriction and reduced portal blood flow. Vasopressin and its synthetic analogue terlipressin are direct vasoconstrictors and can be given systemically or locally during angiography. These two medications, however, can cause significant coronary, cerebral, and splanchic ischemia and are typically used in patients who fail somatostatin or octreotide therapy.

**CONTRAINDICATIONS**

Because gastroesophageal balloon tamponade devices are typically placed as a final attempt to control hemorrhage and prevent imminent death, contraindications to the device are few. They are limited primarily to conditions that predispose patients to esophageal rupture with balloon inflation and include a history of esophageal stricture and recent esophageal or gastric surgery.
PROCEDURE

Patients with an acute variceal hemorrhage that requires a balloon tamponade device are critically ill. Because these patients are at high risk for vomiting, aspiration, and airway compromise, endotracheal intubation should be strongly considered in all patients before placement of a balloon tamponade tube. For the rare patient who is not intubated, use of soft restraints and administration of appropriate analgesia and sedation are critical for a successful procedure.

For placement of a balloon tamponade tube, begin by testing the esophageal and gastric balloons for air leaks (Fig. 41.3, step 1). If there is any concern about a leak, submerge the balloons in water during inflation. If time permits, inflate the gastric balloon in 100-mL increments to the maximal recommended volume while measuring the pressure. Importantly, do not exceed a pressure of 15 mm Hg within the gastric balloon with each successive instillation of 100 mL. Note the pressure at full inflation of the gastric balloon. If no air leaks are detected, fully deflate the esophageal and gastric balloons and clamp the inflation ports. If the kit comes with plastic plugs, they may be used in lieu of clamps to occlude the ports and maintain deflation of the balloon during insertion (see Fig. 41.3, step 2). Once fully deflated, coat the balloons with a thin layer of water-soluble lubricating jelly.

When using the Sengstaken-Blakemore tube, it is important to recall that the device does not have an esophageal aspiration port. To construct a makeshift aspiration port, secure a nasogastric (NG) tube to the tamponade tube with silk sutures such that the distal tip of the NG tube is placed approximately 3 cm proximal to the esophageal balloon (see Fig. 41.3, step 3).

Patients should be as heavily sedated as allowed by cardiovascular parameters. Patients are often hypotensive, limiting the use of many agents. Ketamine would be an excellent choice for sedation of most patients. Position the patient properly for insertion of the tube, with the head of the bed elevated to at least 45 degrees. For patients who are unable to tolerate this position, use the left lateral decubitus position. For nonintubated patients, anesthetize the nasopharynx and oropharynx adequately. Accomplish this by using a topical anesthetic spray or jelly combined with a nebulized lidocaine solution.

Orogastric passage of the tamponade tube is the preferred route of insertion, especially in intubated patients. Nasogastric insertion can be attempted in nonintubated patients. Pass the tube at least to the 50-cm mark and preferably to the maximum depth allowed by the length of the tube (see Fig. 41.3, step 4). A 2015 case series reported the successful use of indirect laryngoscopy to rapidly guide the tamponade tube into the esophagus. After the tube is inserted, apply continuous suction to its gastric and esophageal aspiration ports (see Fig. 41.3, step 5). Inflate the gastric balloon with approximately 50 mL of air and obtain a chest radiograph to confirm that the gastric balloon is below the diaphragm (Fig. 41.4; also see Fig. 41.3, step 6). Confirm the location of the gastric balloon; this is essential to reduce the risk for esophageal rupture from inflation of a misplaced gastric balloon.

Case reports have described the use of an endoscope to guide placement of the tamponade tube. Essentially, a snare is placed through the endoscope and used to grasp the end of the tamponade tube. Use of the endoscope allows direct visualization of the tamponade tube and confirmation that the gastric balloon is in the stomach. Added benefits to this approach include the avoidance of inadvertent tube placement in an esophageal diverticulum, dislodgment of previously placed variceal bands, and inflation of the gastric balloon in a hiatal hernia. This approach may be feasible if the emergency provider has experience with the endoscope device.

Once the location of the gastric balloon is confirmed, connect a manometer to the pressure-monitoring outlet of the gastric balloon (see Fig. 41.3, step 7). Inflate the gastric balloon to the recommended total volume of air in 100-mL increments. Compare the pressure at each 100-mL increment with the values obtained during testing of the gastric balloon. If the pressure during inflation is more than 15 mm Hg higher than the testing pressure at an equivalent volume, it is likely that the gastric balloon has migrated to the esophagus. At this point deflate the gastric balloon and advance it further into the stomach. Obtain another chest radiograph before reinflation.

When the gastric balloon is fully inflated, clamp the inflation and pressure-monitoring ports (see Fig. 41.3, step 8). Slowly pull the device back until resistance is encountered (see Fig. 41.3, step 9). Resistance indicates that the gastric balloon has engaged the cardia and fundus of the stomach (Fig. 41.5). To maintain proper position of the gastric balloon, apply continuous traction. To accomplish this, use an overhead frame and pulley system, a football helmet or catcher’s mask, or the sponge rubber cuff (provided in most kits) for patients who underwent nasogastric insertion. Of these methods, the pulley system is preferred to deliver the recommended 0.5 to 1.0 kg of traction. If the emergency physician does not have the weights required for a pulley system, a 1-L bag of crystalloid solution can conveniently provide 1 kg of traction (Fig. 41.6).

After traction is applied, connect the esophageal and gastric aspiration ports to continuous suction (see Fig. 41.3, step 10). If blood is obtained from either port, inflate the esophageal balloon to approximately 35 to 40 mm Hg. Similar to the gastric balloon, monitor inflation of the esophageal balloon with a manometer connected to the esophageal pressure-monitoring outlet (see Fig. 41.3, step 11). In general, do not inflate the esophageal balloon to more than 45 mm Hg. Keep the esophageal balloon pressure at the lowest inflation pressure that achieves hemostasis. Occasionally, esophageal pressure may transiently spike to values approaching 70 mm Hg. This can occur with respiratory variation or esophageal contraction and is not indicative of overinflation. Once hemostasis is achieved, increase the traction gradually to a maximum of 1.2 kg.

AFTERCARE

Although the primary objective is to control variceal bleeding, the emergency physician must continue care of the patient with a balloon tamponade device until transfer to an intensive care unit, an operating room, a radiology suite, or another facility. Patients should be maintained in a position with the head of the bed elevated to approximately 45 degrees. In addition, it is critical to continue administering sedative and analgesic medications. Connect the esophageal and gastric aspiration ports to continuous suction for approximately the first 12 hours.
Test the esophageal and gastric balloons for air leaks, by submerging under water during inflation. If time permits, record pressures during gastric balloon inflation (see text for details).

Fully deflate the esophageal and gastric balloons. Clamp the inflation ports with a tube clamp, or insert the plastic plugs supplied with the tube into the tube lumen. Lubricate the tube and balloons with water-soluble jelly.

Construct a makeshift esophageal aspiration port by securing a standard NG tube to the SB tube with silk sutures. The distal tip of the NG tube should be 3 cm proximal to the esophageal balloon.

Pass the tube orally (preferred) or nasally, to at least the 50-cm mark, or to the maximum depth allowed by the tube.

After the tube is fully inserted, apply continuous suction to the gastric and esophageal aspiration ports.

Inflate the gastric balloon with 50 mL of air and obtain a chest radiograph to confirm the position of the gastric balloon below the diaphragm.

**Figure 41.3** Balloon tamponade of gastroesophageal varices with the Sengstaken-Blakemore (SB) tube. NG, Nasogastric.

Continued
Mucosal ulceration from direct pressure of the balloons can occur within just a few hours after tube placement. Accordingly, examine the tube, nares, mouth, tongue, and lips frequently, and monitor esophageal balloon pressure periodically. Once the bleeding has been controlled for several hours, decrease the pressure in the esophageal balloon by approximately 5 mm Hg every 3 hours until a pressure of 25 mm Hg is reached. Regardless of the pressure, periodically deflate the esophageal balloon for several minutes every 5 to 6 hours to decrease the incidence of mucosal ischemia and necrosis. Once
device, treat respiratory distress as airway obstruction until proved otherwise. In these patients, use surgical scissors to cut across the lumen of the tube just distal to the inflation and aspiration ports. This will result in deflation of both balloons and allow immediate extraction of the device. Given the risk for airway obstruction, always keep surgical scissors at the bedside of patients who have a balloon tamponade device in place.

Esophageal perforation is another catastrophic complication of balloon tamponade that is almost universally fatal. This dreaded complication can occur from a misplaced gastric balloon, an overinflated esophageal balloon, or prolonged inflation of the esophageal balloon and can result in decreased mucosal blood flow, ischemia, and necrosis. To minimize the risk for esophageal perforation, obtain radiographic confirmation

hemorrhage is controlled and the patient is stabilized, balloon tamponade devices are generally left in place for approximately 24 hours.

**COMPLICATIONS**

This is a difficult procedure that is rarely performed; hence, complications from balloon tamponade can be severe and occur in up to 20% of patients. Major complications include airway obstruction, esophageal rupture, and aspiration pneumonitis. Airway obstruction can be catastrophic and usually results from migration of a dislodged esophageal balloon into the oropharynx. Prevent proximal migration of the tube by maintaining adequate inflation of the gastric balloon, radiographic confirmation, and periodic monitoring of inflation pressure. In nonintubated patients with a balloon tamponade device, treat respiratory distress as airway obstruction until proved otherwise. In these patients, use surgical scissors to cut across the lumen of the tube just distal to the inflation and aspiration ports. This will result in deflation of both balloons and allow immediate extraction of the device. Given the risk for airway obstruction, always keep surgical scissors at the bedside of patients who have a balloon tamponade device in place.

Esophageal perforation is another catastrophic complication of balloon tamponade that is almost universally fatal. This dreaded complication can occur from a misplaced gastric balloon, an overinflated esophageal balloon, or prolonged inflation of the esophageal balloon and can result in decreased mucosal blood flow, ischemia, and necrosis. To minimize the risk for esophageal perforation, obtain radiographic confirmation
CONCLUSION

Balloon tamponade is a critical, lifesaving procedure that may be required in the ED management of unstable patients with bleeding gastroesophageal varices. Indications for placement of a balloon tamponade tube include unsuccessful control of hemorrhage with endoscopy and vasoactive medications, unavailability of consultant physicians when bleeding cannot be controlled with vasoactive therapy, and massive hemorrhage preventing endoscopy. Although complications of balloon tamponade can be fatal, their incidence can be markedly reduced through a stepwise approach to tube placement. Once bleeding is controlled, the emergency physician must continue to monitor tube position and measure balloon pressure until the patient is transferred out of the ED.
REFERENCES


Decontamination of the Poisoned Patient

Christopher P. Holstege and Heather A. Borek

CHAPTER 42

INTRODUCTION

In 2014, the National Poison Data System of the American Association of Poison Control Centers reported 2,165,142 human toxic exposures and 1408 resultant fatalities. Of these total exposures, 28.3% were managed in a health care facility. Massive exposure to some toxic agents (e.g., calcium channel blockers, tricyclic and other antidepressants, antipsychotics, β-blockers, colchicine, chloroquine, cyanide, Amanita phalloides mushrooms, and paraquat) will probably result in severe morbidity or fatality regardless of even the most sophisticated and timely medical interventions. With general supportive care and the use of a few specific antidotes, however, the mortality rate in unselected overdose patients is less than 1% if the patient arrives at the hospital in time for the clinician to intervene.

Key management of poisoned patients seen in health care facilities initially focuses on confirming the diagnosis of possible exposure to a toxin, providing standard cardiovascular and respiratory supportive care, and using a small cadre of specific antidotes. In rare and yet undefined selected instances, prevention of further absorption of toxin by various decontamination procedures may theoretically ameliorate the morbidity or reduce mortality. Although a better final outcome after gastric decontamination may seem intuitively reasonable, there is no definitive evidence from prospective clinical trials that the use of various decontamination techniques positively alters the morbidity or mortality of a poisoned patient.

Before the availability of objective or experimental evidence addressing gastric emptying procedures, most clinicians instituted previously performed unproven decontamination procedures in the emergency department (ED) as a reflex response for the majority of patients suspected of drug overdose, often without much forethought and certainly without confirming data. Mounting evidence relegates any benefit from any form of gastric decontamination to selected cases and specific individual scenarios. In fact, because of the lack of demonstrable benefit and the mounting evidence for potential harm, syrup of ipecac, once a mainstay in the management of poisonings, is no longer recommended, and parents have been instructed by the American Academy of Pediatrics to remove it from the home. Parents and health care providers appear to be heeding this recommendation because virtually no (less than 0.01%) poisoned patients received ipecac in 2014.

Nonetheless, a selective role for other methods of gastric decontamination exists, and there will always be a role for real-time clinical judgment. Because compelling circumstances may clinically support gastric decontamination, this chapter discusses specific clinical procedures. These techniques include gastric lavage, oral administration of activated charcoal, and whole-bowel irrigation (WBI). In addition, dermal decontamination as a result of a toxic exposure is also addressed. Before performing these techniques, the clinician responsible for the care of a poisoned patient must clearly understand that these procedures are not without hazard and that any decision on their use must consider whether the benefit of decontamination outweighs any procedure-related harm.

GASTRIC DECONTAMINATION

Gastric Lavage

Background

The use of gastric lavage (Video 42.1) in poisoned patients has decreased significantly since the 1990s, and in the United States it is currently reported to be used in less than 1% of overdose cases. Numerous animal and human volunteer studies have been conducted to examine the effectiveness of gastric lavage in removing toxins from the stomach, especially with respect to other gastrointestinal decontamination methods. The reported efficacy of gastric lavage in removing markers from the stomach varies significantly in these studies. The difference in these study results is due in part to the variability of the methods used (different fluid-instilled markers, animal models, positioning, amount of lavage, and lavage tube size) and the time that elapsed from instillation of the marker in the stomach until gastric lavage was performed. Even within individual studies, the range of effectiveness of gastric lavage in removing the marker varied considerably. For example, Tandberg and co-workers performed gastric lavage 10 minutes after ingestion of the marker and reported that its effectiveness in removing the marker varied from 18.9% to 67.7%.

Many of these studies do not replicate the typical clinical scenario encountered in emergency medicine. The efficiency of gastric lavage in removing a marker significantly decreases with increasing time after ingestion. This is due to the fact that as time increases after ingestion, there is more time for the marker to be absorbed and passed out of the stomach. For example, Shrestha and colleagues reported that more than 70% of the marker used in their study passed out of the stomach in 60 minutes. It is rare that gastric lavage can be performed within the first hour after toxic ingestion. Not only does it take time for these patients to arrive at the ED, but it also takes time for evaluation, stabilization, and performance of gastric lavage. Watson and associates reported that the mean time required by experienced emergency medicine nurses to perform lavage was 1.3 hours. Gastric lavage may also propel the marker from the stomach into the small intestine, thereby decreasing the effectiveness of removing the toxin from the stomach and actually enhancing the rate of absorption.

Three major studies have examined whether gastric lavage positively influences the outcome of poisoned patients. In a study performed by Kulig and co-workers, there was no difference in outcome in patients who received gastric lavage followed by charcoal versus charcoal alone when these interventions were performed more than 1 hour after ingestion. In patients who were treated within 1 hour of ingestion, gastric lavage followed by charcoal provided a small but statistically significant advantage over activated charcoal alone. Merigian and colleagues demonstrated that in symptomatic patients, the rate of intensive care admission and need for intubation was significantly higher in patients who received gastric lavage...
Gastric Lavage

**Indications**
Potentially life-threatening poisonous ingestions, but only if the procedure can be performed within 60 minutes

**Contraindications**
- Compromised airway protective reflexes (unless patient is intubated)
- Ingestion of corrosive substances (acids or alkalis)
- Hydrocarbons (unless containing highly toxic substances such as pesticides)
- Known esophageal strictures
- History of gastric bypass surgery

**Complications**
- Trauma due to tube insertion
- Instillation of lavage fluid into lungs/aspiration
- Cardiac dysrhythmias
- Hypoxia
- Laryngospasm
- Fluid and electrolyte disturbances
- Hypothermia

followed by charcoal than in those who received charcoal alone. This increased admission and intubation rate was directly attributed to the aspiration of gastric contents as a result of gastric lavage. Pond and associates replicated the study by Kulig and colleagues. They found no difference in outcome between patients who received gastric lavage followed by charcoal and those receiving charcoal alone, regardless of the time of performance of gastric lavage. They concluded that “gastric emptying procedures can be omitted from the treatment regimen for adults after acute overdose, including those who present within 1 hour of overdose and those that manifest severe toxicity.” Westergaard and colleagues recently demonstrated that despite current recommendations against routine use of gastric lavage, the vast majority of cases where it was actually performed did not meet specific criteria for the indications for gastric lavage.

**Indications**
Based on the available literature, gastric lavage should not be routinely used in the management of poisoned patients (Fig. 42.1). There is no universally accepted standard of care that can be applied to the use of gastric lavage in unselected poisoned patients in the ED. Under certain circumstances, however, there may be a theoretical benefit from gastric emptying, and the local poison center should be contacted to assist in deciding whether gastric lavage may be of benefit. Whether specific subsets of overdose patients may benefit from gastric lavage has not been clearly defined.

Only patients who have ingested a potentially life-threatening amount of poison, and in whom the procedure can be performed within 60 minutes, are the primary candidates for gastric lavage. Oral charcoal alone is considered superior to gastric lavage if a drug is adsorbed by charcoal.
Gastric lavage is contraindicated in persons who have ingested corrosive substances (acids or alkalis) or hydrocarbons (unless they contain highly toxic substances such as pesticides), have known esophageal strictures, or have a history of gastric bypass surgery. Caution should be exercised in performing gastric lavage in combative patients and in those who have medical conditions such as bleeding diatheses that could be compromised by performing this procedure.

**Equipment and Preparation**

If the decision is made to perform gastric lavage, careful attention to the details of the procedure results in increased safety for the patient and more effective removal of the ingested poison. Before lavage, intravenous access should be secured and continuous cardiac monitoring and pulse oximetry should be initiated (Fig. 42.2, step 1). A large, rigid suction catheter should be available immediately.

If the patient is highly anxious or agitated, give a small dose of an intravenous benzodiazepine (e.g., midazolam). If the patient's level of consciousness is significantly depressed, airway status is questionable, or the airway is likely to be compromised during the procedure, consider rapid-sequence induction and intubation with a cuffed endotracheal tube before initiating gastric lavage. If the patient is fully awake and alert, proceed to lavage without tracheal intubation. The procedure should proceed deliberately without significant patient resistance. It is intended to be therapeutic, not punitive. Antiquated arguments promulgating that a noxious lavage procedure will keep patients from overdosing again should be abandoned.

The position of the patient during gastric lavage is important. Place all patients in the left lateral decubitus, Trendelenburg position (≈20-degree tilt on the table) (Fig. 42.3). This position diminishes the passage of gastric contents into the duodenum during lavage and decreases the risk for pulmonary aspiration of gastric contents should vomiting or retching occur. Restrain the hands of an uncooperative patient to prevent removal of the gastric or endotracheal tube. Intubated patients on a ventilator may be lavaged in the supine position because of logistic reasons (Fig. 42.4). Under no circumstances should a nonintubated patient undergo lavage in the restrained supine position. Such positioning invites aspiration and diminishes the patient’s natural protective maneuvers, such as coughing and sitting up.

Most clinicians prefer the oral route for gastric lavage, but in selected circumstances a standard large-bore nasogastric (NG) tube (Salem sump pump) may be used. Large-diameter gastric hoses with extra holes cut near the tip have traditionally been recommended for gastric lavage. There are no convincing data on humans to refute or support this recommendation, although large-diameter NG tubes may kink and are significantly more difficult to pass. An 18-Fr Salem sump pump is the maximum diameter for liquid ingestions and in children (Fig. 42.6).

In most cases, a 36- to 40-Fr or a 30-English gauge tube (external diameter, 12 to 13.3 mm) should be used in adults and a 24- to 28-Fr gauge (diameter, 7.8 to 9.3 mm) tube in children. Before passage, estimate the length of tube required to enter the stomach by approximating the distance from the corner of the mouth to the midepigastrium. Premedication avoids the curling and kinking of excess hose in the stomach (see Fig. 42.2, step 2). Passage of an excessive length of hose may cause gastric distention, bruising, and perforation, whereas passage of an insufficient length of hose may result in lavage of the esophagus and increased risk for emesis and aspiration. Commercial lavage systems are available and often use either a gravity fill-and-empty system with a Y-connector or a closed irrigation syringe system. Alternatively, an irrigation syringe can be used for intermittent input and withdrawal of lavage fluid.

**Technique**

Lubricate the gastric tube and pass it gently to avoid damage to the posterior pharynx (see Fig. 42.2, step 3). Use of a bite block or an oral airway may prevent the patient from chewing on the orogastric tube and biting the fingers of the inserter. If the patient is obtunded or paralyzed, extend the jaw to facilitate passage. Never use force to pass the tube. Once the pharynx has been entered, put the patient's chin on the chest to facilitate passage of the tube into the esophagus (see Fig. 42.2, step 4). Cough, stridor, or cyanosis indicates that the tube has entered the trachea; withdraw the tube immediately and reattempt passage. Once the tube is passed, confirm that it is in the stomach. Intragastric placement is usually evident on clinical grounds by the spontaneous egress of gastric contents but may be confirmed by auscultation of the stomach during injection of air with a 30-mL syringe followed by successful aspiration of gastric contents (see Fig. 42.2, step 5). However, due to the risk of lavage through a misplaced tube, confirm tube position radiographically before lavage is performed (Fig. 42.7A). A misplaced tube may irrigate the esophagus with a tube that has doubled back on itself during passage (see Fig. 42.7B). The most serious complication, other than esophageal perforation, is inadvertent passage of the tube into the lungs. Tracheal passage of a lavage tube should be readily obvious in an awake patient before lavage, and obtunded patients are intubated, thereby obviating this problem. If an awake patient begins to vomit during lavage, immediately remove the tube to allow the patient to protect the airway.

Before gastric irrigation, remove the gastric contents by careful gastric aspiration with repeated repositioning of the tip of the tube (see Fig. 42.2, step 6). With the Y-connector closed system, perform lavage by clamping the drainage arm of the Y-adapter and infusing aliquots of fluid into the stomach from a reservoir. Clamp the reservoir arm of the Y, and then open the drainage arm to permit drainage of the stomach contents by gravity. Repeat this procedure. Some resistance is produced by the Y-connector and tubing. Apply suction intermittently to the drainage tubing to enhance emptying of the stomach.

Lavage can be performed adequately with tap water in adults. Because electrolyte disturbances have occurred in children who underwent lavage with tap water, prewarmed (45°C) normal saline is generally recommended for children. Warmed lavage fluid increases the solubility of most substances, delays gastric
Before lavage, obtain intravenous access and place the patient on a continuous cardiac monitor and pulse oximeter. Restrain the hands of uncooperative patients.

Lubricate the gastric tube and pass it gently to avoid damage to the posterior pharynx. Use a bite block or an oral airway to prevent the patient from biting the tube or inserter.

Once the tube is passed, confirm that it is in the stomach via auscultation and aspiration of gastric contents. A radiograph may be obtained if deemed clinically necessary.

Repeatedly introduce small aliquots (200–300 mL in adults) into the stomach and then remove them. Perform this step with the patient in the left lateral decubitus position.

Measure and mark the appropriate depth of the gastric lavage tube before passage. This ensures that the tip is in the stomach and that there is no excess tubing that may kink or knot the tube.

Once the tube enters the pharynx, put the patient’s chin on the chest to facilitate passage of the tube into the esophagus. Pass the tube into the stomach.

After gastric aspiration and lavage are completed, administer a slurry of activated charcoal through the gastric tube. When no longer needed, clamp off the gastric tube before removal.

**Figure 42.2** Gastric lavage. In an awake patient the orogastric tube can be passed with the patient in the sitting position, but the lavage procedure is best performed with the patient in the left lateral decubitus position.
should approximate the amount introduced. Manual agitation of the patient’s stomach by gently “kneading” it with a hand placed on the abdomen may increase recovery. Continue lavage until the fluid becomes clear.

After gastric aspiration and lavage have been completed, administer a slurry of activated charcoal through the gastric tube (see Fig. 42.2, step 8). When no longer needed, clamp off the gastric tube before removal to avoid “dribbling” fluid into the airway. With the increasing use of repetitive doses of activated charcoal, an NG tube may be left in place, or passed,
Instillation of lavage fluid and charcoal into the lungs through patients prior to administering lavage fluid (see Fig. 42.7). Misplacement were not evaluated, however.

Too often for benign ingestions. Intuitively reasonable, it is of unproven value and probably done far too often for benign ingestions. The common routine practice of passing an NG tube in a child who is unwilling to drink charcoal is controversial, and though it does exist, the patient to gagging, drooling, or aspiration, it is imperative to confirm correct placement. Scalzo and associates found radiographically that 7 of 14 children had improper tube placement (too high or too low) despite positive gastric auscultation in all cases. The clinical effects of such misplacement were not evaluated, however. Radiographic confirmation of tube placement should be considered in all patients prior to administering lavage fluid (see Fig. 42.7).

Instillation of lavage fluid and charcoal into the lungs through after the lavage procedure is completed. A patient who remains obtunded may receive additional doses via a standard NG tube. Because the large gastric tube is irritating and may predispose the patient to gagging, drooling, or aspiration, it should be removed. Alert patients should take subsequent doses of charcoal orally as necessary.

Complications
A correctly performed procedure in the appropriate environment is generally safe, but numerous complications have been associated with gastric lavage. The complications can be divided into those caused by mechanical trauma and those resulting from the lavage fluid.

Depending on the route selected for insertion of the tube, damage to the nasal mucosa, turbinates, pharynx, esophagus, and stomach has been reported. After insertion of the tube, it is imperative to confirm correct placement. Scalzo and associates found radiographically that 7 of 14 children had improper tube placement (too high or too low) despite positive gastric auscultation in all cases. The clinical effects of such misplacement were not evaluated, however. Radiographic confirmation of tube placement should be considered in all patients prior to administering lavage fluid (see Fig. 42.7).

Activated Charcoal

Use of Endoscopy to Perform Gastric Lavage
The technique of gastric lavage can also be accomplished using an endoscopic approach. Whereas this is not routinely recommended or performed, there is literature that shows favorable outcomes using endoscopy in specific cases, especially when a pharmacobezoar develops. Bezoar formation may occur with overdoses of a large number of pills or certain types of agents (e., salicylates, anticholinergics). There are numerous case reports of bezoar formation following acute overdose, with subsequent endoscopic removal and clinical improvement.

Activated Charcoal

Background
Activated charcoal is a carbon product that is subjected to heat and oxidized to increase its surface area (Fig. 42.8). It has the capacity to adsorb substances onto the porous surface of the charcoal. The use of activated charcoal for poisoning has been recognized for almost two centuries. To demonstrate the effectiveness of charcoal, in 1930, French pharmacist Touery ingested several times the lethal dose of strychnine mixed with 15 g of activated charcoal. He performed this act in front of a class of colleagues and exhibited no ill effects.

Activated charcoal acts both by adsorbing a wide range of toxins present in the gastrointestinal tract and by enhancing elimination of toxins if systemic absorption has already occurred. It enhances elimination by creating a concentration gradient between the contents of the bowel and the circulation, but it also has the potential to interrupt enterohepatic circulation if the particular toxin is secreted in bile and enters the gastrointestinal tract before reabsorption.

Oral activated charcoal
Decontamination of the Poisoned Patient

Charcoal occurs over time, although this has little clinical significance for most patients and can be overcome by administering additional charcoal.

**Indications**

For years, administration of a single oral dose of activated charcoal for essentially all overdoses was routine. However, with the emergence of new guidelines in overdose management the use of charcoal declined to less than 1% of poisoned patients in 2014. Clearly, charcoal binds many toxins in the gut, thereby decreasing some systemic absorption. It has been shown to decrease the total amount of drug absorbed by the body and also the drug’s half-life. Despite a lack of scientific data demonstrating a decrease in morbidity and mortality and firm evidence to support its widespread use, charcoal is a reasonable intervention for most poisoned patients encountered in the ED if it can be administered easily and safely (Fig. 42.9). The exact indications are not established, and no universally accepted standard of care has been promulgated. A single dose of activated charcoal is indicated if the clinician estimates that a clinically significant fraction of the ingested substance remains in the gastrointestinal tract, the toxin is adsorbed by charcoal, further absorption may result in clinical deterioration, and it can be administered safely. This will usually be a clinical decision because adequate historical data may often be lacking. It may also be administered by multiple dosing if the clinician anticipates that the charcoal will result in increased clearance of an already absorbed drug. It is most effective within the first 60 minutes after an oral overdose, however this time frame may

**Figure 42.7** Radiographic confirmation of gastric lavage tube placement. A, This radiograph confirms proper placement of the lavage tube. It can be seen descending below the diaphragm (1), and the interruption in the radiopaque line (which signifies the most proximal hole) is in the stomach (2). The tip of the tube (3) is also visible. B, The lavage tube in this patient is misplaced. The tube can be seen descending beneath the diaphragm (1) but is kinked in the stomach (2) and has doubled back on itself. The gap in the radiopaque line (3) is at the level of the gastroesophageal junction, and the distal tip of the tube is in the midesophagus (4). Proper measurement of the tube before insertion would probably have prevented this misplacement. If there is any doubt about proper positioning of the tube, a radiograph should be obtained before lavage.

**Figure 42.8** Activated charcoal. Because the contents may settle with time, shake the bottle vigorously before administration. Rinse the container with a small amount of tap water as well to ensure that the patient receives the full dose. Plain charcoal, without sorbitol, is preferred.

is given as a single dose or in multiple doses. The adsorptive capacity of charcoal depends on the inherent properties of the toxin and the local milieu, such as pH. Adsorption begins within minutes of contact with a toxin but may not reach equilibrium for 20 to 30 minutes. Desorption of toxins from
be longer for substances that are likely to form bezoars, such as aspirin. Charcoal is generally considered to provide superior gut decontamination over gastric lavage. There is no definitive evidence, however, that administration of activated charcoal improves outcome.

Contraindications
Administration of charcoal is contraindicated in any person who demonstrates compromised airway-protective reflexes unless already intubated. It is absolutely contraindicated in persons who have ingested corrosive substances (acids or alkalis). Not only does charcoal provide no benefit in a corrosive ingestion, but its administration could also precipitate vomiting, obscure endoscopic visualization, and lead to complications if a perforation developed and charcoal entered the mediastinum, peritoneum, or pleural space. Charcoal should be avoided with ingestion of a pure aliphatic petroleum distillate. Hydrocarbons are not well adsorbed by activated charcoal and its administration could lead to further risk for aspiration. Many hydrocarbons are potential systemic toxins (e.g., carbon tetrachloride and benzene) or are mixed with other potentially significant toxins such as pesticides. In these cases, data are lacking, but charcoal administration can be considered. Caution should be exercised in using charcoal in patients with medical conditions that could be further compromised by charcoal ingestion, such as gastrointestinal perforation or bleeding and in those patients with gastric bypass. Charcoal is not indicated for isolated gastrointestinal perforation or bleeding and in those patients who have ingested a toxin that may suddenly induce seizures. Because it is often impossible to determine the exact nature of an ingestion, a liberal-use policy is advocated for potentially mixed overdoses.

Charcoal administration by paramedics and other emergency response personnel should be performed with caution. There is insufficient evidence to recommend for or against administration of activated charcoal in the prehospital setting. The same indications and contraindications apply as for patients who are in the hospital. The motion of the ambulance during transport may make the patient more prone to emesis. Either spilling of charcoal or vomiting of charcoal may result in significant contamination of the transport vehicle and subsequently place that vehicle out of commission until it can be cleaned.

Technique
There is no universally accurate dose for charcoal. A 10:1 ratio (charcoal to toxin) is recommended if the amount of ingestion is known. Dosing of charcoal should be considered in light of the specific ingestion, but the following empirical doses of single-dose activated charcoal (standard aqueous products such as Liqui-Char [King Pharmaceuticals, Bristol, TN]) are recommended:
- Up to 1 year: 1 g/kg of body weight
- 1 to 12 years: 25 to 50 g
- Older than 12 years: 25 to 100 g

If the ingestion were, for example, clonidine (0.1-mg tablets) or diphenoxylate (0.25-mg tablets), this regimen would be more than adequate for even a massive overdose to achieve the desired 10:1 ratio. If the ingestion consisted of a large number of 325-mg aspirin tablets or 240-mg verapamil tablets, the dosing regimen could be insufficient. If toxic medications with a high-milligram dosage are ingested, it would be prudent to administer more charcoal than indicated by these guidelines. There is no known benefit of mixing charcoal with a cathartic (i.e., sorbitol), and the combination is not suggested. Sorbitol may increase the incidence of vomiting.

Because in many charcoal formulations the contents settle with time, shake the preparation vigorously before administering it to the patient. Follow this by rinsing the container with a small amount of tap water before administering it to the patient to allow ingestion of the full dose. Aqueous activated charcoal has a gritty texture that most patients find unpleasant, but attempts have been made to improve its taste and texture. Mixing activated charcoal with chocolate milk, chocolate- or cherry-flavored syrup, or ice cream may increase palatability, but mixing with these additives has been suggested, though not proved, to cause a decrease in the adsorptive capacity of activated charcoal. Rangan and colleagues reported no decrease in adsorption after mixing superactivated charcoal with a noncaffeinated cola. Scharman and associates demonstrated that a regular, sugared cola was favored by children over a diet cola, but only 20% of the time were they able to cajole even nonpoisoned children younger than 3 years to drink a therapeutic amount of flavored charcoal.

Give activated charcoal orally if the patient is awake and cooperative and by NG tube if the patient is unconscious. If an NG tube is inserted, it is imperative that correct placement be verified radiographically before administering charcoal, especially in obtunded or intubated patients (see Fig. 42.7). Installment of charcoal into the lungs has been reported after inadvertent misplacement within the airways, and massive aspiration can be fatal (Fig. 42.10). Intubation is protective, but it is not uncommon to see some charcoal in the airway even if the patient has been intubated.

A clinical conundrum exists when a patient refuses to drink charcoal and an NG tube must be passed without consent if charcoal is deemed advisable. The common tactic of passing an NG tube in an awake but uncooperative patient merely to administer charcoal is controversial, and no standards exist. Such a scenario is more likely to result in trauma from placement of the tube, a misplaced tube, or subsequent emesis from the
rapid administration of charcoal. Given the unproven efficacy of charcoal, the authors advise against routine insertion of an NG tube simply to administer charcoal in an awake and minimally symptomatic patient. Such a decision is, however, a clinical one that must be made by the health care provider and be based on the entire clinical milieu (Fig. 42.11).

**Complications**

Administration of activated charcoal is not without risks and complications. Published reports have demonstrated adverse effects associated with activated charcoal therapy, including childhood deaths.52 The most common complications of charcoal administration include constipation, diarrhea, and vomiting.53,54 Bowel perforation has been described in a patient with diverticular disease.55 Pulmonary aspiration of activated charcoal is a dreaded complication that can result in pneumonitis, obstruction of the respiratory tree, bronchiolitis obliterans, acute lung injury, and barotrauma.56 Risk factors for serious aspiration

Figure 42.10 A, Vomiting can be expected following charcoal administration, especially if sorbitol is added (not recommended). This patient rapidly became drowsy after administration of charcoal, vomited, but fortunately did not aspirate. B, In another patient who aspirated, the charcoal can be seen at the carina with a fiberoptic scope. Massive aspiration can be fatal. Intubation does not totally protect against minimal charcoal aspiration.

Figure 42.11 A, If an overdose patient will voluntarily drink charcoal, there are few reasons to withhold it, even though a definite clinical benefit in routine cases cannot be proved. If a patient will not drink charcoal, patient management becomes controversial. Passing a nasogastric (NG) tube in a struggling patient or in a recalcitrant child merely to instill the unproven, but theoretically useful antidote is not supported by scientific data. Nonetheless, it remains a common procedure. Though not always easy or pleasant, such an intervention is usually safe. Pulmonary aspiration, even in an awake patient, is the major downside. Restrained supine patients are at greatest risk for aspiration, and that position should be avoided, even in an initially awake patient. B, Charcoal that is voluntarily swallowed or instilled via an oral-gastric lavage tube or an NG tube can induce emesis. This occurs in both obtunded and awake patients. In this instance the patient was unconscious from the overdose and the airway was protected with prior tracheal intubation. Although the intubation procedure does not totally exclude pulmonary aspiration and it carries some morbidity in its own right, it is recommended before use of charcoal in patients who are not able to fully protect their airway. Patients who are initially asymptomatic or minimally affected but have ingested drugs that have the potential to produce rapid deterioration, seizures, or loss of airway protection make decisions on the use of charcoal difficult for the clinician. In borderline cases, some experienced clinicians avoid the use of charcoal altogether.
are large amounts of charcoal instilled over a short period, multiple-dose charcoal in the setting of ileus, charcoal administration in a patient who becomes obtunded, charcoal that is inappropriately diluted, or forced administration of charcoal via an NG tube, especially in a restrained supine patient. These complications can be prevented by prudent dosing of charcoal, close monitoring of the airway, performing abdominal examinations with attention to bowel sounds, and focusing on the mechanism of the substance ingested (i.e., if the substance is likely to cause profound sedation or seizures that would increase aspiration risk). Trivial aspiration of charcoal is common and usually innocuous, even if the patient is intubated. Studies show a 4% to 39% incidence of aspiration pneumonia in intubated patients who received activated charcoal. It has been shown that even in patients with a protected airway and a cuffed endotracheal tube, vomiting can lead to pulmonary aspiration of the charcoal. This can result in a significant increase in lung microvascular permeability and lead to pulmonary edema and respiratory compromise.

Multiple Doses of Activated Charcoal

Indications

The use of multiple-dose activated charcoal (MDAC) may be indicated in selected cases (Fig. 42.12). Its use has been advocated for two purposes: first, to prevent continued absorption of a drug that may still be present in the gastrointestinal tract, and second, to increase serum clearance of a drug that has already been absorbed.

MDAC prevents continued absorption by either binding a drug that may be present throughout the gastrointestinal tract or binding a drug that exists as an extended-release or enteric-coated preparation. MDAC enhances elimination of a drug by interrupting enterohepatic recirculation or augmenting enterocapillary exsorption. With interruption of enterohepatic recirculation, charcoal binds to an active drug that is secreted by the biliary system, thus subsequently preventing reabsorption. With augmentation of enterohepatic exsorption, charcoal produces sink conditions that drive diffusion of the drug from capillaries into the intraluminal space, where it is subsequently eliminated. This process is called intestinal dialysis. Drug characteristics associated with enhanced systemic clearance via MDAC include a low intrinsic clearance, a prolonged distributive phase, low protein binding, and a small volume of distribution.

MDAC has been shown to increase total-body clearance of multiple drugs, including carbamazepine, dapson, phenobarbital, quinine, and theophylline. Despite the reported increase in drug clearance associated with the use of MDAC, improved clinical outcomes have not been definitively demonstrated. For example, Pond and co-workers described 10 comatose patients following phenobarbital overdose who were randomized to receive either single-dose activated charcoal or MDAC. Despite the fact that the MDAC group had a significantly shorter phenobarbital serum half-life, no difference was found between the groups with regard to the duration of intubation or hospitalization. A retrospective study of hospitalized patients with phenytoin toxicity showed that those who received single-dose activated charcoal or MDAC did not benefit in the time to clinical improvement or duration of hospitalization compared to those who did not receive charcoal. In a rural, developing-world setting, routine use of MDAC was not found to alter mortality rates.

Contraindications

MDAC is contraindicated in patients with evidence of bowel obstruction. Ileus is a relative contraindication. Many ill patients in whom ileus develops may be candidates for MDAC if the airway is protected. Administration of MDAC is contraindicated in any patient who does not have an intact or protected airway. MDAC should be avoided in patients with repetitive emesis, especially when associated with decreased mental status or a decreased gag reflex. Concurrent use of cathartics with MDAC remains unproved and is not
recommended. MDAC with cathartics should not be administered to young children because of the propensity for laxatives to cause fluid and electrolyte imbalance. For example, MDAC with sorbitol has been associated with hypernatremia and dehydration and MDAC with magnesium cathartics has been associated with hypermagnesemia, neuromuscular weakness, and coma.

Technique

Give 1 g/kg (≤ 100 g) for the first dose of charcoal. If a cathartic is used, administer it only with the first dose of charcoal to decrease the potential risk for cathartic-induced electrolyte abnormalities, especially in children. Follow the initial dose of charcoal with 0.5 g/kg (≤ 50 g) every 4 hours. Stop giving MDAC if repeated examination reveals an absence of bowel sounds or a distended abdomen. In this case, consider placing an NG tube and put it on low intermittent suction. Patients receiving MDAC may be at increased risk for emesis because of the larger total dose of activated charcoal received. The use of antiemetics may help decrease the incidence of vomiting associated with MDAC. Charcoal therapy should be continued until clinical improvement is evident and plasma drug levels have fallen to acceptable levels.

Complications

The complications encountered with single-dose activated charcoal are also encountered with MDAC. In addition, there have been reports of gastrointestinal obstruction and perforation with MDAC therapy, especially in conjunction with the ingestion of drugs that have anticholinergic properties.

Cathartics

Background

The use of cathartics is theoretically intended to decrease the absorption of substances by accelerating expulsion of the poison from the gastrointestinal tract. Cathartics are often used in conjunction with activated charcoal because of charcoal’s side effect of constipation. The mechanism of action of cathartics is such that, theoretically, it would minimize the possibility of desorption of the drug bound to activated charcoal. There is little evidence that a single dose of aqueous activated charcoal is significantly constipating; however, cathartics are often given for this potential problem. The majority of data suggest negligible clinical benefit from the use of cathartics, and the use of cathartics is not supported by the editors.

Indications

Routine administration of a cathartic in combination with activated charcoal is not endorsed by the American Academy of Clinical Toxicology or the European Association of Poison Centers and Clinical Toxicologists. Administration of a cathartic alone has no role in the management of a poisoned patient.

Contraindications

Cathartics are contraindicated in patients with volume depletion, hypotension, significant electrolyte imbalance, ingestion of a corrosive, ileus, recent bowel surgery, and intestinal obstruction or perforation. Administration of cathartics is also contraindicated in patients who do not have an intact or protected airway. They should be avoided in those with repetitive emesis, especially when associated with decreased mental status or a decreased gag reflex. Cathartics should be used cautiously in young children and the elderly because of the propensity for laxatives to cause fluid and electrolyte imbalance.

Technique

There are two types of osmotic cathartics: saccharide cathartics (sorbitol) and saline cathartics (magnesium citrate, magnesium sulfate, and sodium sulfate). The optimal dose of sorbitol or magnesium citrate remains to be determined. The recommended dose of sorbitol is approximately 1 to 2 g/kg of body weight or 1 to 2 mL/kg of 70% sorbitol in adults and 4.3 mL/kg of 35% sorbitol in children (single administration only). Charcoal formulations may come premixed with sorbitol, but the sorbitol content varies considerably. The recommended dose of magnesium citrate is 250 mL of a 10% solution in an adult and 4 mL/kg body weight of a 10% solution in a child. Multiple doses of cathartics should be avoided.

Complications

Administration of sorbitol has been associated with vomiting, abdominal cramps, nausea, diaphoresis, and transient hypotension. Because the sorbitol content varies between different charcoal-sorbitol combination products, pay attention to the sorbitol content in each brand to avoid excessive sorbitol administration. Be aware that multiple doses of sorbitol have been associated with volume depletion. Multiple doses of magnesium-containing cathartics have been linked to severe hypermagnesemia. Children are particularly susceptible to the adverse effects of cathartics, so use caution or totally avoid using cathartics in children.

Whole Bowel Irrigation

Background

WBI involves the enteral administration of an osmotically balanced polyethylene glycol electrolyte solution (PEG-ES) in a sufficient amount and rate to physically flush ingested substances through the gastrointestinal tract, thereby purging the toxin before absorption can occur. PEG-ES (CoLyte [Pendopharm, Montreal, Quebec, Canada], GoLYTELY [Braintree Laboratories Inc., Braintree, MA]) is isosmotic, is not systemically absorbed, and will not cause electrolyte or fluid shifts. The data available suggest that the large volumes of this solution needed to mechanically propel pills, drug packets (such as in body packers or stuffers), or other substances through the gastrointestinal tract are safe, including use in pregnant women and young children.

Clinical data on the efficacy of WBI remain limited. Ly and colleagues found that the effect of WBI on reduction of acetaminophen concentration versus time was not statistically significant. However, WBI did have a mechanical effect on radiopaque markers in the gastrointestinal tract, with 8 of 10 patients’ markers congregating in the right hemicolon after WBI. A retrospective study by Breauadeau Deguigne and colleagues suggested a reduction in the severity of acute-on-chronic lithium poisoning at a given ingested dose and serum lithium level with those undergoing early decontamination (by sodium polystyrene sulfonate and/or WBI) versus those in whom decontamination was delayed. WBI was shown to mobilize lead BB pellets to the large bowel in a child, where less absorption occurs and the foreign bodies could be removed by colonoscopic intervention. In addition, PEG-ES may play a role in the pharmacologic conversion of some toxins. For
example, it has been shown that the relatively high pH of PEG-ES increases the rate of spontaneous conversion of cocaine to its inactive metabolite benzoylecgonine.93

Indications
WBI may be considered for the ingestion of exceedingly large quantities of potentially toxic substances, toxins that are poorly absorbed by activated charcoal (e.g., iron, lithium), delayed-release formulations, late arrival at the ED after ingestion of a toxin, pharmacobezoars, and body stuffers or packers (Fig. 42.13).96 WBI remains a theoretical option for these ingestions and is often performed on body packers who have ingested many times the lethal amount of such drugs as heroin or cocaine (Fig. 42.14). No definitive evidence exists to show that WBI improves the outcome of poisoned patients.96 Though not a proven procedure, WBI is often suggested by toxicologists, and its use in selected cases is intuitively reasonable and supported by the authors. The most common indication for WBI in the ED is for the treatment of toxic sustained-release medications (such as iron, calcium channel blockers, β-blockers, and lithium) and iron tablets (Fig. 42.15).94

Contraindications
WBI is contraindicated in patients with gastrointestinal obstruction, perforation, ileus, or ingestion of a corrosive agent. It should also be avoided in patients with hemodynamic instability or an unprotected airway.94 WBI should likewise be avoided in debilitated patients and in patients with medical conditions that might be further compromised by its use. The concurrent administration of activated charcoal and WBI might decrease the effectiveness of the charcoal.

**POSITION STATEMENT: WHOLE BOWEL IRRIGATION**

American Academy of Clinical Toxicology; European Association of Poisons Centres and Clinical Toxicologists

Based mainly on volunteer studies, whole bowel irrigation (WBI) can be considered for potentially toxic ingestions of sustained-release or enteric-coated drugs particularly for those patients presenting later than 2 h after drug ingestion when activated charcoal is less effective. WBI can be considered for patients who have ingested substantial amounts of iron, lithium, or potassium as the morbidity is high and there is a lack of other potentially effective options for gastrointestinal decontamination. WBI can be considered for removal of ingested packets of illicit drugs in “body packers.” However, controlled data documenting improvement in clinical outcome after WBI are lacking. WBI is contraindicated in patients with bowel obstruction, perforation, or ileus, and in patients with hemodynamic instability or compromised unprotected airways. WBI should be used cautiously in debilitated patients and in patients with medical conditions that might be further compromised by its use. The concurrent administration of activated charcoal and WBI might decrease the effectiveness of the charcoal.


Figure 42.14 A, This body packer attempted to smuggle more than 50 packets of heroin. All packets were passed intact after 12 hours of whole-bowel irrigation. B, Note the integrity of the carefully wrapped packets that were passed.

Figure 42.15 Whole-bowel irrigation (WBI) is commonly recommended for the treatment of iron ingestion. These radiographs depict the effect of 5 hours of WBI. Note the marked decrease in radiopaque pills (arrow) in the gastrointestinal tract. Intact pills were recovered in the rectal effluent.
with patients who have repetitive emesis, especially when associated with decreased mental status or a decreased gag reflex. WBI should be used cautiously in debilitated patients. WBI should be avoided in gastric bypass patients, due to the large volume of fluid that must be administered and risk of perforation.

**Technique**

PEG-ES is marketed in a powder form. Add tap water to make a total volume of 4 L. The recommended rate of administration is as follows:

- 9 months to 6 years: 500 mL/hr
- 6 to 12 years: 1000 mL/hr
- Older than 12 years: 1500 to 2000 mL/hr

Cooperative patients with intact airway-protective reflexes may drink the solution. The large volume and taste often limit even the most motivated patient's ability to comply. If the patient is unable or unwilling to drink this solution, administer it through a small-bore NG tube after placement is confirmed. Because it is common for WBI to be delayed while the patient and medical personnel attempt to administer the large volumes of oral WBI solution required to be effective, it is suggested that NG instillation be instituted early in the ED course (Fig. 42.16). Unconscious patients with protected airways may receive WBI via an NG tube. In one study, patients vomited shortly after beginning WBI infusion at a rate of 1.5 to 2 L/hr. Antiemetics such as ondansetron, as well as gradually advancing the infusion rate over a 60-minute period, can help ease this side effect. Prewarming the irrigant to a temperature of approximately 37°C avoids the potential complication of hypothermia. To collect the waste products, ask an awake patient to sit on a commode. In an obtunded patient, insert a rectal tube to collect the waste. Toxicologists have recommended adding two to three bottles of activated charcoal to each liter of WBI solution. The benefit is unproved, but there is little theoretical downside to this technique and it is supported by the authors. The binding capacity of charcoal is decreased when combined with PEG-ES, but the clinical consequences of this observation are unknown and probably minimal. Empirically, metoclopramide (10 to 20 mg intravenously) may be coadministered to decrease nausea and facilitate gastrointestinal passage.

The end point of WBI is the arrival of clear rectal effluent or resolution of the toxic effect (or both). There are rare case reports of late purging of drug packets, plant parts, and tablets after the arrival of clear effluent. Radiographic studies may also be beneficial to determine the end point in body packers or in patients who have ingested radiopaque medications.

**Complications**

Few complications from WBI therapy, especially pertaining to acute poisonings, have been reported. Nausea, vomiting, abdominal cramps, bloating, and aspiration have been described. Nausea and vomiting may make administration of WBI difficult. Antiemetics and a 15- to 30-minute break followed by a slower rate may allow readministration. As discussed with the other methods of decontamination, attention should be directed to the airway and the potential for aspiration. Administration of a large amount of chilled or room-temperature WBI fluid to pediatric patients could potentially cause hypothermia. Consider warmed fluids in these patients. If activated charcoal is administered concurrently with WBI, desorption of toxin from charcoal might occur.

---

**Figure 42.16 A and B**. It is very difficult for even the most motivated patient to drink an effective volume of whole-bowel irrigation (WBI) solution. To enhance compliance and to decrease vomiting, polyethylene glycol electrolyte solution (PEG-ES) may be slowly and continuously administered via a nasogastric (NG) tube. An empty bag of saline is hung on an intravenous pole, the corner of the bag is removed, and the PEG-ES is poured into the bag. Standard intravenous tubing is connected to the proximal end of an NG tube (arrow) and the solution is infused continuously via a pump (1 L/hr). In this picture (A), charcoal has been added to the WBI solution. Metoclopramide (intravenous) was coadministered to reduce nausea, a common side effect of WBI.
DERMAL DECONTAMINATION

Background

Numerous hazardous material (HAZMAT) incidents occur each year in the United States. Although many of these events involve little morbidity or mortality, hospitals must prepare for the inevitability of caring for the chemically contaminated patient. Recent conflicts and terrorist attacks in the Middle East highlight the availability and use of chemical weapons, despite international chemical weapons treaties. The Joint Commission (TJC) requires that each US facility perform a hazard vulnerability analysis specific for the community in which that facility is located. Due to this variability in resource need for HAZMAT response, TJC gives limited guidance regarding the specifics of HAZMAT response. Guidelines from TJC involve developing an emergency operations plan (EOP) that describes how that hospital will handle chemical decontamination, as well as handle hazardous materials and waste (Standard EM.02.02.05). This leads to wide variability in HAZMAT response and decontamination practices.

Both the US Occupational Safety and Health Administration and TJC advocate that hospitals work with local emergency planning committees to best utilize community resources. In addition, TJC mandates that each hospital practice their emergency operations plan at least twice yearly, with one of these drills being a community-wide event. However, even in those communities with excellent integration of resources, resource utilization and communication during a HAZMAT incident is expected to be limited. Information regarding the event is often delayed or inaccurate, necessitating care of patients with limited knowledge of the extent or type of exposure. Whereas recommendations of specific personal protective equipment (PPE) or decontamination techniques may vary depending on the specific chemical exposure, this information is unlikely to be known in the initial stages of decontamination. Deployment of community decontamination resources to the scene may be deficient. In addition, despite the best communication, patients are expected to present to health care facilities with little or no warning following a HAZMAT incident. Those requiring more hospital resources will be interspersed with those patients with little or no chemical exposure but present for precautionary evaluation nonetheless. The end result is that in spite of preparation, in mass-casualty incidents the influx of patients will quickly overcome the health care system. Whereas some facilities may develop an EOP that utilizes local fire departments as a facility decontamination strategy, this is an unrealistic expectation in a HAZMAT incident; health care facilities must be prepared to be self-sufficient.

HAZMAT events frequently result in injuries to medical personnel. After the Tokyo attack, 13 of 15 clinicians (87%) reported symptoms while treating patients in the ED, and 23% of involved hospital staff complained of acute poisoning symptoms. Burgess and associates reported that 13% of Washington state emergency care facilities had evacuated their ED or another part of the hospital because of contamination during a 5-year period. Ghilarducci and co-workers surveyed level 1 US trauma centers and reported that only 6% had the necessary equipment required for safe decontamination. Less than 36% of emergency medicine staff had received appropriate training in handling contaminated patients, and 5.6% had experienced injuries to their staff from contact with contaminated patients during a 1-year period. It is imperative that EDs have plans in place to handle patients who have been exposed to potential toxins, provide adequate decontamination facilities, and ensure the safety of the treating medical staff.

Technique

There are a number of key components in the management of HAZMAT incidents and the care of contaminated patients seen in the ED. These components should include early recognition of a HAZMAT event, rapid activation of a plan to manage contaminated patients, initiation of primary triage, appropriate patient registration, patient decontamination, secondary triage, and final treatment.

First, the ED must be able to recognize that an event has occurred before contaminated patients gain entrance into the health care facility (Fig. 42.17, step 1). Communication with the local fire, police, and paramedic systems provides early detection of such events and allows preparation before patients arrive. Security should be arranged to prevent contaminated patients from entering the hospital, and “lockdown” of the facility should be considered.

Second, the ED should have the authority to activate a plan expeditiously to prepare the decontamination facility and allow appropriate preselected personnel to don PPE (see Fig. 42.17, step 2). If necessary, the hospital disaster plan should be activated quickly at the discretion of the ED clinician who is in contact with scene operations and incoming patients. Specific data to determine the appropriate level of PPE to maintain protection of hospital workers remain limited. The minimum PPE for hospital-based decontamination (level C) consists of a splash-proof, chemical-resistant suit with tape, double-layer protective gloves, and a powered air-purifying respirator per the National Institute of Occupational Safety and Health. Higher levels of protection, such as a level A self-contained breathing apparatus (SCBA), fully encapsulated chemical-resistant suit, or level B SCBA chemical-resistant suit, are recommended with unknown chemical and biologic exposures and for entering “hot” zones, but these levels of protection are not readily available in EDs. Fortunately, most chemical exposures are known. For those that occur in the workplace, Material Safety Data Sheets can be obtained and either the local poison center or the Agency for Toxic Substances and Disease Registry (ATSDR) can be contacted for advice on what level of protection is appropriate.

Third, appropriate primary triage should take place (see Fig. 42.17, step 3). Contaminated patients should not enter the ED until proper decontamination has occurred to ensure that the hospital staff will not be subjected to secondary contamination. Appropriate triage should then take place, with experienced personnel performing an initial brief assessment of each patient. The triage and decontamination areas should be organized into several “zones” to prevent further contamination. The hot zone is the location with the highest level of contaminant or where the incident occurred. In most cases of hospital-based decontamination there is no hot zone because patients have been removed from the initial chemical insult. On average, patients arrive at the ED 20 minutes after the event and have had significant off-gassing by this time; however, the majority of patients will have transported themselves and will not have received any prehospital decontamination by emergency medical services/HAZMAT personnel. Basic lifesaving treatments, airway and hemorrhage control, antidote administration (e.g., for cyanide or nerve agents), and
decontamination occur in the “warm” zone. The “cold” zone is safe from contaminant.111

Fourth, a brief sign-in process in the warm zone should capture the patient’s name and date of birth, with full registration to occur after decontamination. Contaminated clothing and valuables should be placed in an impervious bag to avoid potential off-gassing.

Fifth, decontamination should be performed (see Fig. 42.17, step 4). The hospital ED should have preexisting HAZMAT incident protocols that designate the decontamination area and the triage and decontamination team. Ideally, a hospital should have a permanent decontamination facility capable of handling a small number of chemically exposed patients and, in addition, a large portable unit for mass casualties. The decontamination area should meet several qualifications: (1) it should be secured to prevent spread to other areas of the hospital, (2) the ventilation system should be separate from the rest of the hospital or it should be shut off to prevent airborne spread of contaminants, and (3) provisions must be made to collect the rinse water from contaminated patients to prevent contamination of the facility and water supply. At most facilities, the best place to begin initial treatment and evaluation is outdoors. Portable decontamination facilities are available, but their cost may be prohibitive for many institutions. A practical alternative is to have a warm shower nozzle, soap, and a wading pool available outside the entrance to the ED. A tent or screen can provide privacy.

The first priority in decontaminating patients is to remove their clothing while both maintaining privacy and preventing hypothermia. This step is the most important in the decontamination process and can reduce a significant level of contaminant. Clothes should be cut off rather than pulling them
off if possible. All clothing and valuables should be placed in labeled bags. Solids should be brushed off with a soft brush or towel.

Water irrigation has been shown to be efficacious in numerous clinical trials, it is widely available, and it is inexpensive. Skin should be irrigated with copious amounts of warm water and cleansed with soap. Although some agents (e.g., metallic sodium, potassium, cesium, and rubidium) may react with water, it is still more beneficial to immediately irrigate and decontaminate the skin than to delay treatment. Scrubbing with a soft surgical sponge is advocated, while being careful to not abrade the skin. Wounds should be irrigated for an additional 5 to 10 minutes with water or saline. Remove contact lenses and irrigate the eyes for 10 to 15 minutes with saline. Direct irrigation away from the medial canthus to avoid forcing contaminants into the lacrimal duct. With strongly alkaline substances, irrigate for longer times. Irrigate the nares and ear canals, with frequent suctioning if contamination is suspected. Clean underneath the fingernails with a brush. Avoid using stiff brushes and abrasives because they may enhance dermal absorption of the toxin and can produce skin lesions that may be mistaken for chemical injuries. Sponges and disposable towels are effective alternatives.

Products that do not require rinsing constitute a new approach in cases of emergency situations with limited access to water. There are neutralizing products already deployed in various militaries for this exact purpose. For example, the Canadian and the US militaries use Reactive Skin Decontamination Lotion (Emergent BioSolutions, Inc., Rockville, MD), a solution of potassium 2,3-butanediol monoximate and diacetylmonoxime (2,3-butanediol monoxime) dissolved in a solvent of polyethylene glycol monomethyl ether and water and absorbed on a synthetic pad. It is a water miscible lotion, distributed as a sealed pack containing a single-use pad. This product has been shown to neutralize many chemical warfare agents and toxic industrial chemicals. As chemicals are rendered inactive, used pads can be directly disposed of in traditional trash, avoiding the problem of large volumes of contaminated irrigation water. Neutralizing products have been shown to be superior to traditional decontamination methods in head-to-head trials.

Secondary triage should occur after decontamination. Transfer patients with major or moderate injuries to areas designated for such cases. Send patients with minor or no injuries to appropriate holding areas for further evaluation. Medical care at this stage depends on the toxin to which the patient has been exposed and the potential toxicity of that agent. Wounds, after copious irrigation, may need thorough exploration and possibly surgical removal of the contaminant.

For the ED to care for contaminated patients, protocols should be in place and be regularly rehearsed by the facility. Train staff in the procedures and protocols, establish communication between community agencies and hospitals, regularly inspect equipment, and rehearse setups. Obtain template protocols from both the peer-reviewed medical literature and the government literature if needed. For example, guidelines for managing HAZMAT incidents are available from ATSDR. In addition, prompted by the 2001 terrorist attacks, the US Department of Veterans Affairs and several policy experts have developed a “comprehensive hospital-wide emergency mass casualty decontamination program.” This program has been applied at most Veterans Affairs medical centers and has been demonstrated to be a cost-effective protocol, suitable for implementation at other US hospitals.


Paracentesis and diagnostic peritoneal lavage (DPL) constitute the two primary intraperitoneal procedures. They are fundamentally similar in purpose and design; however, the former is generally reserved for medical concerns and the latter for evaluation of traumatic pathology.

DPL

Root and colleagues introduced DPL in 1964.1 It has withstood the passage of several decades and remains a useful diagnostic adjunct for the management of penetrating torso trauma. Following a blunt mechanism of injury, its greatest utility is as a triage tool in the assessment of hemodynamically unstable, multiply injured patients. The intent is to rapidly discover or exclude the presence of intraperitoneal hemorrhage (IPH). The advent and availability of ultrasound (US) in the emergency department (ED) have rendered this purpose complementary to that of US in the diagnostic armamentarium of emergency clinicians evaluating blunt trauma patients.2

Though commonly referred to as diagnostic peritoneal lavage, this procedure has two distinct components: peritoneal aspiration and peritoneal lavage. Peritoneal aspiration, in which an attempt is made to retrieve free intraperitoneal blood, precedes lavage. A finding of intraperitoneal blood is a marker for intraperitoneal organ injury and obviates the need for subsequent lavage. In the lavage portion, normal saline is introduced by catheter into the peritoneal cavity, recovered by gravity, and analyzed for evidence of significant intraperitoneal injury.

Peritoneal lavage can be used as a therapeutic tool in patients with hypothermia and as a means of removing toxins.3 It has also been used as a diagnostic instrument for suspected intraabdominal infection and nontraumatic sources of hemorrhage.3 4 Although the steps of the procedure are the same regardless of the indication, the primary use of DPL is to determine the need for laparotomy after trauma, and this chapter focuses on that indication.

Indications

**Blunt Trauma**

Before the advent of computed tomography (CT) and US, DPL was the sole diagnostic option to supplement physical examination for predicting the need for operative intervention (Table 43.1). It was integral to both reduction of unnecessary laparotomies and discovery of unsuspected and life-threatening intraabdominal hemorrhage in patients with significant closed-head injury.5 6

This procedure may be undertaken by the emergency physician, but its use is not mandated nor considered standard of care, and is often relegated to the trauma service or consulting surgeon. In a number of respected centers in the United States, DPL continues to be a focal diagnostic instrument.2 6 It serves two primary functions.9 First, it can be used to rapidly determine or exclude the presence of IPH. Thus, a patient with a critical closed-head injury, an unstable motor vehicle crash victim with multiple potential sources of blood loss, or a patient with pelvic

---

**Review Box 43.1** Diagnostic peritoneal lavage (semi-open and closed techniques): indications, contraindications, complications, and equipment.
Clinical success in this role has been mixed, with reported regard to hollow viscus injury, it is when serial clinical evaluations cannot be performed that gut perforation leads to preventable mortality. This is especially true in patients with severe closed-head injury or high spinal cord injury, in whom physical assessment of the abdomen is quite compromised. It is for these express scenarios that some authorities recommend DPL as a readily available procedure that can be conducted rapidly in the safe confines of the ED. The ability to undertake DPL in particular or, to a lesser extent, US in a similar manner requires careful consideration of the clinical circumstances, location of equipment, and capabilities of the personnel available.

**Penetrating Trauma**

The advent of DPL was seminal in the promotion of selective management for penetrating abdominal injury. Here its role is more dominant than for blunt trauma because of the far greater likelihood of occult injury to hollow viscera and the diaphragm after a penetrating mechanism.

Instruments and missiles may penetrate the abdominal cavity via the anterior abdominal wall, flank, back, or low chest region. The intraperitoneal space is vulnerable if penetration occurs as high as the fourth intercostal space anteriorly and the sixth or seventh space laterally and posteriorly because the diaphragm may rise to these levels in the expiratory phase of respiration. Coincident thoracic penetration occurs in up to 46% of patients with abdominal injuries. The likelihood of retroperitoneal injury increases when the entry site is over the flank or back, but the prospect of intraperitoneal pathology remains considerable, with cited incidences of up to 43% for the flank and 14% for the back.

**Stab Wounds**

Because only one fourth to one third of patients who sustain stab wounds to the anterior aspect of the abdomen require laparotomy, diagnostic algorithms are used to decrease the rate of unnecessary surgery. An optimal approach would not sacrifice sensitivity for morbid intraperitoneal injury. A pathway using a combination of clinical mandates, local wound exploration, and DPL is well established. These clinical mandates are reasonably accurate predictors of significant intraperitoneal injury (Table 43.2).

DPL fills three roles in the evaluation of patients with abdominal stab wounds: (1) rapid determination of the presence of hemoperitoneum, (2) discovery of intraperitoneal injury requiring surgery in stable patients, and (3) establishment of diaphragmatic violation. As is the case in blunt trauma patients, DPL can be invaluable as a rapid triage tool when the source of hemodynamic instability is not known. Pericardial tamponade, intrathoracic hemorrhage, and IPH may be contributory to hemodynamic instability or wholly causal. Again, as for blunt trauma evaluation, US is the only bedside diagnostic modality for IPH that is competitive in this role, and it carries the added advantage of scanning for intrapericardial and intrathoracic hemorrhage. In determining injury after stab wounds, DPL has 90% accuracy. Serial examinations CT, and laparoscopy are alternative modalities in specific circumstances and centers. The diaphragmatic rents created by stab wounds are generally small; thus, at the outset they do not

**TABLE 43.1 Clinical Indications for Laparotomy After Blunt Trauma**

<table>
<thead>
<tr>
<th>MANIFESTATION</th>
<th>PITFALL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unstable vital signs with strongly suspected abdominal injury</td>
<td>Alternative sources of shock</td>
</tr>
<tr>
<td>Unequivocal peritoneal irritation</td>
<td>Unreliable</td>
</tr>
<tr>
<td>Pneumoperitoneum</td>
<td>Insensitive; may be due to a cardiopulmonary source or invasive procedures (diagnostic peritoneal lavage, laparoscopy)</td>
</tr>
<tr>
<td>Evidence of diaphragmatic injury</td>
<td>Nonspecific</td>
</tr>
<tr>
<td>Significant gastrointestinal bleeding</td>
<td>Uncommon, unknown accuracy</td>
</tr>
</tbody>
</table>


fractures and retroperitoneal hemorrhage can be appropriately routed to lifesaving laparotomy. Furthermore, given its exquisite sensitivity, a negative peritoneal aspiration allows the clinician to proceed to alternative management steps and the patient to forego unnecessary laparotomy. Second, DPL has been used in less exigent circumstances as a means of predicting solid or hollow visceral injury requiring laparotomy. However, in this venue its sensitivity to the presence of hemorrhage may prompt unnecessary laparotomy in patients with self-limited lacerations of the liver, spleen, or mesentery. CT specifically evaluates all intraperitoneal structures, as well as the retroperitoneum, a region inaccessible to DPL. Because the revolution and the speed with which it can be undertaken have vastly improved, CT has become an invaluable adjunct in the management of blunt trauma and has largely replaced DPL in stable patients. It is most useful in identifying injury to solid organs with accompanying IPH and greatly assists nonoperative management of these injuries. The ability of CT to discern hollow viscous and pancreatic pathology has continued to improve as the modality has evolved. With regard to hollow viscous injury, it is when serial clinical evaluations cannot be performed that gut perforation leads to preventable mortality. This is especially true in patients with severe closed-head injury or high spinal cord injury, in whom physical assessment of the abdomen is quite compromised. It is for these express scenarios that some authorities recommend the performance of DPL. The clinician’s concern for hollow viscous injury should be heightened if US or CT demonstrates minimal amounts of free intraperitoneal fluid without evidence of solid organ damage.

Two paradigms have brought US to the forefront. First, this modality has been adopted as the primary triage instrument, in lieu of DPL, for the detection of IPH on the basis of identifying which pouches and gutters are filled with fluid. Clinical success in this role has been mixed, with reported sensitivity for IPH of 65% to 95%. In addition, to be useful in this role, a competent technician and interpreter and the appropriate equipment must be present in real time. It has been demonstrated that emergency clinicians and surgeons can be trained in this technique to a level of competence sufficient for this need. In centers that rely on US, DPL should serve as a reliable study when US equipment is unavailable, performance of US is technically difficult, or the results of US are indeterminate, especially when the patient demonstrates hemodynamic compromise.

DPL is a readily available procedure that can be conducted rapidly in the safe confines of the ED. The ability to undertake CT in particular or, to a lesser extent, US in a similar manner requires careful consideration of the clinical circumstances, location of equipment, and capabilities of the personnel available (Fig. 43.1).

The advent of DPL was seminal in the promotion of selective management for penetrating abdominal injury. Here its role is more dominant than for blunt trauma because of the far greater likelihood of occult injury to hollow viscera and the diaphragm after a penetrating mechanism.

Instruments and missiles may penetrate the abdominal cavity via the anterior abdominal wall, flank, back, or low chest region. The intraperitoneal space is vulnerable if penetration occurs as high as the fourth intercostal space anteriorly and the sixth or seventh space laterally and posteriorly because the diaphragm may rise to these levels in the expiratory phase of respiration. Coincident thoracic penetration occurs in up to 46% of patients with abdominal injuries. The likelihood of retroperitoneal injury increases when the entry site is over the flank or back, but the prospect of intraperitoneal pathology remains considerable, with cited incidences of up to 43% for the flank and 14% for the back.

Because only one fourth to one third of patients who sustain stab wounds to the anterior aspect of the abdomen require laparotomy, diagnostic algorithms are used to decrease the rate of unnecessary surgery. An optimal approach would not sacrifice sensitivity for morbid intraperitoneal injury. A pathway using a combination of clinical mandates, local wound exploration, and DPL is well established. These clinical mandates are reasonably accurate predictors of significant intraperitoneal injury (Table 43.2).

Thus, the presence of one or more mandates the need for urgent laparotomy and precludes the undertaking of other diagnostic studies.

DPL fills three roles in the evaluation of patients with abdominal stab wounds: (1) rapid determination of the presence of hemoperitoneum, (2) discovery of intraperitoneal injury requiring surgery in stable patients, and (3) establishment of diaphragmatic violation. As is the case in blunt trauma patients, DPL can be invaluable as a rapid triage tool when the source of hemodynamic instability is not known. Pericardial tamponade, intrathoracic hemorrhage, and IPH may be contributory to hemodynamic instability or wholly causal. Again, as for blunt trauma evaluation, US is the only bedside diagnostic modality for IPH that is competitive in this role, and it carries the added advantage of scanning for intrapericardial and intrathoracic hemorrhage. In determining injury after stab wounds, DPL has 90% accuracy. Serial examinations CT, and laparoscopy are alternative modalities in specific circumstances and centers. The diaphragmatic rents created by stab wounds are generally small; thus, at the outset they do not.

**TABLE 43.1 Clinical Indications for Laparotomy After Blunt Trauma**

<table>
<thead>
<tr>
<th>MANIFESTATION</th>
<th>PITFALL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unstable vital signs with strongly suspected abdominal injury</td>
<td>Alternative sources of shock</td>
</tr>
<tr>
<td>Unequivocal peritoneal irritation</td>
<td>Unreliable</td>
</tr>
<tr>
<td>Pneumoperitoneum</td>
<td>Insensitive; may be due to a cardiopulmonary source or invasive procedures (diagnostic peritoneal lavage, laparoscopy)</td>
</tr>
<tr>
<td>Evidence of diaphragmatic injury</td>
<td>Nonspecific</td>
</tr>
<tr>
<td>Significant gastrointestinal bleeding</td>
<td>Uncommon, unknown accuracy</td>
</tr>
</tbody>
</table>

create apparent clinical or radiologic abnormalities. However, morbidity from delayed herniation of bowel is common and substantive. Physical examination is notoriously insensitive, and DPL is currently the most sensitive means of discerning this injury in the immediate posttrauma phase. There is some evidence that coronal reconstruction of CT images provides greater sensitivity for detecting small diaphragmatic tears, and as CT technology continues to evolve, it may surpass DPL for evaluation of these subtle injuries. For these small wounds, magnetic resonance imaging may be diagnostic, but because of safety and accessibility concerns, it should be reserved for the nonacute phase of management. Laparoscopy has demonstrated promise in experienced hands.

**Gunshot Wounds**

Injury to multiple organs is the rule after gunshot wounds, and mortality is significantly greater than after stab wounds. The diagnostic approach is more conservative for gunshot wounds because in some studies the likelihood of intraperitoneal injury requiring operative intervention has exceeded 90% when the projectile has entered the intraperitoneal cavity (Fig. 43.3). If clinical mandates are met (see Table 43.2) or if peritoneal...
violation has occurred, most centers proceed to laparotomy.41 One series, however, cited intraabdominal injury in 70% to 80% of cases, thus supporting the contention that nonoperative management could be applied to a substantial percentage of patients.58 In a separate cohort of 152 patients sustaining solid organ injury from penetrating abdominal trauma (70% gunshot wounds and 30% stab wounds), 27% were successfully managed without laparotomy after selection by a protocol combining clinical examination and CT scanning.59 DPL is reserved for two circumstances: (1) the wound tract is neither obviously superficial nor intraperitoneal, and (2) penetration occurred in the low chest region, where diaphragmatic injury is more likely, yet the possibility of intraperitoneal injury also exists.

Contraindications

DPL can be undertaken in virtually any patient irrespective of age or comorbid illness. Adjustment of the technique and site of performance allows relative contraindications to be overcome. Relative contraindications include previous abdominal surgery or infections, obesity, coagulopathy, and second- or third-trimester pregnancy. The sole absolute contraindication is when clinical mandates for urgent laparotomy already exist.

Procedure

Prior to DPL, decompress the stomach and bladder to prevent inadvertent injury. Place the patient in the supine position and administer sedatives and analgesics as appropriate. Perform DPL according to compliance with standards for body fluid precautions. Observe sterile technique throughout the procedure. Before making the skin incisions described later, prepare the site with standard skin antiseptics and drape appropriately. Prophylactic antibiotics are not indicated for routine DPL because local and systemic infections are rare.60 Infiltrate the area for incision and dissection with a local anesthetic such as 1% lidocaine with epinephrine (Fig. 43.4, step 1). Delay the incision for more than 30 seconds after infiltration of local anesthetic to permit local vasospasm, which minimizes bleeding of the wound during the procedure.

---

**TABLE 43.2 Clinical Indications for Laparotomy After Penetrating Trauma**

<table>
<thead>
<tr>
<th>MANIFESTATION</th>
<th>PREMISE</th>
<th>PITFALL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemodynamic instability</td>
<td>Major solid visceral or vascular injury</td>
<td>Thorax, mediastinum</td>
</tr>
<tr>
<td>Peritoneal signs</td>
<td>Intraperitoneal injury</td>
<td>Unreliable, especially immediately after injury</td>
</tr>
<tr>
<td>Evisceration</td>
<td>Additional bowel, other injury</td>
<td>No injury in one fourth to one third of stab wound cases</td>
</tr>
<tr>
<td>Diaphragmatic injury</td>
<td>Diaphragmatic herniation</td>
<td>Rare clinical, radiographic findings</td>
</tr>
<tr>
<td>Gastrointestinal and vaginal hemorrhage</td>
<td>Proximal gut or uterine injury</td>
<td>Uncommon, unknown accuracy</td>
</tr>
<tr>
<td>Impalement in situ</td>
<td>Vascular impalement</td>
<td>High operative risk, pregnancy</td>
</tr>
<tr>
<td>Intraperitoneal air</td>
<td>Perforation of a hollow viscus</td>
<td>Insensitive; may be caused by intraperitoneal entry only or be due to a cardiopulmonary source</td>
</tr>
</tbody>
</table>


![ABDOMINAL GUNSHOT WOUND ALGORITHM](image)

Figure 43.3 Algorithm for abdominal gunshot wounds. *Can be assessed by the path of the missile, plain films, local wound exploration, ultrasonography, and LAP. †Most centers proceed to LAP if peritoneal entry is suspected. ‡Patients with documented superficial and low-velocity injuries can be discharged; high-velocity injuries or those of unknown depth require further tests or observation. ‡DPL, CT, LPY, or SPEs can be used in singular or complementary fashion depending on the clinical scenario. Expectant management of injuries caused by gunshot wounds is rarely attempted. CT, Computed tomography; DPL, diagnostic peritoneal lavage; LAP, laparotomy; LPY, laparoscopy; SPE, serial physical examination. (From Marx J, Isenhour J: Abdominal trauma. In Marx JA, Hockberger RS, Walls RM, et al, editors: Rosen's emergency medicine: concepts and clinical practice, ed 6, St. Louis, 2006, Mosby p 506.)
Diagnostic Peritoneal Lavage: Semi-Open Technique

1. Anesthetize the skin with lidocaine with epinephrine.

2. Make a 4- to 6-cm infra-umbilical midline skin incision with a No. 11 scalpel.

3. Using Army-Navy retractors, bluntly dissect to the linea alba.

4. Make a small 2- to 3-mm opening in the linea alba with a No. 15 scalpel blade.

5. Place towel clips through this opening to grasp and lift each side of the rectus.

6. Advance the catheter and trocar 45° to 60° caudally into the peritoneum.

7. Retract the trocar within the catheter, and advance the catheter toward the pelvis.

8. Withdraw the trochar and aspirate for blood using a 10-cc syringe.

9. If lavage is necessary, infuse 1 L of warm normal saline and recover fluids under gravity.

Figure 43.4 Semi-open technique of diagnostic peritoneal lavage. Note that the stomach and bladder have been decompressed under most circumstances.
Placement of the Catheter

There are two basic methods for DPL: open and closed. The two open techniques are semi-open and fully open, and both typically require an assistant. DPL is clearly within the diagnostic armamentarium of the emergency clinician and surgeon. Either may undertake it in keeping with clinical policies established at the particular trauma center.

Semi-open Technique

Make a skin incision 4 to 6 cm in length with a No. 11 scalpel blade. Using Army-Navy retractors, proceed with blunt dissection to expose the rectus fascia (see Fig. 43.4, steps 2 and 3). With the infraumbilical incision in the midline, continue blunt dissection until the linea alba is seen. Its crossing bands of crural fibers may be apparent.61 Make a small 2- to 3-mm opening in the linea alba with a No. 15 scalpel blade (see Fig. 43.4, step 4). You may notice a tough, gritty sensation when cutting the linea alba with the scalpel. Place towel clips through this opening to grasp each side of the rectus fascia (see Fig. 43.4, step 5). Ask an assistant to lift the two towel clips and carefully advance the catheter and trocar in a 45- to 60-degree caudal orientation. Proceed through the peritoneum into the peritoneal cavity (see Fig. 43.4, steps 6 and 7).62

One method to decrease the likelihood of penetrating any underlying viscera is to hold the fingers low on the catheter/trocator instrument such that on entering the abdominal peritoneum, the fingers prevent deep penetration. Excessive pressure during penetration with the trocar is a common error. Apply steady one-finger pressure to the handle sufficient to “pop” through the peritoneum. After controlled peritoneal penetration of 0.5 to 1.0 cm in the midline, retract the trocar 1.0 to 2.0 cm within the catheter, and advance the catheter carefully toward the pelvis. Some operators advance the catheter toward the right or left side of the pelvis. Use a slight twisting motion during advancement to minimize visceral or omental injury.

The fully open technique extends the semi-open technique by one step. Lengthen the opening in the linea alba to open the peritoneum, and use direct visualization to advance the catheter into the peritoneal cavity. The trocar is unnecessary with the open technique.

A single technician can accomplish the two open techniques, but it is useful to have an assistant help with retraction and handling of the instruments. The fully open method is the more technically demanding and time-consuming. Reserve this method for clinical circumstances in which neither the closed nor the semi-open technique is deemed safe, or they have been attempted and failed. Examples of such circumstances include pelvic fracture, pregnancy, previous abdominal surgery, adhesions, infections, and obesity.

Closed Technique

For the closed technique, introduce the catheter into the peritoneal space in a blind percutaneous fashion.63 Use the simple Seldinger (guidewire) method, in which a small-gauge guide needle is inserted into the peritoneal cavity in the midline just inferior to the umbilicus (Fig. 43.5, step 2). Pass a flexible wire through the needle (see Fig. 43.5, step 3), and remove the needle but not the wire. Make a small stab with a No. 11 scalpel blade at the entry site of the wire to allow easier passage of the catheter through the abdominal wall (see Fig. 43.5, step 4), and then advance a soft catheter over the wire and into the peritoneal cavity. Rotate the catheter while pushing it over the guidewire to facilitate entry into the peritoneal cavity if the catheter does not advance easily. Place the catheter into the right or left pelvic gutter.

Always control the guidewire to avert intraabdominal migration of the wire. Withdraw the wire and aspirate for blood with a 10-mL syringe. Follow this with peritoneal lavage when necessary. Proponents of the guidewire technique promote its ease and rapidity.64–68 Those who prefer the semi-open method argue that the time until peritoneal aspiration, the more critical interval, is minimally different and that this method may have fewer complications and thus be more accurate than the guidewire technique.69–73 Note that for both the semi-open and closed approaches, the time until aspiration is performed should be no more than 2 to 5 minutes.

Site

The optimum location for DPL is at the infraumbilical ring at the inferior border of the umbilicus (Table 43.3). Here, between the rectus abdominis muscles there is adherence of the peritoneum and relative lack of vascularity and preperitoneal fat.61 Closed DPL should always be conducted here. In the event of second- or third-trimester pregnancy, a suprauterine approach is used. If midline scarring is present, a fully open technique at the lateral border of the rectus abdominis in the left lower quadrant may be necessary. The left side is preferred to avoid later confusion about whether an appendectomy has been performed. It is interesting to note that Moore and associates found no increase in complications or misclassified lavage when the closed technique was used in a small series of patients with previous abdominal surgery.74 In the presence of a pelvic fracture, use a fully open supraumbilical approach. This greatly decreases the likelihood of passing the catheter through a retroperitoneal hematoma that has dissected from the fracture anteriorly and across the abdominal wall.75 In patients with penetrating trauma, do not perform DPL through the stab or missile entry site. This approach can contaminate the intraperitoneal cavity, potentially exacerbate the abdominal wall bleeding, and lead to a false-negative result.

Aspiration and Lavage

Once the catheter has been placed successfully into the peritoneal cavity, attach the right-angle adapter, extension tubing, and a non–Luer-Lok syringe and attempt aspiration (see Fig. 43.4, step 8). If 10 mL of blood is aspirated, the test is positive and the procedure is terminated. With penetrating

<table>
<thead>
<tr>
<th>TABLE 43.3 Preferred Site for DPL</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CLINICAL CIRCUMSTANCE</strong></td>
</tr>
<tr>
<td>-----------------------------</td>
</tr>
<tr>
<td>Standard adult</td>
</tr>
<tr>
<td>Standard pediatric</td>
</tr>
<tr>
<td>Second- and third-trimester pregnancy</td>
</tr>
<tr>
<td>Midline scarring</td>
</tr>
<tr>
<td>Pelvic fracture</td>
</tr>
<tr>
<td>Penetrating trauma</td>
</tr>
</tbody>
</table>

*The stab wound or gunshot wound site should be avoided. C, Closed; DPL, diagnostic peritoneal lavage; FO, fully open; SO, semi-open.
infusion tubing (e.g., urologic irrigation tubing sets, such as the Abbott No. 6543 cystoscopy/irrigation set [Allegiance, Michigan]) also shortens fluid influx time. Infuse 1 L of fluid in adults or 15 mL/kg in children. When possible, roll or shift the patient from side to side after the infusion to increase mixing. Place the IV bag or bottle on the floor (or below abdominal level), and allow the fluid to return by gravity.

If the fluid does not return, there may be several reasons. Some IV tubing contains a one-way valve. If tubing with a

trauma, acquisition of lesser amounts may be meaningful because of the tendency for the diaphragm and bowel to hemorrhage minimally when injured. However, no rules have been established in this regard.

If little to no blood is aspirated, lavage the peritoneal cavity with either normal saline or lactated Ringer’s solution (see Fig. 43.4, step 9). Apply a blood pressure cuff or blood infusion pump around the plastic intravenous (IV) bag to speed the influx (i.e., decrease lavage time) if necessary. Large-bore infusion tubing (e.g., urologic irrigation tubing sets, such as the Abbott No. 6543 cystoscopy/irrigation set [Allegiance, Michigan]) also shortens fluid influx time. Infuse 1 L of fluid in adults or 15 mL/kg in children. When possible, roll or shift the patient from side to side after the infusion to increase mixing. Place the IV bag or bottle on the floor (or below abdominal level), and allow the fluid to return by gravity.

If the fluid does not return, there may be several reasons. Some IV tubing contains a one-way valve. If tubing with a

Figure 43.5 Closed technique of diagnostic peritoneal lavage. Note that the stomach and bladder are decompressed under most circumstances.
Complications

Local and Systemic
Local wound complications, including infection, hematoma, and dehiscence, occurred in only 0.3% of patients in two large series. Dehiscence with evisceration is an even rarer condition. Systemic infection has been described rarely (Table 43.4).

Intraperitoneal
The trocar, wire, and rarely, the catheter can inflict iatrogenic intraperitoneal injury. Virtually any structure in the peritoneal cavity can be breached, including the small and large bowel, the bladder, and major vessels. Typically, if the needle is the culprit and even if the trocar is responsible, injury to these structures is minimal and self-limited, and observation of the patient is sufficient.

Technical Failure
Inability to recover peritoneal aspirate or lavage fluid can result in a false-negative interpretation. This can occur in several circumstances. It follows unwitting placement of the catheter into the preperitoneal space, which is less likely to occur with either open technique. Compartmentalization of fluid by adhesions or obstructing omentum can impede the egress of fluid. When a fully open supraumbilical or suprauterine technique is used, the catheter may be too short to access the depths of the intraperitoneal cavity. Finally, the large diaphragmatic tears typical of blunt pathophysiology allow flow of lavage fluid from the intraperitoneal to the thoracic cavity. Saunders and coworkers compared percutaneous DPL and the open technique in a prospective, randomized trial. Fluid obtained by the two techniques had similar test performance for intraabdominal pathology. The open technique took, on average, more than 4 minutes longer, but the percutaneous approach had an 11.2% technical failure rate (versus 3.8% with the open approach).

False-positive findings can occur in two ways. First, iatrogenic misadventure may be responsible. Second, in penetrating trauma, particularly stab wounds, bleeding from the abdominal wall injury site into the peritoneal cavity can lead to positive findings when no injury to intraperitoneal structures has occurred.

Interpretation

Gross Blood
Recovery of 10 mL or more of blood via aspiration is considered a positive finding. Aspirates with lesser volume are generally discarded and are not factored into analysis of the lavage fluid. Grossly bloody aspirates are typically indicative of solid visceral or vascular injury, with a positive predictive value of greater than 90%. Aspiration of blood is responsible for approximately 80% of true-positive DPL findings with blunt trauma and for 50% with stab wounds.

A positive aspiration in a blunt trauma patient who is hemodynamically stable or has been resuscitated to apparent

### Table 43.4: Complications of DPL

<table>
<thead>
<tr>
<th>CATEGORY</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local and Systemic</td>
<td></td>
</tr>
<tr>
<td>Hematoma incision site</td>
<td>Local wound care</td>
</tr>
<tr>
<td>Dehiscence incision site</td>
<td>Local wound care</td>
</tr>
<tr>
<td>Local wound infection</td>
<td>As indicated</td>
</tr>
<tr>
<td>Systemic infection</td>
<td>As indicated</td>
</tr>
<tr>
<td>Intraperitoneal Injury</td>
<td></td>
</tr>
<tr>
<td>Bowel</td>
<td>Observe, usually self-limited</td>
</tr>
<tr>
<td>Bladder</td>
<td>Observe, usually self-limited</td>
</tr>
<tr>
<td>Vascular</td>
<td>Observe, usually self-limited</td>
</tr>
<tr>
<td>Technical Failure</td>
<td></td>
</tr>
<tr>
<td>Inability to Recover Fluid*</td>
<td>Repeat DPL</td>
</tr>
<tr>
<td>Preperitoneal catheter placement</td>
<td></td>
</tr>
<tr>
<td>Compartmentalization of fluid</td>
<td>US, CT</td>
</tr>
<tr>
<td>Obstructed catheter</td>
<td>Gentle catheter manipulation</td>
</tr>
<tr>
<td>Diaphragm injury</td>
<td>Reverse Trendelenburg; consider US, CT</td>
</tr>
<tr>
<td>“Short” catheter (supraumbilical or suprauterine approach)</td>
<td>Trendelenburg</td>
</tr>
<tr>
<td>Intraperitoneal Hemorrhage*</td>
<td></td>
</tr>
<tr>
<td>Iatrogenic injury</td>
<td>As indicated by clinical markers</td>
</tr>
<tr>
<td>Stab wound, abdominal wall bleeding</td>
<td>As indicated by clinical markers</td>
</tr>
<tr>
<td>Pelvic fracture (RBC count)</td>
<td>Complementary CT</td>
</tr>
</tbody>
</table>

*May lead to false-negative DPL results.

*May lead to false-positive DPL results.

CT, Computed tomography; DPL, diagnostic peritoneal lavage; RBC, red blood cell; US, ultrasound.
angiography and embolization will not be delayed unnecessarily should active intraperitoneal bleeding not be found (Fig. 43.6).

WBC Count

An inflammatory peritoneal response to a multitude of stimuli can occur, including stool, blood, and enzymes.\(^ \text{99} \) The white

---

### TABLE 43.5 Diagnostic Peritoneal Lavage

<table>
<thead>
<tr>
<th></th>
<th>POSITIVE</th>
<th>INDETERMINATE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blunt trauma</strong> (RBC/mm(^3))</td>
<td>100,000(^a)</td>
<td>20,000–100,000</td>
</tr>
<tr>
<td><strong>Stab wound</strong> (RBC/mm(^3))</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anterior abdomen</td>
<td>100,000</td>
<td>20,000–100,000</td>
</tr>
<tr>
<td>Flank</td>
<td>100,000</td>
<td>20,000–100,000</td>
</tr>
<tr>
<td>Back</td>
<td>100,000</td>
<td>20,000–100,000</td>
</tr>
<tr>
<td>Low chest</td>
<td>5000–10,000</td>
<td>1000–5000</td>
</tr>
<tr>
<td><strong>Gunshot wound</strong> (RBC/mm(^3))</td>
<td>5000–10,000</td>
<td>1000–5000</td>
</tr>
<tr>
<td><strong>LAM (IU/L)</strong></td>
<td>&gt;20</td>
<td>10–19</td>
</tr>
<tr>
<td><strong>LAP (IU/L)</strong></td>
<td>≥3</td>
<td>NA</td>
</tr>
<tr>
<td><strong>WBCs (per mm(^3))</strong></td>
<td>&gt;500</td>
<td>250–500</td>
</tr>
</tbody>
</table>

\(^a\)In a hemodynamically stable patient with a pelvic fracture and a positive or equivocal red blood cell count, computed tomography should be obtained to corroborate or refute intraperitoneal injury.

DPL, Diagnostic peritoneal lavage; LAM, lavage amylase; LAP, lavage alkaline phosphatase; NA, not applicable; RBC, red blood cell; WBCs, white blood cells.


stability need not mandate urgent surgery. Unnecessary laparotomy will occur if there has been minimal and self-limited damage to the liver, spleen, bowel serosa, or mesentery.\(^ \text{91} \) In this situation, CT and clinical indicators should be used in concert with the findings on DPL.

**RBC Count**

The recommended red blood cell (RBC) threshold varies according to the mechanism and, in the case of stab wounds, the external site of injury (Table 43.5). The optimum criterion will deliver excellent sensitivity, a high positive predictive value, and a minimal incidence of unnecessary laparotomy. Negative laparotomy incurs a prolongation of hospitalization and increases the cost of care, in addition to creating the potential for procedural complications.\(^ \text{82,83} \) RBC counts greater than 10\(^5\) \(\mu\)L are generally considered positive with a blunt mechanism or after stab wounds in the anterior part of the abdomen, flank, or back. Counts of 20,000 to 100,000/mm\(^3\) should be considered indeterminate.\(^ \text{41,45,84,85} \) For stab wounds in the low chest region, the diaphragm at increased risk for injury, the RBC criterion should be lowered to 5000/mm\(^3\) to maximize sensitivity for isolated injury to this structure.\(^ \text{16,43,86,87} \) With gunshot wounds involving the abdomen or low chest region, the same RBC criterion of 5000/mm\(^3\) is applied. This is intended to increase the sensitivity of the test because intraperitoneal entry by a missile carries a 90% or greater likelihood of intraperitoneal injury.\(^ \text{16,60,88} \) An uncomplicated DPL should not result in more than several hundred to several thousand RBCs in the peritoneal lavage fluid.

The incidence of false-positive RBC interpretation in the setting of pelvic fracture is considerable. However, aspiration of free blood in patients with critical pelvic fractures predicts active IPH in more than 80% of cases. A positive RBC count should generally prompt corroboration or refutation of intraperitoneal injury by CT. In this fashion, needed pelvic angiography and embolization will not be delayed unnecessarily should active intraperitoneal bleeding not be found (Fig. 43.6).
blood cell (WBC) count in lavage effluent was formerly touted to predict small bowel injury but has since been proved unreliable. It is insensitive in the immediate postinjury period because 3 to 5 hours is necessary before the test becomes positive (see Table 43.5). Moreover, a positive finding is likely to be falsely so. Therefore, the WBC level in and of itself should not determine the need for laparotomy.

### Enzymes
Alkaline phosphatase is contained intramurally in the small bowel, as well as in hepatobiliary secretions released into the proximal part of the intestine. Amylase is contained in the latter only. Perforation of the small bowel allows access of these two markers to the peritoneal cavity, where they can be recovered by peritoneal lavage. Although levels of the two markers usually rise in tandem, lavage amylase has been shown to be a more accurate marker than lavage alkaline phosphatase (see Table 43.5). In contradistinction to the WBC count, these tests will be positive in the immediate postinjury period. However, they may not be economical if used on a mandatory rather than a selective basis. Neither is helpful in discerning the presence of pancreatic pathology.

### Miscellaneous
Routine bile staining, Gram staining, and microscopy to identify vegetable fibers are rarely productive and are of untested accuracy. Deck and Porter reported that finding urine in the lavage fluid, as evidenced by a straw color, and creatinine in the peritoneal fluid should suggest an intraperitoneal bladder or collecting system injury.

### Conclusion
In the current era of readily available advanced imaging techniques such as CT and US, DPL maintains a diminished, but important role in the evaluation of injured patients. It can identify life-threatening IPH in unstable patients when US is unavailable, indeterminate, or negative for free fluid. In addition, DPL can identify hollow viscus injury when CT is nondiagnostic in patients in whom serial clinical evaluations are impractical or unreliable, as in those with severe traumatic brain and spinal cord injuries. Finally, DPL remains more accurate than CT in identifying occult diaphragmatic injuries and is useful in high-risk patients without other indications for surgical abdominal exploration. It should be used in commonsense fashion. The noted laboratory parameters are guidelines and should not be embraced to the exclusion of pertinent clinical features. Optimal strategies depend largely on the capability of an institution’s resources and personnel in each clinical scenario.

### PARACENTESIS
Therapeutic abdominal paracentesis, the removal of intraperitoneal fluid, is one of the oldest medical procedures and dates to approximately 20 BC. Paracentesis was first described in the modern medical literature by Saloman at the beginning of the 20th century, and it became a valued decompressive therapy. With the advent of diuretics in the early 1950s, paracentesis fell out of favor as a treatment option. Controlled clinical trials in the late 1980s up to the present have restored its reputation by demonstrating the safety and efficacy of large-volume paracentesis (LVP) in adults and children. Because this mode is invasive and consumes clinician hours, it is generally reserved for the treatment of patients with chronic ascites who have tense ascites or whose condition is refractory to diuretic therapy. However, paracentesis (Videos 43.1 and 43.2) remains an important diagnostic agent for patients with new-onset ascites or to determine the presence of worrisome conditions, notably infection, in those with preexistent ascites.

### Clinical Features
#### Determination of Ascites
Small amounts of ascites may be asymptomatic. Larger collections typically cause a sense of abdominal fullness, anorexia, early satiety, and perhaps nausea and abdominal pain. Substantial accumulations create symptoms of respiratory distress by restricting lung capacity. The most predictive history and physical examination findings for excluding the diagnosis of ascites are the absence of ankle swelling and increased abdominal girth and an inability to demonstrate bulging flanks, flank dullness, or shifting dullness. Positive predictors for the diagnosis are a positive fluid wave, shifting dullness, or peripheral edema, with flank dullness having the greatest sensitivity at 94% and positive fluid wave having the greatest specificity.

Patients who lack obvious clinical markers may benefit from the performance of US, which can discern the presence of as little as 100 mL of fluid. Endoscopically guided US may detect as little as 10 mL. It is more sensitive than CT in this respect and can assist in the identification of malignancy. In addition, it is a useful adjunct for determining the location of fluid that may be compartmentalized by preexistent infection or surgical adhesions.

#### Differential Diagnosis
Causes of ascites can be categorized in several ways. On a structural basis, the causes are divided into diseases of the peritoneum and diseases not involving the peritoneum. The former group includes infection, neoplasm, collagen vascular disease, and idiopathic causes. The latter includes cirrhosis, congestive heart failure, nephrotic syndrome, protein-losing enteropathy, malnutrition, myxedema, pancreatic disease, ovarian disease, chylous effusion, Budd-Chiari syndrome, and hepatic venous occlusive disease. Pathophysiologic categories are listed in Box 43.1. In the United States, parenchymal liver pathology is overwhelmingly the most likely cause. Within this group, alcoholic liver disease is responsible for approximately 80% of cases (Table 43.6). Finally, ascites can be classified on the basis of a serum-ascites albumin gradient, that is, the difference between albumin values obtained simultaneously from serum and ascites samples (Box 43.2).

### Indications and Contraindications
Therapeutic paracentesis is often undertaken in the ED setting to relieve the cardiorespiratory and gastrointestinal manifestations of tense ascites. LVP, or removal of more than 5 L, ameliorates the shortness of breath and early satiety that these patients experience. It may also be associated with collateral advantages, such as a reduction in hepatic venous pressure gradients, intravariceal pressure, and variceal wall tension. These parameters are considered important predictors of variceal bleeding, and the improvement after LVP may decrease
Abdominal Paracentesis

**Indications**
- New onset ascites
- Suspected spontaneous bacterial peritonitis
- To relieve the cardiorespiratory and gastrointestinal manifestations of tense ascites

**Contraindications**
- Uncorrected coagulopathy AND clinically evident fibrinolysis or disseminated intravascular coagulation
- Bowel dilation or obstruction
- Pregnancy (technique should be altered as noted below)
- Abdominal hematoma, engorged veins, or superficial infection at puncture site

**Complications**

<table>
<thead>
<tr>
<th>Systemic</th>
<th>Local</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyponatremia</td>
<td>Persistent ascitic fluid leak</td>
</tr>
<tr>
<td>Renal dysfunction</td>
<td>at the wound site</td>
</tr>
<tr>
<td>Hepatic encephalopathy</td>
<td>Abdominal wall hematoma</td>
</tr>
<tr>
<td>Hemodynamic compromise</td>
<td>Localized infection</td>
</tr>
<tr>
<td>Significant bleeding</td>
<td></td>
</tr>
<tr>
<td>Death</td>
<td></td>
</tr>
</tbody>
</table>

**Intraperitoneal**
- Perforation of vessels and viscera
- Generalized peritonitis
- Abdominal wall abscess

**Review Box 43.2** Abdominal paracentesis: indications, contraindications, complications, and equipment.

---

**Box 43.1** Pathophysiologic Classification of Ascites

I. Elevated hydrostatic pressure
   - A. Cirrhosis
   - B. Congestive heart failure
   - C. Constrictive pericarditis
   - D. Inferior vena cava obstruction
   - E. Hepatic vein obstruction (Budd-Chiari syndrome)

II. Decreased osmotic pressure
   - A. Nephrotic syndrome
   - B. Protein-losing enteropathy
   - C. Malnutrition
   - D. Cirrhosis or hepatic insufficiency

III. Fluid production exceeding resorptive capacity
   - A. Infections
     1. Bacterial
     2. Tuberculosis
     3. Parasitic
   - B. Neoplasms

---

**Table 43.6 Causes of Ascites**

<table>
<thead>
<tr>
<th>Cause</th>
<th>Patients (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parenchymal liver disease</td>
<td>80</td>
</tr>
<tr>
<td>Mixed</td>
<td>5</td>
</tr>
<tr>
<td>Malignancy</td>
<td>10</td>
</tr>
<tr>
<td>Heart failure</td>
<td>5</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>2</td>
</tr>
<tr>
<td>Pancreatic</td>
<td>1</td>
</tr>
<tr>
<td>Nephrogenous (“dialysis ascites”)</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Chlamydia</td>
<td>&lt;1</td>
</tr>
<tr>
<td>Surgical peritonitis in the absence of liver disease</td>
<td>&lt;1</td>
</tr>
</tbody>
</table>

*Based on a series of 1500 paracenteses performed in a predominantly inpatient hepatology/general internal medicine setting (BA Runyon, unpublished observations).


BOX 43.2 Classification of Ascites by the Serum-Ascites Albumin Concentration Gradient

HIGH GRADIENT (≥1.1 G/DL)
- Cirrhosis
- Alcoholic hepatitis
- Cardiac ascites
- Massive liver metastases
- Fulminant hepatic failure
- Budd-Chiari syndrome
- Portal vein thrombosis
- Venous occlusive disease
- Fatty liver of pregnancy
- Myxedema
- Mixed ascites

LOW GRADIENT (<1.1 G/DL)
- Peritoneal carcinomatosis
- Tuberculous peritonitis
- Pancreatic ascites
- Biliary ascites
- Nephrotic syndrome
- Serositis in connective tissue diseases


the risk for bleeding. Diagnostic paracentesis, often relegated to inpatient services, is indicated in any patient whose ascites is of new onset or to disclose the presence of infection in patients with known or suspected ascites, particularly in the context of alcohol-related cirrhotic liver disease.\textsuperscript{117,118} Diagnostic paracentesis is also useful in the management of patients with acquired immunodeficiency syndrome (AIDS), in whom the etiology of ascites will be non–AIDS related in three-quarters of cases.\textsuperscript{119} There are few relative contraindications to abdominal paracentesis. Certain systemic and anatomic risks should be considered, however.

Systemic
Given the predominance of alcohol-related cirrhotic liver disease as the cause of ascites, as many as two thirds to three quarters of patients who undergo paracentesis will have a coagulopathy. However, the only prospective study that evaluated the complications of paracentesis determined that transfusion-requiring abdominal hematomas occurred in less than 1% of cases despite the fact that 71% of the patients had an abnormal prothrombin time (PT).\textsuperscript{120} Because transfusion-requiring hematoma is so unlikely, even in this population, prophylactic administration of fresh frozen plasma or platelets is not standard, nor mandated, and imposes considerable cost, in addition to the risk for posttransfusion hepatitis, with little net gain.\textsuperscript{121} Therefore, for patients undergoing repeated therapeutic paracentesis, in the absence of previous problems or obvious clotting issues, obtaining a platelet count and international normalized ratio (INR) before the procedure is not routine. In an investigation of 628 patients undergoing outpatient LVP, the procedure was safely performed when the INR was as high as 8.7 (mean value, 1.7) and the platelet count as low as 19,000/mm\textsuperscript{3} (mean value, 50,000/mm\textsuperscript{3}).\textsuperscript{122} These data countermand older and more conservative recommendations to administer platelets to patients with levels of less than 50,000/mm\textsuperscript{3} or to give fresh frozen plasma to those with a PT exceeding 20 seconds (1.5 times the therapeutic level).\textsuperscript{123} These blood products should be reserved for clinically evident fibrinolysis and disseminated intravascular coagulation.

Anatomic
Structural impediments to the safe introduction of a paracentesis needle can include the bladder, bowel, and pregnant uterus. The bladder is normally tucked into the recess of the pelvis. However, bladders that are neuropathologically distended as a result of pharmacologic agents or medical conditions should preferably be emptied by voiding or by catheterization to avoid puncture. The intestines typically float in ascitic fluid and will move safely away from a slowly advancing paracentesis needle.\textsuperscript{123} Therefore, US guidance may be indicated in cases of suspected adhesions or bowel obstruction. Even if penetrated by an 18- to 22-gauge needle, leakage of intestinal contents will not occur unless intraluminal pressure is five- to tenfold greater than normal conditions.\textsuperscript{124} In second- and third-term pregnancy, an open supraumbilical or US-assisted approach is preferred. The abdomen should be inspected carefully for evidence of abdominal hematoma, engorged veins, or superficial infection, and these sites should be strictly avoided.

Technique

Preliminary Actions
Paracentesis should be performed after the patient has voided. It is not standard to decompress the stomach with a nasogastric tube or the bladder with a catheter before paracentesis. Place the patient in the supine position. Some clinicians prefer the supine lateral decubitus position. Perform paracentesis according to compliance with standards for body fluid precautions. Observe sterile technique throughout the procedure to prevent the iatrogenic introduction of bacteria into the abdominal wall tract or peritoneal cavity. Before making the skin incisions described later, prepare the site with standard skin antisepsics and drape appropriately.

Site of Entry
The best site of entrance for repeated paracentesis is determined by the patient’s previous experience, so this question should be asked of the patient. Theoretically, most sites on the abdominal wall can be used, but in absence of previous experience with the individual patient, two sites are preferred. One site is approximately 2 cm below the umbilicus in the midline (Fig. 43.7), where the fasciae of the rectus abdominis muscle join to form the fibrous, thin, avascular linea alba. Large collateral veins may occasionally be present and should be avoided (Fig. 43.8), as should suspected areas of skin infection. If the patient has midline scarring or if previous experience has been positive, the preferred alternative site is in either the right or left lower quadrant, approximately 4 to 5 cm cephalad and medial to the anterior superior iliac spine (see Fig. 43.7). The importance of remaining lateral to the rectus sheath is to avoid the inferior epigastric artery. Patients with a large quantity of ascites can readily undergo the procedure in the supine position with the head of the bed slightly elevated. Those with lesser amounts of fluid may benefit from a lateral decubitus position with introduction of the needle into the
midline or dependent lower quadrant (Fig. 43.9). Some clinicians prefer to use the lateral decubitus position routinely because the bowel tends to float upward and away from the path of the needle. Hence, the site of needle entrance is in the midline or on the side closest to the bed. Rarely, patients may need to be placed in a facedown, hands-on-knees position. In patients with multiple abdominal scars or suspicion of compartmentalized abdominal fluid for any reason, US guidance is prudent.

Procedure

A prepackaged kit, such as the Safe-T-Centesis system (CareFusion, Vernon Hills, IL), is a convenient way to perform the procedure (Fig. 43.10). Following sterile preparation of the skin, inject local anesthetic at the paracentesis site (Fig. 43.11, step 4). Use a standard 3.8-cm (1.5-inch) metal needle in most cases. If necessary, use a longer 8.9-cm (3.5-inch) spinal needle in obese patients. Plastic sheath cannulas tend to kink and run the risk of being sheared off into the peritoneal cavity, but a...
**Abdominal Paracentesis**

1. Localize the fluid collection by percussion or by ultrasound. Mark the entry site.

2. With ultrasound, identify free pockets of ascitic fluid, as well as organs such as the liver, spleen, kidney, or bowel. See details of ultrasound techniques at the end of this chapter.

3. Cleanse the overlying skin with antiseptic and apply a sterile drape.

4. Anesthetize the skin and the proposed tract of the paracentesis needle with local anesthetic.

5. Insert the needle perpendicular to the skin and slowly advance. See text and Fig. 43.12 for details on the Z-tract method of paracentesis.

6. Advance the needle in 5-mm increments until fluid is returned in the syringe. Advance the catheter over the needle and into the peritoneal cavity (if a catheter-over-the-needle system is being used).

7. Aspirate peritoneal fluid into a 20- to 60-mL syringe for a diagnostic sample.

8. If large-volume (therapeutic) paracentesis is desired, attach the high-pressure tubing to the catheter hub.

9. Insert the high-pressure tubing needle into the evacuated container to withdraw the fluid.

10. After fluid collection is completed, remove the catheter and apply an adhesive bandage (or a pressure dressing if there is persistent fluid leakage).

**Figure 43.11** Abdominal paracentesis.
steel needle can be left in the abdomen during a therapeutic tap for intervals of an hour or longer without injury. The 15-gauge, 3.25-inch Caldwell needle/cannula is an alternative for LVP that has been shown to perform similar to a steel needle in terms of procedural complications and the volume of fluid removed. Use a smaller-gauge (20- to 22-gauge) needle for diagnostic taps because such needles lessen the likelihood of leakage of ascitic fluid through the wound site after the procedure. However, for therapeutic LVP, use an 18-gauge needle because it permits expeditious outflow.126

Insert the needle directly perpendicular to the skin at the preferred site (see Fig. 43.11, step 5). Alternatively, use the “Z-tract” method. For this method, pull the skin approximately 2 cm caudad to the deep abdominal wall with the non–needle-bearing hand while slowly inserting the paracentesis needle (Fig. 43.12). Release the skin when the needle has penetrated the peritoneum and fluid flows. This technique also holds the draining needle in place without suture or tape. Remove the needle after the procedure, and the skin will slide to its original position, helping seal the tract. In any case, insert the needle slowly in 5-mm increments to detect undesired entry of a vessel and to help prevent unnecessary puncture of the small bowel. Avoid continuous suction because it may attract bowel or omentum to the end of the paracentesis needle with resultant occlusion. Once fluid is flowing, stabilize the needle to ensure a steady flow. If flow ceases, gently rotate the needle and advance it inward in 1- to 2-mm increments. When fluid removal is complete, remove the needle and place an adhesive bandage over the puncture site. If there is persistent leakage of fluid, a pressure bandage may be required. Cyanoacrylate adhesive can also be utilized for persistent leakage. First place the patient so the site is in a non-dependent position and apply pressure to the site with gauze for ten minutes. Then withdraw the gauze and apply the adhesive over the puncture site. Dry air applied to the adhesive using nasal cannula can improve drying of the adhesive.

Ultrasound Guidance

US-guided paracentesis may be performed by a radiologist or an experienced emergency clinician ultrasonographer (see Ultrasound Box 43.1). This technique clearly delineates the pocket of ascitic fluid and allows visualization of loculated collections and avoidance of bowel adherent to the anterior abdominal peritoneum. The ultrasonographer scans the abdomen and marks the skin at the point overlying the optimal puncture site (see Fig. 43.11, step 2). Once the entry site is marked, keep the patient immobile and perform the procedure (as detailed previously) as soon as practical to avoid shifting of the fluid, which may decrease the utility of US guidance.

![Figure 43.12 A](image1.png)  Z-tract method of needle paracentesis. Pull the skin approximately 2 cm caudad in relation to the deep abdominal wall with the non–needle-bearing hand while slowly inserting the paracentesis needle perpendicular to the skin.  

![Figure 43.12 B](image2.png)  After penetrating the peritoneum and obtaining return of fluid, release the skin. Note that the needle is now angled caudally.  

![Figure 43.12 C and D](image3.png)  Use of the Z-tract method helps seal the tract and prevent persistent fluid leaks.
Volume of Fluid Removed

Many patients with chronic ascites are well versed on the procedure and have experienced it many times. Some undergo paracentesis on a regular basis in the outpatient setting. Therefore, the best guide to the volume of fluid to be removed for recurrent ascites is based on the patient’s previous experience, and this question should be asked. Removal of 5 or 6 L is routine and well tolerated, and for therapeutic purposes, at least this volume should be removed. Patients seen in the ED are probably less compliant with outpatient regimens and seek care only when in extremis. Hence, their ascites is likely to be much more voluminous than in those treated regularly. In general, the paracentesis volume consists of as much fluid as can be removed without excessive manipulation of the patient. Volumes greater than 5 L are termed LVP. Up to 10 to 12 L may be removed safely in most patients with chronic ascites (Fig. 43.13). For first-time paracentesis and for diagnostic purposes (ruling out bacterial peritonitis, screening for cancer), 200 to 500 mL is usually sufficient, but more can be drained if it flows easily.

Removal of large amounts of ascitic fluid during paracentesis can be accomplished with the assistance of continuous wall suction. This technique should be used for patients with large amount of ascites (chronic liver failure patients) and who regularly have large volumes removed during paracentesis. Materials required include continuous wall suction, three-way stopcock, tubing elbow, syringes (3 mL, 5 mL, and 10 mL), suction tubing, three or more suction canisters, and usual paracentesis supplies (Fig. 43.14). The patient is prepped for paracentesis. Standard tubing is attached to the wall suction, with the other end connected to the first suction canister. Tubing is then utilized to attach the first and second canister. Each canister must have one port with self-sealing filter, which is needed to close the system (see Fig. 43.14B). Seal and cap all ports. This process can be repeated creating a suction train (see Fig. 43.14C). Once the suction canisters are connected, take the final suction tubing end and place into a syringe, with plunger removed. After the paracentesis catheter has been placed and flow obtained, attach the suction syringe to the catheter or first to a 3-way stopcock (see Fig. 43.14D). Turn the paracentesis catheter to the open position and turn on wall suction. Fluid should begin to drain into the first canister, and once full, fluid will continue into the other canisters in order. If flow stops, a stopcock can be helpful to flush the catheter with sterile saline or the patient’s ascitic fluid. The final setup is shown in Fig. 43.14E.128

![Figure 43.13 Large-volume paracentesis in a patient with a history of recurrent ascites. See text for guidelines on the volume to remove and potential complications of removing large volumes. Note the use of blood pressure monitoring.](Image)
Although paracentesis has traditionally been performed “blindly,” use of ultrasound allows the physician to confirm the presence of ascites. In addition, it allows the clinician to evaluate the most optimal location to attempt the procedure, which can be less straightforward in patients with smaller fluid collections or in whom physical examination alone is insufficient to make the diagnosis.

**Equipment**

Use a low-frequency transducer (2 to 5 mHz) to obtain a sufficient depth of penetration to visualize the area of interest. It is frequently helpful to set the initial depth to 20 to 25 cm to ensure that the entire abdomen is visualized.

**Image Interpretation**

The location of ascites is variable and depends on the amount of fluid present, as well as the position of the patient. The dependent areas of the abdomen, those evaluated with the focused abdominal sonography for trauma (FAST) examination, are ideal initial areas of evaluation. Evaluate the right upper quadrant by placing the transducer in either the transverse or the longitudinal orientation in the 8th to...
ULTRASOUND BOX 43.1: Abdominal Paracentesis—cont’d

11th intercostal space at the anterior axillary line (Fig. 43.US1). The transducer will probably need to be adjusted slightly to allow an optimal view of this area, either by moving up or down a rib space or by adjusting the angle of the transducer. Ascitic fluid will appear anechoic (black) and may be seen in Morison’s pouch (in the hepatorenal space), near the inferior aspect of the liver, or at the superior aspect of the liver (Fig. 43.US2). Once this area has been evaluated in detail, evaluate the left upper quadrant in much the same fashion. Because the spleen is typically slightly more superior and posterior than the liver, place the transducer in the 8th to 11th intercostal space in the posterior axillary line (Fig. 43.US3). Again, make slight adjustments as needed to fully evaluate the area in question. Ascitic fluid will appear as anechoic (black) areas in the splenorenal space, as well as the potential space between the diaphragm and the spleen (Fig. 43.US4). Once these areas have been evaluated, look at the pelvis and lower part of the abdomen by placing the transducer just superior to the pubic symphysis (Fig. 43.US5). First identify the bladder to avoid any confusion. The bladder is typically seen as a well-demarcated area that is rectangular in shape in the transverse orientation and slightly triangular in shape in the longitudinal orientation (Fig. 43.US6). In contrast, free fluid will appear to have irregular borders and may seem to “seep” into the crevices of the lower part of the abdomen and pelvis. Additionally, bowel may be seen to float freely within free fluid (Fig. 43.US7). If

Figure 43.US1 Placement of the ultrasound transducer in the right upper quadrant to evaluate for the presence of free fluid.

Figure 43.US2 Free fluid in the right upper quadrant. The liver can be seen as the hypoechoic (dark gray) object on the left of the image, with anechoic (black) fluid seen between the liver and kidney (on the right of the image).

Figure 43.US3 Placement of the ultrasound transducer in the left upper quadrant to evaluate for the presence of free fluid.

Figure 43.US4 Free fluid in the left upper quadrant. Anechoic (black) free fluid can be seen surrounding the spleen at the right of the image. The diaphragm is seen as the bright white arcing structure on the far right of the image.
ULTRASOUND BOX 43.1: Abdominal Paracentesis—cont’d

**Figure 43.US5** Placement of the ultrasound transducer to evaluate the pelvis for free fluid.

**Figure 43.US6** In this ultrasound image of the pelvis, the bladder can be seen as the rectangular object filled with anechoic (black) urine. An area of free fluid can be seen posterior to the bladder.

**Figure 43.US7** Large area of free fluid seen with ultrasound. Bowel loops are seen to “float” within the fluid.

**Figure 43.US8** Place the paracentesis needle at the site that ultrasound had previously identified as being free of bowel loops and containing a large amount of fluid.

Procedure and Technique

Once free fluid has been seen, identify the area with the largest collection of fluid. Use ultrasound to identify the areas that contain the least number of loops of bowel. If the fluid collection appears to be large, use ultrasound to "mark the spot" and then put it aside. Proceed with the procedure in the typical sterile fashion in the area that has been identified and marked (Fig. 43.US8).

In cases in which the fluid collection is smaller, cover the transducer with a sterile cover to guide direct placement of the catheter.

Complications

Complications of paracentesis can be divided into systemic, local, and intraperitoneal categories.

**Systemic**

Retrospective studies have suggested a risk for deterioration in hospitalized patients following LVP. However, the risk attributable to paracentesis is unclear given the difficulty in adjusting for the underlying severity of illness and comorbid conditions in these patients. The risk in ambulatory ED patients has not been investigated but is clearly less because paracentesis is often an outpatient procedure. Most systemic complications of paracentesis are merely temporally related or seen in seriously ill patients with end-stage liver failure who have a plethora of comorbidities.
Oft-cited but poorly documented concerns include hemo-
dynamic compromise and other physiologic conditions caused
by the overzealous removal of large volumes of ascitic fluid
(>5 L). This has been termed postparacentesis circulatory
dysfunction (PCD), which may occur in 15% to
20% of patients undergoing LVP. It may not occur for several
hours or days after paracentesis and is characterized primarily
by hypovolemia (often asymptomatic), hyponatremia, and
impaired renal function. The actual clinical significance of the
associated disorders is unclear. Hypotension, for example, is
customary after paracentesis and is often asymptomatic.

Because upward of 6 L has been reportedly removed in less
than 15 minutes without complication, certain authorities decry
this issue as folklore. Others believe that rapid total para-
centesis is accompanied by marked cardiovascular and humoral
changes, some of which are explained by mechanical factors
directly or indirectly related to relief of abdominal pressure.
Other changes, including systemic vasodilation and humoral
deactivation, are of a nonmechanical nature. Hepatic enceph-
apathy, hyponatremia, hepatorenal syndrome, and rapid
reaccumulation of ascitic fluid have also been ascribed to LVP.
In the editor's clinical experience, PCD of varying degrees is
a real syndrome, and it may occur more often than suspected
because many patients are discharged after the procedure. The
consequences of LVP in ED patients have not been well studied,
and the implications are somewhat obscure in this subset
of patients. Because many patients require therapeutic para-
centesis on a regular basis, ask the patient about previous
experience and the usual volume of fluid removed to help
guide treatment.

Because fluid and electrolyte shifts tend to be minimal after
the removal of large amounts of fluid, colloid infusion is
considered strictly optional by some for patients with para-
centesis of more than 5 L and is universally not recommended
for paracentesis of lesser volume. Others support the
routine administration of albumin if more than 5 L is
removed. When colloid is indicated, albumin has been the de
facto choice. Being a blood product, albumin has been associated
with rare complications, including anaphylaxis. The recom-
manded infusion is 6 to 8 g of IV albumin per liter of ascitic
fluid removed, or 50 g. However, colloid dextran 70 is
favored by some authorities because of cost and concern for
infection.

The necessity of routine plasma expansion after LVP remains
controversial. No study has shown a direct survival advantage
of one expander over another or in comparison to no expander.
Postparacentesis plasma volume expansion does not prevent
asymptomatic laboratory abnormalities. The mere fact that
controversy still exists suggests that no clear indications can
be promulgated to the emergency physician. The authors
suggest not using IV albumin after taps of less than 5 L. We
suggest that it may be used (6 to 8 g of albumin per liter of
fluid removed, or 50 g) when more than 6 to 8 L is removed.
Twenty-five percent albumin can be given if the patient is
hypervolemic, whereas 5% albumin can be given if dehydration
is suspected. If used, albumin is generally given immediately
after the procedure, although administering it immediately
before the procedure also seems reasonable.

Local
Local complications include persistent leakage of ascitic fluid
at the wound site, abdominal wall hematoma, and localized
infection. Persistent fluid leaks can be corrected with a single
suture at the site of puncture. Just as in DPL site persistent
leakage, cyanoacrylate glue can be utilized to stop leakage.
The patient is positioned so the site is non-dependent and
then pressure applied to the site with gauze for ten minutes.
The gauze is then withdrawn and adhesive applied over the
puncture site. To assist with drying, supplemental air can be
applied to the site using nasal cannula or tubing connected to
wall air or oxygen supply. An abdominal wall hematoma
requiring transfusion is very uncommon, but careful observation
in such cases is necessary.

Intraperitoneal
Intraperitoneal complications include perforation of vessels
and viscera. In experienced hands these complications are
uncommon, and in most circumstances, they are self-sealing
and clinically inconsequential. However, generalized peritonitis
and abdominal wall abscess have been reported after paracentesis
in rare cases. The most common cause of postparacentesis
IPH is bleeding as a result of a coagulopathy rather than
large-vessel injury per se.

Interpretation
Ascitic fluid should undergo gross inspection. Routine laboratory
testing includes a differential cell count, albumin assay, and
cultures (Box 43.3). The prevalence of occult infection of ascitic
fluid in asymptomatic outpatient undergoing therapeutic LVP
for resistant ascites is low. As a result, in the absence of concern-
ing symptoms or cloudy fluid, routine laboratory tests and
culture of fluid during outpatient or ED paracentesis are not
warranted.

Inspection
Ascitic fluid is typically translucent and yellow. A dark greenish
brown hue may reflect biliary perforation. Cloudy fluid gener-
ally indicates particulate matter, including neutrophils; fluid
with WBC counts greater than 5000/µL (i.e., > 5000/mm³)
are cloudy, and those greater than 50,000/µL are purulent.
An opaque, milky appearance may indicate elevated triglycer-
eride levels. A blood-tinged appearance requires at least
10,000 RBCs/µL. This may reflect an iatrogenic complication,
malignancy, hemorrhagic pancreatitis, or tuberculous peritonitis, although the last diagnosis creates hemorrhagic-appearing fluid in less than 5% of cases.120

**Cell Count**

Several milliliters of ascitic fluid are sufficient to obtain a differential cell count. Cirrhotic ascites should generally contain less than 250 WBCs/µL (Table 43.7). However, because cells may exit through the peritoneal cavity more slowly than fluid does, the WBC count can rise in the ascitic fluid during the procedure. Thus, an upper limit for uncomplicated cirrhotic ascites is reported as 500 cells/µL.145−147 Lymphocytes should predominate, and clinical signs or symptoms of peritoneal infection should be absent.148 In cases in which spontaneous bacterial peritonitis (SBP) is a clinical consideration, the WBC criterion is 250/µL with greater than 50% polymorphonuclear leukocytes.147

**Albumin**

A serum-ascites albumin gradient can be obtained by simultaneously measuring albumin in ascites and serum and calculating the gradient. A serum-ascites albumin gradient greater than 1.1 g/dL indicates portal hypertension with greater than 95% accuracy (see Box 43.2).148−150

---

**TABLE 43.7** Ascitic Fluid Characteristics in Various Disease States

<table>
<thead>
<tr>
<th>CONDITION</th>
<th>GROSS APPEARANCE</th>
<th>SPECIFIC GRAVITY</th>
<th>PROTEIN (g/dL)</th>
<th>Cell Count</th>
<th>OTHER TESTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cirrhosis</td>
<td>Straw colored or bile stained</td>
<td>&lt;1.016 (95%)</td>
<td>&lt;25 (95%)</td>
<td>1%</td>
<td>&lt; 250 (90%); predominately mesothelial</td>
</tr>
<tr>
<td>Neoplasm</td>
<td>Straw colored, hemorrhagic, mucinous, or chylous</td>
<td>Variable, &gt;1.016 (45%)</td>
<td>&gt;25 (75%)</td>
<td>20%</td>
<td>&gt; 1000 (50%); variable cell types Peritoneal biopsy, cell block, peritoneal biopsy</td>
</tr>
<tr>
<td>Tuberculous peritonitis</td>
<td>Clear, turbid, hemorrhagic, or chylous</td>
<td>Variable, &gt;1.016 (50%)</td>
<td>&gt;25 (50%)</td>
<td>7%</td>
<td>&gt; 1000 (70%); usually &gt; 70% lymphocytes Peritoneal biopsy, stain and culture for acid-fast bacilli</td>
</tr>
<tr>
<td>Pyogenic peritonitis</td>
<td>Turbid or purulent</td>
<td>If purulent, &gt;1.016</td>
<td>If purulent, &gt;2.5</td>
<td>Unusual</td>
<td>&gt; 250; mainly polymorphonuclear leukocytes Positive Gram stain, culture</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td>Straw colored</td>
<td>Variable, &lt;1.016 (60%)</td>
<td>Variable, 15–53</td>
<td>10%</td>
<td>&lt; 1000 (90%); usually mesothelial, mononuclear</td>
</tr>
<tr>
<td>Nephrosis</td>
<td>Straw colored or chylous</td>
<td>&lt;1.016</td>
<td>&lt;25 (100%)</td>
<td>Unusual</td>
<td>&lt; 250; mesothelial, mononuclear If chylous, ether extraction, Sudan staining</td>
</tr>
<tr>
<td>Pancreatic ascites (pancreatitis, pseudocyst)</td>
<td>Turbid, hemorrhagic, or chylous</td>
<td>Variable, often &gt;1.016</td>
<td>Variable, often &gt;25</td>
<td>Variable</td>
<td>Increased amylase in ascitic fluid and serum</td>
</tr>
</tbody>
</table>

*Because the conditions of examining fluid and selecting patients were not identical in each series, the percentages (in parentheses) should be taken as an indication of the order of magnitude rather than as the precise incidence of any abnormal finding.

**Culture and Gram Stain**

The most valuable method for determining the presence of infection is culture. The sensitivity of this test is markedly increased by direct inoculation of blood culture bottles at the bedside as opposed to simply delivering the ascitic fluid to the laboratory.131,152 Gram staining of peritoneal fluid is rarely helpful in the evaluation of ascites.118 Approximately 10 bacteria/µL of fluid is required for a positive Gram stain. Thus, the Gram stain is notoriously insensitive in patients with SBP, in whom the medium concentration of bacteria is 10³ organisms/µL of fluid.151 Gram stain can be expected to be helpful only in cases of free gut perforation.

**Miscellaneous**

Optional tests include measurement of total protein, glucose, lactate dehydrogenase, and amylase. These tests will be beneficial in selected circumstances and need not be obtained on a routine basis. Immunosuppressed patients, including those with HIV/AIDS, should undergo microbiologic testing for opportunistic infections, including tuberculosis.118 Cytologic analysis is recommended in patients with suspicious constitutional symptoms and signs.151,154 Triglyceride and bilirubin studies are indicated if the gross appearance of the fluid is suggestive of increased levels.155

---

Medical Therapy and Disposition

Total paracentesis may be performed safely in the ED, even in cirrhotic patients with large volumes of ascitic fluid (> 5 L). However, the immediate relief provided by the procedure is temporary, and medical therapy is indicated to prevent or slow the reaccumulation of fluid. Measures include reduction of dietary sodium intake (< 2000 mg/day) and the use of diuretics (spironolactone and furosemide) to promote natriuresis. It is prudent to observe patients undergoing LVP in the ED for 2 to 4 hours for hemodynamic compromise. In the absence of other indications for hospital admission, these patients may then be managed in the outpatient setting with close follow-up to ensure adequacy of their medical regimen. Overall, the clinician should base the decision to admit or discharge on the initial scenario, individual patient characteristics, and response to paracentesis.

Spontaneous Bacterial Peritonitis

SBP is an ascitic fluid infection, occurring almost always in patients with cirrhosis and ascites. It is often associated with fever, abdominal pain, altered mental status, abdominal tenderness, diarrhea, paralytic ileus, hypotension, or hypothermia. It may be clinically quite subtle in some patients. Patients suspected of SBP, and after paracentesis, should be treated with empiric antibiotic therapy to improve patient morbidity and mortality. The majority of cases are due to E. coli and Klebsiella. Unfortunately, studies comparing antibiotics in patients with SBP are limited. Cefotaxime 2 g intravenously every 8 hours provides optimal blood and ascitic fluid levels. Other options include cephalosporins and fluoroquinolones, such as ceftazidime, cefepime, or carbapenem. For patients with secondary bacterial peritonitis or polymicrobial infections, broader coverage is warranted with the addition of metronidazole.

Renal failure develops in up to 40% of patients with SBP, which is a significant cause of mortality. This risk of death may be decreased with albumin infused at 1.5 g/kg body weight within six hours of diagnosis, and albumin should be provided with creatinine greater than 1 mg/dL (88 mmol/L), blood urea nitrogen greater than 30 mg/dL (10.7 mmol/L), or total bilirubin greater than 4 mg/dL. Albumin infusion is associated with decrease in incidence of renal impairment and mortality.

Chronic Ambulatory Peritoneal Dialysis

Patients undergoing chronic ambulatory peritoneal dialysis are at an increased risk for peritonitis because of the presence of a chronic indwelling peritoneal catheter. Culture yield is maximized by obtaining a sample of greater than 10 mL of the peritoneal effluent under sterile conditions after a dwell time of at least 2 to 4 hours. Peritonitis is defined by cloudy fluid with more than 100 WBCs/mm³ and greater than 50% polymorphonuclear cells. Although Gram stain is often negative in patients with bacterial peritonitis, it may reveal the presence of yeast and prompt timely initiation of antifungal therapy. Intraperitoneal antibiotics are superior to IV dosing, and removal of the catheter may be required for refractory or recurrent infections. Initial empirical intraperitoneal therapy usually includes a first-generation cephalosporin along with an aminoglycoside, ceftazidime, cefepime, or carbapenem. Vancomycin should be considered in patients with a previous history of methicillin-resistant Staphylococcus aureus colonization or infection, in those who are seriously ill, or in areas with an increased local rate of methicillin resistance. The optimal treatment strategy should be discussed with the consulting nephrologist.

Acknowledgment

This chapter is dedicated to the memory of John A. Marx, MD, editor, author, educator, researcher, leader, humanitarian, and friend.
REFERENCES

896.e2 SECTION VII GASTROINTESTINAL PROCEDURES


When a patient is seen in the emergency department (ED) with a suspected abdominal hernia, the emergency clinician should consider three issues: (1) Is a palpable mass truly a hernia? (2) Is the hernia easily reducible or incarcerated? (3) Is the vascular supply to the bowel strangulated?

Any patient with symptoms of bowel obstruction should also be evaluated for the possible presence of an abdominal hernia. Changes in symptoms is not necessarily a surgical emergency. Some seemingly incarcerated hernias can be reduced by careful manipulation in the ED. A patient seen in the ED with a chronically incarcerated hernia without obstruction or significant changes in symptoms is not necessarily a surgical emergency. Any patient with symptoms of bowel obstruction should also be evaluated for the possible presence of an abdominal hernia (Fig. 44.1).1

Hernias in the groin area have been the subject of medical diagnosis and treatment as long ago as 1550 BC. Throughout history, treatment of this condition has been the focus of ongoing discussion and debate.2–7 This chapter addresses abdominal and groin hernias, both of which are amenable to diagnosis and potential manual reduction in the ED.

BACKGROUND

A hernia is defined as: a protrusion of any viscus from its normal cavity through an abnormal opening. Abdominal hernias are characterized by protrusion of intraabdominal contents (usually bowel or omentum) through an abnormal defect in the abdominal wall musculature. Hernias can develop along a congenital tract that fails to close (e.g., indirect inguinal or umbilical hernias), or along an area of weakness in the muscular and fascial layers (e.g., direct inguinal, ventral, or incisional hernias). This weakness may be the result of aging and the accompanying loss of tissue elasticity, increased intraabdominal pressure, failure of proper healing, or trauma involving the abdominal wall. It is estimated that hernias develop in 5% of men and 2% of women,8,9 and that 75% of them occur in the groin.10 In children and young adults the majority of hernias are indirect inguinal hernias of congenital origin,11 whereas most direct hernias are acquired and become more common as a patient ages.12 In patients with abdominal surgery, incisional hernias can occur in up to 20% of cases.

CLASSIFICATION

One of the first priorities for an emergency clinician is to determine whether a suspected hernia is reducible, incarcerated, or strangulated. A reducible hernia is one whose contents can be returned through the fascial defect back into the abdominal cavity without surgical intervention. Patients often have large reducible hernias for years and are able to reduce them easily, but such hernias can also become strangulated or incarcerated. An incarcerated hernia is one whose contents are not reducible without surgical intervention. These hernias often have associated swelling of the hernia sac contents. A strangulated hernia is an incarcerated hernia whose blood supply to the herniated structures is compromised. Hernias with a small neck are more likely to become incarcerated or strangulated. A strangulated hernia is a surgical emergency because tissue ischemia and necrosis will result if adequate blood flow is not restored.

A primary ventral hernia of the abdominal wall may be umbilical, epigastric, or spigelian, depending on its location. An incisional hernia is found along or near a previous surgical scar. An inguinal hernia is found within the inguinal triangle, which is formed by the inguinal ligament on the inferior side, the inferior epigastric artery on the superior lateral side, and the lateral edge of the rectus abdominis muscle on the medial side. Direct and indirect inguinal hernias occur superior to the inguinal ligament, whereas a femoral hernia is located inferior to the inguinal ligament. A spigelian hernia (lateral ventral hernia) is located in the abdominal wall just lateral to the rectus abdominis muscle.

Indirect Inguinal Hernia

An indirect inguinal hernia passes through the internal (deep) inguinal ring and into the inguinal canal (Fig. 44.2). It is located lateral to the inferior epigastric vessels. During fetal development, the processus vaginalis allows descent of the testes into the scrotum. Failure of this to close before birth can lead to a hernia or hydrocele.

An indirect inguinal hernia is the most common type of hernia. These occur more frequently in males and are commonly diagnosed in children and young adults. Approximately 5% of full-term infants and 30% of preterm infants will have an inguinal hernia.13,14 Incarceration occurs more commonly in patients younger than 1 year, and 30% of hernias in children younger than 3 months become incarcerated.15,16 When an incarcerated inguinal hernia is successfully reduced in a child, surgical repair within 24 to 48 hours should be considered because of the risk for recurrent incarceration.17 When an inguinal hernia is diagnosed, even without incarceration or strangulation, it is important to make a referral for elective repair. Asymptomatic and painless inguinal hernias can progress to cause symptoms over time if they are not surgically repaired,4 although watchful waiting may also be appropriate in some patients.18 Clinical studies demonstrate increased morbidity with emergency versus elective repair of inguinal hernias.19

Direct Inguinal Hernia

A direct inguinal hernia comes directly through the muscular and fascial wall of the abdomen. It is located medial to the inferior epigastric vessels within the inguinal triangle (Fig. 44.3). These can be differentiated from indirect inguinal hernias as direct hernias do not travel along the inguinal canal.

A direct inguinal hernia is the second most common groin hernia. It is an acquired weakening of the myofascial wall caused by aging and the repetitive stress of increased abdominal pressure. This hernia carries a lower risk for incarceration because the hernia orifice is typically wide.
Femoral Hernia
A femoral hernia occurs inferior to the inguinal ligament through a defect in the transversalis fascia. Abdominal contents protrude into the potential space medial to the femoral vein and lateral to the lacunar ligament in the femoral canal, and appear as a medial thigh mass below the area where direct and indirect hernias are typically identified (Fig. 44.4). A small fascial defect leading to constriction by the inguinal ligament means that this hernia becomes incarcerated in up to 45% of cases. A femoral hernia is relatively uncommon, occurs more frequently in women, and is an uncommon condition in children.

Incisional Hernia
An incisional hernia may occur in as many as one out of every five patients following abdominal surgery (Fig. 44.5). Poor wound healing (e.g., because of infection) increases the likelihood of developing an incisional hernia. After repair,
incisional hernias carry a recurrence rate of 20% to 50%. Incarceration risk depends upon the size of the hernia, with larger defects having a lower risk.

**Umbilical Hernia**

An umbilical hernia traverses the fibromuscular ring of the umbilicus (see Fig. 44.5B). This hernia is most commonly found in infants and children, is congenital in origin, and often resolves without treatment by the age of 5. If the hernia persists beyond this age, is larger than 2 cm, or becomes incarcerated or strangulated, it may be repaired surgically. An acquired umbilical hernia may also be seen in an adult, particularly with increased abdominal pressure (such as with obesity, ascites, or pregnancy). An umbilical hernia is more prone to incarceration and strangulation in an adult than in a child.

**Epigastric Hernia**

This hernia occurs in the midline through the linea alba of the rectus sheath. (see Fig. 44.5C). It is usually located in the epigastric region between the xiphoid and the umbilicus. Though previously considered rare in infants, one study found epigastric hernias in 4% of all pediatric patients evaluated for hernias. In adults, these hernias are usually small and contain preperitoneal fat.

**Spigelian Hernia**

A spigelian hernia is rare and occurs through a defect in the lateral edge of the rectus muscle at the level of the semilunar line and near the arcuate line (see Fig. 44.5D). It is caused by a partial abdominal wall defect in the transverse abdominal aponeurosis, the spigelian fascia. Patients are typically 40 to 70 years of age, but the hernia has also been reported in younger patients. Incarceration rates (often with omentum) have been reported to be as high as 20% with these uncommon hernias. Some reports suggest that ultrasound may be a valuable adjunct for the diagnosis of these hernias and may be helpful during attempted reduction procedures.

**DIAGNOSIS**

**History and Physical Examination**

A patient with a symptomatic hernia may seek treatment in the ED because of swelling or pain in the region of the hernia or abdomen. Ask whether the patient has a history of heavy lifting. Inquire about signs of infection and systemic illness, such as fever, chills, and malaise. Seek symptoms associated with sustained increases in intraabdominal pressure such as chronic cough, chronic constipation, or straining to urinate. Determine whether the patient has signs or symptoms of bowel obstruction, including nausea and vomiting. Document a record of previous surgeries and hernia repairs, including the presence of synthetic mesh.

On physical examination, palpate the inguinal canal in males by inverting the scrotal skin and passing a finger into the external ring. Ask the patient to cough or perform a Valsalva maneuver, which increases intraabdominal pressure and facilitates detection of a hernia. Examining the patient in both standing and supine positions can be helpful. Palpation of the external ring is more difficult in females because it is narrower. An indirect inguinal hernia is manifested as a swelling in the area of the inguinal ligament or as scrotal swelling in male patients. It is often painless and may be noted as an incidental finding. On examination, an indirect hernia can be differentiated from a direct hernia in several distinct ways: (1) an indirect hernia begins lateral to the inferior epigastric arteries; (2) the contents of an indirect hernia will strike the top of the finger instead of the volar pad during examination, as the hernia protrudes down the canal instead of directly across a fascial defect; (3) applying pressure over the internal ring after hernia reduction will block recurrence of the hernia during Valsalva, without impacting the bulge of a direct hernia during straining; and (4) a hernia that fills the scrotum is most likely an indirect hernia.

An asymptomatic hernia may be manifested as a mass that is found incidentally on physical examination of the abdomen or groin. If a hernia is easily reducible, no specific intervention is required in the ED, but instructions should be given for appropriate outpatient surgical follow-up for potential elective repair. This is particularly important for inguinal hernias because elective repair is associated with much less morbidity than emergency repair for strangulation.

A child with a groin hernia may have a reducible inguinal or scrotal mass that occurs with straining or crying. Consider the possibility of an incarcerated or strangulated hernia in young children with nonspecific complaints such as vomiting, poor eating, lethargy, or irritability.

**Radiologic Imaging**

When findings on physical examination are equivocal and the emergency clinician suspects an occult hernia, several options are available for diagnostic imaging.
Diagnosis of Incarcerated Versus Strangulated Hernias

When the patient or emergency care provider cannot manually reduce the contents of the hernia back into the abdominal cavity, the hernia is described as incarcerated. Although hernias are a leading cause of bowel obstruction, patients with incarcerated hernias do not necessarily have associated bowel obstruction. Incarceration is more common with femoral hernias, small indirect inguinal hernias, and ventral or incisional hernias. Incarceration can be caused by the presence of a small fascial defect, by constriction of the defect by surrounding musculature, adhesions, or by swelling of the hernia contents.

A strangulated hernia is one in which the vascular supply to the herniated bowel is compromised, thus leading to ischemia. Strangulated hernias will most commonly also be incarcerated, but this is not a universal finding. Ischemic injury of the bowel is suggested by a red, purple, or bluish discoloration of the skin over the hernia, significant abdominal tenderness with peritoneal signs, and radiographic findings of extraluminal air, or poor perfusion of herniated bowel on a CT scan with IV contrast. Patients with strangulated hernias may exhibit bowel obstruction, peritonitis, viscus perforation, intraabdominal abscess, or septic shock. Associated symptoms may include nausea, vomiting, fever, or abdominal distention.

In rare instances a strangulated or incarcerated hernia may inadvertently be reduced en masse to a preperitoneal location (Fig. 44.8), thus making the contents no longer palpable. In this case, the hernia has not been reduced fully into the peritoneal cavity, and the incarceration and ischemia therefore have not been relieved. Because the clinician believes that the hernia has been appropriately reduced, this can result in delay in the diagnosis of ischemic bowel. Fortunately, this occurs in less than 1% of hernias. Persistent pain after reduction of a hernia, especially more than at the orifice of the fascial defect, should alert the physician to the possibility of either preperitoneal reduction or reduction of ischemic bowel.

scanning is the diagnostic modality used to confirm the presence of hernias and associated complications such as bowel obstruction or perforation. Magnetic resonance imaging has a high positive predictive value for patients with clinically uncertain herniations, but its utility in the urgent setting is limited. Ultrasound examination has been shown to have good sensitivity and specificity for the diagnosis of groin hernias, and may decrease the rate of emergency surgery by improving the ability to reduce hernias. Ultrasound may also have good specificity and a high positive predictive value for diagnosing postoperative incisional hernias. However, ultrasound results can be operator dependent and a difficult modality to interpret.
Differential Diagnosis

The differential diagnosis for a groin mass is large. Box 44.1 lists a number of disease processes that may masquerade as hernias. For example, testicular torsion can be mistaken for a hernia, especially if there is an associated reactive hydrocele. Examine the testicle for tenderness, swelling, lie, and cremasteric reflex. If there is concern for testicular torsion, urology should be notified immediately, while diagnostic studies are undertaken. A hydrocele can also be confused with a hernia because both can occupy the same anatomic space (Fig. 44.9). A hydrocele may transilluminate, whereas a hernia generally does not. Differentiation can be difficult and may require ultrasound to define the contents of the scrotum.

REDUCTION

Indications and Contraindications

The indications for reducing a hernia are the presence of a hernia and the absence of strangulation. If manual reduction proves to be difficult, limit repetitive attempts as this may increase the swelling and limit the chance of nonoperative reduction by a surgical consultant. Some patients may require sedation to facilitate a successful manual reduction, and thus it may be helpful to have a surgeon available while reduction is attempted and the patient is under sedation in the ED.
Procedure

The first step in successful hernia reduction is to position the patient properly to reduce intraabdominal pressure. Place the patient in a position so that gravity can facilitate hernia sac reduction back into the peritoneal cavity. Ensure patient comfort to decrease voluntary or involuntary muscle contraction and guarding, as this can increase intraabdominal pressure and make reduction more difficult. For ventral abdominal hernias, place the patient in the supine position. The Trendelenburg position (supine with the head 20 degrees downward) may facilitate reduction of inguinal hernias. Many hernias can reduce spontaneously if the patient is left comfortably in this position for 10 to 20 minutes. In children, spontaneous reduction has been reported in up to 80% of inguinal hernias over a 2-hour period using appropriate patient positioning and relaxation without manipulation.

A cool compress or ice pack may help to reduce swelling and facilitate reduction of the hernia. Before attempting reduction, strongly consider sedation and/or analgesia to facilitate patient relaxation and minimize pain associated with the procedure. Options for procedural sedation include etomidate, propofol, midazolam, and fentanyl. If manual reduction is necessary, approach slowly with soft, ongoing dialogue, and warm hands. This method encourages patient relaxation and minimizes muscular contractions resulting from pain, cold, or other physical discomfort.

Before beginning hernia reduction, identify the components of the hernia that will be manipulated during the procedure. A hernia consists of a defect in the existing wall of tissue (muscle and fascia) that makes up the neck of the hernia sac. If the neck is small, the hernia will be more difficult to reduce, and a higher incidence of incarceration and strangulation will result. When attempting to reduce the hernia, take care to not allow the contents of the hernia sac to override the edge of the hernia orifice because this will cause “ballooning” of the contents of the hernia sac around the hernia neck. Attempt to find the edge of the hernia defect and position your hand or fingers along that edge to help reduce ballooning and stabilize the edge of the fascial defect.

When available, bedside ultrasound may facilitate specifically locating the fascial defect and the hernia sac to be reduced. A high frequency linear probe can be used to identify the extent of the fascial defect, thus allowing the clinician to guide the hernia sac back through in the plane of maximal opening in the fascial defect under direct visualization. This real-time imaging allows guidance of the proximal end of the hernia contents through the defect and limiting the “ballooning” that may occur without the benefit of visualization.

Begin the reduction procedure by gently guiding the proximal contents of the hernia sac back through the neck of the hernia first. In other words, reduce the hernia in the opposite order from which the contents protruded. Guiding the distal end of the contents of the hernia through the fascial defect first may cause the proximal contents to be displaced around the opening (ballooning) and prevent reduction. Apply gentle, steady pressure on the tissue at the neck of the hernia to overcome this problem and then gradually reduce the hernia (Figs. 44.10 and 44.11). Failure to perform this important procedure is a common error that precludes reduction.

When attempting to reduce inguinal hernias in children, place the patient supine in an approximate 20-degree Trendelenburg position, which may allow spontaneous reduction. Another option is to place the patient in the “unilateral frog-leg” position (Fig. 44.12). Stabilize the patient by grasping the anterior superior iliac spines to prevent lateral movement of the pelvis. Abduct the ipsilateral leg, externally rotate and flex the hip, and flex the knee to obtain the classic frog-leg position. The purpose of this position is to allow the greatest reapproximation of both the internal and external rings. After achieving this position, use the fingers of one hand to prevent the hernia contents from overriding the external ring, while using the other hand to provide steady but gentle pressure on the contents of the hernia sac. Repeated forceful attempts are not recommended.
INTERPRETATION

In general, successful reduction can be identified by the absence of a mass, palpation of the hernia ring, and relief of pain. Continued significant pain suggests the possibility that strangulated bowel has been reduced back into the peritoneal cavity. Inability to palpate the hernia ring after reduction suggests that the hernia is in the preperitoneal position, is not completely reduced, or is not palpable because of body habitus.
REFERENCES

Anorectal Procedures

Wendy C. Coates

Patients with anorectal disorders frequently seek care in the emergency department (ED). The condition may be isolated or the anorectal complaint may be an outward manifestation of a serious underlying illness. A thorough history and physical examination must precede any procedure. Because of the nature of these conditions, extreme sensitivity and professionalism must be applied.

Patients may be anxious about anorectal examination or associated procedures, including the simple digital rectal examination (DRE). The results of DRE may lead to performing diagnostic or therapeutic procedures such as anoscopy, excision of thrombosed external hemorrhoids, drainage of anorectal abscesses or pilonidal cysts, reduction of rectal prolapse, or removal of rectal foreign bodies (FBs). For these procedures, analgesia, sedation, or both may be useful adjuncts.

ANATOMY

The rectum and anus compose the most distal portion of the gastrointestinal tract. The rectum begins at the level of the third sacral vertebra and extends distally 12 to 15 cm. Blood supply to the anorectum is derived from the superior, middle, and inferior hemorrhoidal arteries. Venous drainage from the rectum and anus returns to both the portal and systemic systems (Fig. 45.1). The dentate (pectinate) line marks the transition from the rectum to the anus and contains submucosal glands in anal crypts. Occlusion with subsequent infection of these glands is the cause of anorectal abscesses. Sensory innervation to the rectum is primarily visceral, whereas the anus is innervated by cutaneous fibers. Therefore patients are often unaware of rectal pathology because the pain associated with it may be vague or absent. By contrast, anal lesions are usually very painful and well localized.1

DIGITAL RECTAL EXAM (DRE)

Indications and Contraindications

The physical examination should be performed in a private location and the patient should be completely draped and relaxed. Generally, a calm atmosphere and caring examiner preclude the need for analgesic or anxiolytic agents, although they may be needed to facilitate a thorough examination. In some extremely painful conditions, such as thrombosed or gangrenous hemorrhoids, DRE may be postponed until the patient is anesthetized. If a sharp-edged FB (e.g., metal blade or broken glass) is suspected, performing a DRE may cause injury to the clinician, as well as the patient. In these cases, radiologic evaluation with subsequent operative management may be indicated.

Procedure

Place the patient in the lateral decubitus position and wear protective gloves. Begin the examination with a preliminary visual inspection of the perianal area for important information regarding patient hygiene, trauma, or sexually transmitted diseases. Next, ask the patient to perform a Valsalva maneuver and look for prolapsing rectal mucosa or hemorrhoids. When the patient relaxes, prolapsed structures may retreat or remain external to the anus. By placing a gloved finger firmly against the anal sphincter, it will relax and allow entry of the examiner’s gloved, lubricated finger (Fig. 45.2A). Once inserted into the anus, make a 360-degree sweep to identify any irregularities in the anorectum and prostate. After withdrawing the finger, examine stool remaining on the glove for the presence of visible or occult blood (see Fig. 45.2B and C).6 Testing for blood in stool is discussed in detail in Chapter 67.

Complications

Although DRE causes some vasovagal depression, it is safe to perform in patients with acute myocardial infarction.6 Although it is not a firm contraindication in patients whose absolute neutrophil count is dangerously low, many would defer routine DRE, especially avoiding vigorous prostate manipulation, to minimize the likelihood of bacteremia.

ANOSCOPY

Indications and Contraindications

When evaluating a patient for anal pathology, some practitioners use anoscopy as an adjunct to DRE. Internal hemorrhoids, tears in the distal rectal mucosa, FBs, and distal anorectal masses may be visualized. An imperforate anus is the only absolute contraindication to anoscopy; however, severe rectal pain, a common complaint in the ED, may preclude awake anoscopic examination in anxious patients or those suffering from thrombosed hemorrhoids or other painful conditions. Internal visual inspection in these patients is better performed with sedation. In many cases a more definitive study such as sigmoidoscopy or colonoscopy is being planned by the treating physician, and thus anoscopy can be deferred in the acute care setting.

Equipment and Setup

The anoscope is a plastic or stainless steel tube with a removable obturator (Fig. 45.3). It may have an integrated light source or require an external light or head lamp. An appropriate examination table, topical anesthetics, lubricant, gauze, and forceps may also be required.

Positioning

Ideally, place the patient in a prone position on a proctoscopic examination table. The lateral decubitus position with the knees and hips flexed may also be adequate and is sometimes tolerated better. In the lateral decubitus position, place the patient on the left side if the examiner is right-handed (Fig. 45.4, step 1).
**Digital Rectal Examination**

**Figure 45.2** Digital rectal examination. A, Insert a gloved, lubricated finger into the anus and perform a 360-degree sweep to check for any irregularities in the anorectum and prostate. B, Examine the stool for visible and occult blood. This patient had melena, which is dark and tarry in consistency and indicative of upper gastrointestinal hemorrhage. C, This sample is positive for blood. The large arrow points to the blue coloration on the test area of the card, whereas the small arrow points to the control. The control should turn blue. See also a discussion of testing stool for blood in Chapter 67.

**Procedure**

Although most patients do not require intravenous sedation and analgesia, administer these agents as needed to keep the patient relaxed and comfortable. Before anoscopy, perform a routine DRE to identify sources of bleeding or pain and to locate any palpable masses. Next, lubricate the anoscope. After DRE and with the obturator inserted completely into the anoscope, carefully introduce the scope into the anus. Use gentle, constant pressure to overcome resistance from involuntary contraction of the external anal sphincter. Gently advance the anoscope while asking the patient to bear down slightly. Pass the anoscope gently into the anorectum (see Fig. 45.4, step 2). The examiner should keep a thumb pressing on the obturator to prevent it from being dislodged. If the obturator falls back during insertion, remove the anoscope completely.
**Figure 45.3**

A, Reusable stainless steel and B, disposable plastic anosopes are available. Regardless of the type of anoscope used, the obturator must be fully inserted and held in place with the thumb during advancement of the device into the anorectum.

---

**Anoscopy**

1. Left lateral or Sims’ position

2. With the obturator fully inserted into the anoscope, carefully introduce the scope into the anus. Hold the obturator in place with your thumb. Use gentle, constant pressure to pass the anoscope into the anorectum. Advance until the outer flange impinges on the anal verge.

3. Place the patient in the left lateral position (A), prone position (B), or ideally prone on a proctoscopic examination table (C).

4. Remove the obturator when the anoscope is fully inserted. Gradually withdraw the anoscope and visualize the anal canal. Swab away blood or debris to aid in visualization, culture any discharge, and inspect for hemorrhoids, anal fissures, ulcerations, abscesses, or tears.

---

**Figure 45.4** Anoscopy. Remember to keep your thumb on the obturator during passage of the instrument until it is fully inserted (Fig. 45.2). (Step 1, From Hill GJ II: *Outpatient surgery*, ed 3, Philadelphia, 1988, Saunders.)
MANAGEMENT OF HEMORRHOIDS

Hemorrhoids are a common affliction and have been described and treated for more than 4000 years. The refined, low-fiber diet of Western nations makes hemorrhoids extremely common in the United States, where 1 in 25 to 30 individuals is afflicted. One million patients annually seek medical attention for this condition.

Hemorrhoids are normal vascular structures in the submucosal layer of the anal canal, arising from a channel of arteriovenous connective tissues that drains into the superior and inferior hemorrhoidal veins. Hemorrhoidal tissue is composed of vascular, mucosal, and muscular tissue (Fig. 45.5). There are two types of hemorrhoids: internal and external. Internal hemorrhoids originate above the dentate line, are covered with mucosa, and lack sensory innervation. They are easily identified by noting that their covering differs in appearance from the surrounding perianal skin. Internal hemorrhoids are rarely felt by digital palpation unless they are very large or thrombosed. They are usually painless unless gangrenous, strangulated, extruded, or thrombosed, and then

Complications

Although complications are rare, patients often complain of increased pain after the examination. Local mucosal irritation with subsequent bleeding is the most common complication. To prevent transmission of infectious diseases, dispose of or sterilize instruments after each use.

![Figure 45.5](image)

**A**, Anatomic location of internal and external hemorrhoids. **B**, Thrombosed external hemorrhoid. **C**, Thrombosed prolapsed internal hemorrhoids. These hemorrhoids cannot be permanently reduced and are quite painful; occasionally, partial relief can be obtained by manual reduction if they are not gangrenous. They should not be incised in the emergency department; formal hemorrhoidectomy is required if conservative measures are not successful. They are often mistaken for a partial “rectal prolapse.” Sitz baths and stool softeners are frequently futile in such severe cases. **D**, This small external hemorrhoid ruptured and produced minor but persistent bleeding and pain. Conservative treatment consisting of topical corticosteroids or Preparation H (Pfizer Inc., New York, NY) and frequent sitz baths will be curative in 5 to 7 days. (A, From Hill GJ II: Outpatient surgery, ed 2, Philadelphia, 1980, Saunders.)
they may be extremely painful. Anal pain in the absence of such pathology suggests a problem other than simple internal hemorrhoids.5,5

Internal hemorrhoids can be further classified as first through fourth degree. First-degree hemorrhoids do not prolapse but may be identified on anoscopy. Second-degree hemorrhoids prolapse on straining but reduce spontaneously. Third-degree hemorrhoids prolapse on straining and can be reduced manually, whereas fourth-degree hemorrhoids prolapse and are irreducible. Fourth-degree hemorrhoids are prone to thrombosis, strangulation, and eventually gangrene (see Fig. 45.5C).

External hemorrhoids originate below the dentate line and are covered with squamous epithelium. This makes them easily recognizable because their covering matches the surrounding skin. A thrombosed external hemorrhoid appears as a bluish mass covered by epidermis. Acute thrombosis occurs suddenly and is generally very painful because external hemorrhoids are innervated by the inferior rectal nerve. Many patients feel a tender mass and are unable to sit comfortably. Significant bleeding is uncommon but may occur with spontaneous rupture. Increased pressure from straining, or trauma from constipation or diarrhea, may exacerbate external hemorrhoids. Distention and trauma predispose the hemorrhoidal venous plexus to stasis with ensuing clot formation and edema.2,4,6

Conservative Treatment

ED management of minor internal hemorrhoids is conservative. Prolapsing internal hemorrhoids will not benefit long-term from conservative intervention and should receive surgical consultation. A useful mnemonic for managing hemorrhoids is WASH: water (increase fluid intake, warm water contacting the hemorrhoid via bath or directed shower), analgesics, stool softeners, and a high-fiber diet.3 Psyllium (e.g., Metamucil [The Procter & Gamble Company, Cincinnati, OH]) is often prescribed as a dietary supplement to increase fiber. Hemorrhoids that fail to respond to medical management may be treated on an outpatient basis with rubber band ligation, sclerotherapy, and thermotherapy consisting of an infrared beam, electric current, CO2 laser, or ultrasonic energy. Patients who must push hemorrhoids back in after a bowel movement may be identified on anoscopy examination. Second-degree hemorrhoids that fail to respond to medical management may be treated on an outpatient basis with rubber band ligation, whereas fourth-degree hemorrhoids prolapse and are irreducible. Fourth-degree hemorrhoids are prone to thrombosis, strangulation, and eventually gangrene (see Fig. 45.5C).

Indications and Contraindications

Indications for surgical excision of acutely (<48 hours) thrombosed external hemorrhoids in the ED include relief of symptoms and prevention of the formation of permanent perianal skin tags. These appendages remain as loose skin after the body reabsorbs the thrombosis and serve as a nidus for poor perianal hygiene and local irritation. Surgical consultation should be obtained in the ED for multiple painful external hemorrhoids and for profuse bleeding that is hemodynamically significant. Bleeding disorders, serious systemic illness, and hemodynamic instability are all relative contraindications to excision in the ED.

Procedure

Place the patient in the prone or lateral decubitus position. Tape the buttocks apart to aid in visualization (Fig. 45.6, step 1). An assistant is often needed. Administer parenteral analgesics and sedatives as an adjunct to local anesthesia if necessary. Infiltrate with a local anesthetic (such as 0.5% bupivacaine or buffered 1% lidocaine with epinephrine at 1:100,000) just under the skin and over the dome of the hemorrhoid (Fig. 45.7; also see Fig. 45.6, step 2). The overlying skin should blanch, which indicates that anesthesia has been introduced at the appropriate depth. Inject additional anesthetic through the incised tissue into the base of the hemorrhoid, if needed, rather than through the intact skin (see Fig. 45.7C).

Grasp the skin overlying the thrombosis with forceps. Make an elliptical incision around the clot and direct it radially from the anal orifice (Fig. 45.8; also see Fig. 45.6, step 3). Elevate the edges of the skin with forceps and excise from the edges to expose the underlying thrombosis. Remove the clot with forceps or by applying digital pressure (see Fig. 45.6, steps 4 and 5). Frequently, multiple individual clots will be present. If any skin ulceration is noted over the hemorrhoid, include it in the excised portion. Pack the wound loosely with standard cotton gauze to prevent the skin edges from reapproximating prematurely.

For the trip home, place a gauze pad between the buttocks and tape the buttocks together to hold the gauze in place (see Fig. 45.6, step 6). Advise the patient to avoid prolonged standing or straining for the next few days. Minor bleeding may occur. Hygiene with a directed stream of warm water or sitz baths can be started a few hours after the procedure. The gauze may fall out or can be removed at the first sitz bath. After the packing has been removed, the patient may apply a soothing cream to the area for a couple of days (such as Preparation H, Anusol HC, or witch hazel pads). Instruct the patient to avoid using toilet paper after a bowel movement for a few days but to wash the area with mild soap and water in the shower. Most patients are relatively asymptomatic in 48 hours and do not need a routine wound check unless the pain or bleeding persists. Once the clot has been removed recurrent thrombosis is unlikely, but these patients are predisposed to future episodes. Long-term therapy should be directed toward avoiding constipation by increasing dietary fiber and fluid intake. Antibiotics are not routinely indicated.2,4,5,7
Although complications are rare, bleeding and infection do occur. The bleeding usually stops with direct pressure. When simple incision plus drainage is performed instead of an elliptical excision, or when the ellipse of skin is not removed completely, the edges of the skin can close prematurely and result in

**Figure 45.6** Excision of thrombosed external hemorrhoids.
vibrators, sex toy phalluses, aerosol cans, lightbulbs, glass bottles, billiard balls, fruits, vegetables, and small animals (Fig. 45.9). Many of these objects can be removed successfully in the ED. By following some simple guidelines, outpatient treatment can be practical and cost-effective.

Diagnosis of a rectal FB is usually made from the history. The physical examination should therefore concentrate on excluding anorectal or intestinal perforation and determining which objects will be accessible in the ED. DRE will identify objects that are low lying or palpable. Such objects are most likely to be removed successfully in the outpatient setting. Plain radiographs can supplement the examination by delineating the shape, position, and number of objects. If an FB with a sharp edge is suspected from the history or radiograph, omit the DRE to prevent injury to the provider.8,9

Indications and Contraindications
Although some objects may pass spontaneously, delayed removal may lead to obstipation, pain, infection, and perforation. For these reasons FB removal is indicated according to the algorithm included in this chapter (Fig. 45.10). Rectal FB removal can lead to rectal perforation. Although many FBs can be removed safely in the ED, complicated or prolonged attempts may be better performed under general anesthesia.

Removal of FBs in the ED is contraindicated in patients who have severe abdominal pain or signs of perforation, a nonpalpable FB, or sharp objects or broken glass in the rectum. Other situations precluding ED removal include a rectal FB that is unusually difficult to remove (a set time limit has elapsed, or the patient cannot tolerate removal), or when there is insufficient experience or equipment to perform the procedure. Patients who arrive at the ED under these conditions require surgical consultation.8,9

**Figure 45.8** Schematic technique as described in Fig. 45.6. A, For the unroofing technique, make an elliptical incision to remove a piece of the overlying skin. To prevent skin tags, do not use a simple linear incision. B, Blood clots may extrude spontaneously, but C, remove the remaining ones with forceps or express them with the fingers. D, Frequently, multiple clots are present, and they should all be removed. Ask an assistant to provide exposure with forceps if necessary.

**Figure 45.7** A, For surgical treatment of a thrombosed external hemorrhoid in the emergency department, anesthesia can usually be obtained with a single injection of buffered long-acting bupivacaine with epinephrine. In elective cases, apply a eutectic mixture of local anesthetics (EMLA [Actavis Pharma, Inc., Parsippany, NJ]) cream for 1 hour before the procedure. Parenteral sedation is optional. Using a 25-gauge needle, inject an anesthetic solution into the middle of the swollen hemorrhoid just below the surface of the skin. B, Do not move the tip of the needle. With slow injection, the anesthetic will spread over the surface of the dome and into the surrounding tissue. Avoid deep field blocks at the base of the hemorrhoid because they are unnecessary and very painful. C, If pain persists, inject additional anesthetic into deeper tissues through the cut edges, not through the intact skin.

**Figure 45.8** Schematic technique as described in Fig. 45.6. A, For the unroofing technique, make an elliptical incision to remove a piece of the overlying skin. To prevent skin tags, do not use a simple linear incision. B, Blood clots may extrude spontaneously, but C, remove the remaining ones with forceps or express them with the fingers. D, Frequently, multiple clots are present, and they should all be removed. Ask an assistant to provide exposure with forceps if necessary.

**Figure 45.7** A, For surgical treatment of a thrombosed external hemorrhoid in the emergency department, anesthesia can usually be obtained with a single injection of buffered long-acting bupivacaine with epinephrine. In elective cases, apply a eutectic mixture of local anesthetics (EMLA [Actavis Pharma, Inc., Parsippany, NJ]) cream for 1 hour before the procedure. Parenteral sedation is optional. Using a 25-gauge needle, inject an anesthetic solution into the middle of the swollen hemorrhoid just below the surface of the skin. B, Do not move the tip of the needle. With slow injection, the anesthetic will spread over the surface of the dome and into the surrounding tissue. Avoid deep field blocks at the base of the hemorrhoid because they are unnecessary and very painful. C, If pain persists, inject additional anesthetic into deeper tissues through the cut edges, not through the intact skin.
Figure 45.9 Rectal foreign bodies. A, Two large tungsten spheres, one on top of the other, gave a snowman-like appearance on the radiograph. They were removed manually in the emergency department (ED) with the aid of a Foley catheter and large-volume balloon. B, A vibrator that migrated into the sigmoid colon. It required removal in the operating room under general anesthesia. C, A small glass jar that the patient had tried to remove multiple times over a period of hours at home. On arrival at the ED, he had signs of peritonitis and was taken to the operating room, where perforation of the distal colon was discovered. Diverting colostomy was required. D, A toothbrush that was removed manually in the ED.

Figure 45.10 Emergency approach to the removal of rectal foreign bodies. Foreign bodies that are fragile or associated with rectal spasm are generally managed with regional or general anesthesia. The use of supplemental analgesic, anxiolytic, and local anesthetic medications is recommended.
If DRE reveals that the object has an accessible edge or lip, use an instrument to extract it under direct visualization (Fig. 45.12). First, insert an anoscope, rigid sigmoidoscope, vaginal speculum, or retractor into the anus as described previously in the section on Anoscopy. If an intact object is visualized clearly, use a blunt instrument to secure it. Apply gentle traction to remove the object, the instrument, and the anoscope or speculum as a single unit. Grasp the object under direct visualization to avoid pinching or tearing the mucosa.

Rigid sigmoidoscopes offer a unique advantage in that air can be insufflated into the rectum around the FB. This technique can be particularly helpful when retrieving glass objects. Glass rectal FBs often create a vacuum in the segment of bowel just proximal to where they lie. This makes removal with simple traction almost impossible. The vacuum can be released by distending the rectal wall around the object with air. If a sigmoidoscope cannot be used to retrieve a glass object, pass one or two Foley catheters or an endotracheal tube beyond the FB and inflate the balloon or cuff. Then remove the object with the inflated balloons and gentle traction. Specific equipment is often not available to remove all FBs, and the clinician must improvise depending on the circumstances. Something as simple as two large spoons or an endotracheal tube may be used in lieu of complicated forceps and clamps.

Besides creating a vacuum, glass objects are especially difficult to remove because they can break and cause a tear or perforation in the rectal wall. If forceps are used for retrieval of a glass object, coat the grasping edge with rubber or plastic or pad it with gauze. Plaster of Paris has been used to remove a hollow glass object if the object has an open end facing distally. Insert a hollow tube (e.g., an endotracheal tube, or small chest tube) into the open end. Fill the FB with plaster by injecting it through the hollow tube with a large irrigation syringe. Once the plaster cools and hardens around the tube, it can be used as a handle to remove the object with gentle traction. Be careful not to leak plaster onto the mucosa. In addition, heat is released as the plaster hardens and may cause the glass to crack or shatter. After removal, perform sigmoidoscopy to evaluate for edema and possible perforation of the mucosa. Patients with normal findings on post-extraction examination and no evidence of perforation may be released home safely after a period of observation.8,9

FBs that are positioned proximal to the rectum warrant surgical consultation. Management options include observation to enable passage to the rectum or surgical removal. Enemas or cathartics should not be used because they may increase the impaction of a rectal FB or cause it to move higher into the colon.

**Complications**

The most common complication is an inability to remove the rectal FB, which should prompt surgical consultation. The most serious complication of rectal FB retrieval is perforation or a deep mucosal tear, which may necessitate surgery. Cracking or shattering of glass may also require surgical exploration and retrieval. Mild mucosal edema and rectal bleeding are common sequelae of prolonged rectal FB presence and retrieval. These complications may not require any specific treatment. However, the presence of postprocedural abdominal pain, fever, sustained or profuse rectal bleeding, or discharge warrants surgical consultation.
inside the anal canal. The differential diagnosis includes hemorrhoids, polyps, cystocele, and carcinoma.

There are three types of prolapse. (1) Complete prolapse, or procidentia, involves all layers of the rectum protruding through the anal orifice (Figs. 45.13 and 45.14). (2) Incomplete, or occult, prolapse describes internal prolapse that does not reach the orifice. This type is difficult to diagnose in the ED and requires no emergency intervention. (3) Mucosal prolapse is limited to protrusion of mucosa through the anal opening.

Complete and partial prolapse can be distinguished from each other by digital palpation. A thick muscular layer of tissue

---

**Figure 45.12** Rectal foreign body (FB) removal techniques. *ED,* Emergency department. Difficult-to-remove FBs require operating room techniques.
Rectal Prolapse Reduction

1. Complete (recurrent) rectal prolapse in a nursing home patient. To reduce the prolapse, place the patient in the prone or lateral decubitus position. Parenteral sedation may be required.

2. Tape the buttocks apart (or enlist the help of an assistant to spread the buttocks) to aid in reduction.

3. Apply constant, gentle circumferential pressure to the prolapsed area, beginning with the portion closest to the lumen (the most distal segment). Apply pressure with the thumbs while rolling the walls inward to force the prolapse back through the anus.

4. Successful reduction of the rectal prolapse.

Figure 45.13 A, Type I procidentia (rectal prolapse). B, Intussusception of the sigmoid colon beyond the anus. (A and B, From Kratzer GL, Demarest RJ: Office management of colon and rectal disease, Philadelphia, 1985 Saunders.)
between the examiner’s thumb and forefinger suggests complete prolapse. With partial or mucosal prolapse, radial rectal folds may be seen protruding through the rectum. This type of prolapse rarely extends more than 3 to 4 cm from the anus. Complete prolapse can extend 10 to 15 cm outside the anal verge.

Rectal prolapse is most common in children and older adults. Prolapse in children is typically incomplete, or mucosal. It usually affects children younger than 3 years and is often associated with cystic fibrosis, parasitic infection, chronic diarrhea, malnutrition, or as a sequela of chronic neurologic disease. Prolapse is usually self-limited; outpatient management (after manual reduction) includes correcting constipation, avoiding straining, and referring for testing to exclude cystic fibrosis. Rectal prolapse in adults occurs most often in older women and may be recurrent. The cause is poorly understood, but it is associated with chronic constipation, chronic neurologic conditions, or pudendal neuropathies that weaken the anal sphincter.10,11

**Indications for Reduction**

Rectal prolapse may be reduced in the ED. If unsuccessful, surgical referral is appropriate. Definitive surgery may be attempted, but occasional prolapses in debilitated patients are usually treated conservatively. Patients should be referred for outpatient proctoscopy to search for a polyp or malignancy that may have acted as a lead point. If the prolapse is incarcerated, surgical consultation should be obtained.

**Procedure**

Reduce a mucosal prolapse by applying gentle, constant pressure on the mass for a few minutes. In children, intravenous sedation may be necessary to allow reduction. Children are often more relaxed if they are allowed to remain on the parent’s lap during the procedure. After reduction, send the child home with a pressure dressing and stool softeners. Counsel the parents on the use of dietary fiber and increased fluid intake to prevent constipation and straining. Refer the child for outpatient follow-up.12

For reduction of a complete prolapse, place the patient in the prone or lateral decubitus position (see Fig. 45.14, step 1). Parenteral sedation should be provided if the patient is anxious or has difficulty relaxing the sphincteric muscles. Tape the buttocks apart or have an assistant aid in reduction (see Fig. 45.14, step 2). Apply constant, gentle circumferential pressure to the prolapsed area, beginning with the portion closest to the lumen (the most distal segment). Place the thumbs on either side of the lumen while grasping the exterior walls with the fingers. Apply pressure with the thumbs while rolling the walls inward to force the prolapse back through the anus (see Fig. 45.14, step 3). Care should be taken to avoid poking at the tissue with the fingertips. If substantial tissue edema has developed, application of gauze soaked in sugar water may promote shrinkage and subsequent manual reduction.10

**Complications**

Complications after successful reduction are uncommon but may include bleeding and ulceration. Failure to reduce a prolapse requires surgical consultation. Apply saline-moistened gauze over the rectal tissue while awaiting consultation. A persistently prolapsed rectum can result in ulceration, strangulation, and perforation of the bowel wall. Moreover, the possibility of loss of anal sphincter tone and incontinence increases with delays in reduction of rectal prolapse.

**ANAL FISSURE**

An anal fissure is a small laceration or ulcer at the anal verge.5 Anal fissures are the most common cause of sudden, searing anorectal pain. Though appearing trivial on examination, fissures can be extremely painful, even hours after a bowel movement, because of persistent spasm (Fig. 45.15). The condition is quite difficult to eradicate, is debilitating, and can last for months. It is occasionally associated with bright red rectal bleeding. An anal fissure is most commonly found in young adults, men, and women equally. In children, it can be a sign of child abuse. Anal fissures are usually associated with constipation, a hard or strained stool, or chronic diarrhea, but the exact cause is unknown. The diagnosis is relatively easy to make, and the fissure is readily seen by spreading the buttocks. The vast majority of anal fissures occur in the posterior midline, 10% to 15% occur in the anterior midline, and less than 1% occur in lateral positions. Fissures occurring in atypical locations should prompt consideration of other diseases. Multiple or recurrent fissures are associated with Crohn’s disease, tuberculosis, syphilis, human immunodeficiency virus infection, and malignancy. Conservative therapy with the WASH regimen (see the section on hemorrhoids) may promote gradual healing in 4 to 6 weeks.

Most patients with acute anal fissures and almost half of patients with chronic fissures will experience healing with medical therapy. Therapy is aimed at breaking the cycle of pain, spasm, and ischemia, factors thought to be responsible for the development of the fissure. Therapies include relaxation of the internal sphincter, institution and maintenance of atraumatic passage of stool, and relief of pain. Simple measures include bulk agents, stool softeners, and probably most helpful,
warm sitz baths following bowel movements to relax the sphincter. Based on the theory that anal fissures are caused by ischemia through a spasmodic internal sphincter, pharmacological agents, including glyceryl trinitrate (GTN), diltiazem, and botulinum toxin, may be useful as alternatives to surgical sphincterotomy for chronic fissures. GTN ointment applied two to four times per day to the anus results in various healing rates, but a major side effect is dose-related headaches. Nitroglycerin 0.2% ointment applied twice daily may heal chronic ulcers via a reduction in resting anal pressure and an increase in anodermal blood flow. Diltiazem ointment (2%) appears to have efficacy similar to that of GTN but may cause fewer side effects. Diltiazem may be associated with the development of pruritus. Both diltiazem and GTN are first-line therapies. Botulinum toxin causes temporary muscle paralysis by preventing the release of acetylcholine from presynaptic nerve terminals, thereby decreasing pressure in the internal sphincter. Surgical treatment is generally reserved for fissures that have failed medical therapy and is usually curative.
REFERENCES

Modern emergency medical service (EMS) was created in 1966 as a result of the National Highway Safety Act. Since then, provision of medical care in the prehospital setting has undergone considerable change. Today’s EMS providers perform many advanced lifesaving procedures. Nonetheless, the task of immobilizing potential injuries to the spine, pelvis, and extremities has remained a primary EMS function. This chapter reviews basic prehospital immobilization techniques and equipment, including spinal motion restriction, pelvic stabilization, extremity splinting, and removal of protective equipment.

SPINAL MOTION RESTRICTION

Background

The first widely accepted recommendations for “spinal immobilization” following blunt trauma came from the American Academy of Orthopaedic Surgeons in 1971.1 These guidelines called for spinal immobilization of patients with symptoms or physical findings suggestive of spinal injuries.1,2 Since then, recommendations for spinal immobilization have evolved considerably.

During the 1980s and 1990s, indications for spinal immobilization were based primarily on the mechanism of injury, regardless of the presence or absence of symptoms or physical findings suggestive of a spinal injury.2–4 This resulted in routine and often unnecessary prehospital spinal immobilization for all but the most trivial injuries.6

In 1998, Hauswald and colleagues7 published the results of a 5-year retrospective review comparing patients from Malaysia, where cervical spine immobilization was nonexistent, to patients from New Mexico, where cervical spine immobilization based on the mechanism of injury was standard practice. The authors concluded that out-of-hospital immobilization has little or no effect on neurologic outcome in patients with blunt spinal injuries.7 This study, and others demonstrating increased complications8–11 and significant spinal movement, despite the use of cervical collars (c-collars) and long spine boards,14,15 prompted the development and evaluation of prehospital clinical decision rules to selectively immobilize patients after blunt trauma.16–19

A 4-year prospective study in two Michigan counties found that the use of a selective immobilization protocol resulted in spine immobilization for most patients with spinal injury without causing harm to patients in which spine immobilization was withheld.18 A larger study in Maine demonstrated that selective immobilization based on a statewide protocol resulted in only one nonimmobilized unstable cervical spine fracture in more than 32,000 patient encounters.19

Data demonstrating the safe application of selective spinal immobilization protocols by prehospital providers, a lack of scientific evidence demonstrating improved outcomes with the use of long spine boards, and a keen awareness of the complications associated with spinal immobilization has resulted in new recommendations from the National Association of EMS Physicians and the American College of Surgeons Committee on Trauma20 and the American College of Emergency Physicians.21 These recommendations include the selective use of long spine boards following blunt trauma and elimination of their use following penetrating trauma in patients with no evidence of spinal injury (Box 46.1).20

At the present time, the effects of spinal immobilization on mortality, neurologic injury, spinal stability, and adverse effects in trauma patients remains uncertain. Due to the low incidence of actual spinal cord injury (SCI) and associated neurologic sequelae, prospective trials are difficult to safely design and perform. Nevertheless, available data suggests that we should reduce or eliminate routine prehospital spinal immobilization in favor of using validated clinical rules to determine which patients may have sustained a spinal column injury and applying spinal motion restriction strategies only to these patients. However, these new approaches will take time to promulgate through the EMS community. Until then, emergency physicians must be knowledgeable about prehospital spinal immobilization. Therefore the remainder of this section provides a detailed review of immobilization devices and techniques.

Epidemiology

According to the National Spinal Cord Injury Statistical Center, an estimated 12,500 new, survivable SCIs occur in the United States annually;22 and 276,000 people were living with SCIs in the United States in 2014.22 Since 2010 the most common cause of SCI is motor vehicle collision, which accounts for almost 40% of cases, followed by falls and acts of violence, primarily gunshot wounds.22 Sports such as American football, rugby, swimming and diving, gymnastics, ice hockey, track and
The cost of care in both the immediate and extended care setting can be exorbitant, especially among the young. The average lifetime cost of medical care for patients with SCI varies depending on the level of injury and age at time of injury, and ranges from $1 million to more than $4.7 million, with annual costs from $42,000 to $1 million.22

**Pathophysiology**

The direction and strength of the injurious force may help predict the type of injury sustained. Generally speaking, the basic forces that can be exerted on the spine are flexion, extension, rotation, lateral bending, distraction (stretching), and compression (axial loading).23,24 Of course, complex mechanisms may exert multiple forces. For example, high-speed rollover motor vehicle collisions could easily exert all the aforementioned forces.

Injuries to the upper cervical spine (C1 and C2) (Fig. 46.1) occur more often in older, osteoporotic patients than in younger patients. The spectrum of injuries in the cervicocranium includes occipital condyle fractures, occipitoatlantal dislocations, dislocations and subluxations of the atlantoaxial joint, fractures of the ring of the atlas, odontoid fractures, fractures of the arch of the axis, and fractures of the lateral mass of the axis (Fig. 46.2). Involvement of the spinal cord at this high level can cause devastating neurologic injury, and it is reasonable to believe that many of these injuries are not reported because they result in death.25 Subaxial cervical spinal injuries involving C3–C7 have broad clinical implications. Approximately two-thirds of cervical injuries causing quadriplegia occur within

---

**Box 46.1 EMS Spinal Precautions and the Use of the Long Backboard: Position Statement of the National Association of EMS Physicians and the American College of Surgeons Committee on Trauma**

The National Association of EMS Physicians and the American College of Surgeons Committee on Trauma believe that:

- Long backboards are commonly used to attempt to provide rigid spinal immobilization among EMS trauma patients. However, the benefit of long backboards is largely unproven.
- The long backboard can induce pain, patient agitation, and respiratory compromise. Furthermore, the backboard can decrease tissue perfusion at pressure points, leading to the development of pressure ulcers.
- Utilization of backboards for spinal immobilization during transport should be judicious, so that potential benefits outweigh risks.
- Appropriate patients to be immobilized with a backboard may include those with:
  - Blunt trauma and altered level of consciousness;
  - Spinal pain or tenderness;
  - Neurologic complaint (e.g., numbness or motor weakness);
  - Anatomic deformity of the spine;
  - High-energy mechanism of injury and:
    - Drug or alcohol intoxication;
    - Inability to communicate; and/or
    - Distracting injury.
- Patients for whom immobilization on a backboard is not necessary include those with all of the following:
  - Normal level of consciousness (Glasgow Coma Scale 15);
  - No spine tenderness or anatomic abnormality;
  - No neurologic findings or complaints;
  - No distracting injury;
  - No intoxication.
- Patients with penetrating trauma to the head, neck, or torso and no evidence of spinal injury should not be immobilized on a backboard.
- Spinal precautions can be maintained by applying a rigid cervical collar and securing the patient firmly to the EMS stretcher. They may be most appropriate for:
  - Patients who are found to be ambulatory at the scene;
  - Patients who must be transported for a protracted time, particularly prior to interfacility transfer; or
  - Patients for whom a backboard is not otherwise indicated.
- Whether or not a backboard is used, attention to spinal precautions among at-risk patients is paramount. These include application of a cervical collar, adequate security to a stretcher, minimal movement/transfers, and maintenance of in-line stabilization during any necessary movement/transfers.
- Education of field EMS personnel should include evaluation of risk of spinal injury in the context of options to provide spinal precautions.
- Protocols or plans to promote judicious use of long backboards during prehospital care should engage as many stakeholders in the trauma/EMS system as possible.
- Patients should be removed from backboards as soon as practical in an emergency department.

EMS, Emergency medical service.

the lower cervical spine, with fractures occurring most often in C6 and C7 and dislocations most commonly occurring between C5–C6 and C6–C7. 26

The orientation of the facets in the thoracic spine allows significantly less flexion and extension than in the cervical or lumbar spine. In addition, the free space between the thoracic spinal cord and the borders of the spinal canal is relatively small, and the blood supply is less robust. These factors increase the susceptibility of the spinal cord to injuries at this level. At the thoracolumbar junction there is an acute transition in stability because of the loss of rib restraint, which increases the risk for flexion-extension and rotational injuries. Disc size and shape also change, thus making this section of the spine particularly susceptible to injury. Approximately half of all vertebral body fractures and 40% of all SCIs occur between T11 and L2. 27

The lumbar spine is protected only by the abdominal and paraspinous musculature, making it subject to distracting and shearing forces, such as those seen with lap belt injuries. There is also a higher prevalence of compression and burst fractures in the lumbar spine. These fractures commonly occur when axial loading forces straighten the natural lordosis at the moment of impact. 28

The sacrum forms both the terminal portion of the spine and the central portion of the pelvis, which gives it added stability and makes isolated sacral fractures uncommon. Such fractures are usually caused by direct trauma or falls from a height, or occur as a result of sacral insufficiency secondary to osteopenia, chronic steroid use, or previous pelvic irradiation. More often, sacral fractures occur as a result of high-energy mechanisms and are associated with major pelvic disruption. 29

Indications
Spinal motion restriction should be considered for victims of blunt trauma who sustain an injury with a mechanism that has the potential for causing spinal injury and who have at least one of the following criteria: altered mental status, intoxication, a distracting painful injury (e.g., long-bone fracture), a neurologic deficit, and spinal pain or tenderness (see Box 46.1). 20

Extremes of age and the presence of communication barriers (e.g., language, hearing impairment) may affect the ability to accurately assess the patient’s perception and communication of pain and should lower one’s threshold for spinal precautions. 2,5 It is also important to remember that serious cervical cord injuries can occur in the absence of demonstrable fractures. SCI is common in elderly patients with cervical spondylosis, in whom an arthritic osteophyte may sever a portion of the cord as permanently as a fracture or dislocation. In such cases there may be little subjective pain, and the mechanism of injury may appear seemingly minor. 30

Although data is sparse, patients who have had a seizure in the field are at low risk for spinal injury. 31 Nevertheless, immobilization of postictal patients due to a presumed risk for spinal injury and inability to adequately assess the patient is commonplace. Unfortunately, attempts at immobilization often prompt further patient confusion and agitation, and struggling may exacerbate injuries that do exist (Fig. 46.3). In these situations, when there is concern for a spinal injury based on the mechanism of injury, physical findings (e.g., focal neurologic deficit, facial injuries), or patient complaints, administer sedation rather than, or in addition to, spinal immobilization.

In summary, there is no good evidence that cervical immobilization restricts harmful movement, and the use of c-collars may cause harm. There is evidence that c-collars reduce venous return and hence may cause an increase in intracranial pressure (ICP). Taking a patient out of a comfortable position and placing them in a collar that extends the neck does not make them safer.
**Contraindications**

Spinal immobilization with a c-collar or a backboard following blunt trauma is not necessary in patients with a normal mental status (Glasgow Coma Scale 15), no spinal tenderness or anatomic abnormality, no neurologic findings or complaints, no distracting injuries, and no intoxication (see Box 46.1).19

In general, spinal immobilization with a c-collar and a backboard is contraindicated (or may require modification) when its use could harm the patient, when it is logistically impossible, or when the scene is unsafe (Box 46.2). Good clinical judgment, not blind application of protocols, is essential. For example, if application of a c-collar will cause or mask airway compromise secondary to swelling, an expanding hematoma, or other process, it should not be used. Obviously, if a patient requires a surgical airway, the EMS provider will need immediate, unencumbered access to the anterior aspect of the neck. Sometimes preexisting airways (e.g., tracheostomy tube) and associated equipment prohibit proper application of a c-collar. These situations can often be managed with an improvised cervical immobilizer, such as a collar fashioned from a towel roll or prolonged manual stabilization without traction.

Other conditions that may prevent spinal immobilization or require modification of standard techniques and equipment (e.g., towel roll and manual in-line stabilization) include obesity, impaled objects, underlying respiratory problems or acute respiratory distress, altered mental status (e.g., combative patients due to intoxication or psychiatric illness), and cervical dislocation with fixed angulation or anatomic limitations from preexisting conditions such as ankylosing spondylitis and kyphosis.18

There are also scenarios when spinal motion restriction is logistically difficult or impossible. In a mass-casualty incident or wilderness accident, maintaining spinal motion restriction may be impractical or impossible (Fig. 46.4). Moreover, spinal motion restriction may need to be delayed or modified when the scene poses a significant threat to the patient or providers (see Box 46.2).23 In these situations, the prehospital provider may opt for rapid extrication of the patient from the scene without immobilization of the spine (Fig. 46.5).

In general, victims of penetrating trauma to the head, neck, or torso, such as gunshot wounds, with no evidence of spinal injury should not be immobilized.20 No study has demonstrated worsening neurologic outcomes related to a lack of prehospital spinal immobilization,33 whereas delays

---

**Box 46.2 Contraindications to Prehospital Spine Immobilization**

May be potentially harmful (e.g., prevents identification of airway compromise)

Need for a surgical airway

Presence of a preexisting airway (e.g., tracheostomy tube)

Obesity

Impaled objects

Chronic respiratory diseases (e.g., congestive heart failure) or acute respiratory distress from any cause (e.g., ascites)

Altered mental status or agitation (e.g., intoxicated patients)

Cervical dislocation or anatomic limitation because of preexisting conditions (e.g., ankylosing spondylitis)

Logistically impractical (e.g., mass-casualty incident, hazmat incident)

Unsafe scene:

- Exposure to hazardous material, fire, or smoke
- Building explosion or collapse
- Deep or fast moving water that poses a risk for drowning
- Risk for injury from assault (e.g., gunshot, stabbing, blunt trauma)
- Any other circumstance that the emergency medical service provider deems an immediate danger to the life or health of the patient, provider, or both

*Situations listed in this table can often be managed with modified spinal precautions (see text for details).*

---

**Figure 46.4** During a mass-casualty incident such as a train accident, spinal motion restriction of multiple victims with a low probability of spinal injury may not be practical and may delay other more pressing interventions if resources are limited.

**Figure 46.5** Spinal motion restriction may need to be delayed, modified, or omitted when the scene poses a significant threat to the patient or providers, such as evacuating a wounded soldier during a sniper attack. Rescue from a toxic or poisonous environment, such as hydrogen sulfide or carbon monoxide, is a similar situation.
the best c-collars (when used independently) restrict flexion and extension by only 70% to 75% and overall neck movement by 50% or less.41 A number of studies have evaluated neck motion in volunteers immobilized supine on a backboard with various collars in place.38,39,42–45 Although these studies demonstrated small differences among some of the collars, overall they merely confirm the fact that c-collars do not completely prevent motion of the cervical spine. Interestingly, some cadaver studies have shown the potential for c-collars to actually increase motion and force in the unstable cervical spine.46 Despite these limitations, c-collars remain a widely used component of most spinal motion restriction strategies (Video 46.1).

Cervical Extrication Splints
A large variety of short spine boards and intermediate-stage extrication devices are available for prehospital use (Fig. 46.8). Generally, these devices are manufactured from rigid lightweight material. They have a narrow board design that permits easy application in automobiles or confined spaces and are constructed with multiple openings along the edges to allow for a variety of strapping options. Ideally, these devices should also be translucent so that radiographs can be readily obtained...

Figure 46.7 Cervical collars. A, Philadelphia collar. This two-piece, high-type collar comes in four sizes. The collar supports the head in a dish-shaped contour that is formed when the front and rear halves are joined by Velcro fasteners. When properly sized for a patient, this collar provides excellent support. When applied too tightly, it tends to force the mandible backward and can cause compression of the thyroid in some patients. B, Stifneck collar. This collar is made of high-density polyethylene (a hard material) and padded with semiflexible foam margins. Note the low-reaching anterior panel, which contacts the sternum for additional support.

Figure 46.8 Cervical extrication splints. A, Rigid short boards. B, Kendrick Extrication Device. (Courtesy Ferno-Washington, Inc., Wilmington, OH.)
in the emergency department (ED), and they should allow repeated use and easy cleanup.

Application of a cervical extrication splint (Video 46.2) should not produce unnecessary movement or change the position of the head, neck, shoulders, or torso. In conjunction with a good c-collar, a properly applied cervical extrication splint should effectively limit flexion, extension, lateral motion, and rotational motion of the head, neck, and torso.

One commonly used device that meets all these criteria is the Kendrick Extrication Device (KED) (see Fig. 46.8B). This device consists of two layers of nylon mesh impregnated with plastic and sewn over plywood slats to provide rigidity. It has a nylon loop behind the patient’s head that is continuous with the pelvic support straps for additional strength. Part of its anterior thoracic panels can be folded backward to fit obese, pregnant, or pediatric patients. When properly applied, the KED is a snug-fitting, highly adaptable immobilizer that can be used in even the most adverse circumstances.

When patients require immobilization or extrication (or both) in more difficult or treacherous environments, many EMS providers prefer the LSP half-back (Allied Healthcare Products, Inc., St. Louis, MO), which resembles a KED, but is more rugged and durable. In addition to providing spinal motion restriction, it also acts as a harness and can be used for hauling patients over flat surfaces and for vertical lifts (Fig. 46.9).48

Mosesso and coworkers49 compared six prehospital cervical immobilization devices and concluded that the devices were similar in their ability to limit motion of the cervical spine.

Full-Body Spinal Restriction
Long Spine Boards (Backboards)

Backboards are made of wood or plastic composites and can be either rectangular or tapered in shape (Fig. 46.10B). Most rescuers prefer the tapered type because it takes up less horizontal room when angled into a narrow opening or doorway. In addition, the slight narrowing of these boards on either end enhances the effectiveness of strapping.

Most backboards (Video 46.3) have strategically placed openings along the edges that can be used to secure head-stabilizing devices, strap the patient to the board, or lift the patient. Many also feature runners, usually approximately 2.5 cm thick, on their undersides that serve both as stiffeners and as spacers. They raise the board slightly off the ground so that rescuers can get their fingers under the board during lifting. The runners, however, may make it more difficult to slide a patient onto the board.

Advantages of backboards over full-body splints include their ease of storage, low cost, and extreme versatility. The backboard can be used to slide a victim out of an automobile or to protect a victim during removal of a windshield.

Board splints, as a class, are the least comfortable of all immobilizers. Studies have demonstrated that spinal immobilization on a hard backboard causes head, back, and jaw pain.5,13 Pain in these areas may become severe if patients are left immobilized on these boards for extended periods.9–11 In addition, the pain caused by application of a backboard may be difficult to separate from other sources of pain in a trauma patient and might lead to costly radiographs and unnecessary radiation exposure to the patient.2,50 Discomfort may be minimized by using padding at points of contact between a bony prominence and the board. This concept was reaffirmed by Hauswald and colleagues,51 who found that increasing the amount of padding on a backboard decreases the amount of ischemic pain caused by immobilization. In some cases, tissue ischemia can lead to frank pressure ulcers, particularly in the elderly or nutritional deficient populations.52 Other concerning, well-described risks of long spine board immobilization include respiratory compromise,7 aspiration in the event of vomiting, delay to care for management of emergency injuries, and restraint-related “combativeness.”

Scoop Stretcher

In many cases, a litter that separates longitudinally into two halves, commonly called a scoop stretcher, is an ideal field immobilizer (see Fig. 46.10B). In fact, one study found that using the scoop stretcher caused less spinal motion than did a traditional long backboard and logroll technique.54

The scoop stretcher is designed to split into two or four pieces. It is comfortable, rigid, and adaptable to patients of various lengths and provides unobstructed radiographic transparency of the entire spine. If necessary, it can be applied almost instantly or removed without disturbing the position of the victim. The scoop stretcher also provides good lateral stability because of the troughlike shape of its top surface, and it is stable enough to be used for carrying. When cervical motion restriction is desired, a c-collar can be used with the scoop stretcher. In keeping with current recommendations regarding rigid long boards (see Box 46.1), the scoop should be removed once the patient is transferred to the ambulance cot and the practice of placing a long spine board under the scoop or transferring a patient to a spine board for transport should be abandoned. If the patient must be transported to

---

**Figure 46.9** The LSP half-back (Allied Healthcare Products, Inc., St. Louis, MO). This cervical extrication splint resembles a Kendrick Extrication Device but is more rugged and durable. In addition to providing spinal immobilization, it also acts as a harness that can be used for hauling patients over flat surfaces and for vertical lifts.
body splint, which consists of a polyethylene shell injected with closed-cell foam that is radiographically translucent and provides buoyancy in water (see Fig. 46.10E). This full-body splint has a removable head harness and a thoracic harness, as well as pelvic and lower extremity belts. The space between the lower extremities facilitates wrapping with bandage material in the event of fractures. In addition, it is shaped so that it can easily fit into a basket-type rescue stretcher. Similar spine immobilization systems are available for pediatric patients (e.g., Pedi-Pac, Ferno-Washington, Inc.).

One limitation of the scoop stretcher is the potential for trapping clothes, skin, or other objects between interlocking parts. It also interferes slightly with the ischial section of a half-ring traction splint, but works well with Sager-type devices. The Ferno-Washington model 65 scoop (Ferno-Washington, Inc., Wilmington, OH) is the most widely used stretcher of this type. Other devices such as the CombiCarrier (Hartwell Medical, Carlsbad, CA) and the Scoop EXL (Ferno-Washington, Inc.) offer lightweight polymer construction and additional spinal support (see Fig. 46.10C and D).

Full-Body Splints
A variety of full-body splints are available and may be used by some prehospital providers. One popular device is the Miller body splint, which consists of a polyethylene shell injected with closed-cell foam that is radiographically translucent and provides buoyancy in water (see Fig. 46.10E). This full-body splint has a removable head harness and a thoracic harness, as well as pelvic and lower extremity belts. The space between the lower extremities facilitates wrapping with bandage material in the event of fractures. In addition, it is shaped so that it can easily fit into a basket-type rescue stretcher. Similar spine immobilization systems are available for pediatric patients (e.g., Pedi-Pac, Ferno-Washington, Inc.).

Another full-body splint designed to reduce the pain associated with many of the immobilization devices described previously is the vacuum mattress splint (e.g., EVAC-U-SPLINT, Hartwell Medical, and Immobile-Vac, MDI, Gurnee, IL) (see Fig. 46.10F). It consists of a vinyl-coated polyester envelope filled with thousands of 1.1-mm-diameter polyester foam spheres. A manual or electric vacuum pump is used to evacuate
The movement of the neck if the board is suddenly tilted (e.g., to decrease the risk for aspiration in a vomiting patient). Therefore their use is no longer recommended. Additionally, tightly securing the head to any device without similarly securing the body has the potential to increase forces on the cervical spine during transportation and transfer. Because any manner of strapping is unlikely to completely prevent movement of the torso, the head should always be able to shift axially as a unit with the body.

**Foam Padding**

Padding increases comfort and can help prevent further injury. It can also help support an injured extremity or impaled object or allow an obese or kyphotic patient to lie supine on a long backboard.42 Pregnant women may benefit from padding under the right hip to help shift the gravid uterus off the inferior vena cava and increase venous return.58 Padding applied under the neck and shoulders prevents hyperflexion in children with large occiputs (Fig. 46.13) or in individuals wearing certain types of helmets (e.g., bicycle, motorcycle, rock climbing) that cannot be removed in the field.

**Procedure**

**Cervical Spinal Motion Restriction**

The first priority in cervical spinal motion restriction is maintaining the head and spine in the neutral position. If patients are able to cooperate, instruct them to keep their head and neck in the neutral position and remain still. Next, place both hands on the sides of the patient’s head to manually stabilize the cervical spine and minimize flexion, rotation, or bending. Be sure to avoid cervical traction as it can theoretically increase the risk for SCI. Once the cervical spine has been manually stabilized, examine the neck for swelling, ecchymosis, deformity, bony tenderness, or penetrating wounds. Application of a c-collar follows and is generally a straightforward procedure (Fig. 46.14). The rescuer’s intentions should be thoroughly explained to the patient throughout the procedure.

Once the collar is in place, caution conscious patients against movement of the head. Investigate any persistent complaints...
of pain or dyspnea by removal and possible replacement of the collar while manual stabilization is maintained. The size of collar should be determined from the manufacturer’s suggested guidelines. For example, the Stifneck collar (Laerdal Medical Corporation, Wappingers Falls, NY) is available in various sizes and uses the distance from the top of the shoulder to the chin to determine the appropriate size. Use the tallest collar that does not cause hyperextension. For extremely short necks, a special c-collar such as the No-Neck (Laerdal Medical Corporation) is recommended. In cases in which a c-collar of the proper size is not available, construct an improvised device from available materials (Fig. 46.15).

It should also be remembered that application of a c-collar should not be attempted until the patient’s head has been brought into a neutral position and manual in-line stabilization has been applied.41 If the patient experiences cervical muscle spasm, increased pain, neurologic complaints (e.g., paresthesias, weakness), or airway compromise, immediately halt any further movement of the head and neck. In these situations, immobilize the head and neck in the position they are found or in the position of comfort by using an alternative technique (e.g., blanket, towel roll, manual in-line stabilization).

**Figure 46.13** A, Young child immobilized on a standard backboard. Note how the large head forces the neck into flexion. Backboards can be modified by an occiput cutout (B) or a double mattress pad (C) to raise the chest, the actual clinical consequences of which are unknown. (A–C, Adapted from Herzenberg JE, Hensinger RN, Dedrick DK, et al: Emergency transport and positioning of young children who have an injury of the cervical spine, *J Bone Joint Surg Am* 71:15, 1989.)

**Cervical Collar Application**

**A Posterior-First Method**

1. While one provider applies in-line stabilization (*not* traction!), slide the posterior portion of the collar behind the patient’s neck. Maintain in-line stabilization in the neutral position until the patient is fully immobilized.

2. Bring the front portion of the collar around, under the patient’s chin. Ensure that the chin is well supported by the chin piece. Difficulty positioning the chin piece may indicate the need for a shorter collar.

3. Attach the loop Velcro from the posterior portion of the collar to the hook Velcro on the anterior portion. Recheck the position of the patient’s head for proper alignment. Tighten the collar as needed until proper support is obtained.

**B Anterior-First Method**

1. While in-line stabilization is provided, position the chin piece under the patient’s chin.

2. Slide the posterior portion of the collar behind the patient’s neck.

3. Secure the collar and assess proper placement as described above.

**Figure 46.14** Cervical collar application.
The pelvic straps are not applied properly, considerable slippage may occur when the patient is lifted. Connect the free end of each pelvic strap to buckles located at the patient's hip on the outside of the splint. Once a strap is ready to be buckled, it can be either attached to the buckle on its own side or moved across the patient's lap and engaged with the opposite buckle. Many prehospital care providers prefer the latter method because it allows the patient's knees to remain together without discomfort. It is also a good idea to pad the groin area when placing the pelvic support straps because these straps may cause the patient considerable discomfort.

This procedure may need to be modified for certain injuries and preexisting conditions. For example, patients with pelvic fractures may not tolerate placement of the pelvic support and bottom straps, and the gravid abdomen of a pregnant patient may prevent placement of the middle strap.

Next, secure the head to the device (see Fig. 46.16, step 3). When using the KED, wrap the head panels snugly around the head and neck while another rescuer applies the diagonal head straps. It may be necessary to place padding behind the head to maintain a neutral position. Use the forehead as a point of engagement for one strap and the c-collar for the other.

As a final step, tighten all buckles until the entire splint is firmly in place. The patient can now be moved. If the patient is to be lifted from a vehicle, bring the ambulance cot (with a spine board if needed to facilitate extraction) as close to the patient as possible (see Fig. 46.16, step 4). While one rescuer supports the patient's knees, the other rescuer uses the handholds on the splint to lift the patient. The patient should be rotated and laid in a supine position onto a backboard or cot as needed. Loosen the pelvic straps to allow the legs to be lowered onto the backboard. The legs can then be extended and secured or left in the flexed position with a pillow placed under the knees for support.

If needed, apply a lateral immobilizer to help prevent movement of the head and neck and secure the body with straps. Once the patient is on the stretcher, the thoracic straps may need to be readjusted. The device should be removed as soon as possible, with care to avoid unnecessary movement of the spine until significant injuries are ruled out.

**Thoracolumbar Spinal Motion Restriction**

Studies have shown that healthy individuals are able to self-extricate from a vehicle with less spinal motion than when extricated by emergency medical technicians. Consequently, ambulatory patients or those who are able to self-extricate without undue pain should have a c-collar applied and be allowed to move themselves to an EMS cot. If used, backboards should be removed as soon as possible and patients should not be unnecessarily placed on rigid boards during interfacility transfer (see Box 46.1).

**Sitting Position**

To immobilize patients who require extrication and are found in a sitting position, providers can use a short backboard or commercially available cervical extrication device (e.g., KED) or perform rapid extrication using manual in-line stabilization, a c-collar, and a long spine board.

At least two rescuers should be present to apply an extrication splint to a sitting patient.

Open the device butterfly style and gently slide it behind the victim via a rocking motion (Fig. 46.16, step 1). If necessary, carefully rock the patient forward a few degrees to facilitate placement of the splint. Once behind the victim, free the splint's pelvic support straps from their retainers and allow them to dangle at the patient's sides. Next, bring the lateral thoracic panels around the chest just beneath the patient's shoulders. Grasp these panels and slide the splint upward until the top edges of the panels firmly engage the patient's axillae. Now use the thoracic straps to secure the splint, beginning with the middle strap, then the bottom strap, and finally the top strap (see Fig. 46.16, step 2). The straps should be snug, but not so tight that they interfere with respiration.

Fasten the pelvic support straps next. They can be slipped one at a time beneath the patient's lower extremities and brought directly beneath the pelvis with a back-and-forth motion. If the pelvic straps are not applied properly, considerable slippage may occur when the patient is lifted. Connect the free end of each pelvic strap to buckles located at the patient's hip on the outside of the splint. Once a strap is ready to be buckled, it can be either attached to the buckle on its own side or moved across the patient's lap and engaged with the opposite buckle. Many prehospital care providers prefer the latter method because it allows the patient's knees to remain together without discomfort. It is also a good idea to pad the groin area when placing the pelvic support straps because these straps may cause the patient considerable discomfort.

This procedure may need to be modified for certain injuries and preexisting conditions. For example, patients with pelvic fractures may not tolerate placement of the pelvic support and bottom straps, and the gravid abdomen of a pregnant patient may prevent placement of the middle strap.

Next, secure the head to the device (see Fig. 46.16, step 3). When using the KED, wrap the head panels snugly around the head and neck while another rescuer applies the diagonal head straps. It may be necessary to place padding behind the head to maintain a neutral position. Use the forehead as a point of engagement for one strap and the c-collar for the other.

As a final step, tighten all buckles until the entire splint is firmly in place. The patient can now be moved. If the patient is to be lifted from a vehicle, bring the ambulance cot (with a spine board if needed to facilitate extraction) as close to the patient as possible (see Fig. 46.16, step 4). While one rescuer supports the patient's knees, the other rescuer uses the handholds on the splint to lift the patient. The patient should be rotated and laid in a supine position onto a backboard or cot as needed. Loosen the pelvic straps to allow the legs to be lowered onto the backboard. The legs can then be extended and secured or left in the flexed position with a pillow placed under the knees for support.

If needed, apply a lateral immobilizer to help prevent movement of the head and neck and secure the body with straps. Once the patient is on the stretcher, the thoracic straps may need to be readjusted. The device should be removed as soon as possible, with care to avoid unnecessary movement of the spine until significant injuries are ruled out.

**Recumbent Position**

A patient who is found supine does not require the use of a cervical extrication splint. They should, however, receive initial manual in-line cervical stabilization and a c-collar. The patient can then be secured to a full-body spinal immobilizer, such as a scoop stretcher, backboard, or full-body splint if indicated (see Box 46.1).

**Scoop Stretcher.** A patient who is in a supine position can be moved by means of a scoop stretcher. In a conscious patient, rescuers should explain that they are about to apply a scoop-type stretcher beneath the patient's body. Prior to beginning, apply a c-collar and maintain manual in-line cervical stabilization until the patient is completely secured to the stretcher.

Place the scoop on the ground next to the patient and open the latches that regulate its length. Adjust the length to fit the full length of the patient's body and reengage (lock) the latches.
Next, release the latches at each end and separate the stretcher into two halves. Place each half next to the patient. One rescuer then gently pushes half the stretcher under one side of the patient. In some cases it may be necessary to have another rescuer rock the patient to allow proper positioning. Repeat the procedure with the opposite half of the scoop until both halves are aligned beneath the patient. Engage the latch at the head of the device first. Then bring the lower ends together and engage the foot latch to complete the integrity of the stretcher. Strap the patient’s torso into place and immobilize the head with a suitable lateral neck stabilizer. The patient can then be lifted onto another device for transport, such as a stokes basket or ambulance litter. After placement on another device (e.g., ambulance cot), the scoop stretcher can be removed without disturbing the patient’s position.

Full-Body Spine Boards (Backboards). There are several ways of placing a patient onto a spine board. The precise technique used will depend on the space available and the position of the patient within that space. Apply a c-collar or maintain manual cervical in-line stabilization throughout the procedure and avoid spinal compression or traction. For lengthwise extrication, as from an automobile seat, the patient can be slid, either feet first or head first, onto the backboard. It is important that the patient be moved as a unit during this process. Place one end of the backboard on the seat or doorsill of the automobile. One rescuer stabilizes the opposite end while at least two other rescuers lift and slide the patient’s body onto the backboard. Loosen the pelvic straps, and secure the patient to the board.

Figure 46.16 Kendrick Extrication Device. Use of other short boards follows the same principles as depicted here.
Full-Body Spine Board (Backboard): Logroll Maneuver

Position one rescuer at the patient’s head to apply manual in-line stabilization. This rescuer oversees and directs all body movement throughout the procedure.

Position the backboard next to the patient’s body. Note that a lateral neck stabilizer has been preapplied to the board.

When the rescuer at the patient’s head gives the command, roll the patient onto his side, examine the patient’s back, and slide the backboard under the patient.

Roll the patient back onto the board when the head rescuer gives the command. Center the patient on the board before applying the straps.

Strap the patient to the board. Proper strap placement (chest, pelvis, and legs) and firm contact between the straps and the patient are important in limiting lateral motion.

Apply a lateral neck stabilizer, like the Ferno Universal Head Immobilizer shown above. Secure it in place using the supplied straps or tape.

Figure 46.17 Full-body spine board (backboard): the logroll maneuver.

patient’s head to apply manual in-line cervical stabilization (Fig. 46.17, step 1). It is this person’s responsibility to oversee and direct body movement throughout the procedure. Next, position the backboard alongside the patient’s body (see Fig. 46.17, step 2). To minimize thoracolumbar movement, extend the patient’s arms at the sides with the palms resting on the lateral aspect of the thighs. To keep the patient from reaching for a rescuer or object during transfer, some rescuers prefer to have patients cross their arms across the thorax. Place the backboard against the injured (more painful) side of the body.
so that the patient can be rolled onto the uninjured (less painful) side. Position the other rescuers on the side that the patient will be rolled toward, with one rescuer at the midchest and the other at the legs. The rescuer at the chest should reach across the victim and take hold of the shoulder and hips while the other rescuer grasps the hips and lower part of the legs. When everyone is ready, the rescuer at the head gives the command to roll the patient onto the side (see Fig. 46.17, step 3). If possible, examine the patient’s back at this point. Slide the backboard under the patient, and when everyone is ready, the rescuer at the head gives the command to roll the patient back onto the board (see Fig. 46.17, step 4). Before applying straps, it is often necessary to center the patient on the board.

To center the patient on the long spine board, support the cervical spine, shoulders, and hips and slide the patient first caudally, then cephalad, in a zigzag pattern to avoid applying uneven lateral or rotational force onto the spine. Alternatively, during lateral extraction, a recumbent patient can be slid sideways onto the spine board. This improvised technique also requires the presence of three or four rescuers, one of whom can maintain control of the patient’s head and neck.

The lift-and-slide technique, sometimes referred to as a six-person lift or bridge lift, is a technique which may reduce the amount of motion at an unstable vertebral segment when compared with the logroll. Del Rossi and coworkers demonstrated this in cadaver studies, but the difference has never been shown to be clinically relevant and the logroll remains nearly ubiquitous. One disadvantage of this technique is the inability to visually assess the spine during the lift.

To perform the lift-and-slide, rescuers line up opposite each other on both sides of the patient, ideally with at least two rescuers per side. An additional rescuer is located at the patient’s head and provides in-line stabilization and coordinates the lift. When all are ready, they place hands underneath the patient in an alternating fashion and lift the patient directly upward a few inches. At this point, another rescuer inserts the carrying device (e.g., backboard, stokes basket, vacuum mattress) and the patient is lowered and secured. For heavier patients, additional rescuers may be needed. Rescuers may place their head on the shoulder of the person across from them to form an arch and reduce the strain on the lower back. In some cases, such as in environments with limited space, standing over the patient and lifting straight upward may be required.

When necessary, various techniques can be used to secure a patient to the backboard. In addition to the standard thoracic, pelvic, and lower extremity straps, use of an abdominal strap may reduce lateral motion without compromising respiration. Proper strap placement and firm contact between the straps and the patient are also important in limiting lateral motion (see Fig. 46.17, step 5). Commercial products such as Spider-Straps are also available or webbing may be used in a zigzag alternating fashion.

After the body has been strapped to the board, the head can be secured. If necessary, padding should be placed under the occiput to maintain the head in the neutral position. Apply a lateral neck stabilizer (e.g., foam blocks, HeadBed device [Laerdal Medical Corp.]) and secure the head in place with tape or straps (see Fig. 46.17, step 6). Most taping techniques involve the use of one piece across the forehead and one piece across the c-collar. Remember, taping across the neck may make airway management difficult. Placing straps across the head and neck may cause hyperextension or axial stretching of the neck. For this reason, omitting the lower strap is recommended. The head should only be secured as tightly as necessary to minimize motion; it should be able to easily slide axially if the torso shifts during transportation. Thus, if the body is not secured to the board, the head should similarly not be fastened to the board.

**Standing Position**

Traditionally, standing patients were placed on a backboard by positioning the board behind the patient, stabilizing the head, and lowering the patient and the backboard backward until the patient was lying on it. However, there is no evidence that this protects the spine from further injury. Current recommendations are to apply a c-collar and allow ambulatory patients to walk to the stretcher and assist them into supine position on the mattress. Nevertheless, there may still be a role for using a backboard or other rigid device placed behind the patient when a stretcher is not available (as in a wilderness or technical rescue environment) and the patient is unable to comfortably lower themselves to a ground-level extrication device, such as a long board or scoop stretcher. In this case the device should be placed behind the patient while at least one rescuer on each side (more may be needed for large patients) grasps the handles underneath the patient’s arms and slowly tips the patient and board backward until the board is completely on the ground. There is no need to provide manual head stabilization during this maneuver, but the patient should be reminded to keep his head still.

**Pediatric Patients**

Little information is available on the proper selection and application of spinal immobilization devices for children. Most of the data available were derived from studies of adults and might not be applicable to children, especially those under 8 years of age. Half the total growth in head circumference is achieved by the age of 18 months, giving children a disproportionately large head in comparison to the rest of the body. Before 8 years of age, these anatomic and developmental differences result in a higher incidence of upper cervical spine injuries (C1–C2). Because injuries in this area are frequently unstable, proper cervical immobilization in the neutral position is critically important.

In the neutral position, the pediatric cervical spine is normally lordotic or extended. However, because the occiput is large, positioning the child’s body on a standard backboard may force the neck into flexion or a relative kyphosis. The clinical significance of this is currently unclear, but theoretically it may be hazardous for young children. Therefore the standard backboard should be modified to adapt to the child’s larger head size. As a rough guide, the external auditory meatus should be on the same level as the midpoint of the shoulder. Suggested modifications include a cutout in the backboard that accommodates the occiput or a pad under the back at the level of the chest (see Fig. 46.13). If not modified, the standard backboard in conjunction with the disproportionately large head of a child may force the neck into hyperflexion and potentially aggravate an underlying cervical spine injury. Nylander and Treloar showed that all children required elevation of the back (mean height, 25.4 ± 6.7 mm) for correct neutral position on a spine board. Children younger than 4 years required greater elevation than did those 4 years or older. It must be pointed out, however, that there have been no published reports of an SCI resulting from the use of standard...
immobilization techniques and equipment in children. Current guidelines recommend spinal precautions in children under 8 years who have suffered a sufficient mechanism of injury.

Complication

Cervical Immobilization

Improper application of a c-collar can occur if the wrong size is used or too little care is exercised during placement. The best means of preventing either error is strong clinician involvement in training and continuing education of EMS crews together with vigorous feedback regarding correct and incorrect application. In addition, adherence to the manufacturer's collar-specific recommendations for size and application should be emphasized.

A collar that is too small for a patient may be either too tight for the girth of the neck or too short to provide adequate immobilization. Too large a collar commonly results in hyperextension, which can exacerbate a preexisting spinal injury. Underlying spinal abnormalities from conditions such as ankylosing spondylitis, rheumatoid arthritis, or kyphosis can also contribute to exacerbation of injuries with c-collar application.

Improper or prolonged application of an extrication collar may impede venous return and raise ICP. Although the clinical significance of increased ICP produced by cervical immobilization is still unknown, two studies have confirmed that application of a rigid c-collar causes a statistically significant and sustained rise in ICP. Kolb and colleagues reported a 24.8-cm H₂O increase in cerebrospinal fluid pressure in 20 adult patients undergoing lumbar puncture. Hunt and associates reported a 4.6-mm Hg mean rise in ICP in 30 patients with severe traumatic brain injury. The largest rise in ICP was noted in patients with a baseline ICP higher than 15 mm Hg. The authors of these studies concluded that the elevation in ICP produced by cervical immobilization might have deleterious effects in patients with acute or sustained intracranial hypertension.

Long-term use of the Philadelphia extrication collar (Philadelphia Cervical Collar Co., Westville, NJ) as part of the treatment plan for an underlying cervical spine injury has been associated with pressure ulcers of the scalp. Because some collars (e.g., Philadelphia and Stifneck [Laerdal Medical Corporation]) have been shown to exert higher capillary closing pressure at contact points, it is suggested that only collars with favorable skin pressure patterns and superior patient comfort (e.g., Nec-Loc, Jerome Medical, Moorestown, NJ) be used for long-term application.

One final complication should be mentioned. A patient who, for whatever reason, actively resists placement of a c-collar or other splint should not be forced to wear it (see Fig. 46.3). Postictal and intoxicated patients may present such a dilemma. Immobilization of a combative patient cannot be accomplished without considerable muscular exertion, not only by rescuers, but also by the patient. If fractures do exist, it is possible that struggling and using muscles attached to these fractured fragments may actually cause further injury. If the patient permits manual stabilization, this should be used as an alternative. Sedation should be used judiciously to enhance compliance.

Thoracolumbar Immobilization

In general, complications are more likely to occur from failure to immobilize spinal injuries rather than from the process of immobilization itself. When complications do arise, they may be related to improper choice or use of equipment or prolonged immobilization.

Victims are generally strapped in place on a spine board to prevent sliding during transport. If too few straps are used or if the straps are loosely applied, motion during transport can occur. Even when applied correctly, spinal immobilization on a hard board may be extremely uncomfortable for patients and may induce pressure-related tissue damage. In one study, 100% of healthy volunteers reported significant pain after only 30 minutes on a long spine board. Occipital headaches, as well as mandibular, lumbar, and sacral pain, developed in these individuals. Other studies have demonstrated elevated tissue interface pressure in patients on spine boards without air mattress padding. These studies underscore the need to use adequate padding when patients require immobilization on a long board, as well as removal from the board as soon as possible.

Excessive strapping can interfere with respiratory function in both children and adults. In healthy children 6 to 15 years old, forced vital capacity (FVC) has been shown to decrease from 4% to 5% during spinal immobilization. Totten and Sugarman evaluated the effect of two spinal immobilization methods (wooden backboard and vacuum mattress) on eight respiratory function measurements in healthy volunteers. In comparing baselines for each method, six of the eight measures (FVC, FVC%, forced expiratory volume in 1 second [FEV1], FEV1%, peak expiratory flow [PEF], and forced expiratory flow [FEF₂₅₋₇₅%]) were reduced an average of 15%. Although this may not be a problem in healthy volunteers, the effects on patients with chest trauma or preexisting respiratory disease may be significant.

Patients immobilized on a backboard are also at risk for aspiration if they vomit. If vomiting does occur, logroll the patient and board as a unit and suction the airway as needed. Although this procedure may be associated with some spinal movement, airway protection takes priority.

Conclusion

Although spinal immobilization has not been shown to decrease the likelihood of spinal injury and is associated with a variety of complications, prehospital spinal immobilization remains commonplace. Nevertheless, recent evidence supports a widespread shift to selective spinal immobilization by prehospital providers. Until these new recommendations become universal, emergency medicine practitioners should know when spinal immobilization is indicated, recognize properly and improperly applied devices, minimize unnecessary immobilization time, and work with prehospital providers to ensure appropriate indications and proper use.

EXTREMIT Y IMMOBILIZATION

Upper Extremity

Background

The earliest evidence of upper extremity splinting comes from the Egyptians circa 300 BC. In 1903 archaeologists from the Hearst expedition discovered two specimens whose open fractures had been treated with wooden splints and bandages. Their bones showed no signs of healing, so the individuals probably died soon after their injuries; however, numerous
other ancient Egyptian specimens have demonstrated evidence of well-healed forearm fractures.\textsuperscript{76}  

The purpose of splinting is to prevent motion of broken or dislocated bone ends. Carefully applied splints decrease pain while minimizing further damage to muscles, nerves, and blood vessels. Splinting also reduces the risk of converting a closed injury to an open one.\textsuperscript{77} Nevertheless, injuries to adjacent structures may still occur, so circulation, motor function, and sensation distal to the injury must be assessed early and monitored closely during transport.

Over the last 3 decades, prehospital splinting materials, equipment, and techniques have evolved considerably. Today’s prehospital providers can choose from a wide assortment of ready-to-use splints made from a variety of sturdy lightweight materials.

**Indications and Contraindications**

Indications for splinting an extremity are usually clear. Pain after trauma, with or without deformity, should arouse suspicion of underlying bone or joint injury. Other signs include swelling, discoloration, deformity, crepitus, and loss of neurovascular function. However, absence of these findings does not rule out an underlying fracture or dislocation. Thus, in most cases where a musculoskeletal injury is suspected, a splint should be applied and maintained.

There are no absolute contraindications to splinting suspected extremity fractures or dislocations. However, in the setting of multisystem trauma with life-threatening injuries, rapid transport may be more important than extremity splinting. Averting loss of life takes precedence over loss of limb.

**Equipment**

Various types of splints are currently available for immobilizing upper extremity injuries. Emergency care providers should be well trained and familiar with their equipment. The type of splint used is less important than the expertise of the provider. Upper extremity splints can be divided into two basic types: rigid and soft.\textsuperscript{78}

**Rigid Splints**

Rigid and semirigid splints are the mainstays of EMS fracture care (Fig. 46.18-A–C). These splints are made of many different materials, including cardboard, plastic, aluminum, wire, and wood. They are fastened to the injured extremity with tape, gauze, cravats, or Velcro straps. Rigid splints are generally nonflexible (some commercially available splints may have some flexibility in their design) and, when applied properly, immobilize the limb in a rigid fashion to maintain stability. Although most commercial rigid splints are prepadded, many will still benefit from additional soft padding to cushion the splint and increase comfort. This is particularly true over bony prominences. When applying rigid splints, leave the fingertips exposed so that the distal circulation can be continuously monitored.

Cardboard splints are excellent for long-bone fractures in the upper part of the arm. They can be formed into many shapes and are easy to apply, inexpensive, lightweight, radiolucent, and compatible with magnetic resonance imaging. Splints made from wax-impregnated cardboard are also water resistant. Plastic, aluminum, wire, and wood splints, though less malleable, are also good choices. An inexpensive aluminum splint that is popular with emergency care providers and is found in many wilderness medical kits is the SAM Splint (Sam Medical Products, Portland, OR) (see Fig. 46.18D). The SAM Splint is built from a thin core of soft aluminum alloy sandwiched between two layers of closed-cell foam. The SAM Splint is extremely pliable. Bent into any of three simple curves, it is extremely strong and provides support for any fractured or injured extremity. In addition, it is water resistant, lightweight, radiolucent, reusable, and not affected by extreme temperatures or altitudes. These characteristics make it an ideal tool for emergency care providers and outdoor enthusiasts.

Vacuum splints (see Fig. 46.18E) are a special type of rigid splint in which the air is evacuated from a closed bag containing tiny foam beads. Flexibility of the splint before removal of the air allows molding of the splint to conform to the patient’s position. Removal of the air compresses the contents into a solid mass and results in a rigid splint that encases the extremity, immobilizing it in the position it was found. As a result, vacuum splints in general are more comfortable than other rigid splints. Vacuum splints are radiolucent and do not apply external pressure, thus ensuring maximum circulation to the injured extremity.

**Soft Splints**

Soft splints include air splints, pillows, slings, and swaths. Depending on the type of injury being treated, immobilization with pillows, slings, or swaths alone may be inadequate because they allow significant motion. Therefore, when treating fractures and dislocations, these splints are most effective when used in conjunction with some form of a rigid device. However, even when used alone, pillows, slings, or swaths provide cushioning and some limitation of motion that may be adequate for less severe injuries such as sprains and contusions. Table 46.1 provides splint recommendations for a variety of upper extremity injuries.

Air splints are soft splints that become rigid when inflated. Besides providing immobilization, they help compress the underlying soft tissue, reducing local hemorrhage. These devices are sensitive to differences in atmospheric pressure and temperature. Therefore their inflation must be closely

**TABLE 46.1 Management of Specific Upper Extremity Orthopedic Injuries**

<table>
<thead>
<tr>
<th>SITE</th>
<th>SUGGESTED IMMOBILIZATION TECHNIQUES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clavicle</td>
<td>Sling and swath</td>
</tr>
<tr>
<td>Shoulder</td>
<td>Sling and swath as it lies</td>
</tr>
<tr>
<td>Humerus</td>
<td>Cardboard or vacuum splint with a sling and swath</td>
</tr>
<tr>
<td>Elbow</td>
<td>Cardboard or vacuum splint as it lies, realign if circulation deficit exists</td>
</tr>
<tr>
<td>Forearm</td>
<td>Cardboard, malleable metal, air, or vacuum splint with a sling and swath</td>
</tr>
<tr>
<td>Wrist</td>
<td>Pillow, cardboard, malleable metal, or vacuum splint applied in the position found</td>
</tr>
<tr>
<td>Hand</td>
<td>Pillow, cardboard, or malleable metal splint in the position of function</td>
</tr>
<tr>
<td>Finger</td>
<td>Tongue depressor or small malleable metal splint</td>
</tr>
</tbody>
</table>

---

931
GENERAL SPLINTING GUIDELINES:
- Remove unnecessary clothing and visualize the extremity.
- Check neurovascular status before and after splinting.
- Cover open wounds with a dry sterile dressing.
- Immobilize the joint above and below the fracture site.
- Cool and elevate the injured area if possible.
- Frequently reassess neurovascular status and pain.

Figure 46.18 Upper extremity splints. A–C, Padded rigid splints. D, SAM Splint. E, EVAC-U-SPLINT vacuum splint. F, Soft towel splint (a pillow can be used in a similar manner). G, Sling and swath. (D, Courtesy SAM Medical Products, Portland, OR; E, courtesy Hartwell Medical, Carlsbad, CA.)
monitored to ensure that the underlying tissue is not being subjected to pressure-induced ischemia and the development of a compartment syndrome. One study suggested a maximum splint pressure of 15 mm Hg to reduce the risk for ischemia.79 With long ambulance transport times, deflate the splint for 5 minutes every 1.5 hours.

Disadvantages of most air splints include an inability to continuously monitor pulses and susceptibility to puncture. Air splints are designed to conform to a specific shape when inflated and should not be used on angulated fractures. In addition to being radiolucent, some types can be inflated with a refrigerant to provide concurrent cooling.

Pillow splints (see Fig. 46.18F) can be fashioned from any soft bulky material and are excellent choices for hand or wrist injuries. These splints are extremely comfortable and are easily applied.

Slings and swaths are generally used in combination with a rigid or a soft splint. When used alone they can effectively immobilize injuries to the shoulder, clavicle, and humerus (see Fig. 46.18G).

 Procedures
To properly apply a splint to an injured extremity, employ the following general guidelines. Communicate with patients to ensure that they understand what is being done at all times. When possible, administer an appropriate analgesic to make splint application less painful. Remove any unnecessary clothing to adequately visualize the injured extremity. Stabilize the fracture site manually to help limit unnecessary movement and prevent further injury. Check the patient’s neurovascular status (i.e., pulse, motor function, and sensation) before and after the application of a splint. For a severely angulated extremity with neurovascular compromise, reduce the deformity with gentle longitudinal traction (not exceeding 10 lbs of pressure) before splinting. Make only one attempt at fracture reduction. If resistance or pain is encountered, splint the extremity in the position found. Cover open wounds with a dry sterile dressing before applying the splint. When possible, immobilize the joint above and below a suspected fracture site. Once the splint has been applied, cool and elevate the injured area to help reduce swelling and pain. Assess distal neurovascular status frequently. Any deterioration requires immediate evaluation of the splint to determine whether excessive pressure is being applied. In addition, frequently assess and treat pain.

 Rigid Splints
To apply a rigid splint, have an assistant provide support and gentle traction above and below the injury. Apply the splint on the side of the extremity away from any open wounds. The splint should be large enough to immobilize the joint above and below a suspected fracture or the bone above and below a dislocation and should be well padded to reduce the risk for pressure necrosis. Secure the splint to the extremity with gauze, tape, cravats, or Velcro straps (see Fig. 46.18A–C).

Apply vacuum splints in much the same manner as other rigid splints. While an assistant stabilizes the injured site and applies traction, wrap the splint around the extremity and secure it in place with the attached straps. Evacuate the air from the splint with a hand pump until the splint becomes rigid.

 Soft Splints
The application procedure for an air splint depends on whether the splint is equipped with a zipper. If the splint does not have a zipper, first place the splint over the rescuer’s arm until the bottom edge lies above the wrist. Use this hand to grasp the hand of the patient’s injured extremity and use the free hand to provide support and gentle traction above the injury (Fig. 46.19A). An assistant then slides the splint onto the patient’s arm (see Fig. 46.19B). Smooth out any wrinkles and inflate the splint until finger pressure makes a slight indentation (see Fig. 46.19C). Open a zippered air splint and place it around the injured area. Close the zipper and inflate the splint as described previously. With air splints that completely enclose the hand, continuously assess the distal circulation by checking fingertip color, temperature, and capillary refill.

Apply pillow splints by encasing the injury in the pillow and securing it with tape, cravats, or gauze (see Fig. 46.18F). If possible, keep the nail beds exposed to allow frequent neurovascular checks.

To apply a sling, have an assistant support the injured arm in a flexed position across the patient’s chest (Fig. 46.20). Place the long edge of the triangular bandage lengthwise along the patient’s side opposite the injury with its tip over the uninjured shoulder. Bring the other tip over the injured shoulder to enclose the arm in the sling. Adjust the sling so that the arm rests comfortably with the hand higher than the elbow. Tie the ends of the sling together at the side of the neck and pad the knot for comfort. Finally, wrap the point of the sling
Injuries to the lower extremities, including sprains, fractures, and dislocations, are also commonly encountered by prehospital care providers. As with upper extremity injuries, application of a splint is an essential part of the prehospital management of lower extremity injuries (Fig. 46.21). Many of the principles, techniques, and complications discussed for upper extremity splinting also apply to injuries to the lower extremity; the SAM Splint (Sam Medical Products) (see Fig. 46.18D) and pillow splint (see Fig. 46.18F) are just two examples. Table 46.2 provides recommendations for immobilizing a variety of lower extremity injuries. One fundamental difference between splinting upper and lower extremity injuries is use of a traction splint in the management of femoral fractures. The remainder of this section focuses on the use of traction splints (Video 46.4).

**Conclusion**

Injuries to the upper extremities, though not life-threatening, may be limb-threatening and can have significant immediate or long-term effects. Maintain a high index of suspicion for underlying neurovascular injury; check neurovascular status before and after applying a splint and frequently during transport. When possible, administration of appropriate analgesia will be greatly appreciated by the patient.
Traction splints are designed to align fracture segments and restore the cylindrical shape of the thigh (Fig. 46.23). This in turn increases tissue pressure, decreases the potential space for blood loss, and inhibits further hemorrhage. In addition, traction splints help reduce pain, prevent further damage to neurovascular structures, and reduce the incidence of fat embolism.82,83

During World War I, the modified splint was credited with reducing the mortality rate associated with fractured femurs from 80% to 15%.81 Since then, several additional modifications that carry the name of their inventors (e.g., Glenn Hare, Joseph Sager, Allen Klippel) have furthered the development of lower extremity traction splints. Today, traction splints are a nearly universal piece of equipment found on most ambulances, and their application is an integral part of the prehospital care provider’s skill set. The main purpose of the traction splint is to immobilize a fractured femur.82,83

In the setting of a fractured femur, muscle spasm and overlap of fragments may cause the thigh to lose its cylindrical shape and adopt a more spherical appearance (Fig. 46.22).81 The resultant decreased tissue pressure and increased volume may allow 1 to 2 L of blood to accumulate at the fracture site.

### TABLE 46.2 Management of Specific Lower Extremity Orthopedic Injuries

<table>
<thead>
<tr>
<th>SITE</th>
<th>SUGGESTED IMMOBILIZATION TECHNIQUES</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hip</td>
<td>Secure the injured leg to the uninjured leg, ideally padded between, and/or with a long external splint secured proximally and distally. Full body immobilization (backboard, Reeves flexible stretcher [Fisher Scientific, Hampton, NH], scoop stretcher, vacuum mattress, etc.), with the limb padded and secured.</td>
</tr>
<tr>
<td>Midshaft femur</td>
<td>Traction splint, contralateral leg stabilization, or rigid splint</td>
</tr>
<tr>
<td>Knee</td>
<td>Cardboard or vacuum splint in the position found</td>
</tr>
<tr>
<td>Tibia/fibula</td>
<td>Cardboard, air, or vacuum splint</td>
</tr>
<tr>
<td>Ankle</td>
<td>Pillow, cardboard, or air splint</td>
</tr>
<tr>
<td>Foot</td>
<td>Pillow or air splint</td>
</tr>
<tr>
<td>Toe</td>
<td>Tape to the adjacent toe</td>
</tr>
</tbody>
</table>

Traction splints are designed to align fracture segments and restore the cylindrical shape of the thigh (Fig. 46.23). This in turn increases tissue pressure, decreases the potential space for blood loss, and inhibits further hemorrhage. In addition, traction splints help reduce pain, prevent further damage to neurovascular structures, and reduce the incidence of fat embolism.82,83
Indications
Application of a lower extremity traction splint is indicated whenever an isolated midshaft femur fracture is suspected.\textsuperscript{78,84,85} Clinical signs of a fractured femur include thigh pain associated with limb shortening, angulation, crepitus, swelling, or ecchymosis.

Contraindications
Do not use traction splints on patients with pelvic fractures, hip injuries with gross displacement, significant injuries involving the knee, or avulsion or amputation of the ankle or foot.\textsuperscript{85} Use of traction splints is not recommended in the presence of an associated distal tibia-fibula or ankle fracture in the same extremity. In these circumstances, the amount of traction required to realign the fractured femur can distract the distal fracture site. A variety of rigid splints may be considered in these settings.

There has been some controversy over whether a traction splint should be applied to an open femoral fracture for fear that the use of traction may allow contaminated bone fragments to retract into the wound. A 2011 update to the International Trauma Life Support guidelines recommends use of traction splints on open femur fractures only in the austere environment or when access to definitive care will be significantly delayed.\textsuperscript{86} When possible, providers should use copious irrigation to remove foreign materials prior to application of the splint. If traction does force contaminated bone fragments into the wound, this information must be relayed to the receiving clinician and antibiotics initiated as soon as possible. In those situations where a traction splint is not applied, a variety of rigid splints can be used to immobilize the bony fragments in the position that they were found. In either case, stabilization of the fracture site to prevent further hemorrhage, neurovascular damage, or soft tissue injury should take precedence, while making every effort to minimize the risk for increased contamination.

Equipment
Regardless of the type or manufacturer, the basic traction splint consists of a metal frame that extends from the proximal end of the thigh to an area distal to the heel. The padded proximal end fits against the ischial tuberosity and serves as the anatomic fixation point. The proximal portion of the splint may be a ring that encircles the proximal end of the thigh, a partial ring, or a padded bar. At the distal end of the splint there is typically a ratchet-type device that when engaged, creates traction on the distal end of the femur. All traction splints also have several soft elastic straps that support the thigh and leg.\textsuperscript{85}

Commonly used traction splints include the Hare Traction Splint (Dyna-Med, Lexington, KY), Kendrick Traction Device (Ferno-Washington, Inc.), Sager Emergency Traction Splint (Minto Research and Development, Redding, CA), and the Ferno-Trac Traction Splint (Ferno-Washington, Inc.) (Fig. 46.24). Each of these splints has its own advantages, disadvantages, and unique method of application. For example, traction splints that use a half-ring design apply countertraction to the ischial tuberosity from below the shaft of the femur. This produces flexion at the hip joint of up to 30 degrees and will not allow complete fracture alignment unless the patient is in a reclining position approximately 30 degrees from horizontal or the injured extremity is elevated to create the same angle. Traction splints that do not use a half ring do not cause hip flexion.

Procedure
Application of the Ferno-Trac Traction Splint (Ferno-Washington, Inc.) and the Sager Emergency Traction Splint
Prehospital Immobilization

Chapter 46

approximately 6 inches (15 cm) beyond the ankle. With the extremity slightly elevated, place the traction splint (e.g., Ferno-Trac Traction Splint [Ferno-Washington, Inc.]) under the injured leg and rest it firmly against the ischial tuberosity. To ensure that the injured extremity will remain elevated once manual traction has been released, unfold the heel stand and lock it into place. Place the Sager Emergency Traction Splint (Minto Research and Development) between the legs and against the ischial tuberosity, carefully avoiding or protecting the genitalia. Secure only the most proximal ischial strap at this point.

Next, place the ankle harness immediately above the medial and lateral malleoli and attach it to the distal end of the traction splint. Ferno-Trac (Ferno-Washington, Inc.) and other similar traction splints use a ratchet mechanism to apply constant (static) traction on the ankle strap. For dynamic quantifiable splints such as Sager-type devices, gently extend the inner shaft of the splint until the desired amount of traction is achieved, either by comparing leg length or using the quantitative scale to set the desired force. Both static and dynamic traction mechanisms may lose traction as muscle spasm decreases.

Figure 46.25 Application of the Ferno Traction Splint. Note that the splint should be adjusted so that it is 12 inches longer than the uninjured leg before application. (Reproduced and modified with permission from Ferno-Washington, Inc., Wilmington, OH.)
Prior to application of the splint, get a rough measure of the length of splint needed. Extend the splint so that the wheel is at the heel.

Grasp the Kydex buckle and slide the thigh strap up under the leg so that the perineal cushion is snug against the perineum and ischial tuberosity.

Tighten the Kydex buckle thigh strap to draw the perineal-ischial pad to the lateral portion of the crotch.

Apply the ankle harness tightly around the ankle above the medial and lateral malleoli of the ankle. Check the posterior tibial and dorsalis pedis pulses before hitch application and after traction is established.

Shorten the loop of the harness connected to the cable ring by pulling on the strap threaded through the square “D” buckle.

Extend the inner shaft of the splint by opening the shaft lock and pulling the inner shaft out until the desired amount of traction is noted on the calibrated wheel. As a rough guide to determine the amount of traction needed, apply 10% of body weight to a maximum of 22- to 25-lb (10- to 25-kg) traction.

Apply the second longest thigh strap around the knee. Use padding as needed. Next, apply the shortest 6-inch wide strap over the ankle harness and lower the leg.

Apply figure-eight strap around both ankles. The patient’s leg is now secured, traction is controlled, medial and lateral shift of the distal fragment and internal and external rotation is prevented.

Figure 46.26 Application of the Sager traction splint. Note that the splint can alternatively be strapped to the outer portion of the leg. In addition, it is possible to splint both legs with a single splint.
and leg length increases, requiring reevaluation and possible adjustment.\textsuperscript{51,87,88} Regardless of which splint is being used, apply traction gradually to approximately 10\% of body weight or a maximum of 7 kg (15 lb) (15 kg [30 lb] for bilateral fractures). Only rarely will more traction be required.\textsuperscript{84,85} The goal is to stabilize the fracture and maintain proper limb alignment. Use the least amount of force necessary to accomplish this task.\textsuperscript{89}

Before moving the patient, place supportive straps around the thigh, knee, and distal end of the leg to stabilize the extremity. After application of the splint, be sure to recheck distal neurovascular status and then secure the patient and splint firmly to the ambulance stretcher. Take extra care while moving the patient and when closing any transport vehicle door to avoid unnecessary movement and further injury. If the splint extends beyond the dimensions of the stretcher, additional support for the splint may be needed (e.g., short spine board or cardiopulmonary resuscitation board) to ensure that the injured extremity remains elevated throughout transport. Loss of pulses with application of a traction splint requires that the position of straps and the amount of traction be reassessed immediately. It is also important to recheck the position of the splint and the patient’s neurovascular status after moving the patient. Finally, remove the traction splint in reverse order of application.

Special Considerations
The Sager Models S304 (Form III Bilateral) and SX404 (Extreme Bilateral) (Minto Research and Development) offer a unique advantage in confined spaces in that they can be used to immobilize both legs simultaneously with only one splint. In addition, neither splint extends beyond the patient’s heels, thus making them more versatile for use in helicopters, fixed-wing aircraft, and smaller van-type ambulances.

Complications
Complications are generally the result of incorrect application and include pain, ongoing hemorrhage, peroneal nerve injury, perineal injury, movement at the fracture site, ligamentous injury, injury to unrecognized fractures elsewhere, or further neurovascular compromise.

Conclusion
When properly applied, traction splints reduce the pain, hemorrhage, and potential injury to adjacent structures that are associated with femoral fractures. Close attention to the manufacturer’s application instructions and frequent patient reassessment will help reduce complications.

PELVIC IMMOBILIZATION

Background
Pelvic fractures cause significant morbidity and mortality, especially in the elderly. The decreased bone density in older individuals contributes to the likelihood of sustaining pelvic fractures from low- to moderate-energy mechanisms, such as falls from a standing height. The findings on physical examination may be subtle, such as slight asymmetry of the lower extremities, so a high index of suspicion is important. In younger individuals, pelvic injuries are usually caused by high-energy mechanisms, most commonly motor vehicle collisions.\textsuperscript{90} Major pelvic trauma can lead to severe, uncontrollable hemorrhage, hypovolemic shock, and multisystem organ failure (Fig. 46.27).\textsuperscript{91}

In the prehospital setting, early stabilization of fracture segments to control hemorrhage, gentle handling, and immediate transport to a trauma center are the cornerstones of treatment.\textsuperscript{92} However, even the most experienced prehospital provider may find it difficult to identify pelvic injuries in the field (the exception being an open-book fracture). Hence, prehospital providers should not hesitate to stabilize the pelvis if a fracture is suspected.

Indications and Contraindications
Apply a pelvic circumferential compression device to any patient with significant pelvic pain, tenderness, or evidence of pelvic instability after trauma, especially if the patient is hypotensive.\textsuperscript{93,94} There are no contraindications to pelvic stabilization in patients with presumed pelvic fractures.

Procedure
Stabilization of the pelvis consists primarily of compressing the bony ring. This can be accomplished using a variety of devices, including a simple sheet, commercially available pelvic circumferential compression devices (also known as pelvic binders) such as the SAM Sling (SAM Medical Products), the T-POD Pelvic Stabilization Device (Pyng Medical Corp., Richmond, British Columbia, Canada), vacuum “beanbag” mattress splints, and pneumatic antishock garments (PASGs) (Fig. 46.28). With the advent of commercial pelvic binders, use of a PASG for pelvic fracture stabilization is no longer recommended.\textsuperscript{95,96} When using any pelvic stabilizing device remember that aggressively moving patients or aggressively palpating or rocking the injured pelvis can precipitate further vascular injury or clot disruption and lead to life-threatening hemorrhage.

To apply a sheet, place it over a long spine board and then gently logroll the patient by no more than 15 degrees into position on the board.\textsuperscript{97} Alternatively, slide the sheet under the patient by passing it below the lumbar or thigh area and gently maneuvering it into position. In some cases it may be desirable to lift the patient using a scoop stretcher and then

Figure 46.27 Traumatic internal hemipelvectomy. Major pelvic trauma such as this can lead to uncontrollable hemorrhage and hypovolemic shock. Early stabilization and immediate transport are mandatory. (From Asensio JA, Trunkey DD, editors: Current therapy of trauma and surgical critical care, St. Louis, 2008, Mosby.)

![Image of a pelvic fracture]
Complications

Application of any pelvic compression device can damage the underlying soft tissue, as well as organs enclosed within the bony pelvis, including the bladder, urethra, and vagina. However, the potential benefits of pelvic stabilization far outweigh these rare potential complications and should not discourage one from applying a pelvic stabilization device.

Conclusion

Pelvic fractures can be quickly and easily stabilized with a sheet or commercially available pelvic circumferential compression device. Early recognition and stabilization of pelvic injuries, gentle manipulation of the patient, and rapid transport will help reduce morbidity and mortality.

REMOVAL OF HELMETS AND PROTECTIVE EQUIPMENT

Background

Though originally developed for protection of the head during combat, helmets are commonly worn by motorcyclists, athletes (e.g., football, hockey, lacrosse, motor sports), and individuals participating in a host of recreational activities (e.g., cycling, skiing, snowboarding, in-line skating, skateboarding). Emergency care providers must therefore be equipped and trained to remove a helmet safely.

The majority of modern sports helmets consist of a hard polycarbonate shell lined with foam padding, adjustable air cells, or both. Density, strength, and rigidity vary depending on the type of impact that the helmet is designed to protect against. Many helmets are considered multisport and meet the requirements for several potential mechanisms of injury and varying degrees of impact.

Many helmets are modified with additional padding so that they conform tightly to the individual’s head. In addition, helmets used in contact sports (e.g., football, hockey, lacrosse) typically have some type of face mask secured to the front of the helmet. Regardless of which method is used, wrap the sheet around the greater trochanters and tie the ends into a knot to compress the bony pelvis (Fig. 46.29). Pelvic binders are designed for quick and easy application in the field or ED. They are easy to maintain, reusable, and sized to fit 95% of the adult population. The pelvic binder acts similar to a bed sheet and, when placed properly over the area of the greater trochanters, provides a safe and effective force to stabilize pelvic fractures. Application of one such device, the SAM Sling (SAM Medical Products), is illustrated in Fig. 46.30.

![Figure 46.28](image1.png) The SAM Sling may be used to stabilize a fractured pelvis. (Courtesy SAM Medical Products, Portland, OR.)

![Figure 46.29](image2.png) Pelvic stabilization using a sheet. Note that the sheet is wrapped firmly around the greater trochanters and tied to compress the bony pelvis.
SAM Sling Application

1. Remove objects from the patient’s pockets or pelvic area. Unfold the SAM Pelvic Sling with the nonprinted side facing up. Keep the strap attached to the buckle.

2. Place the nonprinted side of the SAM Pelvic Sling beneath the patient at the level of the buttocks (greater trochanters).

3. Wrap the nonbuckle side of the SAM Pelvic Sling around the patient.

4. FIRMLY WRAP the buckle side of the sling around the patient and position the buckle in the midline. Secure by pressing the flap to the sling.

5. Lift the black strap away from the sling by pulling upward.

6. With or without assistance, firmly pull the orange and black straps in opposite directions until you hear and feel the buckle "click." MAINTAIN TENSION!

7. IMMEDIATELY press the black strap onto the surface of the SAM Pelvic Sling to secure it. Note: Do not be concerned if you hear a second "click" after the sling is secured.

8. To remove it, lift the black strap by pulling upward. Maintain tension and slowly allow the SAM Pelvic Sling to loosen.

Figure 46.30 Application of the SAM Sling. (Courtesy SAM Medical Products, Portland, OR.)
the helmet to provide additional protection for the athlete’s eyes and face. These athletes also use a variety of additional pads that may complicate or hinder helmet removal, such as shoulder pads and neck orthoses. Shoulder pad configuration varies from sport to sport and even from player to player. Cervical orthoses limit neck motion when paired with helmets and shoulder pads and protect against spinal and brachial plexus injuries. Commonly used cervical orthoses include neck rolls, cowboy collars, and butterfly restrictors.

Motorcycle helmets may or may not have a full face guard, but in either case they have been shown to reduce the incidence of severe head injury and death and are associated with shorter hospital stays and reduced hospital cost. Although early studies suggested that the use of motorcycle helmets might be associated with an increased incidence of cervical spine injuries, this concern has not been substantiated.

Early research in the area of helmet removal focused primarily on the removal of motorcycle helmets. More recently, the spotlight has turned to football helmet removal because football players commonly sustain head and neck trauma, and their care is frequently complicated by the presence of additional protective equipment. The proper management of a helmeted, spine-injured athlete has prompted much debate and disagreement among various health care professionals. In 1998, the National Athletic Trainers’ Association formed the Inter-Association Task Force for the Appropriate Care of the Spine-Injured Athlete, which in 2001 released its first comprehensive set of guidelines and recommendations titled Prehospital Care of the Spine-Injured Athlete. This consensus document was updated in 2009 and recommended transportation with the helmet and shoulder pads in place for the majority of athletes, to be removed at the hospital following spinal imaging. It did allow for removal of helmet and shoulder pads in limited circumstances where immobilization was inadequate or access to airway was compromised. These guidelines were widely adopted but remained controversial because of the difficulties in accessing the airway, neck and chest, the degradation of imaging quality, and the lack of experience of hospital personnel with removing equipment. New guidelines are in development at the time of publication, but the executive summary for the 2015 revision notes allows for removal prior to transport to mitigate these issues.

Helmet removal requires a careful, methodical approach to avoid compounding a possible injury to the spinal cord. Fluoroscopic studies have detected spinal motion even in the best of circumstances when removing hockey and football helmets. As with all trauma victims, the initial management of an injured athlete is to address the ABCs (airway, breathing, circulation). However, it is important to note that a properly fitting sports helmet holds the head securely in the neutral position, provided the athlete is also wearing shoulder pads. In this case, removing the face mask but leaving the helmet and shoulder pads in place may allow adequate airway control while maintaining proper alignment of the cervical spine and reduce the risk for further injury. Generally, removal of helmet and pads as a unit in the field is reasonable provided that several providers are available and have sufficient experience in the technique.

Motorcycle and motor sport helmets do not usually have a removable face mask, do not always fit properly, and are worn without shoulder pads. Thus motorcycle and motor sport helmets should be removed routinely to properly restrict motion of the spine.

**Indications**

In general, in a highly functioning EMS system, there are few downsides to removal of helmet and shoulder pads (if present) at the scene. This equipment needs to be removed eventually, and hospital staff are nearly always less experienced and practiced than trainers and prehospital personnel.

If the athlete’s helmet is not removed, maintain cervical spine immobilization using tape, commercially available foam blocks, and a lifting device designed to maintain spinal motion restriction. Do not remove the helmet or shoulder pads (if present) individually: either remove both or neither, to avoid hyperextension of the cervical spine. If in doubt regarding the need for removal of a sport helmet, the National Athletic Trainers’ Association suggests remembering two overarching principles: (1) exposure and access to vital life functions (e.g., airway, chest for cardiopulmonary resuscitation or use of an automated external defibrillator) must be established or easily achieved in a reasonable and acceptable manner, and (2) neutral alignment of the cervical spine should be maintained while allowing as little motion of the head and neck as possible.

The potential for helmets to interfere with plain films of the spine was demonstrated in two studies, though nonmetallic helmets were less likely to interfere with computed tomography. Thus removal prior to imaging may be indicated depending on the imaging mode available or desired. Motorcycle and motor sport helmets should be removed in the prehospital setting. Motorcycle helmets with a full face guard make it very difficult to assess and manage the airway and to evaluate injuries to the head and neck. The helmet’s large size and design may cause significant neck flexion if left in place. While decreasing the likelihood of injury, the
increasing use of head and neck restraint devices (e.g., the HANS device [Fig. 46.32A] and the SHR Flex device [Fig. 46.32B]) in professional motor sports has further complicated cervical spine management and helmet removal.

### Contraindications

The only absolute contraindications to helmet removal are neck pain or paresthesias associated with the procedure or an inability to remove the helmet because of an impaled object (e.g., deer antler through a motorcycle helmet). Relative contraindications to helmet removal include unfamiliarity with the technique and lack of sufficient assistance.

### Procedure

#### Sport Helmet Removal

When a face mask is present, if not immediately removing the helmet, remove the mask at the earliest opportunity, before transportation and despite the absence of any respiratory complaints. A variety of tools (e.g., FM Extractor [Collins Surgical, Brockton, MA], Trainer’s Angel [Trainer’s Angel, Riverside, CA], anvil pruner, polyvinyl chloride pipe cutter, or power screwdriver) and techniques are used for face mask removal. All emergency care providers should have tools for face mask removal readily available and be familiar with their use. The choice of equipment is less important than the skill and experience of the personnel using it. To remove the face mask of a football helmet, cut the four plastic loop straps that secure the face mask to the helmet. Remove hockey and lacrosse face masks by unscrewing the external screws holding them in place. Maintain in-line stabilization to keep the head and neck in the neutral position during the entire procedure. With practice, the face mask of any helmet can be quickly and safely removed with minimal movement of the cervical spine.

The National Athletic Trainers’ Association protocol for helmet and shoulder pad removal discussed in this chapter has been shown to effectively limit motion of the cervical spine during removal of equipment. Proper removal of a helmet and shoulder pads requires at least two (and preferably three or four) individuals, one of whom maintains manual in-line stabilization of the cervical spine throughout the procedure.

One rescuer manually stabilizes the head and neck in the neutral position by placing his or her hands on each side of the helmet with the thumbs pointing up (Fig. 46.33, step 1). A second rescuer then removes the chin strap by cutting or unsnapping it (see Fig. 46.33, step 2). Next, this rescuer removes the left or right cheek and jaw pads from the helmet by first slipping the flat blade of a screwdriver or bandage scissors between the pad snaps and the helmet’s inner surface and twisting slightly. Once separated from the helmet, remove the pads by sliding them out firmly and slowly. Remove the opposite side in the same manner. Note that some helmet models (e.g., Riddell Evolution [Riddell, Rosemont, IL]) are padded with a number of air-filled bladders that must be deflated (rather than removed) before removal of the helmet (Fig. 46.34). In this case, the second rescuer deflates the air inflation system by releasing the air at the external ports with the inflation needle that comes with the helmet. Alternatively, an 18-gauge needle or air pump needle may be tried. If an inflation needle is not available (or an 18-gauge needle or air pump needle does not work), directly puncture the bladders with an 18-gauge needle.

The second rescuer then takes over in-line immobilization of the head by using one hand to grasp the patient’s mandible between the thumb and the first two fingers while placing the...
Football Helmet and Shoulder Pad Removal

1. One rescuer manually stabilizes the patient’s head and neck in the neutral position by placing his or her hands on each side of the helmet, with the thumbs pointing up.

2. A second rescuer then removes the chin strap by cutting or unsnapping it.

3. The second rescuer then takes over in-line immobilization of the head by using one hand to grasp the patient’s mandible between the thumb and the first two fingers while placing the other hand under the occiput.

4. The first rescuer then places a thumb inside each ear hole of the helmet and curls his or her fingers along the bottom edge of the helmet. Without pulling laterally, the helmet is removed by gently rotating it off the head.

5. The shoulder pads are removed by cutting the straps underneath the arms and the anterior straps holding the pads together.

6. The hands of the second rescuer stabilize the head as the shoulder pads are removed. When possible, the helmet (with the face mask removed) and shoulder pads should remain in place until the initial clinical and radiographic evaluation is completed in the ED.

Figure 46.33 Football helmet and shoulder pad removal. Proper removal of a helmet and shoulder pads requires at least two rescuers. (Note: this figure does not demonstrate removal of the face mask and cheek/jaw pads, which is recommended before helmet removal. See text for details.)

ED, Emergency department. (Courtesy AtlantiCare Regional Medical Center, Emergency Medical Services, Atlantic City, NJ.)

other hand under the occiput (see Fig. 46.33, step 3). The first rescuer then places a thumb inside each ear hole of the helmet and curls his or her fingers along the bottom edge of the helmet (see Fig. 46.33, step 4). At this point some experts recommend easing the helmet off by pulling laterally and longitudinally in line with the head and neck. However, this maneuver may actually tighten the helmet at the occiput and the forehead. The Inter-Association Task Force recommends rotating the helmet off the head in a gentle fashion without pulling laterally. Remove shoulder pads by cutting the straps underneath the arms and the anterior straps holding the pads together (see Fig. 46.33, step 5). If a cervical orthosis (e.g., neck roll, cowboy collar, or butterfly restrictor) is present, disconnect it from the helmet and shoulder pads before removal.
Figure 46.34 Riddell Revolution football helmet. These helmets are padded with a number of air-filled bladders that must be deflated (rather than removed) before removal of the helmet. A, Frontal view. B, Inside view. See text for details. (Courtesy Riddell, Elyria, OH.)

Motorcycle Helmet Removal

1. One rescuer maintains in-line immobilization by placing his or her hands on each side, of the helmet with the fingers on the victim’s mandible. This position prevents slippage if the strap is loose.

2. A second rescuer cuts or loosens the strap at the D-ring.

3. The second rescuer places one hand on the mandible at the angle, the thumb on one side and the long and index fingers on the other. With the other hand the rescuer applies pressure from the occipital region. This maneuver transfers the in-line immobilization responsibility to the second rescuer.

4. The rescuer at the top moves the helmet. Three factors should be kept in mind:
   - The helmet is egg-shaped and therefore must be expanded laterally to clear the ears.
   - If the helmet provides full facial coverage, glasses must be removed first.
   - If the helmet provides full facial coverage, the nose may impede removal. To clear the nose, the helmet must be tilted backward and raised over it.

5. Throughout the removal process the second rescuer maintains in-line immobilization from below to prevent unnecessary neck motion.

6. After the helmet has been removed, the rescuer at the top replaces his or her hands on either side of the victim’s head with the palms over the ears.

Figure 46.35 Motorcycle helmet removal. (Reproduced with permission from Norman E. McSwain, Jr., MD, FACS, and Richard L. Garrnelli, MD, FACS—American College of Surgeons Committee on Trauma, April 1997.)
When possible, remove the shoulder pads and helmet simultaneously to prevent the head from falling into extension. If the shoulder pads cannot be removed simultaneously, manually stabilize the head in the neutral position during the procedure. The hands of the second rescuer can be moved superiorly as the helmet is being removed so that the thumb and first fingers grasp the maxilla at each side of the nose in the maxillary notch (see Fig. 46.33, step 6).

After removal of the helmet, place a c-collar while maintaining in-line stabilization. Remember that if the helmet cannot be removed and access to the chest area is required, remove the anterior half of the shoulder pads and leave the posterior portion to maintain cervical position with the helmet in place.

Motorcycle Helmet Removal
For the reasons described earlier, motorcycle helmets should be removed in the prehospital setting. The removal method endorsed by the American College of Surgeons in 1997 is the method most often used today (Fig. 46.35)." Providers should be aware that significant cervical motion can occur during removal of a motorcycle helmet if the shoulders are not properly elevated; a folded sheet or jacket placed behind the patient’s shoulders may help limit any cervical motion associated with the procedure.130

If attempts to remove the helmet result in pain or paresthesias, the advanced trauma life support guidelines recommend removing the helmet via a technique described by Aprahamian and coworkers.111 This technique uses a cast saw to bivalve the helmet in the coronal plane. Following division of the outer rigid shell, incise the inner foam material and remove it while maintaining the head and neck in neutral alignment. Although this approach does provide an alternative method of removing the helmet, the intense vibrations produced during use of the cast cutter may exacerbate an underlying spinal injury. In addition, this technique may be slow and difficult with modern, well-fitting, high-quality helmets.121,132,133

The preferred method of helmet removal in organized motor sports is the Eject Helmet Removal System (formerly the Hats Off System), which consists of an inflatable air bladder that can be placed into the helmet before use or introduced into the crown of the helmet with a specially designed insertion tool. When inflated with a hand pump or CO2 cartridge, it purportedly loosens the helmet to allow easy removal (Fig. 46.36).132 Despite little scientific data supporting its use, the IndyCar Racing League, American Speed Association, and American Motorcycle Association have made the Eject Helmet Removal System mandatory at all of their events. In addition, the Championship Auto Racing Teams and National Association for Stock Car Auto Racing strongly recommend its use.

Complications
It is theorized that underlying cervical spine injuries may be exacerbated by failure to adhere to proper helmet removal techniques, though clinical data is lacking.111 In a cadaveric model, Donaldson and colleagues133 demonstrated that even when using proper technique, there is a significant amount of motion during removal of helmets and shoulder pads. Larger studies are needed to determine whether there is a real and significant risk in the clinical setting.

Conclusion
Prehospital care providers must take caution when evaluating and treating an athlete with a suspected head and neck injury. In general, athletic helmets and shoulder pads should be removed at the scene when adequate trained personnel are available, though it is reasonable to transport without removal when equipment is maintaining the spine in neutral alignment, and when there are no other emergency conditions that require their removal. Face masks, however, should be removed early in the management of these patients. Motorcycle and motor sport helmets, as well as sports and other helmets worn without shoulder pads, should be removed routinely to provide access to the patient’s airway and allow proper spinal motion restriction. When indicated, prehospital helmet removal can be accomplished in a safe and effective manner by well-trained prehospital care providers.

The primary concern of removing helmets and protective padding is causing additional injury to the patient. It may also prolong on-scene time and delay transport to the hospital.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Rapid and appropriate emergency care of a patient with an amputated body part is crucial to the salvage and preservation of function. This chapter discusses the emergency management priorities of patients with amputation injuries, the acute care of amputated parts before they are replanted, and the management of distal digit amputations and dermal “slice” wounds.

Amputation may be partial or complete. Injuries with any interconnecting tissue between the distal and proximal portions, even if it is only a small piece of bridging skin, are considered incomplete (or partial) amputations. The peak incidence of traumatic amputations occurs between the ages of 15 and 40 years and approximately 80% of injuries occur in males. Motor vehicle collisions are the leading mechanism of injury, followed by industrial and agricultural accidents. Local crush injuries occur most commonly, and sharp guillotine amputations are least common. Partial amputations occur as often as total amputations. Seventy percent of amputations overall occur in the upper extremity and distal amputations are more common than proximal.

Successful revascularization of amputated parts may ensure viability, but neurologic, osseous, and tendinous healing are critical for ultimate function. If there is incomplete neurologic recovery, limited range of motion, and intolerance to cold, the replanted part may have little functional value for the patient. Rehabilitation from replantation surgery may be prolonged, (often >1 year) and frequently repeat surgical procedures are required. A patient’s preconceived perceptions of medical ability to save an amputated part and well-intentioned promises made by a transferring physician may set up unrealistic expectations for replantation. The emergency clinician should be aware of the limitations of replantation surgery and should not encourage unrealistic expectations in injured patients or their families.

**BACKGROUND**

Replantation attempts (with rare success) have been described for hundreds of years but the ability to consistently replant amputated parts is now possible with advances in microvascular surgical techniques. Upper limb and hand replants were first described in 1962. The first successful microvascular anastomosis of a digital vessel was described in 1965, and now replantation success rates range from 50% to 90%. Survival of the replanted tissue was the criterion for success for the original pioneers in replant surgery, but with further technological and surgical refinements, today’s surgeons emphasize functional recovery as well as viability. Replantation of a part that is painful or useless or that interferes with function is a disservice to the patient and is less desirable than early restoration of function.

### Management of Amputations

**Indications**
- Young stable patient
- Thumb
- Multiple digits injured
- Sharp wounds with little associated damage
- Upper extremity (children)

**Contraindications**

**Absolute**
- Associated life threats
- Severe crush injuries
- Inability to withstand prolonged surgery

**Relative**
- Single digit, unless thumb
- Avulsion injury
- Prolonged warm ischemia (>12 hr)
- Gross contamination
- Prior injury or surgery to part
- Emotionally unstable patient
- Lower extremity

**Complications**
- Infection
- Poor function

**Equipment**
- Saline
- Gauze
- Kerlix
- Styrofoam cooler with ice
- Plastic bag

Review Box 47.1 Management of amputations: indications for reimplantation, contraindications, complications, and equipment. *If the victim is a child or if there are multiple losses, salvage reimplantations are attempted and the relative contraindications are ignored.*
without replantation. With advancements in technology, there are also alternatives to replantation such as targeted reinnervation, transplantation of composite tissue, or the potential for upper extremity transplantation to restore upper extremity function. The operative plan must be tailored to each individual patient with consideration of the patient’s underlying health, ability to undergo and comply with rehabilitation, support network, and likely functional outcome.

**INDICATIONS**

The mechanism of injury is the most important determinant of implant survival. Replantation is more likely to occur with sharp/penetrating injury, distal level of amputation, lack of multilevel involvement, and when the injury is isolated to the extremity. Preservation of the amputated part is indicated whenever there is a potential for replantation. Revascularization and reanastomosis of partially and completely amputated parts should be provided when there is hope of preservation or restoration of function. Aesthetic considerations, patient avocations, and occasionally the patient’s religious or social customs may also influence the decision to proceed with surgery. Ultimately, the microsurgical team and patient must reach the decision together after a rational explanation of the potential results.

Replantation is more commonly performed for upper extremity amputation because a lower extremity prosthesis often provides a good functional outcome and is often superior to outcomes with replantation. Upper extremity prostheses are less able to achieve fine motor activity and a “bad hand” may be more functional than a “good amputation” in the upper extremity. Additionally, there is less muscle (which is less vulnerable to crush injury and allows a longer ischemia time) and increased collateral circulation in the upper extremity, which improves replantation success. Review Box 47.1 summarizes the general indications for replantation. Single-digit amputations that are both proximal to the distal interphalangeal (DIP) joint and distal to the flexor digitorum superficialis may be replanted successfully with good functional recovery (Fig. 47.1). The thumb accounts for 40% to 50% of hand function with its utility in pinch and grip. Even if a replanted thumb has a limited range of motion, it is more functional than any other available reconstructive procedure or prosthetic device. The underlying health of the patient is more important to outcomes than chronologic age. Children consistently have better outcomes due to their superior neuroplasticity, regenerative capacity, and adaptability to rehabilitation, though there are no fixed age limits for replantation. The decision to replant is made on a case-by-case basis by the microsurgical team, who must weigh all the factors involved. In general, patients can expect to achieve 50% of original sensation and motor function of the replanted part, with younger patients and more distal amputations having the best outcomes.

**CONTRAINDICATIONS**

Absolute contraindications to replantation include life-threatening injury or significant comorbidity that prohibits a lengthy operation. Relative contraindications include severely crushed or mangled parts or profuse contamination. Contraindications to replantation are discussed in the following sections. The amputated part should be treated carefully because portions may be used as a graft even if the entire part will not be replanted. For example, an amputated fingertip not suitable for replantation may be an ideal donor source for a skin graft on the stump. In addition, even when replantation is contraindicated, tissue (e.g., skin, bone, tendon) from the amputated part may be useful in restoring function to other damaged parts. In general, never discard amputated tissue until all possible uses of the severed parts are considered.

**Figure 47.1** A and B, Single-digit amputations are not usually replanted unless the thumb is involved. This sharp amputation with little associated damage in a 20-year-old may qualify, but it is best to discuss this with the hand surgery team before giving unrealistic expectations to the patient, who may anticipate a normal result by simply “sewing it back on.” Avulsion of the tendons proximal to the amputation greatly complicates this case.

**GENERAL CONSIDERATIONS**

**Mechanism of Injury**

Severe extremity trauma is a significant cause of morbidity, and the potential for successful replantation in terms of survival, as well as useful function, is directly related to the mechanism of injury. Guillotine-type injuries are the least common but have the best prognosis because of the limited area of destruction. Crush injuries are the most common, but produce more tissue injury and therefore have a poorer prognosis. Avulsion injuries have the worst prognosis because a significant amount of vascular, nerve, tendon, and soft tissue injury invariably occurs.
Ischemia Time
Irreversible ischemic injury occurs in muscle tissue after 2 to 4 hours of warm ischemia and 6 to 8 hours of cold ischemia. Digits have less muscle tissue and are therefore less susceptible to ischemic damage (increasing warm ischemia time to 6 to 12 hours and cold ischemia time to 12 to 24 hours). However, ischemic time alone does not preclude replantation attempt as successful replantation after prolonged ischemia time has been reported (hand reimplantation after 54 hours of cold ischemia; digit reimplantation after 33 hours of warm and 94 hours of cold ischemia). Techniques such as interosseous muscle stripping to prevent necrosis may improve replantation success in scenarios of prolonged ischemia time.

PATIENT ASSESSMENT AND MANAGEMENT
The initial care and treatment of a patient with a traumatic amputation begins with a primary survey and stabilization of the airway, breathing, and circulation. Traumatic limb amputations are less likely to occur in isolation and management of a patient’s injuries must be prioritized such that life-threatening head and torso injuries take precedence over limb-threatening injuries; some even become suicidal. Treat patients with supportive and realistic reassurance, and avoid promises of outcome. It is important that the emergency clinician (or other nonreplantation specialist) not speculate on the specifics of the ultimate prognosis.

CARE OF THE STUMP AND AMPUTATED PART
The goals of initial care of the stump and amputated part include control of hemorrhage and prevention of further injury or contamination.

Stump
Examination of the stump may be brief and should primarily be an assessment of the degree of damage to surrounding tissue (Fig. 47.4). Remove all jewelry from the stump and irrigate with normal saline to remove gross contamination; any manual débridement and dissection should be performed only by the replantation team. Do not use local antiseptics, especially hydrogen peroxide or alcohol, because they may damage viable tissue. Do not clamp arterial bleeders. Similarly, surrounding tissue should not be manipulated, clamped, tagged, or further traumatized in any way. Cover the stump wound with a saline-moistened sterile dressing to prevent further contamination and to limit damage from desiccation. Splint immobilize associated fractures and evaluate the patient for associated conditions such as rhabdomyolysis and compartment syndrome.

Obtain a history regarding the exact mechanism, time, and duration of the injury, handedness, general medical status (notably diabetes, vascular disease, and smoking), allergies, medications, illness, previous injury to the affected part, care of the stump and amputated part before arrival in the emergency department (ED), occupation, avocations, tetanus history, and the patient’s expectations.

Initiate tetanus prophylaxis (if needed) and broad-spectrum systemic antibiotic therapy (e.g., cephalosporins). Intravenous opioids are recommended for pain; titrate the dose to the clinical condition. Digital or regional nerve blocks are effective adjuncts for pain relief but limit ongoing neurologic evaluation (see Chapters 31 and 32). Obtain preoperative labs and imaging, including radiographs of the amputated part and proximal stump that include at least one joint proximal to the injury site (Fig. 47.3). Contact the regional replantation resource center as soon as possible to arrange transportation and provide adequate time for mobilization of the replantation team.

Patients who have suffered an amputation often experience denial, shock, disbelief, and feelings of hopelessness about their injury; some even become suicidal. Treat patients with supportive and realistic reassurance, and avoid promises of outcome. It is important that the emergency clinician (or other nonreplantation specialist) not speculate on the specifics of the ultimate prognosis.
Figure 47.2 Management algorithm for patients with mangled extremities. CTA, Computed tomography angiography. (Used with permission from Scalea TM, DuBose J, Moore EE, et al: Western Trauma Association critical decisions in trauma: management of the mangled extremity, J Trauma Acute Care Surg 72(1):86–93, 2012.)
the stump for protection and to prevent further injury from concomitant fractures or compromised blood flow from a change in position. Splinting and elevation may also reduce swelling and help control bleeding.

**Amputated Part**

Remove all jewelry and irrigate the amputated part with normal saline to remove gross contamination, similar to care of the stump described previously (Fig. 47.3). The amputated part should be wrapped in saline-moistened gauze and placed in a dry and closed waterproof plastic bag (see Fig. 47.5, step 5). Do not place the amputated part directly in ice or cold saline. The bag containing the specimen wrapped in saline-moistened gauze should be immersed in ice water or refrigerated at 4°C. The amputated part itself should NOT come into direct contact with ice or saline or reach freezing temperatures. Do not discard any tissue as any amputated fragment could be potentially used for skin, bone, or nerve grafting. Label the tissue containers with the patient’s name, the amputated part contained within, the time of the original injury, and the time that cooling began.

Treatment is similar for partial amputations with vascular compromise. Irrigate the wound with normal saline. Place a saline-moistened sponge on the open tissue and wrap the injury in a sterile dressing in which a splint has been incorporated to protect from further injury. Place ice packs or commercial cold packs over the dressing to cool the devascularized area (see Fig. 47.5, step 8).

**SPECIAL CONSIDERATIONS**

**Lower Extremity Amputations**

Lower limbs are not commonly replanted because replantation is fraught with potential complications and lower limb prostheses, especially those used below the knee, are well tolerated and functional. Prostheses provide a secure stance and permit locomotion. Lower extremity replantation generally requires skeletal shortening, and distal nerve regeneration is often imperfect, with both deficits producing dysfunction. A patient with a replanted lower extremity, with significant shortening and without sensation, typically has superior functional outcome with a prosthesis. Children are an exception and replantation may be attempted in a child with a lower limb amputation due to their superior replantation functional outcomes.

**Hand Function**

Fine motor hand function involves pinch and grasp functions. If the index finger is amputated, the middle finger can often adequately perform the pinching function formerly provided by the index finger. Power in grasping and gripping is primarily a function of the fourth and fifth digits. An effective grip that provides the ability to hold a variety of objects is a central function of the ring and middle fingers. In addition to its function in pinching, the thumb is the major opposing force for successful grip and grasp. The thumb is the most important digit for adequate hand function, and its loss results in 40% to 50% disability. Such disability requires aggressive attempts to replant amputated thumbs. If this is impossible or unsuccessful, secondary alternatives are pollicization (creation of a thumb by using another digit) or toe transfers.
Care of the stump and amputated part

1. Irrigate the wound with saline solution.
2. Wrap the wound with Kerlix or Kling for pressure, then elevate.
3. Rinse the amputated part with saline solution.
4. Wrap the amputated part in a moist sterile gauze or towel, and place it in a plastic bag or container. Do not immerse the amputated part in saline.
5. Place the amputated part in a container (preferably Styrofoam) and cool with separate plastic bags containing ice or in a container of ice water.
6. For partial amputations, begin by irrigating the wound with saline solution.
7. Place parts in a functional position, then apply a saline-moistened sterile dressing over the wound.
8. Apply a coolant bag outside of the dressing.
9. Splint and elevate the extremity.

NOTE: Do not scrub or apply antiseptic solution to the wound.
Control any bleeding with pressure.
If a tourniquet is required, place it close to the amputation site.

Figure 47.5 Care of the stump and amputated part. Do not place the amputated part directly in ice or saline.
CHAPTER 47  Management of Amputations  953

significant soft tissue loss on the volar aspect of the distal finger, completion amputation and closure by a hand specialist provides the quickest recovery. In these cases, ablation of the nail bed and digital nerve transection are generally required to prevent hook nail deformity and painful neuroma formation, respectively.22 Alternatives to completion amputation include partial-thickness skin grafts, full-thickness skin grafts, Atasoy-Kleinert V-Y flap,

Fingertip Amputations and Dermal Slice Wounds

There are several classification systems for distal fingertip amputations. The Tamai classification is most commonly used, though the Allen classification is the most clinically useful for hand surgeons (Fig. 47.6).

Treatment of distal fingertip injuries remains somewhat controversial, but good results are often achieved with conservative management. Factors that impact management include the amount of soft tissue loss, the integrity of the nail bed, and the patient's age, occupation, and management preference. The basic goals of treatment are to provide tissue coverage, an acceptable cosmetic result, and early functional recovery. Primary closure or healing by secondary intention is preferred when there is adequate soft tissue coverage and no exposed bone (Fig. 47.7). If there is no exposed bone, but significant pulp loss, treatment options include primary closure, healing by secondary intention, completion amputation, full-thickness skin grafting, or split-thickness skin grafting.22

In general, uncomplicated fingertip wounds less than 1 cm in size heal well with conservative treatment. In children under 2 years of age, fingertip wounds up to 2 cm may be managed conservatively, as children in this age group have superior regenerative capacity.7-10 Management is more challenging when there is significant loss of skin and soft tissue from the finger pad.

Volar skin is unique in its combination of toughness and sensitivity. Wounds with a significant loss of volar tissue frequently require additional treatment (Fig. 47.8). If there is

Figure 47.6 The Tamai and Allen classifications of distal fingertip amputations. DIPJ, distal interphalangeal joint. (From Lee DH, Mignemi ME, Crosby SN: Fingertip injuries: an update on management, J Am Acad Orthop Surg 21(12):756–766, 2013.)

Figure 47.7 This very minimal amputation of the fat pad with a mandolin slicer does not involve bone or the nail bed and is a common injury. It will heal very well with conservative therapy. No skin grafting is required. It will require a few weeks to heal, with periodic cleaning and débridement. Note: The ring should be removed as soon as possible.

Figure 47.8
S E C T I O N  V I I I  M U S C U L O S K E L E T A L  P R O C E D U R E S

Figure 47.8 This patient with significant volar tissue loss required microsurgical replantation. A, Tamiel level 2 volar oblique pulp amputation. B, After completion of microsurgical replantation. C, Four-month follow-up. (Used with permission from Peterson SL, Peterson EL, Wheatley MJ: Management of fingertip amputations, J Hand Surg Am 39(10):2093–2101, 2014.)

Kutler lateral V-Y flap, Moberg flap, or other local and distal flap coverage techniques.22–23 A composite graft is another reconstructive option in children, where the amputated part is sutured back to the stump without formal microvascular repair. The tip may potentially necrose, but serves as a biological dressing. In adults, composite graft may be attempted after the amputated portion is defatted but outcomes are inferior to those in children.26,27 Although traditionally some type of grafting or advancement techniques have been used for distal guillotine amputations, conservative management is now more common, even if some portion of bone is exposed (Fig. 47.9).

For more significant distal amputations, replanting the amputated tip preserves the nail (which enhances fingertip sensation and two-point discrimination), digit length, and joint motion, and often results in better overall functional outcomes.28 Moreover, survival rates of finger replantation are high (approaching 90%).6,26,27 Even amputations distal to the insertion of the flexor digitorum superficialis are often attempted because digit function with preserved proximal interphalangeal joint motion is enhanced by the additional length, even if the DIP joint is stiff or fused.6,26,32 Additionally, active range of motion is increased if this insertion site is preserved (82 degrees compared to 25 degrees with a more proximal amputation).6,26,32 Young children potentially regenerate the distal tip, and functional outcomes are also excellent after replantation.31

As discussed previously, incomplete transections and small distal amputations without significant soft tissue loss may heal well with conservative therapy initiated by the emergency clinician. It may require a number of weeks to totally heal, but nonoperative treatment in appropriate patients provides excellent functional and cosmetic results, minimizes recovery time, and has few complications. Débride necrotic and grossly contaminated tissue and irrigate the wound thoroughly with normal saline. If bone is left exposed without soft tissue coverage, the patient will probably need an operative procedure. Alternatively, the bone may be rongeured (shortened) to allow soft tissue coverage and primary healing with better functional recovery. The nail bed tissues should be preserved, if possible, because the presence of a nail improves hand sensation and affects the cosmetic appearance. After cleansing and careful débridement, apply an occlusive dressing directly over the wound and then bandage and splint the finger for protection. Provide tetanus prophylaxis if needed. Amputations that involve the distal phalanx are frequently treated as contaminated open fractures. However, there is no evidence that antibiotics prevent infection or are superior to excellent wound care, and prophylactic antibiotics are generally not recommended.14–16 Wound contraction and healing usually result in acceptable cosmetic and functional recovery in 2 to 3 weeks. Arrange appropriate follow-up to ensure adequate healing and recovery.

Partial Fingertip Amputations

The emergency clinician can also manage partial fingertip amputations distal to the DIP joint. These wounds are treated in a manner similar to complete amputations. However, when the amputation has a substantial amount of undamaged tissue connecting the fingertip, use sutures or bandaging and protective splinting to carefully align and stabilize the injured fingertip. Partially amputated fingertips, especially in children, may occasionally survive and regain vascularization and sensation. The patient must be warned that the distal tip may potentially become necrotic and progress to a complete amputation. Injury to the nail bed requires special attention to ensure proper alignment. If the nail bed tissues are not aligned properly, permanently disfigured nails may result. Complete or partial nail removal may be required for the placement of sutures.

Manage dermal slice wounds (see Fig. 47.7) by gently cleansing the wound and applying antibiotic ointment and a nonadherent dressing, followed by a pressure dressing. The patient should return in 48 to 72 hours for the wound to be inspected and the dressing changed. At that time, instruct the patient on the use of nonadherent dressings that should be changed daily until functional epithelialization of the wound takes place (usually approximately 10 to 14 days). A protective finger splint or guard also minimizes the risk for further injury and pain from trauma to the sensitive wound area. Protection also allows earlier return to functional use and employment. Wounds larger than 10 mm² and those with deep loss of digit pulp tissue may be candidates for skin grafting.

Penis, Ear, and Nose Amputations

Replantation of the penis, ear, and nose generally results in better function and cosmesis than a prosthesis or reconstructive surgery. The amputated parts and wounds should be handled in the same manner as digital replantations.

Penile amputations are uncommon; most cases are self-inflicted trauma in patients who are severely psychologically disturbed. Successful replantation with microsurgical techniques has been reported. Preservation or reconstruction of the urethra to maintain a competent urinary stream is critical for success.37,38 Ears and noses are frequently partially amputated and
complications include cold intolerance, paresthesias, tendon adhesions, bone malunion/nonunion, neuroma formation, and contractures. Despite advanced microsurgical techniques, patients typically recover only 50% of their original function and sensation of the replanted parts, which may have negative psychosocial and economic consequences.

FIELD AMPUTATIONS

Emergency physicians may be called on to perform field amputation of an extremity as a lifesaving intervention. The typical scenario is a victim who is trapped or unable to be extricated from the scene because an extremity is covered or held immobile due to debris or other heavy objects or is lodged in a narrow space. The procedure is not detailed in this text but is contained in the video library (Videos 47.1-47.3).

Acknowledgment

The contributions by Dean Moore II, MD, to the previous edition are appreciated.
REFERENCES


Chapter 48

Extensor and Flexor Tendon Injuries in the Hand, Wrist, and Foot

Peter E. Sokolove and David K. Barnes

Extensor Tendons

Extensor tendons are quite superficial, covered only by skin and a thin layer of fascia, and are thus highly susceptible to injury by commonly experienced trauma. Such injuries may result from lacerations, bites, or burns, but they may also be caused by closed injury or with even seemingly superficial lacerations. Whereas some extensor tendon injuries must be managed by a hand surgeon, others may be treated in the emergency department (ED). The emergency provider must understand the anatomy, principles of treatment, repair technique, and postrepair care of these injuries to ensure the best possible patient outcome. The authors suggest that all flexor tendon lacerations be referred to a hand surgeon.

Functional Anatomy

There are 12 extrinsic extensor muscles of the wrist and digits, all of which are innervated by the radial nerve. The muscles that give rise to these tendons originate in the forearm and elbow (Fig. 48.1). The extrinsic extensor tendons reach the hand and digits by passing through a fibro-osseous tendon sheath (retinaculum) located at the dorsal surface of the wrist. This synovium-lined sheath provides smooth gliding of the tendons and prevents bowstringing when the wrist is extended. The dorsal retinaculum contains six compartments or subdivisions (Fig. 48.2). These compartments are numbered from the radial to the ulnar side of the wrist.

The first compartment contains two tendons, the abductor pollicis longus (APL) and the extensor pollicis brevis (EPB). The APL tendon is the most radial of the extensor tendons and inserts on the base of the thumb metacarpal. It can be palpated just distal to the radial tubercle. The APL tendon causes thumb abduction and extension and some radial wrist deviation. The EPB travels with the APL through the first compartment but inserts at the base of the proximal phalanx of the thumb. The EPB tendon can be palpated over the dorsum of the first metacarpal when the thumb is extended against resistance. Both tendons can be tested by having the patient spread the fingers apart against resistance. The second compartment also contains two tendons: the extensor carpi radialis brevis (ECRB) and the extensor carpi radialis longus (ECRL). These two tendons arise from the lateral epicondyle of the elbow. The ECRL inserts on the base of the index finger metacarpal, and the ECRB inserts on the base of the long (middle) finger metacarpal. Both tendons are powerful wrist extensors, and the ECRL also allows some radial wrist deviation. Wrist extension plays an especially important role in the mechanics of the hand because hand-grip strength is maximal only when the wrist is extended.

The third compartment contains only one extensor tendon, the extensor pollicis longus (EPL). This tendon crosses over the ECRB and ECRL and travels along the dorsum of the thumb to insert on the distal phalanx. The EPL forms the top of the anatomic “snuffbox,” and the bottom is formed by the EPB. The EPL can be visualized when the thumb is extended, and its strength can be tested by having the patient hyperextend at the interphalangeal (IP) joint against resistance. The intrinsic extensor of the thumb can provide some degree of extension at the IP joint. Therefore, if an EPL injury is suspected, it is important to compare extension at the IP joint with that of the unaffected thumb.

The fourth and fifth compartments contain the six tendons that extend the index through the little fingers. Each finger has its own extensor digitorum communis (EDC) tendon. The index and little fingers have an additional independent extensor tendon, the extensor indicis proprius (EIP) for the index finger and the extensor digiti minimi (EDM) for the little finger. The fourth compartment contains the EIP and EDC tendons, and the fifth compartment contains only the EDM tendon. These six tendons can be seen over the dorsum of the hand, where they are poorly protected and prone to injury. In this region the tendinous, ligamentous, and fascial connections between these tendons are known as the juncturae tendinum. Because of these redundant interconnections, a patient may be able to extend a digit, albeit weakly, even when there is a complete laceration of its EDC tendon. To avoid missing a tendon injury on the dorsum of the hand, it is important that the examiner test for tendon strength and not just for active extension.

The course of the extensor tendons along the fingers is more complex, but a basic understanding of this anatomy is essential for the emergency provider to evaluate and treat extensor tendon injuries (Fig. 48.3). The EIP tendon joins the EDC tendon at the level of the metacarpophalangeal (MCP) joint in the index finger; the EDM tendon parallels the course of the EDC tendon. The four EDC tendons eventually insert at the base of the proximal, middle, and distal phalanges. The most proximal insertion of the EDC tendon is at the level of the base of the proximal phalanx. The tendon actually inserts in two ways. First, there is a loose dorsal insertion just distal to the MCP joint. In addition, the EDC tendon inserts into the volar plate via the sagittal band. The sagittal band is a circumferential ligament at the level of the metacarpal head that serves to keep the EDC tendon centered over the metacarpal head, as well as to provide a stable connection with the volar plate located on the palmar side of the hand. After its primary insertion at the level of the MCP joint, the EDC tendon then extends dorsally along the digit. The EDC trifurcates over the proximal phalanx (Fig. 48.4). Its major central slip inserts on the base of the middle phalanx (Fig. 48.5). The lateral branches of the EDC tendon join with the lateral bands from the interossei and lumbricals to form the conjoined lateral bands. The two conjoined lateral bands then fuse together over the middle phalanx to form the terminal extensor mechanism (TEM) that inserts on the base of the distal phalanx (Fig. 48.6). The triangular ligament is a connection between the two conjoined lateral bands that assists in keeping these structures on the dorsal aspect of the digit.

The sixth dorsal compartment of the wrist contains only one tendon, the extensor carpi ulnaris (ECU). This tendon...
CHAPTER 48  Extensor and Flexor Tendon Injuries in the Hand, Wrist, and Foot

**Extensor Carpi Ulnaris – Compartment 6**

**Extensor Digiti Minimi – Compartment 5**

**Extensor Digitorum**

**Extensor Indicus**

**Extensor Pollicis Longus – Compartment 3**

**Extensor Carpi Radialis Brevis**

**Extensor Carpi Radialis Longus**

**Abductor Pollicis Longus**

**Extensor Pollicis Brevis**

**Radial Artery in Anatomical Snuffbox**

**Dorsal Interosseous Muscles**

**Compartment 4**

**Compartment 5**

**Compartment 6**

**Ulna**

**Radius**

**Posterior (Dorsal) View**

**Cross-Section of Most Distal Portion of Forearm**

**Posterior View**

**Figure 48.1** Extensor muscles and tendons of the right wrist and hand. A, Posterior (dorsal) view. B, Cross-section of the most distal portion of the forearm. C, Radial nerve in the forearm: posterior view. (A–C, Netter illustrations used with permission of Elsevier Inc. All rights reserved.)
Injuries to extensor tendons from lacerations are quite common, especially on the dorsum of the hand where they are located superficially. All dorsal wrist, hand, and digit lacerations should be assumed to have an underlying tendon laceration until proved otherwise. Digital extension, albeit weak, can still occur with partial tendon lacerations of up to 90%, so visualization of the tendon and careful strength testing are required to definitively rule out a partial injury. In some cases the specific diagnosis simply cannot be made on the first examination (see later). Complete laceration of an EDC tendon on the dorsum of a hand can still allow digital extension through the juncturae tendinum.

After assessing the strength and neurovascular status of the injured hand it is imperative that the emergency provider visually inspect the wound thoroughly. Inspection should include an assessment of the degree of wound contamination, as well as a search for foreign bodies and occult tendon lacerations. It is often necessary to extend the skin laceration to aid in the visualization of a possible tendon injury. Because an extensor tendon is a mobile structure, it is imperative to visualize it in its entirety through a full range of motion if it is exposed. It is especially important to examine the tendon in the position it originates at the lateral epicondyle of the elbow and inserts at the base of the small finger metacarpal. The ECU functions as a wrist extensor and ulnar deviator. It can be palpated just distal to the tip of the ulna, and its strength can be tested by forced ulnar deviation of the wrist.

**General Approach to Extensor Tendon Injuries**

The key to detecting extensor tendon injuries in the ED is to perform a careful and thorough history and physical examination. These injuries are easy to miss with a cursory examination. Closed injuries may appear innocuous at first but can result in tendon injuries that may lead to severe deformities or dysfunction if undetected (Figs. 48.7–48.9). Closed injuries are also commonly associated with fractures. A hand radiograph is recommended for closed-hand injuries when a fracture is suspected or for open-hand injuries in which a fracture or foreign body is suspected. It is generally accepted that all open injuries that result from glass should be radiographed. Plain radiographs have a sensitivity of approximately 98% for detecting radiopaque foreign bodies (e.g., gravel, glass, metal). Injuries to extensor tendons from lacerations are quite common, especially on the dorsum of the hand where they are located superficially. All dorsal wrist, hand, and digit lacerations should be assumed to have an underlying tendon laceration until proved otherwise. Digital extension, albeit weak, can still occur with partial tendon lacerations of up to 90%, so visualization of the tendon and careful strength testing are required to definitively rule out a partial injury. In some cases the specific diagnosis simply cannot be made on the first examination (see later). Complete laceration of an EDC tendon on the dorsum of a hand can still allow digital extension through the juncturae tendinum.

After assessing the strength and neurovascular status of the injured hand it is imperative that the emergency provider visually inspect the wound thoroughly. Inspection should include an assessment of the degree of wound contamination, as well as a search for foreign bodies and occult tendon lacerations. It is often necessary to extend the skin laceration to aid in the visualization of a possible tendon injury. Because an extensor tendon is a mobile structure, it is imperative to visualize it in its entirety through a full range of motion if it is exposed. It is especially important to examine the tendon in the position

---

**Figure 48.2** A, The extensor mechanism at the wrist and dorsum of the right hand. The six extensor compartments at the wrist contain (1) the abductor pollicis longus (APL) and extensor pollicis brevis (EPB), (2) the extensor carpi radialis longus (ECRL) and extensor carpi radialis brevis (ECRB), (3) the extensor pollicis longus (EPL), (4) the extensor digitorum communis (EDC) II to V and the extensor indicis proprius (EIP), (5) the extensor digiti quinti (EDQ), and (6) the extensor carpi ulnaris (ECU). An important anatomic detail is the presence of a synovial sheath around each tendon unit within each fibro-osseous canal. Note that the EDQ is also called the extensor digiti minimi by some authors. B, Note that the juncturae tendinum allow some weak extension of the finger when the proximal extensor is completely lacerated. (A, Adapted from Thomas JS, Peimer CA: Extensor tendon injuries: acute repair and late reconstruction. In Chapman MW, editor: Operative orthopaedics, ed 3, Philadelphia, 2001, JB Lippincott, p 1487.)
In a study of 34 patients with upper extremity tendon injuries treated by emergency physicians, ultrasound was faster and more accurate than exploration, magnetic resonance imaging (MRI), and specialty consultation. Accuracy of sonography for detection of hand and digit tendon injuries greatly depends on operator experience and is not a routine component of emergency medicine ultrasound practice.

Definitive examination of any wound must occur under the best possible conditions, with a good light source, a bloodless

that it occupied at the time of injury because the tendon injury frequently does not lie directly under the external skin wound (see Fig. 48.8).

Some investigators have advocated the use of ultrasound in the diagnosis of suspected extensor (and flexor) tendon lacerations in the hand. This is a potentially attractive tool because it is easy to use, noninvasive, and provides point-of-care analysis. Ultrasound has the added advantage of facilitating dynamic evaluation of tendons through their range of motion.

In a study of 34 patients with upper extremity tendon injuries treated by emergency physicians, ultrasound was faster and more accurate than exploration, magnetic resonance imaging (MRI), and specialty consultation. Accuracy of sonography for detection of hand and digit tendon injuries greatly depends on operator experience and is not a routine component of emergency medicine ultrasound practice.

Definitive examination of any wound must occur under the best possible conditions, with a good light source, a bloodless

Figure 48.3 Flexor and extensor tendons in the fingers. A, Posterior (dorsal) view. B, Finger in extension: lateral view. C, Finger in flexion: lateral view. Note the directions of force when the extensor tendon is activated (arrows). (A–C, Netter illustrations used with permission of Elsevier Inc. All rights reserved.)
Figure 48.4 Zone of convergence of the digital extensor mechanism, which begins at approximately the midportion of the proximal phalanx and ends at the level of the insertion of the central slip into the dorsal base of the middle phalanx. Proximal to the zone of convergence, the extrinsic and intrinsic components of the extensor mechanism are separate: the central slip is extrinsic, whereas the lateral slips are intrinsic. Within the zone of convergence there is complete reciprocal crossover of fibers from the central slip and lateral slips. The products of the completed convergence are the central slip insertion and the conjoined lateral bands, both of which have dual muscular activity. E, Extrinsic contribution to the conjoined lateral bands; I, intrinsic contribution to the insertion of the central slip; ORL, oblique retinacular ligament; PIP, proximal interphalangeal; TRL, transverse retinacular ligament. (From Thomas JS, Peimer CA: Extensor tendon injuries: acute repair and late reconstruction. In Chapman MW, editor: Operative orthopaedics, ed 3, Philadelphia, 2001, JB Lippincott, p 1500.)

Figure 48.5 The extensor mechanism on the dorsum of a finger. Arrows point to the radial and ulnar lateral band portions of the extensor mechanism, and the probe is lifting the entire structure up off the phalanx.

Figure 48.6 The terminal extensor mechanism.

Figure 48.7 Because of their superficial location, it is difficult to avoid at least partial injury to the extensor tendons with even superficial lacerations of the dorsum of the wrist, hand, or fingers. A, This complete extensor tendon laceration is obvious because the index finger cannot be extended. B, This partial tendon laceration was not appreciated on initial examination, which seemingly demonstrated full tendon function. The entire tendon could not be visualized because of an uncooperative patient. The unappreciated partial laceration progressed to a complete rupture by the time of suture removal. Expeditious delayed primary repair resulted in a good outcome.
field, adequate local anesthesia, and a cooperative patient. It may be impossible to adequately assess some patients completely during the first ED visit. In this case, final diagnosis must be delayed until the proper circumstances permit the required conditions. Occasionally, patient noncompliance thwarts even the most carefully planned follow-up. Frequently, the patient's pain, swelling, anxiety, or degree of intoxication or altered sensorium limits the clinician's diagnostic acumen. Therefore it would not be considered standard to diagnose the presence or the full extent of all extensor tendon injuries immediately.

Whenever logistically possible, emergency providers should consult a specialist when an extensor tendon injury is suspected by mechanism, location of the wound, or tendon dysfunction. Under most circumstances, however, there is no value in obtaining an immediate on-site consultation with a hand or orthopedic surgeon because the scenario would similarly limit any clinician's diagnostic acumen.

If the examining clinician suspects but is unable to locate a tendon laceration, or if a patient is uncooperative with the examination and the circumstances prohibit ideal initial care, patients should be referred for follow-up in 1 to 3 days for a repeat examination. Close the skin and apply a splint for interim wound care. A delay of a few days for definitive diagnosis, surgical repair, or both does not result in any significant alteration in the final outcome. Delayed primary repair, without

---

**Figure 48.8** To examine for a tendon injury, use a bloodless field. Note the sterile glove on the patient to maintain a clean field. It is almost impossible to cut the dorsum of the hand or fingers and avoid at least a partial tendon injury. **A,** The location and depth of this laceration suggest an extensor tendon injury. On examination, the patient had full extension. **B,** No tendon injury is visualized when the laceration was examined with the fingers in extension. When the laceration was extended and probed with the finger flexed, a 60% laceration of the extensor tendon could be viewed in the depths of the wound. **C** and **D,** Note the typical shiny white appearance of the fully exposed tendons (arrows).

**Figure 48.9** Given the superficial location of extensor tendons, suspect a tendon injury even with seemingly superficial lacerations of the dorsum of the hand and fingers. Full function is possible with a significant tendon laceration, and delayed total rupture can occur days to weeks later if the injury is not repaired or splinted. Most partial extensor tendon lacerations do well with 3 to 4 weeks of splinting and no surgical repair. When in doubt, clean the laceration, suture the skin, splint, and refer for subsequent examination in a few days.
the need for tendon grafting or tendon transfer, is a well-accepted technique. In fact, many hand surgeons are reluctant to immediately repair even a complete extensor tendon laceration in a contused, potentially contaminated wound. The exact time frame under which such delayed repair results in an outcome similar to that of immediate repair is not well defined and depends on the clinical scenario. Usually, repair delayed for up to 10 days will still ensure an outcome similar to that of an immediate repair, but this varies depending on the injury. Providers should clearly document the inability to rule out a tendon injury in the ED and the mandate for follow-up within a specified time frame on the medical record and discharge instructions.9

Use of Antibiotics
There are no data to support or refute the use of prophylactic antibiotics as a routine adjunct after tendon injury. In general, prophylactic antibiotics have not been demonstrated to reduce infection rates after soft tissue injury in the setting of proper wound cleaning. Nor have they been proved to reduce infection rates in the absence of gross contamination, retained foreign material, extensive contusion, or a delay in cleaning. Many clinicians opt for antibiotics with gram-positive (e.g., antistaphylococcal) coverage if the tendon has been injured or sutured, but no universally accepted standard of care exists. An individualized approach is advocated. Prophylaxis is generally used for only 3 to 5 days after injury unless there are extenuating circumstances such as immunocompromise, diabetes, a human bite, an unusual source of contamination, or peripheral vascular disease. If the sterility of a wound is in doubt, do not attempt tendon repair.

Preparation for Repair
Before attempting repair of an open extensor tendon injury in the ED, be prepared and have the proper equipment available. Place the patient supine on a gurney that ideally has an arm board attached. Bright overhead lighting is important for wound exploration so that the presence of tendon injuries and foreign bodies can be adequately assessed. Instruments should include, at a minimum, a needle holder, two skin hooks and retractors, sharp (i.e., iris) and blunt-nosed scissors, several small hemostats, and one pair of small single-toothed (i.e., Adson) forceps.

The choice of suture material depends on the location of the tendon injury. For repair of complete tendon injuries on the dorsum of the hand, non-absorbable, synthetic braided suture is preferred. Polyester suture, such as Ethibond (Ethicon, Somerville, NJ) or Mersilene (Ethicon), is recommended. Nylon suture is acceptable but is less ideal because colored nylon may be visible under the skin. Chromic and plain gut should be used to avoid tearing the tendon. Partial tendon injuries in smaller tendons. Small, “plastic repair” tapered needles should be used to avoid tearing the tendon. Partial tendon injuries in the digits are best repaired with fine, synthetic absorbable suture such as polyglactin (Vicryl [Ethicon]). Complex lacerations that involve tissue loss and fraying of the tendon margins (e.g., table-saw injuries) represent a particularly challenging clinical scenario that may make an otherwise straightforward tendon repair very difficult. In these cases, Lalonde and Kozin recommend closing the lacerated skin and tendon together (i.e., dermatotenodesis). Take large, composite bites of skin and tendon together, 5 to 10 mm on either side of the wound, with 3-0 or 4-0 nylon suture tied outside the skin. Tighten the suture until the digit is in full extension.9

Before repairing a tendon injury, it is imperative that the provider use adequate anesthesia so that thorough wound exploration can occur. A field block or regional nerve block can be used on the dorsum of the hand, whereas local anesthesia or a digital nerve block can be used on the fingers. The choice of anesthetic composition has been the subject of long-standing controversy. Traditional teaching admonishes the use of epinephrine-containing anesthetic for fear of digital ischemia; however, many clinicians readily use lidocaine with epinephrine in the hand and fingers without complications. There is ample anecdotal and clinical evidence supporting the safety profile of epinephrine in digital anesthesia. Epinephrine has the benefit of prolonging the anesthetic effect and promoting a bloodless field during wound exploration and repair.10 It is important that the digits be fully anesthetized or, in the case of more proximal wounds on the hand, that the area around the wound be liberally anesthetized because many lacerations must be extended to afford access to the surgical field. It is a common error to avoid extending a laceration and to attempt examination, cleaning, or repair through a small initial skin laceration.

Following the administration of an anesthetic, place a tourniquet on the involved limb if hemostasis is problematic (see Fig. 48.8). It is absolutely essential that adequate control of blood flow be obtained before attempting to repair a tendon laceration. It is very difficult to find the proximal end of a retracted tendon in a bloody field. Before applying a tourniquet, wrap the patient’s arm in several layers of cast padding as a comfort measure and elevate the arm for at least 1 minute to allow blood to drain by gravity. Place a blood pressure cuff on the middle to upper part of the arm, wrap several more layers of cast padding around the cuff, and then inflate it to 260 to 280 mm Hg. Once inflated, clamp the tubes tightly with a hemostat. The use of cast padding during inflation helps avoid premature opening of the cuff. Use of a hemostat to clamp the blood pressure cuff tubes helps avoid cuff deflation. A blood pressure cuff tourniquet is generally well tolerated by patients for approximately 15 to 20 minutes. If tendon repair cannot be accomplished in this time it is likely that the injury is too complex for repair in the ED. When necessary, use parenteral analgesia or anxiolysis to help the patient tolerate a longer tourniquet time. For finger examination, placing a rubber ring tourniquet at the base of the finger should give excellent hemostasis.

Atraumatic technique is essential for minimizing adhesions and scar tissue formation. Tendons should be handled delicately, avoiding crushing force or excessive punctures with forceps and needles. Forceps should be used only on the exposed, cut end of the tendon whenever possible.11

Patterns of Injury and Management
Treatment of extensor tendon injury (Video 48.1) depends primarily on whether the injury is open or closed, as well as the anatomic location of the injury. The most widely accepted classification system is that developed by Verdan,12 which divides the hand and wrist into eight anatomical zones (Fig. 48.10). It is quite useful for emergency providers to become familiar with this classification because in many instances the zone of injury can help determine whether tendon repair should be attempted in the ED. One must keep in mind that repair of
Although this tissue allows smooth gliding of tendons during normal activity, the presence of synovium increases the risk for adhesions after tendon repair. In addition, lacerated tendons in the wrist and distal part of the forearm may retract away from the site of initial injury. This may make tendon retrieval and repair quite difficult and necessitate incision of the retinaculum and exploration of one or more compartments.

As a result of the potential complexity of these injuries, all tendon lacerations in zones 7 and 8 require formal surgical exploration and repair. ED management of these patients includes local wound care with primary repair of the skin and placement of a volar splint in 35 degrees of extension at the wrist and 10 to 15 degrees of flexion at the MCP joints. Promptly refer these patients to a hand surgeon so that repair may be undertaken within 1 week of injury.

**Zone 6 Injuries**

Zone 6 consists of the area over the dorsum of the hand. Extensor tendon injuries in this region are frequently caused by lacerations from broken glass or another sharp object. Common pitfalls in ED management of these injuries are usually related to failure to recognize that the tendon has been injured. It is important to remember that these tendons are superficially located, partial tendon lacerations may occur, and weak extension of a digit is possible with a complete tendon laceration because of transfer of extensor function through the jucturae tendinum. Lacerations of the EIP or EDM tendons are evidenced by an inability to extend independently the index or little finger, respectively. In most cases, missing zone 6 injuries can be avoided if a careful physical examination is performed, including thorough wound exploration under sterile conditions using a tourniquet, adequate local anesthesia, and good lighting.

Extensor tendon injuries in zone 6 are generally acceptable for repair in the ED. Because of the juncturae tendinum, extensor tendons in zone 6 are less likely to retract than those in zones 7 or 8; however, the severed tendon may retract when the injury is more proximal. The distal end of a severed tendon is usually easy to find by passively extending the patient’s affected digit to bring the end into view. Retrieval of the proximal portion of a severed tendon is sometimes required and can usually be accomplished in the ED. Before searching for the proximal end of the tendon, the clinician should have a 4-0 nylon suture loaded onto a needle holder. When the proximal end is located, place this suture as a holding suture as far proximal as possible so that the tendon is not lost again. It is often necessary to use a scalpel to extend the skin wound proximally in a direction parallel to the course of the injured tendon to obtain adequate exposure. One should then begin to search for the tendon by lifting up the overlying skin with forceps and inspecting the proximal portion of the wound. Sometimes the blood-stained end of a tendon tunnel can be seen; this may contain the proximal end of the tendon. By gently placing a small hemostat or toothed forceps up this tunnel, the tendon stump can often be pulled into view.

Once both ends of the injured tendon have been located, the technique used for repair depends on the size and shape of the tendon. Whereas larger, round tendons can accommodate sutures that pass through the core of the tendon, smaller or flat tendons are difficult to repair with this technique. Most of the tendons in zone 6 can be repaired using one of several published techniques that maximize the strength of repair and minimize the risk of adhesions and catching. A detailed description of all available tendon repair techniques is beyond the
scope of this text. Emergency providers should be familiar with some of the more common approaches such as the modified Kessler, modified Bunnell, modified (or augmented) Becker (aka Massachusetts General Hospital), and Krackow-Thomas techniques, and be prepared to apply one or more depending on the injury pattern.6,14,15

Both the modified Bunnell and modified Kessler techniques involve first placing a single suture in one end of the cut tendon, usually made of 3-0 or 4-0 non-absorbable material. Place the suture in the tendon core by inserting the suture needle into the exposed, cut end and then weaving the suture through the lateral tendon margins. Next, place the same suture through the core of the opposite half of the cut tendon. Tie the suture ends in a square knot in between the cut ends of the tendon to bring the two halves together. To improve the tensile strength of the repair, a number of other suture techniques may be used.8 One option is to increase the number of suture strands that cross the repair site (e.g., four strands rather than two). A cadaver study comparing various four-strand tendon repair techniques concluded that the Massachusetts General Hospital technique was more resistant to gap formation than either the Krackow-Thomas or the four-strand modified Bunnell techniques.16 However, this cadaver model could not assess tendon shortening or subsequent range of motion.17 Another way to improve tensile strength is to place a peripheral suture in addition to the core suture. Place a running cross-stitch suture of synthetic, absorbable material (e.g., 5-0 polyglycolic acid, polyglactin, or polydioxanone) circumferentially around the repair site or just on the dorsal surface of the tendon across the laceration site. Alternatively, place sutures laterally along both sides of the tendon starting at approximately 1 cm of either side of the repair site.

The running interlocking horizontal mattress (RIHM) is a newer technique recommended by some authors for primary extensor tendon repair (Fig. 48.12). Using 4-0 non-absorbable suture, the RIHM begins with a simple running suture directed away from the surgeon. Then, using the same suture, the operator places a running mattress directed towards the examiner, interlocking each returning throw underneath the previously crossed suture to secure each pass. Tie the suture at the near end on the outside of the tendon to complete the repair. In a cadaver model of zone 6 injuries, the RIHM repair was significantly stiffer, resulted in significantly less shortening, required less tendon exposure, and took less time to perform than both the augmented Becker and modified Bunnell techniques. There was no significant difference in failure rates. This technique has the added benefit of being able to withstand the forces of early active motion during the postoperative period. It also preserved more tendon length and decreased flexion loss, potentially improving grip strength after repair.18,19

Smaller tendons may be repaired with a figure-of-eight or horizontal mattress suture (Fig. 48.13). Small, tapered needles should be used to avoid tearing the tendon. In a cadaver study comparing multiple suture techniques, it was found that the modified Bunnell technique provided the strongest extensor tendon repair.20 In addition, this technique produced no gapping between the repaired tendon ends and minimized the postrepair restriction of flexion at the MCP and proximal interphalangeal (PIP) joints. Alternatively, some experts recommend the RIHM for extensor tendon repairs. There are no studies that clearly show superiority for any particular technique, and not all hand surgeons agree on the repair approach, even for the same or similar injuries. The ultimate choice of repair technique will depend largely on the treating providers’ familiarity with extensor tendon repair, as well as the size of the tendon. Regardless of the chosen suture technique, it is important to test the degree of flexion passively at the MCP joint after a zone 6 tendon repair to be certain that the tendon has not been excessively shortened.

The approach to partial extensor tendon lacerations is not well defined and no definitive standard of care exists. One evidence-based analysis identified 141 relevant papers, but none
... were germane to the question of repair of partial extensor tendon injuries. The authors concluded that there is no direct evidence to assist in answering this question. Given the lack of literature on the subject, a reasonable approach may be to extrapolate from data on flexor tendon injuries. It has been demonstrated that many partial flexor tendon lacerations do well without repair, but hand surgeons still disagree on the need for repair of these injuries. In a survey of hand surgeons, 30% of respondents repaired all partial flexor tendon lacerations and 45% repaired only lacerations with greater than 50% involvement of the cross-sectional area. Except at the wrist level, extensor tendons are not covered with synovium and are therefore less likely than flexor tendons to form adhesions after repair. This encourages some authors to recommend repair of most partial extensor tendon lacerations. Although the ideal approach to these injuries is not known, it is reasonable to consider repair of partial extensor tendon lacerations to be optional if less than 50% of the cross-sectional area is involved. However, if not repaired, such injuries must be splinted for 3 to 4 weeks to ensure that a partial laceration is not converted into a complete injury. Skin closure, splinting, and referral for follow-up is a standard approach to unrepaired partial extensor tendon lacerations.

After repair of a lacerated EDC tendon in zone 6, apply a volar splint so that the wrist is in 30 to 45 degrees of extension, the affected MCP joint is neutral (0 degrees of flexion), and the unaffected MCP joints are in 15 degrees of flexion. The PIP and distal interphalangeal (DIP) joints should be allowed full range of motion. After 10 days, the MCP joints are allowed 20 to 30 degrees of flexion. If there is an isolated EIP or EDM tendon injury, only the index or little finger must be included in the splint. Dynamic extension splinting may be used as early as 2 days after tendon repair, so close follow-up is recommended.

**Zone 5 Injuries**

Zone 5 consists of the area over the MCP joint. Open injuries in this region should be considered secondary to a human tooth bite until proved otherwise (Fig. 48.14), especially if the injury occurs over the first or second MCP joint because this is frequently the location of a clenched-fist (“fight-bite”) injury. ED evaluation must begin with a careful and persistent history and physical examination, although patients may be reluctant to admit to punching someone in the mouth. The wound should be inspected through its full range of motion because the positions of the EDC tendons change with hand position. It is generally recommended that radiographs be obtained for all these injuries to evaluate for metacarpal head fractures, air in the joint space, or the presence of a foreign body such as a tooth fragment (Fig. 48.15). If, after a thorough evaluation, it is determined that a human bite in this region has resulted in a superficial skin laceration only, without injury to the underlying tendon or joint, outpatient management is appropriate. The wound should be copiously irrigated and left open, a volar splint applied with the wrist in 45 degrees of extension, the MCP joints in neutral position (0 degrees of flexion), and the hand dressed with a bulky dressing. The use of prophylactic antibiotics for these “low-risk” human bites on the hand is controversial and clinical trials have yielded mixed results. Despite the lack of compelling clinical evidence for either approach, many authors recommend...
that 3 to 5 days of prophylactic antibiotics be given to these patients. Regardless of whether antibiotics are prescribed, patients should be seen in 24 to 36 hours for a repeat examination to evaluate for wound infection.

If a human bite results in tendon damage, including partial or complete laceration, some clinicians opt for admission and intravenous antibiotics. However, no specific standard of care exists. Outpatient therapy is acceptable in a reliable patient who has access to follow-up. Delayed closure with evaluation and repair of the tendon should be undertaken by a hand surgeon after 7 to 10 days of antibiotic therapy. Primary closure of even seemingly clean and well-irrigated human bites in this region is not advisable because of the increased risk for wound infection, as well as the potential for septic destruction in cases of traumatic MCP joint arthrolysis. If an open joint is noted on physical examination, a more aggressive approach is warranted. Such patients are generally admitted for intravenous antibiotics after copious irrigation.

If a patient suffers a zone 5 tendon injury, and it can be determined with complete certainty that it was caused by a relatively clean, sharp object rather than a human bite, primary closure is appropriate. Referral of these injuries to a hand surgeon is common practice given the complexities of the injury and possible sequelae. Careful repair of lacerations involving both the EDC tendon and the sagittal band is necessary to prevent subluxation of the EDC tendon away from the center of the metacarpal head. Initial ED management of non–human bite injuries is often limited to skin closure, splinting as described earlier, and referral to a hand surgeon within 1 to 5 days.

Closed extensor tendon injuries in zone 5 usually result from the acute or recurrent application of compressive force to the MCP joint capsule. Closed injuries in this region are sometimes referred to as a boxer’s knuckle (Video 48.2). Repetitive closed injury to the MCP joint region can produce small tears in the EDC tendon, the sagittal band, or the joint capsule. These patients tend to have chronic and recurrent pain and swelling in the MCP joint region but usually have normal radiographic findings. Acute trauma may result in the same injuries or cause more severe damage to the extensor hood. Such patients may have complete disruption of the extensor mechanism including damage to the central tendon and the sagittal band. The MCP joint is swollen, has decreased mobility, and may exhibit an extensor lag. Traumatic subluxation of an EDC tendon may be present and usually involves the middle finger with subluxation to the ulnar side (Fig. 48.16). Dislocation to the radial side is less common, probably because of the juncturae tendinum on the ulnar side that can compensate for injuries to the sagittal band. The subluxation becomes more prominent with flexion at the MCP joint. Controversy exists regarding the initial management of closed injuries in this region. Whereas some authors prefer initial surgical repair, others use an initial trial of extension splinting in some or all cases. Splinting the MCP joint in neutral or slight flexion for 6 weeks has been recommended for dislocations initially seen within 3 weeks of injury. Operative repair is usually reserved for more delayed manifestations or patients who fail splint therapy.

**Zone 4 Injuries**

Zone 4 consists of the area over the dorsal aspect of the proximal phalanx between the MCP and PIP joints. The extensor tendon is a broad, flat structure in this region and is relatively easy to repair. Because the extensor tendon is flat and conforms to the roundness of the proximal phalanx, tendon injuries in this area generally result from a laceration and are almost always incomplete. As a result, extension at the PIP joint is not usually impaired. It is therefore imperative that all these wounds be explored carefully while remembering that the extensor tendon lies immediately beneath the thin overlying skin. Tendons tend to not retract in this area, so close inspection will usually result in location of the injured tendon.

Hand surgeons generally repair central slip lacerations or any laceration that results in an extension lag at the PIP joint. The decision to repair a partial tendon laceration in zone 4, and whether it should be repaired by the emergency provider, is best discussed with the consulting hand surgeon. In general, because of the duality of the extensor system in this region, lacerations of a single lateral slip can either be repaired with 5-0 nonabsorbable suture or left unrepaired and splinted.
Placement of running sutures or simple interrupted sutures with buried knots is appropriate for this area. As placement of core sutures is technically difficult or impossible here, other techniques such as the RIHM are recommended. Postrepair splinting depends on the presence of tension at the repair site. Minor lacerations in zone 4 that do not result in tension on the repair site can be treated with a finger guard for 7 to 10 days and early range of motion. Treat larger lacerations or those that result in tension at the repair site with a splint that extends from the forearm to the digit for 3 to 6 weeks. Apply the splint so that the wrist is in 30 degrees of extension, the MCP joint is at 30 degrees of flexion, and the PIP joint is in neutral position. Group the fingers so that either the index and long fingers, or the long, ring, and little fingers are immobilized.

It is important to recognize that complex partial tendon lacerations (e.g., laceration of a lateral slip with a saw) in zone 4 may result in damage to the gliding layer located between the tendon and the bone. If the patient is still able to actively extend the digit at the PIP joint, these complex partial tendon lacerations are best managed by débriding the frayed tendon ends and splinting the digit in extension rather than attempting to suture the damaged tendon. The splint should be worn for 10 days, followed by active range of motion.

**Zone 3 Injuries**

Zone 3, the area over the PIP joint, is a common site of both closed and open injuries. An open injury usually results from laceration with a sharp object. It is imperative that these wounds be carefully explored in the ED to rule out penetration of the joint capsule (i.e., arthrotomy). Patients with wounds that are suspected of penetrating the joint are generally taken to the operating room for surgical exploration, irrigation, and treatment with intravenous antibiotics, but protocols vary.

Zone 3 tendon lacerations can result in long-term deformity if not carefully repaired, and patients with such injuries are commonly referred to a hand surgeon. Partial lacerations of the central slip or lateral bands are managed variably, and it is advisable to discuss these injuries with the consulting hand surgeon. Lacerations in this area may sometimes result in a complete central slip injury. This may manifest as an acute boutonnière (“buttonhole”) deformity in which the PIP joint rests in 60 degrees of flexion. The signs may be subtle, however, and may be noticeable only by weakened extension at the PIP joint or incomplete extension by only a few degrees.

A boutonnière deformity develops when the central slip is ruptured by an open or closed mechanism that leads to unopposed action of the flexor digitorum superficialis tendon. This results in flexion at the PIP joint, protrusion of the head of the proximal phalanx between the two lateral bands, and disruption of the triangular ligament. When this occurs, the lateral bands are displaced volar to the axis of motion of the PIP joint. The lateral bands then paradoxically become flexors of the PIP joint (Fig. 48.17). In addition, the extensor hood mechanism is pulled more proximally, which results in increased tension on the TEM and hyperextension at the DIP joint. Thus, a boutonnière deformity consists of flexion of the PIP joint with hyperextension of the DIP joint.

Open central slip injuries are usually managed operatively, and complex injuries may require direct attachment of the tendon to bone or tendon reconstruction. If the consulting hand surgeon chooses not to repair the tendon injury immediately, close the skin and apply a splint in the same fashion as described for zone 4 injuries. Thermoplastic splints allow immobilization of the hand without involvement of the wrist but may not be available in the ED setting. Promptly refer these patients to a hand surgeon so that repair may be undertaken within 1 week of injury.

Patients with closed injuries in zone 3 are commonly encountered in the ED. They may complain of a direct blow to the dorsal PIP joint or a “jammed” finger. This injury occurs when an object such as a ball delivers a sudden axial loading force with forced flexion of the PIP joint while it is extended. These patients commonly complain of a painful, swollen PIP joint, which often makes the examination difficult. Some of these injuries may represent PIP joint dislocations that were spontaneously or manually reduced before the arrival of the patient at the ED. The tendon injury that is important to recognize in this setting is an occult isolated central slip rupture. Patients may have decreased extension at the PIP joint, but extension is generally normal because the lateral bands are the primary extensors of this joint. With forced extension against resistance, patients usually have pain and may have decreased strength. To eliminate pain as the cause of the decreased mobility, it may be helpful to test PIP extension against resistance after performing a digital block. With acute central slip rupture, PIP joint extension may be particularly weak when the MCP and wrist joints are held in maximal flexion. In this position, a 15-degree or greater loss of active extension is highly suggestive of a central slip injury. The Elson test may also help identify this injury (Fig. 48.18).

A boutonnière deformity does not usually develop in patients with closed zone 3 injuries until 10 to 21 days after injury unless the central slip and triangular ligament are both disrupted and the lateral bands subluxate toward the volar plate. To prevent the deformity from occurring, the provider should have a high index of suspicion for its presence and treat these patients conservatively. It is advisable that all patients with a swollen, tender PIP joint and pain with flexion or extension be splinted and referred for close follow-up. Apply a dorsal splint overlying the PIP joint while keeping it in full extension. This can be accomplished with an aluminum foam-backed splint or a Bunnell (“safety pin”) splint, although the latter

**Figure 48.17** Boutonnière deformity. This can be an open or a closed injury. Note the flexion of the proximal interphalangeal joint and extension of the distal interphalangeal joint from a laceration of the central slip mechanism.
in part, by the triangular ligament. The TEM inserts on the base of the distal phalanx and allows extension at the DIP joint. Complete disruption of the TEM results in an inability to extend at the DIP joint. Because of the unopposed action of the flexor digitorum profundus (FDP) tendon, the DIP joint rests in the flexed position. The most common closed tendon injury of the hand is known as a mallet deformity of the finger (Fig. 48.20). When evaluating DIP motion, it is important to isolate the function of the extensor tendon by holding the PIP joint in full extension (see Fig. 48.20B and C). Normally, full active extension is possible.

Tendon lacerations in zones 1 or 2 that result in a partial or complete mallet deformity generally warrant discussion with a hand surgeon (Fig. 48.21). Management consists of repair of the lacerated tendon and postrepair immobilization. Some surgeons will use only an external splint; others prefer placement of a Kirschner wire (K-wire) through the distal phalanx into the middle phalanx to help stabilize the joint. One technique for tendon repair is dermatotenodesis, which involves placement of a single, running roll-type suture through the tendon and overlying skin (Fig. 48.22). The DIP joint is then splinted in full extension for at least 6 weeks. Occult partial tendon lacerations are important to recognize to prevent the development of a mallet deformity. If there is a partial tendon laceration in zone 1 or 2 that does not result in an extension lag, the approach to repair is variable, and it is advisable to discuss the repair with the consulting hand surgeon. In general, partial tendon lacerations involving less than 50% of the tendon area that do not result in an extension lag may be splinted in extension for 7 to 10 days with or without repair of the tendon itself. Partial tendon lacerations involving more than 50% that do not result in an extension lag may be repaired by a hand surgeon or an emergency clinician who is experienced.

**Figure 48.18 A**, Diagram of a boutonnière deformity. **B–E**, The Elson test for early diagnosis of an acute rupture of the central slip of the extensor digitorum communis tendon. Such rupture results in a boutonnière deformity in which the distal interphalangeal joint is hyperextended, as shown. **B**, With the patient’s finger flexed (over a straight edge) at the proximal interphalangeal (PIP) joint, palpate the dorsal surface of the middle phalanx. **C**, If the central slip is intact, PIP joint flexion causes the slip to tighten distally, thereby relaxing the lateral bands and leaving the distal phalanx flail (arrows). Thus, when the patient is asked to extend the digit, the examiner feels pressure that is being exerted by an intact central slip. **D** and **E**, If the central slip is disrupted, however, the examiner feels no pressure on the dorsum of the middle phalanx as the patient tries to extend the digit. It is possible for the patient to extend the injured finger successfully only by hyperextending (by action of the lateral bands) (arrows).

**Figure 48.19 A**, Boutonnière splint. **B**, This splint allows active flexion at the metacarpophalangeal and distal interphalangeal joints.
be approximated but not pulled too tightly; otherwise, joint stiffness and limitation of flexion will occur. After repair of a partial tendon laceration, splint the DIP joint in extension for 6 to 8 weeks followed by 2 to 4 weeks of night splinting and active range-of-motion exercises. Patients should be warned after tendon repair that there is likely to be some residual loss of flexion at the DIP joint, even in the best case.

Closed injuries in zones 1 and 2 may result in a partial or complete mallet deformity, depending on the injury pattern. These injuries are usually caused by an axial loading force with forced flexion of the DIP joint while it is being held in extension. A common ED scenario involving this injury is a patient who complains of pain and swelling at the DIP joint after a ball strikes the fingertip.

Closed injuries in zones 1 and 2 may result in a partial or complete mallet deformity, depending on the injury pattern. These injuries are usually caused by an axial loading force with forced flexion of the DIP joint while it is being held in extension. A common ED scenario involving this injury is a patient who complains of pain and swelling at the DIP joint after a ball strikes the fingertip.

Closed injuries in zones 1 and 2 may result in a partial or complete mallet deformity, depending on the injury pattern. These injuries are usually caused by an axial loading force with forced flexion of the DIP joint while it is being held in extension. A common ED scenario involving this injury is a patient who complains of pain and swelling at the DIP joint after a ball strikes the fingertip.

Closed injuries in zones 1 and 2 may result in a partial or complete mallet deformity, depending on the injury pattern. These injuries are usually caused by an axial loading force with forced flexion of the DIP joint while it is being held in extension. A common ED scenario involving this injury is a patient who complains of pain and swelling at the DIP joint after a ball strikes the fingertip.
found inadequate data to establish the most effective type of splint, but the stack splint is the editor's preference for ease of application and patient comfort. Be careful to avoid excessive sustained pressure from the splint on the DIP joint area because skin necrosis may occur. Strictly maintain the DIP joint in full extension for 6 to 8 weeks, including during sleep and splint changes. Adherence to this instruction is essential because patients have a tendency to test its function on their own, thus tearing the healing tendon fibers. The most common reason for treatment failure is patient noncompliance with prolonged splinting. Patients should support the distal fingertip in full extension whenever the splint is removed. Should DIP joint extension be lost at any point during the initial treatment period, reset the treatment clock for an additional 6 weeks.

The third type of closed injury is an intraarticular avulsion fracture of the dorsal lip of the distal phalanx with volar displacement of the remaining portion of the distal phalanx (Fig. 48.24). Such injuries are best referred for definitive treatment consisting of either surgery or more complex splinting. Normally, the DIP collateral ligaments hold the distal phalanx in place; however, if there is a large enough fracture fragment (usually >50% of the articular surface), the remaining distal phalanx fragment displaces in the volar direction secondary to unopposed action of the FDP tendon. When volar displacement of the distal phalanx occurs, this injury may require more aggressive treatment to achieve an optimal outcome. Unfortunately, there are no adequate randomized, controlled trials comparing operative versus conservative treatment of these injuries. Operative repair usually involves open reduction (Video 48.3) and internal fixation of the fracture with placement of a K-wire for additional stabilization. It is important to remember that it is the presence of volar subluxation, not the size of the avulsion fracture, that is most often considered when determining the need for operative management.

Any injuries, whether open or closed, that result in complete disruption of the TEM may result in a swan neck deformity (Figs. 48.25 and 48.26). This deformity consists of flexion at the DIP joint (a mallet finger) and hyperextension at the PIP joint. It results from increased extension force on the middle phalanx caused by dorsal and proximal displacement of the lateral bands. This complication can often be avoided if disruption of the TEM is diagnosed and treated correctly in the ED.

**Complications**

All extensor tendon repairs are subject to the usual complications of wound infection and skin breakdown secondary to prolonged

---

**Figure 48.23** Mallet finger splints. Be careful to avoid direct sustained pressure from the splint on the skin in the area of the distal interphalangeal (DIP) joint. Excessive pressure or hyperextension can cause skin necrosis. The splint should allow easy motion of the proximal interphalangeal joint with no flexion of the DIP joint. Immobilize the DIP joint in full extension or slight hyperextension (5 to 10 degrees). Splints are maintained continually for 6 to 8 weeks, including during sleep, with strict avoidance of any flexion during hand washing or splint changes. A, Commercially available and an ideal and preferred volar plastic splint (Stack mallet finger splint). B, Dorsal aluminum foam splint. C, Volar aluminum foam splint. D and E, Kleinert mallet finger splint. The Kleinert splint provides modest hyperextension and avoids pressure on the skin by removing the middle third of the foam padding, thereby eliminating all direct pressure on the injury site. All give good results.

**Figure 48.24** A mallet fracture with volar subluxation of the distal phalanx. Avoid attempts to reduce any displaced fractures before splinting because any reduction is unlikely to be maintained without surgery; mallet fingers with associated fractures are best referred. Treat both type 1 and type 2 injuries by splinting in full extension for 6 to 8 weeks. Either a dorsal or palmar splint should hold the DIP joint in extension or slight hyperextension (5 to 10 degrees) while allowing free range of motion of the PIP joint (Fig. 48.23). With a properly fitted splint, no flexion of the DIP joint should occur. The splint can be constructed from aluminum, foam-backed splint material, or from a prefabricated stack splint (Stax, North Coast Medical, Inc. Gilroy, CA). A Cochrane review of mallet finger injury treatments
splinting. Tendon rupture is a rare complication after tendon repair and may result from inadequate suture technique or premature motion against resistance. When extensor tendons are repaired, it is important to use at least five throws and to tie a square knot. All extensor tendon repairs require some period of complete immobilization during tendon healing, and the emergency provider must stress the necessity for patient compliance.

Extensor tendon injuries in zone 7 tend to have the worst prognosis. Because of the presence of a synovial lining, postrepair adhesions may develop. Adhesions may lead to decreased excursion of the extensor tendons with resultant decreased mobility at the wrist. There may also be limitation of finger flexion when the wrist is flexed, as well as limitation of finger extension when the wrist is extended. Because of the lack of synovium, the low risk for adhesions, greater tendon excursion, the relatively simple anatomy, and the usual lack of associated injuries, zone 6 tendon injuries tend to have fewer complications than other areas of the hand. The tendons in zone 6, however, do have a tendency to shorten if the ends are approximated too tightly. This may result in restriction of PIP and MCP joint flexion. In addition, worse outcomes may occur with complex zone 6 tendon injuries when additional soft tissue or bony injuries are present.

Zone 5 injuries are particularly prone to infection because injuries in this region commonly occur from a human bite. In addition, if the extensor hood covering the MCP joint is not repaired carefully, subluxation of the EDC tendon may occur. Tendon shortening and stiffness may result if complex partial tendon lacerations in zone 4 are managed too aggressively. As discussed previously, these injuries are often best managed by splinting alone. A common complication of zone 3 extensor tendon injury is the development of a boutonnière deformity, which usually results from failure to diagnose or adequately immobilize a central slip injury. Similarly, undiagnosed or improperly treated extensor tendon injuries in zones 1 and 2 may lead to either a swan neck or a chronic mallet deformity of the digit. DIP joint splinting itself may result in skin ulceration or tape allergy, often occurring in the second week of treatment. Skin breakdown may ensue if the DIP joint is splinted in hyperextension because of decreased skin perfusion.

Postrepair Care and Rehabilitation

Proper care after diagnosis and repair of an extensor tendon injury is extremely important for optimal patient outcome. Even the best initial tendon repair can have a poor result if subsequently treated improperly. Rehabilitation of tendon injuries has evolved since 1980 to include dynamic splinting and active range-of-motion exercises to achieve maximal motion of the affected digit.

Zone 1 and 2 injuries are usually treated with static splinting, as described previously. After 6 to 8 weeks, active range-of-motion exercises should begin. Night splinting is recommended for an additional 2 to 6 weeks after initial immobilization. Some authors also recommend wearing the splint during the day when performing heavy tasks. It is advisable to give the patient a number of extra splints so that the patient (or family) can change the splint frequently to avoid pressure injury. During splint changes it is important that the DIP joint be held in full extension either by using the other hand or by placing the finger against a table. If an extension lag develops at any time, continuous splinting must be repeated. Closed injuries of the central slip (zone 3) are often treated with a boutonnière splint for 4 to 6 weeks, followed by 2 to 6 weeks of gradual flexion exercises and night splinting. During the initial period of immobilization, the patient should be instructed to passively flex the DIP joint every hour to maintain gliding and proper position of the lateral bands.

Lacerations in zones 3 and 4 have traditionally been treated with static splinting from the forearm to the digits. An alternative approach is to splint only the DIP and PIP joints in extension and begin a “short-arc-motion” protocol within 1 to 2 days of repair. This consists of active motion at the PIP joint progressing from 0 to 30 degrees in the first 2 weeks, then to 0 to 50 degrees in the third and fourth weeks. When compared with static splinting, this protocol may lead to better PIP and DIP joint flexion without resulting in tendon rupture or a boutonnière deformity. Dynamic extension splints are also proving to be useful for rehabilitation of zone 3 and 4 tendon injuries.

Early motion after extensor tendon repair has been found to be most useful in zones 5 through 7. A dynamic extension splint in which the wrist is extended 45 degrees and all finger joints rest in the neutral position is commonly used. A volar block allows 30 to 40 degrees of MCP joint flexion, whereas a dynamic traction mechanism passively extends the digits. Dynamic splinting is started 1 to 3 days after repair. Active motion is added at 3 to 4 weeks, and resistance is added at 7 weeks. A randomized, controlled trial of zone 5 and 6 extensor tendon repairs found total active motion with dynamic splinting was superior to static splinting at 4 to 8 weeks, but not at 6 months. However, grip strength in the affected hand was
improved at 6 months with dynamic splinting. A short-arc-motion protocol with controlled active motion at the MCP joint has also been shown to be safe and effective when started 24 to 48 hours after repair. One comparative trial reported that dynamic extension splinting and controlled active mobilization worked equally well for zone 5 and 6 tendon injuries. All early range-of-motion protocols are most beneficial when managed closely by a skilled hand therapist. Patient age, associated injuries, suture type, and repair technique all affect the choice of rehabilitation protocol. Most importantly, patients must be reliable and motivated to take advantage of early range-of-motion techniques. It is best to refer patients to a hand surgeon or hand therapist as soon as possible after repair so that rehabilitation can begin in a timely manner.

EXTENSOR TENDON INJURIES OF THE FOOT

The extensor tendons of the foot are less commonly injured than the extensor tendons of the hand and wrist. The most important extensors of the foot and ankle that may be injured and encountered in the ED are the tibialis anterior, extensor hallucis longus (EHL), and extensor digitorum longus (EDL) tendons.

The tibialis anterior muscle originates on the shaft of the tibia and interosseous membrane and inserts on the medial cuneiform and the base of the first metatarsal. The tibialis anterior extends (dorsiflexion) the foot at the ankle joint and inverts the foot at the subtalar and transverse tarsal joints. Spontaneous rupture of the tibialis anterior tendon may be seen in both elderly and young patients who have been injured during athletic activity. Injury to this tendon commonly results from forceful attempted dorsiflexion of the ankle while it is held fixed in the plantar-flexed position. Patients generally have decreased strength of foot dorsiflexion because the toe extensors are used to accomplish this motion. Rupture or laceration of the tibialis anterior tendon should be promptly referred to an orthopedic surgeon for consideration of formal operative repair. In some cases, closed injuries of the tibialis anterior tendon may be managed nonoperatively depending on the extent of the patient’s symptoms and functional impairment.

The EDL and EHL tendons both originate from the shaft of the fibula and interosseous membrane. The EHL tendon inserts on the base of the distal phalanx of the great toe, whereas the EDL tendon divides into four branches that insert on toes 2 through 5 (Fig. 48.27). Both the EHL and the EDL tendons primarily result in extension of the toes and dorsiflexion at the ankle. The extensor digitorum brevis (EDB) and extensor hallucis brevis (EHB) muscles originate from the upper part of the calcaneus. The EHB tendon joins the lateral aspect of the EHL tendon before inserting on the great toe. The EDB

**Figure 48.27** Muscles of the dorsum of the foot: superficial dissection. (Netter illustrations used with permission of Elsevier Inc. All rights reserved.)
Injury to the EHL and EDL tendons may result from a sharp object lacerating the dorsum of the foot. Patients may have weakness of, or an inability to extend, the involved toe or toes. The examiner may be unable to palpate the injured tendon. Whether one should repair EHL or EDL tendon lacerations is controversial. However, many authors favor repair because failure to repair EDL tendons may result in a claw deformity of the adjacent toes. Lacerations of the EHL and EDL tendons at the level of the ankle are usually repaired, whereas lacerations on the dorsum of the foot and the toe are variably managed. Repair is favored if the patient has significant pain or any flexion deformity of the involved toe. Repair is also favored when both ends of the tendon are easily visualized in the wound and the patient is willing to undergo prolonged immobilization after repair. Because management of these injuries is controversial, it is advisable to discuss the care of these patients with the consulting orthopedic surgeon or podiatrist. Extensor tendon repair of the foot is not usually performed in the ED setting. Superficial cutaneous nerves are easily injured on the dorsum of the foot during wound exploration, which can lead to the formation of a chronic, painful neuroma. If the injury is repaired in the ED, the technique for repair is similar to that used for the dorsum of the hand (zone 6). A posterior splint that includes the toes should be applied after tendon repair. Splint the ankle at 90 degrees with the toes in the neutral position, and recommend non-weight bearing status until specialty follow-up.

FLEXOR TENDON INJURIES

Flexor tendon injuries are more difficult to diagnose and more challenging to treat than extensor tendon injuries. In general, emergency providers do not repair flexor tendons. Anatomic and biomechanical issues, the physiology of flexor tendons and tendon healing, and follow-up rehabilitation and physical therapy are complex and formidable. A satisfactory outcome is more difficult to achieve with an injured flexor tendon than with a similar degree of injury to an extensor tendon. Unlike extensor tendons, flexor tendons are influenced by a number of complex pulley mechanisms. Flexor tendons glide through delicate tendon sheathes, so even a minor defect in tendon integrity is physiologically magnified (Fig. 48.28). In addition, flexor tendon injuries are often associated with nerve and vascular injuries.

The main clinical mandates for emergency providers are to diagnose or consider flexor tendon injuries, provide initial proper wound care, and expedite appropriate consultation and follow-up. Unlike the more superficial extensor tendons, flexor tendons are often buried deep within the hand and forearm, and it is frequently not possible to visualize the tendon in the recesses of a wound. Puncture wounds of the palm often injure flexor tendons, but deep puncture wounds prohibit visualization of injured structures (Fig. 48.29). Therefore a partial flexor tendon injury may be clinically silent until rupture occurs days or weeks later. Delayed repair of undiagnosed flexor tendons may be complicated by tendon retraction or scar formation, and tendon transfer and grafting may be necessary.

It may not be possible on the initial visit for the emergency provider to diagnose the presence of all flexor tendon injuries, nor the full extent of such injuries. Help may be obtained from a specialist if logistically possible, but generally there is no mandate for such immediate on-site examination when questions about tendon integrity exist. Even though consultation is advised before definitive disposition, the same limitations in the examination would similarly confront a specialist. Individual scenarios and local protocols will guide the timing and degree of consultation in the ED.

Notwithstanding the previous discussion, complete flexor tendon injuries are often apparent on physical examination, either by testing individual tendons or by the resting posture of the injured hand. In contrast, partial tendon lacerations are commonly clinically unappreciated because no functional deficit is evident. Clinical clues to a potential flexor tendon injury are weakness of flexor tendon function (difficult to evaluate in an acutely injured extremity), pain at the site of injury when performing active range of motion against resistance, and an abnormal resting position of the hand (Fig. 48.30A and B). The emergency provider may not be able to arrive at a complete or accurate diagnosis without surgical exploration. However, it is counterproductive and potentially harmful to attempt extensive exploration of the deep recesses of the hand or forearm in the ED merely to visualize a suspected flexor tendon injury.

A specialist, usually on an elective outpatient basis, surgically repairs completely transected flexor tendons. Most hand surgeons are reluctant to perform primary repair of a flexor tendon injury on ED patients and prefer to have the wound cleaned, the skin closed, and the patient scheduled for subsequent definitive repair. The final outcome of flexor tendon surgery depends on multiple factors; however, surgical repair of most flexor tendons accomplished within 10 to 21 days of injury (delayed primary repair) generally produces final outcomes similar to those repaired immediately. Therefore, if a partial tendon laceration is not diagnosed at the initial visit and rupture is noted at the time of removal of the skin sutures or inspection of the wound, immediate referral to a hand surgeon would be expected to provide a similar result had the injury been diagnosed at the time of the initial ED visit.

It is appropriate to treat partial flexor tendon lacerations, if appreciated, with careful wound cleaning, skin closure, splinting, and referral for reevaluation in 1 to 5 days. Definitive treatment of partial lacerations remains quite controversial. Some surgeons will repair all partial tendon lacerations, whereas others take a more conservative approach. The conservative approach is supported by experimental evidence suggesting that surgical repair of partially lacerated tendons results in weaker tendons than if the tendons were not surgically repaired. Wray and colleagues suggest forgoing suturing in favor of splinting, followed by early mobilization of tendons with lacerations involving 25% to 95% of the cross-sectional area. Without conclusive evidence either way, a reasonable approach would be to suture tendon lacerations involving greater than 50% of the cross-sectional area with special surgical techniques, suture tendon lacerations involving 25% to 50% of the cross-sectional area with simple or special suture techniques, and simply trim injuries that affect less than 25% of the cross-sectional area to promote normal gliding function. All decisions concerning the type and timing of repair should be made in concert with a consultant, while keeping in mind that some decisions regarding surgical repair of partial injuries cannot be made for weeks or months.
Figure 48.28 A, Muscles of the forearm (deep layer): anterior view. B, Muscles of the forearm (intermediate layer): anterior view.
will probably improve the final outcome, but flexor tendon injuries often produce lifelong disability despite even ideal care in the ED.7,48

ACHILLES TENDON RUPTURE

An Achilles tendon rupture can lead to serious morbidity. Although definitive care of such injuries is not performed in the ED, it is important to make the correct diagnosis and institute proper and prompt referral. This injury is easy to miss, and it is not always diagnosed on the first visit. Initially considered a minor ankle sprain by both the patient and provider, the diagnosis was missed in more than 20% of cases in a 2008 case series.49 Rupture often occurs with steroid use, with degenerative conditions, and in the elderly, but Achilles tendon rupture can also occur in healthy athletic patients with no history of heel pain and often with seemingly minor trauma.
Fluoroquinolone antibiotics have been implicated in Achilles tendon rupture, especially in the elderly. This led the Food and Drug Administration to issue a “black box” warning on the use of all fluoroquinolones for this condition in 2008. The common mechanism of rupture is sudden overload of the tendon by forceful plantar flexion of the foot, as in recreational sports involving jumping (basketball), pushing a heavy object, or stepping up. The injury is usually a complete as opposed to a partial tear, and rupture commonly occurs in a region 2 to 6 cm proximal to the tendon's insertion on the calcaneus. Occasionally, the patient may appreciate an audible snap or pop. Pain may not be perceived in the tendon itself; instead, heel or diffuse ankle pain may be experienced. Because multiple structures plantar flex the foot, the initial result is weakness of flexion or pain with attempts at flexion against resistance, but many partial lacerations are clinically silent. Despite full function, this wound’s location and depth suggest the possibility of at least a partial tendon injury. The prudent course would include meticulous wound care, splinting, skin closure, and contact with a hand specialist to arrange reexamination in a few days, while cautioning the patient that a flexor tendon injury may be present and delayed repair for up to 1 to 3 weeks yields results comparable to immediate repair. Immediate repair is often eschewed because of swelling and wound contamination. Further care may be required. This palm laceration from the sharp top of a metal can seemed superficial. Function was normal. When examined with the fingers in extension, the tendon was readily visualized, a surprise to the clinician given the benign and superficial appearance of the laceration. The visualized tendon was intact. However, when the fingers were flexed (arrow), the position of the hand when the injury occurred, a 20% to 30% laceration of the tendon was demonstrated. This injury will do well with 3 weeks of splinting and no tendon repair. Follow-up with a hand surgeon in a few days is prudent. Note the outrigger aluminum splint incorporated into a short-arm plaster splint (arrow; see Fig. 48.11).
Sonography is appealing because of its relatively low cost, portability, safety profile, the ability to perform static and dynamic evaluations, and the ability to compare the contralateral side easily. The sensitivity and specificity of ultrasound for Achilles tendon rupture, as reported in the radiology literature, are 96% to 100% and 83% to 100%, respectively. Although no studies have directly evaluated the diagnostic accuracy of ultrasound performed by

**Figure 48.30**

A. Note the obvious abnormal resting posture of the hand. This boy’s palm laceration involved the flexor tendons to his index finger. With his hand at rest, his index finger lies in extension, in contrast to his other fingers, which are partially flexed. B. Loss of the digital cascade in the middle finger illustrated here should be indicative of a flexor tendon laceration without further examination. The small glass laceration in the palm accounts for the profundus laceration, apparent only at follow-up. Children are difficult to fully examine in the emergency department. Splinting and follow-up in a few days are prudent based on the injury mechanism and location. C. The flexor digitorum profundus tendon is examined by immobilizing the digit in question and asking the patient to flex the distal interphalangeal joint against resistance. D. The flexor digitorum superficialis tendon is examined by immobilizing the digits not being tested and asking the patient to flex the proximal interphalangeal joint against resistance. Pain and weakness associated with flexion against resistance may suggest a partial tendon laceration, but this is often a very subtle or inaccurate evaluation that must be repeated when the pain and swelling have subsided. (A, Courtesy Robert Hickey, MD, Children's Hospital of Pittsburgh.)
emergency physicians for Achilles tendon ruptures, bedside ultrasound performed by emergency physicians for tendon injuries in other body areas had a sensitivity, specificity, and accuracy of 100%, 95%, and 97%, respectively.5 A sonographic Thompson’s test can be performed by directly visualizing the Achilles tendon with the high-frequency linear ultrasound probe while the calf muscle of the prone patient is gently squeezed. The Achilles tendon is assessed, proximally to distally, for synchronous movement, using dynamic sonography during passive dorsiflexion and plantar flexion of the ankle with the probe over the area of interest. Complete tears are recognized by retraction of the proximal tendon end, and echogenic adipose tissue (Kager’s fat) may be seen to herniate between the torn ends of the tendon (Fig. 48.33).49 The provider will see continuous tendon movement at the rupture site in the case of a partial rupture because some intact fibers remain.54 Treatment is variable, ranging from conservative splinting to surgical repair. Splinting the foot in mild plantar flexion (gravity equinus) can protect the tendon until follow-up in 1 to 5 days.

KNEE EXTENSOR TENDON RUPTURE41

The extensor mechanism of the knee is composed of the four strong quadriceps muscles, the femoral quadriceps tendon, the patella, the patellofemoral and patellotibial ligaments, the medial and lateral retinacula, the patellar tendon, and the tibial tubercle. Both the quadriceps and patellar tendons are subject to rupture, and the condition occurs predominantly in males. Quadriceps tendon rupture is more common in the elderly and in those with systemic degenerative disease, arthritis, and steroid use, and is associated with significant morbidity regardless of treatment. It may also be seen in younger patients, such as after taking a basketball jump shot. Performance-enhancing steroid use is likewise a risk factor in these patients. Patellar tendon rupture is also a serious injury but occurs more commonly in healthy patients, younger than 40 years, participating in sporting events.43

As with Achilles tendon rupture, rupture of the knee extensor mechanism is not always initially suspected or diagnosed; it is missed on initial evaluation in nearly 40% of cases.56 The mechanism of quadriceps tendon rupture is usually a deceleration injury with the knee partially flexed, coupled with a strong quadriceps muscle contraction when the foot is fixed. The trauma may be seemingly minor, such as missing a step or jumping from a low height. A common history is an elderly patient who is descending steps or walking off a curb, misses a step, and attempts to keep from falling. A popping or tearing
sensation may be elicited. The pain may be deceptively minor. The rupture may be partial but is more often complete. Quadriceps tendon rupture usually occurs transversely, just proximal to the patellar insertion, with or without an avulsion fracture of the superior pole of the patella. A suprapatellar gap may be palpated. The mechanism of patellar tendon rupture is usually an excessive load on the flexed knee during athletic activities. The patient complains of pain and inability to extend the knee or ambulate. Patellar tendon rupture generally occurs at the inferior pole of the patella. Athletic patients may continue to play with a partial tear, but a complete rupture does not allow ambulation. Bilateral complete rupture has been described, but the condition is generally unilateral.

With either type of rupture, a large hemarthrosis is usually produced and often prompts the incorrect diagnosis of a ligamentous injury such as an anterior cruciate ligament rupture. A palpable defect superior or inferior to the patella may be appreciated, but diffuse swelling can mask this finding (Fig. 48.34). Lack of the expected defect can be misleading in the presence of a large hemarthrosis. When the history consists of only minor trauma, and the findings on physical examination are subtle, the provider may incorrectly suspect malingering or noncooperation with the examination. With complete rupture, a supine patient is unable to actively extend the knee or lift a straightened leg off the stretcher, and the knee flexes when posterior thigh support is removed from the raised leg. Weak extension, especially in the sitting position, may be possible if portions of the medial and lateral retinaculae are intact, even with a complete rupture of the central rectus femoris. With complete rupture, the patient cannot walk and the knee gives way immediately. As one would intuit, however, a knee immobilizer allows the patient to apparently walk normally. Partial tears may allow the patient to walk with a peculiar forward-leaning gait that helps support the knee in extension.

Plain radiographs may show the occasional patellar avulsion fracture. With quadriceps tendon ruptures, a low-riding patella (patella baja) may be present as the patella falls inferiorly. Conversely, a high-riding patella (patella alta) is often seen with patella tendon ruptures as the quadriceps tendon and patella retract superiorly. MRI is definitive for identifying nuances of the process. As with Achilles tendon rupture, sonography is an emerging diagnostic imaging tool used by some ED providers to assess for ruptures of the quadriceps and patellar tendons.51 Complete tendon ruptures yield hypoechogenicity over the entire length and thickness of the tendon, whereas a hypoechoic line with a wedge-shaped separation may be seen with a partial laceration. The use of dynamic ultrasound will demonstrate the separated tendon ends moving away from each other with intervening echogenic or echolucent fluid.57 Partial tears of either tendon may be treated surgically, or nonoperatively if there is no functional impairment. Complete tears require surgical repair, usually as soon as possible after the diagnosis is made.55 Although definitive treatment is not undertaken in the ED, early diagnosis may improve the long-term outcome. There is no consensus regarding the timing of repair, although delays are correlated with tendon retraction and atrophy. Therefore most authors recommend early repair.58 The postoperative period of recovery for the elderly is prolonged and difficult. Acute injuries diagnosed in the ED should be treated with the knee fully extended in a knee immobilizer and crutches with a 1- to 2-day follow-up, but admission is often warranted to expedite definitive intervention.45

Figure 48.34 A, This elderly man lost his footing going down the stairs and missed only a single step. He felt a pop and was unable to walk. Arthrocentesis yielded a grossly bloody effluent. An obvious suprapatellar soft tissue defect and an inability to perform a straight-leg raise while supine made the diagnosis of a complete quadriceps tendon rupture obvious. B, This man landed on both feet while jumping off a low curb and then collapsed. Pain was minimal and malingering was suspected. Diffuse soft tissue swelling, bilateral knee effusions, normal radiographic findings, and the ability to walk with bilateral knee immobilizers delayed the diagnosis of bilateral quadriceps tendon rupture until follow-up. C, A step-off above the knee readily identified a complete quadriceps tendon rupture. The patient was unable to lift the leg off the stretcher, an activity that made the soft tissue defect obvious.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Management of Common Dislocations

Robin M. Naples and Jacob W. Ufberg

Joint dislocations are frequently encountered in patients seen in the emergency department (ED). They can range from a simple finger joint dislocation to limb- or life-threatening consequences of high-energy trauma. Keys to clinical assessment and radiographic evaluation of these injuries are discussed along with methods of reduction. The emphasis of the chapter is on simple dislocations that should be diagnosed and initially managed in the ED (Videos 49.1 to 49.22). Fracture-dislocations that commonly require operative intervention and emergency orthopedic consultation are not discussed.

PREPARATION OF THE PATIENT

Although many authors claim that their reduction method is well tolerated without premedication, they have not usually measured the discomfort of their patients quantitatively. There are no rigid, generally accepted guidelines for the use of pharmacologic adjuncts in the management of dislocations. Each patient and each dislocation is unique, and the treating clinician must use judgment regarding whether premedication is required, which agent or agents to use, and what dose to give. In general, the authors suggest the routine judicious use of analgesia with or without sedation for the majority of reductions performed in the ED. A calm, cooperative patient may tolerate attempts at gentle reduction of a major joint such as the shoulder, but even the most stoic of patients may be quite uncomfortable with the manipulations necessary for reduction of a dislocated finger. A radial head dislocation in a child is usually easily treated without analgesia; however, reduction of a hip dislocation is unlikely to be successful without a significant amount of sedation and analgesia. Attempting any reduction technique in an extremely anxious patient without premedication will generally frustrate the operator and further upset the patient and may hinder a successful outcome. When multiple attempts are required and significant force must be exerted because of muscle spasm or an uncooperative patient, there is an additional chance of producing complications during the reduction.

Verbal techniques for alleviating anxiety and discomfort are not to be discounted because they can be of great assistance during joint reduction. In field settings, simple hypnosis techniques have been used successfully for major joint dislocations. In the ED, verbal reassurance and distracting conversation are useful adjuncts. Additionally, the use of handheld tablets has shown to aide a variety of painful procedures in the pediatric population and may be a useful adjunct.

In most circumstances, analgesia or sedation of some sort, or both, will be required; the intravenous (IV) route of drug administration is usually the method of choice because it allows rapid relief of patient discomfort and facilitates repetitive dosing for titration to the desired effect (see Chapter 33). Alternatives to procedural sedation and analgesia include intraarticular injection of local anesthetics, hematoma blocks, peripheral nerve blocks, and regional anesthesia (see Chapters 29, 31, and 32).

GENERAL PRINCIPLES

Clinical assessment of a patient with a dislocation must include a search for fractures or other serious injuries, especially if the mechanism involved high energy. This is most important for hip, knee, and posterior sternoclavicular dislocations. For all dislocations, perform a detailed neurovascular examination of the extremity before focusing attention on the injured joint.

Although many dislocations are clinically obvious, some may escape detection for some time as other injuries or issues dominate the clinical picture. A knee dislocation may be quite obvious in a 170-lb man who displays a deformity of the knee, but in a 400-lb patient, the knee may look deceivingly normal on first glance. The history and mechanism of injury can be quite helpful in certain circumstances. For example, a painful shoulder joint in a seizure patient should prompt assessment for a posterior shoulder dislocation, whereas a history of the knee striking the dashboard is a clue to a potential hip dislocation.

Some dislocations can be difficult to identify on plain radiographs. One should keep a high index of suspicion for dislocation when indicated. Carpal dislocations in the wrist may be subtle, but are clinically suggested by severe pain and swelling. Similarly, superior dislocation of the patella may be mistaken for high riding patella typical of patellar tendon rupture. Unlike anterior shoulder dislocations, which are often easily detected on plain films, posterior shoulder dislocations may be missed on initial radiographs.

Some dislocations will have been reduced before clinician assessment. A careful history will uncover these injuries and prompt the necessary assessment of the ligamentous integrity and the possibility of an associated vascular injury and guide proper immobilization and follow-up care. A dislocated and then spontaneously reduced knee is a severe injury that often escapes detection by even a seasoned clinician’s initial evaluation. Other dislocations that are commonly first seen in a reduced state include finger dislocations, patellar dislocations, and radial head subluxations.

Although the chance that a gentle attempt at reduction will cause a fracture or neurovascular injury is extremely low, careful evaluation before and after reduction, as well as documentation of the patient’s neurovascular status, is important. Frequently, the initial pain of the dislocation is distracting, and paresthesias or a weak pulse may not be readily apparent until the joint has been reduced. When the integrity of the pulse is in question, blood pressure at the wrist or foot may be compared with that in the uninjured extremity, or a pulse oximeter may be applied to the distal end of the fingers (Fig. 49.1).

Pre-reduction radiographs are generally recommended. Reasons include difficulty distinguishing a fracture-dislocation by clinical examination and the potential for medicolegal problems if the fracture is not identified before attempts at reduction. More importantly, certain associated fractures predict a poor outcome with closed reduction and make orthopedic consultation a consideration before such attempts. Exceptions
Occasionally, a fracture is detected on post-reduction radio-
can raise questions about the adequacy of the initial procedure.
without radiographic corroboration of a successful reduction
post-reduction period. Re-injury after release from the ED
not remember the actual successful reduction or the immediate
the ED. Patients who have received sedatives and opioids may
virtually all patients who have had a dislocation reduced in
vascular injuries.

The uninjured extremity may give some helpful clues to underlying
oximeter distal to the injury and comparing the results with those of
applying a pulse
A,
most arterial injuries.

standard techniques to assess vascular injury are the strength of the pulse and capillary refill; this should detect
most arterial injuries. A, Taking a blood pressure reading distal to
the injury with a cuff and Doppler ultrasound, or B, applying a pulse
oximeter distal to the injury and comparing the results with those of the uninjured extremity may give some helpful clues to underlying vascular injuries.

Figure 49.1 Significant vascular injuries from dislocations, such as knee, elbow, or ankle dislocations, are usually obvious, but impaired distal circulation may be subtle or delayed because of a slowly increasing intimal flap arterial lesion. Standard techniques to assess vascular injury are the strength of the pulse and capillary refill; this should detect most arterial injuries. A, Taking a blood pressure reading distal to the injury with a cuff and Doppler ultrasound, or B, applying a pulse oximeter distal to the injury and comparing the results with those of the uninjured extremity may give some helpful clues to underlying vascular injuries.

to this rule include suspected radial head subluxation in young
children and clinical circumstances in which radiographs are
not readily available (e.g., in the wilderness). Clinical conditions
(i.e., vascular compromise or threatened skin penetration) may
dictate the need for immediate reduction without radiographs;
however, the few minutes required for initial radiographic
evaluation rarely increases vascular or neurologic complications
and provides very useful information to the consultant.

Some authors question the need for pre-reduction films in
certain patients with obvious or recurrent anterior shoulder
dislocation.9,10 Although post-reduction radiographs are tra-
ditionally obtained, the need for this in a clinically obvious
successful shoulder joint relocation has been questioned.10,11
The authors suggest that post-reduction films be taken in
virtually all patients who have had a dislocation reduced in
the ED. Patients who have received sedatives and opioids may
not remember the actual successful reduction or the immediate
post-reduction period. Re-injury after release from the ED
without radiographic corroboration of a successful reduction
can raise questions about the adequacy of the initial procedure.
Occasionally, a fracture is detected on post-reduction radio-
graphs that was not obvious on the initial films, or a previously
noted minor fracture may be found to reside in an intraarticular
location.10 Point of care ultrasound is being used more frequently
by emergency medicine physicians for musculoskeletal com-
plaints. A growing body of research is revealing the safety and
efficiency of using this modality for diagnosing dislocations
and confirming successful reduction and will be discussed later
in this chapter. As practitioners gain confidence and experience
with ultrasound in this capacity, the need for both pre- and
post-reduction radiographs may be supplanted by the perfor-
mance of ultrasound.

In general, dislocations of all types are less common in
children than in adults because of the relative weakness of the
epiphysial growth plate with respect to the ligamentous support
of the joint. Therefore, in children the epiphysis will tend to
fracture before dislocation occurs, except in the case of radial
head dislocation (nursemaid’s elbow). Reduction techniques
for pediatric dislocations are generally similar to those used
for adults.

The proper terminology for dislocations describes the
relationship of the distal (or displaced) segment relative to the
proximal bone or the normal anatomic structure. The terms
anterior and posterior are used for most dislocations. Therefore,
if the head of the humerus lies anterior to the glenoid fossa,
the injury is an anterior shoulder dislocation. Similarly, if
the olecranon lies behind the distal end of the humerus, the
injury is a posterior elbow dislocation. In the hand, wrist,
and foot, one uses the terms dorsal and volar. Palmar and
plantar are sometimes used in place of volar to describe the
position of the dislocated part. Dislocations can be open or
closed and may have associated fractures, which require a separate
description.

It is generally accepted that the sooner a dislocation is
reduced, the better. This alleviates the patient’s discomfort
and corrects distortion of the surrounding soft tissues. In some
studies the success rate of reduction is higher when attempted
closer to the time of injury.1 However, there is no reason to
forego an attempt at closed reduction of an “old injury” in
the vast majority of dislocations. Although chronic dislocations
persisting several days, weeks, or longer are often difficult to
reduce in a closed manner, such cases are infrequent and should
not deter the physician from attempting closed reduction.

A certain percentage of all types of dislocations are not
amenable to closed reduction. Inability to complete a closed
reduction is generally the result of interposition of soft tissue
structures or fracture fragments and not necessarily due to
improper technique. If sedation and analgesia are adequate to
permit relaxation of the patient’s muscle tone, reduction should
be relatively straightforward. When reduction under adequate
sedation and analgesia is unsuccessful after several attempts,
 further attempts at closed reduction are inappropriate. Gener-
ally, orthopedic consultation should be obtained after two or
three failed attempts.

Once an attempt at reduction is completed, recheck the
neurovascular status that was documented before the reduction
was performed. For the elbow, hand, and forefoot joints, perform
passive range of motion to assess the stability of the reduction
and to ensure a smoothly gliding joint that is free of intraar-
ticular obstruction. For the shoulder, one must be cautious
after reduction as full passive range of motion may cause
repeated dislocation. Testing the ability to place the palm of
the ipsilateral hand on the contralateral shoulder can safely
assess successful range of motion and confirmation of reduction.
In addition to close monitoring of the medicated patient, proper aftercare involves adequate immobilization of the injured joint for comfort and to prevent repeated dislocation. Recommendations for follow-up care depend on the injury and its severity.

**Timing of Reduction**

Questions often arise concerning the necessity of immediate versus delayed reduction, with the clinician fearing disastrous neurovascular consequences if a dislocation is not manipulated immediately on arrival at the ED. In reality, there is rarely an instance in which pre-reduction radiographs, even portable films, cannot be obtained before treatment. Even if the pulse is weak or the fingers are numb, a few minutes’ delay is usually acceptable to gain important radiographic information on the type of dislocation and the presence of an associated fracture and to provide documentation for follow-up clinicians. Important clinical information may be difficult to obtain, or the specific initial injury may be impossible to reconstruct once the joint has been reduced (Fig. 49.2). Of equal importance, a dislocation with concomitant neurovascular injury should be reduced with the least amount of trauma possible, which often requires a few minutes for induction of analgesia and sedation, a time during which radiographs can be obtained. If a vascular or neurologic abnormality is documented before reduction, the joint should be reduced by the timeliest and least traumatic procedure available. Each case should be handled individually by taking the specific injury, available resources, and the clinician’s experience into account. Although multiple unsuccessful or forceful attempts at reduction in the ED should be avoided with all dislocations, this is especially important in patients...
with vascular or neurologic compromise. Occasionally, the more prudent course is reduction under general anesthesia, but this decision must take into consideration the availability of consultation and other resources.

The remainder of this chapter covers dislocations of the various joints with the exception of carpal (wrist) dislocations, which are complex and require orthopedic consultation, and temporomandibular joint dislocations, which are discussed in Chapter 63.

**SHOULDER DISLOCATIONS**

The human shoulder joint is remarkable for its degrees of motion. The glenohumeral joint has the greatest range of motion of any joint in the body, largely because of the loose joint capsule and the shallow nature of the glenoid fossa. However, these anatomic features that allow for great mobility contribute to its instability, making it the most frequently encountered large joint dislocation. Posterior dislocation is uncommon, mainly because of the anatomic support of the scapula and the thick muscular support in this area. Anterior support is less pronounced, with the inferior glenohumeral ligament serving as the primary restraint to anterior dislocation. The depth of the glenoid fossa is somewhat increased by the fibrocartilaginous glenoid labrum, which forms the rim of this structure.

Most shoulder dislocations are anterior (i.e., the humeral head becomes situated in front of the glenoid fossa). Posterior dislocations are the next most common, but they generally account for less than 4% of shoulder dislocations. Less common variations include inferior (luxatio erecta), superior, and intrathoracic dislocations.

**Anterior Shoulder Dislocations**

As stated earlier, shoulder dislocations are the most common major joint dislocation encountered and reduced in the ED, with anterior dislocations comprising the vast majority. The usual mechanism of injury is indirect and consists of a combination of abduction, extension, and external rotation. Only rarely is the mechanism a direct blow to the posterior aspect of the shoulder. Occasionally, especially with recurrent dislocations, the mechanism is surprisingly minor and can be puzzling to the clinician. An anterior dislocation can be induced by mere external rotation of the shoulder while rolling over in bed or reaching behind oneself for the seat belt. When the first dislocation occurs at a younger age, the recurrence rate is higher. If the first dislocation occurs before 20 years of age, there is an 80% to 92% rate of recurrence. If the first dislocation takes place after 40 years of age, the rate of recurrence is 10% to 15%. Rotator cuff injuries, however, occur more frequently in older patients with anterior shoulder dislocations.

The four types of anterior dislocations are classified according to where the humeral head comes to rest. Subcoracoid dislocations account for more than 75% of anterior dislocations. The other anterior shoulder dislocations include subglenoid dislocation and the uncommon subclavicular and intrathoracic dislocations (Fig. 49.3).

**Clinical Assessment**

The presence of an anterior shoulder dislocation is usually obvious (Fig. 49.4). A posterior dislocation is more subtle in terms of both clinical and radiographic findings. It can be misdiagnosed as a severe contusion (Table 49.1). A patient with an anterior shoulder dislocation supports the injured extremity and leans toward the injured side while holding the arm in abduction and slight external rotation. The patient cannot adduct or internally rotate the shoulder. Visual inspection reveals loss of the rounded appearance of the shoulder because of absence of the humeral head beneath the deltoid region. The acromion is prominent and an abrupt drop-off below the acromion can be seen or palpated. An anterior fullness in the subclavicular region is visible in thinner individuals and is easily palpable in most others. Comparison with the uninjured side is a useful aid for both visual examination and palpation. Any attempt at internal rotation is quite painful and resisted by the patient. An inability to place the palm from the injured extremity on the uninjured shoulder is consistent with an anterior shoulder dislocation; after reduction, this maneuver should be possible.

Careful assessment of the neurovascular status of the affected extremity is essential (Figs. 49.5 and 49.6). Injury to the axillary artery is rare. It usually occurs in the elderly and can be quickly assessed by a decreased or absent radial pulse or by the appearance of an expanding hematoma. It is important to evaluate the status of the axillary nerve because this is the most common nerve injury resulting from anterior dislocations. Assess the sensory component of the axillary nerve by testing for sensation over the lateral aspect of the upper part of the arm in the “regimental badge” area over the deltoid muscle. Testing the motor component of the axillary nerve is a difficult undertaking in a patient with a dislocated shoulder, as it requires activation of the deltoid muscle. Less commonly, the brachial plexus may be injured by a stretch injury and produce variable...
nerve deficits. Perform a complete assessment of all the major nerves to the arm because other nerve injuries may occur, such as injuries to the ulnar and radial nerves. A neurologic deficit does not preclude closed reduction, but in patients with a nerve injury, avoid multiple forceful attempts at reduction.

Brachial plexus injuries require an especially atraumatic reduction. If generous sedation and analgesia do not permit easy reduction in the ED, reduction of a dislocation with a nerve injury may be more prudently performed in the operating room with the patient under general anesthesia. Nerve injuries in this setting generally have a good prognosis, but the patient should be informed of the findings and the need for follow-up. The symptoms may require many months to resolve.

Vascular injuries, such as axillary artery disruption, are rare but usually quite obvious because of dysesthesias and coolness of the involved arm. An expanding axillary hematoma, pulse deficit, peripheral cyanosis, and pallor can be seen. Collateral circulation may produce a faint pulse in the extremity, so comparison with blood pressure on the uninjured side may be helpful. Specific lesions include complete disruption, linear tears, and thrombosis. Axillary artery injuries can occur at all ages, but they are more common in the elderly. The artery is particularly at risk with anterior dislocations, and dislocation with spontaneous reduction can produce the injury. Arteriography with surgical repair of the artery is required, occasionally with fasciotomy of the forearm if the ischemia is long-standing.

In many shoulder dislocations, some portion of the rotator cuff will be injured. Rotator cuff tears are easier to evaluate after reduction, when the pain and swelling have subsided.

**Radiologic Examination**

Associated fractures are detected in 15% to 35% of anterior shoulder dislocations, with fractures of the greater tuberosity being the most common. The presence of a fracture of the greater tuberosity does not change the initial management of anterior shoulder dislocations, and these fractures usually heal well after closed reduction in routine fashion. The Hill-Sachs deformity, a sign of repeated dislocations, produces a groove.

---

**Figure 49.4** A, Typical manifestation of an anterior right shoulder dislocation. The shoulder is very painful, and thus the patient resists movement. The outer round contour of the shoulder is obviously flattened, and the displaced humeral head may be appreciated in the subcoracoid area. Frequently, the patient abducts the arm slightly, bends the torso toward the injured side, and supports the flexed elbow on the injured side with the other hand. B, Obvious left shoulder dislocation. This chronic dislocation occurred frequently with minimal trauma, and the patient was able to dislocate it at will, feign a new injury, and obtain narcotics from multiple emergency departments.

**TABLE 49.1** Comparison of Anterior and Posterior Shoulder Dislocations: Classified According to Displacement of the Humeral Head

<table>
<thead>
<tr>
<th>TYPE OF DISLOCATION</th>
<th>FEATURES</th>
<th>OTHER CLINICAL CLUES</th>
<th>RADIOGRAPHS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anterior</td>
<td>Arm held in abduction and slight external rotation (abduction more prominent with subglenoid dislocation) The patient cannot adduct or internally rotate the shoulder</td>
<td>Seen from the front, the shoulder appears “squared off” Distal acromion prominent on a side view</td>
<td>On AP view: obvious dislocation On lateral or “Y” view: humeral head appears anterior to the glenoid fossa</td>
</tr>
<tr>
<td>Posterior</td>
<td>Arm held in the sling position with adduction and internal rotation Attempts at abduction and external rotation cause extreme pain</td>
<td>Coracoid process prominent, glenoid fossa empty anteriorly, and humeral head bulging posteriorly</td>
<td>On AP view: vacant glenoid sign, 6-mm sign, light bulb sign On lateral or “Y” view: humeral head appears posterior to the glenoid fossa</td>
</tr>
</tbody>
</table>

*AP,* Anteroposterior.
Management of Common Dislocations

offered by the patient) can be reduced without prior radiographs because fracture is quite unlikely in these situations. Hendey and coworkers performed a prospective validation study of an algorithm for selective radiography that incorporated the mechanism of injury, previous dislocations, and the clinician’s certainty of joint position. In this study, 24 patients with recurrent atraumatic anterior shoulder dislocations who received neither pre-reduction nor post-reduction radiographs had no clinically significant fractures found on follow-up. These patients had much shorter ED lengths of stay than did patients who received only pre-reduction or post-reduction films, or both.

One retrospective case-control study found that the presence of any of three risk factors (age > 40 years, first episode of dislocation, traumatic mechanism of injury defined as a fall greater than one flight of stairs, a fight or assault, or a motor vehicle collision) predicted clinically important fractures with a sensitivity of 97.7%. However, in a prospective study by Atef and colleagues, associated injuries were found in 60% of patients with anterior shoulder dislocation, a much higher frequency than previously thought; most commonly noted was a rotator cuff tear but the injuries ranged from axillary nerve injury to greater tuberosity fracture.

Anterior dislocations are not subtle on routine anteroposterior (AP) radiographs, and this view detects the most important fracture to identify, that of the humeral neck (Fig. 49.8). An adequate AP view, when combined with the typical clinical examination, allows successful detection of most anterior shoulder dislocations. A true AP view of the shoulder is taken at a right angle to the scapula and requires rotation of the patient to 30 to 45 degrees.

Fractures of the humeral neck are frequently displaced with attempts at closed reduction, which can lead to avascular necrosis of the humeral head. The Hill-Sachs deformity is caused by impaction of the humeral head against the glenoid rim after dislocation. It rarely has any clinical significance but may result in a loose body within the joint. Impaction of the humeral head against the glenoid during dislocation may also cause disruption of the anteroinferior portion of the cartilaginous labrum of the glenoid or the inferior aspect of the bony glenoid, an injury known as a Bankart lesion. It has been implicated as one source of recurrent dislocations but does not affect immediate ED management.

Fractures of the humeral neck are a known complication of shoulder relocation supports the value of pre-reduction radiographs of anterior shoulder dislocations. However, some argue that clinically obvious recurrent dislocations and first-time anterior dislocations without a blunt traumatic mechanism (information usually offered by the patient) can be reduced without prior radiographs because fracture is quite unlikely in these situations.

Hendey and coworkers performed a prospective validation study of an algorithm for selective radiography that incorporated the mechanism of injury, previous dislocations, and the clinician's certainty of joint position. In this study, 24 patients with recurrent atraumatic anterior shoulder dislocations who received neither pre-reduction nor post-reduction radiographs had no clinically significant fractures found on follow-up. These patients had much shorter ED lengths of stay than did patients who received only pre-reduction or post-reduction films, or both.

One retrospective case-control study found that the presence of any of three risk factors (age > 40 years, first episode of dislocation, traumatic mechanism of injury defined as a fall greater than one flight of stairs, a fight or assault, or a motor vehicle collision) predicted clinically important fractures with a sensitivity of 97.7%. However, in a prospective study by Atef and colleagues, associated injuries were found in 60% of patients with anterior shoulder dislocation, a much higher frequency than previously thought; most commonly noted was a rotator cuff tear but the injuries ranged from axillary nerve injury to greater tuberosity fracture.

Anterior dislocations are not subtle on routine anteroposterior (AP) radiographs, and this view detects the most important fracture to identify, that of the humeral neck (Fig. 49.8). An adequate AP view, when combined with the typical clinical examination, allows successful detection of most anterior shoulder dislocations. A true AP view of the shoulder is taken at a right angle to the scapula and requires rotation of the patient to 30 to 45 degrees.

The typical lateral views obtained include a scapular Y view (see Fig. 49.8), a transthoracic view, and an axillary view. These views rarely add to the AP film in patients with an obvious anterior dislocation, but they are of value in posterior dislocations (Fig. 49.9). The usefulness of additional views for anterior shoulder dislocations is primarily to detect fractures, and the previously mentioned lateral views (especially the transthoracic view) are quite limited in this respect. The apical oblique view has been found to be more valuable than the oblique scapular projection for acute shoulder trauma. This view is obtained by angling the beam 45 degrees caudad with the patient in a 45-degree oblique position (Fig. 49.10).
Post-reduction radiographs are obtained to document the success of the reduction. Occasionally, they will reveal a fracture not detected on pre-reduction radiographs. In one series, 8% of patients with anterior shoulder dislocations had Hill-Sachs deformities noted only on post-reduction films. A more recent prospective observational study (2007) found 37.5% of detected fractures associated with anterior shoulder dislocation were only seen on post-reduction radiographs; none however changed clinical management.

Recent advances in the use of point-of-care ultrasound by emergency physicians have resulted in its growing use for diagnosis and confirmation of successful reduction of shoulder dislocations and for administration of local anesthetic to facilitate closed reduction. A study by Abbasi and colleagues found that ultrasound had identical results as plain radiographs for both the detection of shoulder dislocation as well as complete reduction, with a sensitivity of 100%. Whereas further validation studies, as well as studies to determine the ability of ultrasound to recognize associated fractures, still need to be performed, we expect ultrasound to continue to become an ever-increasingly useful tool in the treatment of shoulder dislocations. One disadvantage of using only ultrasound diagnosis is difficulty in maintaining a permanent record of a successful reduction.

**Reduction Techniques**

Hippocrates (450 BC) is generally credited with the first detailed description of reduction techniques, and it is believed that a drawing in the tomb of Upuy (1200 BC) is the earliest depiction of such a method. The Hippocratic technique involves placement of the operator's foot in the axilla to allow countertraction. This technique is problematic and not recommended by some authors. Likewise, the Kocher method, which involves forceful leverage of the humerus, is associated with an increased rate of complications and is generally discouraged in favor of other techniques.

This section discusses several methods of reduction that are well studied, proven to be safe, and easy to master. Regardless of the reduction technique used, gradual, gentle application of the technique is essential. Although all the techniques discussed are generally acceptable and many authors state that their techniques are quite painless, few studies have quantified the actual pain reported by patients. In studies by Matthews and Roberts and Kosnick and colleagues, intraarticular injection of lidocaine was found to offer significant relief of pain during reduction of anterior shoulder dislocations. In addition, a 2014 meta-analysis showed that intraarticular lidocaine had similar success rates as procedural sedation and led to decreased length of stay and complication rates, thus making it a useful alternative to procedural sedation and analgesia.

A previous prospective study and Cochrane review reported similar results. Ultrasound may be used to assist intraarticular lidocaine injection in both adults and pediatric patients. When using intraarticular lidocaine, any blood present should be aspirated from the glenohumeral joint before injecting the anesthetic. Note that 10 to 20 mL of 1% lidocaine has been used with the intraarticular technique and that it may take as long as 15 to 20 minutes for adequate analgesia.

The use of regional nerve blocks (suprascapular and scalene) under ultrasound guidance have also been used to provide analgesia for closed reduction of anterior shoulder dislocations. Studies have demonstrated similar analgesia and reduction
success rates when compared to procedural sedation, but with significantly less one-on-one practitioner time and a lower ED length of stay.37–39 (see Chapter 31).

It is important to note that neither local nor regional anesthesia produces muscle relaxation, but they may obviate the need for IV access and prolonged observation. Operator judgment is an important part of the decision as to whether reduction should be attempted without premedication. Advantages of such an approach include avoidance of potential complications from drug therapy, reduced staff requirements, and theoretically, more rapid patient disposition. Certainly, a patient who is markedly intoxicated may require little, if any supplemental sedative therapy. However, all patients who are reluctant or too anxious to cooperate with an attempt at reduction without medication should receive premedication. Generally, only one attempt is made; if unsuccessful, further attempts at reduction are made after the IV administration of sedatives. When in doubt, it is best to use pharmacologic adjuncts (see Chapter 33).

Several factors will help in deciding which reduction technique is best in each clinical situation. Do not use forceful techniques such as traction-countertraction in patients who...
The Stimson maneuver is a classic technique that offers the advantage of not requiring an assistant. Place the patient prone on an elevated stretcher and suspend approximately 2.5 to 5.0 kg (5 to 10 lb) of weight from the wrist. The weights can be strapped to the wrist, or a commercially available Velcro wrist splint can be placed and the weights hung from the strap with a hook. The slow, steady traction produced with this method often permits reduction, but 20 to 30 minutes may be required. If needed, facilitate reduction by externally rotating the extended arm.

Variations of this method include the recommendation for flexion of the elbow to further relax the biceps tendon and the application of manual traction instead of weights. Rollinson are not being sedated. The clinician’s comfort level with a given technique is always a factor as the greatest success rates will probably result from techniques with which the clinician is most familiar. The time and resources available to the clinician must also be considered because methods such as the Stimson maneuver require more time and the availability of weights and straps. In addition, certain reduction techniques can be performed without assistance, whereas others require an additional person to apply countertraction or to help with manipulation of the scapula or humeral head. Ideally, the emergency clinician should become familiar with a number of different techniques for reducing anterior shoulder dislocations because no single method has a 100% success rate nor is any technique ideal in every situation.
allowed the arm to hang under its own weight after a supraclavicular block and reported a 91% success rate with usually no more than a gentle pull on the arm after 20 minutes in this position. Each variation of the Stimson method can be used in combination with the scapular manipulation technique described later. Indeed, a success rate of 96% has been reported with the combined prone position, hanging weights, IV analgesia and sedation, and scapular manipulation.40

Disadvantages of the Stimson method include the time required to achieve reduction and possible dangers to the patient associated with the positioning required for this technique. There is a potential danger of patients slipping off the elevated bed. A “seat belt” strap or bedsheet may be placed around the patient and stretcher to avoid movement of the patient off the stretcher. Airway access to a patient in the prone position may be more difficult in the setting of an overly intoxicated patient or one who becomes overly sedated. Logistically, the Stimson method may be difficult. One must have a bed that elevates to a suitable height for length of the patient’s arm and a convenient method to hang the weights. Sometimes locating the weights themselves can be challenging in a busy ED.

Scapular Manipulation Technique (See Fig. 49.12B)

This method is popular because of its ease of performance, reported safety, and acceptability to patients. To date, no complications from the scapular manipulation technique have been reported in the literature.29,40,44 Shoulder reduction via this method focuses on repositioning the glenoid fossa rather than the humeral head, and less force is required than with many other methods.36 Its success rate is high, generally greater than 90% in experienced hands.40,44,45 Some studies report higher success rates in patients who have had repeated dislocations and lower rates in patients with an associated greater tuberosity fracture.35 The initial maneuver for scapular manipulation is traction on the arm as it is held in 90 degrees of forward flexion. This may be performed with the patient prone and the arm hanging down, as described for the Stimson method, with or without flexion of the elbow to 90 degrees. Alternatively, this traction may be applied by placing an outstretched arm over the seated patient’s midclavicular region while pulling the injured extremity with the other arm. Regardless of the means of arm traction, slight external rotation of the humerus may facilitate reduction by releasing the superior glenohumeral ligament and presenting a favorable profile of the humeral head to the glenoid fossa.36

The prone patient position is recommended for those not familiar with the technique because it facilitates identification of the scapula for manipulation (medial rotation of the tip). Nonetheless, the technique can be performed with the patient supine given that the patient’s shoulder is flexed to 90 degrees and the scapula is exposed during gentle upward traction on the humerus.47 Although seated scapular manipulation offers the advantage of not requiring the patient to go through the awkward and potentially uncomfortable assumption of the prone position, it is a technically more difficult variation of scapular manipulation, especially if sedation is going to be necessary. When using the prone position, place the injured shoulder over the edge of the bed to allow the arm to hang perpendicularly for the application of traction (see Fig. 49.12A).48

After the application of traction, manipulate the scapula to complete the reduction. Anderson and associates44 recommended manipulation of the scapula after the patient’s arm is relaxed; however, success is possible with no delay in performing this second step.29 Manipulate the scapula by stabilizing the superior aspect of the scapula with one hand and pushing the inferior tip of the scapula medially toward the spine. Place the thumb of the hand stabilizing the superior aspect of the scapula along the lateral border of the scapula to assist the pressure applied by the thumb of the other hand. A small degree of dorsal displacement of the scapular tip is recommended as it is being pushed as far as possible in the medial direction.44

When the patient is properly positioned with the affected arm hanging perpendicularly, the lateral border of the scapula may be difficult to find in larger patients. This border is generally located quite laterally with the patient in this position, and it must be properly located before any attempt at reduction. The reduction itself is occasionally so subtle that it may be missed by both the patient and operator. A minor shift of the arm may be the only clue that reduction has been successful.

Figure 49.11  A, Intraarticular injection for the reduction of an acute anterior shoulder dislocation can be very effective. After aspirating blood from the joint, 10 to 20 mL of 1% plain lidocaine is injected slowly through the lateral sulcus. Allow 15 to 20 minutes for the lidocaine to take effect. B, The empty glenoid fossa is easily identified when considering an injection of lidocaine into the joint to facilitate reduction of a dislocated shoulder.
Careful palpation of the subclavicular area to locate the position of the humeral head before repositioning the patient may be used to determine the success of the reduction.

**Best of Both (BOB) Technique (See Fig. 49.12C)**

A variation of the seated scapular manipulation technique is the “best of both” (BOB) maneuver. To perform the BOB maneuver, position the patient seated sideways on the stretcher with the unaffected shoulder and hip against the fully elevated head of the stretcher. Stand on the foot end of the gurney at the patient’s affected side and use one hand to apply downward force on the proximal end of the patient’s bent forearm. Use the other hand to grasp the patient’s hand and gently rotate the arm internally or externally as needed. Once downward force is being applied, ask an assistant to perform the scapular manipulation maneuver described earlier.

**External Rotation Method (See Fig. 49.12D)**

This method offers the advantage of requiring only one person and no special equipment. The technique needs no strength or endurance on the part of the operator and is well tolerated by patients. The actual pain experienced by patients with this technique has not been quantified, but Plummer and Clinton stated that it can be performed with “little, if any sedation.” In this technique, the basic maneuver is slow, gentle external rotation of the fully adducted arm. In 1957, Parvin described a self-reduction external rotation technique in which the patient sits on a swivel-type chair and grasps a fixed post positioned waist high and slowly turns the body to enact external rotation. Parvin reported that the reduction usually takes place at 70 to 110 degrees of external rotation.

Since Parvin’s initial study, this method has been described with the patient supine and the affected arm adducted tightly to the patient’s side. Flex the elbow to 90 degrees and hold it in the adducted position with the operator’s hand closest to the patient. Use the other hand to hold the patient’s wrist and gently rotate the arm into slow and gentle external rotation. The procedure may require several minutes because each time the patient experiences pain, the procedure is halted momentarily. Although the report of Mirick and coworkers mentioned using the forearm as “a lever,” a later description clearly recommends allowing the forearm to “fall” under its own weight. No additional force should be applied to the forearm, and no traction is exerted on the arm.

The end point of the reduction may be difficult to identify because reduction is frequently very subtle. It is therefore recommended that external rotation be continued until the forearm is near the coronal plane (lying on the bed, perpendicular to the body), a process that usually takes 5 to 10 minutes. If the patient’s dislocation persists after full external rotation, apply steady gentle traction at the elbow. Reduction may occasionally be noted when the arm is rotated back internally. The success rate of this technique in three series performed by emergency clinicians was approximately 80%.

**Milch Technique (See Fig. 49.12E)**

Proponents of this method praise its gentle nature, high success rate, lack of complications, and tolerance by patients. It can be described as “reaching up to pull an apple from a tree.” The basic steps of this technique are abduction, external rotation, and gentle traction of the affected arm. Finally, if needed, the humeral head can be pushed into the glenoid fossa with the thumb or fingers.

In describing this technique, Milch wrote that the fully abducted arm was in a natural position in which there was little tension on the muscles of the shoulder girdle. He postulated that this was related to our ancestral “arboreal brachiation” (swinging from trees). The primary step in this technique is to abduct the affected arm to an overhead position. Russell and colleagues had their patients raise their arm and put their hand behind their head as a first step. Although this seems odd, patients can usually do this quite readily with little assistance and be quite comfortable in this position. Alternatively, abduct the arm by grasping the patient’s elbow or the wrist. Lacey and Crawford found that the prone position, with the patient’s shoulder close to the end of the bed, facilitated this step.

Once the arm is fully abducted, apply gentle longitudinal traction with slight external rotation. If reduction does not occur quickly, push the humeral head upward into the glenoid fossa with the thumb or fingers of the other hand. Beattie and associates reported a success rate of 70% with the Milch technique, but others have achieved success rates of 90% or greater. In a comparison study of external rotation and the Milch technique for reduction of anterior shoulder dislocations without sedation, success rates were equivalent but external rotation was noted to be easier and less painful for the patient.

**Traction-Countertraction (See Fig. 49.12F)**

This method is commonly used in the ED, largely because of tradition, clinician comfort, and a high success rate. Clinician familiarity is an advantage of this technique, but it requires more than one operator, some degree of force, and occasionally, endurance. This technique is usually quite uncomfortable for the patient, and premedication is recommended before any attempt. In a 2014 comparison study of traction-countertraction versus modified scapular manipulation, modified scapular manipulation was better tolerated by patients and had a better success rate on first attempt; traction-countertraction was more successful on second attempt.

With the patient supine, wrap a sheet or strap around the upper part of the patient’s chest and under the axilla of the affected shoulder. Ask an assistant to hold the sheet, preferably by wrapping it around the assistant’s waist to take advantage of body weight rather than arm strength to apply the countertraction. Traction may then be applied onto the extended arm by the clinician, but this generally results in operator fatigue, especially if the operator relies on biceps strength to provide continuous traction. Instead, flex the elbow of the affected side to 90 degrees and wrap a sheet or strap around the proximal part of the forearm and then around the operator’s waist. Elevate the bed to the point at which the sheet can sit at the level of the operator’s ischial tuberosities. This allows the operator to comfortably lean back and use body weight to supply the force of traction, thereby reducing the possibility of operator fatigue. The portion of the sheet that is positioned on the patient’s forearm has a tendency to ride up; flexion of the elbow beyond 90 degrees will minimize this problem. Alternatively, the operator merely leans backward with the arms fully extended, again using the continuous weight of the body rather than the strength of the biceps to provide constant traction.

Once traction has been applied, the operator must be patient because the procedure may take a number of minutes to be successful. The premedication is probably inadequate if the patient resists the procedure or is notably uncomfortable during
**Anterior Shoulder Dislocation Reduction**

**A. Stimson Maneuver**

Place the patient prone on the edge of the stretcher. Be careful that a sedated or intoxicated patient does not fall off the table. Belts or sheets can be used to secure the patient to the stretcher, 5-kg weights are attached to the arm, and the patient maintains this position for 20 to 30 minutes, if necessary.

The addition of scapular manipulation and/or gentle external and internal rotation of the shoulder with manual traction may aid in reduction.

**B. Scapular Manipulation**

Rotate the inferior tip of the scapula medially and dorsally toward the spine with the tips of your thumbs.

The procedure can take place with the patient prone (as in the Stimson technique) or with the patient seated. For the latter, have an assistant apply traction on the arm while applying countertraction on the ipsilateral clavicle.

**C. Best-of-Both Technique**

Position the patient seated sideways with the unaffected shoulder and hip against the upright head of the stretcher. Apply downward force on the patient’s flexed forearm, and gently rotate the arm internally or externally as needed.

Once downward force is applied, instruct an assistant to perform scapular manipulation as described previously.

Figure 49.12 Anterior shoulder dislocation reduction methods. *ED*, emergency department.

*Continued*
**Anterior Shoulder Dislocation Reduction**

**D. External Rotation**

Fully adduct the arm and flex the elbow to 90 degrees. Hold the patient’s wrist and guide the arm into slow and gently external rotation. Halt momentarily if the patient experiences pain. Continue the rotation until the forearm is laying on the bed. No traction is applied.

**E. Milch Technique**

First abduct the arm to an overhead position by grasping the patient’s arm at the elbow or wrist. Once fully abducted, apply gentle longitudinal traction with slight external rotation. If reduction does not occur quickly, push the humeral head upward into the glenoid fossa.

**F. Traction-Countertraction Method**

Wrap one sheet around the affected axilla and the assistant’s waist. The assistant leans back to apply countertraction. Wrap another sheet around the patient’s flexed arm and your waist. Lean back to apply traction.

Reduction can be facilitated by gently adducting the arm (after traction is applied) while a second assistant provides gentle lateral traction on the humerus.

**G. Spaso Technique**

With the patient supine, gently lift the arm toward the ceiling while applying gentle vertical traction. Instruct an assistant to apply countertraction. Apply gentle external rotation during the procedure.

**H. Eskimo Method**

While the patient lies on the unaffected side, lift him a short distance off the ground by grasping the abducted arm of the injured side. This is a field technique and should not be used in the ED.

Figure 49.12, cont’d
Attempts at reduction. Do not hesitate to order supplementary medications. Gentle, limited external rotation is sometimes useful to complete the reduction.\textsuperscript{12} Applying traction to an arm that is slightly abducted from the patient’s body is often successful, but some operators prefer to slowly bring the arm medial to the patient’s midline while maintaining traction or to have an assistant apply a gentle lateral force to the midhumerus to direct the humeral head laterally. Successful reduction is usually presaged by slight lengthening of the arm as relaxation occurs, and a noticeable “clunk” may occur at the point of reduction. A brief wave of fasciculations in the deltoid may also be seen at the time of reduction.

**Spaso Technique (See Fig. 49.12G)**

This technique was first reported by Spaso Miljesic as a simple, single-operator technique requiring minimal force.\textsuperscript{56} One published series reported an 87.5% success rate in premedicated patients when performed by junior house officers.\textsuperscript{37} A 2015 study comparing the Spaso technique to external rotation found both to be equally efficacious, however the Spaso technique was faster, more efficient, and more easily performed.\textsuperscript{39} Place the patient in the supine position and grasp the affected arm around the wrist or distal end of the forearm. Gently lift the affected arm vertically toward the ceiling and apply gentle vertical traction. While maintaining traction continuously, externally rotate the arm. Reduction may be subtle but is generally signaled by hearing or feeling a “clunk.” Completion of this technique may require several minutes of gentle traction to allow the muscles of the patient’s shoulder to relax.\textsuperscript{37}

**Other Methods**

Poulsen\textsuperscript{59} reported a method termed the Eskimo technique that may be performed in field settings (see Fig. 49.12H). In this technique the patient lies on the unaffected side and is lifted a short distance off the ground by grasping the abducted arm on the injured side. The patient’s body weight acts to effect the reduction. Poulsen’s success rate was 74% in a series of 23 patients, all of whom were premedicated. However, the author\textsuperscript{59} also postulated that this technique could place undue stress on the brachial plexus or axillary vessels. Use of this technique, when other options are available, should probably be reserved until more data are obtained.

Noordeen and coworkers\textsuperscript{60} reported a simple method in which the patient sits sideways in a chair with the affected arm draped over the backrest. The operator holds the arm with the wrist supinated, and the patient is instructed to stand up. The success rate was 72% in 32 patients treated in this manner. A variation of the chair technique, which was successful in 97% of 188 anterior shoulder dislocations, involves operator-applied traction on the patient’s flexed elbow by means of a cloth loop or stockinette.\textsuperscript{61} Standing beside the patient, the operator holds the involved elbow in 90 degrees of flexion while stepping down on the cloth loop. The patient sits in the chair, and an assistant may help support the patient by applying countertraction under the involved arm.

**Post-reduction Care**

After an attempt at reduction, the neurovascular status of the affected extremity should be rechecked and the results documented in the patient’s medical record. Indirect evidence that the reduction has been successful includes an immediate decrease in pain, restoration of the round shoulder contour, and increased passive mobility of the shoulder. No harm is done by putting the joint through a limited range of motion. If the patient can tolerate placement of the palm from the injured arm on the opposite shoulder, it is quite likely that the shoulder reduction was successful (Fig. 49.13A). For patients with possible axillary nerve injury, close to 90% will recover with expectant management. Nevertheless, it is prudent to refer these patients for early orthopedic follow-up.\textsuperscript{62}

Post-reduction radiographs are often recommended to make a careful search for new fractures. Although most greater tuberosity fractures do not alter patient management, patients with greater tuberosity fractures displaced more than 1 cm after closed reduction almost always have an associated rotator cuff tear\textsuperscript{63} and should receive prompt orthopedic consultation because they may require operative repair.

Traditional post-reduction treatment focuses on the importance of preventing the shoulder from dislocating again after the patient is discharged. This is best accomplished by immobilizing the joint with a commercially available shoulder immobilizer or a sling and swath to limit external rotation and abduction (see Chapter 50). Orthopedic follow-up is recommended for all anterior shoulder dislocations because the
incidence of rotator cuff injury is as high as 38% and it might complicate restoration of normal function. As a general rule, the older the patient, the shorter the recommended time of immobilization. Those older than 60 years should have early follow-up (e.g., 5 to 7 days) to allow early mobilization and avoid persistent shoulder joint stiffness or adhesive capsulitis. Younger patients are usually immobilized for approximately 3 weeks and can be instructed to follow up within 1 or 2 weeks of the event. As mentioned earlier, young age confers risk of recurrent dislocation, but it is not clear if prolonged immobilization decreases recurrence rates. A 2010 meta-analysis found that immobilization for more than 1 week did not reduce recurrence rates.

Since the early 2000s, the wisdom of immobilization in internal rotation has been questioned. Several studies have shown that placing the arm in internal rotation actually increases labral detachment from the glenoid rim, whereas some degree of external rotation maximizes contact between the detached labrum and the glenoid rim. In one study, cadavers were used to measure the force of contact between the labrum and the glenoid rim in different arm positions. The authors of this study found that maximum contact force was actually generated in 45 degrees of external rotation, whereas no contact force was generated with the arm in internal rotation. Unfortunately, the question of which position (e.g., internal versus external rotation) is superior for post-reduction immobilization remains unanswered. Clinical studies thus far have shown conflicting results. Nevertheless, in the most recent systemic review and meta-analysis comparing recurrence rates and patient-based quality of life assessments between external and internal rotation immobilization (2014), Liu and colleagues found that immobilization in external rotation did not reduce recurrence rates or improve quality of life. As a result, it is not unreasonable to immobilize the extremity in a manner consistent with the recommendations of the orthopedic surgeons at one’s institution until further evidence is presented. When in doubt, a simple sling or the traditional shoulder immobilizer will certainly suffice pending 5- to 7-day follow-up (see Fig. 49.13B).

Discharge the patient with oral analgesics (either nonsteroidal antiinflammatory drugs or narcotics) to minimize discomfort and instruct them to return if the clinical condition worsens. Periodically, one may encounter a return visit from a successfully treated patient who is in severe pain from hemarthrosis. In one series of patients older than 60 years, Trimmings reported excellent pain relief by aspirating the hemarthrosis 24 to 48 hours after shoulder reduction. This can be accomplished by using the technique of arthrocentesis described in Chapter 53. In addition, intraarticular instillation of 10 to 20 mL of 1% lidocaine (or a longer-acting local anesthetic) may be helpful for further pain relief.

**Posterior Shoulder Dislocations**

Posterior shoulder dislocations account for less than 4% of all shoulder dislocations. Because they are so uncommon, posterior dislocations have the potential to be missed. In fact, delays in diagnosis for weeks to months have been reported. This may lead to increased rates of dislocation arthropathy and chronic pain. The mechanism of injury is almost always indirect and consists of a combination of internal rotation, adduction, and flexion. Classic precipitating events include seizure, electrical shock, and falls. Some patients may not be seen until a point well past the original event. In addition, patients with seizures may not complain of pain or a limited range of motion in the immediate postictal period because of their altered mental status.

**Clinical Assessment**

Though clinically less obvious than anterior dislocations, posterior shoulder dislocations do occur in a typical, recognizable manner. Mistakes may be made if the clinician is overly reliant on AP radiographs, which are potentially misleading and may result in misdiagnosing the injury as a soft tissue contusion or acromioclavicular (AC) strain. The principal sign of a posterior dislocation is an arm that is somewhat fixed in adduction and internal rotation and the patient’s inability to rotate the shoulder externally with the elbow flexed at 90 degrees.

**Radiologic Examination**

The key point regarding radiographs for posterior shoulder dislocations is the subtle nature of this dislocation on a single AP film (Figs. 49.15A and 49.16A) and the diagnostic importance of the scapular Y view (see Fig. 49.15B) or the axillary view (see Fig. 49.16B). Diagnosis of posterior shoulder dislocation is quite easy with the axillary view, whereas the routine AP and lateral views are difficult to interpret in approximately half the cases. The axillary view is generally available in the radiology department and can be obtained with as little as 20 to 30 degrees of abduction and the plate placed on the shoulder. In addition to easy visualization of the posteriorly situated humeral head, the axillary view often reveals an impression fracture of the humeral head (see Fig. 49.16B). The scapular Y view is produced by superimposing the head of the humerus over the coracoid, acromion, and body of the scapula, which form a Y shape. In the event of a posterior shoulder dislocation,
shaped appearance created by the greater tuberosity (Fig. 49.17). With posterior dislocation, the space between the articular surface of the humeral head and the anterior glenoid rim is widened, and there is a decrease in the half-moon–shaped overlap of the head and the fossa. There may also be a compression fracture on the medial aspect of the humeral head of the humerus will lie posterior to the glenoid (away from the chest wall) (see Figs. 49.9B and 49.15B).

Even though axillary and scapular Y views are diagnostic, clues to posterior dislocation do exist on AP films. The internally rotated humeral head appears symmetric on an AP film and is in the shape of a light bulb, as opposed to the normal club-shaped appearance created by the greater tuberosity (Fig. 49.17). With posterior dislocation, the space between the articular surface of the humeral head and the anterior glenoid rim is widened, and there is a decrease in the half-moon–shaped overlap of the head and the fossa. There may also be a compression fracture on the medial aspect of the humeral head of the humerus will lie posterior to the glenoid (away from the chest wall) (see Figs. 49.9B and 49.15B).

Even though axillary and scapular Y views are diagnostic, clues to posterior dislocation do exist on AP films. The internally rotated humeral head appears symmetric on an AP film and is in the shape of a light bulb, as opposed to the normal club-shaped appearance created by the greater tuberosity (Fig. 49.17). With posterior dislocation, the space between the articular surface of the humeral head and the anterior glenoid rim is widened, and there is a decrease in the half-moon–shaped overlap of the head and the fossa. There may also be a compression fracture on the medial aspect of the humeral head of the humerus will lie posterior to the glenoid (away from the chest wall) (see Figs. 49.9B and 49.15B).

Even though axillary and scapular Y views are diagnostic, clues to posterior dislocation do exist on AP films. The internally rotated humeral head appears symmetric on an AP film and is in the shape of a light bulb, as opposed to the normal club-shaped appearance created by the greater tuberosity (Fig. 49.17). With posterior dislocation, the space between the articular surface of the humeral head and the anterior glenoid rim is widened, and there is a decrease in the half-moon–shaped overlap of the head and the fossa. There may also be a compression fracture on the medial aspect of the humeral head of the humerus will lie posterior to the glenoid (away from the chest wall) (see Figs. 49.9B and 49.15B).
Reduction Technique (Fig. 49.18A)
Reduce an acute posterior dislocation by applying traction on the internally rotated and adducted arm combined with anteriorly directed pressure on the posterior aspect of the humeral head.12,79 Generous premedication is usually indicated, and countertraction may be applied with a sheet looped in the affected axilla, similar to the procedure described for anterior dislocations. Kwon and Zuckerman12 recommended applying head, as indicated by a dense line. This is known as the trough sign.80 A fracture of the lesser tuberosity should always prompt a search for the presence of a posterior shoulder dislocation.

As mentioned earlier in this section, ultrasound is being used increasingly for the diagnosis of shoulder dislocations, including posterior shoulder dislocations and may be useful as an adjunct to the physical exam when dislocation is clinically suspected but not radiographically apparent.22,25,26

**Figure 49.17** Anteroposterior views comparing posterior dislocation and a normal shoulder joint. Posterior shoulder dislocation causes internal rotation of the humeral head, which makes the head appear as a light bulb rather than its normal club-shaped appearance. In addition, note that the space between the articular surface of the humeral head and the anterior glenoid rim is widened (arrows), and there is a decrease in the overlap between the head and the fossa.

**Figure 49.18** Posterior and inferior shoulder dislocation reduction methods.

**Posterior and Inferior Shoulder Dislocation Reduction**

**A. Posterior Dislocation**

- Apply traction, internal rotation, and adduction to the affected arm.
- Instruct one assistant to apply countertraction (with a sheet wrapped around the waist) and another assistant to apply anteriorly directed pressure on the posterior aspect of the humeral head.

**B. Inferior Dislocation (Luxatio Erecta)**

- Apply overhead traction with the arm in abduction. With your free hand, exert cephalad pressure over the humeral head. Instruct an assistant to apply countertraction toward the patient’s feet with a sheet placed over the injured shoulder.

**Figure 49.18** Posterior and inferior shoulder dislocation reduction methods.
lateral traction on the upper part of the humerus if the humeral head is locked on the posterior glenoid. Hawkins and colleagues suggested that posterior dislocations with an impression defect of the humeral head greater than 20% of the articular surface require open reduction. Posterior dislocations that have been diagnosed late are difficult to reduce in a closed manner, but an attempt with adequate premedication is generally indicated.

Post-reduction Care
As with anterior dislocations, neurovascular examination and radiographs should be repeated after attempts at reduction. Successful reduction is suggested by a patient’s ability to place the palm of the injured arm on the opposite shoulder. Given the rarity of these injuries, orthopedic consultation is often sought early in the care of these patients. In a training environment, involvement of orthopedic residents is of benefit to their education and should be considered early. An analysis and review of the literature of posterior dislocations suggests the majority (65%) of posterior shoulder dislocations will have an associated injury (fracture, reverse Hill-Sachs injury, or rotator cuff tear) underscoring the importance of orthopedic consultation.

Unusual Shoulder Dislocations
Luxatio Erecta
Inferior dislocations of the shoulder, known as luxatio erecta, are quite rare but also quite obvious. The patient has the arm locked in marked abduction with the flexed forearm lying on or behind the head. Occasionally, the humerus may have less abduction, thus potentially obscuring the diagnosis. The humeral head can be palpated along the lateral chest wall. With this injury, the inferior capsule is almost always torn. Associated injuries include fractures of the greater tuberosity, acromion, clavicle, coracoid process, and glenoid rim. Neurovascular compression may be present, but this is usually reversed once reduction is accomplished. Long-term complications include adhesive capsulitis and recurrent dislocations.

To reduce an inferior shoulder dislocation, apply overhead traction (generally with the arm in full abduction) in the longitudinal direction of the arm, and exert cephalad pressure over the humeral head much as with the Milch technique (see Fig. 49.18B). If needed, apply countertraction toward the patient’s feet by using a sheet placed over the injured shoulder. After reduction, bring the abducted arm into adduction against the body and supinate the forearm.

Alternatively, use the “two-step” maneuver described by Nho and associates, in which the luxatio erecta is first converted to an anterior dislocation. To perform this maneuver, place one hand on the medial condyle of the elbow and the other hand around the shaft of the humerus. Push anteriorly on the shaft of the humerus while stabilizing the medial condyle of the elbow, and rotate the humeral head from an inferior to an anterior position. The authors then describe using the external rotation method to reduce what is now a typical anterior dislocation.

Scapular dislocation or “locked scapula” is a rare condition characterized by obvious protrusion of the lateral border of the scapula and significant swelling of the medial border because of tearing of the musculature. To reduce the scapula, apply traction on the abducted arm and apply medial pressure on the scapula.

AC Joint Subluxation and Dislocation
The AC joint is a true diarthrodial joint that consists of a synovial cavity surrounded by a relatively lax capsule and the weak AC ligament. This structure allows the gliding motion necessary for shoulder movement. The major stability of the AC joint comes from two ligaments. The AC ligament is primarily responsible for joint stability in the AP direction. The coracoclavicular ligament, which has posterior (conoid) and anterior (trapezoid) components, anchors the distal end of the clavicle to the coracoid process of the scapula and therefore supports the joint in a superior-inferior direction. In general, AC injuries arise from a direct force such as a fall on the point of the shoulder with the arm adducted. AC joint injuries are categorized according to the Rockwood classification (types I to VI) (Fig. 49.20).

First Degree (Type I)
This injury consists of a minor tear in the AC ligament. The coracoclavicular ligament is intact. The clinical findings are
limited to tenderness in the area of the AC joint. Radiographs show little, if any change in position of the clavicle in relation to the acromion. Management of this condition consists of a sling for comfort, ice, and mild analgesics. Generally, the symptoms subside with 7 to 10 days of rest. Orthopedic referral is not usually necessary unless return to normal function is delayed beyond 2 weeks.

**Second Degree (Type II)**

In addition to a complete tear of the AC ligament, the coracoclavicular ligament is stretched or incompletely torn. The patient generally supports the injured arm and has slight swelling and definite tenderness over the AC joint. Radiographs demonstrate a definite change in the relationship of the distal end of the clavicle to the acromion. However, in type II injuries the inferior edge of the clavicle should not be separated from the acromion by more than half its diameter, and the coracoclavicular distance is the same as that on the uninjured side. This injury can be treated in closed fashion with a sling. Orthopedic referral is recommended, and some still recommend a sling-strap device that elevates the arm and depresses the clavicle for these injuries.

**Third Degree (Type III)**

In this injury, the distal end of the clavicle is essentially free floating because both the AC and coracoclavicular ligaments are completely disrupted. The arm is supported by an uncomfortable-appearing patient, and the distal end of the clavicle is usually seen to be riding high above the acromion. The diagnosis is generally obvious, and radiographs are used mainly to rule out an associated fracture. Radiographic criteria for this degree of injury include the inferior border of the clavicle raised above the acromion or a discrepancy in the coracoclavicular distance between the normal and affected sides. These injuries require orthopedic referral, and a fair bit of controversy exists regarding their subsequent management. Larsen and coworkers conducted a prospective, randomized trial of conservative versus operative management of significant AC separations and concluded that conservative management was generally better, with possible exceptions made for patients with significant cosmetic deformity and for those who frequently keep their arm at 90 degrees of abduction. Even though optimal therapy is still unclear, a logical approach includes ED treatment with a sling and early orthopedic referral.

**Fourth, Fifth, and Sixth Degrees (Types IV to VI)**

In type IV injury, the distal end of the clavicle is free floating and posteriorly displaced into or through the trapezius muscle. Type V injury is characterized by inferior displacement of the scapula with a marked increase (two to three times normal) in the coracoclavicular interspace. Type IV and V dislocations generally require surgery, and orthopedic referral is necessary. Type VI injury involves dislocation of the distal end of the clavicle inferiorly. Because this is usually the result of major trauma, other fractures are often present and should be sought.

**Radiographic Examination**

The diagnosis of AC dislocation is usually made clinically by noting pain and local tenderness at the AC joint in the absence of other findings. Radiographs are generally indicated to rule out associated fractures and to aid in assessing the degree of injury. A single radiograph of the injured shoulder often suffices,
but some clinicians prefer to obtain views of the opposite shoulder for comparison. Although their efficacy has never been proved, it has traditionally been recommended that “weighted” films be obtained for suspected type I or II injuries. Weighted films are generally performed after routine “unweighted” radiographs and are obtained by strapping approximately 4.5 to 7.0 kg (10 to 15 lb) of weight to the patient’s wrists and repeating the radiographs. Weighted films are of questionable value in mild injuries and superfluous in obvious type III to VI injuries. In a prospective study of 70 type I or II injuries, the use of weights was associated with less evident separation in 7 cases, essentially producing a false-negative study in comparison to plain unweighted films. Only three injuries were re-categorized as type III after the performance of weighted films. The authors of this study recommend abandoning the use of weighted films in patients with AC dislocation.

**STERNOCLAVICULAR DISLOCATIONS**

Despite the fact that the sternoclavicular joint is the least stable joint in the body, sternoclavicular dislocations are rare. The primary supports of this joint are the sternoclavicular and costoclavicular ligaments. Anterior dislocations are much more common and usually the result of an indirect mechanism involving a blow thrusting the shoulder forward, or they may be atraumatic, caused by ligamentous laxity in teens and young adults. Posterior dislocations also usually result from a blow to the shoulder but can also be caused by a direct superior sternal or medial clavicular blow. Athletic injuries and those resulting from motor vehicle accidents account for the vast majority of sternoclavicular dislocations. Posterior sternoclavicular dislocation (also known as retrosternal dislocation because the medial end of the clavicle dislocates both posteriorly and medially) is potentially life-threatening because injury to the great vessels or compression of the airway might occur. Patients may complain of dyspnea, choking, or hoarseness with tracheal compression; dysphagia with esophageal compression; ipsilateral upper extremity pain and swelling with subclavian vessel occlusion; or paresthesias if the brachial plexus is compromised. Any suggestion of these complications should prompt immediate surgical consultation.

The clinical manifestation of these injuries is usually straightforward and consists of pain, swelling, tenderness, and deformity of the joint. Patients may complain of pain that is worse with arm movement and when they are supine. Plain radiographs of this joint are difficult to interpret and generally include an apical lordotic-type view with the radiographic tube angled 45 degrees cephalad. Confirmation of the diagnosis is best made with a thoracic computed tomography (CT) scan, which may also identify high rib fractures, pulmonary contusion, or pneumothorax (Fig. 49.22). CT angiography should be obtained to identify associated vascular injury when indicated. Children may have epiphyseal disruption with retrosternal displacement of the medial aspect of the clavicle.

To reduce both types of sternoclavicular dislocation, place a rolled blanket or a sandbag between the scapula and spine to separate the medial aspect of the clavicle from the manubrium. Apply traction on the 90-degree abducted, 10-degree extended arm in line with the clavicle and then push (anterior dislocation) or lift (posterior dislocation) the clavicle back into position. Posterior dislocations may be difficult to reduce and to maintain via closed reduction. Therefore some authors recommend reduction in an operating suite unless complications necessitate immediate reduction. Given the rarity of this injury and the potential for major underlying complications, early consultation is recommended for suspected posterior sternoclavicular dislocations. Once reduced, a clavicle strap may be used to immobilize both anterior and posterior dislocations for up to 6 weeks.

**ELBOW DISLOCATIONS**

The elbow is second only to the shoulder as a site of major joint dislocations in adults; it is the most commonly dislocated joint in children. Anatomically, the principal articulation of
the humerus and ulna is a stable hinge joint with the intercondylar groove of the distal end of the humerus nestled in the olecranon fossa. Because of the stability of the elbow, any dislocation is expected to be accompanied by considerable soft tissue damage. Associated fractures of the radial head and coronoid process of the ulna are common. Elbow dislocations are usually divided into posterior and anterior dislocations (Fig. 49.23). However, there are actually several additional types of elbow dislocations, including lateral, divergent, and isolated dislocations of the radius.95 In the rare divergent dislocations, the radius and ulna are dislocated in opposite directions, either anterior and posterior or medial and lateral.95 Studies have shown that many patients report long-term problems including residual pain and joint stiffness after elbow dislocation.96,97 However, the most serious complication of elbow dislocation is injury to the brachial artery. This injury is possible with any type of elbow dislocation and is a frequent occurrence with open dislocations.95 Vascular compromise can be delayed in onset and result from either unsuspected arterial injury or progressive soft tissue swelling. The circulatory status of the arm must be carefully monitored even after successful reduction.95 In most cases, orthopedic consultation should be sought before disposition. Patients with any variety of elbow dislocation who have significant or immediate soft tissue swelling or hematoma formation or those who have questionable vascular integrity or neurologic findings are often admitted to the hospital or ED observation unit.

Injury to the median and ulnar nerves may be the result of stretch, severance, or entrapment. It is difficult to clinically distinguish these causes; therefore management of nerve injuries is frequently expectant.95 It is imperative to conduct a careful neurologic examination before and after reduction because any increase in findings may indicate entrapment and the need for surgical intervention.95 Myositis ossificans is also a potential complication of elbow dislocations secondary to hemarthrosis, which underscores the advisability of orthopedic consultation early in the course of care.

**Posterior Dislocations**

Posterior dislocations make up the vast majority of elbow dislocations.61 The usual mechanism is a fall on an outstretched hand with the elbow in extension. Findings on clinical examination are usually diagnostic unless severe soft tissue swelling is present. The patient has a shortened forearm that is held in flexion, and the olecranon is prominent posteriorly. The normally tight triangular relationship of the olecranon and the epicondyles of the distal end of the humerus is disturbed in a posterior dislocation. A defect may also be palpated above the prominence of the olecranon.

**Radiologic Examination**

Two radiographic views, an AP and a true lateral, should be obtained (Fig. 49.24). The diagnosis is obvious with proper radiographs. A careful search for fractures of the distal end of the humerus, radial head, and coronoid process must be undertaken because they commonly occur in this injury.95 In children younger than 14 years, the fracture is usually a separation of the medial epicondyle because the epiphyseal plate gives way before the medial collateral ligament of the elbow. Post-reduction radiographs are also necessary to confirm reduction and disclose any associated fractures.98

**Reduction Techniques and Post-Reduction Care**

As with shoulder reductions, some authors claim that their method of elbow reduction is virtually painless,49,99 but this has not been objectively documented. In general, patients with posterior elbow dislocations are quite uncomfortable, and it is beneficial to administer IV analgesics early in the course of care, preferably before positioning for radiographs. In addition to or in lieu of parenteral sedation and analgesia, some clinicians inject the elbow joint with a local anesthetic (e.g., 3 to 5 mL of 2% plain lidocaine) before attempting reduction. Before injection, aspirate the joint to remove any blood.

**Recommended Initial Approach**

A prone technique is advantageous because patients tolerate this position quite well. Hang the flexed elbow over the edge of the bed and position an assistant with his or her back toward the patient such that the patient’s humerus can be encircled...
with both hands and pressure applied with the thumbs on the posterior aspect of the olecranon (Fig. 49.25A). This pressure on the olecranon is intended to lift it up and away from the humerus. Apply longitudinal traction on the arm with the elbow in slight flexion. If reduction is not accomplished, an attempt may be made to further flex the elbow, or the assistant can be instructed to lift the humerus. Reduction is generally noted by a definite “clunk.”

Traditional Traction Method
Place the patient in the supine position and have an assistant stabilize the humerus with both hands (see Fig. 49.25B). Grasp the wrist and apply slow and steady in-line traction. Slightly flex the elbow to keep the triceps mechanism loose, and hold the wrist supinated while applying traction. Reduction is usually signified by a “clunk” that is heard or felt. If this method is not successful after a reasonable period of traction (10 minutes), gently flex the forearm to effect reduction. Alternatively, apply downward pressure on the proximal volar surface of the forearm to free up the coronoid process.

Alternatives
Several authors have described variations of a prone method of reduction that are reportedly well tolerated by patients. In the method described by Minford and Beattie, position the patient with the arm hanging over the padded back of a chair or over the edge of the bed (see Fig. 49.25C). Apply pressure to the prominent posterior aspect of the olecranon to achieve reduction. Alternatively, apply traction with the elbow flexed over the edge of a chair. Pull down on the hand while using the thumb to guide the olecranon into place.

Figure 49.25 Posterior elbow dislocation reduction methods.
Anterior Elbow Dislocation Reduction

Apply in-line traction and backward pressure on the proximal end of the forearm while an assistant provides countertraction on the humerus. A clunk usually indicates that reduction is achieved.

Flex the arm beyond 90 degrees to ensure that the joint has been reduced. Splint the elbow in at least 90 degrees of flexion with a posterior long-arm splint.

Figure 49.26 Anterior elbow dislocation reduction method.

Post-Reduction Care
Once reduction is achieved, put the elbow through a gentle range of motion to ensure that the reduction is stable and there is no mechanical block to movement. An inability to move the elbow through a smooth range of motion after reduction is often caused by an entrapped medial epicondyle fracture fragment, which requires operative intervention. Immobilize the elbow in at least 90 degrees of flexion with the forearm in slight pronation by using a long arm posterior splint (see Chapter 50). After immobilizing the joint, recheck the neurovascular status of the extremity and obtain post-reduction radiographs.

After reduction, any signs of delayed vascular compromise are first addressed by loosening the splint and decreasing the degree of flexion. This may restore adequate blood flow. If not, immediate surgical consultation is necessary for emergency arteriography, exploration of the brachial artery, or both. Brachial arterial injury may not be immediately apparent because of the presence of collateral circulation. The risk for vascular compromise is a reason to consider in-hospital observation. Alternatively, observe the patient in the ED or ED observation unit for 2 to 3 hours after reduction to evaluate for delayed neurovascular compromise before discharge.

The optimal duration of immobilization is unknown. However, a recent randomized multicenter trial (2017) found early range of motion to be safe and associated with faster recovery time and return to work when compared to plaster immobilization for 3 weeks.

Anterior Dislocations
Anterior dislocations of the elbow are quite rare; they usually result from a direct posterior blow to the olecranon with the elbow flexed. On physical examination, the arm is elongated and extended with anterior tenting of the proximal end of the forearm and prominence of the distal end of the humerus posteriorly. These injuries are the result of a great deal of force; they are frequently open and accompanied by significant neurovascular injury. An avulsion of the triceps mechanism may also occur.

To reduce anterior dislocations of the elbow, apply in-line traction and backward pressure on the proximal end of the forearm (Fig. 49.26). An assistant provides countertraction by grasping the humerus with both hands. Given the infrequent nature of anterior dislocations and the high probability of a severe associated injury, the emergency clinician should consider early orthopedic consultation.

Radial Head Subluxation (Nursemaid’s Elbow)
Radial head subluxation is a common pediatric orthopedic issue that generally occurs between the ages of 1 and 3 years. The mean age is just older than 2 years, but this entity has been reported in infants younger than 6 months and in older children up to the preteen years. There is a slight predilection for this injury to occur in girls and in the left arm. The classic mechanism of injury is longitudinal traction on the arm with the wrist in pronation, as occurs when the child is lifted up by the wrist (Fig. 49.27A). There is no support for the common assumption that the relatively small head of the radius in relation to the neck of the radius predisposes the young to this injury. The pathologic lesion is generally a tear in the attachment of the annular ligament to the periosteum of the radial neck, with the detached portion becoming trapped between the head of the radius and the capitellum (see Fig. 49.27B).

Clinical Assessment
The history offered by the caretaker may not be that of the classic pulling-type mechanism. Schunk, in a series of 83 patients, reported that only 51% described such a mechanism...
In a matter of minutes. Once the subluxation is reduced, full activity is generally regained. The affected arm hangs down at the side, slightly flexed and pronated. Minimal discomfort as long as the elbow is not manipulated.

This child will not use the injured right arm but has the radial head (nursemaid’s elbow). It may be difficult to determine exactly where the pathology exists, and the wrist is often thought to be the area of injury. This child will not use the injured right arm but has minimal discomfort as long as the elbow is not manipulated. The affected arm hangs down at the side, slightly flexed and pronated. Once the subluxation is reduced, full activity is generally regained in a matter of minutes.

Radiographs are not generally needed with the classic findings of a child with an arm in the nursemaid’s position that is not tender (or minimally tender in the radial head area) on palpation and an appropriate history. In these cases, findings on radiographs are generally normal, and if obtained, the position of the child’s arm by the x-ray technician often effects reduction. Nevertheless, Frumkin described three cases of nursemaid’s elbow in which a line drawn through the longitudinal axis of the radius did not normally bisect the capitellum on pre-reduction radiographs but did so after reduction. Radiographs are sometimes recommended if the child is not moving the arm normally 15 minutes after reduction. However, this time frame may be too short because reuse can be delayed for more than 30 minutes, particularly in children initially seen some time after the injury. Quan and Marcuse recommended an approach in which no radiographs are obtained on the first visit, including in children released from the ED before regaining full use of the arm. At the time of a 24-hour follow-up visit, radiographs are obtained only if repeat attempts at manipulation are not successful.

Even though this condition does not generally require radiographs, they can be valuable if external signs of trauma are present (e.g., swelling, abrasions, ecchymoses) or if the child does not use the arm normally within 24 hours after the subluxation is thought to be reduced. Other less common conditions that can have similar findings are fractures, joint infections, tumors, and osteomyelitis.

Ultrasound is an additional tool that may be used to help exclude an alternate etiology of elbow pain or confirm the presence of a suspected radial head subluxation. A study performed by Rabiner and colleagues evaluated the use of ultrasound to exclude a posterior fat pad, a finding associated with supracondylar fracture, in patients with suspected radial head subluxation. All patients with normal elbow ultrasound had successful reduction of the radial head subluxation with return to full function. Ultrasound may also be diagnostic in radial head subluxation. The “hook sign,” caused by the hyperechoic J-shaped supinator muscle above the radial head, has been shown to be 100% sensitive for the diagnosis of radial head subluxation. As with other conditions, ultrasound can be a useful radiation-sparing adjunct to the physical exam and may additionally assist in reassuring an anxious caregiver of the diagnosis prior to reduction attempts.

Although tenderness about the elbow has been reported occasionally, there is often little tenderness or swelling in the elbow region. In a cooperative child, the arm and shoulder are carefully palpated to discern any tenderness. Areas of focus on palpation should include the clavicle and distal end of the radius because these are common sites for pediatric fractures. When patient anxiety interferes with reliable assessment of tenderness in a child whose arm is in the classic nursemaid’s position, the examiner can stand at a distance and have the parent or caretaker palpate the extremity to ascertain tenderness. This may also be done with a cooperative patient to reassure a doubtful parent regarding the absence of a fracture. If no tenderness is noted on palpation, it is appropriate to attempt reduction without prior radiographs.

Although resistance to or pain with supination is a frequent finding in patients with radial head subluxation, one need not test for this finding until the time of reduction.

**Radiographic Examination**

**Radial head subluxation (nursemaid’s elbow).** A, The classic mechanism for this injury is longitudinal traction on the arm with the wrist in pronation. B, The pathologic lesion is usually a torn piece of the annular ligament becoming trapped between the radial head and the capitellum of the humerus. (From Fleisher GR, Ludwig S, editors: Textbook of pediatric emergency medicine. Baltimore, 1988, Williams & Wilkins p 1322. Reproduced by permission.)

**Figure 49.27**

**Figure 49.28** Typical findings in a child with subluxation of the right radial head (nursemaid’s elbow). It may be difficult to determine exactly where the pathology exists, and the wrist is often thought to be the area of injury. This child will not use the injured right arm but has minimal discomfort as long as the elbow is not manipulated. A, The affected arm hangs down at the side, slightly flexed and pronated. B, Once the subluxation is reduced, full activity is generally regained in a matter of minutes.

Whereas 22% reported a fall. In patients younger than 6 months, the mechanism in the majority of cases is simply rolling over in bed. Therefore isolated radial head subluxation in children younger than 6 months does not automatically necessitate a child abuse investigation. The typical patient with radial head subluxation is in no distress and holds the involved arm slightly flexed and the wrist pronated at the side (Fig. 49.28). This has been termed the nursemaid’s position. The exact area of pain is often difficult to locate. The child will refuse to use the arm, and this may be the chief complaint. An older child will usually point to the dorsal aspect of the distal end of the forearm when asked where it hurts, which may mislead the clinician to suspect a distal radial buckle fracture.

Although fractures involving the distal radius are commonly referred to as radial buckle fractures, there is often minimal or no obvious deformity. A classic mechanism for this injury is longitudinal traction on the arm with the wrist in pronation. Frumkin described three cases of nursemaid’s elbow in which a line drawn through the longitudinal axis of the radius did not normally bisect the capitellum on pre-reduction radiographs but did so after reduction. Radiographs are sometimes recommended if the child is not moving the arm normally 15 minutes after reduction. However, this time frame may be too short because reuse can be delayed for more than 30 minutes, particularly in children initially seen some time after the injury. Quan and Marcuse recommended an approach in which no radiographs are obtained on the first visit, including in children released from the ED before regaining full use of the arm. At the time of a 24-hour follow-up visit, radiographs are obtained only if repeat attempts at manipulation are not successful.

Even though this condition does not generally require radiographs, they can be valuable if external signs of trauma are present (e.g., swelling, abrasions, ecchymoses) or if the child does not use the arm normally within 24 hours after the subluxation is thought to be reduced. Other less common conditions that can have similar findings are fractures, joint infections, tumors, and osteomyelitis.

Ultrasound is an additional tool that may be used to help exclude an alternate etiology of elbow pain or confirm the presence of a suspected radial head subluxation. A study performed by Rabiner and colleagues evaluated the use of ultrasound to exclude a posterior fat pad, a finding associated with supracondylar fracture, in patients with suspected radial head subluxation. All patients with normal elbow ultrasound had successful reduction of the radial head subluxation with return to full function. Ultrasound may also be diagnostic in radial head subluxation. The “hook sign,” caused by the hyperechoic J-shaped supinator muscle above the radial head, has been shown to be 100% sensitive for the diagnosis of radial head subluxation. As with other conditions, ultrasound can be a useful radiation-sparing adjunct to the physical exam and may additionally assist in reassuring an anxious caregiver of the diagnosis prior to reduction attempts.
Multiple studies have found a variable combination of improved success and less pain with this method. Similarly, a 2012 Cochrane review concluded hyperpronation to be more successful and less painful than the supination method.

**After Attempted Reduction**

If a click is detected, the child will generally regain use of the affected arm quickly (almost always by 30 minutes). Therefore, if a definite click is detected, it is reasonable to observe the child for up to 30 minutes before further intervention. If there is still no use at 30 minutes, the operator may try to determine whether supination is still painful, which would suggest the need to repeat the attempt. In children in whom a click is not detected, the majority will not use the arm by 30 minutes. Therefore a repeated attempt at reduction is recommended after 10 to 15 minutes of nonuse in children who did not have a click during the primary reduction. Two or more attempts are required to produce the click in up to 30% of patients. If the child has not regained use of the arm after a few attempts and a reasonable period of time, some authors recommend that radiographs be obtained. Radiographs may also help relieve parental anxiety. Alternatively, instructions may be given for 24-hour follow-up if normal function is not restored, with consideration of radiographs at the time of follow-up. In two series of patients with nursemaid’s elbow, 6 of 10 patients released without normal arm use had spontaneous restoration of function, whereas the other 4 required remanipulation (which successfully restored function). Delayed use of radiographs may decrease overall radiation exposure in a child with a radial head subluxation.

The use of a posterior splint to protect the elbow of a child who refuses to use the affected arm after a presumed reduction is of uncertain value. However, some form of immobilization (e.g., splint, sling, or both) may be valuable in a child with

**Reduction Techniques**

**Supination Method**

Reduction of nursemaid’s elbow (Fig. 49.29) is generally performed without premedication. If the subluxation has been present for hours, oral or nasal midazolam can be a useful adjunct to overcome the child’s anxiety related to manipulation (see Chapter 33). It is important to explain to the caretaker that the reduction will probably cause the child discomfort and precipitate crying, but that this is transient and a clue to the diagnosis. Position the child seated on the lap of an assistant (often a parent) who stabilizes the arm by holding the humerus adducted to the side. Grasp the elbow with one hand and place the thumb over the region of the radial head. Although it has been stated that the thumb can apply pressure on the radial head, this positioning is mainly useful for palpation of the reduction “click.” Grasp the child’s wrist with the other hand and supinate the extended forearm in a steady, deliberate manner. Slight traction before supination is generally recommended, but it is unclear whether this increases the likelihood of successful reduction. Once supinated, flex or extend the arm; however, flexion is the most common maneuver and may actually be somewhat more successful than extension. An audible or palpable click signifies successful reduction but is not always noted. Once the reduction has been performed, the child usually cries for a few minutes. Generally, the operator should leave the room and then return in 10 to 15 minutes to repeat the examination. Full use of the arm should be evident (see Fig. 49.28B).

**Pronation Method**

This technique is performed with the child positioned as for the supination method. However, the forearm is not supinated. Instead, the forearm is rapidly hyperpronated and flexed.

![Figure 49.29 Nursemaid’s elbow (radial head subluxation) reduction. (From Fleisher GR, Ludwig S, editors: Textbook of Pediatric Emergency Medicine. Baltimore, 1988, Williams & Wilkins, p 1322. Reproduced by permission.)](image-url)
significant residual discomfort after a prolonged period of subluxation or in whom recurrent subluxations have taken place. On occasion, a successful reduction painfully resubluxates with movement; in this case, immobilization and referral may be necessary. If reduction has been achieved clinically and maintained in the ED, analgesics or a follow-up visit is unnecessary. Because other pathology can rarely mimic this condition (e.g., occult fracture, osteomyelitis, joint infection, tumor), full, unrestricted, and painless use of the arm must be evident by 24 hours. If not, further workup is warranted.

The emergency clinician should consider parental education regarding prevention of radial head subluxation. Parents or caregivers should be instructed to lift toddlers who are at an age most at risk for these injuries from beneath their axilla, as opposed to forcibly lifting or pulling them by the wrist or hand.

**HAND INJURIES**

The hand is extremely susceptible to dislocation injuries because of its frequent use and exposed nature. Proper motion and function of the hand are intimately related to normal anatomic alignment. The emergency clinician must therefore be skilled in the diagnosis and management of dislocations involving the hand. An improperly managed hand injury can result in significant long-term or permanent disability.

Anatomically, the joints of the digits are quite similar and consist of a hinged joint with a tongue-in-groove–type arrangement. The soft tissue support includes two collateral ligaments attached to a volar plate (Fig. 49.30). The volar plate is composed of dense fibrous connective tissue that is thickened at its distal attachment and thinner at its proximal attachment to allow folding with joint flexion. Dorsal dislocation of a digit requires failure of the volar plate, whereas lateral dislocation disrupts a collateral ligament and induces at least a partial tear in the volar plate.

Radiographic examination of all hand injuries is relatively straightforward and includes at least two views (AP and lateral) of the injured area (Fig. 49.31). The most important radiographic error in evaluating joint injuries in the hand is failing to get a true lateral view of the injured joint. This may lead to missing a fracture or a loose body in the joint.

Anesthesia is generally required for proper management of dislocations involving the hand. It is most often accomplished with a finger or wrist block (see Chapter 31), although a more proximal regional or Bier block may be used on occasion (see Chapter 32). Getting a secure grip on the digits may be difficult and could complicate the reduction. Wearing rubber gloves or wrapping gauze around the fingers may be useful.

**Figure 49.30** A and B, The collateral ligament–volar plate relationship. The metacarpophalangeal and interphalangeal joints derive their strength from a combination of the two collateral ligaments and the volar plate. Dislocations of these joints require tearing of at least two parts of this three-part structure. C, Lateral view demonstrating the collateral ligament–volar plate relationship. (A and B, From Carter P, editor: Common hand injuries and infections. Philadelphia, 1983, Saunders, p 114. Reproduced by permission; C, redrawn from Eaton RG: Joint injuries of the hand. Springfield, IL, 1972, Charles C Thomas.)

**Figure 49.31** Phalangeal dislocations. A, Dorsal dislocation of the metacarpophalangeal (MCP) joint of the thumb. B, Dorsal dislocation of the proximal interphalangeal (PIP) joint of the middle finger. Note the presence of a small volar plate fracture (arrow). C, Dorsal dislocation of the distal interphalangeal (DIP) joint of the little finger. Dorsal dislocations are much more common than volar ones. AP, Anteroposterior; LAT, lateral.
**Thumb Dislocations**

The opposable thumb is an essential structure for countless activities. Despite its strong ligamentous and capsular support, the exposed positioning of the thumb makes it a frequent site of dislocations and subluxations. The metacarpophalangeal (MCP) joint in the thumb is similar to the MCP joints in the fingers but has a stronger volar plate and collateral ligaments.\(^{118}\)

**Interphalangeal Joint Dislocation of the Thumb**

The single IP joint of the thumb has strong cutaneous-periosteal attachments, and dislocations of this type are therefore frequently open.\(^{118}\) Dislocations at the IP joint are generally dorsal and can be reduced in a manner similar to that for IP dislocations of the finger (Fig. 49.32A and B). First, re-create the mechanism of injury by applying longitudinal traction and hyperextension to distract the phalanges. Next, flex the IP joint with continued traction and apply direct pressure on the base of the distal phalanx to complete the reduction.\(^{118}\)

After reduction, test the range of motion and stability of the joint and obtain post-reduction radiographs. If adequate reduction has been achieved, splint the thumb in slight flexion for 3 weeks.\(^{118}\) Orthopedic or hand specialist referral is advisable.

**Metacarpophalangeal Joint Injury of the Thumb**

Dorsal Dislocation

The MCP joint of the thumb can be dislocated dorsally by a hyperextension injury. The proximal phalanx will come to rest in a position dorsal to the first metacarpal (Fig. 49.33; also see Fig. 49.31A). There are two basic types of MCP dislocation

---

**Phalangeal Joint Dislocation Reduction**

**A. Traction Method**

- **Dorsal PIP dislocation**
  - Apply axial traction to the finger, and then push anteriorly on the base of the dislocated phalanx.
  - Flex the finger while continuing to apply traction and anterior pressure.
  - After reduction, test for range of motion and stability. Obtain a postreduction x-ray and apply a splint.

**B. Exaggeration Method**

- **Dorsal PIP dislocation**
  - Exaggerate the dislocation to distract the phalanges, and then apply pressure to the base of the dislocated phalanx.
  - Flex the finger while continuing to apply traction and anterior pressure.
  - After reduction, test for range of motion and stability. Obtain a postreduction x-ray and apply a splint.

**C. Thumb MCP Dislocation**

- If a simple thumb MCP dislocation is treated with traction alone, the forces will often interpose the volar plate and result in an irreducible complex dislocation. To reduce the joint properly, (1) firmly grasp the thumb, (2) exaggerate the dislocation by hyperextending the thumb, and (3) apply pressure to the base of the dislocated phalanx.
  - To complete the reduction, flex the thumb forward. Post-reduction, check the stability of the joint by putting it through the full range of motion, and assess the integrity of the collateral ligaments (see text). Apply a thumb spica splint with the thumb held in flexion.

---

*Figure 49.32 Phalangeal dislocation reduction. MCP, Metacarpophalangeal*
from the entrapped metacarpal head. On radiographs of simple dislocations, the joint surfaces are in close contact, whereas they are separated in complex dislocations. The presence of a sesamoid bone in the joint space is diagnostic of a complex MCP dislocation (Fig. 49.35).

To reduce a simple MCP dislocation, hyperextend the joint as far as possible with the wrist in flexion to relax the tendons (see Fig. 49.32). Once maximal hyperextension is achieved, push the base of the proximal phalanx distally while bringing the joint back into flexion. Applying simple traction alone as an initial maneuver risks trapping the volar plate and creating a complex dislocation. After reduction, test the stability of the joint by putting it through a full range of motion. Assess the integrity of the collateral ligaments with the MCP joint in flexion (see later). Simple MCP dislocation injuries generally require casting for 3 weeks with the joint in moderate flexion.

Volar Dislocation
Volar dislocations are rare and generally associated with collateral ligament ruptures. They are commonly irreducible because of interposition of one or both extensor tendons and the dorsal capsule. Orthopedic consultation is required.

Ulnar Collateral Ligament Rupture
Also known as gamekeeper’s or skier’s thumb, this injury results from a laterally directed force at the thumb MCP joint causing rupture of the ulnar collateral ligament (Fig. 49.36). The usual mechanisms include falling with a ski pole in the hand or having the thumb alone draped over the steering wheel in an auto crash. These injuries are most often initially seen in the reduced state with just a complaint of pain in the area. Early recognition of this injury is essential to prevent further disability because this ligament is important for the grasping function of the thumb. A strain or partial tear probably cannot be diagnosed in the ED. It is therefore prudent to immobilize all significantly “sprained thumbs” in a thumb spica splint for a few days and reexamine those with significant injuries. A complete or severe ligamentous tear is generally diagnosed by stress testing of the MCP joint (Fig. 49.37). Radiographs occasionally demonstrate an avulsion-type fracture. The exact positioning of the thumb for stress testing is debatable, but the metacarpal should be stabilized with the thumb and index
SECTION VIII  MUSCULOSKELETAL PROCEDURES

Figure 49.36 Rupture of the ulnar collateral ligament (gamekeeper’s thumb). 1. This injury is caused by forcible abduction. If unrecognized and untreated, progressive metacarpophalangeal (MCP) subluxation may occur (2), which interferes with grasping and causes significant permanent disability. Suspect this injury when there is a complaint of pain in this region. Look for tenderness on the medial side of the MCP joint. (Modified from McRae R: Practical fracture treatment. Edinburgh, UK, 1981, Churchill Livingstone, p 162. Reproduced by permission.)

Figure 49.37 Stress testing of the ulnar collateral ligament of the thumb. This is done both clinically and with an anteroposterior radiograph. A, A line is drawn on the skin with a pen. The line is along the long axis of the metacarpal and the proximal phalanx. B, Deviation of the straight line during stress indicates instability. The metacarpal is fixed with the operator’s other hand. Note: With acute injuries of the thumb, a simple sprain may be diagnosed when an ulnar collateral ligament injury is partial or severe and stress tests are negative because of swelling and spasm. Therefore it is prudent to splint all significantly sprained thumbs and reexamine them in a few days if the symptoms persist.

Figure 49.38 Carpometacarpal (CMC) dislocation of the thumb (arrows in A and B). A, Anteroposterior (AP) and lateral (LAT) radiographs of a CMC dislocation of the thumb, which is an uncommon injury. The patient sustained this injury when an air bag deployed during a motor vehicle collision. These dislocations are not generally amenable to closed reduction and usually require Kirschner wires. B, Pre- and post-reduction films of a CMC dislocation.

finger of one hand while applying stress with the other hand. Louis and associates\textsuperscript{119} recommended stressing the joint in full flexion because virtually no lateral movement of the MCP joint should be noted in this position. Instability in full flexion of greater than 35 degrees is indicative of a complete rupture. Hossfeld and Uehara\textsuperscript{118} suggested testing the MCP joint in 20 to 30 degrees of flexion to lessen the stabilizing effects of the volar plate; the results should be compared with stability on the other side.

Cast partial injuries to the ulnar collateral ligament for 3 weeks; complete rupture usually requires operative repair.\textsuperscript{119} An associated nondisplaced fracture may be treated in closed fashion, whereas a displaced fracture is an indication for operative repair.\textsuperscript{119}

CMC Dislocations of the Thumb

CMC dislocations of the thumb are uncommon (Fig. 49.38); when present, they often occur with an associated fracture. Because closed reduction is generally unstable, operative stabilization by percutaneous placement of Kirschner wires (K-wires) is usually required.\textsuperscript{118}

Finger Dislocations

The basic anatomic structure of the fingers is similar to that of the thumb with the exception that the fingers have more lateral support from the MCP joints, which makes collateral ligament injury here much less common than in the thumb. The treatment principles are also similar. It is advisable to...
order radiographs for a specific finger (not just “hand” films). Complete views of the finger will include an AP view, a true lateral view, and an oblique view. A true lateral view is extremely important for detection of subtle dislocations or small avulsion fractures on the volar surfaces (Fig. 49.39).

**Proximal Interphalangeal (PIP) Joint Dislocations**

The PIP joint is extremely important, and any loss of motion in this joint may severely restrict normal function.117 This joint is also prone to stiffness, so careful treatment of injuries in this area is essential. Injuries to the PIP joint are generally slow to heal and often result in an increase in joint size as a result of scar tissue formation.117 Because of this propensity for a less-than-perfect outcome, it is advisable to refer PIP injuries after emergency care.

Proper examination of the PIP joint (Fig. 49.40) includes the application of ulnar and radial stress to test the integrity of the collateral ligaments and hyperextension to determine the integrity of the volar plate. Inability to actively extend the flexed PIP joint against resistance suggests a central slip rupture, which may progress to a boutonnière deformity (see Chapter 48).117 Such an examination should be carried out after any successful joint reduction. This examination should also be conducted on a painful PIP joint that is radiographically normal to detect soft tissue injury in a spontaneously or field-reduced dislocation. This is extremely important in athletes because coaches often reduce these injuries.117

**Dorsal PIP Dislocations**

Dorsal PIP dislocations are among the most common types encountered in the ED (Fig. 49.41; also see Fig. 49.31B). The mechanism is usually a blow to the end of the finger, such as from a thrown ball, that creates an axial load and hyperextends the finger.118 The middle phalanx comes to rest dorsal to the proximal phalanx (Fig. 49.42). An associated disruption of the volar plate is generally present.117,118 The deformity is obvious

**Figure 49.39** Fracture-dislocations of the proximal (PIP) and distal interphalangeal (DIP) joints. True lateral radiographs are required to maximize sensitivity for small avulsion fractures. **A**, A volar avulsion fracture (arrow) is readily seen on this dorsal DIP dislocation. **B**, The fractures are more difficult to appreciate on this unfortunate patient who sustained dislocations of both the DIP and PIP joints in a fall (arrows).

**Figure 49.40** Post-reduction stress applied to a dislocated proximal interphalangeal (PIP) joint. **A**, If the volar plate has been completely disrupted, the PIP joint will hyperextend with both passive and active motion. **B**, Inability to actively extend the PIP joint indicates rupture of the central slip of the extensor tendon. **C**, Passive lateral stress is applied to check the integrity of the collateral ligaments. (A–C, From DePalma AF: Management of fractures and dislocations: an atlas. Philadelphia, 1970, Saunders, pp 1203–1204. Reproduced by permission.)

**Figure 49.41** Dorsal proximal interphalangeal joint dislocation.

**Figure 49.42** A dorsal proximal interphalangeal joint dislocation **A**, may involve rupture of the volar plate itself or **B**, may involve avulsion of varying amounts of bone from the middle phalanx. If a large fragment of bone is avulsed from the base of the phalanx, the dislocation is unstable after reduction. The collateral ligaments will tear in varying degrees and should be assessed with stress testing after reduction.
on clinical examination, and radiographs clearly demonstrate the injury. An associated fracture of the volar lip may be detected. If this fracture affects greater than 33% of the joint surface, closed reduction will be unstable because the collateral ligament is attached to the bony fragment. In these cases, operative repair will be necessary.118

A dorsal PIP dislocation can be reduced after a digital block. The usual method (see Fig. 49.32A and B) is to exaggerate the injury by slight traction and hyperextension, thereby distracting the middle phalanx. Apply pressure to the base of the middle phalanx as the finger is brought into flexion. These injuries usually reduce fairly easily, and failure of routine attempts should raise suspicion of interposed soft tissue, for which orthopedic consultation should be sought.

After reduction is completed, place the joint through a range of motion to ensure stability of the reduction. If stable, splint the joint in 20 to 30 degrees of flexion for 3 weeks.117,118 Alternatively, buddy taping to an adjacent finger for 3 to 6 weeks allows early active motion and prevents hyperextension, which should be avoided.118 Because PIP injuries can be slow to heal and are complicated by stiffness, it is advisable to refer these patients for orthopedic follow-up.

Volar PIP Dislocations
Volar PIP dislocations are uncommon injuries and virtually always accompanied by injury to the central slip of the extensor tendons. If the dislocation is reduced before the ED visit and there is no indication that the dislocation was volar, this injury may be incorrectly treated by splinting in mild flexion as though it were the more common dorsal dislocation. However, with disruption of the central slip of the extensor tendons, immobilization in flexion will lead to the development of a boutonnière deformity.117,120 Even when this injury is recognized and treated properly, some impairment of mobility may occur. It is generally best to seek early orthopedic consultation for these injuries because some require operative repair. If the emergency clinician accomplishes a closed reduction, post-reduction films must demonstrate normal congruity of the joint surfaces, and a central slip attachment fracture must be excluded.120 If so, splinting of only the PIP joint in full extension should be undertaken for 3 weeks and early orthopedic follow-up ensured.117

Lateral PIP Dislocations
Lateral PIP dislocations are fairly common and often reduced in the field. The patient will frequently describe how the finger was pointing in an unnatural manner (Fig. 49.43). The injury can be detected by applying ulnar and radial stress to the PIP joint. If still dislocated, re-create the injury and then apply longitudinal traction to reduce the dislocation. Treat partial tears of the collateral ligaments by buddy taping the finger for 3 to 6 weeks.117 Management of complete tears is controversial, with some advocating operative therapy for all such injuries and others using varying durations of immobilization or buddy taping.117 Referral is suggested for all but the mildest of PIP collateral ligament injuries.

Distal Interphalangeal (DIP) Dislocations
As in the thumb, the distal phalanx is attached firmly to skin and subcutaneous tissue by osteocutaneous fibers. For this reason, dislocations of the DIP joint are frequently open.117 A DIP dislocation is usually dorsal, and the mechanism is a blow to the end of the finger. Despite the dislocation, the DIP joint may retain some range of motion, so it is important to not overlook these injuries.121 Lateral radiographs are diagnostic.

Management of a dorsal DIP dislocation involves reduction in a fashion similar to that described for other IP joint injuries (see Fig. 49.32). Hyperextend the finger and apply traction to distract the joint, and then apply pressure on the base of the distal phalanx during flexion. Following reduction, check the joint for stability and place the finger in a dorsal splint for 10 to 12 days.117

An injury to the DIP joint that may be confused with a dislocation is the mallet finger (Fig. 49.44). This injury is often caused by forced flexion of an extended finger (e.g., struck by a baseball). The patient is unable to extend the fingertip, but the joint appears normal on passive extension by the examiner. The injury is a rupture of the extensor tendon, with or without avulsion of a small piece of bone. Unless the injury is properly splinted or surgically immobilized, permanent deformity will occur (see Chapter 48).

MCP Dislocations
The pathology and management of finger MCP dislocations are identical to those of the thumb, as discussed earlier. The same classification of simple and complex applies; the complex type requires operative repair. Dimpling on the palmar surface suggests the presence of a complex dislocation. It is important to remember that application of traction alone for a simple MCP dislocation may convert it to a complex dislocation. For dorsal dislocations, flex the wrist to relax the tendons and hyperextend the joint as far as possible. Then apply pressure on the base of the proximal phalanx to effect reduction. After reduction of a simple dorsal MCP dislocation, buddy taping is generally sufficient to secure the reduction.117 Volar dislocations are rare and require orthopedic consultation.
CHAPTER 49  Management of Common Dislocations

1011

long-term disability, such as chronic pain and weakness, premature arthritis, and avascular necrosis. Specific intervention to establish anatomic alignment is usually undertaken on diagnosis or within a few days of the injury, so identification in the ED, initial splinting, and proper referral are paramount. Definitive treatment is beyond the scope of this chapter and is performed by a consultant, often a hand surgeon.

The most common carpal injuries are scapholunate dissociation and lunate and perilunate dislocation, although a number of variations exist. These injuries are often sustained by a fall on an outstretched hand. Carpal dislocations and dissociations produce pain, weakness, decreased range of motion, and often significant soft tissue swelling, with many symptoms seemingly out of proportion to the radiographic findings. Specific radiographic diagnosis is frequently difficult, and it is not uncommon for definitive diagnosis to be missed on initial ED evaluation. CT or magnetic resonance imaging may be required to delineate the specifics of the injury. Marked swelling on the dorsal surface of the hand or wrist, in the absence of a definitive fracture on plain radiographs, is a common scenario, and such findings should prompt consideration of carpal dislocation or dissociation (Fig. 49.45). If a diagnosis is not forthcoming, splinting and reevaluation of a markedly swollen and painful hand or wrist in 2 to 3 days is prudent.

Scapholunate dissociation is a serious, frequently missed injury that is characterized by a widened space (>3 mm) between the scaphoid and lunate bones on a plain radiograph. This is best visualized on a posteroanterior view with the hand closed in a fist and the wrist in ulnar deviation (Fig. 49.46). It is indicative of disruption of ligaments stabilizing the two carpal bones. Even though the symptoms and swelling may be impressive, the radiographic findings are subtle and often overlooked, with initial and subsequent complaints being attributed to a severe bruise or sprain. When identified, the hand and wrist are immobilized in the ED by a splint in neutral position or in 10 to 15 degrees of wrist dorsiflexion. Referral should also be initiated.

Lunate dislocation and perilunate dislocation (Figs. 49.47 and 49.48) can also occur as a result of trauma and produce pain, swelling, and disability. Plain radiographs will usually

Figure 49.44  A, A mallet finger injury is not a dislocation; it is a rupture of the extensor tendon to the distal phalanx. It can occur with or without (demonstrated here) an avulsion fracture after seemingly minor trauma. B, This mallet deformity was caused by a baseball striking the fingertip end on and producing acute flexion of the joint. C, A stack splint, with the proximal interphalangeal joint left free to allow relaxation of the distal interphalangeal joint, is kept in place for 6 to 8 weeks. Caution the patient to avoid flexing the joint to “test it out” during splint changes. Wire fixation may be required.

CMC Dislocations

CMC dislocations are rare injuries that are frequently misdiagnosed. The usual site of injury is the fifth CMC joint, which is dorsally dislocated.118 The injury is generally the result of a high-energy mechanism, such as a motor vehicle crash or a fall. The diagnosis can be quite difficult to make because the appearance may be subtle even on a lateral radiograph. Associated fractures and other injuries are frequently present, and percutaneous fixation is usually required.118

Carpal Dislocation/Dissociation

Dislocation and dissociation of the carpal bones are significant injuries that require identification in the ED and expeditious immobilization and reduction to ensure the best ultimate outcome. Nonetheless, these injuries often produce significant

Figure 49.45 With this much soft tissue swelling 2 days after a fall on an outstretched hand, something more than a sprain or bruise is likely. If no fracture is seen on radiographs, consider carpal bone fracture or dissociation as the culprit. Scaphoid fracture, scapholunate dissociation or lunate or perilunate dislocation should be considered. Computed tomography or magnetic resonance imaging may be required to define the exact pathology. Note that patients with such injuries hold the wrist in flexion, worsening the edema by compression of dorsal hand veins.

Figure 49.46 Scapholunate dissociation is a serious, frequently missed injury that is characterized by a widened space (>3 mm) between the scaphoid and lunate bones on a plain radiograph. This is best visualized on a posteroanterior view with the hand closed in a fist and the wrist in ulnar deviation. It is indicative of disruption of ligaments stabilizing the two carpal bones. Even though the symptoms and swelling may be impressive, the radiographic findings are subtle and often overlooked, with initial and subsequent complaints being attributed to a severe bruise or sprain. When identified, the hand and wrist are immobilized in the ED by a splint in neutral position or in 10 to 15 degrees of wrist dorsiflexion. Referral should also be initiated.

Lunate dislocation and perilunate dislocation (Figs. 49.47 and 49.48) can also occur as a result of trauma and produce pain, swelling, and disability. Plain radiographs will usually
mentous and muscular support is very strong. Therefore hip dislocations are usually the result of significant force, and a careful search for other limb- or life-threatening injuries must be undertaken. Common mechanisms of hip dislocation include motorcycle or car accidents, vehicles striking pedestrians, and falls.122

Associated fractures are quite common with hip dislocations. In fact, up to 88% of hip dislocations are accompanied by an associated fracture.123 If a fracture complicates the dislocation,
Analgesia and Anesthesia

Dislocation of a prosthetic hip can usually be managed with moderate amounts of IV premedication in the ED. Premedication recommendations for other acute traumatic dislocations run the gamut from general anesthesia for all reductions to orthopedic consultation is generally indicated. However, the emergency clinician should be able to reduce simple hip dislocations, which are dislocations without an associated fracture or with a very minor fracture. Hip dislocations may occasionally be missed in the setting of severe trauma because other injuries garner more attention. A missed diagnosis can also occur when a fracture of the femur obscures the clinical picture of hip dislocation. Common complications of hip dislocation include sciatic nerve injuries and avascular necrosis of the femoral head. Sciatic nerve injuries are seen with 10% to 14% of posterior hip dislocations. Avascular necrosis of the femoral head is one of the more disabling complications associated with hip dislocation. Although it is generally stated that early reduction will reduce the frequency of this complication, evidence for this statement is hard to find. Dreinhofer and coworkers noted poor outcomes despite early (i.e., <6 hours) reduction of type I hip dislocations (dislocation without a significant associated fracture). Yang and colleagues found that reduction beyond 24 hours was associated with a worse prognosis, but they could not find a significant time factor for those reduced in less than 24 hours. Nevertheless, it is still advisable to reduce hip dislocations as soon as feasible to decrease soft tissue distortion. If evidence of nerve injury exists, the dislocation should be treated as an emergency and reduced as early as possible.

Radiographic Examination

Dislocation of the hip is generally obvious on the standard AP pelvic film that is often taken during trauma resuscitation (Fig. 49.49). A lateral or oblique view of the hip may help clarify the type of dislocation and allow detection of associated fractures.

Analgesia and Anesthesia

Dislocation of a prosthetic hip can usually be managed with moderate amounts of IV premedication in the ED. Premedication recommendations for other acute traumatic dislocations run the gamut from general anesthesia for all reductions to
involved in closely monitoring a critically ill patient in the hospital. It is preferable in multiply injured patients because of the difficulty of reducing hip dislocations. The supine position may be used to immobilize the patient and prevent further injury before reduction is attempted.

Several basic methods of hip reduction have been reported. Reduction Techniques

Reduction Techniques

Several basic methods of hip reduction have been reported. Some techniques involve placing the patient in the prone position (e.g., the Stimson technique), whereas others require that the patient be supine with downward stabilization of the pelvis performed by an assistant. The supine position may be preferable in multiply injured patients because of the difficulty involved in closely monitoring a critically ill patient in the prone position.

Stimson Technique (Fig. 49.51A)

In the prone or gravity method described by Stimson, the patient is positioned so that the distal portion of the pelvis overhangs the edge of the stretcher. Flex the hip, knee, and ankle to 90 degrees and apply downward pressure on the posterior aspect of the proximal end of the tibia. Gently rotate the hip internally and externally to facilitate reduction. If needed, an assistant may apply direct downward pressure on the femoral head. An alternative and more comfortable way to provide downward pressure on the tibia is to grasp the patient’s ankle, place your knee on the patient’s calf, and use your body weight to apply pressure. This method is believed to be the least traumatic; however, associated thoracoabdominal injuries or a need for deep sedation, which may pose an airway risk, may make the prone position risky for the patient.

Allis Technique (See Fig. 49.51B)

For the Allis technique, place the patient supine and have an assistant stabilize the pelvis by applying downward pressure at the anterior superior iliac spines. Exert upward traction in line with the deformity and flex the hip to 90 degrees. Gently rotate the hip internally and externally until it is reduced. Some prefer to stand on the patient’s stretcher so that body weight can be used for leverage. Howard suggested modifying this technique by applying lateral traction on the flexed upper part of the femur to disengage the head of the femur from the outer lip of the acetabulum.

Whistler Technique (See Fig. 49.51C)

Another method known as the Whistler technique was described by Walden and Hamer. Place the patient in the supine position with both knees flexed to 130 degrees. While an assistant stabilizes the pelvis, stand beside the affected limb. Place an arm under the affected knee and grab the unaffected knee. With the other hand, anchor the ankle of the affected leg firmly against the stretcher. Using the arm placed under the knee as a lever, raise your shoulder to elevate the affected knee. This allows the femoral head to move anteriorly around the acetabular rim and relocate. Although experience with this technique is limited, it appears to be a promising, gentle reduction method.

Captain Morgan Technique (See Fig. 49.51D)

Another gentle method for reducing posterior hip dislocations is similar to the Whistler technique but uses the clinician’s knee as opposed to the arm. To perform the Captain Morgan technique, place the patient supine with the affected knee flexed. Stand on the affected side and place your flexed knee underneath the patient’s flexed knee. Apply force with your leg in an upward direction. Internal or external rotation can be added as needed. One case series described an extremely high success rate with this method of reduction, with only one failed attempt secondary to bony fragments in the joint space, which required open reduction.

Once reduction is achieved, the legs are immobilized in slight abduction by placing an abduction pillow or another object between the knees. Radiographs should be repeated to confirm reduction, and the patient should be admitted to the hospital.

Dislocations of Hip Prostheses

Dislocation of a hip prosthesis is a separate issue (Fig. 49.52). Unlike primary dislocations, which require significant trauma, a prosthetic hip may dislocate with minimal force, such as rolling over in bed or trying to get out of a chair. Most dislocations occur in the first 3 to 4 months after surgery, but recurrent dislocations may take place much later. The majority of dislocations are posterior. The emergency physician should consider

Figure 49.50 A, Posterior hip dislocation in a patient with a hip prosthesis. The right leg is adducted, flexed at the knee, shortened, and internally rotated. B, The much less common anterior dislocation. The left leg is shortened, abducted, flexed at the knee, and externally rotated, similar to the appearance of a hip fracture.

Figure 49.51 A, B, Posterior dislocation. Anterior dislocation.
consultation with the orthopedic surgeon who placed the prosthesis.

The three major causes of prosthetic hip dislocations include (1) assumption of a position that exceeds the stability of the prosthesis, (2) soft tissue imbalances, and (3) component malposition.130

Emergency clinicians have been shown to be highly successful in reducing prosthetic hip dislocations.131 Reduction techniques are similar to those described earlier; however, the urgency is not as paramount because problems with bone necrosis do not exist. Although complications are occasionally unavoidable, the clinician must be aware that forceful reduction of a dislocated hip prosthesis may dislodge the acetabular cup, fracture underlying osteoporotic bone, or loosen the prosthesis. Unlike other hip dislocations, patients with prosthetic hip dislocations will often not require hospital admission and may be discharged after discussion with the consulting orthopedic surgeon. The most common way to reduce such dislocations is shown in Fig. 49.53.

**Anterior Hip Dislocation**

Anterior hip dislocations are less common than posterior dislocations and account for 10% to 15% of all hip dislocations.134 There are three general types of anterior hip dislocations that are defined by the place where the femoral head comes to rest (Fig. 49.54): iliac or subspinous, pubic, and inferior or obturator dislocation. Anterior hip dislocations generally
Although the knee is a simple hinge joint, dislocations are quite rare because of its strong ligamentous support. The major ligaments include the anterior and posterior cruciate and the collateral ligaments. The usual mechanism of a knee dislocation involves a great deal of force, such as a motor vehicle crash resulting from forced abduction of the thigh, which may occur with a fall or motor vehicle crash. The clinical picture varies with the type of dislocation. With the obturator (inferior) type, the leg is abducted and externally rotated with varying degrees of flexion. In the other types, the hip is usually extended and externally rotated.

Reduction Techniques (Fig. 49.55)
The Stimson gravity method may work for anterior hip dislocation, although it is not recommended for the pubic type. Alternatively, the Allis maneuver can be applied in a modified fashion. Place the patient in a supine position and have an assistant stabilize the pelvis while applying lateral countertraction on the thigh. Flex the hip slightly and apply traction along the long axis of the femur. Gently adduct and internally rotate the leg to affect reduction.

For the reverse Bigelow technique, place the hip in partial flexion and abduction. Apply traction in the line of the deformity, and adduct, sharply internally rotate, and extend the hip. Caution should be exercised when using this technique because the sharp internal rotation may result in fracture of the femoral neck in patients with osteoporotic bone. As with posterior dislocations, admission to the hospital is required for patients with these injuries.
or a sporting injury. However, knee dislocation has been reported after minor mechanisms, such as stepping off a curb or into a hole, usually in association with a twisting action.\textsuperscript{132} Obese patients may be more likely to dislocate a knee with surprisingly minor trauma, coined low-velocity or ultra-low velocity injuries, such as when stepping into a hole causes a twisting motion (Fig. 49.56). These patients are more commonly women and are at increased risk of vascular injury when compared to high-energy mechanism multi-ligament disruptions.\textsuperscript{133-135} As with other joint dislocations, knee dislocations are described with respect to the position of the distal bone in relation to the proximal one (i.e., tibia in relation to...
A severely unstable knee can be defined as one that has greater than 30 degrees of recurvatum (hyperextension) on lifting the leg off the stretcher by the heel\textsuperscript{138} or one that exhibits gross instability after reduction.\textsuperscript{136} Because pain and muscle spasm may limit the physical examination for stability, a knee hemarthrosis, usually a large one with signs of posterior or calf hemorrhage, is a potential tip-off to a reduced dislocation. An impressive effusion may not be present with a knee dislocation because the joint capsule is often disrupted and extravasation occurs into the surrounding tissue, usually posteriorly. The most important part of the clinical assessment is the vascular status of the extremity (see the next section). Nerve injury is less common, but peroneal nerve injury is a recognized complication, particularly with a posterolateral dislocation.\textsuperscript{136} Peroneal nerve injury should be suspected if the patient is unable to dorsiflex the ankle (footdrop) or has decreased sensation on the dorsum of the foot. Posterolateral dislocations may be irreducible because the medial femoral condyle buttonholes through the joint capsule.\textsuperscript{136} A clue to this injury is the presence of a dimple sign at the medial joint line.

**Clinical Assessment**

Knee dislocations are usually clinically obvious; however, in some cases the dislocation may have spontaneously reduced before evaluation in the ED and be manifested only as severe knee pain with hemarthrosis. When a spontaneously reduced knee dislocation is associated with other major trauma, the diagnosis can be missed. Obese patients may have a seemingly normal appearance of their knee, but an obvious deformity will be visible on the initial radiographs (see Fig. 49.56). A grossly unstable knee that does not appear to be dislocated is probably a reduced dislocation and carries the same risk for vascular and other complications as a dislocated knee.\textsuperscript{138} A

**Vascular Injury**

The most feared complication of a knee dislocation is severance or internal injury of the popliteal artery (Fig. 49.59). Injury

---

**Figure 49.57** The three most common types of knee dislocations: anterior (1), posterior (2), and lateral (3). (From DePalma AF: Management of fractures and dislocations: an atlas. Philadelphia, 1970, Saunders, p 1621. Reproduced by permission.)

**Figure 49.58** Anterior knee dislocation seen on a lateral radiograph. The patient sustained this injury when he was struck by a vehicle.

**Figure 49.59** Complete occlusion of the popliteal artery after posterior knee dislocation. This injury can be subtle or delayed. (From Valji K, editor: Vascular and interventional radiology, ed 2, Philadelphia, 2006, Saunders.)
to the popliteal artery may complicate both anterior and posterior knee dislocations and occurs because the artery is relatively fixed both proximally and distally.\textsuperscript{136} If popliteal artery injury occurs, it is often due to dissection with posterior dislocations or intimal tearing due to traction forces during anterior dislocations.\textsuperscript{137} In addition, Varnell and associates\textsuperscript{138} noted that vascular injury was as common in a severely unstable knee (e.g., field reduced) as in an acutely dislocated knee. The incidence of popliteal artery injury in a dislocated knee is approximately 20\% in most series.\textsuperscript{139–140} However a 2014 database review of 8050 patients with knee dislocation found an overall frequency of vascular injury of 3.3\%.\textsuperscript{141} The seriousness of this complication is largely due to the fact that the collateral circulation about the knee is poor,\textsuperscript{135} and the end result of injury to the popliteal artery (or vein) may be amputation, particularly if recognition is delayed.\textsuperscript{142,143} It should also be noted that nerve injuries are more common in patients with a vascular injury.\textsuperscript{144,145}

It has previously been stated that popliteal artery disruption may be present despite a normal pulse.\textsuperscript{146} Such statements have led to recommendations to perform arteriography or exploration of all knee dislocations.\textsuperscript{146} However, some studies question this perspective. Varnell and associates\textsuperscript{138} reported a pulse deficit or absent pulse in all patients with vascular injury. Kendall and coworkers\textsuperscript{139} also reported clear clinical evidence of all popliteal artery injuries in knee dislocations. This group recommended exploration for obvious ischemia, angiography for patients with ischemia whose pulse is restored after relocation, and observation for all others.\textsuperscript{139} Dennis and colleagues\textsuperscript{147} reported that physical examination alone had 100\% accuracy in predicting the need for surgical intervention in patients with posterior knee dislocations. Miranda and associates\textsuperscript{148} reported that popliteal artery injury can be safely and reliably predicted by a physical examination that includes specific evaluation for active posterior hemorrhage (expanding hematoma, absent pulse, or the presence of a thrill or bruit). However, it is noted that the hard physical signs of arterial injury might be delayed for 24 to 48 hours. Thus, although focused clinician examination may be quite accurate in the vast majority of cases, any dislocated knee should prompt serious concern about the vascular integrity of the leg given the sometimes subtle or delayed manifestation of vascular injuries.

Simple palpation of the artery may not be sensitive enough to detect a decreased pulse. An ankle-brachial arterial pressure index (ABI) should be considered to compare blood pressure in the ankle with that in the arm. In addition, consider digit pulse oximetry to compare the uninjured leg with the injured one. Mills and coworkers\textsuperscript{145} reported the results of a prospective study of 38 patients with knee dislocation to evaluate the accuracy of ABI in identifying vascular injury. Patients with an ABI of less than 0.90 underwent arteriography, whereas those with an ABI of 0.90 or greater underwent serial examination and delayed arterial duplex evaluation. Eleven (29\%) of the patients had an ABI of less than 0.90, and all of them had arterial injuries requiring surgical intervention. Of the remaining 27 patients with an ABI of 0.90 or greater, none had a vascular injury noted on serial examination or duplex ultrasonography. No patient in this group was found to have vascular compromise at follow-up (range, 4 to 36 months).\textsuperscript{147}

Knee dislocations are usually clinically obvious or easily visible on plain radiographs. Therefore it is occult knee dislocations (i.e., dislocated and spontaneously reduced) that are problematic for the emergency clinician to diagnose. Internal derangement with a knee hemarthrosis (often of the size noted with a torn anterior cruciate ligament) is a common first sign that the knee had previously been dislocated and spontaneously reduced. Therefore all knee injuries with significant swelling, hemarthrosis, or a dislocating mechanism of injury should be evaluated with the specific intent of ruling out vascular injury.

If vascular compromise is detected on clinical assessment, it is appropriate to reduce the knee dislocation without obtaining radiographs, although a few minutes to obtain portable radiographs and to administer IV medication would be unlikely to make a difference in the final outcome.\textsuperscript{136} Use of Doppler ultrasound for pulse checks and ABI should also be considered with these injuries (see Chapter 1). Early consultation should be sought for knee dislocations because of the high incidence of complications and the frequent need for operative intervention. The decision to pursue angiography in a patient with a dislocated knee is best made in consultation with an orthopedic surgeon.

**Reduction Technique (Fig. 49.60)**

The need for IV sedation and analgesia depends on the clinical situation, but it should be considered routine whenever possible (see Chapter 33). The basic initial approach for all types of knee dislocation is to apply traction to the extremity. This
Anterolateral dislocation is the most common type. It is accompanied by obvious prominence of the fibular head anteriorly; no associated neurovascular problems are noted. The less common posterior dislocation may be accompanied by peroneal nerve injury. Patients have varying degrees of disability, and some may walk on the leg with only mild discomfort. On radiographic examination the three cardinal signs of anterolateral dislocation are lateral displacement of the fibula on an AP film, a widened proximal interosseous space, and anterior displacement of the fibular head on a lateral view (Fig. 49.61). If high clinical suspicion exists for fibular head dislocation and radiographs are nondiagnostic, CT is the next study of choice.

Reduction Technique
To reduce an anterior fibular head dislocation, place the patient supine and flex the affected knee to 90 degrees to relax the biceps femoris tendon. Dorsiflex and externally rotate the foot and apply direct pressure to the fibular head; reduction is usually signified by a snap. The method for reduction of a posterior dislocation is the same except that direct pressure is applied in a forward direction. Patients should not bear weight for 2 weeks and should receive orthopedic referral. Immobilization is probably unnecessary.

Post-reduction Care
Appropriate aftercare for knee dislocations requires serial reassessment of the neurovascular status of the extremity, post-reduction films, and admission to the hospital. Use a knee immobilizer to provide stabilization and comfort. These injuries cause severe ligamentous and other derangements in the knee and generally require operative stabilization with a long period of recovery and physical therapy.

DISLOCATIONS OF THE FIBULAR HEAD
The fibula can be dislocated at its proximal articulation in the knee joint. This is most commonly an anterolateral dislocation. The fibular head is normally nestled in a stable manner behind the lateral tibial condyle with two supporting tibiofibular ligaments. The tibiofibular joint has a separate synovial cavity, and therefore a typical knee joint effusion will not be seen with this dislocation. When the knee is flexed, the stability of this joint is decreased because of relaxation of the fibular collateral ligament. The typical mechanism of injury is a fall on a flexed, adducted leg, often combined with ankle inversion. This mechanism is seen in sports and parachute landings. Posterior dislocations can result from a twisting mechanism or a direct blow to the area while the knee is flexed.

Anterolateral dislocation is the most common type. It is accompanied by obvious prominence of the fibular head anteriorly; no associated neurovascular problems are noted. The less common posterior dislocation may be accompanied by peroneal nerve injury. Patients have varying degrees of disability, and some may walk on the leg with only mild discomfort. On radiographic examination the three cardinal signs of anterolateral dislocation are lateral displacement of the fibula on an AP film, a widened proximal interosseous space, and anterior displacement of the fibular head on a lateral view (Fig. 49.61). If high clinical suspicion exists for fibular head dislocation and radiographs are nondiagnostic, CT is the next study of choice.

Reduction Technique
To reduce an anterior fibular head dislocation, place the patient supine and flex the affected knee to 90 degrees to relax the biceps femoris tendon. Dorsiflex and externally rotate the foot and apply direct pressure to the fibular head; reduction is usually signified by a snap. The method for reduction of a posterior dislocation is the same except that direct pressure is applied in a forward direction. Patients should not bear weight for 2 weeks and should receive orthopedic referral. Immobilization is probably unnecessary.

PATELLAR DISLOCATION
Patellar dislocations are fairly common, especially in adolescents. The usual mechanism is a powerful quadriceps contraction combined with a strong valgus and external rotation component. This may be seen in activities such as making a “cut” in sports or with dancing. The patella may also dislocate with a direct blow to the flexed knee. Factors predisposing to patellar dislocation include chronic patellofemoral abnormalities such as genu valgum and femoral anteversion. Patellar dislocation is described by the relationship of the patella to the knee joint. Lateral dislocations are the most common by far. Other types include superior, medial, and intercondylar (Fig. 49.62).
The patella may be spontaneously reduced in the field with simple leg straightening. The patient will report that the leg “went out” and may describe actually seeing the lateral deformity caused by the displaced patella. Clinical clues to a spontaneously reduced patella include the presence of a knee effusion and tenderness along the medial edge of the patella. Fairbank’s test or the patellar “apprehension” sign is elicited when the patella is pushed laterally and the patient grabs for the knee, a response indicative of the sensation of repeated injury.63

Radiographs
Pre-reduction films are difficult to obtain because the patient’s knee is usually in flexion. Some recommend pre-reduction films when possible in all patients63; however, it is easy to reduce these injuries before radiographs are obtained. The diagnosis is usually obvious, and there are no reports in the literature of complications from gentle reduction. Osteochondral fractures are detectable in approximately half the patients with patellar dislocations, but many of these fractures are visible only on arthroscopy.152 Post-reduction radiographs are recommended, as are pre-reduction studies, when the diagnosis is uncertain. The clinical diagnosis of patellar dislocation in an older patient should be made with caution because these are primarily injuries involving the young.

Reduction Technique and Post-Reduction Care
Reduction of a lateral patellar dislocation is usually quite simple. Premedication is often not required if the patient can be verbally reassured. If the patient is anxious or in great discomfort, premedication should be considered (see Chapter 33). The two basic maneuvers for patellar relocation are extension of the knee and gentle medial pressure applied to the patella while lifting the most lateral edge of the patella over the femoral condyle (Fig. 49.64).63

Immobilize the leg in extension with a splint or commercially available knee immobilizer (see Chapter 50). Orthopedic follow-up is necessary because of the need for physical therapy and the high rate of persistent instability.152 However, hospitalization is not required for routine lateral dislocations of the patella. Recurrent dislocations and those associated with an osteochondral fracture might require operative repair.

Patellar dislocations in other locations are often irreducible, and orthopedic consultation should be sought. Intracondylar, superior, and vertical axis (dorsal fin) dislocations are extremely rare and require operative reduction. The rare horizontal dislocation may relocate with closed reduction, but surgical reduction is often necessary.

ANKLE DISLOCATIONS
The ankle joint is a modified saddle joint in which the talus is nestled in the mortise formed by the distal ends of the tibia and fibula.61 The ligamentous support of the ankle is quite strong, and pure dislocations are uncommon. Usually, there are associated fractures of the ankle joint (Fig. 49.65). Ankle dislocations are described by the relationship of the talus to the tibia. Posterior dislocations of the ankle are more common than anterior dislocations, and they usually result from a fall on a plantar-flexed foot. Patients with posterior dislocations often
have an associated fracture of one or more of the malleoli, occasionally seen only after reduction (Fig. 49.66). The clinical picture usually consists of significant deformity and disability.

Anterior dislocations generally result from forced dorsiflexion or a blow directed posteriorly onto the distal end of the tibia while the foot is fixed. The talus is prominent anteriorly, and the dorsalis pedis pulse may be lost secondary to pressure from the talus. Superior dislocations are uncommon and result in diastasis of the tibiofibular joint. These injuries are usually the result of a significant axial force, such as a fall from a significant height. Therefore clinicians should search for concomitant calcaneal or low spine fractures. Lateral dislocations of the ankle are always associated with fractures of the malleoli or distal end of the fibula.

Radiographic Examination

Because of the high rate of associated fractures and the clinical difficulty in assessing for the presence or the exact nature of a dislocation, it is recommended that pre-reduction radiographs be obtained for all suspected ankle dislocations (see Figs. 49.2 and 49.65). It is acceptable to reduce the dislocation without a radiograph if severe vascular compromise is present but, for the vast majority of cases, the few minutes taken to obtain

**Figure 49.65** Types of ankle dislocation. Note that there are usually associated fractures of the tibia, fibula, or both.
Lateral dislocations are really fracture-dislocations, and orthopedic consultation is generally required as part of the ED course. The emergency clinician will often need to reduce these injuries because of the extreme lateral deformity and occasional compromise of the dorsalis pedis artery by stretch. Open dislocations (in the absence of vascular compromise) may be better handled by operating room washout before attempting reduction (Fig. 49.68). If a lateral fracture-dislocation is to be reduced in the ED, the approach is quite similar to that for posterior ankle dislocations. However, instead of pressure in the AP direction, the foot is moved medially after the application of traction.61

**Post-reduction Care**

After reduction, splint the ankle at 90 degrees with a long leg posterior splint. Application of a stirrup splint in addition to the posterior splint will provide additional stability (see Chapter 50). The necessity of admission to the hospital must be determined in consultation with an orthopedic surgeon. Many patients with these injuries have associated fractures that require surgical intervention.

**DISLOCATIONS OF THE FOOT**

The importance of the foot is recognized by anyone who has had to spend time ambulating with an injury in this area. For purposes of this discussion, dislocation injuries of the foot can be divided into those of the hindfoot and those of the forefoot.

**Hindfoot Injuries**

Injuries to this area are uncommon and usually result from high-energy transfer. The major dislocations are subtalar and talar dislocations and midtarsal fracture-dislocations (Lisfranc injury). Although x-ray findings are often subtle and easily overlooked, a Lisfranc injury is complex and always a fracture-dislocation because of the rigid nature of the region (Figs. 49.69 and 49.70). These injuries require orthopedic management and are not discussed here.

**Subtalar Dislocation**

This uncommon injury generally occurs secondary to sports, falls from heights, or motor vehicle crashes. The calcaneus, navicular, and forefoot are displaced from the talus (Fig. 49.71). The primary mechanisms are severe inversion causing medial dislocation or severe eversion resulting in lateral dislocation. The medial type occurs so commonly during basketball that it has been termed *basketball foot*. This injury is usually seen in young adult males. Medial dislocations constitute the majority (85%) of these injuries, with lateral dislocations making up the rest.

The diagnosis is usually obvious because the talus is prominent and often tents the skin of the proximal part of the foot. The medial type has been termed an *acquired clubfoot*, the lateral type as an *acquired flatfoot*. Some authors recommend spinal or general anesthesia for all such injuries; however, it is usually possible to reduce these injuries with IV premedication. Position the patient in a supine fashion and flex the hip and knee much as for reduction.

**Reduction Techniques (Fig. 49.67)**

Unless a strong contraindication is present, it is advisable to administer IV sedation and analgesia to patients with an ankle dislocation early in their care, preferably before conducting any manipulations or radiologic studies. Reduction is always painful in an awake patient, and sufficient premedication must be administered. For posterior dislocations, place the patient supine and flex the knee slightly to relax the Achilles tendon. An assistant can do this, or the patient can be positioned such that the knee hangs over the end of the bed. Grasp the foot with both hands; place one hand on the heel and the other on the forefoot. Apply traction on the slightly plantar-flexed foot. Have a second assistant apply downward pressure on the distal end of the tibia and move the heel anteriorly to affect reduction.

For anterior dislocations, the initial steps and positioning are the same as those for posterior dislocation. However, instead of plantar flexion, dorsiflex the foot to free the talus. The second assistant applies upward pressure on the distal end of the tibia while the operator applies traction and pushes the foot in a posterior direction.61
Dislocation of the Talus

In this extremely rare injury, the talus is essentially extruded from its normal position and comes to lie anteriorly. This injury is generally open, is not amenable to closed reduction, and virtually always progresses to avascular necrosis. Talec-tomy and arthrodesis are often required, and orthopedic referral should be undertaken on an emergency basis if vascular compromise of the foot is present.

Forefoot Dislocations

Much of what is pertinent to the diagnosis and management of forefoot dislocations has already been discussed in the management of dislocations of the finger and hand MCP joints. Anatomically, the joints are quite similar.

Metatarsophalangeal (MTP) Dislocations

These uncommon injuries are generally the result of hyperextension resulting in dorsal dislocation of the great toe MTP joint (Fig. 49.72A). Among the lesser toe MTP joints, lateral or medial displacement of the digit on the metatarsal head is more common and usually the result of jamming the toe on a piece of furniture. As with MCP dislocations, they can be simple or complex. Complex dislocations of the first toe can be suspected by the presence of sesamoid bones in the joint.
space on pre-reduction radiographs. Complex MTP dislocations are often irreducible.

For simple MTP dislocations, reduction is accomplished by increasing the deformity through hyperextension and then applying traction while exerting thumb pressure over the base of the dislocated proximal phalanx. Plantar flexion of the foot may be used to relax the flexor tendons. Operative intervention is required after reduction if crepitus is present on motion, the joint is unstable, or an intraarticular loose body is noted on post-reduction radiographs.

**Interphalangeal Dislocations**

In the foot, IP dislocations result from an axial load on the toe, such as from kicking a wall (see Fig. 49.72B). These dislocations are generally dorsal and can be reduced as in the hand. Specifically, the toe should be dorsiflexed to exaggerate the
deformity and then undergo traction followed by plantar flexion. Dislocations of the first toe IP joint are usually buddy-taped to the second toe for 2 to 3 weeks, whereas those of lesser toes can be taped for 10 to 14 days.\textsuperscript{154} Reduction of dorsal dislocations may be difficult because of entrapment of the plantar plate inside the joint.\textsuperscript{157} As in the hand, complex dislocations or those failing closed reduction may occur. Such dislocations require open reduction and internal fixation.\textsuperscript{157}

CONCLUSION

The following points are important regarding the assessment and management of dislocations:
1. A search for other more serious injuries should be undertaken when there is a high-energy mechanism of injury.
2. Neurovascular assessment should be performed early in the evaluation and appropriately documented.
3. Radiographs and IV premedication are generally indicated before attempts at reduction.
4. Reduction attempts should involve gentle, gradual application of force and patience on the operator’s part.
5. After completing reduction, the operator should recheck the patient’s neurovascular status, request post-reduction radiographs (except with radial head subluxations), and in certain circumstances, assess the stability and range of motion of the joint.
6. A definite percentage of dislocations are irreducible, and the need for multiple attempts should halt prolonged and forceful attempts in the ED and prompt orthopedic consultation.
7. Dislocations accompanied by neurologic injury should be reduced by the most expeditious and least traumatic method.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
CHAPTER

REFERENCES
1. Mirick MJ, Clinton JE, Ruiz E: External rotation method of shoulder
and Kocher techniques for acute anterior dislocation of the shoulder.
3. Plummer D, Clinton J: The external rotation method for reduction of
acute anterior shoulder dislocation. Emerg Med Clin North Am 7:165,
1989.
4. Thakur AJ, Narayan R: Painless reduction of shoulder dislocation by
5. Garnavos C: Technical note: modifications and improvements of the
Milch technique for the reduction of anterior dislocation of the shoulder
7. McQueen A, Cress C, Tothy A: Using a tablet computer during pediatric
procedures: a case series and review of the “apps.” Pediatr Emerg Care
10. Hendey GW, Kinlaw K: Clinically significant abnormalities in post
reduction radiographs after anterior shoulder dislocation. Ann Emerg
11. Harvey RA, Trabulsy ME, Roe L: Are postreduction anteroposterior
and scapular Y views useful in anterior shoulder dislocations? Am J
12. Kwon YW, Zuckerman JD: Subluxations and dislocations about the
glenohumeral joint. In Bucholz RW, Heckman JD, Court-Brown CM,
et al, editors: Rockwood and Green’s fractures in adults (vol 2). ed 6,
Philadelphia, 2006, Lippincott, Williams & Wilkins, p 1285.
13. Riebel GD, McCabe JB: Anterior shoulder dislocation: a review of
14. May VR, Jr: Posterior dislocation of the shoulder: habitual, traumatic,
dislocations and humeral neck fractures: a prospective clinical and EMG
17. Gates JD, Knox JB: Axillary artery injuries secondary to anterior dislocation
18. Hersche O, Gerber C: Iatrogenic displacement of fracture-dislocations
19. Hendey GW, Chally MK, Stewart VB: Selective radiography in 100
associated with an anterior shoulder dislocation. Acad Emerg Med 11:853,
2004.
after anterior shoulder dislocation: a prospective study. Int Orthop 40(3):
22. Kahn JH, Mehta SD: The role of post-reduction radiograph after shoulder
ultrasonographic examination in the management of shoulder dislocation
portable ultrasound in the management of shoulder dislocation. Emerg
29. McNamara RM: Reduction of anterior shoulder dislocations by scapular
30. Matthews DE, Roberts T: Intra-articular lidocaine versus intravenous
analgesic for reduction of acute anterior shoulder dislocations: a prospective

49   Management of Common Dislocations 1026.e1

of acute shoulder dislocations: a prospective, randomized study comparing
intraarticular lidocaine with intravenous analgesia and sedation. Am J
intravenous analgesia and sedation for manual closed reduction of acute
33. Ng VK, Hames H, Millard WM: Use of intra-articular lidocaine as
analgesia in anterior shoulder dislocation: a review and meta-analysis of
34. Wakai A, O’Sullivan R, McCabe A: Intra-articular lignocaine versus
intravenous analgesia with or without sedation for manual reduction of
acute anterior shoulder dislocation in adults. Cochrane Database Syst Rev
35. Stone MB, Sutijono D: Intraarticular injection and closed glenohumeral
36. Aronson PL, Mistry RD: Intra-articular lidocaine for reduction of shoulder
37. Blaivas M, Lyon M: Ultrasound-guided interscalene block for shoulder
block and procedural sedation analgesia in shoulder dislocation reduction.
sedation and ultrasound-guided interscalene nerve block for shoulder
42.	Lippert FG, 3rd: A modification of the gravity method of reducing
43. Rollinson PD: Reduction of shoulder dislocations by the hanging method.
44. Anderson D, Zvirbulis R, Ciullo J: Scapular manipulation for reduction
45. Pishbin E, Bolvardi E, Ahmadi K: Scapular manipulation for reduction
of anterior shoulder dislocation without analgesia: results of a prospective
shoulder dislocations using the Milch technique: a study of ski injuries.
J Trauma 21:802, 1981.
47. Doyle WL, Ragar T: Use of the scapular manipulation method to reduce
an anterior shoulder dislocation in the supine position. Ann Emerg Med
27:92, 1996.
48. Sagarin MJ: Best of both (BOB) maneuver for rapid reduction of anterior
49. Parvin RW: Closed reduction of common shoulder and elbow dislocations
subcoracoid shoulder dislocation: evaluation of an external rotation
52. Milch H: The treatment of recent dislocations and fracture-dislocations
53.	Lacey T, 2nd, Crawford HB: Reduction of anterior dislocations of the
shoulder by means of the Milch abduction technique. J Bone Joint Surg
55. Ghane MR, Hoseini SH, Javadzadeh HR, et al: Comparison between
traction-countertraction and modified scapular manipulation for reduction
56. Miljesic S, Kelly AM: Reduction of anterior dislocations of the shoulder:
dislocation: a prospective randomized study comparing Spazo and Kocher
59. Poulsen SR: Reduction of acute shoulder dislocations using the Eskimo
dislocation of the shoulder: a simple method of reduction. Injury 23:479,




Splints are used frequently in the emergency department (ED) for temporary immobilization of fractures and dislocations and for definitive treatment of soft tissue injuries. Patients with a variety of nontraumatic musculoskeletal disorders (e.g., gout, inflammatory joint diseases, infections, burns) also benefit from short-term immobilization. Immobilization is the mainstay of fracture therapy, but though intuitively beneficial, it is difficult to find firm scientific data that support the use of splinting for soft tissue injuries. Although the general principle of immobilizing sprains and contusions is strongly supported by custom and personal preference, its exact influence on healing, number of complications, and ultimate return to normal activity is not known. In most studies of ankle sprains, for example, the function and pain of the injured joint are similar at 6 weeks’ follow-up, regardless of whether treatment consisted of ad lib walking, a simple elastic bandage, a posterior splint, or a formal cast. A systematic review of 22 clinical trials comparing various treatments of acute lateral ankle sprains (cast, splint, or early mobilization with support) found no favorable effect of immobilization. The current data support functional management for most acute ankle sprains.

Patients are routinely seen in the ED with injuries that are amenable to splinting to relieve pain and augment healing (see Review Box 50.1). Although a strict standard of care cannot be promulgated, the use of short-term splinting in the ED for acutely painful conditions remains a common practice.
Emergency clinicians have virtually abandoned the use of circumferential casts in favor of premade commercial immobilizing devices or splints constructed of plaster of Paris or fiberglass. The impetus for this change is primarily related to the complications occasionally associated with circumferential casts, liability issues, and ease of application brought about by new technology. In most instances, properly applied splints provide short-term immobilization equal to that of casts while allowing continued swelling and thus reducing the risk for ischemic injury. Other obvious advantages of splints are that patients can take them off when immobilization is no longer needed or can remove them temporarily to bathe, exercise the injured part, or perform wound care.

Most splinting techniques (Videos 50.1 to 50.6) are handed down from house staff or experienced clinicians, but the procedure is often suboptimal and haphazard. This chapter presents guidelines for adequate immobilization of injuries commonly encountered by emergency clinicians. Details of the construction and application of commonly used custom-made plaster splints are provided.

### INDICATIONS AND CONTRAINDICATIONS

Theoretically, immobilization facilitates the healing process by decreasing pain and protecting the extremity from further injury. Other benefits of splinting are specific to the particular injury or problem being treated (Fig. 50.1). For example, in the treatment of fractures, splinting helps maintain bony alignment. Splinting deep lacerations that cross joints reduces tension on the wound and helps prevent wound dehiscence. Immobilizing tendon lacerations may facilitate the healing process by relieving stress on the repaired tendon. The discomfort of inflammatory disorders such as tenosynovitis or acute gout is greatly reduced by immobilization. Deep space infections of the hands or feet, as well as cellulitis over any joint, should similarly be immobilized for comfort. Limiting early motion may also reduce edema and theoretically improve the immune system’s ability to combat infection. Hence, selected puncture wounds and mammalian or human bites of the hands and feet may be immobilized until the risk for infection has passed. Splinting large abrasions that cross joint surfaces prevents movement of the injured extremity and reduces the pain produced when the injured skin is stretched. Finally, victims of multiple trauma should have fractures and reduced dislocations adequately splinted while other diagnostic and therapeutic procedures (e.g., fluid resuscitation, airway control, computed tomography [CT] scans, tube thoracostomy) are completed. Immobilization decreases blood loss, minimizes the potential for further neurovascular injury, reduces the need for opioid analgesia, and may decrease the risk for fat emboli from long-bone fractures.

![Figure 50.1 Indications for splinting. Splinting is traditionally thought of as a treatment of fractures, such as the comminuted distal radius fracture depicted in A. However, splinting is also beneficial in a wide variety of other conditions, such as B, acute gout, C, human and mammalian bites of the hand, and D, tendon injuries. Other indications include inflammatory disorders such as tenosynovitis, deep lacerations that cross joints, and deep space infections of the hands and feet.](image-url)
Support Materials

Plaster of Paris
Plaster of Paris is the most widely used material for ED splinting. Its name originated from the fact that it was first prepared from the gypsum of Paris, France. When gypsum is heated to approximately 128°C, most of the water of crystallization is driven off and a fine white powder is left behind—plaster of Paris. When water is added to the plaster, the reaction is reversed, and the plaster recrystallizes or “sets” by incorporating water molecules into the crystalline lattice of the calcium sulfate hydrate molecules.

Today, plaster is impregnated into strips or rolls of a crinoline-type material. The crinoline allows easy application, helps keep the plaster molded to the proper form during the setting process, and adds support to the finished splint. Plaster rolls and sheets are available in a variety of setting times and widths (2-, 3-, 4-, or 6-inch widths). The distinct advantage of plaster over commercially available premade splints is that plaster can more easily be molded and tailored to the individual’s anatomy, thereby negating the “one-size-fits-all” approach. In addition, plaster rolls and strips are generally less expensive than premade splints.

Prefabricated Splint Rolls
The use of plaster splints in the form of prefabricated splint rolls (e.g., OCL, BSN Medical, Hamburg, Germany) is very popular among emergency clinicians. These splint rolls have 10 to 20 sheets of plaster enclosed between a thick layer of protective foam padding on one side and a thin layer of cloth on the other. Like custom-constructed splints, they are secured to the extremity with an elastic bandage. The major advantage of prefabricated splint rolls is that significant time is saved because the splint and padding come ready to apply. In addition, prefabricated splint rolls are ideal for intermittent splinting and can be removed and reapplied by the patient as needed. However, prefabricated plaster splint rolls are more expensive than simple plaster rolls, and they lack some of the versatility and custom-fit qualities of self-made plaster splints.

Prefabricated splint rolls composed of layers of fiberglass between polypropylene padding (e.g., Ortho-Glass [BSN Medical], 3M Scotchcast One-Step [3M St. Paul, MN]) are now commonplace in many EDs. Fiberglass splint rolls offer the same time-saving aspect as prefabricated plaster splint rolls but require only 3 minutes to set, thus making application faster. In addition, splints made of prefabricated fiberglass rolls cure more rapidly (20 minutes), have no messy residue (i.e., they can be hydrated in a conventional sink without a special trap), can be washed and reapplied, and are stronger and lighter than splints constructed of prefabricated plaster rolls. Another advantage is the polypropylene padding, which wicks moisture away from the skin better than polyester, nylon, or cotton padding does. Prefabricated fiberglass splint rolls are more expensive than both simple plaster rolls and prefabricated plaster splint rolls and, like prefabricated plaster splints, lack some of the versatility and custom-fit qualities of self-made plaster splints.

Woodcast Composite Splinting Material
A new synthetic material that has had a limited presence in the United States, but looks promising in European studies is Woodcast Composite Splinting Material. Woodcast splints are made from spruce tree chips and biodegradable plastic, making it nontoxic and environmentally friendly. The material comes in slabs ranging from 1-to-4 mm thick from which the provider can cut out the desired shape for a specific splint. Woodcast splints are applied in the same manner as conventional splints (Fig. 50.2) and have comparable stiffness. Like splints made from plaster of Paris and fiberglass, increasing the thickness of the splint increases its ultimate strength. The composite material is easily molded to the contour of an extremity or digit by heating for 15 minutes to a temperature of 65°C using one of the Woodcast heating devices (Fig. 50.3). The material takes approximately 15 minutes to set, providing sufficient time for applying and molding the splint. If needed, setting time can be reduced to 5 to 10 minutes with application of ice packs. Woodcast splints can be rewarmed and remolded after application and gloves are not required when handling the material. Compared to conventional splinting materials, Woodcast splints cause fewer skin reactions, and are associated with similar rates of healing and patient satisfaction.

Protective and Miscellaneous Equipment

Stockinette
A single layer of stockinette is commonly used over the skin and under cotton (Webril, Medtronic, Minneapolis, MN) padding, circumferential casts, and splints. It protects the skin and, when folded back over the ends of the plaster, creates a smooth, professional-looking, padded rim. Stockinette is available in 2-, 3-, 4-, 8-, 10-, and 12-inch widths.

Padding
Padding under the splint protects the skin and bony prominences and accommodates swelling of the injured extremity. Most commercially available splints contain adequate padding in the premade product, but in some instances, additional padding is prudent. In general, the older thin cotton padding known as sheet wadding has been replaced by newer material such as Webril (Curity) and Specialist (Johnson & Johnson, New Brunswick, NJ) cast padding. Webril is soft cotton with a much coarser weave than sheet wadding; consequently, it has greater tensile strength, adheres better, and can be applied more evenly. Specialist padding uses micropleated cotton fibers...
that relax when moistened. This results in uniform, felt-like padding that conforms to the surface being wrapped. Felt (0.5-inch thick) may also be used to pad bony prominences.

**Elastic Bandages**

Elastic bandages are used to secure the splint in place. Elastic bandages are available in 2-, 3-, 4-, and 6-inch widths. Some bandages use metal clips, whereas others use a Velcro-type mechanism to secure the bandage in place. Metal clips should be taped in place to avoid inadvertent removal.

Patients often use or request an elastic bandage for many soft tissue injuries. Although applying an elastic bandage to an injured part is popular, it is of minimal benefit alone. The downside is that the bandage may be wrapped too tightly and cause additional injury or distal swelling (Fig. 50.4).

**Adhesive Tape**

Use adhesive tape to prevent slippage of the elastic bandages, to line the cut edges of a bivalved cast, and to buddy-tape digits. Coban tape can be used in a similar manner and has the advantage of adhering only to itself.

**Utility Knife, Scalpel, and Plaster or Trauma Scissors**

Use a utility knife, a No. 10 scalpel blade, or plaster or trauma scissors to cut and shape dry plaster.

**Bucket**

Use a large bucket (preferably stainless steel) for wetting plaster. Do not prepare the plaster in the sink unless it is equipped with a special drain designed to accept plaster residue without clogging. A bucket is not required for the minimal amount of water used to soften premade fiberglass splints. They can be activated by placing them directly under the faucet.

**Protective Gear**

Use gowns or sheets to prevent soilage of both the patient’s and the clinician’s clothing. Use gloves (vinyl or latex) and safety glasses to prevent skin or eye damage from plaster dust, wet plaster, or uncured fiberglass polymer. Wearing gloves also decreases clean-up time for the clinician.

**GENERAL PROCEDURE OF CUSTOM SPLINT APPLICATION**

This following text refers to the application of custom-made plaster splints unless otherwise stated (Figs. 50.5 and 50.6). If periodic wound care is required, apply a more easily removable splint (e.g., OCL [BSN Medical], Ortho-Glass [BSN Medical], Velcro-type splint [described later]) in lieu of the standard splint (Fig. 50.7). Address the issue of removability before the
Plaster Splint Application: Standard Method

1. Measure the length of plaster to be used against the area to be splinted. Roll out the appropriate layers of plaster (roughly 8 layers for upper extremities, 12–15 for the lower).

2. Place a single layer of stockinette over the extremity. The stockinette should extend 10–15 cm beyond both ends of the area to be splinted. Generally, use a 3-inch stockinette for the arm, and a 4-inch for the leg.

3. Wrap 2 to 3 layers of Webril around the entire area to be splinted, overlapping each pass by 25 to 50%. Avoid wrinkling, which may cause pressure sores. See text for precautions regarding potential ischemic injury.

4. Submerge the plaster strips in a bucket of water until the bubbling stops. Do not use water hotter than 24°C (75°F). Using warmer water may cause thermal injury, because the plaster releases heat when activated.

5. Smooth the plaster between your fingers and remove all excess water. Lay the plaster out on a table and smooth further to remove all wrinkles and ensure uniform lamination of all layers.

6. Apply the plaster over the Webril and smooth it over the extremity. Avoid using your fingertips, which may leave indentations in the plaster. A layer of Webril may be placed over the plaster to prevent it from sticking to the elastic wrap.

7. Fold the stockinette over the edge of the plaster and Webril. This helps to hold the splint in place, and provides a smooth padded edge.

8. Secure the splint to the extremity with an elastic wrap by proceeding in a distal-to-proximal fashion. Do not wrap the elastic bandage too tightly.

9. If metal clips are used to secure the elastic wrap, cover them with tape to prevent them from falling off. Alternatively, the wrap can simply be secured with tape.

10. Place the extremity in the desired position, and use the palms of your hands to mold the splint to the contour of the extremity. Again, avoid using your fingertips, which may leave indentations that result in pressure sores.

Figure 50.5 Plaster splint application: standard method (volar wrist splint depicted here).
Plaster Splint Application: Alternative Method

1. Measure the splint and apply stockinette as depicted in steps 1 and 2 of Fig. 50.3. Additionally, premeasure 5 to 6 layers of Webril of the same length as the plaster. Soak and prepare the plaster.

2. Position the extremity as desired, and then gently mold the splint using the palms of your hands.

3. Lay out 3 to 4 layers of Webril, which will serve as padding for the splint. Place the plaster (which has already been smoothed) on top of these layers of Webril.

4. Place an additional layer of Webril on top of the plaster, which will prevent it from sticking to the elastic wrap. Essentially, you are sandwiching the plaster between layers of Webril.

5. Apply the splint to the extremity. Enlist the help of an assistant to hold the splint in place.

6. Secure the splint to the extremity with elastic bandages by wrapping in a distal-to-proximal fashion. Remember to fold the stockinette over the edges of the plaster and Webril.

Position the extremity as desired, and then gently mold the splint using the palms of your hands.

Figure 50.6 Plaster splint application: alternative method (sugar-tong splint depicted here). This technique may be used if significant swelling is anticipated as Webril (the same diameter as the plaster) is placed directly over the wet plaster rather than wrapped circumferentially around the extremity.
Prefabricated Fiberglass Splint Application

1. Measure the length of splint to be used against the area to be splinted. These splints are available in a variety of lengths and widths.

2. Cut the splint to length using a pair of trauma shears. Round the corners of the cuts to avoid sharp edges of fiberglass.

3. Stretch the cotton padding several inches beyond the edge of the fiberglass on both ends of the splint to assure that there will be no fiberglass in contact with the skin.

4. Prefabricated fiberglass splints require minimal wetting. Simply open one end of the pouch and run tap water over the fiberglass. When water exits the bottom, it is ready to apply. Excess water may be wrung out.

5. Apply the splint to the extremity. If stockinette is used (optional though recommended), fold the edges of the stockinette over the splint. This provides comfort and also holds the splint in place during application.

6. Secure the splint to the extremity using elastic bandages wrapped in a distal-to-proximal fashion.

7. For splints that involve the hand, cut a hole in the center of the elastic bandage, in the loop that is in the region of the thumb.

8. Bring the thumb through the hole. This provides a perfect fit, reduces bunching of the elastic in the web space, and allows for more thumb mobility.

9. Place the extremity in the desired position, and mold the splint with your palms to the contours of the body. Avoid using fingertips, which may leave indentations in the fiberglass.

10. After splint application, observe for vascular compromise or increased patient discomfort. If either occur, loosen the elastic wrap. Never discharge a patient with increasing pain after splint application.

Figure 50.7 Prefabricated fiberglass splint application (volar wrist splint depicted here).
Box 50.1 Caveats for Proper ED Splinting

- Always use cool, clean water.
- Do not oversaturate the plaster splint. Minimal water is required for fiberglass splints.
- Make the splint smooth when placing it on the patient to avoid bumps and pressure points.
- Smooth and mold the splint without squeezing. Use the palms of the hands, not the fingers, to mold the splint to fit the contour of the body part.
- Place the padded side against the skin. Extra cotton padding is optimal.
- Simply roll elastic bandages over the extremity without undue tension.
- Protect or pad the edges.
- Leave the fingertips exposed to check for circulation and sensation.
- Keep the patient still until the splint has dried and hardened.
- The postsplint check includes function, arterial pulse, capillary refill, temperature of the skin, and sensation (FACTS).
- Emphasize and demonstrate splint elevation to the patient.
- Tape over the metal clips used to fasten the elastic bandage to keep them in place and avoid ingestion by a child.

ED, Emergency department.

Patient Preparation

If the clinical situation permits, cover the patient with a sheet or gown to protect clothing and the surrounding area from water and plaster. Nursing staff and housekeeping also appreciate this courtesy. Properly drape hallway patients if areas of the body are exposed. Carefully inspect and examine the involved extremity before splinting, and clearly document the presence of all skin lesions and soft tissue injuries. Clean, repair, and dress all wounds in the usual manner. When open fractures or joints are to be immobilized, cover the soft tissue defect with saline-moistened sterile gauze.

Padding

When the splint involves the digits, place padding between the fingers and toes to prevent maceration of the skin. This can be done with pieces of Webril or gauze cut to the appropriate length.

Following placement of padding between the fingers and toes in self-made splints, use a stockinette over the skin as the first protective layer (see Fig. 50.5, step 2). Extend the stockinette at least 10 to 15 cm beyond the area to be splinted at both ends of the extremity. Later, after the plaster has been applied, fold the stockinette back over the ends of the splint to create smooth, padded rims and to help hold the splint in place when applying elastic bandages (see Fig. 50.5, step 7). To avoid pressure damage, do not pull the stockinette too tightly over bony prominences such as the heel. In addition, prevent wrinkling over flexion creases by slitting and overlapping the stockinette at bony prominences. One may also use two separate pieces of stockinette (one at each end of the splint) to produce the smooth padded rims. As a general rule, use 3-inch-wide stockinette for the upper extremity and 4-inch-wide stockinette for the lower extremity.

After the stockinette has been properly positioned, wrap Webril around the entire area that will be exposed to plaster. Apply at least two or three layers of Webril, with each turn overlapping the previous turn by 25% to 50% of its width (see Fig. 50.5, step 3). Make sure that the Webril extends 2.5 to 5.0 cm beyond the ends of the splint so that it, too, can be folded back over the splint to help create smooth, well-padded edges. Place extra padding over areas of bony prominence, such as the radial condyle or the malleoli (Box 50.2). Although this can be done with Webril, Mother’s Cotton adds an additional measure of protection without the worry of wrinkling or ischemic injury. If significant swelling is anticipated, use three or four layers of Webril. Be careful to avoid wrinkling because this can result in significant skin pressure when a tight splint is worn for a long period. Prevent wrinkles by proportionately stretching or even tearing the side of the Webril that must wrap around the larger portion of an extremity. Joints that must be immobilized in a 90-degree position, such as the ankle, make continuous Webril wrapping difficult. To avoid wrinkles around the ankle, place the joint in the proper position before padding. Wrap the Webril around the malleolus and midtarsal regions first, and then cover the bare calcaneal region with overlapping vertical and horizontal Webril strips until the entire heel region is evenly padded. Use the same approach in similar areas such as the elbow. Choose a width of Webril appropriate for the extremity to be splinted. In general, use the 2-inch width for the hands and feet, the 3- or 4-inch width for the upper extremity, and the 4- or 6-inch width for the lower extremity.

A final caveat when using Webril is to be aware of the potential for ischemic injury. This rare complication is most likely to occur in an extremity that continues to have significant swelling after the patient is released from the ED. Ischemia may result because the concentrically placed Webril can become a constricting band. If this situation is anticipated, it can easily be prevented. Cut through the Webril along the side of the extremity that is opposite the plaster splint. Then, secure the splint to the extremity in the usual manner. Alternatively, place two or three layers of Webril (the same diameter as the plaster)
and the risk for serious burns.15 Table 50.1 lists the setting time produces less heat, thus reducing both patient discomfort molding the splint. Furthermore, plaster with a longer setting times, the most important variable affecting the rate of crystallization is water temperature. Warm water hardens the production process (Box 50.3). Given plaster with equal setting times, the most important variable affecting the rate of crystallization is water temperature. Warm water hardens a splint faster than cold water does and should not be used when extra time is needed for splint application.

Plaster Preparation
The choice of plaster setting time depends on the nature of the injury and the expertise of the clinician. Use extra-fast–setting plaster when rapid hardening is desired to help maintain alignment of an acutely reduced fracture. However, for the majority of ED splints, plaster with slower setting times (e.g., Specialist Plaster Bandage Fast Setting, BSN Medical) is recommended.14 Plaster that sets more slowly is easier for some clinicians to use because it affords more leeway in applying and molding the splint. Furthermore, plaster with a longer setting time produces less heat, thus reducing both patient discomfort and the risk for serious burns.15 Table 50.1 lists the setting times for commonly used plaster. These setting times can be adjusted by adding different substances to the plaster during the production process (Box 50.3). Given plaster with equal setting times, the most important variable affecting the rate of crystallization is water temperature. Warm water hardens a splint faster than cold water does and should not be used when extra time is needed for splint application.

The ideal length and width of plaster depend on the body part to be splinted and the degree of immobilization required. The best way to estimate length is to lay the dry splint next to the area to be splinted. It is best to use a generous length because wet plaster shrinks slightly from its dry length. In addition, if the wet splint is too long, the ends can be folded back easily. Plaster width varies according to the type of splint being made and the body part that is injured, but generally, it should be slightly greater than the diameter of the limb to be splinted. Specific recommendations regarding splint length and width are discussed in the sections describing individual splints.

The thickness of a splint depends on the size of the patient, the extremity that is injured, and the desired strength of the final product. An ankle splint may crack quickly and become useless if only eight layers are used, but this thickness may be ideal for a wrist splint. In general, it is best to use the minimum number of layers necessary to achieve adequate strength. Thicker splints are heavier and more uncomfortable. It is also important to note that plaster thickness is a major determinant of the amount of heat given off during the setting process. More than 12 sheets of plaster create an increased risk for significant burns, especially when using extra-fast–drying plaster, when using dipping water with a temperature higher than 24°C, or when a pillow is placed under or around the extremity for support during the setting process (Box 50.4). For an average-sized adult, splint the upper extremities with 12 to 15 sheets. Such layering usually provides the strength necessary for adequate immobilization while reducing the patient’s discomfort and the risk for significant burns. In a 136-kg (300-lb) patient, however, up to 20 layers may be required to make a durable ankle splint.

Keep the dipping water clean and fresh. Reusing water that has been used previously for wetting plaster increases the amount of heat given off during crystallization and causes the plaster to set more quickly. As a rule of thumb, keep the temperature of the water at approximately 24°C. This temperature allows a workable setting time and has not been associated with increased risk for significant burns. As the temperature of the dipping water approaches 40°C, the potential for serious burns increases, even with splint thicknesses consisting of fewer than 12 plies. It is interesting to note that water temperature has been shown to be only a minor consideration in heat production in some studies (see Box 50.4).

Splint Application (see Fig. 50.5)
Completely submerge the dry splint in the water bucket until the bubbling stops. Remove the splint and gently squeeze out excess water until the plaster has a wet and sloppy consistency. Place the splint on a hard table or countertop (a protective covering is recommended to prevent water or plaster damage) and smooth out the splint (with gloved hands) to remove any

<table>
<thead>
<tr>
<th>PLASTER</th>
<th>SETTING TIME (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fast drying</td>
<td>5–8</td>
</tr>
<tr>
<td>Extra-fast drying</td>
<td>2–4</td>
</tr>
</tbody>
</table>

**BOX 50.3** Effect of Water Temperature and Different Additives on the Setting Time of Plaster

<table>
<thead>
<tr>
<th>Accelerates Setting Time</th>
<th>Slows Setting Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reusing the dip water</td>
<td>Cool dip water</td>
</tr>
<tr>
<td>Higher dip water temperature</td>
<td></td>
</tr>
<tr>
<td>Salicylic acid</td>
<td>Glue</td>
</tr>
<tr>
<td>Zinc</td>
<td>Gum</td>
</tr>
<tr>
<td>Magnesium</td>
<td>Borax</td>
</tr>
<tr>
<td>Copper</td>
<td></td>
</tr>
<tr>
<td>Iron</td>
<td></td>
</tr>
<tr>
<td>Aluminum</td>
<td></td>
</tr>
<tr>
<td>Salt</td>
<td></td>
</tr>
<tr>
<td>Alum</td>
<td></td>
</tr>
</tbody>
</table>

**BOX 50.4** Variables That Increase Heat Production During Crystallization

**MAJOR**
- Increased splint thickness
- Setting time
- High dip water temperature
- Wrapping the extremity for support while drying

**MINOR**
- High humidity
- High ambient temperature
- Reusing the dip water

* Faster setting times produce more heat.
* Dip water temperature has been a minor determinant of heat production in some studies. Use room-temperature not hot water to make a splint.
wrinkles and ensure uniform lamination of all layers. Lamination helps increase the final strength of the splint. Position the splint over the Webril and gently smooth it over the extremity. Plaster is usually somewhat adherent to Webril, but an assistant may be required to hold the splint in place during positioning. Once the splint has been properly positioned over the extremity, fold back the underlying stockinet and Webril to help hold it in place. Secure the splint with an appropriately sized elastic bandage by wrapping in a distal-to-proximal direction. Finally, place the extremity in the desired position and mold the wet plaster to the contour of the extremity with only the palms of the hand. Finger indentations may cause ridges, which can produce pressure points.

Molding the wet splint to conform to the body’s anatomy is probably the most important, yet the most frequently overlooked step to ensure adequate immobilization. The act of molding may cause some pain, so be sure to forewarn the patient. Mold with the palm or flat side of the fingers to avoid putting ridges or indentations in the underlying plaster. Complete all manipulation of the wet plaster before it reaches a thick, creamy consistency. Any movement after this time, which is known as the critical period, results in an imperfect crystalline network of calcium sulfate molecules and greatly weakens the ultimate strength of the splint. While the plaster is setting, do not wrap a pillow or blanket around the extremity for support because this leads to inadequate ventilation around the splint and greatly increases the amount of heat produced (see Box 50.4).

If an elastic bandage is applied directly over wet plaster, it may be incorporated into the drying plaster, thus making subsequent removal of the bandage difficult. To make it easier for patients to remove and reaply the splint, wrap a single layer of Webril or roll gauze around the wet plaster loosely before applying the elastic bandage. This prevents the wet plaster from becoming stuck to the elastic bandage. Use only one layer of Webril over the plaster because multiple layers are associated with high drying temperatures.

Before the patient is released from the ED, check the extremity for adequate immobilization and evaluate the patient for any evidence of vascular compromise or significant discomfort. If either occurs, loosen the elastic bandage. If the discomfort persists, place additional padding over the painful areas. If this measure, too, is unsuccessful, make a new splint while paying special attention to proper molding so that the wet plaster does not become indented. By resting tender tissue, splinting usually relieves the discomfort quickly, and patients generally say that they feel better soon after the splint has been applied. In general, a splint should decrease the patient’s pain, not increase it; hence, do not readily release a patient who complains of increased pain after a splint has been placed. Such complaints may be due to manipulation during splinting, but increased pain should be further addressed or explained.

After a properly fitting, comfortable splint has been applied, place two strips of tape along each side of the splint to prevent the elastic bandage from slipping. It is also prudent to place tape over any metal fasteners used to secure the elastic bandages because these objects can fall off and be swallowed or aspirated by infants and small children. Use enough tape to include the entire circumference of the area under the fasteners, not just small pieces of tape that may not adhere to the moist splint. Finally, provide a sling for patients with upper extremity injuries and, if required, crutches (and instructions for their proper use) for those with lower extremity injuries.

**Patient Instructions**

Give patients both verbal and written instructions on splint care and precautions. Stress the importance of elevation in helping to decrease pain and swelling. Demonstrate this as well because most patients do not understand the medical definition of elevation. At night, wrap and secure a pillow around the injured extremity to help the patient keep it satisfactorily elevated. If the injury is less than 24 hours old, encourage the application of ice bags or cold packs. It is useless to apply cold packs over plaster, but it can be beneficial to apply them over Webril or an elastic bandage directly over an injury. It may be necessary to remove the splint to ice the injury. In theory, cold therapy stiffens collagen and thus reduces the tendency for ligaments and tendons to deform. Cold therapy also decreases muscle spasm and excitability, reduces blood flow (thereby limiting hemorrhage and edema), raises the pain threshold, and decreases inflammation. Because the thermal conductivity of subcutaneous tissue is poor, apply cold packs for at least 30 minutes at a time. This guideline is in contrast to the popular recommendation of “ice 20 minutes on, 20 minutes off,” which often does nothing more than cool the skin. Do not use cold packs for more than the first 24 to 48 hours because cold can interfere with long-term healing. Instruct the patient to not stress the splint for at least 24 hours because plaster does not approach optimal strength until evaporation has reduced the water content of the plaster to approximately 21% of its initial hydrated level. This process of removing excess water by evaporation is called curing, and it generally takes several days to be completed. However, by 24 hours the water content of the plaster has usually been reduced enough to produce a strong, resilient splint. In addition, because the chemical process involved in the formation of plaster is reversible, the patient should avoid getting the splint wet. If the injury permits, the splint can be removed for showering and then reapplied. Alternatively, one or more plastic bags may be placed over a splint before showering.

Splints may crack, break, or disintegrate with wear, and a useless splint should be removed or replaced. Give patients general guidelines for length of immobilization and appropriate follow-up care. Avoid long-term immobilization, particularly in the elderly, because this can produce permanent disability. It is extremely important for the patient to continue to check for signs of vascular compromise. If the patient experiences a significant increase in pain, numbness or tingling of the digits, pallor of the distal end of the extremity, decreased capillary refill, or weakness, instruct the patient to return to the ED or to see his or her primary clinician without delay. As with casting, increased pain after splinting is a warning sign that should prompt a return visit, not telephone advice. Avoid giving excessive doses of opioids during the first 2 to 3 days after splinting to allow pain to prompt a follow-up visit.

**UPPER EXTREMITY SPLINTS**

**Long Arm Splints**

**Long Arm Posterior Splint**

*Indications.* Use a long arm posterior splint (Figs. 50.8 and 50.9) to immobilize injuries to the elbow and proximal end of the forearm. It completely eliminates flexion and extension of the elbow but does not entirely prevent pronation and supination...
Splinting Techniques

CHAPTER 50

Long Arm Posterior Splint

Application

A. Extend the splint from the posterior aspect of the humerus to the elbow and then along the ulnar aspect of the forearm to the distal metacarpals. Flex the elbow to 90°, maintain the forearm in the neutral (thumb-up) position, and place the wrist in a neutral or slightly extended (10° to 20°) position.

B. An anterior splint that mirrors the posterior splint may be used to increase stability and prevent supination and pronation. An anterior splint is never used alone.

Indications

Injuries of the elbow and distal end of the forearm, including:
- Distal humerus fractures (shown above)
- Supracondylar fractures
- Olecranon fractures
- Elbow dislocations

Figure 50.8 Long arm posterior splint.

Figure 50.9 When preparing a splint (such as a long arm splint) that involves a right angle, cut out a notch (arrow) to allow a smooth bend. Note that padding needs to be applied before splinting.

Long Arm Anterior Splint

Indications

A long arm anterior splint is never used alone but, rather, as an adjunct to a long arm posterior splint to improve immobilization by increasing stability and preventing pronation and supination of the forearm (see Fig. 50.8B).

Construction

Construct a long arm posterior splint with 8 to 10 layers of 4- or 6-inch-wide plaster. Start the splint on the posterior aspect of the proximal end of the arm. Extend it down the arm to the elbow and then along the ulnar aspect of the forearm and hand to the level of the metacarpophalangeal (MCP) joints.

Application

Apply a stockinette and Webril as described previously. Cut a hole in the stockinette to expose the thumb, and place extra padding over the olecranon to prevent pressure injury. Position the arm with the elbow flexed to 90 degrees, the forearm neutral (thumb upward), and the wrist neutral or slightly extended (10 to 20 degrees). Ask an assistant to hold the wet splint in place, particularly when applying both a posterior and an anterior splint. Once the splint has been properly positioned, fold the ends of the stockinette and Webril back and secure the splint in place with 2-, 3-, or 4-inch elastic bandages. Finally, fold the sides of the splint up to create a gutter configuration and carefully mold the plaster around the extremity with the palms of the hand. The fingers and thumb should remain free to prevent stiffness from unnecessary immobilization.
Construction. Construct a long arm anterior splint in the same manner as described for a long arm posterior splint. It mirrors the posterior splint by running down the anterior aspect of the arm to the antecubital fossa, where it continues along the radial aspect of the forearm to the distal end of the radius.

Application. Use stockinette, Webril, and positioning similar to the way described for the application of a long arm posterior splint. When using both an anterior and a posterior long arm splint, have an assistant available to hold the wet splint in place. Place the anterior splint first and then position the posterior splint. Once both splints have been properly positioned, fold the ends of the stockinette and Webril back and secure the splint in place with 2-, 3-, or 4-inch elastic bandages. Finally, fold up the sides of the splint to create a gutter configuration and carefully mold the plaster around the extremity with the palms of the hands. Keep the patient’s fingers and thumb free to prevent stiffness from unnecessary immobilization.

Double Sugar-Tong Splint
Indications. Use a double sugar-tong splint (Fig. 50.10) like a long arm posterior splint to immobilize injuries to the elbow and forearm. However, because it prevents pronation and supination of the forearm, it is preferable for some distal forearm and elbow fractures.

Construction. The splint consists of two separate pieces of plaster, a forearm splint and an arm splint. Construct each piece with eight layers of 3- or 4-inch plaster. The forearm portion of the splint runs from the metacarpal heads on the dorsum of the hand along the dorsal surface of the forearm around the elbow. It continues along the volar surface of the forearm to the palm of the hand and stops at the level of the MCP joints. The arm portion of the splint begins on the anterior aspect of the proximal end of the humerus. It runs down the arm over the forearm splint and around the elbow. It then continues up the posterior aspect of the arm, once again going over the forearm splint, until it reaches the starting point.

Application. Use a stockinette, Webril, and positioning similar to the way described for the application of a long arm posterior splint. When using both an anterior and a posterior long arm splint, have an assistant available to hold the wet splint in place. Place the anterior splint first and then position the posterior splint. Once both splints have been properly positioned, fold the ends of the stockinette and Webril back and secure the splint in place with 2-, 3-, or 4-inch elastic bandages. Finally, fold up the sides of the splint to create a gutter configuration and carefully mold the plaster around the extremity with the palms of the hands. Keep the patient’s fingers and thumb free to prevent stiffness from unnecessary immobilization.

Double Sugar-Tong Splint
Indications. Use a double sugar-tong splint (Fig. 50.10) like a long arm posterior splint to immobilize injuries to the elbow and forearm. However, because it prevents pronation and supination of the forearm, it is preferable for some distal forearm and elbow fractures.

Construction. The splint consists of two separate pieces of plaster, a forearm splint and an arm splint. Construct each piece with eight layers of 3- or 4-inch plaster. The forearm portion of the splint runs from the metacarpal heads on the dorsum of the hand along the dorsal surface of the forearm around the elbow. It continues along the volar surface of the forearm to the palm of the hand and stops at the level of the MCP joints. The arm portion of the splint begins on the anterior aspect of the proximal end of the humerus. It runs down the arm over the forearm splint and around the elbow. It then continues up the posterior aspect of the arm, once again going over the forearm splint, until it reaches the starting point.

Application. Use a stockinette, Webril, and positioning similar to the way described for the application of a long arm posterior splint. When using both an anterior and a posterior long arm splint, have an assistant available to hold the wet splint in place. Place the anterior splint first and then position the posterior splint. Once both splints have been properly positioned, fold the ends of the stockinette and Webril back and secure the splint in place with 2-, 3-, or 4-inch elastic bandages. Finally, fold up the sides of the splint to create a gutter configuration and carefully mold the plaster around the extremity with the palms of the hands. Keep the patient’s fingers and thumb free to prevent stiffness from unnecessary immobilization.

**Figure 50.10** Double sugar-tong splint.
If the splint is going to incorporate one or more digits, insert a piece of Webril or gauze between the digits to prevent skin maceration. Place the forearm in the neutral position (thumb upward) with the wrist extended slightly (10 to 20 degrees). Avoid wrist flexion. After properly positioning the wet plaster, fold back the ends of the stockinette and Webril and use a 3- or 4-inch elastic bandage to secure the splint in place. Fold the sides of the splint up around the forearm to create a gutter effect, and carefully mold the plaster to conform to the contours of the palm and wrist. Some clinicians prefer to extend the splint to the fingertips and then fold the wet plaster back toward the palm, which allows the fingers to “grasp” the rounded distal end when at rest. With either method, keep the thumb and fingers free to move unless they are injured and are being intentionally immobilized by the splint.

**Sugar-Tong Splint**

**Indications.** Use a sugar-tong splint (Fig. 50.12) for fractures of the distal end of the radius and ulna. The advantage of this splint over a volar splint is prevention of pronation and supination of the forearm. In addition, it immobilizes the elbow, which is desirable for the first few days after a distal forearm fracture.

**Construction and Application.** Construct and apply this splint in the same manner as the forearm portion of the double sugar-tong splint described earlier.
**Thumb Spica Splint**

**Indications.** Use a thumb spica splint (Fig. 50.13) to immobilize injuries and fractures of the scaphoid, lunate, thumb, and first metacarpal. It is also used for the treatment of de Quervain's tenosynovitis. Traditionally, a thumb spica splint or cast was thought to be a requirement for properly immobilizing scaphoid fractures; however, there is no totally agreed standard. Clay and coworkers\(^1\) stated that the optimal method of casting scaphoid fractures has not been definitively established. They were unable to prove a difference in patient comfort, recovery of function, or incidence of nonunion between a Colles cast and a traditional scaphoid cast that included the thumb.

The incidence of nonunion of scaphoid fractures is approximately 10%, regardless of the type of immobilization in the ED, but it is greatest with unstable proximal pole fractures. Because some scaphoid fractures heal poorly under the best of circumstances, it seems prudent to provide thumb immobilization in the initial splinting. Failure to do so, such as when a “sprained wrist” is suspected, should not be construed as being beneath the accepted standard of care. Most volar splints will at least partly immobilize the base of the thumb, so the discussion may be moot.

**Construction.** Construct the splint with eight layers of 3-inch-wide plaster. Extend the splint from just distal to the interphalangeal joint of the thumb to the midforearm level.

**Application.** Place the forearm in the neutral position with the wrist extended 25 degrees and the thumb in the wineglass position (Fig. 50.14). Apply a stockinette and Webril from the base of the palm to the midforearm level. It is difficult to place a stockinette around the thumb. Instead, cut a hole in the stockinette to expose the thumb, and then pad the exposed thumb with small vertical strips of Webril or wrap it with 2-inch Webril. Place the dry plaster over the radial aspect of the forearm from just beyond the thumb interphalangeal joint to the midforearm level. Once in position, mark the location of the first MCP joint and make a small (1 to 2 cm) perpendicular cut 1 cm distal to the mark on each edge of the plaster (see Fig. 50.13 inset). This allows the splint to be molded around the thumb without creating a buckle in the plaster. If the plaster distal to the cut notches is too wide to mold around the thumb without overlapping, trim the edges to the desired width. Wet the plaster and secure it in place with a 2- or 3-inch elastic bandage. It is important to carefully mold the wet plaster around the thumb and palm and to maintain the thumb in the wineglass position while the plaster is drying.

**Ulnar Collateral Ligament Injury**

(Gamekeeper’s or Skier’s Thumb)

Forced abduction with hyperextension of the thumb is the usual mechanism for injury to the ulnar collateral ligament. This is a rather disabling injury that requires early recognition because improper treatment can result in chronic pain, pinch
neutral position with the wrist in slight extension (10 to 20 degrees), the MCP joints in 50 degrees of flexion, and the proximal and distal interphalangeal joints in slight flexion (10 to 15 degrees). When immobilizing a metacarpal neck fracture (i.e., boxer’s fracture), flex the MCP joint to 90 degrees. Once in proper position, fold the sides of the splint up to form a gutter. Finally, fold the ends of the stockinette and Webril back to help hold the splint in place while it is secured to the extremity with a 2- or 3-inch elastic bandage.

Radial Gutter Splint

**Indications.** Use an ulnar gutter splint (Fig. 50.16) to immobilize fractures and serious soft tissue injuries of the little and ring fingers and fractures of the neck, shaft, and base of the fourth and fifth metacarpals.

**Construction.** Make the splint with six to eight layers of 3- or 4-inch plaster. It incorporates both the little and the ring fingers. It runs along the ulnar aspect of the forearm from just beyond the distal interphalangeal joint of the little finger to the midforearm level.

**Application.** Apply a stockinette and Webril as usual. Place additional Webril or gauze between the little and ring fingers to prevent maceration of the skin. Position the forearm in the neutral position with the wrist in slight extension (10 to 20 degrees), the MCP joints in 50 degrees of flexion, and the proximal and distal interphalangeal joints in slight flexion (10 to 15 degrees). When immobilizing a metacarpal neck fracture (i.e., boxer’s fracture), flex the MCP joint to 90 degrees. Once in proper position, fold the sides of the splint up to form a gutter. Finally, fold the ends of the stockinette and Webril back to help hold the splint in place while it is secured to the extremity with a 2- or 3-inch elastic bandage.

**Ulnar Gutter Splint**

**Indications.** Use an ulnar gutter splint (Fig. 50.16) to immobilize fractures and serious soft tissue injuries of the little and ring fingers and fractures of the neck, shaft, and base of the fourth and fifth metacarpals.

**Construction.** Make the splint with six to eight layers of 3- or 4-inch plaster. It incorporates both the little and the ring fingers. It runs along the ulnar aspect of the forearm from just beyond the distal interphalangeal joint of the little finger to the midforearm level.

**Application.** Apply a stockinette and Webril as usual. Place additional Webril or gauze between the little and ring fingers to prevent maceration of the skin. Position the forearm in the neutral position with the wrist in slight extension (10 to 20 degrees), the MCP joints in 50 degrees of flexion, and the proximal and distal interphalangeal joints in slight flexion (10 to 15 degrees). When immobilizing a metacarpal neck fracture (i.e., boxer’s fracture), flex the MCP joint to 90 degrees. Once in proper position, fold the sides of the splint up to form a gutter. Finally, fold the ends of the stockinette and Webril back to help hold the splint in place while it is secured to the extremity with a 2- or 3-inch elastic bandage.

**Figure 50.13** Thumb spica splint.
and fingers in the position of function or in the intrinsic position (see Fig. 50.14). Neither position has been proved to be superior for the first few weeks of splinting, which makes them both acceptable for initial immobilization in the ED. In the position of function, the forearm is in the neutral position with the wrist in slight extension (10 to 20 degrees), the MCP joints in 50 to 60 degrees of flexion, and the proximal and distal interphalangeal joints in slight flexion (5 to 10 degrees) (see Fig. 50.14A). When immobilizing a metacarpal neck fracture, the intrinsic position is often used, with the MCP joint flexed to 90 degrees and the fingers extended (see Fig. 50.14B).

Place the dry plaster over the extremity and mark the location of the thumb. Cut a hole in the dry plaster to expose the thumb. Dip the plaster and position the wet splint over the extremity. Fold back the ends of the stockinette and Webril to help hold the splint in place and secure it to the extremity with a 2- or 3-inch elastic bandage.

**Figure Splints**

Use finger splints for sprains, fractures, tendon repairs, or infections. Minor finger sprains can often be managed with dynamic splinting (e.g., buddy taping) (Fig. 50.18A) or a commercially available foam splint with aluminum backing (one-surface splint) (see Fig. 50.18B), but fractures, tendon repairs, and some soft tissue injuries benefit from formal splinting (e.g., thumb spica and ulnar and radial gutter splints). Specific conditions, such as mallet finger, require a specialized splint (plaster or Stack splint) (see Fig. 50.18C). When complete immobilization of a finger is required (e.g., unstable phalangeal fractures), an "outrigger" finger splint that incorporates the wrist may be used (see Fig. 50.18D). Both the position of function and the intrinsic position are acceptable for initial splinting.

**Pitfalls of Hand Dressings and Splints**

The two most common problems with hand dressings and splints are putting them on too tightly and leaving them on too long (Table 50.2). One must be especially careful to avoid wrapping elastic bandages too snugly. Before discharge from the ED, instruct patients to loosen an elastic bandage if it feels too tight, and be sure that they have access to emergency follow-up care. It is often advisable to start patients on a regimen of early protected motion. This means that the patient removes the splint for a specified period, performs a prescribed exercise, and then replaces the splint. A splint is not an all-or-none device, and the patient is generally weaned from it slowly before it is discarded entirely. A stiff hand is a nonfunctional one, and stiffness is often a consequence of prolonged immobilization. It is important for patients to be made aware of their responsibility for the injured hand.
**Ulnar Gutter Splint**

**Application**
The ulnar gutter splint incorporates both the little and the ring fingers. Therefore place Webril or gauze between the digits to prevent maceration of the skin. Run the splint along the ulnar aspect of the forearm from just beyond the distal interphalangeal joint of the little finger to the midforearm level.

Maintain the forearm in the neutral position with the wrist in slight extension (10° to 20°), the metacarpophalangeal (MCP) joint in 50° of flexion, and the proximal and distal interphalangeal joints in slight flexion (10° to 15°). When immobilizing a metacarpal neck fracture, flex the MCP joint to 90°.

**Indications**
Injuries of the ulnar side of the hand, including:
- Fourth and fifth metacarpal fractures (shown above)
- Serious injuries of the ring and little fingers

---

**Sling, Swathe and Sling, and Shoulder Immobilizer**

**Sling**
Use a sling to maintain elevation and provide immobilization of the hand, forearm, and elbow (Fig. 50.19). It is most often used in conjunction with a plaster splint or cast. A number of commercial slings are available to choose from. Many of them are fairly economical and simple to use, whereas others are more expensive and do not allow the versatility of a simple, inexpensive triangular muslin bandage. When applying a sling, make it long enough to adequately support the wrist and hand. A sling that is too short will allow the wrist and hand to hang down (ulnar deviate) and can result in ulnar nerve injury.

**Swathe and Sling**
Use of a swathe and sling is the treatment of choice for most proximal humeral fractures and shoulder injuries, such as reduced dislocations. The sling supports the weight of the arm, and the swathe immobilizes the arm against the chest wall to minimize shoulder motion.

**Shoulder Immobilizer**
In most EDs the swathe and sling have been replaced by commercially available shoulder immobilizers. Its advantage is that it may be removed for showering and range-of-motion exercises and is easily reapplied by the patient (a desirable option in the care of a shoulder dislocation). If the shoulder immobilizer is used for more than a few days, pad the axilla to absorb moisture and decrease skin chafing.

A Velpeau bandage is a sling and swathe device that positions the forearm diagonally rather than horizontally across the chest with the hand elevated to the level of the shoulder. It offers no particular advantage over a standard sling and swathe, is difficult to apply, cannot be removed easily, and is not well tolerated with prolonged immobilization.

**Figure-of-Eight Clavicle Strap**
In the past, clavicle fractures were often treated with an uncomfortable and complex figure-of-eight bandage. Despite its early popularity, this device never proved to be superior to a simple sling (in terms of cosmesis, functional outcome, or pain relief). Moreover, use of a figure-of-eight clavicle strap may actually promote nonunion or increase the deformity at the fracture site. When compared with a simple sling, a figure-of-eight clavicle strap is very uncomfortable, prohibits bathing, often causes chafing and discomfort in the axilla, and may predispose to axillary vein thrombosis. Thus, most emergency clinicians have abandoned the figure-of-eight dressing in favor...
of a simple sling, which is sufficient to treat most clavicular fractures.

LOWER EXTREMITY SPLINTS

Knee Splints

Knee Immobilizer

Indications. A knee immobilizer (Fig. 50.20) is commonly used for mild to moderate ligamentous and soft tissue injuries involving the knee. It is removable and extremely easy to apply, which makes it popular among patients and clinicians alike. In most EDs it has replaced the bulkier plaster splint. Its use should be restricted to injuries that do not require immediate surgical intervention, traction, or casting. For these injuries, in which temporary but more complete immobilization is needed, use a plaster knee splint because it provides better stabilization and costs much less than a knee immobilizer. The exact scientific benefit of a knee immobilizer is poorly studied and difficult to document. However, it clearly helps relieve pain and, at least theoretically, hastens healing.

Application. Knee immobilizers are available in small, medium, large, and extra-large sizes. To choose the appropriate size, place the knee immobilizer next to the injured leg so that the tapered end lies distal to the patient’s knee; if present, the cutout patellar area on the anterior aspect of the splint lies adjacent to the knee joint. In this position, the splint should extend distally to within a few inches of the malleoli and proximally to just below the buttocks crease. To apply the knee immobilizer, slide the open splint under the injured leg, wrap it around the extremity, and secure it in place with the Velcro straps. A knee immobilizer can be applied directly over clothing, thus obviating the need to remove or cut the patient’s pants.

Posterior Knee Splint

Indications. In many EDs the knee immobilizer has virtually replaced the plaster knee splint for mild to moderate injuries to the knee. However, a plaster knee splint can be particularly useful in patients whose extremities are too large for a knee immobilizer, in the treatment of angulated fractures, or for temporarily immobilizing injuries that require immediate operative intervention or orthopedic referral. The posterior (gutter) knee splint (Fig. 50.21A) is the type most commonly applied, but as alternatives, two parallel splints can be placed along each side of the leg and foreleg to create a bivalve effect (see Fig. 50.21B), or a long leg U-splint can be applied (see later). A bilateral knee splint is slightly more difficult to apply.
**Finger Splinting Techniques**

**A. Buddy Taping**
- Webril or gauze padding
- Half-inch adhesive tape

Taping between the digital joints (toes or fingers) allows the normal adjacent finger to protect the collateral ligament of its injured neighbor. Place Webril between the digits to prevent maceration of the skin.

**C. Mallet Finger Splints**

**B. Dorsal Aluminum Foam Splint**

The bone is subcutaneous dorsally, and splints here afford better immobilization of the digit. The dorsal splint also allows preservation and use of tactile sense, which encourages function and better splint acceptance on the part of the patient.

**D. “Outrigger” Finger Splint**
- Foam finger splint with aluminum backing
- Finger splint sandwiched between layers of plaster

A padded aluminum splint is incorporated into the middle of a plaster splint to form an outrigger configuration. The plaster splint is applied to the dorsum of the hand and wrist with an elastic bandage; the finger is then taped to the aluminum splint. This provides complete immobilization of the finger.

*Top:* The dorsal splint immobilizes only the distal interphalangeal joint, which allows use of the finger. Hyperextension of this joint predisposes to skin sloughing and should be avoided. The patient should be advised to not flex the joint during splint changes.

*Bottom:* A Stack splint is designed especially to treat a mallet finger. Long-term immobilization (8 weeks) or surgical fixation is required for this injury.

*Figure 50.18* Finger splinting techniques.
than a posterior knee splint, but it may provide better immobilization of the lateral and medial collateral ligaments and can be used for injuries to these structures.

**Construction.** Construct a posterior knee splint with 12 to 15 layers of 6-inch plaster. It should run from just below the buttocks crease to approximately 5 to 8 cm above the malleoli. The sides of the splint are folded upward to form a gutter configuration.

**Application.** Apply a stockinette in the usual manner, and wrap the leg with 4- or 6-inch Webril. Have an assistant elevate the leg and hold the splint in place while securing it to the extremity with 4- or 6-inch elastic bandages. If no aide is available, place the patient in the prone position and have him use his toes to elevate the lower part of the leg off the bed to allow sufficient room to wrap the Webril and elastic bandages around the injured extremity. Lay the splint along the posterior surface of the extremity, and secure it in place with 4- or 6-inch elastic bandages.

### Jones Compression Dressing

**Indications.** A Jones compression dressing is commonly used for short-term immobilization of soft tissue injuries of the knee. It immobilizes and compresses the knee, thereby reducing both pain and swelling. However, because it does allow slight flexion and extension of the knee, it should not be used for injuries that require strict immobilization. In addition, it is difficult to maintain the splint for more than a few days.

**Construction.** A Jones dressing is fashioned from 6-inch Webril and elastic bandages.

**Application.** To apply a Jones dressing, place the patient supine on the stretcher. If available, ask an assistant to elevate the patient’s leg to facilitate wrapping. If no help is available, place a pillow under the patient’s heel to lift the extremity off the stretcher. Wrap Webril around the extremity from the groin to a few inches above the malleoli. Use two or three layers of Webril and overlap the previous turn by 25% to 50%. Complete the dressing by wrapping 6-inch elastic bandages (two are usually required) around the Webril. If more support is required, repeat the process with another two or three layers of Webril held in place by additional elastic bandages.

### Ankle Splints

#### Posterior Splint

**Indications.** A posterior ankle splint (Fig. 50.22) is one of the most common splints applied in the ED. As noted in the introductory section, the entire concept of splinting an acutely sprained ankle has been questioned, with no firm evidence to support better outcomes with splinting or casting versus functional management (early mobilization with an external support). Nonetheless, an acutely sprained ankle is painful, and if nothing else, splinting for a few days helps alleviate the pain.

Use a posterior splint primarily to immobilize severe ankle sprains, fractures of the distal ends of the fibula and tibia, and reduce ankle dislocations. It can also be used for fractures of the tarsal and metatarsal bones or for other foot conditions that require immobilization. With particularly severe or unstable injuries, an additional anterior splint may be used to provide extra immobilization resembling that of a formal cast (Fig. 50.23). For severe lateral or bilateral ligamentous injuries, add a U-splint or stirrup splint (see later) to the posterior splint for increased immobilization. With minor soft tissue injuries, patients may have partial weight bearing on ankle splints after 24 hours. If the patient will be bearing weight, wearing a cast shoe over the splint makes it easier to walk and increases the longevity of the splint. Generally, though, walking on the splint is prohibited if immobilization for more than 2 or 3 days is desired.

**Construction.** Make the posterior splint with 4- or 6-inch-wide plaster strips. It should extend from the plantar surface of the great toe or metatarsal heads along the posterior surface of the foreleg to the level of the fibular head. If it hurts to move the toes, incorporate them into the splint (after padding is placed between the digits). It is a common mistake to apply a posterior splint that does not extend far enough to support the ball of the foot. Use 15 to 20 layers if partial weight bearing is allowed because this splint frequently breaks or cracks when walked on.19

### TABLE 50.2 Useful Estimates of Splint Times for Various Hand Problems

<table>
<thead>
<tr>
<th>INJURY</th>
<th>SPLINT TYPE</th>
<th>IMMOBILIZATION TIME</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mallet finger</td>
<td>FIN</td>
<td>8 wk</td>
</tr>
<tr>
<td>Boutonnière deformity</td>
<td>FIN</td>
<td>6 wk</td>
</tr>
<tr>
<td>Distal phalanx—soft tissue</td>
<td>FIN</td>
<td>1–2 wk</td>
</tr>
<tr>
<td>Extensor tendon</td>
<td>DHWF</td>
<td>3 wk</td>
</tr>
<tr>
<td>Sprain-strainb</td>
<td>Interphalangeal joint FIN</td>
<td>1–2 wk</td>
</tr>
<tr>
<td></td>
<td>Wrist</td>
<td>DHWF</td>
</tr>
<tr>
<td>Hand burn</td>
<td>DHWF</td>
<td>5–7 wk</td>
</tr>
<tr>
<td>Infection</td>
<td>DHWF</td>
<td>5–7 wk</td>
</tr>
<tr>
<td>Digit</td>
<td>DHWF</td>
<td>5–7 wk</td>
</tr>
<tr>
<td>Wrist</td>
<td>DHWF</td>
<td>5–7 wk</td>
</tr>
<tr>
<td>Severe hand contusion</td>
<td>DHWF</td>
<td>5–7 wk</td>
</tr>
<tr>
<td>Fracture</td>
<td>Distal phalanx FIN</td>
<td>2–3 wk</td>
</tr>
<tr>
<td></td>
<td>Middle phalanx FIN</td>
<td>2–3 wk</td>
</tr>
<tr>
<td>Proximal phalanx</td>
<td>DHWF</td>
<td>2–3 wk</td>
</tr>
<tr>
<td>Metacarpal</td>
<td>DHWF</td>
<td>2–3 wk</td>
</tr>
<tr>
<td>Carpal tunnel</td>
<td>DHWF</td>
<td>Night only</td>
</tr>
<tr>
<td>de Quervain’s disease</td>
<td>DHWF</td>
<td>2–3 wk</td>
</tr>
<tr>
<td>Trigger finger</td>
<td>FIN</td>
<td>Night only</td>
</tr>
</tbody>
</table>

*These are average times only. Every patient is treated as an individual when a splint is used. Clinical judgment is critical.

*Diagnosis of a sprain should be made only after a thorough effort has been made to rule out a fracture or dislocation. This is particularly true in the wrist.

DHWF, Digit-hand-wrist-forearm; FIN, finger.
Application. The easiest way to apply a posterior splint is to place the patient in the prone position with the knee and ankle flexed at a 90-degree angle. Failure to place the ankle in a 90-degree angle results in a plantar-flexed splint. A supine patient may help maintain the ankle in a 90-degree angle by pulling up on the foot with a wide stockinette stirrup. Flexing the knee to a 90-degree angle relaxes the gastrocnemius muscle and facilitates ankle motion. With the knee and ankle in the proper position, apply a stockinette and pad the foot and leg with Webril as described earlier. Use extra padding over bony

Shoulder Slings

The shoulder immobilizer is used for most proximal humerus fractures and shoulder injuries. It may be removed for showering and range-of-motion exercises and is easily reapplied by the patient.

An elastic bandage and sling provide similar shoulder immobilization. Note that the wrist is supported by the slings.

Proximal humerus fractures are common injuries in the elderly, especially after a fall. A shoulder sling is the immobilization device of choice.

Figure 50.19 Shoulder slings.
Knee Immobilizer

**Application**
Choose the appropriate-size knee immobilizer for the patient. The tapered end should be distal to the knee. The splints simply wrap around the leg (and can be applied over clothing) and are secured with Velcro straps.

**Indications**
Ligamentous and soft tissue injuries of the knee.

Above, a *Segond fracture* is demonstrated. Note the small bony avulsion fracture on the lateral aspect of the tibial plateau. Although at first glance this appears to be a rather innocuous injury, a Segond fracture is highly associated with tears of the anterior cruciate ligament. This patient should be treated with a knee immobilizer until orthopedic follow-up can be arranged.

*Figure 50.20* Knee immobilizer.

Posterior Knee Splint

**Application**
A, Extend the posterior knee splint from just below the buttocks crease to approximately 2 to 3 cm above the malleoli.
B, Alternatively, place two parallel splints along each side of the leg and foreleg to create a bivalve effect. Although this variation is more difficult to apply, it may provide better immobilization of the lateral and medial collateral ligaments.

**Indications**
Angulated fractures around the knee (shown above) Temporary immobilization of injuries prior to operative repair Knee injuries in patients with extremities too large for a knee immobilizer

*Figure 50.21* Posterior knee splint.
Anterior-Posterior Splint

**Indications.** An anterior splint is never used by itself, but it can augment a posterior splint to create a bivalve effect (see Fig. 50.23). Use it for serious fractures and soft tissue injuries of the ankle.

**Construction.** Cut a piece of plaster several centimeters shorter than the one used for the posterior splint, but as this splint does not bear weight, only 8 to 10 layers are required.
Anterior-Posterior Ankle Splint

Application
Apply the posterior splint as described in Fig. 50.20.
For the anterior portion, begin on the dorsal surface of the foot at the level of the metatarsophalangeal joints. Extend the splint along the anterior portion of the foreleg to the same height as the posterior splint. Maintain the ankle at a 90° angle.

Indications
Serious fractures and soft tissue injuries of the ankle.
The anterior portion of the splint is never used by itself; it is always used to augment the immobilization provided by the posterior ankle splint.

Figure 50.23 Anterior-posterior ankle splint.

Application. Position the patient and apply padding as described for a posterior splint. After the wet posterior splint has been applied, place the anterior splint over the anterior aspect of the ankle and foreleg parallel to the posterior splint. Hold the two in place with elastic bandages as described earlier for a posterior splint alone. An assistant is needed to apply an anterior-posterior splint because it is extremely difficult to hold both splints in place while wrapping the elastic bandages. Once secured, carefully mold both splints over the instep and ankle joint.

U-Splint (Stirrup Splint)
Indications. Use a U-splint or stirrup splint (Fig. 50.24) primarily for injuries to the ankle. It functions like a posterior splint, and either one provides satisfactory ankle immobilization. In one study that compared these splints in normal volunteers, the U-splint allowed less plantar flexion and broke less often with plantar flexion than did the posterior splint. In addition, because it actually covers the malleoli, a U-splint may protect the medial and lateral ligaments from further injury better than a posterior splint can.

Construction. Construct a U-splint of 4- or 6-inch-wide plaster strips. The splint passes under the plantar surface of the foot from the calcaneus to the metatarsal heads and extends up the medial and lateral sides of the foreleg to just below the level of the fibular head.

Application. Position the patient and apply padding as described for a posterior splint. Lay the wet plaster across the plantar surface of the foot between the calcaneus and the metatarsal heads with the sides extending up the lateral and medial aspects of the foreleg. Secure it in place with 4-inch elastic bandages. Wrap the elastic bandage around the extremity starting at the metatarsal heads and continuing around the ankle in a figure-of-eight configuration. Once the ankle has been wrapped, use another 4- or 6-inch elastic bandage to secure the remainder of the splint in place. Carefully mold the splint around the malleoli. The plaster may overlap on the anterior aspect of the ankle; this overlap does not interfere with the splint’s ability to accommodate further swelling. Note that if a U-splint is combined with a posterior splint, apply the posterior splint first.

Walking Boots
Indications. Use a walking boot (e.g., CAM Walker Boot) for the treatment of moderate to severe soft tissue injuries of the ankle, including second- and third-degree sprains (Fig. 50.25D). In addition, many orthopedic surgeons use it for isolated, nondisplaced lateral malleolar fractures. A walking boot provides a degree of immobilization similar to that of a U-splint but is easier to remove for bathing and dressing, and the Velcro straps allow adjustment for edema.
Refer patients in a walking boot for follow-up with an appropriate specialist. Advise them of the importance of partial
U-Splint (or Stirrup/Sugar-Tong Splint)

**Indications**
Injuries of the ankle, including:
- Fractures of the distal tibia and fibula
- Postreduction stabilization of ankle dislocations (above)

The U-splint can be combined with the posterior ankle splint to provide both anterior-posterior and medial-lateral stability.

**Application**
Apply the splint under the plantar surface of the foot and extend it up the medial and lateral sides of the foreleg to just below the level of the fibular head. Keep the ankle at a 90° angle.

For immobilization of the knee, extend the sides of the splint proximally to the groin to create a long leg splint.

**Figure 50.24** U-splint (also referred to as a stirrup or sugar-tong splint).

Splints for Ankle Sprains

A walking boot can be used for the treatment of moderate to severe soft tissue injuries of the ankle, including second- and third-degree sprains and isolated, nondisplaced lateral malleolar fractures. A walking boot provides a similar degree of immobilization as a U-splint but is easier to remove for bathing and dressing, and the Velcro straps allow adjustment for edema.

Prefabricated semirigid orthoses such as the Aircast (pictured above) are often used for patients with minor ankle sprains. The Aircast can be worn inside the patient’s shoe to provide early mobility along with increased ankle stability.

The Unna boot or an Ace wrap provides effective immobilization of an ankle soft tissue injury. The Unna boot is applied from a semisolid paste roll. The wrap is then covered with gauze or an elastic bandage. The entire dressing can be cut off by the patient at home. For similar short-term immobilization without plaster, a modified Jones dressing can be used. Copious Webril is wrapped around the ankle and foot and covered with an elastic bandage. A cast shoe can be used with this dressing.

**Figure 50.25** Splints for ankle sprains.
or non-weight bearing as indicated by the type and degree of injury. When cleared by the follow-up physician, a walking boot allows easy transition to full weight bearing. Studies have shown that rapid mobilization after ankle injuries improves functional outcome and reduces disability time.

**Application.** Walking boots come in a variety of sizes from extra small to extra large, depending on the manufacturer. The boot should fit comfortably with the patient’s calcaneus snugly in the heel of the boot and the patient’s toes close to but not extending over the front edge of the boot. Once the appropriate size has been determined, place the patient’s bare foot and ankle into the boot, and adjust the Velcro straps for a secure, but comfortable fit.

**Semirigid Orthosis**

**Indications.** In patients with lateral ankle sprains associated with a stable joint, a functional brace with early mobilization is frequently more comfortable and results in earlier return to normal function than complete immobilization in a plaster splint or cast (see Fig. 50.25B). Consequently, functional bracing with early mobilization has become the treatment of choice in most EDs. However, it should be pointed out that there is no documented difference in long-term outcome between the two methods of treatment.

**Application.** Most functional ankle braces resemble a U-splint with air bladders (Aircast, Inc., Summit, NJ) or foam padding (DeRoyal, Inc., Powell, TN) for cushioning the malleoli. The braces are secured about the ankle with Velcro straps. The device can be worn within the patient’s shoe over a sock and helps eliminate ankle instability.

**Hard Shoe (Cast or Reese Shoe)**

**Indications.** Use a hard shoe to help reduce pain on ambulation in patients with fractures or soft tissue injuries of the foot. This device can also be used over a splint or cast to allow partial weight bearing. A hard shoe is commonly used for fractured toes that have been buddy-taped.

**Application.** If a cast shoe is going to be used for a patient with a fractured toe, first buddy-tape the injured digit to the adjacent toe (Fig. 50.26). After this is done, the patient can slip the hard shoe on like a sandal. Fasten the shoe with ties or Velcro straps.

**Ankle Wraps and Bandages**

There are no data supporting the routine use of ankle wraps for simple sprains, but some pain relief may be afforded by a proper wrap. Some type of temporary immobilization is commonly used in the ED and usually requested and expected by patients. For minor ankle injuries, a simple elastic (Ace) bandage can be applied in a figure-of-eight configuration. Apply the wrap to give only lateral support with minimal compression. It should not be tight enough to impair venous drainage, a common problem when patients apply their own elastic wraps (see Fig. 50.4).

---

**Hard Shoe Splint**

![Figure 50.26 Hard shoe splint.](image-url)
An Unna boot placed over Webril is an alternative to a simple elastic bandage (see Fig. 50.25C). The Unna boot is constructed from a semisolid paste roll that hardens as it dries. Apply an Unna boot in a figure-of-eight configuration, similar to a simple elastic bandage. Once in place, wrap the ankle with roller gauze or an elastic bandage. The entire dressing can be cut off by the patient at home.

**Soft Cast**

**Indications.** A soft cast is basically a modified Jones compression dressing for the ankle. Use it for minor ligamentous and soft tissue injuries of the foot and ankle that do not require prolonged or complete immobilization. A soft cast can help reduce the pain and swelling often associated with mild ankle sprains, and it gives support for early weight bearing.

**Construction.** Make the soft cast of 3- or 4-inch Webril and elastic bandages.

**Application.** Place the patient in a supine position with the foot and ankle extending off the end of the stretcher. Alternatively, ask an assistant to elevate the leg or place pillows under the knee and foreleg. Wrap the ankle and foot with five to seven layers of Webril starting at the metatarsal heads and continuing around the ankle in a figure-of-eight configuration. Extend the Webril 5 to 7 cm above the malleoli and overlap each turn by 25% to 50% of its width. After the Webril is in place, wrap an elastic bandage around the foot and ankle in a similar fashion. Additional layers of Webril and elastic bandages are seldom required. A cast shoe can be used with this dressing.

**COMPLICATIONS OF SPLINTS**

**Ischemia**

A compartment syndrome (see Chapter 54) leading to ischemic injury and ultimately to a Volkmann ischemic contracture is the most worrisome complication of cylindrical casts. Although the risk for ischemia is drastically reduced with splinting, Webril or elastic bandages can cause significant constriction. To reduce the likelihood of constriction occurring, do not pull the elastic bandage excessively tight. If the patient has a high-risk injury, cut the Webril lengthwise before the plaster is applied. Stress the importance of elevation, no weight bearing, and application of cold packs, and carefully review the signs and symptoms of vascular compromise with every patient. All patients whose injuries have the potential for significant swelling or loss of vascular integrity should receive follow-up care in the first 24 to 48 hours. *Never ignore complaints of increasing pain under a splint*. Patients with splint-related discomfort must be reevaluated clinically and should not be treated with a telephone prescription for opioid analgesics (Fig. 50.27).

**Heat Injury**

Fiberglass splints produce minimal heat when drying, but plaster generates considerable heat as it hardens. Many clinicians are unaware of the potential for drying plaster to produce second-degree burns. Thermal injury can occur with both cylindrical casts and plaster splints. Some clinicians have reported a higher incidence of burns with the use of plaster splints, although the reasons for this are unclear. Box 50.4 lists factors that can increase the amount of heat produced during plaster recrystallization. Their effects are additive, and this fact should be taken into account when applying a splint. For example, if 15 sheets of plaster are needed for strength in a particular splint, one should not increase the heat production further by using extra-fast–drying plaster or reusing warm dip water. To avoid plaster burns, use only 8 to 12 sheets of plaster when possible, use fresh dip water with a temperature near 24°C, and never wrap the extremity in a sheet or pillow during the setting process. Peak temperatures usually occur between 5 and 15 minutes after plaster wetting.

The patient should be warned that the hardening process produces warmth. The heat of drying may produce pain in patients with hemophilia-related hemorrhages. Splinting these patients may require that the plaster splint be placed only long enough to verify proper fit; the splint is then reapplied after setting (and cooling) of the plaster. If any patient complains of significant burning while the plaster is drying, do not ignore this complaint! Immediately remove the splint, and promptly cool the area with cold packs or cool water. Patients with vascular insufficiency or sensory deficits (e.g., diabetic neuropathy, stroke) are at high risk for plaster burns and require close observation during the drying process.

**Pressure Sores**

Pressure sores are an uncommon complication of short-term splinting (Fig. 50.28). They can result from stockinette wrinkles, irregular wadding of Webril, incorrectly padded or unpadded bony prominences, irregular splint ends, plaster ridges, or indentations produced from using the fingers rather than the palms to smooth and mold the wet plaster. Attention to detail during padding and splinting reduces the incidence of pressure sores. However, whenever a patient complains of a persistent pain or burning sensation under any part of a splint, remove the splint and inspect the symptomatic area closely. The padding incorporated in premade plaster and fiberglass splints is generally all that is needed for safe short-term splinting. However, the life of a splint applied in the ED may
be longer than intended by the clinician; therefore it is prudent to err on the side of additional padding when placing splints on patients who may overuse the splint, such as those who will not use crutches, or for those who may not have ready access to follow-up.

**Infection**

Bacterial and fungal infections can occur under a splint. Infection is more common in the presence of an open wound but may occur with intact skin or in a skin lesion produced by prolonged splinting. The moist, warm, and dark environment created by the splint is an excellent nidus for infection. Toxic shock syndrome has been rarely reported from a staphylococcal skin infection that clandestinely developed under a splint or cast. In addition, it has been shown that bacteria can multiply in slowly drying plaster. To avoid infection, clean and débride all wounds before splint application, and use clean, fresh tap water for plaster wetting. If necessary, apply a removable splint that allows periodic wound inspection or local wound care.

**Dermatitis**

Occasionally, a rash develops under a plaster cast or splint. Allergy to plaster is exceedingly rare, but there are several reports of contact dermatitis when formaldehyde and melamine resins are added to the plaster. The rash is usually pruritic with weeping papular or vesicular lesions. Because these resins are unnecessary for ED splints, their use should be avoided whenever possible. Dermatitis has also been reported with the use of fiberglass splinting material.

**Pruritus**

Itching under a cast can be problematic. Patients, especially children, use various objects such as pencils, coat hangers, or forks to get to the itch. This can cause skin maceration and possible infection, or a foreign body can be left under the cast (Fig. 50.29).

**Joint Stiffness**

Some degree of joint stiffness is an invariable consequence of immobilization (Fig. 50.30). It can range in severity from mild
to incapacitating and can result in transient, prolonged, or in some cases, permanent loss of function. Stiffness appears to be worse with prolonged periods of immobilization, in elderly patients, and in patients with preexisting joint diseases such as rheumatoid arthritis or osteoarthritis. Thus, splints should be left on for only the time necessary for adequate healing. Table 50.3 lists several injuries that commonly require splinting, along with some suggestions for length of immobilization. Fractures, dislocations, or other conditions that require prolonged immobilization (>7 days) should have orthopedic follow-up. Tell patients that a splint is only a short-term device and that prolonged immobilization can be detrimental. For minor injuries, suggest that patients use their own judgment about when to remove the splint, but a definite end point should be set.

**Cast Pain**

Cast-related pain is a common complaint that brings patients to the ED. Because of the potential for ischemia with circumferential casts, fully investigate all complaints, and rule out vascular compromise. If a patient states that a cast is too tight, it probably is (see Fig. 50.27). Do not prescribe narcotics for cast pain until a properly fitting (i.e., one that is not too tight) cast has been ensured.

Perform a detailed history and physical examination on all patients complaining of cast pain. The nature and onset of the pain are of particular importance. A dull, nonspecific pain that has worsened gradually since the time of injury may be the only clue to an early compartment syndrome (see Chapter 54). A sudden onset of throbbing pain associated with swelling and redness suggests a possible deep venous thrombosis. In both these cases, rapid intervention is the key to decreasing morbidity and mortality. During the physical examination, pay particular attention to areas of tenderness and the effect of active and passive movement on the severity of the pain.

With a compartment syndrome, tenderness over the involved compartment is a common finding, and stretching or contracting ischemic muscle elicits significant pain. Evaluate for the presence and quality of distal pulses, the amount of edema fluid present, distal sensation, capillary refill, and color and temperature of the digits. The five Ps (pain, pallor, paresthesias, paralysis, and pulselessness) are pathognomonic for ischemia. Unfortunately, they seldom occur simultaneously, and their presence together is usually a late finding that carries a poor prognosis. Hence, the emergency clinician must maintain a high index of suspicion for possible ischemia and remove the cast if any possibility of vascular compromise exists. Almost any cast can be bivalved, with Webril and stockinette also cut, and the bivalved cast reapplied after inspection without significant loss of short-term immobilization.

To loosen a cast, use an oscillating cast saw to cut along the medial and lateral aspects of the cast (Fig. 50.31). This is called bivalving the cast, and it allows the halves to be spread and reapplied in a less constricting manner while still maintaining proper immobilization. To use an oscillating power saw, proceed in a series of downward cutting movements facilitated by wrist supination, and remove the blade between cuts to prevent it from getting hot enough to burn the skin. This is particularly important if synthetic material has been used in the cast. Additionally, do not allow the blade to slide along the skin, and never use the saw on unpadded plaster. In an apprehensive patient, demonstrate that the cast saw blade only vibrates (it does not turn) and that it does not cut the skin.

After the medial and lateral sides of the cast are completely cut through, separate the two halves with a cast spreader, and cut the padding lengthwise with scissors. Note that the padding may also contribute to the pressure, so once the plaster is bivalved, compression from padding must also be relieved. This may be sufficient to relieve early ischemia if the problem is simple postinjury swelling, but both the padding and the cast can be totally removed to inspect the injured area if necessary. If ischemia cannot be ruled out, measure compartment pressures (see Chapter 54) and obtain an orthopedic consultation.

If vascular integrity is established and no other problems are found, replace the bivalved cast. First, pad the extremity in the usual manner with fresh Webril. Line the cut ends of the bivalved cast with white adhesive tape, and replace the cast around the extremity. Finally, secure the cast in place with elastic bandages.

If plaster sores are causing the patient discomfort, consult the clinician who placed the cast. In some cases, additional padding is all that is needed, but in others, a window should be cut out over the problem area. Because pressure sores can lead to significant tissue necrosis, refer the patient for follow-up care within 24 hours.

If the patient’s problem is plaster (or, more likely, resin) dermatitis, administer topical or oral steroids and antihistamines. Provide therapy in concert with an orthopedic surgeon because the patient may require admission for other forms of immobilization until the cast can be replaced. In mild cases, changing the cast or splint and using antihistamines for symptomatic relief may suffice. All patients should receive close follow-up, and if the condition does not improve, the cast must be removed.

<table>
<thead>
<tr>
<th>TABLE 50.3 Suggested Length of Immobilization for Conditions That Frequently Require Splinting</th>
</tr>
</thead>
<tbody>
<tr>
<td>TYPE OF CONDITION</td>
</tr>
<tr>
<td>Contusions</td>
</tr>
<tr>
<td>Abrasions</td>
</tr>
<tr>
<td>Soft tissue lacerations</td>
</tr>
<tr>
<td>Tendon lacerations</td>
</tr>
<tr>
<td>Tendinitis</td>
</tr>
<tr>
<td>Puncture wounds and bites</td>
</tr>
<tr>
<td>Deep space infections and cellulitis</td>
</tr>
<tr>
<td>Mild sprains</td>
</tr>
<tr>
<td>Fractures and severe sprains</td>
</tr>
</tbody>
</table>

---

*aConsiderable controversy surrounds the length of immobilization for tendon lacerations, and duration is therefore best left to the consultant surgeon.  
<sup>b</sup>Usually requires prolonged immobilization, which is best determined by an orthopedic surgeon.
CONCLUSION

Splinting is an important means of temporary fracture immobilization and provides protection and comfort for a variety of soft tissue injuries. The clinician should be aware of potential complications that can occur with improper splint application, including ischemia, thermal injury, and pressure sores. Use proper technique to minimize the risk for these adverse outcomes. When ischemia is suspected, the emergency clinician should also be facile in the release of circumferential cast and splint material.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES

Podiatric Procedures

Douglas L. McGee

Chapter 51

Common Nontraumatic Conditions of the Foot

Many painful conditions of the foot are chronic and do not usually require definitive treatment in the emergency department (ED); however, patients are often seen with common conditions that require evaluation and proper referral. To accomplish this, the clinician must be cognizant of basic podiatric conditions, including painful lesions over bony prominences, heel pain, foot infections, and pain on the plantar surface of the foot.

Footpad Use

Footpads redistribute pressure over an inflamed, tender area of the foot. The particular type of footpad and its placement depend on the condition being treated (Fig. 51.1). Commercially available aperture footpads are recommended for the temporary relief of warts, corns, hyperkeratoses, and bunions. Verruca virus introduced into the plantar surface of the foot may produce a painful hyperkeratotic lesion, commonly referred to as a plantar wart, on the sole of the foot. A simple callus may be painful and result in the formation of a “hard corn” when formed over the bony prominence of a digit. Once recognized, and after other conditions are ruled out, definitive care of these lesions is rarely indicated in the ED.

When tenderness is elicited over more than one metatarsal head, the diagnosis is metatarsalgia. Pain that is progressively worse while walking but relieved by rest, often beneath the second or third metatarsal head, is typical in this case. A pad placed under the first metatarsal head to raise the second and third metatarsals may provide some relief.

A bunion develops when unbalanced forces applied to the first metatarsal cause lateral displacement of the distal end of the hallux. Bunions typically form in women wearing heeled shoes with narrow toe boxes. The patient may complain of numbness over the distal, medial aspect of the first toe as a result of compression of the terminal branch of the medial dorsal cutaneous nerve. The mechanical forces that precipitate bunion formation may also cause other painful conditions, including intermetatarsal neuromas, hammertoes, ingrown toenails, corns, and calluses. Bursitis may develop over the medial bony prominence of the first metatarsophalangeal (MTP) joint. Self-adherent bunion pads placed over the first MTP joint may provide temporary relief (Fig. 51.2). In the ED, treat patients suffering from these common disorders with analgesics and footpads, followed by referral for definitive care. Recommend that the patient avoid wearing the offending shoes. In men and postmenopausal women consider gout when evaluating pain over the first MTP joint, particularly in the presence of other signs of inflammation (e.g., redness, swelling, warmth).

Heel Pain Syndromes

Bony spurs on the plantar surface of the calcaneus, retrocalcaneal bursitis, calcaneal apophysitis, and other conditions may cause heel pain. Treat most of these conditions with rest, nonsteroidal antiinflammatory drugs (NSAIDs), modification of physical activities or shoe wear, footpads, and orthoses. Some clinicians also include injection of anesthetics or steroids for these conditions.

Heel spur pain can be quite bothersome and chronic or recurrent. This condition is not easily remedied in the ED, and after other conditions are ruled out, such as an unexpected foreign body (FB), minimal intervention with podiatric referral is often the best course of action. Patients typically have pain over the medial border of the plantar aspect of the calcaneus. The pain gradually worsens over a period of months. A bony prominence that begins as periostitis extends from the medial aspect of the calcaneal tuberosity into the central plantar fascia and may be seen on radiographs. Radiographs of the calcaneus that do not demonstrate a bony spur suggest plantar fasciitis (see the later section on Painful Conditions of the Plantar Surface of the Foot), even though many patients with plantar fasciitis have plantar calcaneal and Achilles spurs. Plantar calcaneal heel spurs are found in nearly 15% of the population, only 30% of whom have heel pain. Although many persons with heel spurs are asymptomatic, 75% of patients with heel pain have heel spurs. If however, radiographs have little value in evaluating nontraumatic heel pain because they rarely demonstrate radiographic abnormalities that prompt additional treatment.

Shoe supports with a heel pad or cup, or a doughnut-shaped orthotic often help reduce the discomfort by redistributing weight. Few randomized controlled trials have evaluated steroid therapy; those that have do not provide substantial evidence supporting its long-term efficacy. Treatment for the painful site is 10 to 20 mg of methylprednisolone injected into the medial aspect of the foot while avoiding the sensitive plantar surface. Some evidence suggests that injecting 25 mg of prednisolone acetate into the medial aspect of the heel provides partial pain relief at 1 month in comparison to lidocaine only, but no advantage can be detected at 3 months. A short-leg walking cast may be effective in some patients with recalcitrant heel pain. There is little evidence to suggest that specific interventions aimed at reducing heel pain are superior to conservative, supportive treatment alone.

Retrocalcaneal Bursitis, Achilles Tendinopathy, and Calcaneal Apophysitis

Although retrocalcaneal bursitis and Achilles tendinopathy (formerly referred to as Achilles tendinitis) are anatomically
distinct, the clinical findings are similar. Pain at the insertion of the Achilles tendon is worsened with prolonged standing or walking and is aggravated by passive or active range of motion in both conditions. Directed palpation can distinguish one entity from the other, but both are treated similarly. Tenderness of the Achilles tendon suggests tendinopathy, whereas tenderness between the tendon and the calcaneus suggests retrocalcaneal bursitis. Importantly, Achilles tendinopathy has been noted to develop spontaneously after the use of quinolone antibiotics, occasionally with rupture. The condition may occur during quinolone use or a few weeks after therapy and prompts immediate discontinuation of use of the drug if recognized. Rest, elevation, ice, NSAIDs, heel pads, and an open-backed shoe provide relief in the majority of patients. A corticosteroid injection is not usually performed, but it may provide some relief, although its superiority over conservative measures is unproved. Repeated steroid injection is associated with Achilles tendon rupture. Injection of platelet-rich plasma (PRP) as treatment for Achilles tendinopathy has gained some support among sports medicine physicians, but recently published studies (2016) have failed to demonstrate short- or long-term improvement in pain or function when PRP injection is compared with placebo. Osteochondrosis of the posterior calcaneal apophysis, often referred to as Seaver’s disease, may cause pain that is worsened by activity in children between 7 and 10 years of age. It is thought to represent an overuse syndrome in an athletically active child with tenderness in the posterior heel region. Treat this self-limited condition with rest, ice, and heel pads. Radiographs are not indicated unless other diagnoses are suspected. Activity is resumed when the pain abates.

**Painful Conditions of the Plantar Surface of the Foot**

**Plantar Fasciitis**

Repeated microtrauma to the plantar aponeurosis causes pain on the plantar surface of the foot (Fig. 51.3A). Plantar fasciitis is typically unilateral and found in women who wear high-heeled shoes. The pain is maximally severe in the morning or after prolonged sitting and improves after walking, often referred to as first-step pain. Some patients with plantar fasciitis may also have a calcaneal heel spur, but the presence or absence of this radiographic finding is clinically irrelevant. Pain is elicited with palpation (see Fig. 51.3B), toe walking, or passive stretching of the plantar aponeurosis. Frequently, this annoying condition resolves spontaneously, but resolution is slow, with as long as 6 to 18 months needed. Conservative therapy, including rest, elevation, ice, and NSAIDs, results in a satisfactory outcome after 6 to 8 weeks in 90% of patients. The pain improves over time in most patients, with or without NSAIDs, although the addition of NSAIDs appears to increase pain relief when compared with conservative treatments alone. The emergency clinician can do little to treat this chronic, distressing condition. Stretching exercises each morning and evening can be suggested (see Fig. 51.3D). Night splinting to keep the foot dorsiflexed and custom orthoses made from the patient’s foot impression can be very helpful but usually require referral to podiatry for proper fitting (see Fig. 51.3C). Corticosteroid injection is used by some clinicians; however, its benefit remains unproved. A single injection may be warranted as supplemental therapy in resistant cases. Repeated injections of corticosteroids should be avoided and have been associated with rupture of the plantar fascia and fat pad atrophy. Some authors suggest the injection of autologous PRP when plantar fasciitis is chronic. The benefits of PRP remain unproved. In a 2010 study, 50 units of botulinum toxin type A injected into the plantar fascia decreased pain in comparison to placebo but did not cure the disease.

**Forefoot Neuroma**

A forefoot neuroma, also known as Morton’s neuroma, is a painful condition of the plantar surface of the foot. It most commonly affects women who wear high-heeled shoes. The neuroma forms after chronic irritation to the digital sensory nerve between the metatarsals. A neuroma frequently occurs in the third interspace but may be found in the second (Fig. 51.4A). Patients report the sensation of a lump or cord in the interspace and describe paresthesia or numbness in the third or fourth toes. Direct compression plus release of the forefoot causes pain and often a “click” (Mulder’s sign) (see Fig. 51.4B). Rest, elevation, ice, and NSAIDs may result in some improvement, but surgical excision is often required. Few data support the use of corticosteroid injections. In one study, less than...
Figure 51.3 Plantar fasciitis. A, The plantar fascia spans the plantar aspect of the foot. The origin is the medial tubercle of the calcaneus, which is the most common site of pain. A heel spur may be seen on radiographs, but inflammation of the plantar fascia, not the spur, is the source of the pain. B, Palpation of the tubercle of the calcaneus reproduces the pain of plantar fasciitis. C, A posterior lower leg splint may be created for patients in the emergency department, and it is intended to be worn at night to keep the foot in a position of dorsiflexion. Referral to podiatry for custom-fit orthoses may also be beneficial. D, Rolling the arch of the foot back and forth over a frozen water bottle will stretch the fascia and, over time, may lessen the pain of plantar fasciitis. A tennis ball can also be used.

Figure 51.4 Morton’s neuroma. A, Site of Morton’s neuroma arising from a digital nerve. B, Palpation of the distal metatarsal area may reveal pain from Morton’s neuroma. Squeezing and releasing this area (not shown) can elicit pain and a clicking sensation, termed Mulder’s sign.
50% of patients with a foot neuroma had any benefit from injected corticosteroids.17 Another study demonstrated complete or partial relief in 80% of patients injected with corticosteroids.19 Although this study demonstrated a trend toward improved outcome when injected corticosteroids were compared with footwear modifications, corticosteroid injection therapy alone was not statistically better than footwear modification at 1 year.19

**Ganglion Cyst of the Foot**

A ganglion cyst is histologically similar to the synovial sheath and contains synovial fluid. The diagnosis is easy to make when the cyst is located over a tendon on the dorsum of the foot (Fig. 51.5), but may be difficult when located among the compact structures of the plantar forefoot. A ganglion cyst usually causes edema along the involved tendon sheath. The mass should roll under the examiner’s finger; a painless, immovable mass suggests a soft tissue neoplasm. Painful ganglion cysts are treated by aspiration, with or without injection of a corticosteroid (see Chapter 52). After local or regional anesthesia (see Chapters 29 and 31), insert a 20-gauge needle into the cyst and withdraw yellow, thick, synovial fluid. Manually express any remaining synovial fluid after the needle is withdrawn. Corticosteroid injection is often advocated for ganglion cysts, but recurrence is common after aspiration and corticosteroid injection, as high as 57% in one study.20 Many studies have shown that recurrence, chronic pain, neuritis, stiffness, and infection are not uncommon, even after surgical excision.21 Other authors report improved functional outcome, a 9% occurrence of paresthesia, and only a 5.7% recurrence rate after excision.22

**TRAUMATIC CONDITIONS OF THE FOOT**

Trauma to the feet and toes is common and covers a broad spectrum of injury. Lacerations, fractures, compartment syndromes, and nail bed injuries are described in other chapters of this text. Three specific injuries— toe fractures, sesamoid-bone fractures, and puncture wounds to the plantar surface of the foot—are discussed in detail here.

**Toe Fractures and Fractures of the Sesamoid Bones**

Emergency clinicians often treat toe fractures and can intervene to relieve the pain and encourage healing. As with any other fracture, pay attention to the possibility of disrupted joint cartilage, hypermobility of the fracture segments, and malposition or malunion of the fracture fragments. Fracture displacement greater than 2 mm is uncommon; reduction is rarely needed for most toe fractures.23 However, aggressive reduction is indicated for fractures of the proximal phalanx of the great toe because it represents the main propulsive segment of the forefoot (Fig. 51.6). A plaster cast alone without anatomic reduction is insufficient treatment. Displacement suggests axial rotation or abnormal biomechanical interaction between the hallux and its own interphalangeal or MTP joint.

In the acute setting, a non–weight-bearing ankle splint that extends beyond the great toe provides protection until the patient with a complicated fracture of the great toe obtains follow-up with a foot and ankle surgeon. Open fractures require careful cleaning, usually antibiotic therapy, and close follow-up. Fractures of the lesser toes generally result from jamming the toe into a nightstand or bedpost when barefoot. Radiographs of the lesser phalanges confirm the suspected fracture and may occasionally reveal an unsuspected interphalangeal or MTP dislocation. However, radiographs are not typically required, and the injured digit is easily reduced.

Treat closed, lesser phalangeal fractures with immobilization for 6 weeks. After the fracture is reduced, splint the injured toe against an adjacent noninjured toe. Place a soft corn pad or other suitable material between the toes to prevent skin maceration, and hold the toes together with adhesive tape or a self-adherent wrap such as Coban (3M Company, St. Paul, MN) (Fig. 51.7). Demonstrate the procedure to the patient or family and dispense or prescribe enough material so that the splint can be changed every 2 to 3 days at home. Have the patient wear a less restrictive, stiff-soled shoe (Fig. 51.8). A postoperative shoe (or similar footwear) may be a comfortable alternative for the first several days.
Jumping from a height can result in a fracture of the first MTP joint sesamoid bone (Fig. 51.9A). The great toe sesamoid bones lie in grooves on the bottom of the metatarsal head. Each bone lies within the tendon of its respective flexor hallucis brevis muscle belly. Localized pain on the plantar aspect of the first metatarsal head accompanies a sesamoid-bone fracture. Bipartite sesamoids (tibial more frequently than fibular) are common. Comparison radiographs clarify whether the radiographic abnormality represents a fracture.

For a tibial sesamoid injury, an aperture bunion-type pad, reinforced medially with 0.5- to 0.75-cm-thick felt, protects the sesamoid and transfers weight bearing to the surrounding structures (see Fig. 51.9B). A hard-soled shoe and NSAIDs are also helpful. Subsequent radiographs rarely show bony consolidation, but the fracture interface appears smoother.

Stress Fractures

A metatarsal stress fracture, commonly of the distal second and third metatarsals, can develop in runners, military recruits, or those with repetitive trauma to the foot (Fig. 51.10). Other risk factors are listed in Box 51.1.54 There is pain on walking but minimal to no external findings. A bone scan or magnetic
resonance imaging (MRI) may detect the subtle fracture that often eludes a plain radiograph. Occasionally, an occult FB of the foot is the culprit and can mimic a stress fracture. Women are more prone to stress fractures than men. Treatment is usually rest and alleviation of the precipitant causes.

**Plantar Puncture Wounds**

Plantar puncture wounds present a diagnostic and therapeutic challenge for the clinician. Considerable controversy exists regarding the proper initial management of puncture wounds in the plantar surface of the foot, and no universally accepted standard of care exists. Treatment recommendations range from simple cleaning of the wound to aggressive débridement. No single approach has been demonstrated to be superior. The author supports close inspection for retained foreign material and an initially conservative approach, but an aggressive one is recommended if the patient returns, has an infection, or if the pain persists for more than a few days.

Although nails produce many such wounds, various other objects may cause them, including other metal objects, wood, and glass. Patient response to the injury depends on the penetrating material, location and circumstances of the wound, depth of penetration, footwear, time from injury until initial evaluation, and underlying health. As superficial puncture wounds generally do well, depth of penetration may be a primary determinant of outcome. Because one's reflexes are not fast enough to pull back when stepping on a sharp object, the clinician should assume that the entire length of a protruding nail has entered the foot (minus the thickness of the footwear). Stepping on an unknown object in a field or while walking in...
a stream requires a more cautious approach than a simple puncture from a known object, such as a protruding nail.

The vast majority of patients who step on a nail suffer nothing more than transient pain and never seek medical attention. Because most minor puncture wounds are not seen in the ED, the true risk for infection, including osteomyelitis, is unknown. Consequently, reported infection rates vastly overstate the actual incidence. Probably no more than 2% to 8% of puncture wounds become infected, and in only a small percentage of these wounds is surgical débridement needed or does osteomyelitis develop. A prospective series suggested that only the presence of symptoms (e.g., redness, tenderness, increased swelling) at 48 hours is associated with risk for infection or a potential retained FB. Retained foreign material (e.g., a piece of sock or a portion of a tennis shoe sole) in the wound is an important factor in persistent infection. Because no single test detects all possible FBs, tailor the evaluation of a suspected FB to the suspected object.

**Evaluation**

The approach to the patient depends on several factors, including the time from injury to evaluation, suspicion of an FB, and the presence of infection. The extent to which an FB is pursued depends on the history and findings on physical examination. When the patient clearly states that a needle, pin, or nail was removed intact, radiographs or local wound exploration for a retained metallic FB is not needed. Stepping on an identified intact nail does not always require a radiograph, but if there is any debate regarding a retained metallic object, plain radiographs readily demonstrate their presence. Plain radiographs also demonstrate other radiopaque objects such as glass, gravel, bone, and teeth (Fig. 51.11). Rubber or pieces of a sock will not be visible on a radiograph. If the patient steps on an unknown object, a radiograph is usually indicated unless the entire depth of the wound can be ascertained and inspected. Ultrasonography is noninvasive and does not use radiation, thus making it potentially useful for radiolucent FBs, but soft tissue, air, or calcifications may suggest a retained FB when none is present. Bedside ultrasound has been used to localize radiolucent FBs for removal. Computed tomography (CT) can demonstrate radiopaque and radiolucent objects, but its expense and greater radiation exposure than with plain films make it unsuitable as an initial screening tool. Use CT scanning when other screening tools fail to demonstrate a suspected FB, when infection is present, or when joint penetration is suspected. Fluoroscopy may be used to help localize metallic or radiopaque FBs during exploration and removal. It may be particularly useful for long metallic objects such as needles and pins. MRI provides no additional benefit and cannot be used when metallic objects are retained. Diabetic patients with unexplained foot pain or infection warrant investigation for a retained FB; diabetic neuropathy may mask the initial puncture injury.

**Treatment**

As a general rule, the plantar surface of the foot should be examined under good lighting and in a bloodless field. This is best accomplished with the patient in a prone position, not in a chair (Fig. 51.12). Plantar puncture wounds explored for persistent infection often have foreign material in them. Because persistent infection does not develop in most patients, initial deep exploration in the absence of evidence or a strong suspicion of retained material cannot be advocated. Routine initial deep wound exploration, including coring of the wound, is not supported by scientific research and is not recommended for simple, noninfected puncture wounds. However, selected wounds may benefit from exploration to facilitate a search for FBs and to promote irrigation, cleansing, and drainage. When the wound is large and retained organic material is suspected, local wound exploration is warranted. Patients wearing rubber-soled shoes during plantar puncture wounds may retain a portion of the shoe in the wound. Exploration of the wound is most productive after local anesthesia or a regional anesthetic block and excision of the epidermal flap. An incision may be required to facilitate removal of the FB when known to be present, but it has not been proved that extensive removal of surrounding tissue increases successful removal of retained material.

Some clinicians favor a coring technique when an FB is found or suspected. Although this may be too aggressive for many wounds, it may be the best way to remove particulate matter as a block (Fig. 51.13). To accomplish coring, advance a No. 11 blade to the hilt and excise a 2- to 3-mm core. Use a hemostat to grab the cutout core or to open the tract to better visualize the wound. Alternatively, use a 2- to 3-mm punch biopsy instrument to core out the tract. The tract can be packed with gauze for a few days or left open. There is no...
Initial antibiotic therapy remains controversial, but there is no evidence to suggest that prophylactic antibiotics reduce the already low rate of infection. In fact, some cases of *Pseudomonas* osteomyelitis develop despite initial treatment with anti-*Pseudomonas* antibiotics. The author does not suggest routine prophylactic antibiotics for uncomplicated puncture wounds, and such treatment may theoretically select out resistant organisms.

Symptomatic patients who delay presentation after injury (≥48 hours) may have an increased probability for a retained FB when inflammation or infection develops. Local wound exploration is warranted unless the patient does not have an infection or a clearly superficial cellulitis is expected to respond to simple oral antibiotics. Recurrent infection, deep soft tissue tenderness, and increasing soft tissue swelling suggest a retained FB or deep space infection, such as osteomyelitis. Such patients require prompt specialty referral or additional diagnostic studies in the ED.

**Figure 51.12** Removal of a foreign body in the plantar surface of the foot.
Foreign Body Removal: Coring Technique

A foreign body is difficult to appreciate in this puncture wound of the big toe.

To core out a puncture, use a No. 11 blade and advance it to the hilt.

When the cored-out section was removed with a hemostat, a piece of rubber was found deep in the wound.

After coring out a puncture tract, the wound can be left open (as shown here) or packed (as shown in Fig 51.12).

Figure 51.13 Foreign body removal: coring technique.

Patients with obvious signs of infection within a few days of a puncture wound usually have a simple cellulitis (with or without an FB) with a gram-positive organism (Fig. 51.14). In a patient with persistent pain or swelling days to weeks after a puncture wound, the presence or absence of a deep soft tissue infection or low-grade osteomyelitis cannot be ruled in or ruled out by physical examination, plain radiographs, or laboratory tests (such as a sedimentation rate, complete blood count, or wound cultures). A high index of suspicion, coupled with additional investigation, is the prudent approach to patients with minimal findings on physical examination, normal laboratory test results, and continued pain or swelling after a seemingly simple puncture wound in the bottom of the foot. This usually entails CT, bone scan, or MRI. A study of 80 children with plantar puncture wounds and signs of infection at presentation found simple cellulitis in 59, retained FBs in 11, and osteomyelitis or septic arthritis in 10 children. Because a significant number of children in this study had retained FBs or bony infections, a cautious approach is warranted.31

Figure 51.14 This 5-day-old inflamed and infected puncture wound tract probably harbors a foreign body. Minor lymphangitis (arrow) spread up the ankle, but there were no systemic complaints, adenopathy, or leukocytosis. Excision of a core of the tract found scant pus and a few pieces of the patient’s sock embedded in the wound. The wound was left open and did well with oral antibiotics and warm soaks.
Plantar puncture wounds complicated by *Pseudomonas* osteomyelitis and osteochondritis are clinically clandestine and particularly devastating. They have been described for nearly 40 years. Some investigators have cultured *Pseudomonas* from the soles of tennis shoes, thus suggesting that puncture wounds made through athletic footwear may be inoculated with *Pseudomonas* (Fig. 51.15).  

No evidence suggests that prescribing prophylactic anti-*Pseudomonas* antibiotics on initial evaluation of an uncomplicated wound will prevent infection in patients with subsequent deep space or bone infections. Some argue that prophylactic antibiotics, at the time of puncture, may select out resistant organisms. Surgical débridement and prolonged intravenous antibiotics are often required for established infections.

**INGROWN TOENAIL**

An ingrown toenail is characterized by progressive curving or excessive widening of the lateral margin of the toenail and impingement of the nail into the periungual soft tissue (Fig. 51.16). The toenail normally grows distally in an unimpeded manner, thereby allowing the nail to pass beyond the lateral nail fold. Nail deformity, tight-fitting shoes, and rotational deformity of the toes increase the friction between the nail and the nail fold. Toenails that have been trimmed in a curve increase the likelihood that the lateral nail margin will impinge on the lateral nail fold (Fig. 51.17). The resulting soft tissue injury may lead to hyperkeratosis, edema, and erythema of the nail fold or frank infection (Fig. 51.18). Although an ingrown toenail can be found on any toe, the majority occur on the great toe.

**Evaluation**

The patient has pain, edema, and erythema of the lateral nail fold. Pressure over the nail margins increases the pain. Because
intense pain often precipitates an early visit to the ED, most inflammatory or infectious responses are confined locally. Recurrent ingrown toenails or those in patients with circulatory dysfunction, neuropathy, or diabetes may have underlying osteomyelitis. Cleanse the toe gently to facilitate visualization of the periungual debris. When the free edge of the lateral part of the nail can easily be visualized as separate from the lateral nail fold, consider other painful conditions of the toe such as trauma, gout, paronychia, and cellulitis.

**Treatment**

Because the toe is exquisitely tender, additional treatment will usually require digital block anesthesia. The decision to treat and what course of action to take in the ED depend on the patient’s degree of discomfort. Two general courses of action are often suggested: removal of the offending nail spicule or removal of the spicule and some portion of the nail. Any degree of nail removal is usually followed by ablation of the nail bed. Nail-splitting techniques may also be effective and are less invasive than nail removal techniques.

**Removal of the Nail Spicule and Débridement of Hyperkeratosis for Minor Ingrown Toenails**

When the amount of inflammation and pain and the degree of nail deformity are both minimal, such as involvement of only the distal toenail area, simple removal of the impacted nail spicule is indicated and usually curative. If the condition is advanced, complete removal of a portion of the nail may be required. All procedures are best done with a tourniquet to produce a bloodless field. After a digital block and thorough cleansing, remove an oblique segment of the nail approximately one third to one half the way to the proximal nail fold (Fig. 51.19). The ideal instrument is an English anvil nail splitter, which is designed to cut the nail while minimizing trauma to the underlying nail matrix (Fig. 51.20). Sharp, pointed scissors may be substituted if care is taken to minimize injury to the nail bed by maintaining upward pressure while cutting the nail. Some clinicians use a disposable electric cautery device to cut the nail after softening it by soaking in warm soapy water.

After cutting the nail spicule free from the bulk of the nail, grasp the free edge of the nail with forceps or hemostats and remove it to expose the irritated area. The nail fold typically contains impacted debris that must be removed after the nail fold has been gently retracted away from the nail. Remove debris until the epidermis or dermis is uncovered while taking care to avoid aggressive débridement that causes bleeding. A silver nitrate stick applied to the débrided area may be used to control bleeding. Dress the area with antibiotic ointment and a nonadherent dressing. Soaking the toe in warm water two or three times a day, with home redressing, is explained. Harsh chemicals are avoided in the soaking regimen. The wound should be reinspected for signs of infection at 48 to 72 hours. If the problem has resolved, no further therapy is necessary. If the problem is persistent or recurrent, referral for definitive podiatric care is reasonable. Instruct patients to wear less constricting shoes and to trim the nail straight across. Because the toe is exquisitely tender, additional treatment will usually require digital block anesthesia. The decision to treat and what course of action to take in the ED depend on the patient’s degree of discomfort. Two general courses of action are often suggested: removal of the offending nail spicule or removal of the spicule and some portion of the nail. Any degree of nail removal is usually followed by ablation of the nail bed. Nail-splitting techniques may also be effective and are less invasive than nail removal techniques.

**Removal of the Nail Spicule and Débridement of Hyperkeratosis for Minor Ingrown Toenails**

When the amount of inflammation and pain and the degree of nail deformity are both minimal, such as involvement of only the distal toenail area, simple removal of the impacted nail spicule is indicated and usually curative. If the condition is advanced, complete removal of a portion of the nail may be required. All procedures are best done with a tourniquet to produce a bloodless field. After a digital block and thorough cleansing, remove an oblique segment of the nail approximately one third to one half the way to the proximal nail fold (Fig. 51.19). The ideal instrument is an English anvil nail splitter, which is designed to cut the nail while minimizing trauma to the underlying nail matrix (Fig. 51.20). Sharp, pointed scissors may be substituted if care is taken to minimize injury to the nail bed by maintaining upward pressure while cutting the nail. Some clinicians use a disposable electric cautery device to cut the nail after softening it by soaking in warm soapy water.

After cutting the nail spicule free from the bulk of the nail, grasp the free edge of the nail with forceps or hemostats and remove it to expose the irritated area. The nail fold typically contains impacted debris that must be removed after the nail fold has been gently retracted away from the nail. Remove debris until the epidermis or dermis is uncovered while taking care to avoid aggressive débridement that causes bleeding. A silver nitrate stick applied to the débrided area may be used to control bleeding. Dress the area with antibiotic ointment and a nonadherent dressing. Soaking the toe in warm water two or three times a day, with home redressing, is explained. Harsh chemicals are avoided in the soaking regimen. The wound should be reinspected for signs of infection at 48 to 72 hours. If the problem has resolved, no further therapy is necessary. If the problem is persistent or recurrent, referral for definitive podiatric care is reasonable. Instruct patients to wear less constricting shoes and to trim the nail straight across. Because the toe is exquisitely tender, additional treatment will usually require digital block anesthesia. The decision to treat and what course of action to take in the ED depend on the patient’s degree of discomfort. Two general courses of action are often suggested: removal of the offending nail spicule or removal of the spicule and some portion of the nail. Any degree of nail removal is usually followed by ablation of the nail bed. Nail-splitting techniques may also be effective and are less invasive than nail removal techniques.

**Removal of the Nail Spicule and Débridement of Hyperkeratosis for Minor Ingrown Toenails**

When the amount of inflammation and pain and the degree of nail deformity are both minimal, such as involvement of only the distal toenail area, simple removal of the impacted nail spicule is indicated and usually curative. If the condition is advanced, complete removal of a portion of the nail may be required. All procedures are best done with a tourniquet to produce a bloodless field. After a digital block and thorough cleansing, remove an oblique segment of the nail approximately one third to one half the way to the proximal nail fold (Fig. 51.19). The ideal instrument is an English anvil nail splitter, which is designed to cut the nail while minimizing trauma to the underlying nail matrix (Fig. 51.20). Sharp, pointed scissors may be substituted if care is taken to minimize injury to the nail bed by maintaining upward pressure while cutting the nail. Some clinicians use a disposable electric cautery device to cut the nail after softening it by soaking in warm soapy water.

After cutting the nail spicule free from the bulk of the nail, grasp the free edge of the nail with forceps or hemostats and remove it to expose the irritated area. The nail fold typically contains impacted debris that must be removed after the nail fold has been gently retracted away from the nail. Remove debris until the epidermis or dermis is uncovered while taking care to avoid aggressive débridement that causes bleeding. A silver nitrate stick applied to the débrided area may be used to control bleeding. Dress the area with antibiotic ointment and a nonadherent dressing. Soaking the toe in warm water two or three times a day, with home redressing, is explained. Harsh chemicals are avoided in the soaking regimen. The wound should be reinspected for signs of infection at 48 to 72 hours. If the problem has resolved, no further therapy is necessary. If the problem is persistent or recurrent, referral for definitive podiatric care is reasonable. Instruct patients to wear less constricting shoes and to trim the nail straight across. Because the toe is exquisitely tender, additional treatment will usually require digital block anesthesia. The decision to treat and what course of action to take in the ED depend on the patient’s degree of discomfort. Two general courses of action are often suggested: removal of the offending nail spicule or removal of the spicule and some portion of the nail. Any degree of nail removal is usually followed by ablation of the nail bed. Nail-splitting techniques may also be effective and are less invasive than nail removal techniques.

**Removal of the Nail Spicule and Débridement of Hyperkeratosis for Minor Ingrown Toenails**

When the amount of inflammation and pain and the degree of nail deformity are both minimal, such as involvement of only the distal toenail area, simple removal of the impacted nail spicule is indicated and usually curative. If the condition is advanced, complete removal of a portion of the nail may be required. All procedures are best done with a tourniquet to produce a bloodless field. After a digital block and thorough cleansing, remove an oblique segment of the nail approximately one third to one half the way to the proximal nail fold (Fig. 51.19). The ideal instrument is an English anvil nail splitter, which is designed to cut the nail while minimizing trauma to the underlying nail matrix (Fig. 51.20). Sharp, pointed scissors may be substituted if care is taken to minimize injury to the nail bed by maintaining upward pressure while cutting the nail. Some clinicians use a disposable electric cautery device to cut the nail after softening it by soaking in warm soapy water.

After cutting the nail spicule free from the bulk of the nail, grasp the free edge of the nail with forceps or hemostats and remove it to expose the irritated area. The nail fold typically contains impacted debris that must be removed after the nail fold has been gently retracted away from the nail. Remove debris until the epidermis or dermis is uncovered while taking care to avoid aggressive débridement that causes bleeding. A silver nitrate stick applied to the débrided area may be used to control bleeding. Dress the area with antibiotic ointment and a nonadherent dressing. Soaking the toe in warm water two or three times a day, with home redressing, is explained. Harsh chemicals are avoided in the soaking regimen. The wound should be reinspected for signs of infection at 48 to 72 hours. If the problem has resolved, no further therapy is necessary. If the problem is persistent or recurrent, referral for definitive podiatric care is reasonable. Instruct patients to wear less constricting shoes and to trim the nail straight across. Because the toe is exquisitely tender, additional treatment will usually require digital block anesthesia. The decision to treat and what course of action to take in the ED depend on the patient’s degree of discomfort. Two general courses of action are often suggested: removal of the offending nail spicule or removal of the spicule and some portion of the nail. Any degree of nail removal is usually followed by ablation of the nail bed. Nail-splitting techniques may also be effective and are less invasive than nail removal techniques.
Cleanse the toe with antiseptic and then administer a digital block. Use of a tourniquet to provide a bloodless field is ideal but is not depicted here.

First separate the nail from the nail bed by advancing and separating scissors held parallel to the nail bed (see Fig. 51.23), and then split the nail lengthwise toward the cuticle as depicted above.

Grasp the end of the cut toenail with a hemostat and twist toward the remaining nail.

Use the twisting motion to remove the free piece of nail.Sharply remove any remaining skin and hyperkeratotic debris. Silver nitrate may be applied to the nail bed if desired.

Inspect the remnant to make sure that the entire piece of nail has been removed as desired. Note the prominent spicule present on this ingrown toenail.

Cover the wound with antibiotic ointment, a nonadherent dressing, and a dry sterile wrap.

**Figure 51.22** Removal of an ingrown toenail. Antibiotics effective against gram-positive organisms are prescribed for significant associated infection but are likely not required for this localized process.

Partial nail removal accomplishes two things: removal of the offending portion of nail and destruction of the underlying nail matrix to prevent regrowth of the nail. Phenol, the most commonly applied chemical, causes neurolysis of the nerve endings and necrosis of the nail matrix in a procedure called *matricectomy.* Several studies have demonstrated that 10% sodium hydroxide solution is as effective as phenol and may be associated with less postprocedural pain and faster recovery. As most EDs do not stock phenol or sodium hydroxide solution, nail bed ablation is usually not pursued on the initial visit.

After a digital block, exsanguinate the toe by squeezing or wrapping, and apply a tourniquet at the base of the toe. Stabilize the toe with the nondominant hand. Separate the lateral third of the nail from the nail bed by advancing and separating scissors held parallel to the nail bed (Fig. 51.23). Split the nail
lengthwise toward the cuticle. An English anvil nail splitter is desirable to begin the procedure, but sharp scissors or a No. 11 blade also work. Take care to perform a controlled division along the longitudinal lines of the nail for several millimeters past the proximal nail fold (cuticle). Grasp the end of the cut toenail with a hemostat or forceps. Remove the free piece of nail by twisting it toward the remaining nail. This will pull the nail root out from under the cuticle. Inspect the remnant to be certain that the entire piece of nail has been removed as desired. Sharply remove any remaining or swollen/heaped-up skin and all hyperkeratotic debris. After removal of the nail, most clinicians apply a silver nitrate stick to the nail bed and to granulation tissue for 2 to 3 minutes.38 When finished, the nail bed is open and the heaped-up tissue is now flat.

As an option and if phenol or other ablating solutions are available, the following is suggested to permanently ablate the nail bed so that a new nail will not grow (Fig. 51.24). Apply a 10% sodium hydroxide solution to the nail bed with a cotton-tipped applicator for 1 to 2 minutes to provide effective ablation of the nail matrix.35–37 Alternatively, apply a 1% solution of aqueous phenol to the nail matrix beneath the involved area of the lateral nail groove and proximal nail fold with cotton-tipped applicators. A 1% phenol solution can be prepared by diluting a 70% to 90% aqueous phenol solution in an 80:1 ratio (e.g., 8 mL distilled water to 0.1 mL phenol). Remove some of the cotton if the applicator is too bulky to concentrate

**Figure 51.23** Before cutting the nail lengthwise, separate the nail from the nail bed by advancing and spreading iris scissors held parallel to the nail bed. Arrow identifies the technique described in the figure regarding placing the phenol soaked applicator under the eponychium. Note the use of a tourniquet to provide a bloodless field.

**Figure 51.24** Nail ablation technique for the treatment of an ingrown toenail. The lateral portion of the nail is cut and removed (A) to expose the nail bed. Granulation tissue is curetted (B and C), and the nail matrix is cauterized with hydrogen peroxide or phenol (D) (see text).
Nail-Splinting Technique for Ingrown Toenail

A

Insert a wound closure strip under the corner of the affected nail via a sawing motion. Secure the remaining ends of the tape strip to the toe.

B

Insert a piece of small tubing split lengthwise proximally along the lateral nail edge until the nail edge and nail spicule are encircled by the tube. Secure the tube in place with wound closure strips.

Figure 51.25 Splinting techniques for ingrown toenails.

the solution beneath the proximal nail fold and lateral nail groove. Apply thoroughly moistened (but not saturated) applicators for three 30-second applications. Avoid forcing phenol under the remaining nail by rolling the applicator so that it rolls over the matrix and over the nail surface rather than against the split edge of the nail. Although it is necessary to cover the lateral aspect of the nail bed and lateral nail fold, do not allow excess phenol to contact the exposed nail bed or surrounding healthy tissue. After the third application, the cauterized tissue appears brown-tinged or gray. Alternatively, a 1% phenol solution can be applied for 5 minutes. Thoroughly irrigate the cauterized nail bed with water and rub the area with a gloved finger to remove all traces of phenol. Snip away any remaining debris or dead skin with scissors.

Apply antibiotic ointment (not containing neomycin) and a nonadherent dressing to the wound, followed by a dry sterile wrap. Do not forget to remove the tourniquet after the dressing has been applied. Instruct the patient to wash the wound twice daily followed by dry dressing changes. Systemic antibiotics do not hasten wound healing and are not necessary in most cases. Soaking the open wound in warm water twice a day is soothing and allows the patient to view the healing process. The wound will heal in 2 to 4 weeks and may be accompanied by serous drainage for 2 weeks. The patient should be informed of this possibility. Complications include nail regrowth, infection, growth of an inclusion cyst, or delayed healing. If the condition returns, podiatric referral is recommended for more extensive ablation of the nail bed.

Nail-Splinting Technique

Splinting of the nail spicule at the lateral edge of the affected nail may allow the toenail to grow out without affecting the inflamed soft tissue. This technique provides time for the periungual tissue to heal while the nail continues to grow until it can be trimmed straight across. No portion of the nail is removed when the nail is initially splinted. When the degree of inflammation is minimal, elevation of the nail spicule is easily accomplished with forceps or a hemostat. A cotton pledget inserted under the lateral edge to maintain elevation is often sufficient in minor cases. Alternatively, a wound closure strip can be used to elevate the corner of the offending nail. After cleansing the edge of the nail and providing drainage if an abscess is present, insert a wound closure strip obliquely under the corner of the nail with a to-and-fro sawing motion until the corner is sufficiently elevated. Secure the tape closure around the toe (Fig. 51.25A). Instruct the patient to soak the toe in warm water daily, remove the tape closure, and reinsert a new tape strip. This procedure is repeated until the corner of the nail or the nail spicule has grown out and cleared the periungual soft tissue, at which time it can be cut straight across. When the degree of inflammation is moderate, nail splinting is accomplished by using the flexible tube procedure. Obtain a surgical drain (2 to 3 mm in diameter) used to perform percutaneous drainage procedures, or the small tubing in venipuncture kits. Split a 1-cm piece of the drainage tube lengthwise. Perform a digital block and elevate the lateral edge of the nail with forceps or a hemostat. Insert the split drainage tube along the lateral edge of the nail so that it completely encircles the nail spicule and push it proximally until as much of the lateral nail and nail spicule as possible are covered (see Fig. 51.25B). Some authors suggest that the tube be secured with 2-0 suture passed through the nail, but securing the tube with wound closure strips is more easily accomplished (see Fig. 51.25B). Instruct the patient to wash the affected area daily and replace loosened tape strips when needed. When the inflammation and granulation tissue have subsided and the nail spicule has grown sufficiently to not impinge on the periungual soft tissue, the tube splint is removed by the patient and the nail is cut straight across.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Bursitis and tendinitis are terms frequently used to describe a variety of common and often ill-defined regional musculoskeletal conditions characterized chiefly by pain and disability at the involved site. They are either periarticular or contained within specific soft tissue structures. Myofascial pain syndromes are characterized by sensory, motor, and autonomic symptoms that are associated with a trigger point, a hyperirritable point in skeletal muscle that reproduces the patient’s symptoms. These musculoskeletal conditions largely rely on a clinical diagnosis in that they often cannot be confirmed by objective data such as radiographs or laboratory studies. Use of injection therapy with local anesthetics and corticosteroids for bursitis and tendinitis can relieve pain, reduce inflammation, and improve mobility. Injection therapy may provide definitive treatment of a condition or serve as an adjunct to facilitate rehabilitation therapy. Several invasive and noninvasive techniques can be used for the treatment of trigger points and myofascial pain syndromes. Successful treatment of any of these musculoskeletal conditions depends highly on an accurate diagnosis and the use of appropriate techniques.

GENERAL ANATOMIC CONSIDERATIONS

Bursae and Tendon Sheaths

Bursae are small fluid-filled sacs, subcutaneous and deep, which provide a cushion for bone and the surrounding soft tissue. They develop in relation to friction and facilitate the gliding motion of tendons and muscles. There are approximately 78 bursae on each side of the body as described by Monro and Spalteholz.1,2 Adventitial bursae may form in response to abnormal shearing stress at sites subjected to chronic pressure, such as a bunion over the metatarsal head of the great toe. Tendon sheaths are similar in composition to bursae but differ in overall shape. Tendon sheaths are long and tubular, whereas bursae are usually round and flat.

Inflammation of bursae, as in bursitis, can be seen microscopically as a thickening of the normal thin surface of synovial cells lining the bursal wall.3 This thickening coupled with increased production of synovial fluid leads to localized pain and swelling. Aside from simple overuse, bursitis may be caused by local trauma, infection, and crystal deposition. Underlying systemic and autoimmune disorders, such as lupus, rheumatoid arthritis, ankylosing spondylitis, psoriatic arthropathy, and gout, can also lead to bursitis.

Because of the adjacent location of bursae and tendons, an inflammatory process in one may also involve the other.4 Tendinitis and tenosynovitis are used to describe similar inflammatory reactions in tendons and tendon sheaths, respectively.

Some forms of tendinitis may be caused by factors other than overuse, inflammation, trauma, and degenerative disease. Gonococcemia, for example, is one cause of tenosynovitis that should be considered in the appropriate setting.5 Another rare cause is hemodialysis.6 In 2008 the U.S. Food and Drug Administration (FDA) added a black box warning to fluoroquinolone antibiotics, highlighting the potential for tendinopathy and tendon rupture. Clinicians should avoid tendon sheath injections in patients who are taking this class of drugs.7

Common sites of tendinitis and bursitis in the body are depicted in Fig. 52.1.

Trigger Points

Myofascial pain related to trigger points is probably omnipresent, but it is vague and ill-defined in the literature, the specific syndromes are unknown to many clinicians, and the disorders are difficult to clarify in many patients. Hence, myofascial pain originating from trigger points is often attributed to a plethora of other conditions. Consequently, the true incidence is unknown, and few clinicians practice trigger point injection therapy. Examples of misdiagnosis can include fibromyalgia, overuse syndromes, statin-induced myopathy, and malingering.

Trigger points are focal areas of hyperirritability usually found within a taut band of skeletal muscle or in the muscle fascia that are painful on compression and associated with a characteristic pattern of referred pain, motor dysfunction, and autonomic phenomena (Box 52.1).8 Trigger points can also be identified by the local twitch response, a brisk contraction of muscle fibers elicited by snapping palpation or rapid insertion of a needle into the trigger point itself.9

Trigger points probably develop in response to muscle fiber injury. The injury may be an acute traumatic event or more subtle repetitive microtrauma. The underlying pathophysiology has not been fully elucidated but probably involves chronic muscle stress, excessive release of acetylcholine, and dysfunctional motor end plates.10

Trigger points can occur in any muscle or muscle group; they are generally unilateral, but bilateral trigger points have been reported. As the stress associated with myofascial pain commonly affects both single muscles and whole muscle groups, trigger points tend to cluster. In the upper part of the trunk, a common trigger point cluster involves the muscles of the neck and shoulder area, including the trapezius, levator scapulae, and infraspinatus muscles (Fig. 52.2). In the lower part of the trunk, the quadratus lumborum, gluteus medius, and tensor fasciae latae are commonly affected. Trigger points often affect other muscles innervated by the same spinal segments, and subsequent treatment is usually directed at all muscles innervated by both the anterior and posterior branch of the same spinal nerve. Table 52.1 lists common trigger points and their associated myofascial pain syndromes.

RATIONALE FOR INJECTION THERAPY

Bursitis and Tendinopathies

Management of the pain resulting from bursitis and tendinitis may be greatly enhanced by the proper selection and administration of local injections. Successful application of local injection and intrasynovial (bursa and tendon sheath) therapy requires an understanding of the diagnosis, accurate localization of the

1071
should not be viewed as a single quick fix, but a method to facilitate other modalities.

The precise mechanisms for the lasting analgesia and beneficial therapeutic effects of local injection therapy have not been clarified. Few clinical trials have adequately measured the efficacy of corticosteroid therapy. Although steroids are known to reduce inflammation, it is unclear whether the antiinflammatory effect is responsible for the increased range of motion and relief of pain that the patient usually experiences. Histologic studies of chronic tenosynovitis lesions demonstrate degeneration, but not inflammation. It is therefore possible that the pain experienced with tendinitis and bursitis occurs as a result of mechanisms other than inflammation, such as mechanoreceptor stimulation by shearing, traction, or activation of nociceptive receptors by substance P and chondroitin sulfate.

Though often performed in the emergency department (ED) and ubiquitous therapy by orthopedic surgeons, rheumatologists, and family practitioners, injection therapy may not be definitive care. Hence, follow-up and additional evaluations and interventions should be considered. In short, injection therapy should be considered an adjunct to a variety of treatment modalities including pain control, physical therapy, occupational therapy, relative rest, immobilization, and exercise. Additional pain control can be achieved with such options as nonsteroidal antiinflammatory drugs (NSAIDs), acupuncture, ultrasound, ice, heat, and electrical nerve stimulation. Besides pain relief, early participation in rehabilitative activities and exercises can be an important aspect of patient recovery. Patients receiving only analgesics may have worse outcomes than those who also incorporate exercise as part of their treatment. Any factors that provoke the initial injury should also be identified because failure to eliminate these provoking factors can contribute to the injury developing into a chronic condition.

Although opinions in the literature differ, we recommend that corticosteroid injections not be repeated in the same site pathologic condition, and appropriate injection techniques. Lidocaine and corticosteroid preparations may be injected separately or together as adjuncts for pain control. The goal of corticosteroid injection therapy is relief of pain so that the patient is able to regain function and participate in a physical rehabilitation program. In many cases a single injection may be all that is required to ameliorate a painful condition. However, injection therapy is best viewed as an adjunct in the management of painful tendinitis and bursitis syndromes.

**Figure 52.1** Common sites of tendinitis and bursitis. (Redrawn from Walker LG, Meals RM: Tendinitis: a practical approach to diagnosis and management, J Musculoskel Med 6:24, 1989.)

**Box 52.1 Clinical Characteristics of Myofascial Pain Syndromes**

- Characteristic distribution pattern of the pain
- Restricted range of motion with increased sensitivity to stretching
- Weakened muscle because of pain with no muscular atrophy
- Compression causing pain similar to the patient’s chief complaint
- A palpable taut band of muscle correlating with the patient’s trigger point
- Local twitch response elicited by snapping palpation or rapid insertion of a needle
- Reproduction of the referred pain with mechanical stimulation of the trigger point
- Associated autonomic phenomena

**Figure 52.2** Common trigger point clusters on the back. In the upper part of the trunk, a common trigger point cluster involves the muscles of the neck and shoulder area including the trapezius, levator scapulae, and infraspinatus muscles. The quadratus lumborum, gluteus medius, and tensor fasciae latae are commonly affected lower trunk muscles.
### TABLE 52.1 Myofascial Pain Syndromes

<table>
<thead>
<tr>
<th>MUSCLE</th>
<th>TRIGGER POINT LOCATION</th>
<th>AREA OF PAIN</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levator scapulae</td>
<td>Superior medial aspect of the scapula along the flat muscle belly; insertion sites on the transverse processes of C1-C4</td>
<td>Posterior cervical region, posterior aspect of the scalp, periauricular area</td>
<td>May cause or contribute to headache syndromes in some patients</td>
</tr>
<tr>
<td>Splenius capitis and semispinalis capitis</td>
<td>May occur in any part of these muscles</td>
<td>Over the muscles themselves, head, face</td>
<td>Having the patient point to the area of maximal pain may help localize the trigger point or points; may also cause dizziness</td>
</tr>
<tr>
<td>Trapezius</td>
<td>Angle of the neck, occipital insertion sites</td>
<td>Trapezius muscle itself, occiput</td>
<td>Inject neck trigger points carefully to avoid iatrogenic pneumothorax</td>
</tr>
<tr>
<td>Sternocleidomastoid</td>
<td>Sternal and clavicular origins, occipital insertion site, upper two thirds of the muscle belly</td>
<td>Sternocleidomastoid muscle itself, periauricular area, frontal area, face</td>
<td>May also cause dizziness, ipsilateral ptosis, lacrimation, and conjunctival injection</td>
</tr>
<tr>
<td>Infraspinatus</td>
<td>Anywhere in the infraspinatus muscle</td>
<td>Arm, posterior and lateral aspects of the shoulder</td>
<td>May cause sympathetic hyperactivity; subject to early degeneration</td>
</tr>
<tr>
<td>Rectus abdominis</td>
<td>Most common in the upper 3 segments, less often in the lower muscle segments</td>
<td>Anterior abdominal wall (upper segment), back (lower segments)</td>
<td>Often flare after abdominal surgery</td>
</tr>
<tr>
<td>Pectoralis major and minor</td>
<td>Most common at the insertion site on the anterior medial aspect of the shoulder and in the inferior muscle belly, but may be found anywhere in the muscle</td>
<td>Over the trigger point, upper most part of the muscle, ipsilateral arm</td>
<td>Because trigger points may be found anywhere in the muscle, it is important to search the entire muscle</td>
</tr>
<tr>
<td>Intercostals</td>
<td>Intercostal spaces</td>
<td>Chest (increased during inspiration)</td>
<td>Often flare after chest surgery or trauma; inject carefully to avoid iatrogenic pneumothorax</td>
</tr>
<tr>
<td>Tensor fasciae latae</td>
<td>Muscle belly</td>
<td>Lateral thigh pain to the knee</td>
<td>Tensor fasciae latae trigger points are generally easy to locate</td>
</tr>
<tr>
<td>Anterior tibialis</td>
<td>Upper third of muscle</td>
<td>Anterior aspect of the foreleg and ankle, dorsal surface of the ankle</td>
<td></td>
</tr>
<tr>
<td>Gastrocnemius and soleus</td>
<td>Medial and lateral muscle margins, along the midline of the muscle</td>
<td>Posterior of the knee, Achilles tendon near the heel</td>
<td>Often flare with vascular insufficiency of the lower extremities</td>
</tr>
<tr>
<td>Quadratus lumborum</td>
<td>Along the 12th rib, around the iliac crest, lateral border of the muscle</td>
<td>Area of the 12th rib (especially during deep inspiration), anterior abdominal wall</td>
<td>May accentuate postoperative pain or pain associated with abdominal wall scars</td>
</tr>
<tr>
<td>Gluteus medius</td>
<td>Along the iliac shelf, in severe cases, the entire gluteal ridge may be involved (including the gluteus minimus and maximus from the sacroiliac joint to the anterior superior iliac spine)</td>
<td>Most often cause hip, leg, and lower back pain; may cause remote pain in the cervical region and head</td>
<td>Often associated with sympathetic hyperactivity; common in late-stage pregnancy and in patients with unequal leg lengths</td>
</tr>
</tbody>
</table>

See also Fig. 52.30 for diagrams of trigger points for specific muscles.

---

*TABLE 52.1 Myofascial Pain Syndromes*
unless at least a partial clinical response has occurred. In addition, an injection should not be repeated in the same site more than once every 3 months. Some limit corticosteroid injections at any given site to two or three injections before alternative therapy is pursued. Despite few data on the outcome of repeated injections, these recommendations are generally accepted and may limit the risk for adverse effects.

Though universally practiced and generally considered safe and effective for short-term therapy, there are sparse scientific data defining a true benefit of corticosteroid injections for musculoskeletal conditions. Inflammation is not always the cause of tendinopathy. Although true inflammatory tendinitis may respond quite well to corticosteroid injections, conditions such as posttraumatic shoulder impingement and rotator cuff tears may not benefit from local injection any more than from treatment consisting of rest, time, physical therapy, and NSAIDs. Despite appearing to initially be effective for conditions such as olearon bursitis, lateral epicondylitis, and de Quervain’s tenosynovitis, long-term relief of other conditions is often superior with other modalities. In addition, oral corticosteroids can be as effective as local injection and can be an alternative for emergency clinicians reluctant to perform an injection.

**Trigger Points**

Injection therapy is the most widely accepted and scientifically supported modality for treating trigger point pain. However, because it may place patients at risk for becoming dependent on injection for pain relief, some authors reserve injection therapy for patients who have failed other measures (Box 52.2). Various substances have been used for trigger point injections, including local anesthetics, botulinum toxin, sterile water, and sterile saline. Dry needling, a technique that involves multiple advances of a needle into the muscle at the region of the trigger point, provides as much pain relief as an injection of lidocaine. In fact, in a 2001 systematic review on needling therapies for trigger points, Cummings and White concluded that, based on current medical evidence, “the nature of the injected substance makes no difference to the outcome and wet needling is not therapeutically superior to dry needling.” In support of these findings, it has been proposed that the needle (not the injected substance) reduces trigger point pain by mechanically disrupting dysfunctional activity at the motor end plate. Nevertheless, the addition of a local anesthetic is recommended to reduce the degree of postinjection soreness.

The use of steroids for trigger point injection is controversial and without a clear rationale because there is little evidence to support an inflammatory pathophysiology for trigger point pain. Hence, the use of steroids for trigger point injection is not recommended.

**INDICATIONS AND CONTRAINDICATIONS**

**Bursae and Tendon Sheaths**

The indications for steroid injection are twofold: therapy and diagnosis. Injection therapy offers not only relief of pain, particularly when a local anesthetic is used concurrently, but also a medium to deliver therapeutic agents. In addition to relieving pain, injection therapy may aid in diagnosis. When injecting a bursa, for example, bursal fluid is sometimes collected for laboratory analysis. In addition, relief of pain helps differentiate a localized site of injury from referred or visceral pain.

Absolute contraindications to local injection therapy are limited and include specific infections such as bacteremia, infectious arthritis, periarticular cellulitis or ulceration, and adjacent osteomyelitis (Box 52.3). The procedure is also contraindicated in patients with bleeding disorders. A history of hypersensitivity, either to the corticosteroid or to the vehicle by which it is delivered, is an absolute contraindication. Finally, corticosteroid injections should not be performed in a patient who has a documented osteochondral fracture. Relative contraindications depend on both the clinician’s experience and the indication for the injection. Violation of the integrity of the skin or chronic foci of infection, either locally or in the vicinity of the site of involvement, is a relative contraindication. The procedure is also relatively contraindicated in patients taking anticoagulants, in patients with poorly controlled diabetes, and in those with internal joint derangements or

---

**BOX 52.2** Modalities for Treating Trigger Point Pain

**NONINVASIVE TECHNIQUES**

- Oral medications (NSAIDs, muscle relaxants, neuroleptics, opiates)
- Spray and stretch
- Massage therapy
- Ischemic compression

**INVASIVE TECHNIQUES**

- Injection
- Dry needling

*Physical therapy, transcutaneous electrical stimulation, and ultrasound are adjuncts to both invasive and noninvasive techniques.

**BOX 52.3** Contraindications to Local Injection Therapy

**ABSOLUTE**

- Infection (bacteremia, infectious arthritis, periarticular cellulitis or ulceration, adjacent osteomyelitis)
- Uncontrolled bleeding disorder
- Hypersensitivity/allergy to corticosteroid or vehicle
- Osteochondral fracture

**RELATIVE**

- Anticoagulant therapy
- Joint instability
- Poorly controlled diabetes (steroids raise blood glucose levels)
- Hemarthrosis
- Decubitus ulcers
- Joint prosthesis
- Adjacent abraded skin
- Chronic foci of infection
- Internal joint derangement
- Partial tendon rupture
hemarthrosis. Patients with a preexisting tendon injury may be subject to tendon rupture if the corticosteroid injection relieves the pain and full activity is then resumed. Therefore partial tendon rupture is a relative contraindication.

**Trigger Points**

Consider trigger point injection once a myofascial pain syndrome has been identified (see Box 52.1 and Table 52.1). As mentioned previously, some authors reserve injection for patients who fail noninvasive modalities (see Box 52.2). There are few contraindications to trigger point injection. Overlying infection is an absolute contraindication. Relative contraindications include proximity to sensitive structures, bleeding disorders, anticoagulation, an uncooperative patient, and lack of clinician experience. Dry needling is as effective as injection therapy, so in those with allergy to local anesthetics, the clinician may perform dry needling to avoid the problem.

**HAZARDS AND COMPLICATIONS**

All injection therapies can be expected to cause local effects such as pain at the site, bruising, and hematoma formation. These can be ameliorated with proper technique. Local anesthetics should be mixed with a corticosteroid preparation to increase volume, decrease postinjection pain, and assess the accuracy of bursae and tendon sheath injections. However, some clinicians prefer to inject local anesthetics alone before injecting the corticosteroid. Subcutaneous bleeding may occur occasionally at the site of injection if a venule, an arteriole, or a capillary is penetrated. Warn patients that this may occur and reassure them that the discoloration or hematoma will disappear spontaneously. Advise patients to apply ice packs or cold compresses to the involved area for the first 24 hours. Be aware of the local anatomy and aspirate after injecting every 1 to 2 mL of solution to help prevent inadvertent vascular injection. Penetration into or striking a nerve may cause sharp pain or paresthesia, and the patient should be warned of this possibility in advance.

**Anesthetic Injections.** Injection of a local anesthetic causes discomfort secondary to tissue irritation related to its acidity and from tissue distension caused by infiltration. Mixing the anesthetic solution with sodium bicarbonate 8.4% (9 mL of anesthetic to 1 mL of sodium bicarbonate) can reduce this pain by creating a more physiologic pH. Warming and slowly injecting the anesthetic may also be helpful. Other minor reactions occasionally seen after injection of amide preparations include light-headedness or dizziness, pallor, weakness, sweating, nausea, fainting (rare), and tachycardia. These symptoms usually disappear within a few minutes after the injection and rarely require any treatment except reassurance and application of a cold compress to the patient’s forehead. Frequently, it is difficult to determine whether the symptoms are the result of sensitivity to the drug or a fright (vasovagal) reaction to needles and injections. Always place the patient in a supine, prone, or reclining position during the injection to minimize the effect of a potential vasovagal reaction.

The major hazards with injection of local anesthetics are hypersensitivity reactions and accidental intravenous or intraarterial injection (see Chapter 29). Serious or fatal hypersensitivity to procaine and other regional anesthetic agents is encountered very rarely. This possibility is usually suggested by a previous history of reactions to these compounds. Ester solutions (e.g., procaine, tetracaine) that produce the metabolite para-aminobenzoic acid (PABA) account for the majority of these reactions. Amide solutions (e.g., lidocaine, bupivacaine) are rarely involved, and usually the preservative methylparaben, which is structurally similar to PABA, is responsible. When a definite history of sensitivity is present, use of any agent from that class of anesthetic agents is absolutely contraindicated.

**Corticosteroid Injections.** Corticosteroid injections have been found to be safe procedures with few complications (Box 52.4). Although there is evidence of allergic reactions from corticosteroids given orally and parentally, the possibility of an allergic reaction caused by corticosteroid injection is highly unlikely, and such cases occur infrequently. Nevertheless, the clinician should be aware that anaphylaxis after injection of methylprednisolone acetate has been reported. In addition, an unusual rash after an intraarticular methylprednisolone injection, which appears to be consistent with a delayed type of hypersensitivity, has also been documented. Introducing infection is one of the most serious potential complications, but infections occurring as an aftermath of intrasynovial injections are extremely rare. In a study at the Mayo Clinic involving 3000 injections given in 1 year, no infections were reported. Others have found the risk for infection to be 4.6 per 100,000 intraarticular injections. Although the problem of infection is usually avoided with meticulous attention to aseptic technique, caution the patient to watch for the development of any significant pain, redness, or swelling after local injections. We do not recommend routine prophylactic antibiotic administration after corticosteroid injections.

**BOX 52.4 Potential Side Effects of Corticosteroid and Local Anesthetic**

**INJECTION THERAPY**

**Systemic Side Effects**

- Cushing’s syndrome
- Facial flushing
- Nausea
- Impaired diabetic control
- Menstrual irregularity/uterine bleeding
- Hypothalamic-pituitary axis suppression
- Fall in the erythrocyte sedimentation rate and C-reactive protein level
- Anaphylaxis
- Dysphoria
- Pancreatitis
- Cataracts

**Local Side Effects**

- Postinjection flare of pain
- Skin depigmentation, fat atrophy
- Bleeding, bruising
- Steroid “chalk” calcification
- Steroid arthropathy
- Nerve injury
- Tendon rupture and atrophy
- Joint and soft tissue infection
Local undesirable reactions are usually minor and reversible. After steroid injection, approximately 2% of patients may experience an acute synovitis otherwise known as postinjection flare.\textsuperscript{10,16} This may be slightly more common with methylprednisolone acetate (Depo-Medrol, Pfizer Inc., New York, NY) and less common with triamcinolone acetonide (Kenalog, Bristol-Myers Squibb Company, Princeton, NJ). Characterized by an increase in pain and joint swelling, symptoms usually begin a few hours after steroid injection and can last as long as 3 days. These reactions can be stressful to the patient who has expected relief, rather than exacerbation, from an injection. These reactions may be quite symptomatic but should not be misinterpreted as faulty technique or an infection secondary to the injection. Histologically, steroid crystals have been seen within polymorphonuclear leukocytes, making it a true synovitis.\textsuperscript{13,22} This reaction may be difficult to differentiate from an infection, and infection must be ruled out if the symptoms last longer than 48 hours or are associated with fever, warmth, or other suspicious signs. Postinjection flare appears to be more likely to develop with the more soluble (shorter-acting) steroid solutions and may be related to the carrier in which the steroid is manufactured.\textsuperscript{1,3,24} Limiting activity of the involved area for 2 days after the injection might help reduce the incidence of postinjection flare.\textsuperscript{22} When it does occur, the reaction is usually mild and can be controlled adequately with the application of ice or cold compresses and analgesics as needed.

Rarely, afterpains may occur and last for several hours after an injection. Although the cause is obscure, this phenomenon may result from the trauma of needle insertion, penetration of inflamed tissue, or pressure on adjacent nerves from local swelling or bleeding. Afterpains can usually be relieved by the application of moist or dry heat and analgesics until the pain abates, but it is best prevented by mixing a long-acting anesthetic, such as bupivacaine, with the steroid preparation.

Another relatively minor complication is localized subcutaneous or cutaneous atrophy at the site of the injection.\textsuperscript{11} This problem is chiefly of cosmetic concern and is recognized as a small depression in the skin frequently associated with depigmentation, transparency, and the occasional formation of telangiectases. These changes in the skin occur when injections are made near the surface and some of the injected steroid leaks back along the needle track. The skin depression usually recedes and the skin returns to normal when the crystals of the steroid have been completely absorbed. These changes are usually evident 6 weeks to 3 months after the initial injection and generally resolve within 6 months, although they can be permanent.\textsuperscript{1,11} In the two-syringe technique the anesthetic is injected first, the needle is advanced into the bursa/peritendon area, and the syringe is then exchanged for another to inject the steroid. This technique helps prevent injection site atrophy by avoiding any leakage of the steroid suspension close to the skin's surface.\textsuperscript{26} Use a small amount of lidocaine or normal saline to flush the suspension from the needle before removing it. The Z-tract technique is a method of creating an indirect route from the skin puncture to the ultimate site of the steroid injection.\textsuperscript{26} To perform this technique, insert the needle 0.5 to 1.0 cm from the actual target site. When the needle is halfway through the fat tissue, redirect it to the target site and inject both the anesthetic and the corticosteroid. Injection site atrophy is more likely to occur with preparations that are less soluble and thus longer acting.\textsuperscript{11,13}

Minor skin depigmentation, especially in dark-skinned individuals (e.g., African Americans), occurs in 1.3% to 4% of patients undergoing local corticosteroid injections.\textsuperscript{2} It usually occurs within a few months after the initial injection and resolves several months later.\textsuperscript{28} Causes include leakage of the steroid preparation back along the needle track and poor injection technique. The exact etiology is unclear but may be secondary to reducing the number and activity of melanocytes.\textsuperscript{28} This complication can be limited by applying pressure to the site during withdrawal of the needle. Hydrocortisone would be the preferable agent for superficial injections to minimize depigmentation.

One of the most serious complications after local steroid injection is tendon rupture. In general, the risk is very low (<1%) and appears to be related to the dose used.\textsuperscript{11,13} Some believe that injecting steroids directly into the tendon leads to a decrease in the tendon's tensile strength.\textsuperscript{11,22,29,30} Gray and Gottlieb,\textsuperscript{11} however, noted no cases of tendon rupture after more than 300 tendon sheath injections. We still advise that one be diligent and careful about injecting into the surrounding area of the tendon sheath and not into the tendon substance. Moreover, by using one size of needle and syringe, the operator is more likely to appreciate the increase in resistance when injecting directly into the tendon. We also suggest limiting the frequency of injections to no more than once every 3 months in the same site.\textsuperscript{5,11-11} Tendon rupture is more likely to occur in major stress-bearing tendons in athletes, such as the Achilles tendon and the patellar tendons. For this reason, injection of corticosteroids in these areas should be avoided in the ED.\textsuperscript{23}

There have been reports of accidental nerve injury after corticosteroid injection, particularly of the ulnar nerve (for treating medial epicondylitis) and the median nerve (for treating carpal tunnel syndrome).\textsuperscript{31} In addition, pericapsular calcifications develop in up to 42% of patients undergoing local steroid therapy, although they are generally asymptomatic.\textsuperscript{11,26} Finally, within minutes to hours after injection, approximately 1% of patients may experience facial and neck flushing. This reaction may last a few days, but it is usually a benign and self-limited reaction. Facial flushing seems to be more common with triamcinolone preparations.\textsuperscript{11,13,22}

Systemic absorption of local corticosteroid injections does occur, although at a slower rate than with oral steroids.\textsuperscript{12} As a result patients are at low risk for systemic complications, but they do occur. Specifically, intrasynovial injections of steroids have been shown to suppress the hypothalamic-pituitary-adrenal axis for 2 to 7 days.\textsuperscript{11} This complication is more likely to occur in patients who receive repeated injections in a short period or multiple injections in different sites at one time.\textsuperscript{22} Corticosteroids can also exacerbate hyperglycemia in diabetics.\textsuperscript{22,26} Abnormal uterine bleeding has also been reported.\textsuperscript{11,12,23}

Other potential complications of corticosteroid and local anesthetic injections are outlined in Box 52.4.

### CORTICOSTEROID PREPARATIONS

Commonly used corticosteroid repository preparations for the injection of bursae and tendon sheaths are described in Table 52.2. Local anaesthetics such as lidocaine or bupivacaine can be mixed with the corticosteroid preparation in the same syringe. All corticosteroid suspensions, with the exception of cortisone and prednisone, can produce a significant and rapid antiinflammatory effect (in the synovial spaces). Corticosteroids should not be used for trigger point injections.
Corticosteroid preparations are categorized by their solubility and relative potency. Solutions that are more soluble have a shorter duration of action, primarily because they are absorbed and dispersed more rapidly. The addition of tertiary butyl acetate to the solution causes decreased solubility and therefore a longer duration of action. For example, triamcinolone hexacetonide, the least soluble preparation has the longest duration of action. Because the long duration of action increases its potential for subcutaneous atrophy, some authors use this preparation only for intraarticular injections.

There is little consensus in the literature regarding which corticosteroid to use and what dosage is most appropriate for a given site. Centeno and Moore noted that the choice of injection agent is most dependent on the institution where the clinician trained. In 1995 a survey of 172 rheumatologists found that opinions differ regarding almost every facet of soft tissue and intraarticular injection, including patient preparation, choice of corticosteroid, and postinjection advice. Some clinicians advocate mixing both shorter- and longer-acting corticosteroids in the same syringe with little consideration for the location or type of condition. We do not recommend using longer-acting corticosteroids for soft tissue injections, particularly because of the increased risk for associated atrophy, including atrophy of surrounding structures such as ligaments and fascia. In general, use a short- or intermediate-acting agent for an acute or subacute condition such as bursitis or tendinitis; reserve longer-acting agents such as triamcinolone for chronic and prolonged conditions, including arthritis.

### DOSE AND ADMINISTRATION

#### Bursae and Tendon Sheaths

The dose of any corticosteroid suspension used for intrasynovial injection may be selected arbitrarily. Factors that influence the dosage and expected response include the size of the affected area, the presence or absence of synovial fluid or edema, the severity and extent of any synovitis, and the steroid preparation selected. Dosages may need to be reduced in the elderly.

A useful guideline for estimating dosage is as follows: for relatively large spaces such as the subacromial, olecranon, and trochanteric bursae, use 40 to 60 mg of methylprednisolone acetate or its equivalent. For medium- or intermediate-sized bursae and ganglia at the wrists, knees, and heels, use 10 to 20 mg. For tendon sheaths, such as flexor tenosynovitis of the digits and the abductor tendon of the thumb (de Quervain’s disease), use 5 to 15 mg. Sometimes it may be necessary to give larger doses for an optimal response. Intrabursal treatment of the elbow (olecranon) or the knee (prepatellar) bursae, which contain a considerable amount of fluid, may require 30- to 40-mg doses.

Unlike intraarticular injections for synovitis in patients with chronic joint disease, repeated infiltration for soft tissue conditions such as bursitis and tendinitis is not generally recommended or required. However, if only a partial response occurs or if recurrence develops, a single injection can be repeated as long as one waits at least 12 weeks between injections.

#### Trigger Points

In contrast to injection of bursae and tendon sheaths, use small volumes of local anesthetic, botulinum toxin, sterile water, or sterile saline for trigger point injection. In most cases 1 to 2 mL of the chosen agent is sufficient.

### PREPARATION OF THE SITE

Preparation of the site before injection requires meticulous adherence to aseptic technique. Anatomic landmarks may be outlined with a skin pencil. It is important that the injection site and needle tip remain sterile with use of the no-touch technique, although sterile drapes are not generally considered necessary. For operator protection, universal precautions should be followed.

### TECHNIQUES

#### General Considerations

Before beginning the injection inform the patient of the specific indication or indications for treatment. Describe the procedure, including the risks and complications, and obtain informed consent.
The optimum technique for joint and soft tissue injections has not been firmly established. However, one important aspect of a successful technique is accurate positioning of the needle. Injecting an inflamed synovial space, such as a bursa containing fluid, may be as simple as puncturing a balloon. Aspiration of fluid confirms that the needle has correctly entered the sac. Conversely, direct injection into a painful soft tissue lesion requires consent. Subsequently, document the details of the procedure in the medical record. A written consent form is not standard and its use is best based on institutional or departmental policy.

The material required for local injection procedures includes antiseptic solution, needles, syringes, a hemostat, culture and laboratory tubes, bandages, and sterile gauze (Fig. 52.3). Special trays may be stocked for this purpose. The usual sizes of needles for injection sites and corticosteroid doses are listed in Table 52.3.

### Bursae and Tendon Sheaths

Local skin anesthesia is an option before injection but not universally practiced. For bursae and tendon sheaths, local anesthetics can be injected alone or with corticosteroids mixed together in the same syringe. Use the Z-tract technique to limit the risk for a fistulous tract in the soft tissue. Because the steroid may theoretically precipitate or layer in the barrel of the syringe during the injection, agitate the syringe immediately before using it to optimize its distribution. In addition to minimizing the pain associated with the injection, mixing an anesthetic with the corticosteroid also produces a larger volume for delivery.6

The duration of action of lidocaine is approximately 1 hour, whereas bupivacaine may last 6 to 8 hours. Caution the patient that the local anesthetic effect may wear off within a couple of hours and that the beneficial effects of the corticosteroid may be delayed.

---

**TABLE 52.3** A Guide for Needle Size and Dosage for Injection of Common Regional Disorders

<table>
<thead>
<tr>
<th>DISORDER OR INJECTION SITE</th>
<th>NEEDLE SIZE</th>
<th>USUAL DOSAGE OF METHYLPREDNISOLONE ACETATE (mg)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bicipital tendinitis</td>
<td>1.5–2 inches, 22–25 gauge</td>
<td>20–40</td>
</tr>
<tr>
<td>Calcareous tendinitis</td>
<td>1.5–2 inches, 22–25 gauge</td>
<td>20–60</td>
</tr>
<tr>
<td>Subacromial bursitis</td>
<td>1.5 inches, 22–25 gauge</td>
<td>20–40</td>
</tr>
<tr>
<td>Radiohumeral bursitis</td>
<td>1.5 inches, 22–25 gauge</td>
<td>20–40</td>
</tr>
<tr>
<td>Epicondylitis</td>
<td>1.5 inches, 22–25 gauge</td>
<td>20–40</td>
</tr>
<tr>
<td>Olecranon bursitis</td>
<td>1-1.5 inches, 20 gauge</td>
<td>15–30</td>
</tr>
<tr>
<td>Ganglia on the wrist</td>
<td>1 inch, 22–25 gauge</td>
<td>10–15</td>
</tr>
<tr>
<td>de Quervain's disease</td>
<td>3/4 inch, 22–25 gauge</td>
<td>10–20</td>
</tr>
<tr>
<td>Carpal tunnel syndrome</td>
<td>1–1.5 inches, 22–25 gauge</td>
<td>20–40</td>
</tr>
<tr>
<td>Digital flexor tenosynovitis</td>
<td>3/4 inch, 22–25 gauge</td>
<td>5–10</td>
</tr>
<tr>
<td>Trochanteric bursitis</td>
<td>1.5–2 inches, 22–25 gauge</td>
<td>20–40</td>
</tr>
<tr>
<td>Prepatellar bursitis</td>
<td>1–1.5 inches, 22–25 gauge</td>
<td>15–20</td>
</tr>
<tr>
<td>Anserine bursitis</td>
<td>1–1.5 inches, 22–25 gauge</td>
<td>20–40</td>
</tr>
<tr>
<td>Bunion bursitis</td>
<td>1 inch, 22–25 gauge</td>
<td>5–10</td>
</tr>
<tr>
<td>Calcaneal bursitis</td>
<td>1 inch, 22–25 gauge</td>
<td>10–20</td>
</tr>
<tr>
<td>Superficial trigger point</td>
<td>1.5 inch, 22 gauge</td>
<td>N/A</td>
</tr>
<tr>
<td>Deep trigger point</td>
<td>2.0–2.5 inch, 21 gauge</td>
<td>N/A</td>
</tr>
</tbody>
</table>

*Empirical dose. A larger or smaller dose may be used, depending on the clinical scenario.

bAllows bursa aspiration and steroid injection without removing the needle.

N/A, Not applicable.
Trigger Points

Successful treatment of a myofascial pain syndrome requires accurate diagnosis (see Box 52.1 and Table 52.1) and precise identification of the trigger point (a taut band of muscle fibers). There are three generally accepted methods for identification of trigger points: flat palpation, pincer palpation, and deep palpation.1

To perform flat palpation, slide a fingertip across the muscle fibers of the affected muscle group, and at the same time use the opposite hand to retract the skin to either side until the taut band is identified (Fig. 52.4, 1–3). Snap the band like a violin string to precisely identify the trigger point. Pincer palpation involves firmly grasping and rolling the muscle fibers between the thumb and forefinger until the taut band is found (see Fig. 52.4, 4–6). Use deep palpation when the taut band is obscured by superficial tissue. Place a fingertip over the muscle attachment of the area suspected of housing the trigger point and apply pressure in different directions. Reproduction of the patient’s symptoms identifies the trigger point. It may be helpful to mark the trigger point with a skin marker for easy identification before treatment.

Once the trigger point has been found, therapy can be divided into invasive and noninvasive techniques (see Box 52.2). Noninvasive techniques used in the ED include spray and stretch, massage therapy, and ischemic compression therapy.15 Physical therapy, transcutaneous electrical stimulation, and ultrasound treatments are adjuncts that may be arranged by the patient’s primary care physician. Invasive techniques involve injecting the trigger point with a local anesthetic, botulinum toxin, or dry needling.

Noninvasive Techniques

Spray and Stretch

The technique of spray and stretch was once advocated by some as the single most effective treatment of trigger point pain.1 Place the patient in a comfortable position; ensure that the target muscle is well supported and under minimal tension, and that one end of the trigger point is securely anchored. Anesthetize the skin overlying the trigger point with vapocoolant spray (ethyl chloride, dichlorodifluoromethane, or trichloro- monofluoromethane) over the entire length of the muscle. Apply the spray at a 30-degree angle to the skin. After the first pass of spray, apply immediate pressure on the other end of the muscle to create a passive stretch. Perform multiple slow spray passes over the entire length of the muscle while maintaining passive stretch until the muscle achieves a full range of motion. Do not perform more than three repetitions before rewarming the area with moist warm heat, and do not allow each spray to last more than 6 seconds. Educate patients not to overstretch muscles after each therapy session.

Massage Therapy

This technique, as described by Simons and colleagues, uses deep stroking or stripping massage to allow the affected muscle group to be lengthened and relaxed as much as possible.16

Ischemic Compression Therapy

The principle behind ischemic compression therapy is to use pressure to induce ischemia for ablation of the trigger point.
To perform this technique, apply and maintain pressure on the trigger point with increasing resistance until tension in the muscle is relieved. The patient might perceive mild discomfort but not profound pain. Repeat the process for each trigger point encountered.

**Invasive Techniques**

**Injection Therapy**

Almost any trigger point is suitable for injection therapy. Those that fail to respond to noninvasive treatments should be strongly considered for injection. Historically, various substances have been used, including local anesthetics, botulinum toxin, sterile water, and sterile saline. Despite the different compositions, durations of action, and mechanisms of action of these substances, a common finding is that the duration of pain relief following the procedure outlasts the duration of action of the injected substance. As noted earlier, the authors of a 2001 systematic review concluded that based on current medical evidence, the nature of the injected substance makes no difference on the outcome and that wet needling is not therapeutically superior to dry needling. Nevertheless, the addition of a local anesthetic has been shown to reduce the degree of postinjection soreness and is recommended by most authors.

The technique most often recommended for trigger point injection has been referred to as the universal technique. Position the patient in a recumbent position to assist in relaxation of the affected muscles, overall comfort, and prevention of syncope. Re-identify the previously marked trigger point of interest, and scrub the overlying skin with a topical antiseptic solution. For superficial trigger points, use a 22-gauge, 1.5-inch needle. Deeper muscles may require a 21-gauge, 2- or 2.5-inch needle (see Table 52.3). Grasp the skin overlying the trigger point between the thumb and index or middle finger of the non-dominant hand. Insert the needle approximately 1 to 1.5 cm from the trigger point and advance it into the trigger point at a 30-degree angle. Aspirate to confirm that a blood vessel has not been entered, and inject a small amount of the agent. Withdraw the needle to the skin, redirect it to another area of the trigger point, and inject again. Use a fast-in, fast-out approach to elicit a local twitch response, which has been shown to increase the effectiveness of the trigger point injection and allows the entire trigger point area to be treated. Severe cramping or paresthesia suggest inadvertent nerve entry and mandates withdrawal and redirection of the needle. For best results it is critical to elicit a local twitch response with every injection. Following the procedure, the muscle group that was injected should undergo a full active stretch.

**Specific Regions and Clinical Entities**

**Bursitis and Tendinitis**

**Shoulder Region**

Shoulder pain continues to be a common complaint in both the emergency and primary care settings. The pain can be associated with significant disability and may result from pathology that is intraarticular or extraarticular including bicipital tendinitis, calcareous tendinitis, and subacromial bursitis (Fig. 52.5). Because injection (Videos 52.1 and 52.2) is easy and safe to perform, these areas are frequently injected, especially in patients who have failed more conservative therapy such as ice, rest, and oral antiinflammatory medications. Injections may also offer a diagnostic advantage when evaluating a patient with subacromial pain or a rotator cuff syndrome in differentiating shoulder weakness caused by impingement (shoulder strength improves after injection) and a true rotator cuff tear (no change in strength following injection). A potential long-term complication of untreated persistent inflammation is the development of a frozen shoulder (adhesive capsulitis).

**Bicipital Tendinitis (Tenosynovitis) (Fig. 52.6).** The biceps tendon is a common extraarticular cause of shoulder pain and injuries to the tendon can range from inflammation to complete rupture. Bicipital tendinitis and tenosynovitis represent non-specific low-grade inflammation of the biceps tendon and/or its sheath that is more common in those who repeatedly flex the elbow against resistance, such as weight lifters and swimmers. The elderly are more likely to develop degenerative tendinosis and rupture, whereas isolated tendinitis is more common in younger patients.

The bicipital tendon courses through the joint and along the bicipital (intertubercular) groove (see Fig. 52.5), which can be appreciated when the elbow is held at 90 degrees of flexion and the arm is internally and externally rotated. Patients typically complain of pain over the biceps muscle extending into the anterior shoulder. They may have restricted or normal range of motion and normal strength; however, they usually complain of tenderness on palpation over the bicipital groove. Efforts to elevate the shoulder, reach the hip pocket, or pull a back zipper all aggravate the symptoms. Tenderness over the bicipital groove does not confirm the diagnosis, however, because the supraspinatus tendon is in close proximity to the insertion of the bicipital tendon. Other diagnostic clues include the Lippman test, in which rolling the bicipital tendon produces localized tenderness; the Yergason test, which elicits pain along the bicipital groove when the patient attempts supination of the forearm against resistance while holding the elbow flexed at a 90-degree angle against the side of the body; and the Speed test, in which pain is reproduced when the patient resists forward elevation of the humerus against an extended elbow.
so that it touches the bicipital border gently. Deposit the remainder of the drug at this point. With any of these injections, resistance to injection suggests intratendinous needle placement, which should be avoided. If the two-syringe technique is used, instill 1 to 1.5 mL of an intermediate-acting corticosteroid suspension, such as prednisolone tebutate, at the point of maximum tenderness. Inject the anesthetic (2 to 4 mL of 1% lidocaine, or 0.25% bupivacaine) along the upper and lower borders of the tendon. Ultrasound may aid in locating and injecting the tendon sheath.

Calcareous Tendinitis, Supraspinatus Tendinitis, and Subacromial Bursitis. These inflammations are so clinically similar that their symptoms and signs are difficult to differentiate. The musculotendinous rotator cuff is composed of the supraspinatus, infraspinatus, teres minor, and subscapularis muscles, which insert as the conjoined tendon into the greater tuberosity of the humerus. The subacromial bursa lies just

(see Fig. 52.6). Although helpful, these tests are only moderately specific for bicipital tendinitis. Radiographic findings are normal and they are not required if the clinical diagnosis is supported.

**Approach.** Place the patient in a seated position with both hands resting comfortably in their lap. Locate the point of maximal tenderness along the bicipital tendon. Make entry with a 22- or 25-gauge, 3.9- to 5.0-cm needle through an optional lidocaine skin wheal. Avoid an intratendinous injection, which may cause weakening of the tendon and predispose the patient to tendon rupture. Advance the needle along the side of the tendon at a 30-degree angle and aim at one border of the bicipital groove to perform a *peritendinous infiltration* (see Fig. 52.6). Administer one-third of the injection at this point. Withdraw the needle slightly but keep it subcutaneous and redirect it upward approximately 2.5 cm for injection of another third of the drug. Withdraw it again and redirect it downward so that it touches the bicipital border gently. Deposit the remainder of the drug at this point. With any of these injections, resistance to injection suggests intratendinous needle placement, which should be avoided. If the two-syringe technique is used, instill 1 to 1.5 mL of an intermediate-acting corticosteroid suspension, such as prednisolone tebutate, at the point of maximum tenderness. Inject the anesthetic (2 to 4 mL of 1% lidocaine, or 0.25% bupivacaine) along the upper and lower borders of the tendon. Ultrasound may aid in locating and injecting the tendon sheath.
superior and lateral to the supraspinatus tendon (Fig. 52.7). Both the tendon and the bursa are located in the space between the acromion process and the head of the humerus, and are particularly prone to impingement in this “critical zone.” Impingement can occur when the shoulder moves forward and compresses the cuff and bursa under the anterior third of the coracoacromial arch. Injections into either the bursa or the tendon sheath area are commonly performed to relieve inflammation and overuse.

In calcareous (or calcific) tendinitis of the shoulder, a calcific deposit (hydroxyapatite) is located within the substance of one or more of the rotator cuff tendons (commonly the supraspinatus). These calcium crystals can occasionally rupture into the adjacent subacromial bursa and cause acute pain and tenderness in the deltoid area (Fig. 52.8). The bursae in relation to the greater tuberosity and the subdeltoid (subacromial) bursa are the most common sites of calcific deposits.

During the acute or hyperacute stage the patient holds the arm in a protective fashion against the chest wall. The pain may be incapacitating, and all ranges of motion are disturbed, with internal rotation especially limited. Tenderness is often diffuse over the perihumeral region. The patient may also complain of pain at night when lying on the affected side and with abduction of the arm. Supraspinatus tendon impingement is most apparent when the humerus is abducted and internally rotated. The Hawkins test elicits pain with forcible internal rotation while the patient’s arm is passively flexed forward at 90 degrees, and the Neer test elicits pain with full forward flexion between 70 and 120 degrees (Fig. 52.9). Both tests are fairly sensitive but not specific for supraspinatus tendon impingement. Constitutional symptoms are rare, but swelling may sometimes be apparent with the hyperacute form, and a fever and elevated sedimentation rate may even develop in some patients. When shoulder radiographs demonstrate a calcific deposit, the shadow appears “hazy” with lightening of the periphery caused by the pressure of inflammatory edema (see Fig. 52.8). Night pain may be intolerable and often requires opioids for control.

**ANTERIOR APPROACH.** In calcific tendinitis, or supraspinatus tendinitis without calcification, use the anterior (subcoracoid) approach. With the extremity resting on the patient’s lap, externally rotate the arm approximately 15 degrees. The point of insertion is over the depression that is palpable inferior and lateral to the coracoid process and medial to the head of the humerus (see Fig. 52.9).

![Figure 52.7](image1.png)

**Figure 52.7** The “critical zone.” Note the close relationship of the supraspinatus tendon and subacromial bursa to the humeral head and acromion, which make an exact clinical diagnosis very difficult.

![Figure 52.8](image2.png)

**Figure 52.8** Calcareous tendinitis. **A,** In calcareous tendinitis of the shoulder, calcium hydroxyapatite crystals are deposited in the tendons of the rotator cuff and occasionally rupture into the adjacent bursa. **B,** Abnormal calcific deposits in calcareous tendinitis of the shoulder are usually demonstrated roentgenographically in the suprahumeral region or adjacent to the greater tuberosity. When the bursal calcifications appear to be dense (as on this x-ray film), they are frequently asymptomatic, whereas lightening at the margins of the calcium deposit is compatible with the presence of inflammatory edema in the rotator cuff, which produces pain and tenderness in the shoulder region. The location of the calcific deposit on the radiograph may be a useful guide for the point of entry for aspiration and injection. Direct the needle to the calcareous deposit, aspirate, and deposit a portion of the steroid medication there. The calcium may subsequently disappear. Not infrequently, it does so spontaneously.
**Hawkin's test.** With the patient's arm and elbow flexed to 90 degrees, the examiner rotates the forearm internally (i.e., thumb pointed down so that the palm of the hand is directed as far posteriorly as possible). Pain is reproduced if there is impingement of the coracoacromial ligament.

**Neer's test.** With one hand placed on the patient's scapula, the forearm is slowly flexed forward. The arm should be internally rotated such that the thumb is pointing downward. This test causes pain as the greater tuberosity of the humerus impinges on the acromion. Pain may be reproduced between 70 degrees and 120 degrees of forward flexion.

**Anterior Approach**

Externally rotate the arm to 15 degrees. Insert the needle over the depression that is palpable inferior and lateral to the coracoid process and medial to the head of the humerus.

**Posterolateral Approach**

Insert the needle in the depression just inferior to the posterolateral tip of the acromion and superior to the head of the humerus.

**Lateral Approach**

For the lateral approach, insert the needle over the superior aspect of the humeral head and under the lateral margin of the acromion.

**Figure 52.9** Calcaneous tendinitis, supraspinatus tendinitis, and subacromial bursitis. Subacromial bursitis pain is quite common and responds well to injection. Recurrent pain can be due to a rotator cuff tear, which requires evaluation with magnetic resonance imaging. Pain is felt in the deltoid area and can radiate down the arm. The bursa lies mainly under the acromion, but this is variable. Supraspinatus tendinopathy often occurs together with subacromial bursitis.
POSTEROLATERAL APPROACH. With the patient sitting and the lower part of the extremity resting on the lap, make a lidocaine skin wheal at the depression approximately 1 cm inferior to the posterolateral tip of the acromion, located between the head of the humerus and the acromion. Direct a 3.9- to 5.0-cm, 22- or 25-gauge needle toward the center of the head of the humerus and upward at an angle of approximately 10 degrees. Because the bursa does not extend posteriorly beyond the middle of the acromion, it is important that the needle be positioned sufficiently anterior and inferior to the acromion (see Fig. 52.9). After the site has been penetrated 2 to 3 cm, aspirate for any fluid or calcific material. Remove the syringe but leave the needle in position. Attach another syringe containing 20 to 40 mg of methylprednisolone suspension or an equivalent intermediate-acting steroid, and inject the medication. Little resistance should be encountered when injecting the steroid. If resistance is appreciated, reposition the needle because it may be in the tendon substance of the rotator cuff. Follow this injection with 4 to 6 mL of 1% lidocaine (or a similar volume of 0.25% bupivacaine). Alternatively, combine the local anesthetic and steroid in the same syringe. Be generous with the volume of local anesthetic injected to ensure adequate dispersion of the steroid. A single treatment relieves the majority of acute disorders. An injection into the peritendinous space is similar to that described previously except that the needle is advanced deeper than with a subacromial bursal injection.

If calcific tendinitis is suspected, some recommend attempting to aspirate the calcium deposits. After the bursa has been anesthetized, use an 18-gauge needle to penetrate the calcium, which often creates a gritty sensation. Barbotage can facilitate cleavage of the calcium deposits and obtain greater dissemination of the injection. Use this method as described previously by aspirating and reinjecting the steroid or anesthetic-steroid combination repeatedly. Needleling followed by aspiration and lavage of the subacromial space has also been proven to be effective.

Sometimes a painful reaction develops after the analgesic has worn off. Warn the patient about this possibility and give appropriate analgesia. A sling may provide additional relief. Short-term use of opioids is also appropriate. Although some authors claim that patients who do not also undergo physical therapy after corticosteroid injection have satisfactory results, some evidence supports the importance of close patient follow-up, range-of-motion exercises, and physical therapy for total recovery.

Acromioclavicular Joint Inflammation. Pain arising in the acromioclavicular (AC) joint can be the result of an acute injury, such as falling on an outstretched hand or from repetitive strain injuries and degeneration. With this injury, all ranges of motion of the shoulder cause pain, and the joint is tender but rarely swollen. Tenderness over the AC joint is sensitive but not specific for AC joint pathology. Be aware that an obvious deformity or mechanism of injury may suggest AC separation or dislocation. With AC joint inflammation, crepitus is not uncommon. Adduction of the arm across the body with forward elevation to 90 degrees (the cross-arm test) may exacerbate the pain because the AC joint is compressed with this motion.

In a study by Jacob and Sallay, injection of corticosteroids provided short-term relief of symptoms but did not alter the long-term course of patients with AC joint arthropathy. However, a 2008 prospective study found that improvements in pain and function after injection of the AC joint can last up to 12 months. Nevertheless, because many patients obtain relatively short-term pain relief from injection therapy, some clinicians believe that AC joint injection should be performed only in patients with persistent pain despite a trial of rest, oral antiinflammatory medications, and modification of activity.

Approach. Make an entry through an optional cutaneous lidocaine wheal over the interosseous groove at the point of greatest tenderness (Fig. 52.10). This is usually at the AC joint, which is palpated as a small V-shaped depression posteriorly.
at the most lateral aspect of the clavicle.\(^{40}\) In this area the joint line is relatively superficial. Advance a 2.2- to 2.5-cm, 22- or 25-gauge needle approximately 5 mm and inject 1 to 2 mL of lidocaine and 5 to 10 mg of a prednisolone suspension. It is not necessary to advance the needle beyond the proximal margin of the joint surface.

Elbow Region

The elbow is subject to characteristic extraarticular disorders, including radiohumeral bursitis, lateral and medial epicondylitis (tennis elbow and golfer’s elbow), and olecranon bursitis (barfly’s elbow).

Radioumernal Bursitis, Lateral Epicondylitis, and Medial Epicondylitis. Radiohumeral bursitis occurs at the juncture of the radial head and the lateral epicondyle of the elbow. This condition is commonly found in combination with lateral epicondylitis, which is thought to result from repetitive microtrauma at the insertion of the extensor carpi radialis longus and extensor digitorum longus muscles. The symptoms of the two adjacent problems are indistinguishable, but tenderness overlies the radiohumeral groove with bursitis, whereas tenderness occurs chiefly at the lateral epicondyle with tennis elbow (Fig. 52.11). Although the term epicondylitis suggests an inflammatory cause of the pain, some evidence indicates that the injury in lateral epicondylitis results from a degenerative process causing a “tendinosis.”\(^{55}\) Regardless of the exact pathophysiology, there is often a history of repetitive motion of the wrist (flexion, extension, supination, pronation, or any combination thereof) such as while playing tennis, gardening, or using tools.\(^{6}\) A clinical sign supporting the diagnosis of tennis elbow is provocation of pain when the patient attempts extension of the middle finger against resistance with the wrist and elbow held in extension. Alternatively, pain is reproduced at the elbow when the patient is asked to extend the wrist against resistance.

Medial epicondylitis (golfer’s elbow) is a similar condition, although it occurs on the side opposite that of lateral epicondylitis and is much less common (Fig. 52.12).\(^{4}\) This condition involves the origin of the pronator teres and flexor carpi radialis muscles. On physical examination the patient usually complains of pain when the wrist is flexed against resistance or when the forearm is pronated. Palpation of the medial epicondyle also elicits tenderness.

There is evidence supporting the short-term efficacy of corticosteroid injection for both lateral and medial epicondylitis.\(^{54–62}\) Successful injection of lateral epicondylitis produces a predictable short-term (usually less than 6 weeks) improvement in pain that is superior to that with nonsteroidal drug therapy and physical therapy.\(^{55–59,62}\) However, after 6 weeks, physical therapy reduces symptoms more than corticosteroid injection does.\(^{61}\) Nevertheless, some clinicians still feel corticosteroid injection should be a first-line treatment for lateral epicondylitis.\(^{61}\) A similar effect was noted with medial epicondylitis; at 6 weeks patients injected with corticosteroids also reported significantly less pain than those receiving physical therapy alone.\(^{57}\) However, at 3 months and 1 year there were no significant differences in pain between groups receiving corticosteroids and physical therapy and physical therapy alone.\(^{55}\)

**APPROACH.** For lateral epicondylitis, pronate the patient’s forearm and flex the elbow to 90 degrees (Fig. 52.13). Palpate the radial head as a bony protrusion just distal to the epicondyle (confirm identification of the radial head by rotating the patient’s forearm). The entry site is at the point of maximal tenderness, which is usually found at a location slightly distal to the lateral epicondyle. Using a 3.9-cm, 22-gauge needle, deposit 20 to 30 mg of methylprednisolone or equivalent intermediate-acting steroid mixed with anesthetic through a lidocaine skin wheal. Alternatively, follow the steroid with 1 to 3 mL of local anesthetic. Inject the solution in a fanlike pattern while avoiding direct tendon injection. For radiohumeral bursitis, instill part of the repository preparation into the radiohumeral bursa and

---

**Figure 52.11** Lateral epicondylitis. Commonly known as tennis elbow, this condition is common and quite painful and the result of microscopic rupture and incomplete tendinous repair of the extensor carpi radialis brevis origin on the lateral epicondyle of the humerus. Pain usually occurs over the lateral humeral epicondyle during work or recreation. Although this condition often responds well to an injection, recurrences are common. Physical therapy is indicated for recurring pain.

**Figure 52.12** Medial epicondylitis. Also known as golfer’s elbow, medial epicondylitis is a flexor tendinitis with pain in the medial aspect of the elbow that is elicited by flexing the wrist. This condition is much less common than the lateral variety.
part at the lateral epicondyle (see Fig. 52.11). With medial epicondylitis (golfer’s elbow) use a similar technique, but take care to avoid the ulnar nerve, which lies in the ulnar groove behind the medial epicondyle (Fig. 52.14). Damage to this nerve during steroid injection has been reported. Subcutaneous injections should also be avoided because they can result in skin depigmentation, atrophy, or both.

**Olecranon Bursitis (Aseptic).** Olecranon bursitis is an inflammation of the olecranon bursa of the elbow, located between the skin and the olecranon process (Fig. 52.15). Swelling of the bursa is easy to detect given its superficial location and can be differentiated from an elbow joint effusion by preserved elbow extension and flexion. The most common cause of olecranon bursitis is minor trauma, or activities that involve chronic leaning or repetitive elbow motion. As a result of the latter, olecranon bursitis is also known as **barfly elbow** or **student’s elbow.** It may also be seen after an AstroTurf rug burn of the elbow during sporting activities. Other patients at risk for olecranon bursitis include gardeners, auto mechanics, carpet layers, gymnasts, and wrestlers. More significant trauma, such as a direct blow to the elbow, can also cause olecranon bursitis. In this case, hemorrhage into the bursa results in acute hemorrhagic bursitis. Less common causes of olecranon bursitis include hemodialysis and systemic diseases such as rheumatoid arthritis, lupus, uremia, and gout.

Although most cases of olecranon bursitis are sterile, the olecranon bursa is the most frequent site of septic bursitis. Therefore, it is important to accurately differentiate between the two entities. Steroid injections are absolutely contraindicated in cases of confirmed or suspected septic bursitis. Frequently the diagnosis will be suggested by the history and physical examination, but it may be necessary to aspirate and analyze the fluid if septic bursitis is suspected.

In aseptic olecranon bursitis, findings on radiographs are usually normal, but soft tissue swelling may be evident. Bony spurs or amorphous calcific deposits may also be seen, especially in older patients. Occasionally with rheumatoid arthritis and gout, nodules, or tophi, may be palpated within the bursal sac. The bursa and surrounding structures are not typically tender, and there is full and painless range of motion of the involved elbow. Signs of infection such as warmth and erythema of the overlying skin are usually absent. It should be noted, however, that pain, warmth, tenderness, and erythema might be present in both septic and aseptic olecranon bursitis. If there is any suspicion of septic olecranon bursitis, aspiration should be performed and corticosteroid injection deferred until an infectious cause has been ruled out.
Aseptic olecranon bursitis may be cosmetically bothersome to the patient but does not usually cause discomfort and may resolve spontaneously. In cases of bursal swelling that is neither tender nor tense, treatment is symptomatic and includes NSAIDs, compression, and avoidance of further injury. In cases of acute hemorrhagic bursitis, aspiration of the bursa followed by application of a compression dressing and ice will decrease the incidence of chronic bursitis. In patients with aseptic olecranon bursitis who have large, tense, and inflamed bursae, aspiration with steroid injection (after infection has been excluded) has been shown to hasten the resolution of symptoms. Smith and colleagues demonstrated the superiority of intrabursal methylprednisolone acetate over oral naproxen or placebo at 6 months, and noted faster resolution and less reaccumulation of fluid with the steroid injection. The addition of a course of an oral NSAID after steroid injection did not affect the outcome. Aspiration of the olecranon bursa, followed by injection of corticosteroids into the elbow joint, is an alternate approach that has been associated with good success and fewer complications. Following aspiration and steroid injection, application of a compression dressing and a brief period of relative immobilization may be helpful. Repetitive steroid injections for aseptic olecranon bursitis have been associated with triceps rupture and should be avoided.

Olecranon Bursitis (Septic). Because of its superficial location, the olecranon bursa is a common location for septic bursitis along with the pre- and infrapatellar bursa. Infection of the other bursa is much less common. The infection is most likely caused by direct percutaneous inoculation of common skin organisms into the bursa as a result of trauma or contiguous spread from an overlying cellulitis. Septic bursitis secondary to hematogenous spread is rare. Most cases of septic bursitis are caused by Staphylococcus aureus (80%), followed by streptococcal organisms. Other less common organisms include coagulase-negative staphylococci, enterococci, and gram-negative organisms such as Escherichia coli and Pseudomonas aeruginosa. Isolated cases of bursitis caused by fungi (Aspergillus terreus, Candida lusitaniae), Brucella, and Mycobacterium tuberculosis have also been reported. Such unusual organisms should be considered in cases of septic bursitis that are subacute or chronic, and those discovered in an immunocompromised host.

The most common cause of olecranon septic bursitis is trauma. It has been estimated that as many as 70% of cases of septic bursitis are related to trauma, either chronic and caused by repetitive injury, or acute and often associated with occupational or recreational activities. Other risk factors for the development of septic bursitis include chronic illnesses (e.g., diabetes mellitus, alcoholism), and previous inflammation of the bursa, which occurs with gout, rheumatoid arthritis, and uremia. Infection may follow an injection of corticosteroids into the bursa in up to 10% of cases. At times the diagnosis of septic bursitis can be challenging. Acute gouty olecranon bursitis may have a very similar clinical picture, and often can only be accurately differentiated from septic arthritis by fluid analysis (Fig. 52.16). Other conditions that may mimic bacterial septic olecranon bursitis include acute hematomatoid bursitis, aseptic bursitis secondary to oxalosis induced by dialysis, or infectious bursitis caused by unusual organisms such as Mycobacterium, Serratia marcescens, or fungi.

Approximately one third of all cases of olecranon bursitis are septic. In some cases the diagnosis of septic bursitis is obvious (see Fig. 52.16). The onset of pain and swelling may be quite rapid (over a period of 8 to 24 hours), as opposed to the more gradual onset of aseptic bursitis. The bursa is erythematous, tense, swollen, warm, and very painful. Flexion of the elbow is limited by pain; however, some joint mobility may be present because the bursa does not usually extend into the joint. The patient may report a history of trauma to the area, which may be evident on physical examination. Some patients will also have a fever. Smith and associates found that the infected bursa was generally 2.2°C or warmer than the unaffected elbow.

Aspiration of infected bursal fluid with culture of bacteria from the aspirate confirms the diagnosis of septic bursitis. Fluid is usually easily obtained from the tense bursa and (in
The olecranon bursa is found on the posterior aspect of the elbow and is directly superficial to the olecranon process of the ulna. Trauma is the leading cause of olecranon bursitis, but it may also be related to rheumatoid arthritis, lupus, uremia, and gout. Most cases are sterile, although septic bursitis is a clinical possibility and should be considered.

**Aspiration and Injection of Olecranon Bursitis**

1. Painless swelling over the posterior aspect of the elbow is characteristic of nonseptic olecranon bursitis. The mass is soft, movable, and fluctuant.
2. Using sterile preparation, advance a 20-gauge needle on a 10-mL syringe parallel to the forearm.
3. Aspirate the bursal fluid completely. Compress the bursa during aspiration. Typically, slightly blood-tinged serous fluid is obtained.
4. Change the aspirating syringe while the needle remains in the bursal sac.
5. Inject a long-acting steroid preparation (such as 40 mg of methylprednisolone). Use an elastic bandage to compress the site for 12 hours.

Figure 52.15 Olecranon bursitis.

...
to remember that a low bursal white blood cell count does not exclude a septic cause. Moreover, the sensitivity of Gram stain may be 50% or less. In a study of 200 patients with olecranon bursitis, cell count and Gram stain were not always helpful in the acute evaluation and treatment of new cases of septic bursitis. Therefore if clinical suspicion for septic olecranon bursitis is high, even in the presence of a normal or equivocal fluid cell count or Gram stain, steroid injection should be delayed and empirical antibiotic therapy started until the results of culture are available.

Treatment of septic bursitis includes the use of antibiotics directed against penicillinase-producing Staphylococcus, splinting, warm soaks, and drainage of the bursa. Drainage may be performed by daily needle aspiration until the fluid is sterile. However, open incision and drainage might be required, particularly if the infection is recurrent or refractory. In addition to daily drainage of the bursa as necessary, administer antibiotics for at least 2 weeks. Consider additional antibiotic coverage against methicillin-resistant S. aureus based on the patient’s risk factors and epidemiology. Outpatient therapy with oral antibiotics is generally acceptable, although this approach has been challenged for immunocompromised patients. This decision is generally guided by the clinical appearance of the bursa, in addition to associated comorbid conditions, compliance, and other factors regarding patient care. Response to antibiotic therapy might be slow. Thus it is important to initiate antimicrobial treatment as soon as clinical suspicion of septic bursitis exists. Additional treatment with oral NSAIDs may help reduce the pain. Standard gout medications will generally resolve acute gouty bursitis.

**Wrist and Hand Region**

**Ganglion Cysts of the Wrist or Hand.** These cystic swellings occur frequently on the hands, especially on the dorsal aspect of the wrist (Fig. 52.17). They are easily discovered on physical examination and are usually firm and round, and have a rubbery consistency. When physical examination alone is not diagnostic, transillumination or ultrasound can help differentiate cysts from solid tumors. Ganglion cysts are common and make up approximately 60% of all soft tissue tumors affecting the wrist and hand. They usually develop spontaneously in adults between 20 and 50 years of age, with a female-to-male ratio of 3:1. Ganglion cysts may also be seen on the foot and ankle, generally on the extensor surface.

The etiology of ganglia remains obscure; there is usually no history of trauma. The word ganglion is derived from the Greek word meaning “cystic tumor.” The mesothelium- or synovium-lined cystic structures are attached to or may arise from tendon sheaths or near the joint capsule and do not
15% and are found on the hand and wrist. Flexor tendon sheath ganglia make up the remaining 10% to 15% and are found on the hand and wrist. Figure 52.17 Typical dorsal ganglion cyst of A, the wrist and B, dorsum of the foot.

Extend into the joint itself. Attachment is often by a pedicle. The wall of a ganglion is smooth, fibrous, and of variable thickness. The cyst is filled with a clear, gelatinous, sticky, or mucoid fluid of great density. The viscous fluid in the cyst may sometimes represent almost pure hyaluronic acid. The types of ganglia vary with their location. The most common ganglia are located on the dorsal surface of the wrist and extend into the joint itself. Attachment is often by a pedicle. The wall of a ganglion is smooth, fibrous, and of variable thickness. The cyst is filled with a clear, gelatinous, sticky, or mucoid fluid of great density. The viscous fluid in the cyst may sometimes represent almost pure hyaluronic acid.

The types of ganglia vary with their location. The most common ganglia are located on the dorsal surface of the wrist and extend from the scapholunate joint. These constitute approximately 70% of ganglia. Volar wrist ganglia, which arise over the distal aspect of the radius, constitute another 20% to 25% of ganglia and are often adherent to the radial artery. Flexor tendon sheath ganglia make up the remaining 10% to 15% and are found on the hand and wrist.

Close to 60% of ganglia spontaneously regress and do not require treatment. However, they can sometimes cause pain, loss of function, or weakness (from soft tissue or nerve compression or bone erosion), prompting intervention. In addition, some patients request treatment because of cosmetic concerns. Treatment options include aspiration and surgical excision. Simple aspiration is usually an effective approach, but up to three aspirations may be required for complete resolution. In one study, 69% of 116 patients required only a single aspiration for successful treatment. Two or three aspirations were required in 19%, and only 12% of patients ultimately needed surgical excision. It should be noted, however, that ganglia often recur after aspiration or surgery. In fact, over half of all ganglion cysts that undergo aspiration will recur within one year, and most of those within the first 3 months. Following aspiration there is no apparent benefit to the instillation of steroids, although there may be some value in injecting a combination of a steroid and hyaluronidase. Because aspiration is associated with considerable cost savings and shorter recovery times than surgery, and because there is no difference in recurrence rates, aspiration appears to be the initial treatment of choice. In the event that nonsurgical treatment fails, surgical excision may be indicated. The old treatment of attempting to rupture the ganglion with a heavy book is not advised because of the potential for local injury. Aspiration of volar wrist ganglia should also be undertaken with caution because the radial artery often adheres to the cyst and may be at risk for injury.

**APPROACH.** Following the instillation of 1% lidocaine for local anesthesia, insert a 2.5-cm, relatively large-bore needle (17 to 18 gauge) into the center of the ganglion and aspirate the contents (see Fig. 52.17A). Usually 1 to 2 mL of mucinous fluid can be aspirated. Milk the contents of the cyst toward the aspiration needle to maximize the volume removed, taking care not to stick yourself with the needle. After the cyst is localized by aspiration, if desired (although there is no supporting evidence for this practice) use another smaller needle to administer 10 to 15 mg of methylprednisolone or a combination of methylprednisolone and hyaluronidase (which has been shown to be beneficial). Following aspiration a splint is not usually required, and activity need not be restricted. There is no proven role for routine NSAIDs or oral steroid therapy.

**de Quervain’s Disease and Intersection Syndrome.** de Quervain’s disease, a relatively common disorder, is a stenosing tenovaginitis of the extensor pollicis brevis (EPB) and abductor pollicis longus (APL) tendons of the thumb (Fig. 52.18). Though commonly referred to as a tenosynovitis, which denotes inflammation of the synovial sheaths, this condition is more accurately described as a tenovaginitis, which refers to thickening of the fibrous sheath of the first extensor compartment. In 1912, de Quervain described “thickening of the dense fibrous connective tissues without any fresh sign of inflammation, neither round cell inflammation nor increase in numbers of cells.” Histologically the thickening is caused by the accumulation of mucopolysaccharide within the tendon sheath.

It is commonly thought that the disorder occurs more often after repetitive use of the wrists, especially with a wringing motion. Hence, the syndrome has been called washerwoman’s sprain and often no specific cause is apparent. However, a study by Kay challenged the association between repetitive motion and de Quervain’s disease. In a retrospective study of 100 cases, no strong correlation was found with occupation or history of repetitive activities in patients in whom de Quervain’s disease was diagnosed. Thus it may be possible that repetitive activities exacerbate the pain associated with a condition for which the cause is unclear. Women during pregnancy or within 12 months of childbirth are also frequently affected. Tenderness, enlargement, and occasionally palpable crepitation are elicited just distal to the radial styloid process, where both tendons come together in an osseofibrous tunnel. Patients
or compression (the carpometacarpal grind test) and rotation of the thumb produce pain, the condition is most likely caused by degenerative changes in the carpometacarpal joint of the thumb rather than de Quervain’s disease. It should be noted that gonococcal tenosynovitis of the wrist may mimic de Quervain’s disease, and one should inquire about other symptoms (e.g., sore throat, penile or vaginal discharge, or fever) and carefully look for the characteristic rash of this sexually transmitted disease (see Fig. 52.19).

Patients with mild symptoms may be managed with splinting and NSAIDS, but more severe cases typically require local corticosteroid injection. In one study comparing steroid injection with immobilization and oral NSAID therapy alone, the latter was effective only in a small group of patients with minimal symptoms. In another retrospective study, 84% of 58 patients were effectively managed either with a single injection (60%) or with repeat injections (24%), with only 12% requiring surgical treatment. These data support a meta-analysis of 495 subjects treated for de Quervain’s tenovaginitis in which an 83% cure rate was found with injection alone versus 61% for injection and splinting, 14% for splinting alone, and 0% for rest or NSAIDs. The strikingly favorable response to local injection therapy suggests that surgery to release the tendon sheaths is seldom needed. Interestingly, some evidence suggests that failure of steroid injections may be the result of an anatomic variant in which the EPB is located in a separate synovial compartment.

This is supported by a study in which steroids selectively injected into the EPB tenosynovium resulted in the resolution of symptoms in all 50 patients. Clinical suspicion for this anatomic variant should be raised when previous steroid injections have failed and patients have pain with firm resistance to thumb metacarpophalangeal joint extension (the EPB entrapment test). Consider selective injection into the EPB tenosynovium in these patients.

Accurate injection of the corticosteroid has been found to be an important aspect of the patient’s response to treatment. Using radiographic dye to verify correct corticosteroid placement, researchers have found that when the corticosteroid and anesthetic injection did not successfully reach either the EPB or the APL compartment, the patient did not experience relief of pain. Conversely, when the medication was injected accurately, most patients reported an improvement in symptoms. If available, ultrasound may help guide the injection. Studies from 2012 comparing ultrasound-guided injections with standard blind technique demonstrated higher success rates with the use of ultrasound.

Intersection syndrome is a condition that may easily be confused with de Quervain’s disease. Because the treatment approach and the clinical course of intersection syndrome differ from that of de Quervain’s tenovaginitis, it is important that the clinician also be familiar with this entity. First described in 1841 by Velpea, intersection syndrome describes a clinical entity approximately 4 to 8 cm proximal to the location of de Quervain’s disease (see Fig. 52.18). Although the cause is not yet clear, intersection syndrome is thought to result from inflammation of the second dorsal compartment of the wrist, which houses the extensor carpi radialis longus (ECRL) and extensor carpi radialis brevis (ECRB) tendons. Other possible causes include inflammation of a bursa that develops between the APL and the ECRB tendons, and inflammation of the ECRL and ECRB tenosynovium where they cross the muscle bellies of the APL and EPB.

Figure 52.18 de Quervain’s disease and intersection syndrome. De Quervain’s tenosynovitis occurs in the first dorsal compartment of the wrist secondary to tenosynovitis of the abductor pollicis longus and extensor pollicis brevis tendons. Symptoms, which include pain over the radial styloid, are generally caused by overuse. Intersection syndrome, often confused with de Quervain’s disease, produces symptoms 4 to 8 cm more proximally.

Figure 52.19 This 25-year-old woman had signs and symptoms of tenosynovitis of the wrist that was initially thought to be de Quervain’s disease. A single hemorrhagic papule (arrow) demonstrating a septic embolus was found on the forearm and is a subtle but classic lesion of gonococcal bacteremia. One aspect of this sexually transmitted disease is tenosynovitis. Cervical cultures were positive for Neisseria gonorrhoeae despite the absence of vaginal symptoms.

usually have radial wrist pain that is exacerbated with certain thumb and wrist movements. The condition may be confused with first carpometacarpal arthrosis (osteoarthritis of the thumb) and intersection syndrome. Radiographs will have normal findings in de Quervain’s disease, but it might be appropriate to rule out other pathologic abnormalities. Ultrasound may demonstrate thickening and edema of the synovial sheath, but has not yet been routinely used to make or confirm the diagnosis. Rarely, gonococcal tenosynovitis might simulate this inflammatory condition (Fig. 52.19).

A useful clinical maneuver that indicates de Quervain’s disease is the Finkelstein test (Fig. 52.20). With the patient’s thumb grasped in his or her palm, apply ulnar deviation at the wrist. Severe pain at the site of the affected tendon sheaths indicates a positive test. When performing the Finkelstein test, also palpate the tender area for crepitus. If axial traction
De Quervain’s Disease And Intersection Syndrome

Intersection syndrome is characterized by pain, tenderness, edema, and occasional crepitus 4 to 8 cm proximal to the radial styloid and may be mistaken for de Quervain’s disease. This condition is seen in athletes who play sports that require forceful repetitive wrist flexion and extension, such as rowing, weight lifting, gymnastics, and tennis. Treatment includes rest, NSAIDs, and immobilization with a thumb spica splint in 15 degrees of wrist extension. After a 2- to 3-week trial of splinting, corticosteroid injection therapy is recommended for patients with persistent symptoms. Most patients with intersection syndrome respond well to nonoperative treatment, and surgery is generally reserved for refractory cases.

APPROACH. For injection of de Quervain’s tenosynovitis position the patient’s hand so that the ulnar side of the wrist is on the table and the radial side is facing upward (see Fig. 52.20). Introduce a 2.2-cm, 25-gauge needle at the most tender point (approximately 1 cm distal to the radial styloid) through a lidocaine skin wheal, and inject 10 to 20 mg of prednisolone or an equivalent intermediate-acting steroid suspension mixed with 4 to 5 mL of 1% lidocaine adjacent and parallel to the tendon sheath (peritendinous infiltration). The injection should be under the edge of the first dorsal compartment retinaculum within the first extensor compartment. If firm resistance is met or if needle movement is noted when the patient abducts and extends the thumb, the needle may be in the tendon and should be redirected to prevent intratendinous injection. Because many superficial vessels are present in this area, aspirate before injecting to verify that the needle is not in a blood vessel. Be generous with the injection volume because a common reason for failure is the inability to get medication into both tendon sheaths. Frequently, edema is visible at the radial aspect of the first metacarpal base and at the thumb metacarpophalangeal joint dorsally after successful injection. Most patients with de Quervain’s disease respond well to nonoperative treatment, and surgery is generally reserved for refractory cases.

A, For de Quervain’s disease, insert the needle through the most tender point, usually about 1 cm distal to the radial styloid. B, For intersection syndrome, the target site is again the site of maximal tenderness, which is usually 4 to 8 cm proximal to the radial styloid.

APL, Abductor pollicis longus; EPB, extensor pollicis brevis.

Figure 52.20 de Quervain’s disease and intersection syndrome.
Injection for intersection syndrome is similar to that for de Quervain's disease except that the target site is at the point of maximal tenderness, which is usually 4 to 8 cm proximal to the radial styloid (see Fig. 52.20).

**Carpal Tunnel Syndrome.** Carpal tunnel syndrome (CTS) is the most common nerve entrapment neuropathy of the wrist. Caused by median nerve compression in the fibro-osseous tunnel of the wrist, carpal tunnel syndrome is characterized by pain at the wrist that is associated with pain and paresthesia along the median nerve territory. This includes the medial aspect of the thumb, palmar side of the index and middle fingers, and radial half of the ring finger. Typically, the patient wakes during the night with burning or aching pain, numbness, and tingling. Occasionally the discomfort is extremely severe and causes the patient to seek emergency care. Clinical signs that support this diagnosis include a positive Tinel sign, which is elicited by reproducing the tingling and paresthesia by tapping (with a reflex hammer) over the median nerve at the volar crease of the wrist (Fig. 52.21). In addition, one can perform the Phalen test, described as holding the dorsal sides of the flexed wrists at a 90-degree angle against each other for several minutes to provoke symptoms in the median nerve distribution (Fig. 52.21). The Phalen test is more sensitive than Tinel sign and is more specific for CTS. Severe muscle atrophy of the thenar eminence may develop in advanced or neglected cases. In many patients the disturbance is idiopathic, without a recognizable underlying cause. However, people who participate in repetitive activities of the wrist, such as typing, driving, assembly line work, and racquet sports are at risk for the development of carpal tunnel syndrome. Other conditions associated with CTS include rheumatoid arthritis (sometimes as the initial manifestation), pregnancy, hypothyroidism, diabetes, and acromegaly. CTS is a clinical diagnosis, but nerve conduction studies can help support the diagnosis. Radiography is not indicated.

Initial therapy for carpal tunnel syndrome consists of modification of activity and splinting; the latter is particularly helpful at night. Of the medications sometimes used to treat carpal tunnel syndrome, diuretics, NSAIDs, and pyridoxine have been shown to offer little to no relief. In contrast, the benefits, at least in the short term (e.g., 4 to 6 weeks), of local steroid injections for carpal tunnel syndrome have been well proven. In a 2007 Cochrane review of randomized trials...
by Marshall and colleagues, local steroid injection provided greater clinical improvement 1 month after injection than did placebo and up to 3 months after injection than oral steroid treatment. However, symptoms after local corticosteroid injection were no different from the symptoms after either NSAIDs or splinting at 8 weeks. The benefits of initial steroid injection versus surgical intervention for carpal tunnel syndrome are controversial because there is little scientifically valid information from which to draw any conclusions. In one small study, Hui and associates found that surgery resulted in a better symptomatic outcome (but not grip strength) than did local steroid injection over a 20-week period. It appears that local steroid injection for carpal tunnel syndrome offers improvement in symptoms, but this improvement may not be permanent or long term. Current recommendations include a trial of conservative treatment (including possible steroid injection) for patients who have mild to moderate symptoms and lack thenar wasting. Surgery is indicated for patients with persistent symptoms after conservative treatment and for those with severe weakness of the thumb abductors. Nerve compression should be confirmed by nerve conduction studies before surgery.

**APPROACH.** Insert the needle through a lidocaine skin wheal just ulnar to the palmaris longus tendon and approximately 1 cm proximal to the distal crease at the wrist. The palmaris longus tendon can be appreciated by having the patient pinch all the fingertips together while holding the wrist in a neutral position (see Fig. 52.21). Injecting medial (ulnar) to the palmaris longus is preferred because it avoids accidental injection of the median nerve and superficial veins. Direct a 2.5- to 3.9-cm, 25-gauge needle at a 45-degree angle to the skin surface toward the tip of the middle finger. Advance the needle 1 to 2 cm and inject 20 to 40 mg of methylprednisolone (or another intermediate-acting steroid equivalent) with or without lidocaine along the track and into the tissue space. If the patient complains of paresthesia, or the needle meets resistance during injection, redirect the needle to avoid injecting directly into a nerve or tendon, respectively. Up to 2 weeks may be required for the paresthesia to abate significantly, although it usually takes only a few days for improvement of nocturnal pain. A lightweight wrist splint may hasten recovery. Repeated injections may be given, but if a response is not elicited or permanent after two or three injections decompressive surgery should be considered.

**Trigger Finger (Stenosing Flexor Tenosynovitis).** A trigger or snapping finger is one of the most common problems of the hand, affecting roughly 2% of the general population. It is characterized by a stenosed tendon sheath at the level of the first annular pulley (A1), which is located on the palmar surface over the base of the metacarpal head (Fig. 52.22). In this condition the A1 pulley becomes inflamed, and a nodule develops on the tendon as it gets “pinched” under the constricted sheath. Locking occurs when the involved digit is in flexion and is especially troublesome when the patient awakens in the morning. This can occur in any finger but is seen most frequently in the ring and middle fingers.

Besides thickening and stenosing of the tendon sheath, a trigger finger may also be characterized by flexor tendon synovitis. The tendon sheaths are long and tubular, and the walls are lined with a thin layer of synovial cells. Symptoms develop when the tendon becomes trapped and is unable to glide within the tendon sheath. A nodule or fibrinous deposit may form at a site in the tendon sheath, usually over or just distal to the metacarpal head of the trigger finger. When the digit is flexed, the nodule moves with the tendon proximally, and on extension it gets “stuck” on the pulley thereby leading to intermittent catching of the tendon. The nodule may be palpable on physical examination, but this is not necessary for diagnosis. Carpal tunnel syndrome commonly coexists with a trigger finger and may be caused by tenosynovitis. Common causes of tenosynovitis include trauma, diabetes mellitus, and rheumatoid arthritis, although it may also be a primary and idiopathic disorder.

Conservative treatment including local rest or splinting, application of moist heat, and NSAID therapy is a common initial approach aimed at alleviating pain and allowing for smoother movements of the finger. Splinting involves immobilization of the metacarpophalangeal (MCP) joint with slight flexion for up to 6 weeks and has been shown to be an effective option for those who want to avoid corticosteroid injections. However, if these measures fail to control the symptoms, or if the symptoms are severe, corticosteroid injection is indicated. One double-blind, placebo-controlled, randomized study of trigger finger demonstrated that steroid injections were significantly more effective than placebo. At follow-up 3 weeks after treatment, 9 of the 14 patients receiving a combination of lidocaine and steroid injections were asymptomatic versus 2 of the 10 patients receiving lidocaine injections alone. Similar results were noted in a 2009 Cochrane review, which found that corticosteroid injection with lidocaine was more effective than lidocaine alone at 4 months. Steroid injections are more successful when performed in patients with a palpable nodule or symptoms for less than 6 months. Younger age, insulin-dependent diabetes, involvement of multiple digits, and a history of other tendinopathies is associated with treatment failure. There is also evidence to suggest that the efficacy of steroid injections for trigger fingers diminishes with each subsequent injection at the same site. As a general guideline, if treatment fails after three injections (separated by several weeks or months), consider surgery for treatment of the condition. In addition, patients with insulin dependent diabetes mellitus require surgical release for trigger finger more often than do noninsulin-dependent diabetics. Division of the first annular pulley, digital nerve injury, scarring, and recurrence are well-known complications of surgical release.

**APPROACH.** Preparation of the site before injection requires meticulous adherence to aseptic technique. Rest the patient's hand on a table with the palm facing upward. The injection point is at the base of the finger's flexion crease between the A1 and the A2 tendon pulleys. Using a 2.2-cm, 25-gauge needle, enter the skin at a 30-degree angle and insert the needle into the tendon sheath parallel to the tendon fibers. Inject 0.25 to 0.35 mL of an intermediate-acting corticosteroid suspension mixed with 1.5 to 3 mL of anesthetic (see Fig. 52.22). If resistance is felt on insertion of the needle, an intratendinous location is suggested. Withdraw the needle slightly before injection. Similar injections can be administered in the base of the thumb metacarpal for a snapping thumb. Although injection into the tendon sheath is the goal, Taras and coworkers showed that injection of steroid into the subcutaneous tissue surrounding the tendon sheath provided similar improvement as intrasheath injections. If relapses are frequent or the clinical response is not satisfactory, surgical release is indicated.
CHAPTE 52 Treatment of Bursitis, Tendinitis, and Trigger Points

1095

GTPS is primarily caused by tendinopathy of the gluteus medius or minimus. The principal bursae, when involved, are the subgluteus maximus bursa, the subgluteus minimus bursa, and the gluteus minimus bursa, although other bursae of the hip may be affected (Fig. 52.24). The pain may be acute, but is more often subacute or chronic; frequently, patients have already tried NSAIDs without success and have been assigned a number of incorrect diagnoses. Risk factors include female gender, obesity, and those with underlying lower gait disturbances and scoliosis. The chief locus of the pathologic condition is in the abductor mechanism of the hip. Pain occurs near the greater trochanter and may radiate down the lateral or posterolateral aspect of the thigh and, rarely, into the knee.132 The pain is described as deep, dull, and aching, and often interferes with sleep. Lying on the affected hip, stepping from curbs, and descending steps frequently provoke the pain.

On examination, with the patient lying on the unaffected side, tenderness may be elicited over and adjacent to the greater

Carpal/Metacarpal Inflammation. Overuse and aging can lead to pain at the base of the thumb and fingers. This is especially common in elderly women and can be quite painful. The first metacarpal articulates with the trapezium, a common site for this condition. Injection therapy is usually quite successful (Fig. 52.23).

Hip Region
Greater Trochanteric Pain Syndrome (Formerly Trochanteric Bursitis). Greater trochanteric pain syndrome (GTPS) is the second leading cause of lateral hip pain after osteoarthritis.128 Formerly referred to as trochanteric bursitis, recent imaging and histopathological studies have shown that the actual bursa is rarely involved. GTPS can be confused with other conditions at or near the hip such as sacroiliac disease, hip or femur pathology, and metastases.129-131 The hallmark, though, is the demonstration of discrete tenderness on deep palpation at or adjacent to the greater trochanter.129

Digital Flexor Tenosynovitis (“Trigger Finger”). Trigger finger (stenosing tenosynovitis) can affect any digit, including the thumb, but it is most common in the ring and middle fingers. Palpation of the flexor tendon sheath over the metacarpophalangeal joint often reproduces the symptoms. Injection is often very effective, but the nodule may persist.

In advanced cases, inflammation at the proximal (A1) pulley of the flexor tendon sheath overlying the metacarpophalangeal joint may hold the digit in either a flexed or an extended position. Generally, the tendon becomes thickened either proximal or distal to the pulley, which causes a snapping or a locking phenomenon with finger flexion or extension.

Locate the tendon point at the base of the finger flexion crease, which is located between the A1 and A2 pulleys.

Angle the needle 30 degrees into the involved tendon sheath, parallel to the tendon fibers.
trochanter. In contrast to true hip joint involvement, the Patrick FABERE sign (flexion, abduction, external rotation, and extension) may be negative, and complete passive range of motion is relatively painless. Active abduction when the patient lies on the opposite side typically intensifies the discomfort, and sharp external rotation may accentuate the symptoms. Internal rotation does not usually affect the level of pain. Hip radiographs may demonstrate a calcific deposit adjacent to the trochanter; however, the incidence of this finding is low.

GTPS is a self-limited condition that can improve with conservative measures such as rest, ice, oral acetaminophen and NSAIDS, weight reduction, and correction of length discrepancy. In patients with persistent pain, corticosteroid injection for GTPS can be an effective therapy. In one study, 77% of patients reported improvement of their pain after an injection of betamethasone mixed with lidocaine; 61% of the patients had sustained improvement at 26 weeks. However, in a 2011 randomized controlled trial, the benefits of steroid injection were not seen at 12-month follow-up. Failure of corticosteroid therapy should prompt the clinician to seek alternative diagnoses, such as true hip joint disease, which can easily be confused with GTPS. In addition, although rare, there have been case reports of septic trochanteric bursitis caused by tuberculosis.

**APPROACH.** Place the patient in a supine or lateral recumbent position and identify the site of maximum tenderness for needle entry. Advance a 3.9- to 5.0-cm, 20- or 21-gauge needle perpendicular to the skin until the tip of the needle reaches the trochanter (Fig. 52.25). Withdraw the needle slightly and widely infiltrate the site with 3 to 10 ml of lidocaine and 20 to 40 mg of methylprednisolone or an equivalent steroid.

**Ischiogluteal Bursitis.** *Weaver’s bottom* is a painful disorder characterized by pain over the center of the buttocks with radiation down the back of the leg. This condition is rarely diagnosed initially and is often mistaken for lumbosacral strain, a herniated disk, or a spinal cord tumor. The ischial or ischiogluetal bursa is adjacent to the ischial tuberosity and overlies the sciatic and posterior femoral cutaneous nerves. Patients are usually asymptomatic when standing; however when going to a seated position, the gluteus maximus muscle slides away from the bursa allowing for direct contact between the bursa and subcutaneous tissue. The condition is often associated with a sedentary lifestyle, which is where it gets its nickname. Sitting on hard surfaces, bending forward, and standing or tiptoes may all provoke the pain. Maximal tenderness is present over the ischial tuberosity. At times a soft tissue mass may also be felt in this area. When it is recognized, a skillful intrabursal injection, coupled with a few days rest, usually relieves the extreme pain.
CHAP TER 52  Treatment of Bursitis, Tendinitis, and Trigger Points

1097

Examination may reveal subtle evidence of trauma including abrasions or puncture wounds. Patients with aseptic bursitis may experience the same swelling; however, warmth and tenderness are mild or absent in most patients.

APPROACH. Place the patient in a prone position and insert a 5.0-cm, 20- to 22-gauge needle through a lidocaine skin wheal. Advance the needle cautiously to avoid the sciatic nerve, which lies at a depth of approximately 6.5 to 7.5 cm. If paresthesias occur (indicating contact with a nerve), withdraw and redirect the needle. Inject 5 to 10 mL of lidocaine and 20 to 40 mg of methylprednisolone into the bursa. Because of the close proximity of the bursa to the sciatic nerve, some experts advise against the use of injection therapy in this area, especially as the addition of steroids can cause soft tissue atrophy making sitting uncomfortable.

Knee Region

Prepatellar Bursitis. Housemaid’s knee or nun’s knee is characterized by swelling with effusion of the superficial bursa anterior to the patella (Fig. 52.26). In contrast to intraarticular pathology, passive motion of the knee is fully preserved and the pain is generally mild, except during extreme knee flexion or direct pressure. Although the disorder is usually caused by pressure from repetitive kneeling on a firm surface, it can also develop after direct trauma, and occasionally it is a manifestation of rheumatoid arthritis or gout. Although uncommon, the prepatellar bursa is one of the most frequent sites of septic bursitis. In fact approximately one third of all cases are septic in nature. Moreover, patients with septic prepatellar bursitis may not have the classic signs of infection such as erythema, warmth, or fever, making it difficult to differentiate from aseptic bursitis. The bursal aspirate should therefore always be sent for laboratory analysis.

Acute bursitis is more likely to be septic with the most common bacteria being skin flora; however, gout and direct trauma are also known causes. Chronic bursitis is more likely to be secondary to trauma and less likely infection. Symptoms of septic bursitis include localized redness, swelling, and tenderness over the patella. Fever may be present in 30% of patients. Examination may reveal subtle evidence of trauma including abrasions or puncture wounds. Patients with aseptic bursitis may experience the same swelling; however, warmth and tenderness are mild or absent in most patients.
**Medial Collateral Ligament Bursa.** This bursa is located anterior to the tibia and posterior to the medial collateral ligament. Injury to this bursa often occurs when the patient undergoes a twisting motion with concurrent external rotation of the tibia. Tenderness may be appreciated along the anteroinferior aspect of the medial collateral ligament, and the pain is exacerbated with extension of the knee. This condition may sometimes be confused with a medial meniscus tear, and magnetic resonance imaging might be necessary to differentiate the two conditions. Treatment is usually successful with conservative measures, including relative rest, compression, and NSAIDs.

**Popliteal Cyst.** Baker cysts are herniated fluid-filled sacs of the articular synovial membrane that extend into the popliteal fossa, sometimes through the natural communication between the bursa in the posterior of the knee and the joint itself. These cysts may also be caused by swelling of the medial gastrocnemius or semimembranosus bursae alone. Baker cysts are common with prevalence in the general population of 4.7%. They may occur secondary to trauma and are seen in patients with rheumatoid arthritis, gout, and osteoarthritis. Patients will often complain of *popliteal fossa* tenderness and swelling that may extend into the calf. Activities that involve active flexion of the knee, such as walking or jumping, exacerbate the symptoms.

The clinical manifestation of Baker cysts can mimic that of deep venous thrombosis (DVT). For this reason, symptomatic Baker cysts are also known as *pseudothrombophlebitis syndrome*, and it is important to take great care in differentiating the two. In most studies, 2% to 6% of patients suspected of having DVT actually have a popliteal cyst as the cause of their knee or calf pain. Ultrasound is an important tool in this diagnosis. Many cases of Baker cyst resolve spontaneously over a few weeks. However, treatment may require surgery to correct any articular injury or to remove the cyst.

Smith and colleagues reported safe and effective treatment of symptomatic popliteal cysts with ultrasound-guided aspiration of fluid through a spinal needle, fenestration of the cyst walls and septations, and injection of 1 mL triamcinolone and 2 mL 0.5% bupivacaine into the decompressed remnant. Another approach involves arthrocentesis follow by injection of glucocorticoids directly into the knee joint. In a study of thirty patients with knee osteoarthritis complicated with a symptomatic Baker cyst, a single intraarticular injection of 40 mg triamcinolone acetonide resulted in an improvement in knee pain, swelling, and range of motion, in addition to a reduction in Baker cyst size. Steroid injections should be done with care because of the risk for neurovascular injury.

**Ankle, Foot, and Heel Region**

**Ankle Tendinitis.** This is a relatively uncommon condition that may result from unusual repetitive activity, or rarely from acute trauma. The disorder is differentiated from ankle joint involvement by the lack of pain or restricted motion during passive flexion and extension of the ankle. Active flexion and extension of the toes produces pain and there is tenderness along the involved tendons. Initial treatment consists of rest, NSAIDs, and immobilization, sometimes for several weeks. Some patients may eventually need operative intervention for prolonged symptoms. Local steroid injections have been used successfully in patients who do not respond to conservative measures; however, the risk for tendon rupture is well documented. As a result, tendon sheath injections should be
This patient has had anterior knee swelling for 6 weeks after falling directly onto his knee. This collection is confined to the prepatellar space, and passive range of motion of the knee is preserved.

After careful antiseptic preparation, infiltrate the entry site and the anticipated trajectory of the needle with lidocaine.

Keep the knee extended during the procedure. Insert the needle into the fluctuant area in the space between the skin and the patella; either a medial or a lateral approach may be used.

Once in the bursa, aspirate the fluid. Occasionally, a substantial volume of fluid may be obtained, so use a large syringe.

“Milk” the bursa during aspiration to facilitate fluid removal. The needle may be repositioned if needed because the prepatellar bursa is a multiloculated structure (as opposed to a single cavity).

Note that this patient’s aspirate was grossly hemorrhagic, consistent with his history of knee trauma.

Remove as much fluid as possible. This patient’s knee regained a normal appearance after the procedure.

At the end of the procedure, apply a compressive dressing to reduce recurrence.

**Figure 52.27** Prepatellar bursitis aspiration.
reserved for patients with persistent symptoms despite an adequate trial of conservative therapy, and should be done in consultation with a foot and ankle specialist.

**APPROACH.** Enter the tendon sheath tangentially with a 2.5- to 3.9-cm, 22- or 25-gauge needle and inject approximately 2 to 4 mL of a mixture of corticosteroid (20 to 40 mg of methylprednisolone) and lidocaine.

**Bunion Bursitis.** Hallux valgus deformity of the big toe is more commonly referred to as a bunion. This common medical condition is found in up to 23% of all adults, especially patients over 65 and in women. On rare occasions inflammation develops in the medial bursa of the first metatarsophalangeal joint of the great toe. Initial treatment is conservative and includes shoe modification, stretching, and ice; however, on occasion tense swelling occurs and decompression is required. In these cases, aspiration with culture of the fluid should be performed.

**APPROACH.** If no infection is present, the bursa is injected with 5 to 10 mg of methylprednisolone via a 2.5-cm, 20-gauge needle. Special shoes or an orthopedic correction will be needed if the swelling recurs.

**Heel Pain.** Talalgia may be caused by many different conditions including Achilles tendinitis, retrocalcaneal bursitis, and plantar fasciitis (Fig. 52.29). Additional discussion may be found in Chapter 51. The bursae of clinical significance around the heel include the retroachilles bursa (located in the space between...
the skin and the Achilles tendon), the retrocalcaneal bursa (located between the Achilles tendon and the calcaneus), and the subcalcaneal bursa. Achilles tendinitis or bursitis may be traumatic in origin but is more apt to be part of a systemic disease such as rheumatoid or gouty arthritis. Although a normal Achilles tendon is thick and strong, when affected by an inflammatory arthropathy it is predisposed to degeneration, and because the Achilles tendon is not invested by a full synovial sheath, it is more vulnerable to intratendon instillation. Because of the potential hazard of tendon rupture after local steroid injection, avoid infiltration of steroids into this area. Rather, treat Achilles tendinitis with rest, splinting, and oral NSAIDs.

Retrocalcaneal bursitis is often seen in association with Haglund deformity, a bony ridge on the posterolateral aspect of the calcaneus. The bursa lies anterior to the Achilles tendon and posterior to the calcaneus. Local swelling and tenderness at the posterior aspect of the heel, proximal (and sometimes lateral) to the insertion of the Achilles tendon, characterize this bursitis. Treatment is focused on minimizing pressure on the bony ridge, which includes the use of open-heeled shoes (clogs), bare feet, sandals, or a heel lift. Conservative measures such as ice, oral NSAIDs, and rest are other common treatment modalities. Corticosteroid injections are not recommended because of the risk for Achilles tendon rupture.\(^{152,153}\)

The condition in this region that is most amenable to injection therapy is plantar fasciitis, which is also the most common cause of heel pain in adults.\(^{154}\) The plantar fascia is located deep to the fat layer of the foot and extends from the calcaneus to the base of the digits. It is responsible for support of the arch of the foot, and with passive dorsiflexion of the toes.\(^{154}\) The pain may be relieved with activity. Although patients may have a heel spur, many symptomatic patients do not, and the presence of a heel spur should not be considered pathognomonic for the condition. Although still debated, the pathology is generally thought to originate from microtears in the fascia, often as a result of overuse.\(^{154}\) Obesity may also increase the risk for plantar fasciitis because of the excessive load on the fascia.\(^{154}\)

Most cases of plantar fasciitis eventually resolve with nonsurgical management.\(^{154}\) Treatment begins with elimination of any precipitating activity, relative rest, stretching exercises, arch supports, and night splints. If these conservative measures are not effective, injection of the painful heel will usually provide short-term improvement. In a Cochrane review, steroid injections for plantar fasciitis resulted in significant improvement at 1 month, but not at 3 or 6 months when compared with control groups.\(^{157}\) Ultrasound guidance may improve the accuracy and success rates for injection, but data thus far are conflicting.\(^{158-160}\) It should also be remembered that there is a risk for rupture of the plantar fascia and fat pad atrophy with corticosteroid injections.\(^{152}\) One study reported a rupture rate of close to 10%.\(^{161}\) Therefore caution should be exercised when injecting steroids into a painful heel. In addition, because this condition tends to be chronic or recurring, referral to an appropriate specialist is recommended.

**APPROACH.** Insert a 2.5-cm, 22- to 24-gauge needle at the spot of maximal tenderness on the medial aspect of the heel (see Fig. 52.29). Enter the plantar surface at 90 degrees by sliding into the space at the midpoint of the calcaneus. The tip of the needle should lie in the aponeurosis of the attachment to the os calcis. Inject 1 mL of lidocaine and 10 to 20 mg of methylprednisolone. Injection through the more superficial aspect of the base of the foot should be avoided because it may result in dispersion of medications into the fat pad and produce fat pad atrophy.\(^{155}\)

**Trigger Points**
The technique for trigger point injection (universal technique) is the same regardless of location and is described in detail earlier in the chapter (see section on *Invasive Techniques*).

**Myofascial Headache Syndromes**

Trigger points commonly contribute to the muscle component of many headache syndromes, so it is important to carefully examine the muscles of the entire head, neck, shoulder, and back region for hypersensitive areas when evaluating patients with headaches. Trigger points are often found in the sternocleidomastoid, levator scapulae, and trapezius muscles, and less frequently in the scalp and facial muscles. The posterior strap muscles, including the splenius and semispinalis muscles, are also commonly involved. Trigger points along the muscles of mastication can lead to unilateral headaches, and also ear, tooth, and temporomandibular joint pain.\(^{162}\) Trigger points in the thoracic paraspinal muscles are frequently associated with both migraine and tension headaches, whereas trigger points in the quadratus lumborum and gluteus medius have been associated with unilateral headaches.

**Torticollis.** Myofascial causes of torticollis usually involve the trapezius, sternocleidomastoid, and levator scapulae muscles, either alone or in a synergistic manner. The splenius and semispinalis muscles may also be involved. Carefully inject trigger points in the trapezius muscle to avoid puncturing of a high-rising apical pleura.

**Levator Scapulae Muscle Syndrome.** Painful sensitive foci may occur at the origin of the levator scapulae muscle on the superior medial aspect of the scapula, along the flat muscle belly, or at the insertion of the transverse processes of the first four cervical vertebrae (Fig. 52.30B). Pain is usually referred to the posterior cervical region, the posterior aspect of the scalp, and the periauricular area.

**Splenius Capitis and Semispinalis Capitis Muscle Syndrome.** Pain resulting from the trigger points in the splenius capitis and semispinalis capitis muscles may be located over the muscles themselves or be perceived in the head and face and give rise to headaches and dizziness (see Fig. 52.30B). Trigger points in the splenius capitis and semispinalis capitis muscles may be difficult to pinpoint, so having the patient point to the area of maximal tenderness is extremely helpful.

**Trapezius Muscle Syndrome.** The trapezius muscle is a frequent source of muscle pain and headache, especially at the angle of the neck or at the occipital insertion where trigger points are most commonly located (see Fig. 52.30C). When injecting trigger points at the angle of the neck be careful to not puncture the apical pleura.

**Sternocleidomastoid Muscle Syndrome.** The sternocleidomastoid muscle is also a frequent source of neck pain and headache. Trigger points are most often found at its sternal and clavicular origins and occipital insertions, in addition to
A. Levator scapulae muscle syndrome  
B. Splenius capitis and semispinalis capitis muscle syndrome  
C. Trapezius and sternocleidomastoid muscle syndromes  
D. Infraspinatus muscle syndrome  
E. Rectus abdominis and pectoralis muscle syndromes  
F. Intercostal muscle syndrome  
G. Tensor fasciae latae muscle syndrome  
H. Anterior tibialis muscle syndrome  
I. Gastrocnemius/soleus muscle syndrome  
J. Quadratus lumborum/gluteus medius muscle syndromes  

Figure 52.30 Trigger points. See text for descriptions of the various syndromes.
the upper two thirds of the muscle belly (see Fig. 52.30C). Dizziness, ipsilateral ptosis, lacrimation, and conjunctival injection may accompany trigger points located in the sternoclavomastoid muscle. Pain may involve the muscle itself or be referred to the periauricular, facial, or frontal areas.

**Cervical Myofascial Pain Syndrome.** This syndrome represents a chronic cause of neck pain, sometimes mistaken for radiculopathy. Trigger points can be present in the head, neck, or shoulder area causing referred pain through the cervical area. A nonfocal neurologic exam and sometimes imaging is paramount to exclude more serious causes.

**Myofascial Shoulder Disorders**

Myofascial shoulder pain is frequently misdiagnosed as bursitis. It is important to remember that a painful shoulder may be caused by trigger points, which are most often located in the posterior scapular muscles. Other common sites include the supraspinatus, infraspinatus, and pectoralis major muscles. The teres, deltoid, and triceps muscles are rarely involved. Occasionally, trigger points located in the splenius, semispinalis, and gluteal muscles contribute to shoulder pain syndromes and should be treated if found.

**Scapula Muscles.** When injecting trigger points in the lateral scapular and periscapular muscles, place the patient prone with a pillow under the chest to round the shoulders and facilitate injection. Note the anatomy and boundaries of the shoulder before injection, and warn the patient to not move the shoulder. Stabilize the scapula with the nondominant thumb and fingers to prevent movement of the lower portion of the scapula, which could result in inadvertent puncture of the pleura.

Myofascial pain is also commonly associated with a number of muscle beds at the medial border of the scapula including the rhomboids, the serratus anterior, the subscapular muscles, and the levator scapulae. To inject trigger points in these medially situated muscles, have the patient place the ipsilateral hand behind the back. This will cause winging of the scapula and a safer approach. Direct the needle tangentially for easy access to the serratus anterior and subscapularis muscles. As with the lower portion of the scapula, care should be taken to inject the levator scapulae at an oblique angle that is nearly parallel to the thorax to help reduce the chance of accidentally inducing pneumothorax.

**Infraspinatus Muscle Syndrome.** Because of its multiple functions, this muscle is subject to earlier degeneration than other muscles of the rotator cuff and is more susceptible to trigger points (see Fig. 52.30D). Trigger points in the infraspinatus muscle invariably cause sympathetic hyperactivity and often contribute to dystrophy-like syndromes of the upper extremity. To identify trigger points in the infraspinatus muscle it is important to palpate along the entire length of the muscle bundles, and also across the grain of the muscle (see Fig. 52.4).

**Somatic Visceral Reflex Phenomenon**

Skeletal muscle trigger points may contribute to visceral pain by either induction or continuation of the spinal reflex arc. Visceral sympathetic afferent nerves converge on the same dorsal horn neuron as somatic nociceptive afferent nerves do. Reflex sympathetic efferent nerve activity may result in spasm of the visceral sphincters and cutaneous nociceptors (leading in part to referred cutaneous pain). The rectus abdominis muscle is particularly prone to trigger point development in conjunction with visceral pain. For example, right upper quadrant trigger points are associated with gallbladder disease, left upper quadrant trigger points with esophageal and ulcer disease, right lower quadrant trigger points with dysmenorrhea, and left lower quadrant trigger points with intestinal disorders.

Treatment of abdominal wall trigger points can provide significant relief of somatic and visceral pain in appropriate patients. Consider injecting these trigger points when palpating the trigger point or placing the affected muscle under tension (e.g., performing a sit-up) produces the characteristic visceral discomfort. It should also be noted that abdominal wall trigger points are often accompanied by trigger points in the corresponding posterior paraspinal muscle segment. For example, esophageal spasm is frequently associated with trigger points on the left posterior aspect of the thorax at spinal nerve levels T3 through T6. To provide more effective pain control, treat these trigger points also.

**Rectus Abdominis Muscle Syndrome.** These muscles are frequent sites of anterior abdominal wall pain. They often flare up after abdominal surgery and may be a major cause of postoperative pain in some patients. Abdominal wall trigger points are most commonly found in the upper three segments of the rectus muscle and can be located more easily by placing the patient in a supine position with the head and neck flexed so that the rectus muscles are under tension (see Fig. 52.30F). Pain and tenderness are usually localized directly over the trigger point. Trigger points in the lower rectus segments may be a cause of low back pain. Lower segment trigger points may be associated with trigger points in the corresponding posterior lumbar spinal segments (i.e., L4, L5, S1).

**Pectoralis Major/Pectoralis Minor Muscle Syndrome.** The pectoralis major muscle is a frequent site of myofascial pain, particularly at its insertion on the anterior medial portion of the shoulder (see Fig. 52.30E). The inferior belly of the muscle is also a common area for trigger points, so be sure to carefully search the entire muscle to avoid missing a treatable area. Pain is usually located at the trigger point, but patients with trigger points in the clavicular portion of the muscle may have referred pain in the uppermost part of the muscle, whereas others may experience referred pain in the arm.

**Intercostal Muscle Syndrome.** In patients with musculoskeletal chest pain, palpate the intercostal muscles for areas of tenderness. Pain emanating from the exterior intercostal muscles is usually localized near the site of the trigger point and is emphasized during inspiration (see Fig. 52.30F). Intercostal muscle trigger points often flare after chest surgery or trauma. Exercise extreme care when injecting an intercostal muscle trigger point to avoid entry into the pleural space.

**Knee Region**

**Tensor Fasciae Latae Muscle Syndrome.** Because the tensor fasciae latae muscle is easy to examine, trigger points located here are easy to identify (see Fig. 52.30G). Patients usually complain of pain along the lateral aspect of the thigh as far down as the knee.

**Ankle, Foot, and Heel Region**

**Anterior Tibialis Muscle Syndrome.** Trigger points located in the anterior tibialis muscle usually cause pain along the
anterior aspect of the ankle, but the entire ankle may be involved in severe cases. Trigger points are most commonly found in the upper third of the muscle and typically cause pain in the anterior portion of the leg and dorsal portion of the ankle (see Fig. 52.30H).

**Gastrocnemius/Soleus Muscle Syndrome.** Myofascial pain related to trigger points in the gastrocnemius and soleus muscles is usually located behind the knee and along the Achilles tendon near the heel (see Fig. 52.30J). These trigger points are generally found along the lateral and medial margins of the muscle group or along the midline (or in both areas) and often flare in patients experiencing vascular insufficiency of the lower extremities. One author suggests locating and injecting these trigger points for relief of the pain associated with intermittent claudication.169

**Myofascial Back Pain**
Patients with myofascial back pain syndromes can have presenting symptoms of chronic aching muscle pain, spasms, stiffness, and sometimes weakness. Direct palpation of the back muscles may reveal the presence of trigger points as the true cause, especially in unilateral back pain. The most common trigger points are found in the levator scapulae, quadratus lumborum, and gluteus medius muscles. Syndromes involving the quadratus lumborum and gluteus medius should be considered in all patients presenting with low back pain.

The lumbosacral muscles are also commonly involved in low back pain. Trigger points may occur secondary to nerve root or vertebral spondylosis and cause neuropathic pain. In these cases, patients may experience only minimal or temporary relief from trigger point injections. Hence, if the patient is no better after a reasonable trial of trigger point injections, referral should be made to further evaluate for more invasive treatment.

**Quadratus Lumborum Muscle Syndrome.** The quadratus lumborum is considered a hip hiker and lateral flexor of the spine. It also assists respiratory function by anchoring the 12th rib for the pull of the diaphragm. Trigger points may be found along the 12th rib, around the iliac crest, and along the lateral border of the muscle (see Fig. 52.30J) and are often associated with distress on deep inspiration and 12th rib pain. Pain can be local or referred to the anterior abdominal wall. In addition, these trigger points may accentuate postoperative pain or painful abdominal scars over the lower quadrant.

**Gluteus Pain Syndromes.** The gluteal region is composed of several muscles that allow for extension, rotation, and abduction of the hip joint.170 Trigger points located in the gluteus medius may be the most critical trigger points in the lower extremities (see Fig. 52.30J). They are most commonly found along the iliac shelf, and with extensive involvement along the entire gluteal ridge, including the gluteus minimus and maximus muscles from the sacroiliac joint to the anterior superior iliac spine. Trigger points in the gluteus medius may cause hip pain that mimics greater trochanteric pain syndrome (common during the latter stages of pregnancy), whereas trigger points in the gluteus minimus can cause pain radiating down the lateral and posterior aspect of the leg imitating sciatica.171

Similar to the infraspinatus muscle syndrome, trigger points in the gluteus medius muscle are frequently associated with sympathetic hyperactivity. In addition, when these trigger points flare, they often recruit trigger points in the quadratus lumborum, tensor fasciae latae, and other gluteal muscles, where they induce diffuse low back pain, or in the cervical muscles causing neck pain and headaches. Though less common, isolated gluteus medius pain may also occur and usually projects along the iliac crest into the posterior of the hip, thigh, and calf.

**ACKNOWLEDGMENTS**

The editors and author wish to acknowledge the contributions of Jason Becker and Joshua Markowitz to this chapter in previous editions.
BACKGROUND

Arthrocentesis, the puncture and aspiration of a joint, is an acknowledged, useful procedure that is easily performed in the emergency department (ED). It has been established as both a diagnostic and therapeutic tool for various clinical situations. When performed properly, the procedure offers a wealth of clinical information and is associated with few complications. In the ED it is difficult to make an accurate assessment of an acutely painful, hot, and swollen joint without performing arthrocentesis.

INDICATIONS AND CONTRAINDICATIONS

The indications for arthrocentesis are listed in Review Box 53.1.

Infection in the tissues overlying the site to be punctured is generally considered an absolute contraindication to arthrocentesis. However, inflammation with warmth, swelling, and tenderness may overlie an acutely arthritic joint, and this condition may mimic a soft tissue infection. Once convinced that cellulitis does not exist, the clinician should not hesitate to obtain the necessary diagnostic joint fluid. Known bacteremia is a theoretical relative contraindication because infection can spread to the joint; however, this complication should be weighed against the useful information and culture results gained by fluid analysis. Bleeding diatheses are rarely a relative contraindication, and arthrocentesis to relieve a tense hemarthrosis in bleeding disorders such as hemophilia is an accepted practice after infusion of the appropriate clotting factors. There are few data regarding the safety or dangers of arthrocentesis in patients taking anticoagulants or platelet inhibitors. Studies have demonstrated that the risk for iatrogenic hemarthrosis in patients treated with oral anticoagulants is extremely low, even in those who have international normalized ratios (INRs) as high as 4.5. One prospective trial of 32 patients taking warfarin found no complications after arthrocentesis. Hence, when necessary, arthrocentesis should be performed in patients taking anticoagulants. The value of reversing a coagulopathy with blood components before the procedure is not proved, and clinical judgment should prevail. Prosthetic joints are at high risk for infection, and arthrocentesis should be avoided whenever possible in this situation. However, if an infected prosthesis is suspected, arthrocentesis should be performed.

Articular Versus Periarticular Disease

Periarticular conditions such as trauma, tendinitis, bursitis, contusion, cellulitis, or phlebitis may mimic articular disease and suggest the need for arthrocentesis. Therefore evaluation of acute joint disease requires that the clinician first determine whether the patient’s constellation of signs and symptoms derives from the joint itself or from some other musculoskeletal or periarticular structure. Such a distinction, however, may be difficult, if not impossible to make without analysis of synovial fluid. No specific test or physical finding has high specificity for solving this dilemma; however, some physical findings may prove helpful. A common periarticular structure that can be associated with a joint effusion is a Baker cyst (popliteal cyst).

### Review Box 53.1 Arthrocentesis: indications, contraindications, complications, and equipment.

#### Arthrocentesis

**Indications**
- Diagnosis of septic or crystal-induced arthritis
- Diagnosis of traumatic bony or ligamentous injury
- Instillation of medications for acute or chronic arthritis
- Relief of the pain of acute hemarthrosis
- Determination of communication between the laceration and joint space

**Contraindications**
- **Absolute:**
  - Overlying cellulitis
- **Relative:**
  - Bleeding diathesis

**Complications**
- Introduction of infection
- Bleeding
- Allergy to local anesthetic
- Pain

#### Equipment

- Chlorhexidine or Betadine solution
- Sterile gauze
- Sterling drape
- 18- or 20-gauge needle
- 3-way stopcock
- Lidocaine
- Syringes
the course of a specific tendon. May be elicited with tendinitis, or the pain may be traced along is significant when the patient attempts active motion. Through a range of motion with minimal discomfort, yet pain periarticular inflammation, one can often passively lead a joint specific movements or at specific points around the joint. In localized, and tenderness can be elicited only with certain adequate diagnosis.

commonly radiates into the elbow or the neck. Therefore down the front of the thigh into the knee. Shoulder joint pain may be diffuse or clearly localized to the joint, or it may radiate. Directions, with active and passive motion producing pain. Effusion into the joint occurs posterior to the patella, whereas bursal swelling occurs anterior to it (Fig. 53.1). When considerable articular effusion of the knee is present, the capsule of the joint is distended and an inverted U-shaped swelling of the joint develops. This characteristic shape occurs because the dense patellar ligament prevents distention of the capsule along its inferior border. In addition, with the knee extended a large effusion causes the patella to “float” or lift away from the femoral condyles. Complete extension and flexion are often impossible because of the joint tension produced by the effusion.

Joint effusion causes limited movement of the joint in all directions, with active and passive motion producing pain. The pain arising from a pathologic condition involving a joint may be diffuse or clearly localized to the joint, or it may radiate. Hip pain, for example, frequently radiates into the groin or down the front of the thigh into the knee. Shoulder joint pain commonly radiates into the elbow or the neck. Therefore complete examination of contiguous structures is essential for adequate diagnosis.

In contrast, pain from a periarticular process is often more localized, and tenderness can be elicited only with certain specific movements or at specific points around the joint. In periarticular inflammation, one can often passively lead a joint through a range of motion with minimal discomfort, yet pain is significant when the patient attempts active motion. Crepitus may be elicited with tendinitis, or the pain may be traced along the course of a specific tendon.

**Septic Arthritis**

Acute monoarticular arthritis is a common problem in emergency medicine. Although acute monoarticular arthritis has many causes, septic arthritis is the one requiring most urgent diagnosis and treatment. Infectious arthritis is still relatively frequent, and suspicion of a septic process in the joint is the first step in appropriate management; confirmation requires arthrocentesis and culture of synovial fluid. In the ED, synovial fluid analysis is the diagnostic test most heavily relied on in making the diagnosis of an acute intraarticular infection. Culture remains the most definitive study, although it is not 100% sensitive. Gram stain may be helpful, but a negative Gram stain does not exclude the presence of a joint infection because not all infected joints have a positive Gram stain. Therapeutic arthrocentesis might need to be repeated when treating a septic joint. Such therapy is usually performed on an inpatient basis.

Infection of a joint occurs by one of several mechanisms: hematogenous spread (bacteremia, infective endocarditis, intravenous drug use) from a contiguous source of infection, direct implantation, postoperative contamination, or trauma. Septic arthritis is typically monoarticular with a swollen, erythematous, and painful joint. The noninfectious differential diagnosis includes crystal-induced arthritis, fracture, hemorrhhosis, foreign body, osteoarthritis, ischemic necrosis, and monoarticular rheumatoid arthritis. In addition, osteomyelitis may mimic septic arthritis because of the close proximity of the infected metaphysis to the joint space. In many instances an acutely inflamed joint from gout or other arthritides simply cannot be distinguished from infection clinically. Nonetheless, early diagnosis is essential to prevent complications such as impairment of growth, articular destruction with ankylosis, osteomyelitis, and soft tissue extension.

Because an acutely swollen joint may be indicative of a number of disease entities, a thorough history and physical examination are the cornerstones of evaluation, followed by arthrocentesis (Fig. 53.2). Laboratory findings can be useful in making a diagnosis, as can response to therapy (e.g., the response to empirical antibiotics in gonococcal arthritis is often the only criterion for diagnosis because the organism is difficult to culture from joint fluid). Blood cultures may be positive because joint infections may be due to hematogenous spread. Patients with malignancy (especially leukemia) or those who are immunosuppressed or otherwise debilitated are at particular risk for a septic cause. Infectious arthritis should be considered primarily in these patients, as well as in those with preexisting joint diseases such as rheumatoid arthritis. In general, a swollen joint is not usually injected with corticosteroids until the possibility of infection has been eliminated.

*Neisseria gonorrhoeae*, *Staphylococcus* (including methicillin resistant), and *Streptococcus* are the most frequently identified etiologic agents. *N. gonorrhoeae* is the most common organism causing septic arthritis in adolescents and young adults. Patients older than 40 years and those with other medical illnesses are more likely to have *Staphylococcus* joint infections. In children, *Staphylococcus*, *Streptococcus*, and *Escherichia coli* predominate. *Haemophilus influenzae* was a common cause of pediatric septic arthritis in the past, but widespread use of the conjugate vaccine has reduced *H. influenzae* infection rates to nearly zero. In neonates, staphylococci, Enterobacteriaceae, group B streptococci, and *N. gonorrhoeae* are the most likely organisms. Staphylococcal or pseudomonal infections commonly develop in injection drug abusers. *Salmonella* arthritis is more prevalent in patients with sickle cell disease than in the general population; however, more common organisms still predominate. Prosthetic joints or postoperative infections have high rates of *Staphylococcus aureus*, *Streptococcus epidermidis*, Enterobacteriaceae, and *Pseudomonas* infection.
consists of scattered painless, nonpruritic 0.5- to 0.75-cm macules or papules with necrotic or pustular centers, distributed on the extremities and trunk. Eventually, the infection settles into one or two large joints to yield a purulent arthritis. It would be prudent to consider empirical therapy for MRSA in those suspected of having septic arthritis until the results of culture become available. MRSA-infected joints can be multiple, progress rapidly, and be very destructive of joint tissue and adjacent bone.

Although precise incidence data for nongonococcal septic arthritis have not been established, predisposing factors have been described and include age 60 years or older, diabetes mellitus, rheumatoid arthritis, hip or knee prosthesis, joint surgery, and skin infection. The simultaneous occurrence of gout and septic arthritis is possible, and one should not allow the establishment of a diagnosis of crystal-induced disease to stop a thorough search for infection.

Because N. gonorrhoeae is the most common organism causing septic arthritis, gonococcal arthritis deserves special mention. Disseminated gonococcal infection occurs in 0.5% to 3% of cases of mucosal infection. Gonococcal septic arthritis is more common in women, especially during pregnancy or after menstruation, because women with sexually transmitted gonorrhea infections are more likely to be asymptomatic. The time needed for local infection to disseminate can vary from several days to weeks. Patients will often experience systemic symptoms, including fevers, chills, and malaise, as well as migratory polyarthritis. Gonococcal tenosynovitis without joint involvement occurs in two thirds of patients. Dermatitis is also present in two thirds-of patients (Fig. 53.3). The most common rash consists of scattered painless, nonpruritic 0.5- to 0.75-cm macules or papules with necrotic or pustular centers, distributed on the extremities and trunk. Eventually, the infection settles into one or two large joints to yield a purulent arthritis. Overt urethritis and vaginitis may be absent or overlooked because of concentration solely on the obvious joint pathology. Hence, it is important to realize that disseminated gonococcal infections can be associated with surprisingly minimal or even absent signs and symptoms of a genital infection source. Some joints may become inoculated through hematogenous spread from anal and oral sites of infection. Even though N. gonorrhoeae–infected joint fluid is usually “septic” in character, the yield of positive synovial fluid cultures has ranged from 25% to 50%. Blood cultures appear to be less helpful as they are positive in only 20% to 30% of cases. Because blood and joint fluid culture has a low yield, it would be prudent to culture all possible sites of gonococcal infection, including anal and pharyngeal sites. The organism can often be identified in asymptomatic genitourinary sites, with cultures from the primary mucosal site being positive in up to 80% of infected patients.

A positive Gram stain is immediately diagnostic of septic arthritis. However, Gram stains are positive in only 71% of gram-positive infections, 40% to 50% of gram-negative infections, and 25% of gonococcal infections. An elevated synovial white blood cell (WBC) count and a reduction in synovial fluid glucose may give confirmatory data. However, the synovial fluid WBC count in gonococcal arthritis is often between 10,000 and 20,000 cells/mm. Although mild leukocytosis and an elevated erythrocyte sedimentation rate may occur, normal laboratory values do not exclude infection.

Figure 53.3 A and B, Tophaceous gout. These nodules are painless and full of uric acid crystals. C, The acutely swollen and painful wrist joint in this patient is most likely due to acute gouty arthritis, which can produce fever and leukocytosis. In some cases, joint fluid analysis is the only way to differentiate gout from a septic joint. D, Aspiration of a tophus yields a precipitated, waxy, soft uric acid conglomeration.

The prevalence of community-acquired methicillin-resistant S. aureus (CA-MRSA) mandates special attention. Epidemiologic data on the incidence of CA-MRSA septic arthritis are sparse; however, one 2009 study noted that 50% of synovial fluid cultures in suspected septic arthritis ultimately grew MRSA. It would be prudent to consider empirical therapy for MRSA in those suspected of having septic arthritis until the results of culture become available. MRSA-infected joints can be multiple, progress rapidly, and be very destructive of joint tissue and adjacent bone.

Although precise incidence data for nongonococcal septic arthritis have not been established, predisposing factors have been described and include age 60 years or older, diabetes mellitus, rheumatoid arthritis, hip or knee prosthesis, joint surgery, and skin infection. The simultaneous occurrence of gout and septic arthritis is possible, and one should not allow the establishment of a diagnosis of crystal-induced disease to stop a thorough search for infection.

Because N. gonorrhoeae is the most common organism causing septic arthritis, gonococcal arthritis deserves special mention. Disseminated gonococcal infection occurs in 0.5% to 3% of cases of mucosal infection. Gonococcal septic arthritis is more common in women, especially during pregnancy or after menstruation, because women with sexually transmitted gonorrhea infections are more likely to be asymptomatic. The time needed for local infection to disseminate can vary from several days to weeks. Patients will often experience systemic symptoms, including fevers, chills, and malaise, as well as migratory polyarthritis. Gonococcal tenosynovitis without joint involvement occurs in two thirds of patients. Dermatitis is also present in two thirds-of patients (Fig. 53.3). The most common rash consists of scattered painless, nonpruritic 0.5- to 0.75-cm macules or papules with necrotic or pustular centers, distributed on the extremities and trunk. Eventually, the infection settles into one or two large joints to yield a purulent arthritis. Overt urethritis and vaginitis may be absent or overlooked because of concentration solely on the obvious joint pathology. Hence, it is important to realize that disseminated gonococcal infections can be associated with surprisingly minimal or even absent signs and symptoms of a genital infection source. Some joints may become inoculated through hematogenous spread from anal and oral sites of infection. Even though N. gonorrhoeae–infected joint fluid is usually “septic” in character, the yield of positive synovial fluid cultures has ranged from 25% to 50%. Blood cultures appear to be less helpful as they are positive in only 20% to 30% of cases. Because blood and joint fluid culture has a low yield, it would be prudent to culture all possible sites of gonococcal infection, including anal and pharyngeal sites. The organism can often be identified in asymptomatic genitourinary sites, with cultures from the primary mucosal site being positive in up to 80% of infected patients.

A positive Gram stain is immediately diagnostic of septic arthritis. However, Gram stains are positive in only 71% of gram-positive infections, 40% to 50% of gram-negative infections, and 25% of gonococcal infections. An elevated synovial white blood cell (WBC) count and a reduction in synovial fluid glucose may give confirmatory data. However, the synovial fluid WBC count in gonococcal arthritis is often between 10,000 and 20,000 cells/mm. Although mild leukocytosis and an elevated erythrocyte sedimentation rate may occur, normal laboratory values do not exclude infection.
Isolated nontraumatic hemarthrosis may occasionally be seen by the emergency clinician. An inflammatory reaction may follow intracapsular bleeding, and the proliferative reaction and the hyperplastic synovium formed might predispose patients to recurrent hemorrhage in that joint, especially those with bleeding diatheses. The knee is the most commonly affected joint, followed by the ankle, elbow, shoulder, and hip.\(^1\)

The most common cause of intraarticular hemorrhage in the setting of no or minor trauma is a hereditary clotting factor deficiency such as hemophilia. Hemarthrosis is an infrequent complication of oral anticoagulant therapy but might occur even with prothrombin times within the normal range.\(^19\) Cessation of anticoagulant therapy in these patients must be weighed against the risk for adverse clot formation (e.g., acute cerebrovascular accident). Chronic arthritis does not appear to be a long-term complication in patients with intraarticular bleeding from oral anticoagulant therapy. Hemarthrosis may also be a complication of sickle cell anemia, pseudogout, amyloidosis, pigmented villonodular synovitis, synovial hemangioma, rheumatoid arthritis, and infection.\(^{18,20}\)

Management of acute hemarthrosis depends on the cause. Hemarthrosis associated with oral anticoagulant therapy improves only after use of the oral anticoagulant is discontinued and the prothrombin time returns to normal. Hemarthrosis after trauma is a frequent occurrence. It is most common in the knee and often denotes significant internal damage. A massively swollen knee after trauma is frequently seen with knee dislocation (occasionally with spontaneous relocation) and a tear of the anterior cruciate ligament. Intraarticular fractures can cause a significant hemarthrosis.

Distension of the joint by effusion or hemorrhage causes considerable pain and disability. If the fluid is not removed it is partially absorbed, but part of it may undergo organization and result in the formation of adhesions or bands in the joint. This is one argument for drainage of the joint.\(^2\) Some believe that in an otherwise healthy joint that is subjected to a single traumatic event, even a relatively large hemarthrosis will be spontaneously reabsorbed without significant sequelae and therefore presents no pressing need for drainage. Unfortunately, no literature exists to guide the best approach; hence, there is no universal standard of care regarding the need or lack thereof of draining blood from a traumatic joint.

Nonetheless, a large, tense, traumatic effusion is quite painful and its presence precludes proper evaluation of an injured joint. Therapeutic arthrocentesis to drain a symptomatic traumatic effusion is a common and well-accepted practice.\(^{2,19,21,22}\) The source of blood after trauma is frequently a tear in a ligamentous structure, capsule, synovium, or fracture. Cruciate (especially anterior) ligament injury is the most common cause of significant hemarthrosis after trauma to the knee.\(^{20}\) A joint effusion that develops 1 to 5 days after trauma may be secondary to a slow hemorrhage or reinjury, but the swelling is often caused by a nonhemorrhagic irritative synovial effusion.

Occasionally, one will diagnose an occult fracture by the presence of lipohemarthrosis, or fat globules in the arthrocentesis specimen (Fig. 53.4). This may be appreciated when the bloody effusion is placed in a clear container (e.g., emesis basin) and held to a light. If the history of trauma is vague, arthrocentesis may be required to differentiate hemorrhage from other causes of joint effusion. An occult tibial plateau fracture is an example in which evaluating for lipohemarthrosis

---

**Hemarthrosis**

Isolated nontraumatic hemarthrosis may occasionally be seen by the emergency clinician. An inflammatory reaction may follow intracapsular bleeding, and the proliferative reaction and the hyperplastic synovium formed might predispose patients to recurrent hemorrhage in that joint, especially those with bleeding diatheses. The knee is the most commonly affected joint, followed by the ankle, elbow, shoulder, and hip.\(^1\)

The most common cause of intraarticular hemorrhage in the setting of no or minor trauma is a hereditary clotting factor deficiency such as hemophilia. Hemarthrosis is an infrequent complication of oral anticoagulant therapy but might occur even with prothrombin times within the normal range.\(^19\) Cessation of anticoagulant therapy in these patients must be weighed against the risk for adverse clot formation (e.g., acute cerebrovascular accident). Chronic arthritis does not appear to be a long-term complication in patients with intraarticular bleeding from oral anticoagulant therapy. Hemarthrosis may also be a complication of sickle cell anemia, pseudogout, amyloidosis, pigmented villonodular synovitis, synovial hemangioma, rheumatoid arthritis, and infection.\(^{18,20}\)

Management of acute hemarthrosis depends on the cause. Hemarthrosis associated with oral anticoagulant therapy improves only after use of the oral anticoagulant is discontinued and the prothrombin time returns to normal. Hemarthrosis after trauma is a frequent occurrence. It is most common in the knee and often denotes significant internal damage. A massively swollen knee after trauma is frequently seen with knee dislocation (occasionally with spontaneous relocation) and a tear of the anterior cruciate ligament. Intraarticular fractures can cause a significant hemarthrosis.

Distension of the joint by effusion or hemorrhage causes considerable pain and disability. If the fluid is not removed it is partially absorbed, but part of it may undergo organization and result in the formation of adhesions or bands in the joint. This is one argument for drainage of the joint.\(^2\) Some believe that in an otherwise healthy joint that is subjected to a single traumatic event, even a relatively large hemarthrosis will be spontaneously reabsorbed without significant sequelae and therefore presents no pressing need for drainage. Unfortunately, no literature exists to guide the best approach; hence, there is no universal standard of care regarding the need or lack thereof of draining blood from a traumatic joint.

Nonetheless, a large, tense, traumatic effusion is quite painful and its presence precludes proper evaluation of an injured joint. Therapeutic arthrocentesis to drain a symptomatic traumatic effusion is a common and well-accepted practice.\(^{2,19,21,22}\) The source of blood after trauma is frequently a tear in a ligamentous structure, capsule, synovium, or fracture. Cruciate (especially anterior) ligament injury is the most common cause of significant hemarthrosis after trauma to the knee.\(^{20}\) A joint effusion that develops 1 to 5 days after trauma may be secondary to a slow hemorrhage or reinjury, but the swelling is often caused by a nonhemorrhagic irritative synovial effusion.

Occasionally, one will diagnose an occult fracture by the presence of lipohemarthrosis, or fat globules in the arthrocentesis specimen (Fig. 53.4). This may be appreciated when the bloody effusion is placed in a clear container (e.g., emesis basin) and held to a light. If the history of trauma is vague, arthrocentesis may be required to differentiate hemorrhage from other causes of joint effusion. An occult tibial plateau fracture is an example in which evaluating for lipohemarthrosis

---

**Figure 53.3** A, Often mistaken for insect or spider bites, these multiple embolic skin lesions (ranging from single or multiple petechiae to pustules) may be seen in patients with acutely swollen joints infected with *Neisseria gonorrhoeae*. Skin lesions are usually found on the extremities, especially the feet and hands, and may be present before a large joint effusion accumulates or with gonococcal tenosynovitis. B, A genital manifestation of urethritis (penile discharge) or vaginitis may be seen but can be rather clinically silent. C, Maculopapular rash associated with *N. gonorrhoeae*. 
Intraarticular Corticosteroid Injections

In 1951 Hollander and coworkers first demonstrated that intraarticular corticosteroid injections are useful for relief of symptoms in patients with severe rheumatoid arthritis. The use of steroids has proved to be a dependable method for providing rapid relief of pain and swelling of inflamed joints, although it is strictly local, usually temporary, and rarely curative. It is easily performed in the emergency setting. Acute gout responds well to joint injection, and this may be of particular value. Following therapeutic arthrocentesis, it may be desirable to inject 2 to 15 mL, depending on joint size, of a long-acting local anesthetic (see Table 53.1) into the joint to facilitate examination or provide temporary relief of the symptoms.

Corticosteroid injections are most helpful when only a small number of joints are actively inflamed. The most frequently used corticosteroids for intraarticular injection are shown in Table 53.1. Diminution of joint pain, swelling, effusion, and warmth is usually evident within 6 to 12 hours after injection.

Thoroughly washing the arthrocentesis needle should be done before each withdrawal to prevent the introduction of additional bacteria. Aseptic technique is important. Sterile injectable solutions are used. The needle is inserted at the point where the joint bulges maximally, certain landmarks being important. The most crucial part of arthrocentesis is defining evidence of an effusion. Although one may aspirate successfully at the point where the joint bulges maximally, certain landmarks are important. The most crucial part of arthrocentesis is defining the joint anatomy by palpating the bony landmarks as a guide.

Table 53.1 Intrasyovial Corticosteroid Preparations

<table>
<thead>
<tr>
<th>PREPARATION</th>
<th>LARGE-JOINT DOSE (mg)</th>
<th>SMALL-JOINT DOSE (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Triamcinolone hexacetone</td>
<td>20</td>
<td>2–6</td>
</tr>
<tr>
<td>Triamcinolone acetonide</td>
<td>20</td>
<td>2–6</td>
</tr>
<tr>
<td>Methylprednisolone acetate</td>
<td>40</td>
<td>3.5–10.5</td>
</tr>
<tr>
<td>Triamcinolone diacetate</td>
<td>20</td>
<td>2–6</td>
</tr>
<tr>
<td>Dexamethasone acetate</td>
<td>5</td>
<td>0.5–1.5</td>
</tr>
</tbody>
</table>

Listed in approximate descending order of duration of action.

The dose will depend on joint size, capsular distensibility, and degree of inflammation.

Ultrasound offers a significant advantage when evaluating a patient with a complaint of joint pain. In patients with obesity or significant pain, limited physical examination makes the diagnosis of joint effusion difficult. Attempting blind aspiration in these patients may cause significant pain for the patient and frustration for the clinician. Ultrasound allows the physician to thoroughly evaluate the joint space for the presence of effusion and to plan the best approach for aspiration. The initial evaluation of the major joints is discussed hereafter, followed by a general approach to aspiration.

**Knee**

Although effusions of the knee are frequently diagnosed on physical examination alone and aspirated blindly, ultrasound allows the clinician to distinguish effusion from other conditions that may cause generalized swelling (such as bursitis).

A high-frequency transducer (7.5 to 10 mHz) should be used to allow the greatest resolution. Begin with the indicator pointing toward the patient’s head (in longitudinal orientation) over the anterior aspect of the knee and attempt to locate the patella (Fig. 53.US1). The patella can be seen as a brightly echogenic (white) object with posterior shadowing (Fig. 53.US2). Locating the patella is key to distinguishing prepatellar bursitis, which will appear as a dark, fluid-filled collection superficial to the patella, and a joint effusion, which will appear as a dark, fluid-filled collection deep to the patella. Once the patella has been identified, move the transducer medially or laterally to “look under” the patella into the joint space (Fig. 53.US3). Fluid will appear as a dark gray or black collection between the articular surface of the femur and fibula or tibia (Fig. 53.US4). Once this area has been evaluated, move the transducer superiorly to evaluate the suprapatellar bursa, which lies superior to the patella and deep to the quadriceps tendon. The suprapatellar bursa communicates with the joint space and frequently houses a large amount of fluid (Fig. 53.US5).

**Shoulder**

Either an anterior or posterior approach can be used to evaluate the shoulder. In the anterior approach, place the patient first in a seated position with the elbow adducted and the palm facing up. Then, place the high-frequency transducer in a transverse orientation over the approximate location of the biceps tendon (Fig. 53.US6). The biceps tendon is an extracapsular extension of the joint and will be seen to

**ULTRASOUND BOX 53.1: Arthrocentesis**

by Christine Butts, MD

Ultrasound image of a normal patella. The patella is seen as a brightly echogenic (white) arcing line just beneath the surface (arrow). Prepatellar fluid collections, such as bursitis, will be seen superficial to this area.

Figure 53.US1 Placement of the ultrasound transducer in a longitudinal orientation over the patella. Such placement enables localization of the patella and serves to orient the sonographer.

Figure 53.US2 Ultrasound image of a normal patella. The patella is seen as a brightly echogenic (white) arcing line just beneath the surface (arrow). Prepatellar fluid collections, such as bursitis, will be seen superficial to this area.

Figure 53.US3 Placement of the ultrasound transducer to “look under” the patella into the joint space.

Figure 53.US4 Ultrasound image of the junction of the femur (at the left of the image) and the tibia (at the right of the image) (arrow). When an effusion is suspected, the suprapatellar recess should be evaluated in addition to this space because fluid frequently collects in the potential space superior to this junction.
distend with fluid when a joint effusion is present. A normal tendon can be seen to lie within the biceps groove of the humerus (Fig. 53.US7). When surrounded by fluid, the tendon will be seen to “float” within an anechoic (black) area (Fig. 53.US8).

To evaluate the joint from the posterior approach, place the patient in a seated position with the affected hand on the opposite shoulder to open the joint space. Then, place the transducer at the approximate location of the articulation of the humeral head with the glenoid (Fig. 53.US9). In a normal joint, the humerus can be seen to articulate with the glenoid without any intervening fluid (Fig. 53.US10). When an effusion is present, a dark gray or black collection can be seen medial to the glenoid (Fig. 53.US11).

Ankle

The ankle joint is best evaluated in the longitudinal axis with the transducer placed over the space between the posterior edge of the tibia and the talus (Fig. 53.US12). Slightly plantar-flexing the foot will enable the transducer to “fit” in this space. In a normal joint, the distal edge of the tibia can be seen to articulate with the talus without any intervening fluid (Fig. 53.US13). When an effusion is present, it is seen as a triangular, dark gray or black pocket between the tibia and talus (Fig. 53.US14). Ultrasound can also be used to identify the location of the dorsalis pedis artery before aspiration.

Figure 53.US5 Ultrasound image of a joint effusion within the suprapatellar recess of the knee joint. The recess is distended with anechoic (black) fluid and the femur can be seen as the hyperechoic (white) line at the bottom of the image.

Figure 53.US6 Placement of the ultrasound transducer over the anterior aspect of the shoulder to evaluate the biceps tendon. To obtain the best view, the patient’s arm should be flexed at the elbow, supinated, and slightly abducted.

Figure 53.US7 Ultrasound image of a normal-appearing biceps tendon. The tendon will appear as a hyperechoic (white) bundle within the groove of the humerus as indicated by the arrow.

Figure 53.US8 The biceps tendon is surrounded by anechoic (black) fluid. (Courtesy Verena Valley, MD.)

Figure 53.US9 Placement of the ultrasound transducer over the posterior aspect of the shoulder to evaluate the glenohumeral joint. This joint line can typically be palpated to approximate the best initial position.
Elbow

The elbow is easily evaluated from the posterior approach. With this approach, place the transducer over the olecranon fossa with the elbow flexed and the lower part of the arm supported (Fig. 53.US15). In a normal elbow the olecranon fossa can be seen as a slight "divot" between the olecranon of the ulna and the humerus (Fig. 53.US16). When an effusion is present, dark gray or black fluid can be seen to distend this space (Fig. 53.US17).

Figures 53.US10-53.US14

Hip

The hip joint is unique in that physical examination may suggest the presence of an effusion, but direct confirmation is difficult with traditional examination techniques. Ultrasound will easily confirm the presence of an effusion. To evaluate the hip joint, select a low-frequency transducer (3 to 5 mHz) initially because of the depth of the joint. In very thin patients, the distance from the skin to the joint may be small enough to allow the use of a high-frequency transducer. Align the transducer in a slightly oblique axis (mimicking the orientation of the femoral neck) along the inguinal area. It may be helpful to aim toward the umbilicus. Look for the femoral neck and head. They will appear as brightly echogenic outlines in the expected location.
shape of the bones (Fig. 53.US18). Once the femur has been identified, look for the joint capsule. It will appear as an arcing, hyperechoic line superficial to the bones (Fig. 53.US19). In a normal hip there will be very little tissue between these two structures. There may be a small amount of anechoic or hypoechoic fluid present in this space in a normal hip, and correlation with the unaffected side will aid in evaluation. In the presence of a significant effusion, a large anechoic or hypoechoic fluid collection will be seen between the femoral neck and the capsule (Fig. 53.US20).

**Figure 53.US15** Placement of the transducer in the transverse plane over the area of the olecranon fossa. For best resolution, a high-frequency transducer should be used.

**Figure 53.US16** Ultrasound image of a normal elbow joint. The olecranon fossa can be identified as the echogenic (white) crescent at the bottom of the image (arrow). The area just above the fossa should be evaluated for the presence of fluid indicative of an effusion.

**Figure 53.US17** Elbow joint distended by anechoic fluid (arrow) superficial to the olecranon fossa. (From Allan PL, Baxter GM, Weston MJ, editors: Clinical ultrasound, ed 3, Philadelphia, 2011, Elsevier.)

**Figure 53.US18** Normal ultrasound of the hip. The femoral head can be seen as the hyperechoic (white) line highlighted by the arrow. The area immediately superficial to the femur is devoid of any significant fluid collection.

**Figure 53.US19** Ultrasound image of the hip with demonstration of the joint capsule. The joint capsule can be seen as the hyperechoic (white) arcing structure marked by the arrow.

**Figure 53.US20** Ultrasound image of a hip effusion. As in a normal hip, the femur can be seen at the bottom of the image. Immediately superficial to the femur, an anechoic (black) fluid collection is highlighted by the arrow.
First, select a puncture site and an approach to the joint. Avoid tendons, major vessels, and major nerve branches. In most instances the approach is via the extensor surfaces of joints because most major vessels and nerves are found beneath the flexor surfaces. In addition, the synovial pouch is usually more superficial on the extensor side of a joint. Ultrasound may be particularly helpful in locating small effusions.

Use aseptic technique, including the use of sterile gloves and instruments, to avoid infection. Do not attempt arthrocentesis if there is a definite or suspected infection overlying the joint. Allow antiseptic preparation solution to dry for several minutes because the bactericidal effects of iodine are both concentration and time dependent. Remove the iodine solution with an alcohol sponge to prevent transference of iodine into the joint space, which can lead to an inflammatory process. Although the utility of draping is unproved and it may obscure the site, a sterile perforated drape may be placed over the joint.

With appropriate local anesthesia, arthrocentesis should be a relatively painless procedure; without anesthesia, it may be quite painful and distressing to the patient. The synovial membrane itself has pain fibers associated with blood vessels, and the articular capsule and periosteum are richly supplied with nerve fibers so both are very sensitive. The articular cartilage has no intrinsic pain fibers. It is important to have the patient relax during the procedure. Tense muscles narrow the joint space and make the procedure more difficult, often necessitating repeated attempts or resulting in inadequate drainage. Distraction of the joint may enhance the target area, especially in areas such as the wrist and finger joints. Traction not only increases the chance of entering the joint but also lessens the chance of scoring the articular cartilage with the needle.

To best accomplish anesthesia, infiltrate the skin down to the area of the joint capsule along the entire route of needle penetration. Use a local anesthetic agent such as 1% or 2% lidocaine (Xylocaine [AstraZeneca, Cambridge, United Kingdom]) via a 25- to 27-gauge needle. For extremely painful joints, a regional nerve block is appropriate.

Use the landmarks described in “Specific Arthrocentesis Techniques” later in this chapter. Take care not to bounce the needle off bony structures as a means of finding the joint space because this may cause unnecessary pain. However, in contrast to earlier beliefs, striking bone with the arthrocentesis needle is unlikely to damage articular cartilage. Attach an 18- to 22-gauge needle or intravenous catheter and needle set of suitable length to an appropriately sized syringe. Insert the needle at the desired anatomic point through the skin and subcutaneous tissue into the joint space. Use the largest needle that is practical to avoid obstructing the lumen with debris or clot. Large joints, such as the knee, can accommodate large effusions, so it is suggested that one use a 30- to 60-mL syringe because it may be difficult to change a syringe when the needle is within the joint cavity (Fig. 53.5). A three-way stopcock placed between the needle and the syringe is an option for draining large effusions (Fig. 53.6). If the syringe must be changed during the procedure, grasp the hub of the needle with a hemostat and hold it tightly while the syringe is removed. The authors prefer to use only a rigid needle and not a flexible catheter to perform arthrocentesis; however, a sturdy catheter is used by some clinicians. If an intravenous catheter and needle set is used, remove the needle while leaving the outer atraumatic plastic catheter in the joint space. Then attach the syringe to the catheter for aspiration. Manipulation of the joint or catheter can now occur with little threat of tissue injury.

Aspiration of synovial fluid and easy injection and return of fluid indicate intraarticular placement of the needle tip. As a general rule, try to remove as much fluid or blood as possible. If the fluid stops flowing, it indicates that the joint has been drained completely, the tip of the needle has become dislodged, or debris or clot is obstructing the needle. In this case, slightly advance or retract the tip of the needle, rotate the bevel, or lessen the force of aspiration. Never reintroduce a needle through a plastic catheter that has been left in the joint. Occasionally, reinjecting a small amount of fluid into the joint space confirms placement of the needle and may clear the needle. If fluid flows freely back into the joint and is easily aspirated, one has probably removed all the fluid. If resistance is met, the needle has probably been jarred from the joint space and is lodged in soft tissue. In some instances, minor changes in position produced by flexion or extension of the joint may allow the fluid to flow more freely. Scraping or shearing the articular cartilage with the needle should be avoided. One should enter the joint in a straight line and avoid unnecessary side-to-side motion of the needle.

Send synovial fluid for studies as indicated by the clinical situation. Studies usually obtained include cell count with differential, crystal analysis, Gram staining, and bacterial culture and sensitivity analysis. Synovial protein, glucose, and lactate dehydrogenase determinations have been shown to be unreliable in distinguishing noninflammatory from inflammatory and infectious causes and are no longer recommended. Less frequently obtained studies include rheumatoid factor analysis, lupus erythematosus cell preparation, viscosity analysis, mucin clot, fibrin clot, fungal and acid-fast stains, Lyme titer, fungal and tuberculous culture, and synovial fluid complement analysis. If the arthrocentesis is performed for the relief of hemarthrosis, the fluid need not be sent for analysis. One should be selective in ordering tests. There is no need to order a large battery of tests routinely on all fluids. If the volume of fluid collected is low, Gram stain, culture, and examination of the “wet preparation” under regular and polarizing microscopy have the highest priority. Prompt examination of specimens should be performed

**ULTRASOUND BOX 53.1: Arthrocentesis—cont’d**

**Aspiration**

Once the joint in question has been evaluated, the optimal site for aspiration can be planned. In contrast to the traditional, blind aspiration technique, the use of ultrasound may suggest an alternative approach. Ultrasound will enable the clinician to map the best approach to the effusion. Once this approach has been clarified, one of two techniques can be applied. The site can be marked and the aspiration can then proceed blindly under sterile conditions. In other cases it may be preferable to perform the tap under direct ultrasound guidance. In these circumstances the needle is inserted either from the transverse or from the long-axis approach and guided directly into the joint space.
to avoid misdiagnosing borderline inflammatory fluids, missing crystals that dissolve with time, or overinterpreting the findings because of new artifactual crystals that appear over a prolonged time.24

Complications
Significant complications are rare with arthrocentesis but include the following:
1. Infection. Skin bacteria may be introduced into the joint space during needle puncture. Nevertheless, infection occurs rarely because the bacteria are either quickly cleared or not viable.2 One can further limit this complication by maintaining rigorous sterile technique and avoiding insertion of the needle through obviously (or possibly) infected skin or subcutaneous tissue. Various studies report the incidence of infection after routine arthrocentesis to be in the range of 1 in 10,000.34 However, in immunocompromised patients, particularly those with rheumatoid arthritis, the incidence is higher (1 in 2000 to 10,000 aspirations).35,36 Joint aspiration in the presence of bacteremia was discussed previously. Acute pain, redness, and swelling 12 to 24 hours after steroid
injection can mimic infection but is most likely an inflammatory reaction (steroid flare) to the steroid preparation (often methylprednisolone).

2. Bleeding. Bleeding with subsequent hemarthrosis is rarely a complication, except in patients with a bleeding diathesis. In those with a bleeding diathesis such as hemophilia, arthrocentesis should be delayed until clotting competence has been enhanced by infusing specific clotting factors. In general, spontaneous bleeding into a hemophilic patient’s joint is an indication for replacement of clotting factors. Occasionally, a small quantity of blood may be aspirated along with synovial fluid. This happens most often when the joint is nearly emptied. A small amount of blood-tinged fluid is generally the result of nicking a small synovial blood vessel and is usually inconsequential.

3. Arthrocentesis and joint injections in patients receiving chronic warfarin therapy, with a therapeutic INR, were shown to be safe by Ahmed and Gertner,3 without an increased risk of bleeding complications. In this study of 456 procedures in patients on chronic warfarin therapy, there was no statistically significant difference in the overall complication rate between patients with an INR of 2.0 or greater and patients with an INR less than 2.0. Of note, 103 of 456 procedures (22.5%) were safely performed in patients with an INR greater than 3. Aspirating fluid from small joints is not uncommon and most often the result of vasovagal influences. To help prevent this, perform the procedure with the patient laying down whenever possible.

4. Allergic reaction. Hypersensitivity to the local anesthetic can usually be prevented by thorough history taking. Facial and torso flushing associated with corticosteroid injection may represent an idiosyncratic reaction to preservatives in the steroid preparation.7 Fainting during the procedure is not uncommon and most often the result of vasovagal influences. To help prevent this, perform the procedure with the patient lying down whenever possible.

5. Corticosteroid-induced complications. See earlier section on Intraarticular Corticosteroid Injections.

**SPECIFIC ARTHROCENTESIS TECHNIQUES**

Arthrocentesis of the hip is generally performed by an orthopedic surgeon under fluoroscopic, ultrasound, or magnetic resonance imaging, or computed tomography guidance and is not discussed here. If available, fluoroscopy or ultrasound can also be used to guide aspiration of other joints, but these imaging adjuncts are not generally required. For small joints, application of traction is often very helpful in obtaining fluid. While applying continuous suction to the aspirating syringe, walk the needle over palpated bone until the joint is entered or fluid is obtained. However, it may be quite difficult to obtain fluid from small joints in the hand and foot, so the clinician must often treat empirically. If only one drop of fluid is obtained from small joints, it is best to send it for culture.

**First Carpometacarpal Joint (Fig. 53.7)**

**Landmarks.** The radial aspect of the proximal end of the first metacarpal is the arthrocentesis landmark for this joint. Locate the abductor pollicis longus (APL) tendon by active extension of the tendon.

**Position.** Oppose the thumb against the little finger so that the proximal end of the first metacarpal is palpable. Apply traction to the thumb to widen the joint space between the first metacarpal and the greater multangular (trapezium) bone.

**Needle Insertion.** Insert a 22- to 23-gauge needle at a point proximal to the prominence at the base of the first metacarpal on the palmar side of the APL tendon.

**Comments.** Degenerative joint disease commonly affects this joint. Arthrocentesis is moderately difficult. The anatomic “snuffbox” (located more proximally and on the dorsal side of the APL tendon) should be avoided because it contains the radial artery and superficial radial nerve. A more dorsal approach may also be used.

**Interphalangeal and Metacarpophalangeal Joints (Fig. 53.8)**

**Landmarks.** The landmarks are located on the dorsal surface. For the metacarpophalangeal joints, palpate for the prominence at the proximal end of the proximal phalanx. For the interphalangeal joints, palpate for the prominence at the proximal
end of the middle or distal phalanx. The extensor tendon runs down the midline.

**Position.** Flex the fingers to approximately 15 to 20 degrees and apply traction.

**Needle Insertion.** Insert a 22- to 25-gauge needle into the joint space dorsally, just medial or lateral to the central slip of the extensor tendon.

**Comments.** Synovitis causes these joints to bulge dorsally. Normally, it is unusual to obtain fluid in the absence of a significant pathologic condition.

**Radiocarpal Joint (Wrist) (Fig. 53.9; Videos 53.1 and 53.2)**

**Landmarks.** The dorsal radial tubercle (Lister’s tubercle) is an elevation found in the center of the dorsal aspect of the distal end of the radius. The extensor pollicis longus tendon runs in a groove on the radial side of the tubercle. The tendon can be palpated by active extension of the wrist and thumb.

**Position.** Position the wrist in approximately 20 to 30 degrees of flexion with accompanying ulnar deviation. Apply traction to the hand.

**Needle Insertion.** Insert a 22-gauge needle dorsally, just distal to the dorsal tubercle on the ulnar side of the extensor pollicis longus tendon. Avoid the anatomic snuffbox, located more radially, to prevent injury to the radial artery or superficial radial nerve.

**Radiohumeral Joint (Elbow) (Fig. 53.10; Videos 53.3 and 53.4)**

**Landmarks.** The lateral epicondyle of the humerus and the head of the radius are the arthrocentesis landmarks for the radiohumeral joint. With the elbow extended, palpate the depression between the radial head and the lateral epicondyle of the humerus.

**Position.** With the palpating finger still touching the radial head, flex the elbow to 90 degrees. Pronate the forearm, and place the palm flat on a table.

**Needle Insertion.** Insert a 20-gauge needle from the lateral aspect, just distal to the lateral epicondyle and directed medially.

**Comments.** Elevation of the anterior fat pad or the presence of a posterior fat pad on a lateral soft tissue elbow radiograph signifies blood, pus, or fluid in the elbow joint (see Fig. 53.10B). Effusions in the elbow joint may bulge and be readily palpated (see Fig. 53.10C). Frequently, the effusion appears inferior to the lateral epicondyle. The bulge can then be aspirated from a posterior approach on the lateral side (see Fig. 53.10D). A medial approach is not recommended because the ulnar nerve and superior ulnar collateral artery may be damaged. Gout and septic arthritis commonly affect this joint. The most common cause of elbow hemarthrosis after trauma with no obvious fracture is a nondisplaced radial head fracture. A small hemarthrosis need not be aspirated, but removal of blood from a tense elbow joint will significantly hasten recovery and facilitate range of motion in patients with a radial head fracture.
Knee Joint, Anteromedial Approach (Fig. 53.12; Videos 53.6 and 53.7)

**Landmarks.** The medial surface of the patella at the middle or superior portion of the patella is the landmark for the knee joint.

**Position.** It is usually recommended that the knee be extended as far as possible. Alternatively, some practitioners prefer to flex the knee 15 to 20 degrees by placing a towel under the popliteal region to open the joint space up. Relaxation of the quadriceps muscle greatly facilitates needle placement. Keep the foot perpendicular to the floor.

**Needle Insertion.** Insert an 18-gauge needle or catheter and needle set at the midpoint or superior portion of the patella.

---

Glenohumeral Joint (Shoulder), Anterior Approach (Fig. 53.11; Video 53.5)

**Landmarks.** Anteriorly palpate the coracoid process medially and the proximal end of the humerus laterally.

**Position.** The patient should sit upright with the arm at the side and hand in the lap.

**Needle Insertion.** Insert a 20-gauge needle at a point inferior and lateral to the coracoid process and direct it posteriorly toward the glenoid rim.

**Comments.** Arthrocentesis of this joint is moderately difficult. Other approaches have been suggested but are less well accepted.
Arthrocentesis

approximately 1 cm medial to the anteromedial patellar edge. Direct the needle between the posterior surface of the patella and the intercondylar femoral notch. The patella may be grasped with the hand and elevated to aid entry of the needle into the joint. Keeping the needle/syringe parallel to the bed limits internal injury.

Comments. If the patient is tense, contraction of the quadriceps will greatly hinder entering the joint. However, the knee is probably the easiest joint to enter, and removal of a tense hemarthrosis will relieve pain and facilitate examination for ligamentous injury. If fluid stops flowing, squeeze the soft tissue area of the suprapatellar region to “milk” the suprapatellar pouch of fluid. Alternatively, wrap the patient's thigh with a 6-inch elastic bandage from the groin to the suprapatellar area before beginning the procedure. The knee can easily accommodate 50 to 70 mL of fluid, so use a large syringe. Hold or secure the hub of the needle with a hemostat to remove the syringe without changing the position of the intraarticular needle. Alternatively, a stopcock on the needle will allow complete removal of fluid without changing the position of the needle (see Fig. 53.12B). The knee is a common site for septic arthritis (especially gonococcal) and various inflammatory or degenerative diseases. An anterolateral approach can be accomplished in a similar manner.

Tibiotalar Joint (Ankle) (Fig. 53.13; Video 53.8)

Landmarks. The medial malleolar sulcus is bordered medially by the medial malleolus and laterally by the anterior tibial tendon. The tendon can easily be identified with active dorsiflexion of the foot.

Position. With the patient lying supine, plantar-flex the foot.

Needle Insertion. Insert a 20- to 22-gauge needle at a point just medial to the anterior tibial tendon and directed into the hollow at the anterior edge of the medial malleolus. The needle must be inserted 2 to 3 cm to penetrate the joint space.

Comments. If the joint bulges medially, one may use an approach that is more medial than anterior and enter at a point just anterior to the medial malleolus. The needle may have to be advanced 2 to 4 cm with this approach.

Metatarsophalangeal and Interphalangeal Joints (Fig. 53.14; Videos 53.9 and 53.10)

Landmarks. For the first digit, landmarks are the distal metatarsal head and the proximal base of the first phalanx. For the other toes, the landmarks are the prominences at the proximal interphalangeal and distal interphalangeal joints. The extensor tendon of the great toe can be located by active extension of the toe.

Position. With the patient supine, flex the toes 15 to 20 degrees. Then apply traction.

NEEDLE INSERTION. Insert a 22-gauge needle on the dorsal surface at a point just medial or lateral to the central slip of the extensor tendon.
characteristically results in low-viscosity synovial fluid. The string sign is a simple test for assessing viscosity. The practitioner measures the length of the “string” formed by a falling drop of synovial fluid extruded from a syringe or stretched between the thumb and the index finger of a gloved hand. Normal joint fluid produces a string 5 to 10 cm long (Fig. 53.15). If viscosity is reduced, as with inflammatory conditions, synovial fluid forms a shorter string or falls in drops.

Mucin Clot Test

The mucin clot test also corresponds to viscosity and inflammation. The greater the inflammatory response, the poorer the mucin clot and the lower the viscosity. This test may be useful to define the degree of polymerization of hyaluronate.
Glucose and Protein
Current literature suggests that synovial protein and glucose are highly inaccurate markers of inflammation. In one study of 100 consecutive patients undergoing diagnostic arthrocentesis, the sensitivity of synovial protein and glucose was found to be 0.52 and 0.20, respectively. The authors of this study recommended that ordering chemistry studies on synovial fluid should be discouraged because such studies are likely to provide misleading or redundant information.

Serology
Though available, most serologic tests are not likely to be useful in the emergency setting. Polymerase chain reaction (PCR) is an effective means of identifying septic arthritis, even in the setting of a negative fluid culture or when antibiotics have been administered previously. PCR can also help isolate slowly growing microorganisms. Gas-liquid chromatography, a rapid and sensitive method for detection of short-chain fatty acids, may complement the currently available methods used to diagnose septic arthritis.

Counterimmunoelectrophoresis and latex agglutination are also useful and available in some centers on an emergency basis. Other immunologic markers such as complement, rheumatoid factor, and antinuclear antibodies have little diagnostic value in the acute setting but may be useful to the clinician providing follow-up care when compared with serum levels.

Fluid Processing
Proper collection of joint fluid is essential for examination and testing. Tests for viscosity, serology, and chemistries are done on fluid collected in a red-topped (clot) tube, whereas cytology samples are collected in tubes with an anticoagulant (purple top). One should always transfer the fluid for crystal examination into a tube with liquid heparin (green top) because undissolved heparin crystals from powdered anticoagulant tubes can be seen on microscopy. Early transfer of synovial fluid to this green-topped tube is essential to prevent clotting. Culture requirements for transport and processing should be accessed before the procedure to ensure appropriate processing or plating of specimens.

Polarizing Microscope
No synovial fluid analysis is complete until the fluid has been examined for crystals under a polarizing light microscope. The polarizing microscope used for crystal identification differs from the ordinary light microscope in that it contains two identical polarizing prisms or filters. One filter, called the polarizer, is positioned below the condenser. The other filter is called the analyzer and is inserted at some point above the objective. Examination for crystals is performed by most hospital laboratories.

Cell Count
A leukocytosis consisting predominantly of neutrophils is usually seen with inflammatory arthritides; a WBC count greater than 50,000/mm³ (i.e., >50,000/µL) is highly suggestive of a septic joint. Shmerling and colleagues found a WBC count of greater than 2000/mm³ to be 84% sensitive and 84% specific for all inflammatory arthritides. Of their septic arthritis patients, 37% had a synovial WBC count lower than 50,000/mm³. However, 89% of their patients with a synovial WBC count greater than 50,000/mm³ had a septic joint.

Mucin clots are produced by mixing one part joint fluid with four parts 2% acetic acid. A good clot indicates a high degree of polymerization and correlates with normal high viscosity. In inflammatory synovial fluid, such as that seen with osteoarthritis- and rheumatoid arthritis–related effusions, the mucin clot is poor. This test is rarely performed.

Cell Count
A leukocytosis consisting predominantly of neutrophils is usually seen with inflammatory arthritides; a WBC count greater than 50,000/mm³ (i.e., >50,000/µL) is highly suggestive of a septic joint. Shmerling and colleagues found a WBC count of greater than 2000/mm³ to be 84% sensitive and 84% specific for all inflammatory arthritides. Of their septic arthritis patients, 37% had a synovial WBC count lower than 50,000/mm³. However, 89% of their patients with a synovial WBC count greater than 50,000/mm³ had a septic joint.

Polarizing Microscope
No synovial fluid analysis is complete until the fluid has been examined for crystals under a polarizing light microscope. The polarizing microscope used for crystal identification differs from the ordinary light microscope in that it contains two identical polarizing prisms or filters. One filter, called the polarizer, is positioned below the condenser. The other filter is called the analyzer and is inserted at some point above the objective. Examination for crystals is performed by most hospital laboratories.

Polarization Physics
The polarizer allows passage of light in only one specific orientation. The analyzer acts as a crossed filter by removing all light in the light path unless the material being examined rotates the beam from the polarizer into the plane of the analyzer. The compensator functions by imparting color of a certain wavelength (red at approximately 550 nm). Birefringent
When examining crystals under polarized microscopy, the technician orients crystals on a stage according to two axes, referred to as x and z. If the long axis of the crystals is blue when parallel to the x-axis and yellow when perpendicular to it, it is calcium pyrophosphate and termed positively birefringent. If the long axis of the crystal is yellow when parallel to the z-axis and blue when perpendicular to it, it is monosodium urate and termed negatively birefringent. Urate crystals are 2 to 10 µm and usually needle shaped (Fig. 53.16). Calcium pyrophosphate crystals range from 10 µm down to tiny crystals that have to be examined with the oil objective; they appear as rods, rhomboids, plates, or needle-like forms and are weakly birefringent (Fig. 53.17). Cholesterol crystals are sometimes seen and are large, very bright square or rectangular plates with broken corners.40,41

Items found in synovial fluid that can be confused with sodium urate or calcium pyrophosphate crystals include collagen fibrils, cartilage fragments, cholesterol crystals, metallic fragments from prosthetic arthroplasty, and corticosteroid esters.40 One may also identify fat globules (Fig. 53.18). Note that rare cases of uric acid spherulites in gouty synovia have been reported.40 The spherulites are birefringent and do not take up fat stains.

Table 53.2 summarizes synovial fluid features for the joint diseases commonly encountered and studies commonly performed in the ED.

**TABLE 53.2 Characteristics of Synovial Fluid**

<table>
<thead>
<tr>
<th>APPEARANCE</th>
<th>VISCOSITY</th>
<th>CELLS PER mm³</th>
<th>PMN (%)</th>
<th>CRYSTALS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>Transparent</td>
<td>High</td>
<td>&lt;180</td>
<td>&lt;10 Negative</td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td>Transparent</td>
<td>High</td>
<td>200–2000</td>
<td>&lt;10 Occasional calcium pyrophosphate and hydroxyapatite crystals</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>Translucent</td>
<td>Low</td>
<td>2000–50,000</td>
<td>Variable Negative</td>
</tr>
<tr>
<td>Psoriatic arthritis</td>
<td>Translucent</td>
<td>Low</td>
<td>2000–50,000</td>
<td>Variable Negative</td>
</tr>
<tr>
<td>Reactive arthritis</td>
<td>Translucent</td>
<td>Low</td>
<td>2000–50,000</td>
<td>Variable Negative</td>
</tr>
<tr>
<td>Spondyloarthropathy</td>
<td>Translucent</td>
<td>Low</td>
<td>2000–50,000</td>
<td>Variable Negative</td>
</tr>
<tr>
<td>Gout</td>
<td>Translucent to cloudy</td>
<td>Low</td>
<td>200 to &gt;50,000</td>
<td>&gt;90 Needle-shaped, positive birefringent monosodium urate monohydrate crystals</td>
</tr>
<tr>
<td>Pseudogout</td>
<td>Translucent to cloudy</td>
<td>Low</td>
<td>200–50,000</td>
<td>&gt;90 Rhomboid, negative birefringent calcium pyrophosphate crystals</td>
</tr>
<tr>
<td>Bacterial arthritis</td>
<td>Translucent to cloudy</td>
<td>Low</td>
<td>2000 to &gt;50,000</td>
<td>&gt;90 Negative</td>
</tr>
<tr>
<td>PVNS</td>
<td>Hemorrhagic or brown</td>
<td>Low</td>
<td></td>
<td>Negative</td>
</tr>
<tr>
<td>Hemarthrosis</td>
<td>Hemorrhagic</td>
<td>Low</td>
<td></td>
<td>Negative</td>
</tr>
</tbody>
</table>

PMN, Polymorphic nuclear cell; PVNS, pigmented villonodular synovitis.
JOINT ARTHROGRAPHY

Background

Wounds near joints raise concern regarding joint penetration. Treatment and interventions may be altered significantly if a joint space has been traumatically violated. Plain radiographs may demonstrate air in the joint, which clinches the diagnosis, but in questionable cases the diagnostic approach includes injection arthrograms. Historically, these were performed by injecting methylene blue into the joint in question and assessing leakage from the joint. However, the dye can interfere with arthroscopic evaluation and produce an inflammatory reaction, so saline arthrography (SA) is now preferred. SA, or a “saline load test,” was first described in 1975 but became more popular in the 1990s.\(^{42,43}\)

Indications and Contraindications

SA should be performed in patients with penetrating injuries near a joint in which violation of the joint itself is unclear (Fig. 53.19). Smaller joints such as those in the hand are inspected visually. However, for larger joints, SA is the preferred test.

Figure 53.18 A, Blood aspirated from a traumatized joint is placed in an emesis basin and put under a bright light to search for lipohemarthrosis, which is clinical evidence of an intraarticular fracture. Note the characteristic greasy sheen of floating fat. Do not throw away this blood before assessing for lipohemarthrosis. B, A fibular head fracture (arrow) was initially thought to be responsible for a large knee hemarthrosis, but anatomically it is extraarticular. A tibial plateau fracture is not appreciated on this radiograph. Such injuries are often occult. C, On this view one might appreciate the radiolucent lines suggestive of a lateral tibial plateau fracture (arrows), but such subtle findings are easily missed, as was the case with this patient. Therapeutic arthrocentesis was performed to alleviate pain, and an obvious lipohemarthrosis was noted. D, Magnetic resonance imaging demonstrates an obvious lateral tibial plateau fracture (arrows), which was searched for only after lipohemarthrosis was noted. Computed tomography could also be used.

Figure 53.19 This periarticular laceration raises the question of knee joint penetration. A plain radiograph may demonstrate air in the joint space, but a saline arthrogram may also be used. Methylene blue alone is not generally required and it can cause an inflammatory reaction and obscure arthroscopy. A small amount of methylene blue may be added to color the saline.
cause the patient discomfort. The sensitivity of the test in detec-
ting small traumatic joint injuries is proportional to the
volume injected. Specifically, for knee injuries, injecting 50 mL
of saline was 46% sensitive and injecting 100 mL, 75% sensitive;
to achieve 95% sensitivity required an average of 194 mL of
saline. The following is the recommended volume of injection
per joint:
- Knee: 100 to 200 mL
- Elbow: 20 to 30 mL
- Ankle: 20 to 30 mL
- Wrist: 5 mL
- Shoulder: 40 to 60 mL

Once the injection is complete, do not remove the needle but
close the stopcock to avoid backflow. Examine the joint for
evidence of leakage of fluid from the wound. This is performed
in a static position but, if negative, also with some gentle
passive movement of the joint. Visible leakage of fluid into
the laceration confirms the diagnosis of joint space violation.
A negative test is defined as absence of evidence of leakage
after an appropriate amount of saline has been injected. A slow
loss of fluid may indicate a small insult to the joint, and saline
can be left in the joint for a few minutes to observe for this.
After completion, the fluid should be evacuated for patient
comfort. This is generally performed by leaving the original
needle in place with a closed stopcock attached, which is then
used to aspirate the saline in the joint.

**Complications**

Complications associated with performing SA are essentially
the same as those for performing arthrocentesis. In addition,
some temporary patient discomfort because of joint distention
should be assumed.

**Conclusion**

Small traumatic joint penetration can be difficult to diagnose
clinically. SA can help confirm the diagnosis. To be a sensitive
test, it must be performed with an adequate amount of saline
infusion to truly "load" the joint. If the test is positive, ortho-
pedic consultation is indicated.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
Open fractures, dislocations, and exposed joints are true orthopedic emergencies that must be managed aggressively to prevent morbidity and mortality. Even when managed appropriately, these injuries may be further complicated by a compartment syndrome, a condition of increased pressure within a limited space that results in compromised tissue perfusion leading to tissue ischemia, and ultimately dysfunction of the neural and muscular structures contained within that space. Compartment syndrome is most often associated with significant trauma, particularly long bone fractures (e.g., tibia, radius, ulna). However, it may also occur following less severe trauma that does not cause fractures, such as crush injuries, severe thermal burns, penetrating trauma, injury to vascular structures in the extremities and, in some cases, even minor injuries. Patients on anticoagulants, those with a bleeding diathesis, and those who continue to use an injured limb are at increased risk. Nontraumatic causes of compartment syndrome occur less often, and include ischemia-reperfusion injury, excessive muscular exertion, thrombosis, bleeding disorders and coagulopathies, vascular disease, nephrotic syndrome, certain animal envenomations and bites, extravasation of intravenous (IV) fluids, injection of recreational drugs, and prolonged external pressure (e.g., from a tight circumferential bandage or cast, or from lying on an extremity for a prolonged period secondary to drug or alcohol intoxication).

Numerous drugs and toxins have been reported to cause rhabdomyolysis, possibly because of a direct effect or secondary to agitation and exertion, with the theoretical potential for the development of compartment syndrome (Fig. 54.1). This list is exhaustive but includes heroin, various hydrocarbons, cocaine, amphetamines, antidepressants, antipsychotics, salicylates, propoxyphene, nonsteroidal antiinflammatory drugs, succinylcholine, human immunodeficiency virus (HIV) medications, antimetabolite and cancer drugs, antimalarials, diphenhydramine, baclofen, ecstasy, ethanol, anticoagulants and thrombolytics, strychnine, statins, and phenothiazines.

Causes of compartment syndrome are categorized into those that decrease compartment volume capacity, those that increase the contents of a compartment, and those that create externally applied pressure (Box 54.1).

Subtleties in the early signs and symptoms of compartment syndrome or other clinical priorities render some cases simply impossible to recognize and treat early enough to thwart the ultimate disability. In fact, approximately 10% of cases of acute compartment syndrome secondary to fractures and 20% of non–fracture-associated cases will have muscle necrosis requiring débridement at the time of surgery. These statistics indicate that even with diligent clinical care, it is often difficult to identify all cases before injury to muscles occurs. This is

## Compartment Syndrome Evaluation

### Indications
Suspected compartment syndrome with clinical findings that are equivocal or difficult to interpret:
- Unresponsive patients
- Uncooperative patients
- Children
- Patients with multiple or distracting injuries
- Patients with peripheral nerve deficits attributable to other causes (fracture-associated nerve injury, diabetic peripheral neuropathy)

### Contraindications
**Absolute**
- None
**Relative**
- Coagulation disorders
- Overlying infection, cellulitis, or burns

### Complications
- Pain from needle insertion, fluid injection, or bleeding
- Inaccurate readings due to poor technique, improper needle position, injected fluid or anesthetic, or external compression
- Injury to underlying tissue, nerves, or blood vessels
- Local or systemic infection

### Equipment
- Stryker pressure monitor
- Side-port 18-gauge needle
- Diaphragm chamber
- 3-mL syringe prefilled with saline

Equipment shown is for the Stryker method. The side-port needle, prefilled syringe, and diaphragm chamber are packaged together in the Stryker 295-2 Quick Pressure Monitor Set (Stryker, Kalamazoo, MI).

*Review Box 54.1* Compartment syndrome evaluation: indications, contraindications, complications, and equipment.
The diagnosis of compartment syndrome is not always straightforward. A, This man was initially seen while in a coma from a heroin overdose and had been lying on his arm for a number of hours. He was hypotensive, in renal failure, comatose, and on a ventilator. The entire arm was swollen and rhabdomyolysis was correctly suspected. B, Clear urine, a strongly positive dipstick for blood (arrow), and no red blood cells by microscopy equate to myoglobinuria. Because of the coma, he was unable to voice any complaint of pain. C, When he awakened 20 hours later, the pain was severe, and compartment pressures indicated the need for fasciotomy. Heroin can cause rhabdomyolysis, and hypotension/reperfusion and certainly prolonged pressure on the muscles may have exacerbated the condition. D, The classic wringer washer injury predisposes to compartment syndrome, but industrial rollers are now usually the culprit.
Over the past 40 years or so, various needles and equipment have been developed to measure compartment pressure (Fig. 54.2).5–11 The \textit{wick catheter}, originally developed to measure subcutaneous and brain tissue pressure, was modified during the mid-1970s to provide continuous compartment pressure measurements. This catheter is rarely used today because of the fear of catheter breakdown leading to errors in measurement and retained foreign bodies. The \textit{slit catheter} was introduced in 1980.10,11 As its name implies, this catheter has slits at its distal end that prevent clogging. The proximal end of the catheter is connected to a transducer and infusion system, which permits continuous pressure monitoring. Both the wick and slit catheters have been shown to offer similar accuracy and reproducibility as long as patency of the catheter is ensured.11–13

The Stryker Intracompartmental Compartment Pressure Monitoring System (Kalamazoo, MI) has become the most commonly used commercially available device to measure compartment pressure in the emergency department (ED) (see Review Box 54.1). This device uses a fluid-filled pressure measurement catheter, a pressure monitor, and a fluid infusion mechanism that maintains catheter patency and ensures accurate measurement. In contrast to earlier devices in which relatively large volumes of fluid were injected into the compartment to measure pressure, the Stryker system uses a minimal amount of saline (< 0.3 mL). This helps ensure accurate measurements particularly true in uncooperative, unconscious, or critically injured patients who are unable to report symptoms. Unfortunately, the vagaries of the clinical scenario that result in failure to recognize the early signs and symptoms of compartment syndrome may have severe and irreversible limb- or life-threatening consequences.

Although compartment syndrome is essentially a clinical diagnosis, objective measurement of compartment tissue pressure may assist in confirming the diagnosis and determining if operative management is required. This chapter discusses the indications, complications, and interpretation of compartment pressure monitoring, as well as the equipment and techniques required to measure and monitor compartment pressure.

**BACKGROUND**

Though recognized as a clinical syndrome in the mid-19th century, the pathophysiology of ischemia in extremities was not fully described until more than a century later. Postischemic myoneural dysfunction and its associated contractures were first described in the 1870s by the German surgeon Richard von Volkmann, who recognized the effects of increased pressure causing vascular compromise of the limb.4

**BOX 54.1 Causes of Compartment Syndrome**

<table>
<thead>
<tr>
<th>DECREASED COMPARTMENTAL VOLUME</th>
<th>Intraarterial drug injection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Closure of fascial defects</td>
<td>Cold</td>
</tr>
<tr>
<td>Application of excessive traction to fractured limbs</td>
<td>Orthopedic surgery</td>
</tr>
<tr>
<td>Compressive devices (casts, splints, circumferential dressings)</td>
<td>Tibial osteotomy</td>
</tr>
<tr>
<td></td>
<td>Hauser’s procedure</td>
</tr>
<tr>
<td></td>
<td>Reduction and internal fixation of fractures</td>
</tr>
<tr>
<td></td>
<td>Snakebite</td>
</tr>
</tbody>
</table>

**BACKGROUND**

Over the past 40 years or so, various needles and equipment have been developed to measure compartment pressure (Fig. 54.2).5–11 The \textit{wick catheter}, originally developed to measure subcutaneous and brain tissue pressure, was modified during the mid-1970s to provide continuous compartment pressure measurements. This catheter is rarely used today because of the fear of catheter breakdown leading to errors in measurement and retained foreign bodies. The \textit{slit catheter} was introduced in 1980.10,11 As its name implies, this catheter has slits at its distal end that prevent clogging. The proximal end of the catheter is connected to a transducer and infusion system, which permits continuous pressure monitoring. Both the wick and slit catheters have been shown to offer similar accuracy and reproducibility as long as patency of the catheter is ensured.11–13

The Stryker Intracompartmental Compartment Pressure Monitoring System (Kalamazoo, MI) has become the most commonly used commercially available device to measure compartment pressure in the emergency department (ED) (see Review Box 54.1). This device uses a fluid-filled pressure measurement catheter, a pressure monitor, and a fluid infusion mechanism that maintains catheter patency and ensures accurate measurement. In contrast to earlier devices in which relatively large volumes of fluid were injected into the compartment to measure pressure, the Stryker system uses a minimal amount of saline (< 0.3 mL). This helps ensure accurate measurements particularly true in uncooperative, unconscious, or critically injured patients who are unable to report symptoms. Unfortunately, the vagaries of the clinical scenario that result in failure to recognize the early signs and symptoms of compartment syndrome may have severe and irreversible limb- or life-threatening consequences.

Although compartment syndrome is essentially a clinical diagnosis, objective measurement of compartment tissue pressure may assist in confirming the diagnosis and determining if operative management is required. This chapter discusses the indications, complications, and interpretation of compartment pressure monitoring, as well as the equipment and techniques required to measure and monitor compartment pressure.

**BACKGROUND**

Though recognized as a clinical syndrome in the mid-19th century, the pathophysiology of ischemia in extremities was not fully described until more than a century later. Postischemic myoneural dysfunction and its associated contractures were first described in the 1870s by the German surgeon Richard von Volkmann, who recognized the effects of increased pressure causing vascular compromise of the limb.4

**BOX 54.1 Causes of Compartment Syndrome**

<table>
<thead>
<tr>
<th>DECREASED COMPARTMENTAL VOLUME</th>
<th>Intraarterial drug injection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Closure of fascial defects</td>
<td>Cold</td>
</tr>
<tr>
<td>Application of excessive traction to fractured limbs</td>
<td>Orthopedic surgery</td>
</tr>
<tr>
<td>Compressive devices (casts, splints, circumferential dressings)</td>
<td>Tibial osteotomy</td>
</tr>
<tr>
<td></td>
<td>Hauser’s procedure</td>
</tr>
<tr>
<td></td>
<td>Reduction and internal fixation of fractures</td>
</tr>
<tr>
<td></td>
<td>Snakebite</td>
</tr>
</tbody>
</table>

**BACKGROUND**

Over the past 40 years or so, various needles and equipment have been developed to measure compartment pressure (Fig. 54.2).5–11 The \textit{wick catheter}, originally developed to measure subcutaneous and brain tissue pressure, was modified during the mid-1970s to provide continuous compartment pressure measurements. This catheter is rarely used today because of the fear of catheter breakdown leading to errors in measurement and retained foreign bodies. The \textit{slit catheter} was introduced in 1980.10,11 As its name implies, this catheter has slits at its distal end that prevent clogging. The proximal end of the catheter is connected to a transducer and infusion system, which permits continuous pressure monitoring. Both the wick and slit catheters have been shown to offer similar accuracy and reproducibility as long as patency of the catheter is ensured.11–13

The Stryker Intracompartmental Compartment Pressure Monitoring System (Kalamazoo, MI) has become the most commonly used commercially available device to measure compartment pressure in the emergency department (ED) (see Review Box 54.1). This device uses a fluid-filled pressure measurement catheter, a pressure monitor, and a fluid infusion mechanism that maintains catheter patency and ensures accurate measurement. In contrast to earlier devices in which relatively large volumes of fluid were injected into the compartment to measure pressure, the Stryker system uses a minimal amount of saline (< 0.3 mL). This helps ensure accurate measurements particularly true in uncooperative, unconscious, or critically injured patients who are unable to report symptoms. Unfortunately, the vagaries of the clinical scenario that result in failure to recognize the early signs and symptoms of compartment syndrome may have severe and irreversible limb- or life-threatening consequences.

Although compartment syndrome is essentially a clinical diagnosis, objective measurement of compartment tissue pressure may assist in confirming the diagnosis and determining if operative management is required. This chapter discusses the indications, complications, and interpretation of compartment pressure monitoring, as well as the equipment and techniques required to measure and monitor compartment pressure.
The remainder of this chapter describes the most commonly used devices and techniques for measuring compartment pressures in the acute setting. Each method provides rapid measurements with reasonable accuracy. The method chosen will depend on the availability of the supplies and equipment necessary for a particular procedure and the experience of the operator.

(Video 54.1 and 54.2) While reducing the chance of further increases in compartment pressure, the Styrker system also has the ability to record a single measurement or provide continuous compartment pressure recordings when required.

Noninfusion systems such as the transducer-tipped fiber optic system offer a distinct advantage over conventional fluid-filled systems in that they do not produce hydrostatic pressure artifacts or require the injection of fluid for long-term or continuous measurements. However, the fiber optic transducer is relatively large, must be attached to a sheath approximately 2.0 mm in diameter, and is likely to cause pain during measurements.

A newer approach to predicting the onset of compartment syndrome involves measuring compartment pH as a marker of compromised circulation and decreased tissue perfusion. A rise in lactic acid from ischemic tissue lowers the pH of the compartment and has promise as an early predictor of compartment syndrome.

In recent years, noninvasive, less painful methods of measuring compartment pressure have been studied in patients with both acute and chronic exertional compartment syndromes. Investigations using magnetic resonance imaging (MRI), single-photon emission computed tomography (SPECT), myotonometry, electromyography, ultrasound, near-infrared spectroscopy, and microwave tomography have provided encouraging results in the evaluation of compartment syndrome. In addition, externally applied devices that measure muscle tissue “hardness” are under investigation as an economic alternative to these modalities, although support for their use has been mixed. Though promising, these evolving noninvasive methods have not yet replaced needle-driven techniques as the standard for measuring intracompartmental pressure.

The remainder of this chapter describes the most commonly used devices and techniques for measuring compartment pressures in the acute setting. Each method provides rapid measurements with reasonable accuracy. The method chosen will depend on the availability of the supplies and equipment necessary for a particular procedure and the experience of the operator.

(Please note that the image contains figures 54.1 and 54.2 which are not transcribed here.)

**PATHOPHYSIOLOGY**

Several theories have been proposed to account for the tissue ischemia associated with compartment syndrome. They include the “arteriovenous (AV) gradient” theory, which suggests that reduced AV pressure-perfusion gradients prevent adequate blood supply; the “critical closure” theory, in which blood flow is arrested well before the AV perfusion gradient declines to zero; and the “venous occlusion” theory, which states that externally applied pressure, thrombotic events, and reperfusion contribute to the increased compartment pressure and, ultimately, tissue ischemia. Although the exact mechanism has not been agreed on, inherent in each of these theories is a decrease in blood flow to levels below those required to meet the metabolic demands of the tissue. Hence, the final common pathway is cellular hypoxia and muscle necrosis.

Adequate blood flow to tissues is a function of AV gradients across capillary beds. Once blood flow falls below a critical level, delivery of oxygen to these structures is impaired and aerobic cellular metabolism is no longer possible. Anaerobic metabolism then ensues until energy stores become depleted. Muscles then become ischemic, and a reduction in venous and lymphatic drainage creates increased pressure within this confined space. It is important to note that ischemia and necrosis of the musculature can occur despite an arterial pressure high enough to produce pulses, therefore merely assessing distal pulses is insufficient.

A drop in blood pressure, an increase in compartment pressure, or a combination of the two can reduce AV gradients and lead to insufficient blood flow. Hypotension can occur in a variety of settings, including hypovolemia, acute blood loss, cardiac disease states (e.g., ischemia), and sepsis. An increase in the contents of a compartment, a decrease in its volume capacity, and external constriction of a compartment can all lead to increases in compartment pressure. Thus, the relationship between intracompartmental pressure and the circulatory status of the extremity is an important factor in the development of compartment syndrome.

At rest, normal adult compartment pressures are typically below 10 mm Hg. However, deviations of 2 to 6 mm Hg have been reported. Data suggest that normal compartment pressures in the lower extremity at rest are higher in children. The perfusion pressure of a compartment is defined as the difference between diastolic blood pressure and intracompartmental pressure. A model using legs of volunteers has shown that a progression of neuromuscular deficits occurs when the perfusion pressure drops below 35 to 40 mm Hg. Below this level, tissue perfusion is interrupted. Studies of neuromuscular tissue ischemia have demonstrated that inflammatory necrosis can occur at intracompartmental pressures between 40 and 60 mm Hg.

Whitesides and colleagues demonstrated that when tissue pressure within a closed compartment rises to within 10 to 30 mm Hg of the patient’s diastolic blood pressure, inadequate perfusion ensues and results in relative ischemia of the involved limb. Heppenstall and associates further clarified this relationship by demonstrating that the difference (ΔP) between mean arterial pressure (MAP) and the measured compartment pressure is directly related to blood flow to the tissue. They noted that as compartment pressure approaches MAP, ΔP decreases. Once ΔP falls below 30 mm Hg, tissue ischemia becomes more likely. In normal musculature, a ΔP of less than 30 mm Hg results in loss of normal aerobic cellular metabolism. In

traumatized muscle, a ΔP of less than 40 mm Hg is associated with abnormal cellular function, thus highlighting the importance of maintaining adequate systemic blood pressure in the setting of neuromuscular injury.84

For years, conventional wisdom maintained that immediate reperfusion of traumatized tissue would provide improved motor and neurologic function after injury. In the last decade, research suggests that muscle tissue may remain viable even after prolonged periods of ischemia and that a substantial proportion of the injury is generated during reperfusion.69,70 Tissue acidosis, intracellular and extracellular edema, free radical–mediated injury, loss of adenine nucleotide precursors, and interruption of mitochondrial oxidative phosphorylation by increased intracellular calcium have been implicated in the development of reperfusion-associated compartment syndrome.49-54 A 2010 study has identified the potential role of N-acetylcysteine in the attenuation of tissue injury in compartment syndrome.55 Even in the absence of local trauma, ischemia followed by reperfusion has been shown to increase compartment pressure in canine models of hypovolemic shock.56

Evidence also suggests that elevated compartment pressure itself (in addition to causing ischemia) plays a role in the cellular deterioration seen with compartment syndrome.57 In a study by Heppenstall and associates,57 muscle ischemia caused by placement of a tourniquet was compared with an experimentally derived high-pressure compartment syndrome. The authors found no difference in the degree to which phosphocreatine levels fell between groups. However, levels of adenosine triphosphate (ATP) diminished rapidly in the compartment syndrome group in comparison to the tourniquet group. Moreover, phosphocreatine levels, ATP, and tissue pH normalized within 15 minutes of releasing the tourniquet. In the group with compartment syndrome, these levels remained low even after fasciotomy. These results suggest that elevated tissue pressure plays a synergistic role with ischemia in cellular deterioration.

**CLINICAL FEATURES**

Any compartment limited by fascial planes is potentially at risk for compartment syndrome. However, because of their propensity for injury and the presence of several low-volume compartments, the lower extremities are most commonly affected. In the leg, the anterior compartment is involved most often,58 whereas the posterior compartment is most often associated with a delayed diagnosis. The hands, feet, forearms, upper part of the arms, thighs, thorax, abdomen, gluteal musculature, and back are other locations where compartment syndrome is known to occur.1

Risk factors for the development of compartment syndrome include recent trauma to an extremity (including acupuncture, venipuncture, IV infusions, or IV drug use), vascular injury, bleeding within a compartment, a restrictive cast or splint, a crush or compression injury, a prolonged lithotomy position, prolonged external pressure on an extremity, placement of a tourniquet during an operative procedure, and circumferential burns.55-64

Long-bone fractures account for approximately 75% of traumatic compartment syndromes, and the absence of a fracture in a traumatized extremity is a factor in delayed diagnosis. The tibia is most often involved, followed by bones of the forearm. Supracondylar fractures in children are at risk for compartment syndrome. Comminuted fractures increase the risk. Closed fractures are of greatest concern, but open fractures do not necessarily decompress elevated compartment pressures. Treatment of fractures, by both open and closed reduction, can increase compartment pressures. Compartment pressures may peak immediately after reduction. In addition, some evidence suggests that compartment syndrome may occur in the setting of chronic exertion and muscle overuse.65,66 Although the exact etiology remains elusive, studies have demonstrated elevated lactate and water levels in the tibialis anterior muscle after exercise with a reduction in these levels after fasciotomy.66,67 Increases in muscle mass (related to a rise in blood volume during exertion) and hypertrophy of muscle and fascia with chronic use and exertion have also been reported.68-72

The signs and symptoms of acute compartment syndrome generally appear in a stepwise fashion, with rapid progression over a few hours. This underscores the need for serial examinations, particularly in those with equivocal or subtle findings. The classic findings associated with arterial insufficiency, the “five P’s” (pain, pallor, pulselessness, paresthesias, paralysis), are often described as signs of acute compartment syndrome, but these are seldom all seen together and do not need to be present to suspect the diagnosis. When this constellation of signs and symptoms are present, it is generally late in the course of the disease when tissue death and permanent dysfunction are likely.

Early signs of compartment syndrome include a burning sensation over the involved compartment, nonspecific sensory deficits, or poorly localized deep muscular pain. Common features include pain that seems out of proportion to the apparent injury and clinical examination and pain that intensifies when the musculature is passively stretched. Pallor, a pulse deficit with respect to the opposite limb, paresthesias, paresis, or paralysis are variably seen and lack diagnostic sensitivity. Neurologic complaints such as weakness and paresthesias can be confusing because peripheral nerve injury may also result from the inciting trauma. In addition, these findings are not discernable in young children57,73 and patients with altered mental status. When present, sensory deficits typically precede motor deficits and manifest distal to the involved compartment. Paralysis is always a late finding.

The period between the injury and the onset of symptoms can be as short as 2 hours and as long as 6 days.75 The peak interval appears to be 15 to 30 hours. Frequently, the first symptom described by patients is pain greater than expected given the clinical scenario. Although pain out of proportion to the visible injury may raise the question of drug-seeking behavior, a focused examination for the possibility of limb-threatening disorders must precede this diagnosis of exclusion.

Physical examination may reveal a tense compartment with a firm, “woody” feel, muscles that are weak, and hypoesthesia in the distribution of the involved nerves. Sensory deficits, including loss of two-point discrimination and decreased vibratory sensation, may be present.76-78 The presence or absence of a palpable arterial pulse is not an accurate indicator of relative tissue pressure or the risk for compartment syndrome. Pulses may be present in a severely compromised extremity.77 Table 54.1 lists the signs and symptoms of compartment syndrome specific to each compartment.

**DIAGNOSIS**

Even experienced clinicians may find it difficult to diagnose a compartment syndrome, and no specific standard of care
Notwithstanding the difficulties just described, the diagnosis of compartment syndrome is primarily a clinical one that may be supplemented by direct measurement of compartment pressure. In a study evaluating the utility of clinical findings in making the diagnosis of compartment syndrome, Ulmer noted that the sensitivity and positive predictive value of clinical findings are low, whereas the specificity and negative predictive value of these findings are high. Nonetheless, the study found that although the sensitivity of an individual clinical finding may be low, the probability of compartment syndrome rises considerably when more than one clinical hallmark is present. Other studies have suggested that the absence of clinical evidence is more useful in excluding compartment syndrome than its presence is in confirming the diagnosis. All things considered, compartment syndrome remains largely a clinical diagnosis, and a high index of suspicion is paramount.

The differential diagnosis of compartment syndrome is extensive and includes primary vascular, nerve, and muscle injuries that produce similar findings. Acute arterial occlusion, cellulitis, osteomyelitis, neuropraxia, reflex sympathetic dystrophy, synovitis, tenosynovitis, stress fractures, envenomations, necrotizing fasciitis, deep vein thrombosis, and thrombophlebitis are additional diseases that should be considered. Differentiating

---

**TABLE 54.1 Compartment Syndromes and Associated Physical Signs**

<table>
<thead>
<tr>
<th>COMPARTMENT</th>
<th>SENSORY LOSS</th>
<th>MUSCLES WEAKENED</th>
<th>PAINFUL PASSIVE MOTION</th>
<th>LOCATION OF TENSENESS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foot</td>
<td>Digital nerves</td>
<td>Foot intrinsics</td>
<td>Toe flexion, extension</td>
<td>Dorsal or plantar surface of the foot</td>
</tr>
<tr>
<td>Lumbar</td>
<td>—</td>
<td>Erector spinae</td>
<td>Lumbar flexion</td>
<td>Paraspinous</td>
</tr>
<tr>
<td>Forearm</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dorsal</td>
<td>—</td>
<td>Digital extensors</td>
<td>Digital flexion</td>
<td>Dorsal surface of the forearm</td>
</tr>
<tr>
<td>Volar</td>
<td>Ulnar, median nerves</td>
<td>Digital flexors</td>
<td>Digital extension</td>
<td>Volar surface of the forearm</td>
</tr>
<tr>
<td>Hand</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Interosseus</td>
<td>—</td>
<td>Interosseus</td>
<td>Abduction/adduction (metacarpophalangeal joints)</td>
<td>Dorsum of the hand between the metacarpals</td>
</tr>
<tr>
<td>Leg</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anterior</td>
<td>Deep peroneal nerve</td>
<td>Toe extensors</td>
<td>Toe flexion</td>
<td>Anterior aspect of the leg</td>
</tr>
<tr>
<td>Superficial posterior</td>
<td>—</td>
<td>Soleus and gastrocnemius</td>
<td>Foot dorsiflexion</td>
<td>Calf</td>
</tr>
<tr>
<td>Deep posterior</td>
<td>Posterior tibial nerve</td>
<td>Toe flexors</td>
<td>Toe extension</td>
<td>Distal medial part of the leg between the Achilles tendon and tibia</td>
</tr>
<tr>
<td>Gluteal</td>
<td>(Rarely sciatic)</td>
<td>Gluteals, piriformis, or tensor fasciae latae</td>
<td>Hip flexion</td>
<td>Buttock</td>
</tr>
<tr>
<td>Upper Part of Arm</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flexor</td>
<td>Ulnar and median nerves</td>
<td>Biceps and distal flexors</td>
<td>Elbow extension</td>
<td>Anterior aspect of the upper part of the arm</td>
</tr>
<tr>
<td>Extensor</td>
<td>Radial nerves</td>
<td>Triceps and forearm extensors</td>
<td>Elbow flexion</td>
<td>Posterior aspect of the upper part of the arm</td>
</tr>
</tbody>
</table>
Invasive Compartment Pressure Monitoring

Indications and Contraindications

The earliest objective manifestation of acute compartment syndrome is an elevation in the tissue pressure of one or more compartments. However, signs and symptoms do not generally develop until tissue pressure has reached a critical level (see earlier section on Pathophysiology). In some patients the diagnosis of compartment syndrome is clinically obvious, and one can proceed directly to fasciotomy. When the clinical findings are equivocal or difficult to interpret, measurement of tissue pressure may help guide treatment. However, it must be remembered that the interpretation of such measurements always requires clinical judgment.

There are several groups of patients in whom clinical findings are difficult to obtain or interpret and who would benefit from measurement of compartment pressure. These groups include unresponsive patients, uncooperative patients, children, patients with multiple or distracting injuries, those with peripheral nerve deficits attributable to other causes (e.g., fracture-associated nerve injury, diabetic peripheral neuropathy), and those whose clinical findings are equivocal.

There are no absolute contraindications to measurement or continuous monitoring of compartment pressure. Caution should be taken when performing these procedures on patients with platelet dysfunction or other coagulation disorders. If possible, avoid inserting needles through overlying areas of infection, cellulitis, or burns.

ANCILLARY STUDIES

In general, laboratory and radiographic studies are not helpful in confirming the diagnosis of compartment syndrome. However, they might be useful in identifying other diagnoses, associated conditions, and complications. Box 54.2 lists useful studies for patients in whom compartment syndrome is suspected. A 2013 retrospective study of 97 patients by Valdez and colleagues attempted to determine if a threshold serum creatinine kinase (CK) level was associated with the development of compartment syndrome. They found that a CK level greater than 4000 U/L was associated with compartment syndrome and when serial measurements were obtained, the combination of a maximum CK greater than 4000 U/L, a maximal chloride level greater than 104 mg/dL, and a minimal blood urea nitrogen (BUN) level less than 10 mg/dL had a 100% association with compartment syndrome.

ANCILLARY STUDIES THAT MAY BE HELPFUL IN IDENTIFYING OTHER DIAGNOSES, ASSOCIATED CONDITIONS, AND COMPLICATIONS IN PATIENTS SUSPECTED OF HAVING COMPARTMENT SYNDROME

Laboratory Studies
- Complete metabolic profile (including electrolytes and renal function testing)
- Complete blood count with differential
- Serum and urine myoglobin
- Creatine phosphokinase
- Urinalysis to evaluate for concurrent rhabdomyolysis
- Coagulation studies

Imaging Studies
- Radiography of the affected limb to evaluate for a fracture or foreign body
- Ultrasonography to rule out deep vein thrombosis or Doppler ultrasonography to evaluate blood flow to the extremity

Patient Preparation and Positioning

Explain the procedure to the patient or surrogate. Written informed consent is not a universal standard, but may be an institutional requirement. Positioning the patient and extremity for measurement of the compartment pressure depends on the extremity and the compartment being studied, coexisting associated nerve injury, diabetic peripheral neuropathy, and those whose clinical findings are equivocal.

There are no absolute contraindications to measurement or continuous monitoring of compartment pressure. Caution should be taken when performing these procedures on patients with platelet dysfunction or other coagulation disorders. If possible, avoid inserting needles through overlying areas of infection, cellulitis, or burns.


| TABLE 54.2 Clinical Findings in Patients With Compartment Syndrome, Arterial Occlusion, and Neuropraxia |
|---------------------------------------------------------------|-------------------------------|-----------------|
| Pressure increased in the compartment | + | − | − |
| Pain with stretch | + | + | − |
| Paresthesia or anesthesia | + | + | + |
| Paresis or paralysis | + | + | + |
| Pulses intact | + | − | + |

and fascia. Inadvertent injection into the underlying structures may falsely elevate compartment pressure. Patient movement (as a result of inadequate analgesia or improper positioning) may also falsely elevate compartment pressure, particularly if the patient requires limb restraint. To minimize the discomfort of multiple attempts at localizing the correct compartment, some have advocated the use of ultrasound guidance to improve the accuracy of needle insertion.\(^9\) However, there is currently no consensus regarding the use of ultrasound guidance in this setting.\(^9\)\(^0\)\(^1\) In general, place the extremity being studied at the level of the heart and in a position that permits insertion of the needle perpendicular to the compartment being measured. This may require that an assistant hold the extremity above the stretcher. Remove any obstruction to needle entry and all objects that may constrict or exert pressure on the compartment.

Perform compartment pressure measurements with sterile technique, including standard skin preparation and draping at the planned insertion site (see Chapter 34). Avoid placing the needle through areas of overlying burn or infection. If a circumferential cast is present, bivalve the cast or, if necessary, create a window overlying the desired area with a cast saw.

**Equipment**

Needles commonly used for measurement of compartment pressure include a simple 18-gauge needle, an 18-gauge spinal needle (for deep compartments), and a side-port needle (Stryker Instruments, Kalamazoo, MI). The side-port needle may be more accurate than simple 18-gauge needles.\(^9\) Nevertheless, simple 18-gauge needles are more readily available and more commonly used. The wick catheter and slit catheter described previously generally require specialized, cumbersome equipment not often available in most EDs and are therefore not described.

In an effort to reduce the pain associated with insertion of larger-gauge needles, Mars and colleagues evaluated the accuracy and reliability of measuring compartment pressures with smaller-gauge needles.\(^9\)\(^2\) The authors compared compartment pressures measured with 18- to 25-gauge needles and found that smaller-gauged needles provided results similar to or better than the more traditional 18-gauge needles did. Furthermore, the addition of a side port did not improve accuracy.\(^9\)\(^2\) Unfortunately, no additional studies have been done to corroborate these findings. As a result, most clinicians continue to use 18-gauge needles.

**Pressure Measurement Systems**

**Stryker Intra-Compartmental Pressure Measurement (Fig. 54.3)**

**Equipment**

- Stryker Quick Pressure Monitor System
- Stryker handheld pressure monitoring unit (included)
- One side-port 18-gauge needle (included)
- One diaphragm chamber (included)
- One syringe prefilled with 3 mL of saline (included)

**Setup and Procedure (see Fig. 54.3)**

1. Position the patient. Prepare and anesthetize the needle insertion site according to the guidelines described earlier.
2. Open the Quick Pressure Monitor Set and remove the contents while maintaining sterile conditions.
3. Place the needle firmly on the tapered chamber stem.
4. Remove the cap on the prefilled syringe and screw the syringe onto the remaining chamber stem. Take care to avoid contaminating the fluid pathway.
5. Open the monitor cover and inspect the device for damage or contamination. Place the chamber into the device well (black surface down) and push gently until it seats.
6. Snap the cover closed, but DO NOT FORCE IT. The latch must have “snapped” in place.
7. Hold the needle at approximately 45 degrees from the horizontal plane and slowly force fluid through the disposable system to purge it of air. **Caution:** DO NOT allow saline to roll down the needle into the transducer well.
8. Turn on the unit; it should read between 0 and 9 mm Hg.
9. Hold the device at the intended angle of insertion and press the “ZERO” button to calibrate it. After a few seconds, the display should read “00.” Failure to calibrate the device while in the intended angle of insertion may result in inaccurate readings. **Note:** The display must read “00” before continuing. If it does not, follow the troubleshooting instructions provided by the manufacturer before proceeding.
10. Now insert the needle into the compartment being measured (see later section on Needle Placement Techniques). Slowly inject no more than 0.3 mL of saline into the compartment to equilibrate with the interstitial fluid.
11. Wait for the display to reach equilibrium and record the resulting pressure.
12. **For additional measurements,** turn the unit off and repeat steps 8 through 11. Recalibrate the unit to zero before each measurement.
13. For continuous monitoring with an indwelling slit catheter, refer to the instructions accompanying the system.

**Mercury Manometer System (Fig. 54.4)**

**Equipment**

- Two 18-gauge simple or spinal needles
- Two plastic extension tubes
- One 20-mL syringe
- One three-way stopcock
- One vial of sterile normal saline
- One mercury manometer

**Setup and Procedure**

1. Position the patient. Prepare and anesthetize the needle insertion site according to the guidelines described earlier.
2. Assemble the syringe, tubing, extension tubing, and needle as shown in Fig. 54.4A.
3. Insert the needle into a vented vial of sterile saline. Aspirate a column of saline into the tubing halfway to the stopcock; care should be taken to avoid the formation of bubbles. Close the three-way stopcock to the tube to prevent the loss of saline during insertion of the needle, and remove the needle from the vial of sterile saline.
4. Insert the needle into the muscle of the compartment being measured (see later section on Needle Placement Techniques).
5. Attach the second extension tubing to the monitor and to the third port of the three-way stopcock. Turn the stopcock so that the syringe is open to both extension tubes (see Fig. 54.4B). This closed system has equal pressure in both extension tubes.
6. Increase the pressure in the system gradually by slowly depressing the syringe plunger while simultaneously watching...
Compartment Pressure Evaluation: Stryker Method

1. Place the 18-gauge needle with a side port on the tapered stem of the diaphragm chamber.

2. Screw syringe prefilled with 3 mL of saline onto the back of the diaphragm chamber.

3. Place the diaphragm chamber assembly into the pressure monitor, black side down. Gently push the chamber until it is well seated in the device.

4. Snap the cover closed—do not force it! Listen for the latch to snap into place.

5. Hold the needle at a 45° angle up from horizontal and depress the plunger to force fluid through system and purge it of air.

6. Turn the pressure monitor on. It should read between 0 and 9 mm Hg.

7. Hold the pressure monitor at the intended angle of insertion and press the “ZERO” button to calibrate the unit. The device display should read “00.”

8. Insert the device into the compartment being measured (after skin cleansing and administration of anesthetic). Slowly inject no more than 0.3 mL into the compartment and wait for the device to record and display the pressure.

Figure 54.3 Measuring compartment pressure with the Stryker 295-2 pressure monitor. Normal tissue compartment pressure is between 0 and 8 mm Hg. The diagnosis of compartment syndrome is not based only on compartment pressures. The clinical evaluation must be correlated with measured pressures.
the column of saline. The mercury manometer will rise as pressure in the system increases. When the pressure in the system exceeds that in the tissue, injection of saline into the compartment will occur, which causes the saline column to move. Read the manometer at the moment that the saline is noted to move. This reading corresponds to the tissue pressure in millimeters of mercury.

7. To obtain a second reading, completely remove the needle and repeat steps 4 through 6. A third measurement might be necessary to achieve two readings in agreement. Check the needle for tissue plugs and blood clots between readings.

Procedural Caveats
The most common error with this system is depressing the syringe plunger too quickly. Only when saline is injected slowly into the compartment will the mercury column (which has greater inertia) accurately reflect compartment pressure. Another source of error is obstruction of the needle with a plug of tissue (or blood clot) if the plunger of the syringe is pulled back. Finally, aneroid manometers are prone to inaccuracy, are not well calibrated at lower pressure ranges, and should not be substituted for the more accurate mercury manometers for this procedure.

**NEEDLE PLACEMENT TECHNIQUES FOR SPECIFIC COMPARTMENTS**

**General Principles**
Accurate pressure measurements depend on careful needle insertion and confirmation of correct placement. Proper needle insertion requires (1) reliable placement in the compartment being measured, (2) avoidance of important neurovascular structures, (3) simplicity and reproducibility, and (4) minimal patient discomfort. Most compartments are superficial and easily accessible. Only the deep posterior compartment in the leg and the gluteal compartment may require a spinal needle to reach the required depth. Most approaches require that the needle enters the tissue perpendicular to the skin.

**Lower Extremity**
Because of its high vulnerability to injury and limited fascial compliance, the lower part of the leg is predisposed to compartment syndrome. The foreleg traditionally has four compartments: anterior, lateral, deep posterior, and superficial posterior (Fig. 54.5). The anterior compartment is the most frequent site of compartment syndrome. In some patients the tibialis posterior muscle may occupy its own compartment, separate from the rest of the deep posterior compartment. Keep this possibility in mind when measuring compartment pressure in this area.

The easiest cross-sectional level for placement of the needle in any foreleg compartment is approximately 3 cm on either side of a transverse line drawn at the junction of the proximal and middle thirds of the lower part of the leg (see Fig. 54.5). When measuring compartment pressure in the foreleg, place the patient in the supine position with the leg at the level of the heart. An exception occurs when measuring the superficial posterior compartment. In this case, place the patient in the prone position. Prepare and anesthetize the needle insertion site as described previously in this chapter.

**Anterior Compartment (Fig. 54.6A)**
With the patient supine, palpate the anterior border of the tibia at the junction of the proximal and middle thirds of the lower part of the leg. Insert the needle perpendicular to the skin 1 cm lateral to the anterior border of the tibia to a depth of approximately 1 to 3 cm. A several fold rise in pressure during (1) external compression of the anterior compartment just proximal or distal to the needle insertion site, (2) plantar flexion of the foot, or (3) dorsiflexion of the foot confirms proper needle depth.

**Figure 54.4** A and B, Mercury monitor technique for monitoring compartmental pressure. This system is not usually applicable to use in the emergency department. IV, intravenous. (A and B, From Whitesides TE, Haney TC, Morimoto K, et al: Tissue pressure measurements as a determinant for the need of fasciotomy, *Clin Orthop* 113:43, 1975.)
Compartment Syndrome Evaluation

Deep Posterior Compartment (see Fig. 54.6B)
With the patient supine, have an assistant elevate the leg slightly off the stretcher (if the clinical situation permits). Palpate the medial border of the tibia at the junction of the proximal and middle thirds of the lower part of the leg while simultaneously palpating the posterior border of the fibula on the lateral aspect of the leg at the same level. Insert the needle perpendicular to the skin just posterior to the medial border of the tibia and direct it toward the palpated posterior border of the fibula to a depth of 2 to 4 cm (the final depth depends on the amount of subcutaneous adipose tissue). Confirm proper needle depth by observing a rise in pressure during (1) toe extension or (2) ankle eversion.

Lateral Compartment (see Fig. 54.6C)
With the patient supine and the leg at heart level, have an assistant elevate the leg slightly off the stretcher (if the clinical situation permits). Palpate the posterior border of the fibula at the junction of the proximal and middle thirds of the lower part of the leg. Insert the needle into the skin just anterior to the posterior border of the fibula and direct it toward the fibula to a depth of 1 to 1.5 cm. If the needle contacts bone, withdraw it approximately 0.5 cm. Confirm proper needle depth by observing a rise in pressure during (1) external compression of the lateral compartment just inferior or superior to the needle’s entrance or (2) inversion of the foot and ankle.

Superficial Posterior Compartment (see Fig. 54.6D)
With the patient in the prone position and the leg at heart level, identify an imaginary transverse line (or draw one with a marking pen) between the proximal and middle thirds of the lower part of the leg. Insert the needle perpendicular to the skin at this level, 3 to 5 cm on either side of a vertical line drawn down the middle of the calf. Direct the needle toward the center of the leg to a depth of 2 to 4 cm. Confirm proper needle depth by observing a rise in pressure during (1) digital external compression of the posterior compartment just inferior or superior to the needle insertion point or (2) dorsiflexion of the foot.

Forearm
Traditionally, the forearm has been considered a two-compartment limb. However, some authors place the extensor carpi radialis brevis, the extensor carpi radialis longus, and the brachioradialis muscles in a third compartment called the “mobile wad.” The forearm compartments (particularly the volar compartment) are predisposed to compartment syndrome because of their use during vigorous exercise, accessibility for drug use, IV infiltration of fluid or medication, and vulnerability to injury and burns.

The junction of the proximal and middle thirds of the forearm is the cross-sectional level for insertion of the needle.

Figure 54.5 Fascial compartments of the lower part of the leg with enclosed muscle groups (insert, upper left): 1, anterior; 2, lateral; 3, superficial posterior; and 4, deep posterior compartments.
When measuring forearm compartment pressure, place the patient in the supine position with the arm at the level of the heart. Prepare and anesthetize the needle insertion site as previously described.

**Volar Compartment (Fig. 54.7A)**

Hold the forearm in supination. Identify the palmaris longus tendon by having the patient oppose the thumb and small finger with the wrist flexed against resistance. Follow the tendon to the junction of the proximal and middle thirds of the forearm. Palpate the posterior border of the ulna and insert the needle perpendicular to the skin just medial (ulnar) to the palmaris longus tendon. Direct the needle toward the palpated posterior border of the ulna to a depth of 1 to 2 cm. Confirm proper needle depth by observing a rise in pressure during (1) external compression of the volar compartment just proximal or distal to the needle insertion point or (2) extension of the fingers or wrist.

**Dorsal Compartment (see Fig. 54.7B)**

Hold the forearm in pronation with the elbow flexed and the dorsum of the forearm facing upward. Palpate the posterior aspect of the ulna at the junction of the proximal and middle thirds of the forearm. Insert the needle perpendicular to the upper extremity compartments

---

**Lower Extremity Compartments**

**A. Anterior**

Place the patient in the supine position. Insert the needle at the junction of the proximal and middle thirds of the lower part of the leg, 1 cm lateral to anterior border of the tibia. Direct the needle perpendicular to the skin a depth of 1 to 3 cm.

**C. Lateral**

Place patient in the supine position with leg slightly elevated. Insert the needle at the junction of the proximal and middle thirds of the lower part of the leg, just posterior to the medial border of the tibia. Direct the needle perpendicular to the skin and toward the posterior border of the fibula to a depth of 2 to 4 cm.

**B. Deep Posterior**

Place patient in the supine position with the leg slightly elevated. Insert the needle at the junction of the proximal and middle thirds of the lower part of the leg, and just posterior to the medial border of the tibia. Direct the needle perpendicular to the skin and toward the posterior border of the fibula to a depth of 2 to 4 cm.

**D. Superficial Posterior**

Place the patient in the prone position. Insert the needle at the junction of the proximal and middle thirds of the lower part of the leg, 3 to 5 cm on either side of the anatomic midline. Direct the needle perpendicular to the skin toward the center of the leg to a depth of 2 to 4 cm.

**Figure 54.6 Lower extremity needle placement techniques.** (From Castalow CB: Color atlas of emergency department procedures. Philadelphia, 2005, Saunders.)
Compartment Syndrome Evaluation

Chapter 54

Gluteal Musculature

A two-layer fascia encases the muscle bellies of the tensor fasciae latae anteriorly and the gluteus maximus posteriorly. This fascia divides the musculature into three distinct compartments: maximus, tensor, and medius/minimus (Fig. 54.8A and B). The sciatic nerve is deep to the fascia but lies between the

Figure 54.7 Upper extremity needle placement techniques. (From Custalow CB: Color atlas of emergency department procedures. Philadelphia, 2005, Saunders.)
Although compartment syndrome of the foot is rare, it is being reported with increasing frequency as clinicians become more aware of this entity. Despite a lack of universal agreement on the number or exact location of the anatomic compartments in the foot, it is generally accepted that there are at least nine compartments separated into four groups: the central/calcaneal, intrinsic/interosseous, medial, and lateral compartments. For measurement of pressure in any of the foot compartments, place the patient in the supine position with the foot at the level of the heart and prepare and anesthetize the needle insertion site as described previously. Note that pedal edema has been shown to increase resting tissue pressure in the foot.

**Medial Compartment (see Fig. 54.9B)**

The medial compartment contains the abductor hallucis and flexor hallucis brevis muscles. The compartment is bounded medially and inferiorly by an extension of the plantar aponeurosis, laterally by an intramuscular septum, and dorsally by the first metatarsal. To measure pressure in this compartment, insert the needle perpendicular to the skin at the medial aspect of the foot just inferior to the base of the first metatarsal into the abductor hallucis muscle, which is approximately 1 to 1.5 cm deep. Confirm proper needle depth by observing a rise in pressure during external compression of the medial compartment of the foot.

**Central (Calcaneal) Compartment (see Fig. 54.9B)**

The central compartment contains the flexor digitorum brevis, the quadratus plantae, the lumbricals, and the abductor hallucis muscles. Its boundaries are the plantar aponeurosis inferiorly, the osteofascial tarsometatarsal structures dorsally, and the
INTERPRETATION OF COMPARTMENT PRESSURE MEASUREMENTS

Compartment pressures must be interpreted within the context of the clinical picture. Inaccurate measurements are far worse than no measurement at all, and clinical evaluation is more telling than pressure measurements alone. To be most accurate, compartment pressures should be measured in the area of highest pressure and greatest tissue damage. Inaccurate measurements may result from placement of the needle into tendon or fascia, plugged needles, defective or poorly calibrated devices, injection of fluid into the compartment, and movement during the procedure. Whitesides and colleagues found a 1 mm Hg increase in compartment pressure for every 1 mL of saline infused into the anterior compartment of the lower extremity. It is difficult to assess the relevance of this finding, but recognition of the potential for its occurrence is important.

Reports of normal human compartment pressure vary in the literature. In comparing several techniques, Shakespeare and associates found an average normal pressure of 8.5 mm Hg with slightly higher resting pressure in individuals who were physically fit. Willy and coworkers found a mean of 15 mm Hg (±8 mm Hg) when using an electronic transducer-tipped probe in healthy volunteers. Although the generally accepted range of normal is between 0 and 10 mm Hg, others have noted pressures in normal subjects ranging from 0 to as high as 18 mm Hg. Even though compartment pressure measurements have not been studied extensively in pediatric populations, one study reported higher resting compartment pressure in children than in adults.

When properly performed, each method described in this chapter has acceptable accuracy in the clinical setting. Studies...
have found standard deviations of between 2 and 6 mm Hg with all the techniques described earlier. It is generally agreed that the mercury manometer method of measuring compartment pressure is the least accurate. The arterial line system used with a simple (straight) or side-port needle provides a higher degree of accuracy for simple, episodic readings. The Stryker Intra-Compartmental Pressure Monitoring System provides consistent, accurate readings for both episodic and continuous pressure monitoring. The development of miniature transducer-tipped devices is ongoing. Noninvasive modalities, including MRI, SPECT, myotonometry, electromyography, near-infrared spectroscopy, and ultrasound, continue to be investigated as painless alternatives to needle-driven compartment pressure measurements, and although the early results are promising, they are not currently in widespread use.

Mubarak and coworkers and Hargens and colleagues suggested that an absolute compartment pressure of 30 mm Hg is the “critical pressure” requiring fasciotomy. Others have supported these recommendations. However, it is important to note that a compartment pressure of 30 mm Hg does not necessarily precipitate compartment syndrome in the absence of other factors and must be interpreted within the context of the clinical scenario. Although compartment syndrome may develop in some patients at this pressure, in others it will not, as tolerance to increasing pressure appears to be variable.

Despite a body of literature addressing measurement of compartment pressure, there continues to be no consensus regarding a specific compartment pressure threshold at which fasciotomy should be performed. Obviously, no measured compartment pressure, by itself, is an indication for fasciotomy, but it may guide consultation and further observation. A single pressure reading can be misleading, and trends in pressure provide more information to the clinician. Some argue that an absolute compartment pressure of 30 mm Hg or greater should be the threshold for fasciotomy. Matsen found that no patients with a pressure lower than 45 mm Hg had symptoms of compartment syndrome, whereas all patients with a pressure higher than 60 mm Hg had symptoms. Whitesides and colleagues found that fasciotomy was required when intracompartmental pressure approaches 20 mm Hg below diastolic pressure, whereas McQueen and associates recommended using a perfusion pressure (diastolic pressure minus compartment pressure) of less than 30 mm Hg as a criterion for fasciotomy. Heppenstall and coworkers concluded that use of either the Whitesides or McQueen criteria would produce similar results and recommended using a ΔP (MAP minus the measured compartment pressure) of 30 mm Hg or less in nontraumatized muscle and a ΔP of 40 mm Hg or less in traumatized muscle as a guide for fasciotomy.

In summary, absolute compartment pressures of 30 mm Hg or higher garner concern for consultation or intervention for compartment syndrome. Based on pathophysiology, it appears quite reasonable to consider the difference between diastolic blood pressure and the measured compartment pressure (perfusion pressure) as a guide for diagnosing compartment syndrome and a strong consideration for fasciotomy. A compartment pressure within 30 mm Hg of the patient’s diastolic blood pressure, essentially a perfusion pressure of less than 30 mm Hg, is generally considered the threshold for consideration of fasciotomy.

Factors other than compartment pressure alone are important in the development of compartment syndrome and the need for fasciotomy. In patients with chronic systemic hypertension, higher compartment pressure may be necessary before nerve injury or muscle ischemia occurs. Situations in which MAP is low (e.g., hypovolemia, peripheral vascular disease) might interfere with the patient’s ability to tolerate even mildly elevated compartment pressure. Finally, the duration of increased compartment pressure is also an important factor in the development and severity of compartment syndrome.

COMPLICATIONS

All invasive monitoring systems are associated with some degree of pain during insertion of the needle or catheter. Adequate analgesia and, when necessary, procedural sedation are often necessary to gain the patient’s cooperation and prevent movement during pressure measurements. When using local anesthetics, avoid injections into the compartment because this can increase pain and result in inaccurate (higher) readings. Failure to properly use pressure measurement devices may result in inaccurate results and a delay in the diagnosis of compartment syndrome. As with any invasive procedure, care must be taken to minimize bleeding and additional damage to underlying tissue, nerves, or blood vessels. The risk for both local and systemic infection is similar for all the measurement procedures described in this chapter. Strict adherence to aseptic technique and universal precautions is mandatory. This includes sterilization of catheters and the use of sterile solutions in addition to sterile gloves and supplies whenever possible.

ACKNOWLEDGMENTS

The author recognizes the original contributions of Neal R. Frankel, DO, and L. Albert Villarin, Jr., MD, to a previous edition of this text.
REFERENCES


INTRODUCTION

This chapter addresses urologic conditions that are either initially or eventually associated with an emergency procedure or may need to be performed in the absence of a urologic surgeon.

Testicular torsion is a scrotal emergency that can be challenging to diagnose under the best clinical circumstances. Although surgical exploration is the only definitive diagnostic and therapeutic procedure, many urologists prefer an initial diagnostic imaging study prior to surgical exploration, provided imaging can be obtained expediently while simultaneously preparing for operative intervention. This chapter addresses bedside maneuvers for this entity, including manual testicular detorsion.

Emergency penile conditions include paraphimosis and priapism. This chapter addresses bedside paraphimosis reduction techniques, as well as penile aspiration-irrigation-injection for relief of ischemic priapism. In addition, relief of phimosis complicated by the inability to void by means of a dorsal slit of the penile foreskin will be detailed for completeness (Video 55.1).

Access to and the subsequent evaluation of the bladder and urine are clinically important to all emergency practitioners. Various approaches to urine sampling and bladder drainage, including the techniques and complications of male and female urethral catheterization in various clinical situations and emergency suprapubic bladder access, are addressed.

Finally, a discussion of radiographic imaging of the genitourinary (GU) system is provided, with an emphasis on assessing lower urinary tract injury. Although the timing of GU radiologic examinations within the workup of the critically ill, multitrauma patient must be individualized, general guidelines for successful imaging are provided.

TESTICULAR TORSION

Introduction

Although acute scrotal pain comprises less than 1% of overall emergency department (ED) visits, this presentation may provoke great anxiety for the patient or caretaker, given its highly sensitive nature. One of the most challenging aspects of scrotal complaints is that a wide variety of clinical conditions may present in a similar fashion: a male patient complaining of an acute, painful, swollen, and tender hemiscrotum. Although the differential is extensive, testicular torsion is the principle fertility threat that needs to be ruled out. Definitive management of testicular torsion involves surgical exploration and orchiopexy. Manual detorsion, with or without spermatic cord anesthesia, can be attempted while simultaneously preparing for operative intervention.

Procedure Overview

Indications: testicular torsion.
Contraindications: alternative cause of acute scrotal pain.
Complications: worsening ischemia, increased pain.

Equipment:
- for spermatic cord anesthesia: 1% lidocaine (10 mL for an adult, or age-/weight-appropriate local anesthetic dosing for child), 27-gauge (or similar) needle for anesthetic infiltration.
- for manual detorsion: bedside sonography may be helpful in assessing for pre- and postprocedure intratesticular blood flow.

Key Procedure Sequence

1. Initiate urologic consultation and simultaneous preparation for surgical intervention.
2. Administer spermatic cord anesthesia (optional).
3. Administer systemic analgesia or light sedation (optional).
4. Keep the patient comfortable in a reclining or supine position.
5. Stand beside the stretcher (right side if right-hand dominant, or vice versa).
6. Rotate the testicle from medial to lateral (two thirds of torsion occurs lateral to medial).
7. Rotate 180 degrees initially; it may ultimately require two to three rotations to alleviate pain.
8. If mechanically difficult or there is increased pain, detorse in the opposite direction.
9. The end point is relief of pain or return of intratesticular blood flow on sonogram.
10. Consult urology department for definitive surgical scrotal exploration.

Description

Background
An acute scrotum is defined as an acute, painful swelling of the scrotum or its contents, accompanied by local signs or general symptoms. Although the list of diagnostic possibilities for a patient with an undifferentiated acute scrotum is extensive, early identification and skillful management of testicular torsion is critical as it may threaten testicular viability and future fertility if not managed expeditiously. Acute epididymitis is commonly the cause of acute scrotal pain in adolescents and adults. Testicular (or epididymal) appendage torsion is another frequent cause of acute scrotal pain in prepubertal boys.

Differentiating testicular torsion from alternative conditions takes precedence over a definitive diagnosis. The presence of an intact cremasteric reflex and testicular sonography are frequently utilized, yet imperfect, diagnostic tools in assessing for testicular torsion.

Anatomy and Physiology
A congenital anomaly of fixation of the testis, termed the bell-clapper deformity, is associated with the development of testicular torsion. This occurs when the intrascrotal portion of the spermatic cord lacks firm posterior adhesion to the scrotal wall and remains surrounded by the tunica vaginalis (Fig. 55.1). As a result of the abnormal attachment, the testis may be suspended horizontally. These anatomic features predispose the affected testis to rotation.

Pathophysiology
Testicular salvage rates decrease with time from the onset of ischemia. A meta-analysis of 1140 patients in 22 series demonstrated a greater than 90% salvage rate with surgery within 6 hours of pain onset. Testicular atrophy may lead to subfertility. Furthermore, testicular loss may lead to contralateral testicular dysfunction through immune-mediated or other mechanisms.

The testis may tors, detors, and retors, so clinicians should be cautious about assigning an exact time as the time of onset and deferring urologic consultation based on an erroneous assumption that the testis is not salvageable. The gold standard for determining testicular viability is intraoperative visualization of the affected testis, which dictates early urologic involvement. Although all clinicians recognize the need for expedient surgery in the setting of known torsion, not all consultants will agree on surgical exploration without some adjunctive testing.

When utilized in the appropriate clinical setting, ultrasound remains the most useful diagnostic modality in the evaluation of GU complaints (Fig. 55.2). A patient with compelling historical and examination findings of testicular torsion does not require any preoperative diagnostic tests. Color flow Doppler ultrasonography (CDUS) may be very helpful in all other cases. The classic sonographic finding suggestive of testicular torsion is diminished intratesticular blood flow. In addition, examination of the spermatic cord with high resolution gray-scale ultrasonography may reveal “coiling” or “kinking” of the cord at the site of torsion. Sonography is used not only to exclude testicular torsion, but also to search for alternative causes of acute scrotal pain. In epididymitis, perfusion may be normal (or increased) due to the effects of inflammatory mediators on local vascular beds, although this is a nonspecific finding. An infarcted appendage may be visualized on ultrasound as well. It has been suggested that emergency physicians may be able to accurately assess for intratesticular blood flow in patients presenting with acute scrotal pain using bedside sonography.

Color flow Doppler ultrasound has long been regarded as the diagnostic modality of choice in assessing for testicular torsion. However, false-negative ultrasound results have been reported. Many of these studies are case reports or case series, limited by small numbers and retrospective design. Two larger series reported documented intratesticular blood flow with CDUS in 6 of 23 (26%) and 50 of 208 cases (24%), respectively, of confirmed testicular torsion. Doppler ultrasound may reveal seemingly adequate intratesticular blood flow in partial torsion, which can be very misleading to the practitioner.

Radionuclide scintigraphy and CDUS show similar sensitivity, as well as false-negative rates, for the diagnosis of testicular torsion. However, given the widespread availability and expertise with ultrasound technology, combined with the risks of isotope radiation exposure, radionuclide procedures have fallen out of favor. The use of magnetic resonance imaging has been explored, but limitations include speed of imaging and availability.

Intravaginal testicular torsion is a congenital bilateral abnormality. The ischemic testis must be detorsed and pexed with nonabsorbable (e.g., nylon, polypropylene, or Prolene [Ethicon Inc.]), rather than absorbable (e.g., chromic, polyglandin, or Vicryl [Ethicon Inc.]), suture. The torsed testis that is pexed with absorbable suture remains at risk for subsequent postoperative torsion. Given the bilateral nature of the congenital abnormality, orchiopepexy of the nonischemic contralateral testis is necessary.

Once the diagnosis of testicular torsion is suspected, immediately place a call to notify a urologist of the suspected diagnosis, the perceived need for surgical exploration, and the fact that you will be attempting testicular detorsion while awaiting patient transport to the operating room. At some point before the patient leaves the ED, meticulous charting to document time, suspected diagnosis, notification of the urologist, and any manipulation of the affected testis must be done. All efforts are then focused on attempting testicular detorsion.

Indications
Testicular torsion may be relieved by manual detorsion. A study of 162 cases of testicular torsion revealed that anticipated lateral to medial rotation occurred in 67% of cases, with medial to lateral rotation in the remaining 33%. This challenges the standard dogma of medial to lateral rotation, or “opening the book,” as the standard method for detorsion. There may be a cranial-caudal component to torsion as well. The end point of manual detorsion is relief of pain, or the return of intratesticular blood flow as seen on ultrasound imaging. Although manual detorsion may allow for reperfusion of the testis, a lesser degree of residual torsion may remain. Given that infarction can occur with as little as 180 degrees of torsion, immediate surgical exploration after what is thought to be a successful manual detorsion is still advocated. The bottom line is that specialty consultation and plans for possible immediate...
**Manual Testicular Detorsion**

A. Anatomy of testicular torsion. Testicular torsion occurs when the testis twists within the tunica vaginalis. Patients with the bell-clapper deformity (i.e., incomplete fusion of the tunica along the epididymis, which results in incomplete attachment of the testicle to the scrotum) are at higher risk.

B. Spermatic cord block. Grasp the spermatic cord between your thumb and index finger. Use a 30-gauge needle to infiltrate the entire cross section of the spermatic cord and its surrounding rim with anesthetic. This will cause visual ballooning of the grasped segment of the cord. Gently massage this bulge to disperse the anesthetic. Usually about 10 mL is required.

C. Testicular torsion more commonly occurs in a medial direction. Initially attempt detorsion by rotating the testis outward toward the thigh. This is most successful if attempted within the first few hours of torsion, before the onset of significant scrotal swelling. Intravenous narcotics (such as fentanyl) can be administered or a cord block performed before attempting detorsion.

D. Detorsion maneuver. Detorsion of the testicle may require testicular rotation through two planes. To release the cremasteric muscle, rotate the testis in a caudal-to-cranial direction simultaneously with medial-to-lateral rotation. The right testis is shown.

attempted detorsion is that the anesthesia takes away an important subjective endpoint: relief of the patient’s pain after manipulation of the testis. However, many authors do advocate spermatic cord anesthesia before detorsion, and if anesthesia of the spermatic cord is elected, it can be done in the following manner.

**Spermatic Cord Anesthesia**

Local anesthesia of the spermatic cord using 1% plain lidocaine is usually done at the external inguinal ring (see Fig. 55.1B). First prepare the skin with an antiseptic solution. The cord can usually be grasped between the thumb and the index finger, and 10 mL of 1% plain lidocaine (for an adult) can be directly injected into the cord. If the cord is swollen, as is often the case in testicular torsion, or if the testicle is lying very high in the hemiscrotum as a result of spermatic cord torsion (so as to preclude grasping), the cord may be palpated at the pubic tubercle as it passes over the pubis and the lidocaine injected at this landmark. Lee and colleagues were able to perform surgical exploration need to occur regardless of outcome of the detorsion procedure. Detorsion, however, may convert an emergency procedure to an urgent one.

**Contraindications**

Presence of an alternative cause of acute scrotal pain is a relative contraindication to the detorsion procedure. However, precise diagnosis may be impossible prior to definitive surgical exploration. A nonanomalous, appropriately fixated testicle should not be adversely affected by an initial trial (e.g., 180 degrees) of manual detorsion if circumstances are highly suspect for spermatic cord torsion.

**Procedure: Manual Detorsion and Spermatic Cord Anesthesia**

Manual detorsion is performed in the following manner. Advise the patient that the procedure will be uncomfortable and painful and offer systemic analgesia or light sedation if appropriate. The rationale for not using spermatic cord anesthesia with attempted detorsion is that the anesthesia takes away an important subjective endpoint: relief of the patient’s pain after manipulation of the testis. However, many authors do advocate spermatic cord anesthesia before detorsion, and if anesthesia of the spermatic cord is elected, it can be done in the following manner.

**Spermatic Cord Anesthesia**

Local anesthesia of the spermatic cord using 1% plain lidocaine is usually done at the external inguinal ring (see Fig. 55.1B). First prepare the skin with an antiseptic solution. The cord can usually be grasped between the thumb and the index finger, and 10 mL of 1% plain lidocaine (for an adult) can be directly injected into the cord. If the cord is swollen, as is often the case in testicular torsion, or if the testicle is lying very high in the hemiscrotum as a result of spermatic cord torsion (so as to preclude grasping), the cord may be palpated at the pubic tubercle as it passes over the pubis and the lidocaine injected at this landmark. Lee and colleagues were able to perform
Manual Detorsion

The goal of manual detorsion is to reestablish or increase blood flow to a previously ischemic testis. This should be done in conjunction with preparation of the operating suite. It should never delay operative intervention.

Before initiating detorsion, ensure that the patient is as comfortable as possible in a reclining or supine position. Lithotomy position gives the examiner the most access to the patient’s genitalia and prevents the patient from retreating during the procedure. If light analgesia and/or sedation are selected, implement them at this time.

Manual detorsion begins with the clinician standing comfortably at the side of the bed or stretcher, preferably on the patient’s right side if the clinician is right-handed, or vice versa. Detorsion is begun just as one would open a book (i.e., an initial 180 degree detorsion of the patient’s right testis is done in a counterclockwise fashion) (see Fig. 55.1C and D). The patient’s left testis is detorsed 180 degrees in a clockwise fashion. Pain relief is an objective end point. If one rotation relieves some but not all the pain, continue with another rotation. The degree of torsion may range from 180 to 1080 degrees with medians of 360 to 540 degrees. Therefore many patients require two or three rotations to alleviate pain. If the initial detorsion is mechanically difficult (which it will be if one is detorsing in the wrong direction) or makes the pain worse, detorse the testis in the opposite direction and observe the result. Approximately one third of testicular torsions occur in the lateral, or unexpected, direction. The objective success of any testicular manipulation can be substantiated by an increase in Doppler signal and the patient’s relief of pain.

With successful detorsion, the testicle returns to its normal anatomic position. Resolution of induration and swelling of the spermatic cord, testis, and epididymis will depend on the degree and duration of ischemia. Thus, the more severe the torsion and the longer it has been present, the longer it will take for the edema and induration to resolve. With significant ischemia, the entire epididymis often becomes enlarged like a link sausage (uncommon in epididymitis except in severe cases or those that are initially misdiagnosed or seen late in their clinical course), and the testis becomes quite firm, simulating a testicular tumor. In some authors’ experience, these reversible changes usually resolve over 3 to 4 hours.

Aftercare

Provide systemic analgesics as needed for discomfort. Importantly, even though manual detorsion will save an ischemic testicle, it should not be substituted for definitive scrotal exploration.

Complications

In a portion of cases, the testis will torsed in the opposite direction (medial to lateral) or have multiple twists. This may become apparent as the clinician assesses the results of the detorsion procedure by palpation, relief of edema, and return or increase of the Doppler signal.

Conclusions

Spermatic cord torsion may be relieved by manual detorsion. In two thirds of cases, this is accomplished by medial to lateral rotation of the affected testicle. However, in the remaining one third of cases, lateral to medial rotation is necessary to untwist the spermatic cord. In all cases, manual detorsion, even when successful, only serves as a temporizing bridge to definitive surgical management.

PRIAPISM

Introduction

Priapism is defined as prolonged erection of the penis, generally lasting over 4 hours, in the absence of sexual desire or stimulation (Fig. 55.3). This medical condition was named after Priapus, an ancient Greek god of fertility and horticulture who was endowed with oversized genitalia.

Priapism can be divided into two main categories. Ischemic priapism, also known as low-flow priapism, is the most commonly seen variant and is due to painful venous engorgement of the corpora cavernosa and requires emergency treatment. Non-ischemic (high-flow) priapism is quite rare and is often painless. It is caused by increased arterial inflow to the penis as a result of traumatic arteriocavernosal fistulas and does not require urgent treatment.

Ischemic priapism can be thought of as a compartment syndrome of the penis. The corpora cavernosa becomes engorged with stagnant, oxygen-depleted venous blood due to either an intraluminal obstruction of venous blood flow or an inability of the penile muscle tissue to adequately contract and augment venous outflow.

Procedure Overview

Indications: failure of an initial trial of subcutaneous or oral terbutaline (optional).

Contraindications: high-flow (nonischemic) priapism, stuttering priapism.

Complications: hematoma formation, infection, systemic absorption, and subsequent effects of α-adrenergic agents; permanent impotence may result from priapism regardless of promptness of therapeutic interventions.

Equipment: see later.

Figure 55.3 Typical appearance of priapism. It has many causes, is painful, often recurrent, and can lead to impotence. Many cases can be effectively treated in the emergency department (ED), but definitive ED intervention is not a mandated standard of care.
Key Procedure Sequence

1. Administer parenteral opioid analgesic or light sedation.
2. Give a trial of 0.25 to 0.5 mg subcutaneous terbutaline (repeat in 15 minutes), or 5 mg terbutaline by mouth (optional).
3. Give an intracorporal injection of α-adrenergic agents and, if necessary, repeat every 20 minutes for a total of three doses (minimally invasive approach).
4. If unsuccessful, aspirate 20 to 30 mL of corporal blood.
5. If detumescence is not achieved, irrigate with saline or a dilute α-adrenergic agonist solution (aspirate-irrigate-aspirate cycle, as needed).
6. A visible change from venous (dark red) to arterial (bright red) blood is a marker of success.
7. For persistent erections, consult urology for possible shunt placement (cavernosum-spongiosum shunt).
8. If the procedure is successful, loosely wrap the penis in an elastic bandage.
9. A 3-day course of an oral α-adrenergic agent is reasonable at the time of discharge.
10. Arrange for urgent urology follow-up and administer strict return precautions.

Description

Background

Priapism is manifested by a persistent, usually painful, penile erection, unrelated to sexual stimulation and not relieved by ejaculation.

Over one third of patients with severe priapism may suffer permanent erectile dysfunction despite treatment, with obvious functional and emotional sequelae.15

Anatomy and Physiology

Priapism is characterized clinically by a soft glans penis and spongy urethra in the presence of two erect penile bodies (corpora cavernosa) (Fig. 55.4). Two important concepts are worthy of mention. First, there is communication of blood flow between the corpora cavernosa, therefore in most cases the operator needs to access only one of the corpora. Secondly, introduction of vasoactive or other agents into the corpora is akin to an intravenous injection, so systemic effects may be precipitated, particularly after partial or full detumescence is achieved.

Pathophysiology

The pathophysiology of priapism is complex. The pharmacologic basis for treatment is based on manipulating blood flow via the α- and β-adrenergic receptors. Priapism is believed to result from increased arterial inflow of blood into the corpora cavernosa secondary to dilatation of the cavernosal arteries. Relaxation of the cavernosal tissue occurs and secondary compression of the emissary veins leads to engorgement of both corpora cavernosa during an erection. When the cavernosal pressure approaches the arterial pressure, blood flow is markedly reduced. Ischemic or low-flow priapism results after several hours of continuous painful erection, leading to intracavernosal acidosis and sludging of blood, with subsequent thrombosis of the cavernosal arteries, fibrosis of the corporal tissue, and irreversible impotence.

High-flow priapism is less common than low-flow priapism and usually results from traumatic production of arteriocavernosal fistulas. It is not associated with intracavernosal ischemia or acidosis and is therefore painless and may be treated electively rather than emergently.

In the past, priapism was most often encountered as a complication of a number of medical (e.g., hematologic, neoplastic, and drug-related) conditions (Table 55.1). Today, many cases are iatrogenic, resulting from the current practice of using vasoactive substances (e.g., papaverine and phentolamine) and other newer erectile dysfunction medications to induce penile erections in impotent men. Sickle cell disease continues to be a leading cause of priapism.15 Sickle cell patients may experience such a high rate of recurrence that home self-injection of vasoactive drugs into the penis has been advocated. Cocaine use is one etiology that is likely under-reported. A drug screen may unravel some discrepancies between clinical findings and history. As an end result, vasoactive drugs promote engorgement of the corpora cavernosa and reduction in venous outflow, which may result in low-flow or ischemic priapism. Several phosphodiesterase inhibitors and prostaglandin E1 are the drug treatments for impotence approved by the U.S. Food and Drug Administration (FDA). These medications act by increasing penile blood flow and enhancing smooth muscle relaxation. The incidence of priapism with these medications is quite low, particularly with the phosphodiesterase inhibitors. Penile rigidity due to a non-deflating penile prosthesis (pseudopriapism) or malignant replacement of the corpora in patients with bladder or prostate cancer should not be confused with true priapism.

Indications

The emergency clinician should attempt to identify reversible causes for low-flow priapism and, often in conjunction with a urologic surgeon, initiate specific corrective therapy as soon as possible. Low-flow priapism in children and young adults may be due to sickle cell disease and such cases may respond to noninvasive standard anti-sickling measures. However, the role of transfusion therapy in patients with priapism due to sickle cell anemia is uncertain.
CHAPTER 55  Urologic Procedures

subcutaneous) terbutaline, is the same regardless of the inciting etiology, although its utility and efficacy are debated.40–42 It is thought that terbutaline, a β2-adrenergic agonist, increases venous outflow from the engorged corpora by way of relaxation of venous sinusoidal smooth muscle. Terbutaline is of unproved benefit; however, given its limited propensity for adverse effects, a trial is reasonable in select circumstances while awaiting specialty consultation.43

If terbutaline fails to work rapidly, the next step is penile blood aspiration, saline irrigation of the corpora cavernosum, and intracorporal instillation of an α-adrenergic receptor agonist such as dilute phenylephrine or epinephrine.

Contraindications
A subtype of ischemic priapism is known as stuttering priapism. This entity is typically observed in patients with sickle cell disease. Patients experience recurrent episodes of priapism that often last less than 3 hours and often do not require emergency treatment unless symptoms become markedly prolonged.44 High-flow (nonischemic) priapism is treated surgically (elective repair).

Procedure
A suggested algorithm for the initial treatment of acute nonischemic priapism in the emergency setting is presented in Box 55.1. Options for treatment include:
1. Minimally invasive technique (direct intracorporal injection of α-adrenergic agent, without aspiration or irrigation).
Dilute with saline; final volume, 1 mL.

Dilute with 0.9 mL saline; final volume, 1 mL.

The same procedure may be attempted as an initial approach. This minimally invasive procedure may be performed, less traumatic, and less invasive than aspiration and irrigation. This minimally invasive procedure may be attempted as an initial approach. This same procedure may be used as a self-injection technique for home treatment of priapism.

Aspiration/Irrigation Technique

Minimally Invasive Technique: Simple Injection

Relief of priapism by simple injection of vasoactive solutions into the corpus cavernosum has been reported.44 Inter cavernous injection therapy for the management of priapism is simple to perform, less traumatic, and less invasive than aspiration and irrigation. This minimally invasive procedure may be attempted as an initial approach. This same procedure may be used as a self-injection technique for home treatment of recurrent priapism. With this technique, a 25- to 27-gauge needle is used to inject vasoactive substances into the corpus at the proximal end of the penis (2–4 cm distal to the shaft origin), with the goal of pharmacologically reversing priapism (Fig. 55.5). Often, this can be accomplished without anesthesia. One option is to draw up 0.5 mg (0.5 mL) of phenylephrine and add 0.5 mL of saline diluent for a final volume of 1 mL. (Table 55.2). Puncture the corpus with the needle at the 10 o’clock or 2 o’clock position at the base of the penis (with 12 o’clock being the dorsal vein of the penis), aspirate blood to confirm position, and inject the solution. If detumescence is not achieved in 20 to 30 minutes, give a repeat injection, up to a total of three injections. In one small study, successful detumescence was achieved in eight of nine patients by simple intracorporal injection of phenylephrine in this regimen, with three or fewer injections required.46 Alternatively, 0.1 mg of epinephrine (0.1 mL of 1:1000) diluted with 0.9 mL of saline may be used (1 mL total volume). Regardless of medication used, only one side needs to be injected, but two or three injections might be necessary. Wait at least 20 minutes after each injection before additional interventions. Note that this is essentially an intravenous injection and systemic effects may occur. As such, proceed with caution in patients with cardiovascular disease. Success has also been noted by injecting the corpus cavernosum with 1 mL of the local anesthetic lidocaine (2%) with epinephrine (1:100,000) into each side, or 2 mL into one side.47

Aspiration/Irrigation Technique

Box 55.2 lists the equipment needed for aspiration and irrigation of the corpus cavernosum. This procedure entails drainage of blood from the erect penis, irrigation with saline if necessary (e.g., inadequate return of blood with lack of detumescence), and finally the instillation of a vasoactive medication. Alternatively, irrigation with aliquots of a dilute vasoactive solution may be effective (aspirate-infuse-aspirate cycle as needed).

Place the patient in the supine position. Parenteral analgesia and sedation are suggested. Local anesthesia at the puncture site is recommended. An injection of 1% plain lidocaine placed at the base of the penis for a dorsal penile nerve block or placement of a circumferential penile block can be performed, but are usually not necessary (see Fig. 55.5B). Prepare and drape the penis in sterile fashion. Grasp the shaft of the penis with your nondominant thumb and index finger. Palpate the engorged corpus cavernosum laterally (2 and 10 o’clock positions) and insert a 21- to 19-gauge butterfly needle into the corpus cavernosum. Alternatively, a 20-gauge intravenous catheter may be used, but the butterfly needle is preferred. A dialysis access butterfly needle may also be used (Fig. 55.6). If palpation fails to demonstrate the corpus, blindly inserting the needle at either
Management of Acute Priapism

A. Minimally Invasive Method

• Anesthesia is not required. Use a small-gauge needle to inject a small aliquot of an adrenergic agent into the corpus at 10 or 2 o’clock at the proximal end of the penile shaft. (Only one side need be injected.)
• Aspirate blood (2) to confirm placement prior to injection.
• Consider a repeated injection after 20 minutes if unsuccessful, but be wary of the systemic effects of adrenergic agents (especially if repeated doses or injections are given during partial detumescence).

Adrenergic Agents
• 0.2 to 0.5 mg of phenylephrine per dose (refer to text for instructions on dilution)
• 0.1 mg of epinephrine (0.1 mL of a 1:1000 solution) mixed with 0.9 mL of saline

(Use a 1-mL tuberculin syringe to draw up these small volumes.)

B. Aspiration and Irrigation Method

This patient experienced 18 hours of priapism after penile self-injection of papaverine as therapy for impotence.

For corpus irrigation the irrigation needle can be placed through a simple skin wheal, or perform a penile dorsal nerve block by injecting 1% plain lidocaine at the base of the dorsal aspect of the penis. The dorsal nerves are relatively superficial and deep injections are not required.

Insert no smaller than a 19-gauge butterfly needle into the corpus via the proximal penile shaft at either the 2 o’clock or the 10 o’clock position and aspirate. Only one side need be punctured. Slow steady suction will be most successful, whereas excessive suction may halt the aspiration. Do not puncture the corpus via the glans (see text). After initial aspiration, irrigate (slowly inject and withdraw) 10- to 20-mL aliquots of vasoactive solution until detumescence persists.

After detumescence with the first aspiration or with aspiration-irrigation-aspiration of a vasoactive medication (see text), wrap the penis with an elastic bandage to discourage reengorgement and to compress the puncture site. Note: Acceptable procedures include aspiration alone followed by instillation of a small aliquot of epinephrine (0.1 mg) and combined multiple aspirations and irrigations with a vasoactive solution. The end point is the appearance of bright red arterial blood and/or persistent detumescence.

Figure 55.5 Management of acute priapism. The minimally invasive technique (A) consists of directly injecting the corpus with a small aliquot of an adrenergic agent. This is often successful and can be used at home by a motivated patient with recurrent problems (such as sickle cell disease). It may require multiple injections and 15 to 20 minutes to be effective.
If detumescence is not achieved following aspiration (with or without saline irrigation), irrigation with a dilute vasoactive substance is another option. A number of dilute irrigation solutions have been suggested, but none have been proven to be superior. Some suggest 20 to 30 mL of a phenylephrine/normal saline solution (10 mg of phenylephrine in 500 mL of normal saline) as the exchange for 20 to 30 mL of aspirated corporal blood. Some clinicians add 2500 to 5000 units of heparin to the solution, but the use of heparin is of unproven value. Alternatively, 1 mg of epinephrine can be added to 1 L of saline, with irrigation performed using 20- to 30-mL aliquots. It is important to note that corporal irrigation is performed with a much less concentrated solution (phenylephrine, 20 µg/mL) as compared with that used for the minimally invasive technique or vasoactive instillation of 1 mL aliquots following detumescence detailed earlier (phenylephrine, 500 µg/mL).

Note that the corpus cavernosum has ready access to systemic circulation, and injecting a drug into it is essentially the same as an intravenous injection. When detumescence occurs, the unmetabolized drug enters the systemic circulation; therefore vasoactive drug dosages should be monitored carefully.

**Aftercare**

Observe the patient in the ED for recurrence. Although the ideal observation period is unknown, 2 hours has been suggested. Fig. 55.5B, step 4 demonstrates the entire penis loosely wrapped with elastic (ACE, 3M) bandage to prevent hematoma formation at the injection site(s). Strict return precautions must be provided for priapism recurrence, and urgent urology follow-up should be ensured. A short course of an oral α-adrenergic agent, such as pseudoephedrine for 3 days, is often recommended. However, the value of this intervention is of questionable value and unproven benefit.

**Complications**

Although hematoma formation or infection can occur with properly performed aspiration, these complications are infrequent. Injected or instilled vasoactive agents can be absorbed systemically, with the potential for toxic effects. Therefore the intracavernosal use of vasoactive agents is relatively contraindicated in patients with conditions sensitive to these agents (e.g., severe hypertension, dysrhythmias, monoamine oxidase inhibitor use, etc.). Blood pressure and cardiac rhythm should be monitored throughout the procedure if the patient is at risk. Failure to aspirate blood is a potential complication, usually because of a misplaced needle, applying excessive suction, or if blood has clotted. Because impotence is a well-recognized complication of priapism regardless of the cause or the promptness of therapeutic intervention, the patient should be advised regarding this potential long-term consequence.

**Conclusions**

Because prolonged priapism increases the risk of subsequent erectile dysfunction, an aggressive management strategy is advised. After 4 hours of persistent priapism, there is heightened release of inflammatory cytokines in the acidic and hypoxic corpora cavernosa. Inflammation may include smooth muscle changes including cell death and fibrosis, which may cause permanent erectile dysfunction. Recurrence is not uncommon, and some patients require multiple procedures on a recurring basis. Surgical shunting procedures by the urology team might be required if these other measures are met without success.

---

**Figure 55.6** An 18-gauge dialysis or butterfly catheter can be used for irrigation of the corpora if simple aspiration or injections of medication are unsuccessful. Using needles smaller than 19-gauge is usually counterproductive.

10 o’clock or 2 o’clock will usually gain access to this large vascular structure. Because there is communication of blood flow between both sides, the operator needs access to only one of the corpora. Either side may be punctured. The site of needle placement is typically anywhere from the base to the proximal shaft, 2 to 4 cm distal from the shaft origin. The glans should not be used as a puncture site.

The needle is advanced at a 45-degree angle, using constant suction. Blood is usually readily aspirated; once it is obtained do not advance the needle further. Next, stabilize the needle. Avoid deep penetration to minimize the risk of injury to the cavernosal artery during this procedure.

Aspirate an initial 20 to 30 mL of corporal blood while milking the corpus with the nondominant hand. Do not apply excessive suction because this often halts the aspiration. A common error is to use too much suction with a 60-mL syringe. Using a 10-mL syringe, and changing it if it fills with blood, is preferable. Use of a butterfly reduces the danger of dislodging the needle when changing syringes.

Continue aspiration until the original egress of dark blood ceases and bright red arterial blood returns or when complete detumescence is obtained and persists. Because multiple anastomoses exist between the two corpora cavernosa, bilateral aspiration is not required. If there is inadequate return of blood with a lack of detumescence, consider irrigating with 20 to 30 mL of saline.

If detumescence is achieved after initial aspiration with or without saline irrigation, no further treatment may be required. If this is successful, some clinicians advise instilling an aliquot of vasoactive substance. Phenylephrine is recommended as the agent of choice as it may minimize the risk of cardiovascular side effects more commonly seen with other sympathomimetic agents. Inject 0.1 to 0.5 mg of phenylephrine (in 1 mL of normal saline). It is important to note that the recommended concentration for instillation is similar to that recommended for the minimally invasive technique detailed earlier. Lower concentrations in smaller volumes should be used for patients with cardiovascular risk factors or for children. If irrigation is performed with a dilute vasoactive substance as delineated hereafter, additional medication instillation is not suggested.
or balanoposthitis are also associated with development of paraphimosis. Inflammation can result in contracture of the distal foreskin. Later when the foreskin is retracted proximally over the compressible glans, the contracted foreskin forms a constrictive band and gets stuck in the retracted position.

**Procedure Overview**


Equipment: for simple manual reduction, all that is needed is a topical anesthetic lubricant. Adjunctive equipment for alternative reduction techniques and maneuvers as described in this chapter are optional.

---

**Paraphimosis**

**Introduction**

Paraphimosis is the inability to completely reduce the penile foreskin distally, back to its natural position overlying the glans penis. This condition occurs exclusively in uncircumcised males and is a urologic emergency.

Paraphimosis may occur at any age, but often occurs in the extremes of life (Fig. 55.7). The condition can be quite subtle and may be either unrecognized or misdiagnosed as an allergic reaction, penile trauma, infection, or edema resulting from systemic volume overload (e.g., congestive heart failure, nephrotic syndrome) (Fig. 55.8).

Iatrogenic paraphimosis may occur following urinary catheterization or medical examination if the foreskin is not returned to its native location overlying the glans. Poor hygiene

---

**Figure 55.7** Paraphimosis often occurs in the extremes of life. **A,** Uncircumcised boy with paraphimosis. This may be mistaken for infection or localized trauma, especially if it is unclear whether circumcision has been performed. Always seek this history. **B,** Paraphimosis, pictured here, may be mistaken for penile trauma, angioedema, or infection. Note that the swollen tissue is proximal to the coronal sulcus (asterisk). The cause of paraphimosis in this case was failure to replace the foreskin after a catheter change in an uncircumcised nursing home patient. Before reduction, the catheter is usually removed. When the edema is minimal, the catheter may be left in place during reduction. **C,** Appearance of the penis after catheter removal and foreskin reduction.

**Figure 55.8** **A,** This edematous foreskin may have numerous causes, from benign edema to paraphimosis, and at first glance the cause might not be obvious. This patient has paraphimosis that is appreciated by careful inspection to identify **B,** the normal coronal sulcus and **C,** a phimotic foreskin band proximal to the glans.
Key Procedure Sequence

1. Apply a topical anesthetic lubricant to the distal penis (foreskin and glans).
2. Administer a systemic analgesic (optional).
3. Compress the foreskin and glans with a snugly grasped hand.
4. Use the thumbs to invert the glans penis proximally through the phimotic ring.
5. Use the index and long fingers to reduce the phimotic ring distally.
6. If unsuccessful, try alternate maneuvers to reduce edema before reattempting manual reduction.
7. Consider an optional trial of glans adjuncts: administer an ice pack or a compressive bandage to reduce glans edema.
8. Consider an optional trial of foreskin adjuncts: administer micropuncture wounds, use sugar, or use hyaluronidase to reduce foreskin edema.
9. If unsuccessful, reduce the paraphimosis surgically by dorsal slit or definitive circumcision.
10. Arrange urgent urology follow-up following reduction.

Indications

Emergency reduction is indicated whenever a paraphimosis exists.

Contraindications

There are no contraindications.

Procedure

Manual Reduction Technique

The current standard for reducing paraphimosis is manual reduction. This can be accomplished by using a nonirritating topical anesthetic lubricant applied to the inner surface of the foreskin (not to the shaft of the penis) and the glans to reduce friction and decrease the discomfort of the procedure. A penile block may be performed if necessary (see Fig. 55.9, steps 2 and 3). If there is significant discomfort or patient apprehension, systemic analgesia or procedural sedation may be useful adjuncts. For young children, general anesthesia may be necessary.

Compress the foreskin and glans with a snugly grasped hand for several minutes to reduce as much edema fluid as possible (see Fig. 55.9, steps 4–6). Use the thumb to push the glans through the foreskin that has been incircled by the entire palm (see Fig. 55.9, step 7). Alternatively, place the index and long fingers of both hands in apposition just proximal to the phimotic ring. Align both thumbs on the urethral meatus. Apply constant force with the thumbs to try and invert the glans penis proximally while using the index and long fingers to reduce the phimotic ring distally over the glans penis and into its normal anatomic position (see Fig. 55.9, step 8). Successful reduction results in the appearance of an uncircumcised penis with a phimotic foreskin (see Fig. 55.9, steps 9 and 10). The key to success in both these maneuvers is the application of slow, steady pressure.

Adjunctive Techniques to Assist in Manual Reduction

Several alternative methods for reducing the edema have been described in the literature. These methods can be used before attempting manual reduction, or if simple manual reduction fails.

Babcock clamps (noncrushing tissue clamps) can be used to reduce the paraphimotic foreskin (Fig. 55.10). Apply six to eight Babcock clamps spaced evenly around the foreskin, straddling the phimotic ring (one edge just proximal and the other edge just distal to the phimotic ring). Grasp all clamps and apply simultaneous distal traction to pull the phimotic ring over the glans. After reduction, remove the clamps. It is important to inspect the foreskin for injuries.

Other techniques focus on reduction of glans or foreskin edema (or both), followed by reduction of the paraphimosis. Diminished glans edema facilitates foreskin reduction to its natural position. One suggested maneuver is to apply an ice pack. In the “iced-glove” method, use cold compression to reduce foreskin swelling and induce vasoconstriction in the glans penis (see Fig. 55.10). Half fill a large glove with crushed ice and water, and tie the cuff end securely. Invaginate the thumb of the glove and then draw it over the lubricated paraphimotic penis. Hold the thumb of the glove securely in place over the glans for 5 to 10 minutes. The combination of cooling and compression usually decreases the edema sufficiently to permit manual reduction of the foreskin. Wrapping the penis in a compressive bandage is another alternative. A 2-inch compressive bandage (such as an ACE [3M] bandage or similar)
**Paraphimosis Reduction**

This patient had asymmetric paraphimosis. Note that the glans is edematous, as is the foreskin. A circumferential constricting band is found proximal to the glans and foreskin.

Complete a ring block of the penis by depositing anesthetic circumferentially around the proximal part of the shaft.

Alternatively, wrap an elastic bandage around the distal end of the penis for several minutes. This step is often omitted, which makes the procedure unnecessarily difficult.

Grasp the shaft of the penis with one hand and apply force onto the urethral meatus with the thumb of your other hand. Push the glans forward while sliding the foreskin distally.

The key to success is the application of slow, steady pressure. Note that the glans has retracted into its normal position and only the edematous foreskin is visible.

Perform a penile block in cases that will require significant manipulation. First, block the dorsal penile nerves by depositing anesthetic at the 2 and 10 o’clock positions on the dorsal aspect of the shaft of the penis.

Compress the foreskin and glans by grasping it with the palm of your hand and applying pressure for several minutes.

Note that manual compression substantially reduced the amount of swelling (compare with step 1; the foreskin and glans were taut prior to compression).

Alternatively, grasp the penis with the fingers of both hands just proximal to the phimotic ring, and use your thumbs to apply constant pressure on the glans.

Successful reduction results in the appearance of an uncircumcised penis with a phimotic foreskin.

*Figure 55.9* Paraphimosis reduction. (Step 8, from Neuwirth H, Frasier B, Cochran ST: Genitourinary imaging and procedures by the emergency clinician, *Emerg Med Clin North Am* 7:1, 1989.)
Paraphimosis Reduction: Alternative Techniques

A. Babcock Clamps

1. Babcock clamps will not crush tissue. Place six to eight clamps so that they are evenly spaced on the distal edge of the foreskin and straddling the phimotic ring.
2. Grasp all clamps and apply simultaneous distal traction to pull the phimotic ring over the glans.

B. Iced-Glove Method

1. Fill a large glove halfway with crushed ice and water.
2. Securely tie the cuff end and invaginate the thumb of the glove.
3. Draw the invagination over the lubricated paraphimotic penis and hold it in place for 5 to 10 minutes. The combination of cooling and compression decreases edema and facilitates foreskin reduction.

Figure 55.10 Paraphimosis reduction: alternative techniques.

Alternatively, techniques focusing on reduction of foreskin edema have been advocated (Box 55.3). The Dundee technique involves creating multiple micropunctures of the edematous foreskin and then expressing edema fluid.55 Hyaluronidase has been reported to result in rapid reduction of prepuce edema that facilitates manual reduction of the foreskin. This enzyme, when injected into the swollen retracted foreskin, causes hydrolysis of hyaluronic acid that, in turn, increases tissue permeability so the edema in the foreskin is diffused out into the surrounding penile tissue. There have even been advocates of a noninvasive way to reduce the foreskin edema via the application of granulated sugar to the penis. Sugar forms an osmotic gradient that draws out edema fluid, but this may take several hours.57 Publications on these alternative procedures are generally observational in design with very small numbers. To date, there have not been any large studies of comparative effectiveness; as such it is difficult to recommend any one method as superior.58

If the previous methods are unsuccessful, it may be necessary to incise the constricting phimotic tissue to permit reduction
of the foreskin over the glans by dorsal slit procedure. In adults, this can generally be carried out under penile-block anesthesia, but procedural sedation or general anesthesia may be necessary for young children.

**Aftercare**
Observe the patient to ensure adequate local hemostasis, ability to void spontaneously, and recovery from analgesia or sedation, if utilized. Replacing the foreskin to its native position following examination, catheter placement, sexual activity, or any other manipulation is essential to preventing recurrent episodes. Patients should be referred to a urologist for evaluation of possible surgical options, including circumcision.

**Complications**
Penile shaft laceration or simple tearing of edematous and taut penile skin may occur during manual or surgical paraphimotic reduction. Simple suturing is all that is required for most injuries. If reduction of a paraphimosis cannot be achieved by other means, surgical interventions should be considered.

**Conclusions**
Emergency manual or surgical reduction of the edematous foreskin is mandatory to restore proper circulation, relieve discomfort, and permit resolution of potential serious sequelae: skin ulceration and gangrene. It must be done as soon as a paraphimosis is recognized. Once the foreskin is successfully reduced, urgent referral for dorsal slit or definitive circumcision is necessary.

**PHIMOSIS**

**Introduction**
Phimosis is constriction of the foreskin (prepuce) that limits its retraction proximally over the glans (Fig. 55.11). Uncircumcised infants and young children often have a physiologic phimosis due to adhesions between the prepuce and glans. This is in contrast to a pathologic phimosis, where failure to retract results from distal scarring of the prepuce. Circumcision (removal of the foreskin), rendering phimosis and paraphimosis anatomically impossible, is commonplace in America. However, circumcision rates vary with religious affiliation, racial and ethnic group, as well as socioeconomic status.

Phimosis does not ordinarily require any treatment. However, a patient with phimosis may present acutely to the ED when he is unable to void spontaneously as a result of distal urethral obstruction. It may prevent or make urethral catheterization more challenging. In such situations, the phimotic opening may need to be dilated. An alternative strategy is to crush a portion of the foreskin followed by an incision (dorsal slit), using local anesthesia with or without parenteral analgesia or sedation to allow access to the urethral meatus. This procedure can be readily performed in the acute care setting when necessary.

**Procedure Overview**
**Indications:**
1. Phimosis and the inability to void or perform urethral catheterization;
2. As definitive treatment following successful foreskin reduction in a patient with paraphimosis; or

**Contraindications:** none.

**Complications:** injury to the urethral meatus or glans penis, bleeding at site of tissue injury (when tissue is crushed and then cut).

**Equipment:** see later.

**Key Procedure Sequence**
1. Clean and drape the penis with sterile towels.
2. Inject local anesthesia: lidocaine without epinephrine.
3. Use toothed forceps to test for adequate local anesthesia.
4. If anesthesia is inadequate, consider dorsal nerve block or “ring block” at the base of the penis.
5. Advance both jaws of a hemostat proximally between the inner layer of the foreskin and glans.
6. Use a hemostat to separate preputial adhesions.
7. Remove the hemostat, reinsert one jaw, and use the hemostat to crush interposed foreskin tissue.
8. Cut the resultant crushed foreskin tissue longitudinally with straight scissors.
9. Use absorbable sutures to reapproximate leaves of foreskin resulting from the cut (if necessary).
10. With success, the prepuce is easily retracted for exposure of the glans penis and urethral meatus.

**Description**

**Background**
Phimosis is constriction of the foreskin that limits retraction of the foreskin over the glans. Phimosis does not ordinarily require any treatment. However, phimosis may present acutely to the ED when a patient is unable to void spontaneously as a result of distal urethral obstruction.

**Anatomy and Physiology**
A physiologic phimosis in young uncircumcised boys consists of a pliant, unscarred preputial orifice on physical examination. The foreskin gradually becomes retractile over time as a result of intermittent erections and keratinization of the
inner epithelium. By 3 years of age, the foreskin can easily be retracted in 90% of patients, with nearly all becoming retractile by late adolescence.60,61

Pathophysiology
A pathologic phimosis exists when the failure to retract results from distal scarring of the prepuce. This is typically a subacute condition that may present acutely to the ED when a patient is unable to void spontaneously as a result of distal urethral obstruction. Pathologic phimosis is caused by local trauma, infection, chemical irritation, complications of circumcision (insufficient tissue removal), or poor hygiene.

Patients with acute phimosis complain of penile pain over hours to days. On examination, the physician will discover a tender foreskin that is not easily retracted. Although the prepuce may be gently manipulated to allow a better exam, do not attempt forced retraction. Forceful retraction contributes to future adhesion and stricture formation.

Indications
Dorsal slit of the foreskin is performed in any emergency situation to gain access to the urethral meatus for urethral catheterization. In the setting of paraphimosis, a dorsal slit may be used as a definitive treatment following simple foreskin reduction, or for phimotic ring incision and reduction in a patient with an otherwise irreducible paraphimosis. Elective circumcision rather than dorsal slit of the foreskin is the definitive procedure of choice in nonemergency situations.

Contraindications
None.

Procedure
Box 55.4 lists the equipment needed to perform dorsal slit of the foreskin. With the patient in the supine position, clean and drape the penis with sterile towels. Clipping of the pubic hair is unnecessary. Infiltrate lidocaine without epinephrine into the dorsal midline of the foreskin just beneath the superficial fascia throughout the course of the proposed incision, starting proximally at the level of the coronal sulcus and proceeding distally to the tip of the foreskin (Fig. 55.12, step 1). Consider mixing equal volumes of lidocaine with bupivicaine. After several minutes, grasp the foreskin with toothed forceps to test for anesthesia. The operator must be certain that the inner surface of the foreskin is also anesthetized. If this area is not numb, use a dorsal nerve block or ring block at the base of the penis (Fig. 55.13).62 Systemic analgesia or procedural sedation may be useful adjuncts as well.

After achieving adequate local anesthesia, systemic analgesia, or procedural sedation if needed, take a straight hemostat and carefully advance both jaws of the hemostat proximally to the area of the coronal sulcus between the inner layer of the foreskin and the smooth glans penis, carefully separating any existing preputial adhesions (see Fig. 55.12, step 2). Take care that the meatus and urethra are visualized or palpated at all times to avoid inadvertent injury during this maneuver. Once release of adhesions is complete, open the hemostat, and place one jaw of the hemostat in the recently developed plane between the glans penis, opened to tent the skin to ensure proper placement, and the superior overlying inner layer of foreskin. Advance the hemostat to the level of the coronal sulcus and then close it, effectively crushing the interposed anesthetized foreskin. Leave the closed hemostat in place for 3 to 5 minutes, then remove it and cut the resultant serrated crushed foreskin longitudinally with straight scissors throughout the extent of the crushed tissue.

Normally, the incised, anatomic approximated skin edges bleed and ooze. Not infrequently, these skin edges of the foreskin separate into two layers, the outer foreskin and the inner foreskin (see Fig. 55.12, step 3). Absorbable chromic or Vicryl (Ethicon Inc.) (4-0 to 5-0 for children, 3-0 to 4-0 for adults) running hemostatic sutures may be placed, each beginning proximally at the apex of the dorsal slit and carried distally, reapproximating the two leaves of foreskin. Standard antibiotic ointment may be used to lubricate the suture material and facilitate passage of the suture through the delicate foreskin tissue.

After successful dorsal slit of the foreskin, the prepuce is easily retracted for cleansing of the glans penis or exposure of the urethral meatus. Postprocedural conscientious foreskin reduction to its normal anatomic position must be ensured after any distal penile manipulation to avoid paraphimosis.

If reduction of a paraphimosis cannot be achieved with other means, then surgical interventions should be considered. Fig. 55.14 describes the dorsal slit procedure for paraphimosis reduction, which involves carefully sterilizing the field, anesthetizing the penis on the dorsal aspect, followed by a linear incision.

Aftercare
Ideally, a definitive elective circumcision is recommended after a dorsal slit. Some patients complain about the appearance of their incised foreskin (“beagle ears”) and the relative inconvenience during urination, whereas others are pleased they no longer have their phimosis and decline further treatment (see Fig. 55.12, step 4).

Complications
Injury to the urethral meatus, urethra, and the glans penis may occur if the hemostat or straight scissors are blindly and unknowingly introduced into the urethra. Bleeding may occur if the hemostat has not adequately crushed the foreskin or if the scissors incision is made lateral to the serrated crushed tissue. The latter two problems are easily resolved with the previously described running hemostatic suture.

Conclusions
Dorsal slit of the foreskin is performed in any emergency situation either to gain access to the urethral meatus for urethral catheterization or as definitive treatment after simple foreskin reduction, or phimotic ring incision and foreskin reduction in a patient with an otherwise irreducible paraphimosis. This minor operative procedure can be readily performed in the acute care setting when necessary.

BOX 55.4 Equipment Necessary for the Dorsal Slit Procedure

| 1% lidocaine without epinephrine |
| 5-mL syringe |
| 27-gauge needle |
| 1 straight Crile clamp |
| 1 straight scissors |
| 1 needle holder |
| 4-0 absorbable suture |
or uneventful depending on the operator's expertise, confidence, and gentleness.

The merits of various approaches to urine specimen collection depend on the patient's age and clinical setting.

Urethral catheterization is definitive and routinely used to collect urine for analysis and culture in infants and young children who are not yet potty-trained. For older children
who can follow direction, midstream urine collection is a reasonable alternative approach to specimen collection.

In adult men without anatomic lesions, first-voided midstream specimens can define the presence or absence of culture-proven bacteriuria. This certainly represents a simple, noninvasive and user-friendly approach to urine collection in the busy ED setting.

In adult women, properly collected clean-catch midstream specimens have been found to be as bacteriologically reliable as catheterized specimens. A few caveats are worth mentioning. Ideally, patients must sit backward on the toilet when collecting the specimen (i.e., patient facing the wall behind the toilet, which theoretically promotes labial spreading). Of more concern is the fact that such studies excluded patients with vaginitis, urologic abnormalities, pregnancy, and vaginal bleeding. These diagnoses represent the clinical circumstances for which urine is commonly examined in young women visiting the ED. In this at-risk population, catheterized urine specimens are optimal. However, this needs to be balanced with the reality that catheterization may introduce unnecessary patient discomfort and resource utilization, as well as the risk of introducing bacteria into the bladder.

**Figure 55.13** Regional anesthesia of the penis. Consider the use of supplemental intravenous analgesia or procedural sedation, or both, based on the clinical scenario. (A1, From Soliman MG, Tremblay NA: Nerve block of the penis for postoperative pain relief in children, *Anesth Analg* 57:495, 1978. Reproduced by permission.)
Dorsal Slit (Paraphimosis Treatment)

1. Infiltrate local anesthetic from the constricting band of the paraphimosis proximally along the dorsal aspect of the penis. Make an incision along this line.

2. A diamond-shaped defect will result from the linear incision of the foreskin. Approximate the two apices of the dorsal slit (a and b) after the foreskin is reduced.

Figure 55.14 Dorsal slit treatment of paraphimosis. This procedure may be necessary if manual reduction of paraphimosis (see Fig. 55.9) is unsuccessful.

Urethral Catheterization

Indications
- Acute urinary retention
- Obstructive uropathy
- Urine output monitoring in any critically ill or injured patient
- Collection of a sterile urine specimen for diagnostic purposes
- Intermittent bladder catheterization in patients with neurogenic bladder dysfunction
- Urologic study of the lower urinary tract

Contraindications
- Situations in which a less invasive procedure is sufficient
- Trauma patient with suspected urethral injury

Complications
- Urethral trauma and hemorrhage
- Paraphimosis (if the foreskin is not reduced after the procedure)
- Infection
- Undesirable catheter retention (nondeflating balloon)

Equipment

- Sterile drapes
- Sterile gloves
- Sterile water
- Urethral catheter
- Betadine solution and applicator forceps
- Cotton balls
- Sterile collection bag
- Lubricant

Review Box 55.1 Urethral catheterizations: indications, contraindications, complications, and equipment.

Procedure Overview

Indications: bladder access for urinary drainage or evaluation of bladder urine.
Contraindications: when other less invasive procedures will be equally effective and informative; relatively contraindicated in certain pelvic or GU trauma until urethral integrity is assured.
Complications: mechanical (false passage, iatrogenic paraphimosis following foreskin retraction for urethral access), infection, bleeding, undesired catheter retention, complications of long-term catheter use.

Equipment: the equipment listed in Review Box 55.1 is included in most standard catheterization trays and must be available before attempting urethral catheterization.

Key Procedure Sequence

1. Identify the urethral meatus; expose it with the nondominant hand.
2. Cleanse the meatus and surrounding tissues with antiseptic solution.
3. With the dominant hand, cleanse the area in a circular motion, from the meatus outward.
4. Lubricate the catheter (sterile), and pass it through the meatus towards the bladder.
5. For males: consider instilling 5 mL of 2% viscous lidocaine (or similar) into the urethra prior to passing the catheter.
6. For females: discard any catheter that has inadvertently entered the vagina.
7. Pass the catheter fully into the bladder, confirm its placement by the flow of urine. For men, continue to insert the catheter fully.
8. Inflate the balloon with 10 mL of air or water.
9. If resistance or obvious discomfort is felt, deflate the balloon and attempt repositioning.
10. After successful passage and balloon inflation, gently withdraw the catheter until resistance is met.
11. Connect the catheter to a drainage bag. Always reduce the penis foreskin if present to avoid inadvertent iatrogenic paraphimosis.

Description

Background
Patients are often apprehensive about catheterization. If the clinician shows concern regarding positioning and exposure, the patient will be reassured. Although adequate exposure may be obtained in the frog-leg position, the use of an examination table with stirrups (lithotomy position) is ideal, especially for female catheterization.

Anticipation and preparation of all materials necessary for urethral catheterization beforehand are reassuring to the patient. It is frustrating for the health care provider and upsetting to the patient when the patient is told “not to move or touch anything” while a search is made for additional equipment. Most catheterizations are performed using a standard catheterization tray. Often, these trays contain more equipment than is truly needed. This necessitates opening the tray and establishing a sterile field at the bedside, selecting those items that will be needed, and discarding the rest of the equipment. Once the penis or labia has been touched in preparation for the procedure, the touching hand (usually nondominant) is contaminated and ideally should not be handling any of the sterile equipment. When a standard catheterization tray is not used or is not available, go through the anticipated procedure mentally and secure all the appropriate equipment before actually starting the procedure.

Anatomy and Physiology

Female Catheterization
The female urethra is a short (~4 cm) straight tube, usually of wide caliber, lying on top of the vagina. It must be approached between the double labia, and the urethral meatus is occasionally hidden and not obvious. This is in contrast to most males, except those with hypospadias. If the female patient nervously adducts her legs, successful catheterization will be very difficult if not impossible.

The female urethral meatus is oval but may appear as an anteroposterior slit with rather prominent margins situated directly superior to the opening of the vagina and approximately 2.5 cm inferior to the glans clitoris (Fig. 55.15). It is the first of three orifices encountered when examining any female genitalia cephalad to caudal in the lithotomy position. The urethral meatus might be especially difficult to find in the very young infant and the older, postmenopausal woman. Anticipation of this and knowledge of the anatomic variances will help to modify any patient discomfort associated with needless catheter tip probing, which is an unsettling experience for both the patient and the catheterist.

Occasionally, the urethral meatus recedes superiorly into the vagina and is not immediately visible because of either prior surgical procedures or atrophic postmenopausal changes. Anticipation of such cases will allow the examiner to gently advance a nondominant index finger into the vagina in the superior midline. The urethral meatus can usually be palpated then visualized as a soft mound surrounded by a firmer ring of supporting periurethral tissue. Rarely, the meatus will have receded so far superiorly and intravaginally that it cannot be visualized at all, and catheterization must be carried out by palpation alone. From the meatus (if the patient assumes a supine position), the urethra proceeds straight back to slightly downward as it advances into the bladder just behind the symphysis pubis (Fig. 55.16A).

In women with a urethrocele or cystourethrocele, in whom the urethra or the bladder falls into the vagina, the “normal” urethral course might be significantly more posterior. The normal anatomic relationships in these situations may be re-created by spreading the index and long fingers and placing them along the superior vaginal wall and gently applying upward support (see Fig. 55.16B). This reconstitutes the normal anatomic relationships and permits straight, rapid urethral catheterization. Because the female urethra is so short, only half the total length of the catheter has to be inserted before it is safe to inflate the balloon.

Male Catheterization
Because the urethral meatus is usually evident in most males, it might seem a simple matter to insert a urethral catheter, yet catheterization can be quite difficult. The normal male urethra is approximately 20 cm long from the external urethral meatus to the bladder neck (Fig. 55.17). The posterior prostatic urethra is approximately 3.5 cm long, and the contiguous external sphincter or urogenital diaphragm that encompasses the membranous urethra is 4 cm from the bladder neck. In males, any catheter must be fully inserted to the balloon-inflating side-arm channel before it is safe to inflate the balloon. At the first egress of urine from the catheter, the balloon is just passing through the membranous urethra. The catheter balloon still has approximately 3 to 4 cm to go before clearing the bladder neck. Inflation of the Foley balloon at any point before full insertion of the catheter might result in iatrogenic urethral injury.

The male urethra is relatively fixed at the level of the urogenital diaphragm and symphysis pubis; traction downward on the penis kinks and promotes urethral folding at the level of the penile suspensory ligament (Fig. 55.18A), which creates a level of spurious obstruction. For this reason, the penis should always be held taut and upright during any urethral instrumentation, including catheterization (see Fig. 55.18B). The catheter then needs to make only a single curve rather than a complex S-curve as it traverses into the bladder.

Pathophysiology
See Table 55.3: Approach to Difficulties in Male Catheterization.
Figure 55.15 External female genitalia. The urethral meatus is directly superior to the vaginal introitus and approximately 2.5 cm inferior to the clitoris. (Netter illustration from https://www.netterimages.com. Elsevier, Inc. All rights reserved.)
**Indications**
Urinary catheterization and instrumentation are rarely a primary cause of urinary infection in otherwise healthy patients who urinate normally and carry small amounts of postvoid residual urine. As with any procedure, catheterization needs to be limited to those clinical situations in which the benefits outweigh the risks. The following are considered to be indications for urethral catheterization:

1. Acute urinary retention.
2. Urethral or prostatic obstruction leading to compromised renal function.
3. Urine output monitoring in any critically ill or injured patient.
5. Intermittent bladder catheterization in patients with neurogenic bladder dysfunction.

**Contraindications**
Urethral catheterization should be avoided when other less invasive procedures will be equally effective and informative. A traditional relative contraindication to urethral catheterization is the trauma patient with suspected urethral injury as evidenced by blood at the urethral meatus, an abnormal-feeling or high-riding prostate on rectal examination, penile, scrotal, perineal hematoma, or radiographic evidence of urethral/bladder trauma. In the setting of a severely fractured pelvis or diastasis of the pubic symphysis, urethrogram should always precede attempts at catheterization. Although some authors, and more liberal...
### TABLE 55.3 Approach to Difficulties in Male Catheterization

<table>
<thead>
<tr>
<th>CONDITION</th>
<th>ISSUE</th>
<th>WORK-AROUND STRATEGY</th>
</tr>
</thead>
</table>
| Phimosis                 | Inability to identify the urethral meatus because of a tight phimotic opening | Dilate the phimotic opening sufficiently to identify the urethral meatus and blindly pass the catheter  
Dorsal slit of the foreskin to expose the glans and urethral meatus in extreme cases |
| Foreskin edema           | Edema may obscure the glans penis and urethral meatus                  | Attempt to identify the cause of the edema; exclude paraphimosis and foreign body strangulation  
Manually compress the swollen foreskin by hand or between opposing cold packs  
If unsuccessful, snugly wrap the distal penis in a compressive dressing for 10 min |
| Meatal stenosis          | The urethral meatus may be either congenitally or secondarily narrowed by scarring | Trial of a smaller-caliber catheter  
If a larger-caliber catheter is necessary, meatal dilation or meatotomy might be necessary under the direction of a urologist |
| Urethral stricture       | May develop as a result of trauma, infection (especially STDs), lower urinary tract instrumentation, or long-term indwelling catheter drainage | Trial with a coudé catheter  
However, manual force should not be used to negotiate or to dilate urethral strictures because it may lead to false passages, bleeding, and future increased scarring  
Consideration of urethral dilation with the use of filiforms and followers under the direction of a urologist |
| External urethral sphincter spasm | Voluntary or involuntary contraction of the urogenital diaphragm (external sphincter) produces spurious urethral resistance at approximately 16 cm from the meatus | Encourage the patient to lay supine and take slow, deep breaths while consciously trying to relax the perineum and rectum  
Plantar flexion of the toes and ankles may also aid in relaxation of the pelvic floor  
Gentle but steady pressure exerted on the syringe or the catheter along with the aforementioned steps may aid in passage  
The external sphincter is composed of striated muscle and fatigues within a few minutes  
If continued resistance, assume an anatomic abnormality and consider RUG before further instrumentation |
| High bladder neck        | Enlarged intravesical portion of the prostate with a secondary high-riding bladder neck  
Typically resistance encountered after the catheter has been passed 16–20 cm into the urethra | Slow instillation or injection of 20–30 mL of sterile lubricating jelly  
If unsuccessful, consider a trial with a coudé catheter; may be enhanced by digital compression of the prostate during digital rectal examination |
| Pelvic trauma, straddle injury | Partial or complete urethral injury might occur  
Injudicious catheterization has the potential for conversion of a partial injury (may heal with little or no scarring) to a complete urethral injury (often requires surgical repair and usually results in some degree of postoperative morbidity)  
Blood at the urethral meatus, a high-riding or abnormal-feeling prostate on rectal examination, and penile, scrotal, or perineal ecchymoses might be evidence of potential urethral or bladder injury but have poor sensitivity |

*RUG, Retrograde urethrography; STDs, sexually transmitted diseases.*
clinical protocols, allow for gentle attempts at catheter passage in the presence of the previously discussed traditional contraindications, these findings necessitate strong consideration of retrograde urethrography (RUG) to delineate urethral integrity before any attempted urethral catheterization.68

**Procedure**

**Equipment**
The equipment listed in Review Box 55.1 is included in most standard catheterization trays and must be at hand before attempting urethral catheterization. Check the list of contents before opening the tray because some trays do not include certain items. For most routine adult in-and-out catheterizations, a 14-Fr red rubber catheter or Foley balloon catheter is adequate (Fig. 55.19). In neonates or infants, a small feeding tube taped in place produces the least amount of urethral trauma. In older boys, a red rubber catheter or Foley balloon catheter may be used. Table 55.4 lists age-appropriate sized catheters or feeding tubes for children.69

A 14- to 18-Fr coudé catheter should be considered after unsuccessful passage of a straight Foley balloon catheter or in any male patient with known enlargement of the median prostatic lobe (Fig. 55.20). If a coudé catheter is not available, a larger 18- to 22-Foley balloon catheter can be attempted. In a male patient with a urethral stricture in whom attempts at catheterization with a straight Foley or coudé catheter have failed, retrograde cystoscopy, passage of a guidewire, or passage of filiforms/followers under the direction of a urologist is the next step. If immediate bladder access is required in any emergency, suprapubic placement of a Foley balloon catheter using the Seldinger technique can be performed (as described in the later section on Suprapubic Cystostomy).70

**General Procedure (Figs. 55.21 and 55.22)**

As stated previously, urethral catheterization must be done using sterile technique. Both male and female patients have special urethral meatal considerations. In the uncircumcised or partially circumcised male, control of the penile foreskin is paramount to ensuring success. Before establishing a sterile field, retract the available foreskin to its fullest extent proximal to the glans penis (Fig. 55.23A). Unfold a standard 4 × 4 gauze pad, then refold it in its longest dimension and carefully wrap it around the retracted foreskin at the level of the coronal sulcus (see Fig. 55.23B). This will prevent the tendency for lubricant-associated normal anatomic foreskin reduction during catheterization and provide a continuously dry and sterile field. Secure the folded 4 × 4 gauze pad surrounding the foreskin between the nondominant long and ring fingers when beginning the procedure and do not release until the procedure is completed. This position leaves the nondominant index finger and thumb available for manipulating the catheter. After catheterization, remove the 4 × 4 gauze pad and reduce the penile foreskin to its normal anatomic position to prevent the development of an iatrogenic paraphimosis.

After exposing the urethral meatus in the female and the glans penis and urethral meatus in the male, use an antiseptic solution soaked into cotton balls or oversized cotton-tipped applicators to cleanse the exposed meatus and surrounding tissues. This is best done by hand but can also be done using the plastic forceps in the catheterization tray. Begin the cleansing in a circular motion on the urethral meatus and proceed outward, intentionally moving any debris toward the periphery and thus creating a sterile field.

**Table 55.4** Pediatric Urethral Catheter Size

<table>
<thead>
<tr>
<th>AGE GROUP</th>
<th>PEDIATRIC CATHETER SIZE*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infants (yr)</td>
<td>5- to 8-Fr feeding tube</td>
</tr>
<tr>
<td>1–3</td>
<td>8-Fr feeding tube</td>
</tr>
<tr>
<td>4–6</td>
<td>10-Fr feeding tube</td>
</tr>
<tr>
<td>7–12</td>
<td>10- to 12-Fr red rubber catheter (Robinson)</td>
</tr>
<tr>
<td>&gt; 12</td>
<td>14-Fr red rubber catheter (Robinson)</td>
</tr>
</tbody>
</table>

*Sizes are approximate; 6-Fr is the smallest balloon catheter and is quite flimsy, and therefore 8 Fr is recommended; 12-Fr is the smallest coudé catheter.

---

**Figure 55.19** Urethral catheters. A wide variety of types and sizes of catheters are available. For most adult catheterizations, a 14-Fr or 16-Fr Foley catheter is sufficient. For male patients with prostatic enlargement, a 14-Fr to 18-Fr coudé catheter may be used (see Fig. 55.20). For patients with gross hematuria, a large (20 Fr and up) three-way catheter may be used. Table 55.4 lists age-appropriate sized catheters or feeding tubes for children.

**Figure 55.20** Coudé catheter. A coudé catheter is used for men with prostatic enlargement. It features a semi-rigid curved tip that helps the catheter traverse the median lobe of the enlarged prostate gland. When inserting the coudé, the curved tip should be pointing upward toward 12 o’clock, that is, pointing toward the dorsal aspect of the penis.
**Male Urethral Catheterization and Bladder Irrigation**

Prepare your equipment and place a sterile fenestrated drape around the penis. If the patient is uncircumcised, retract the foreskin prior to the procedure (see Fig. 55.23).

Inject the urethra with 5 to 10 mL of 2% viscous lidocaine to distend the urethra and provide topical anesthesia. A commercially available Urojet syringe (depicted here) is ideal. If possible, wait 5 to 10 minutes for maximum anesthetic effect.

Advance the catheter to the hilt before inflating the balloon with the recommended amount of air or water. If there is obvious resistance or patient discomfort, immediately deflate and reevaluate the position of the catheter.

Affix the catheter to the thigh with tape or a catheter-specific attachment device. Note that a three-way catheter was placed in this patient with gross hematuria.

Continuously infuse the irrigant into the bladder. Rates of 1 to 2 L/hr are acceptable, provided that the amount of irrigant drained equals the amount infused.

Use your nondominant hand to hold the penis, and cleanse the meatus and surrounding tissue with antiseptic. Your nondominant hand is now contaminated and should not let go of the penis or handle sterile equipment until the insertion has been completed.

Hold the penis taut and upright with your nondominant hand while you pass the catheter into the urethra (see Fig. 55.18).

After balloon inflation, slowly withdraw the catheter until the balloon is against the bladder neck and precluding further withdrawal. Connect to a drainage system if one is not preattached.

To irrigate the bladder, attach the irrigation port to a 2- to 4-L bag of saline irrigation fluid. (Hand irrigation with a 60-mL catheter-tipped syringe is an alternative.) Attach the drainage port to a large-volume collection bag.

Monitor output carefully during the procedure. The goal of irrigation is to obtain clear urinary effluent.

---

**Figure 55.21** Male urethral catheterization and bladder irrigation.
Female Urethral Catheterization

Lubricate an appropriately sized catheter (14- to 16-Fr is commonly used in adults) with viscous rather than inspissated lubricating jelly. Injection of the male urethra with 5 mL of 2% viscous lidocaine or a syringe filled with anesthetic lubricant can be helpful for topical anesthesia and urethral distention. Regardless, advise the patient of mild urethral discomfort and the potential urge to void. Gently pass the catheter by hand, or with the aid of a hemostat or plastic forceps, into the urethra and upward into the bladder. During slow, gentle passage of the catheter, be aware of the anatomic considerations discussed previously. Any catheter that inadvertently enters the vagina should be discarded.

After passing the catheter fully in all male patients, slowly inflate the balloon with 10 mL of air or tap water. Sterile water or saline is not required. Most 5-mL balloons will easily accommodate 30 to 50 mL of air or water without bursting. Obvious resistance or patient discomfort on balloon inflation should signal potential erroneous urethral positioning and mandates reevaluation. If this occurs, immediately deflate the Foley balloon and reposition or slightly withdraw the catheter.

Figure 55.22 Female urethral catheterization.
Then connect the catheter to either a sterile leg bag or a closed-system bedside drainage bag. If the patient will be released with an indwelling Foley catheter, it can be initially connected to a leg bag, which is then comfortably fastened to the lower thigh and upper calf. The patient and family must be instructed regarding proper care of the catheter and drainage device. In most other cases, the catheter may be secured to either the thigh or the lower abdomen (preferred in males) with adhesive tape or simply placed under the knee and left to drain dependently into the bedside drainage bag.

then pass it fully again before balloon reinflation. If this is unsuccessful a second time, remove the catheter. At this point, RUG is recommended to evaluate the urethra for a potential obstructive problem or false passage (see later section on RUG). It may be difficult or impossible to pass a Foley catheter in a male with a markedly edematous penis (Fig. 55.24). If this cannot be accomplished, a suprapubic catheter may be required.

After successful catheter passage and Foley balloon inflation, slowly withdraw the catheter until the approximation of the balloon with the bladder neck precludes further withdrawal.

Figure 55.23 A, In this uncircumcised male, the foreskin has been fully retracted for placement of a catheter. B, A gauze pad allows a firm grasp on the foreskin to prevent movement during catheterization. Reduce the foreskin to its original position once the catheter is in place.

Figure 55.24 A, The urethral meatus cannot be found in the markedly edematous penis in this patient with massive anasarca. B, Wrap the penis in an elastic bandage for 8 to 10 minutes. C, Much of the edema is reduced with compression alone. D, An assistant stabilizes the penile shaft with the thumb and index finger (arrow) while the operator (asterisk) spreads the redundant tissue with forceps to visualize the meatus deep within the swollen tissue and advances the catheter.
Bladder Irrigation
After prostate surgery (transurethral resection) or in other clinical scenarios, blood clots may form in the bladder and cause acute urinary retention. Blood loss can be significant. This condition is usually easily relieved by the use of a large three-way irrigation catheter. A 22F to 26F Foley three-way catheter is preferred. Infuse saline irrigation fluid continuously into the bladder, while allowing the egress of fluid and blood clots. Continue the procedure until the bladder remains decompressed or bleeding stops (see Fig. 55.21, steps 8-10). Gravity alone usually provides adequate ingress–egress of fluid, but gentle syringe irrigation with 60-mL aliquots of saline may be used. Irrigation can be brisk, 1 to 2 L/hr or more, as long as the volume of drained saline is equal to the volume infused. Infuse saline (supplied in 2- or 4-L bags) by gravity. As stated previously, syringe irrigation of a Foley catheter with 60-mL aliquots is an acceptable alternative. The goal is to obtain clear urine.

Difficult Urethral Catheterization
Difficult urethral catheterization (DUC) is a common urologic problem. In females, this is due primarily to vaginal atrophy with retraction of the meatus into the vagina. This can typically be overcome by placing the patient in the lithotomy position, using a speculum, or guiding the catheter blindly with the provider’s index and middle fingers. However, DUC in males may present greater challenges. Common causes in males include urethral strictures, bladder neck contracture, false passage, and prostate disease. Trial of a couéd catheter is generally the first-line approach to DUC. Alternatives utilized by urologists include flexible cystoscopy, passage of a guidewire, or filiforms/followers. Suprapubic cystostomy should generally be considered an option of last resort given its inherent complications. Various causes and approaches to DUC in the emergency care setting are presented in Table 55.3.

Aftercare
Systemic antimicrobial prophylaxis should not be used routinely in patients with short- or long-term catheterization to reduce catheter-associated bacteruria or urinary tract infection because of concern about selection of antimicrobial resistance.

Complications
Although urethral catheterization performed by skilled personnel in appropriate circumstances has an acceptable complication rate, untoward sequelae of catheterization are not unusual.

Mechanical
False passages might be established in any area of the urethra when force is exerted on the catheter. In an uncircumcised patient, negligence in reducing the retracted foreskin to its normal anatomic position after urethral catheterization or instrumentation might lead to painful paraphimosis and associated complications.

Bleeding
Hematuria is common immediately, even after an atraumatic catheterization. Careful monitoring of symptom trajectory is prudent, as most often, postprocedure hematuria resolves rapidly and spontaneously.

Infection
The incidence of bacteriuria associated with indwelling catheterization is 3% to 8% per day, and the duration of catheteriza-
A flaplike defect in the inflating channel of a balloon catheter may prevent balloon deflation. A wire stylet (from a central venous catheter kit) can be passed down the inflating channel to raise the flap and deflate the balloon.

Note that the entire inflation channel is filled with debris (arrow), thus preventing egress of fluid from the retaining balloon.

Cut the catheter, remove the inflation port, and advance the stiff end of a guidewire through the inflation channel. The wire may puncture the balloon or clear the channel so that fluid escapes.

Once the channel has been cleared, insert a 20-gauge needle gently into the inflation channel to aspirate the balloon fluid.

In the case of a nondeflating suprapubic balloon, you can try to insert a 25-gauge spinal needle parallel to the catheter to puncture the balloon. Maintain traction on the catheter to stabilize the balloon during the procedure.

This balloon demonstrates “cuffing” where the balloon deflated asymmetrically and got caught up during withdrawal.

Figure 55.25 Removal of a nondeflating catheter. (1, From Eichenberg HA, Amin M, Clark J: Nondeflating Foley catheters, Int Urol Nephrol 8:171, 1976. Reproduced by permission.)
Traumatic Foley Catheter Removal

Occasionally, uncooperative or demented patients will pull out their Foley catheter with the balloon still inflated (Fig. 55.26). One would intuit that urethral injury is possible; however, there remains limited data on the incidence or type of injury from this maneuver or how to deal with this situation. Usually, there is blood at the meatus, and it may be difficult to know whether the patient can spontaneously urinate. If the patient can spontaneously urinate, it would seem reasonable to gently pass another Foley catheter to avoid urethral obstruction by tears or clots and allow healing of urethral trauma with a new catheter in place. Most of the time, however, the clinician has minimal data yet is faced with the decision on how to approach this conundrum. Surprisingly, this incident does not usually result in significant urethral injury. It is assumed that a new catheter will be required, either to continue the original catheter indication or to provide a stent while any urethral injury heals.

In the absence of available urologic consultation, gentle attempts at passage of another Foley catheter are reasonable, without a routine retrograde urethrogram (described later in this chapter). If the catheter does not pass easily, stop and perform a retrograde urethrogram to assess the nature and extent of the pathology. Most of the time, simply replacing the catheter will be the most prudent approach. Prophylactic antibiotics are reasonable under these circumstances. Complete eversion and prolapse of the bladder have been rarely associated with this misadventure.7

Conclusions

Access to and the subsequent evaluation of bladder urine are at times clinically important to emergency practitioners. The merits of various approaches to urine specimen collection depend on the patient’s age and clinical setting. Urethral catheterization generally provides a relatively safe and effective means of bladder access.

SUPRAPUBIC ASPIRATION

Introduction

Suprapubic aspiration of the bladder, first reported as a method of collecting urine for bacteriologic study in 1956,74 offers the clinician a relatively simple means of obtaining uncontaminated bladder urine. It is utilized primarily in young children, although there are limited indications for this procedure in adults as well. It has been demonstrated that with proper technique, suprapubic aspiration consistently provides an uncontaminated urine specimen capable of distinguishing true bacteriuria from contamination.79,80 However, the perception and demonstration of increased discomfort and procedural failure rate has led to a precipitous decline in its use, particularly outside of the neonatal intensive care unit.80–82 Nevertheless, suprapubic aspiration continues to be considered the gold standard method of urine collection.83

Despite the lack of experience by most currently practicing clinicians, the procedure may still be necessary in girls with labial adhesions and boys with phimosis. The relatively recent introduction of ultrasound technology has led to a significant improvement in success rates, and likely in the comfort of the clinician performing the procedure.84–86

Procedure Overview

Indications: provides the clinician with a urine sample that is reliable for bacteriologic interpretation.

Contraindications: infection at proposed puncture site.

Complications: infection or hematoma at puncture site; simple penetration of the bowel with a needle is considered an innocuous event and requires no specific treatment.

---

**TABLE 55.5 Alternative Methods of Removing Nondeflating Catheters**

<table>
<thead>
<tr>
<th>METHOD</th>
<th>COMMENTS</th>
</tr>
</thead>
</table>
| Needle puncture of the balloon (see Fig. 55.25, step 5) | Overinflate the balloon (50–100 mL of saline)  
Draw the balloon against the bladder neck  
Use local anesthesia for the suprapubic approach  
Use a 25- to 27-gauge spinal needle to puncture the balloon  
Ultrasound guidance may be helpful |
| Hyperinflation of the balloon | High rate of balloon fragmentation: 100 of 100 in one report74  
Postprocedure cystoscopic inspection of the bladder essential  
Consider adding 50–100 mL of saline into the bladder to cushion the effects of subsequent balloon rupture if no urinary retention is present  
Inject up to 200 mL of air into a 5-mL balloon to induce rupture |
| Injection of an erosive substance into the balloon port (e.g., mineral oil) | This technique has fallen out of favor  
High rate of balloon fragmentation: 95 of 100 in one report74  
Associated with chemical cystitis |
**Anatomy and Physiology**

See following figures.

**Pathophysiology**

Results in urine specimen collection capable of distinguishing true bacteriuria from contamination.

**Indications**

In the neonate or the young child, suprapubic aspiration or urethral catheterization can provide the clinician with a sample that is reliable for bacteriologic interpretation. Although disconcerting to some parents, suprapubic aspiration is not a dangerous procedure, and the sensitivity of urinalysis for bacteriuria approaches 100%. However, for many young children, urine can generally be more readily collected by urethral catheterization.

For adult patients, the indications for suprapubic aspiration are more limited because these patients can usually cooperate with the clinician. Men with condom catheters or phimosis, however, may require suprapubic aspiration to minimize urethral contamination. Aspirated cultures, rather than catheterized specimens, may help assess for infection versus contamination in patients with asymptomatic bacteriuria on routine urine collection. In infections caused by organisms that in other circumstances are often discounted as contaminants (e.g., *Staphylococcus epidermidis* or *Candida albicans*), suprapubic
aspiration or a catheterized specimen is required to confirm the presence of such pathogens. In patients in whom the possibility of intravesical infection is a concern (e.g., patients with chronic infections of the urethra or the periurethral glands), suprapubic aspiration may help distinguish a bladder source from a urethral source.

Contraindications
Skin or soft tissue infection in the area of the proposed anterior abdominal wall puncture site.

Procedure
Place the child supine with the legs in a frog-leg position and restrain as necessary (Fig. 55.27A, step 1). First locate the bladder. A full, palpable, or percussible bladder should be readily apparent, but this can be difficult to discern in all but the thinnest patients. If there is any question about the location or the amount of bladder urine, a quick ultrasound examination is informative. The point of entry in the skin should be 1 to 2 cm above the superior edge of the symphysis pubis. Pass the syringe and needle perpendicular to the abdominal wall toward the bladder, usually at a 10- to 20-degree angle from the true vertical, somewhat cephalad in children (see Fig. 55.27A, step 2) and somewhat caudad in adults (see Fig. 55.27B). Note that the bladder of a newborn is an abdominal organ and that it will be missed if the needle is inserted too close to the pubis or angled toward the feet.

After draping the prepared skin and choosing the point of entry, raise a skin wheal of local anesthesia to reduce discomfort. After anesthetizing the skin, advance a longer, mid-caliber needle (usually 22-gauge, 3.75 to 8.75 cm in length) in the midline through the skin and quickly into the bladder. Advance the needle attached to a syringe and aspirate during advancement. As soon as the bladder is entered, urine will appear in the syringe. A short needle is adequate for virtually all pediatric patients. After collecting the urine, withdraw the syringe and needle. Microscopic hematuria always follows the procedure but gross hematuria is uncommon.

If urine is not obtained, do not remove the needle but withdraw it to a subcutaneous position and redirect it at a different angle. Often, a child may spontaneously start to void after any type of invasive stimulus (e.g., bladder irrigation by a probing needle, venipuncture, or lumbar puncture). Hence, prepare to collect a spontaneously voided specimen, should that option arise. Anticipate this before beginning blood or spinal fluid collection during the bacteremic workup of the febrile neonate.

In most patients, an acceptable urine sample can be obtained with the first needle pass. If the needle points too caudad, in an effort to avoid entering the peritoneum, it is possible to enter the retropubic space, skimming the bladder muscle and never penetrating the bladder mucosa.

Aftercare
Following withdrawal of the needle, place a simple bandage over the puncture site.

Complications
Bacteremia does not typically result from this procedure.\(^8\) Bowel penetration has occurred in children with distended abdomens from gastrointestinal disturbances.\(^9\) The combination of gaseous bowel distention and relative hypovolemia might displace and flatten the relatively empty bladder against the pelvic floor. Even when the large bowel has been penetrated, most patients typically recover uneventfully. As a general rule, simple penetration of the bowel with a needle is considered an innocuous event and requires no specific treatment.

Conclusions
Although it can be performed in a patient of any age, suprapubic aspiration is utilized for urine collection in young children only when alternative collection methods are not feasible. It consistently provides a urine specimen that is capable of distinguishing true bacteriuria from contamination. However, in many cases, alternative methods of urine collection are preferred over this procedure.

SUPRAPUBIC CYSTOSTOMY
Introduction
In general, any patient requiring a urethral catheter, but in whom a catheter cannot be safely passed, is a candidate for a suprapubic cystostomy tube (Video 55.3). Among the most user-friendly devices for suprapubic bladder access is the Cook peel-away sheath unit (Cook Medical, Bloomington, IN).\(^9\) It uses the Seldinger (guidewire) technique to gain bladder access and allows suprapubic placement of a Foley balloon catheter for definitive bladder drainage. This device is readily suitable for ED use when compared with other suprapubic bladder access approaches, such as trocar-type devices.

When emergency bladder drainage is required and a transurethral Foley catheter cannot be placed, any device suitable for central venous access can be inserted suprapublically using the Seldinger technique (Fig. 55.28).

Procedure Overview
Indications: need for emergency bladder access and drainage. Contraindications: bladder not readily definable or low bladder volume; relative contraindications include history of previous lower abdominal surgery, intraperitoneal surgery, irradiation, bleeding diathesis, or concurrent antithrombotic use. Complications: a wide variety of complications have been reported, including bowel or other organ injuries, which serve as reminders that suprapubic cystostomy is not innocuous.

Equipment: Chiou Suprapubic Tube Insertion Kit (Cook Medical) and Foley catheter.

Key Procedure Sequence
1. Prep and drape the skin for a sterile procedure.
2. Inject local anesthesia ~2 to 3 cm above the pubic symphysis.
3. Locate the bladder by palpation (or ultrasound if available).
   While aspirating, advance a 22-gauge spinal needle with an attached syringe until urine returns.
4. Keep the needle in place, remove the syringe, and thread the guidewire through the needle into the bladder.
5. Once the guidewire is threaded into the bladder, remove the needle.
6. Make a stab incision at the guidewire/skin interface.
7. Pass a peel-away sheath and indwelling fascial dilator device over the wire.
Description

Background
In situations where emergency bladder access is necessary, the Seldinger (guidewire) technique allows for suprapubic placement of a Foley balloon catheter for definitive bladder drainage.

A. Pediatric

1. Restrain the infant and place her in a frog-leg position. Prepare and drape the skin, and raise a skin wheal with local anesthetic to reduce discomfort. Despite the safety of this procedure, it may be disconcerting for worried parents, and they may wish to leave the room during the aspiration.

B. Adult

2. Puncture the abdominal wall with a 22-gauge needle in the midline approximately 1 to 2 cm cephalad to the superior border of the pubic bone. Keep the syringe perpendicular to the plane of the abdominal wall (usually 10 to 20 degrees from the true vertical). The bladder is an abdominal organ in infants, and placing the needle too close to the pubic bone or angling toward the feet might cause the needle to miss the bladder. Localizing the bladder with bedside ultrasound facilitates this procedure.

In adults, the peritoneum is pushed cephalad by the filled bladder during suprapubic aspiration. Direct the needle slightly caudad.

Figure 55.27 Suprapubic aspiration.

8. Remove the guidewire and fascial dilator, leaving only the sheath in place.
9. Pass a Foley balloon catheter through the sheath, and inflate the catheter.
10. Withdraw the peel-away sheath, leaving only the suprapubic catheter in place.
Suprapubic cystostomy with a Foley catheter inserted through a Cook peel-away sheath introducer (Cook Medical, Bloomington, IN).

**Figure 55.28** Suprapubic cystostomy. A, Any catheter that can be used as a central venous catheter can be inserted suprapubically via the Seldinger technique. B, The more traditional and more permanent suprapubic cystostomy with a Foley catheter inserted through a Cook peel-away sheath introducer (Cook Medical, Bloomington, IN).

**Anatomy and Physiology**

See following figure.

**Pathophysiology**

In emergency situations, the majority of patients requiring suprapubic cystostomy are trauma patients with urethral disruption, or men with severe urethral stricture or complex prostatic disease.

**Indications**

If time allows, it is prudent to discuss the case with a urologist to explore alternative methods of bladder access and urine drainage prior to initiation of a suprapubic cystostomy. Even in the case of major trauma, many affected patients will require laparotomy for associated injuries, so the suprapubic catheter can be placed intraoperatively.

**Contraindications**

Because placement of a suprapubic tube involves some risk, proper patient selection is important. Do not perform the procedure in a patient whose bladder is not readily definable. Although no absolute reported minimum bladder volume has ever been established, there must be enough urine in the bladder to allow the needle to fully penetrate the bladder dome without immediately exiting through the base. There must also be enough urine in the bladder to displace the bowel away from the anterosuperior surface of the bladder and the entrance of the needle. Ultrasound can be helpful in defining bladder anatomy.

Individuals who have a history of previous lower abdominal surgery, intraperitoneal surgery, or irradiation may have developed adhesions or adherence of the bowel to the anterior bladder wall. They are potentially at greater risk for bowel injury during percutaneous suprapubic cystostomy tube placement than those without previous abdominal surgery. Blind suprapubic cystostomy tube placement should be avoided in these patients. The absence of any of these risk factors does not totally exclude the risks of bowel or intraperitoneal injury, but it reduces them significantly. Patients with bleeding diatheses are at greater risk for postinsertion bleeding, either into the bladder, into the retropubic space, or at the site of skin entry.

**Procedure**

The following comments concern the placement of the Cook peel-away sheath, as described by Chiou and colleagues. With modifications, the same principles apply for any type of suprapubic catheter placement.

If necessary, shave the lower abdomen and apply antiseptic topically. Fill a 10-mL syringe with 1% lidocaine, and attach it to a 22-gauge, 7.75-cm or similar spinal needle. Raise a skin wheal in the proposed site (~2–3 cm above the pubic symphysis), and infiltrate the subcutaneous tissue and rectus abdominis muscle fascia at a 10- to 20-degree angle toward the pelvis.

Locate the bladder by advancing the needle in the prescribed direction while aspirating the syringe. Urine is easily aspirated when the bladder is entered (Fig. 55.29, step 1).

Once the bladder has been located, remove the syringe from the needle and advance a guidewire through the needle into the bladder (see Fig. 55.29, step 2). Withdraw the needle, leaving only the guidewire traversing the anterior abdominal wall and positioned inside the bladder. Use a No. 11 or similar scalpel blade to make a stab incision directly posterior to the wire through the skin, subcutaneous tissue, and superficial anterior abdominal wall fascia. Then pass the peel-away sheath and indwelling fascial dilator together over the wire into the bladder (see Fig. 55.29, step 3). Remove the guidewire and fascial dilator, leaving only the peel-away sheath inside the bladder (see Fig. 55.29, step 4). Then pass a Foley balloon catheter through the indwelling intravesical sheath into the bladder (see Fig. 55.29, step 5). Aspirate urine to confirm proper placement. Inflate the Foley balloon with a minimum of 10 mL of air, water, or saline (see Fig. 55.29, step 6). A 5-mL balloon will accommodate 10 mL easily and make accidental catheter distraction less likely. Withdraw the peel-away sheath from the bladder and anterior abdominal wall and literally peel it away from the catheter, leaving only the indwelling suprapubic Foley catheter (see Fig. 55.29, step 7). Withdraw the catheter slowly until the inflated balloon approximates the cystostomy site (see Fig. 55.29, step 8).

**Aftercare**

Connect the catheter to a drainage bag, and then dress the wound with 4 × 4 gauze pads to complete the procedure. The major difficulty with cystostomy tubes of all designs is securing them to the patient’s skin. Those with retention balloons, such as the standard Foley urethral catheter, are the most secure and need only tape to attach them to the anterior abdominal wall.
Suprapubic Cystostomy (Peel-Away Sheath Technique)

1. Enter the bladder with a syringe and needle. Confirm location by aspirating urine.

2. Remove the syringe and pass the guidewire through the needle into the bladder.

3. Remove the needle and then pass the dilator and peel-away sheath over the wire into the bladder. A small stab wound in the anterior abdominal fascia may be required to accommodate the dilator and sheath.

4. Remove the dilator and wire and leave only the sheath inside the bladder.

5. Pass the preselected Foley balloon catheter through the sheath into the bladder. Aspirate urine to confirm location.

6. Inflate the balloon with a minimum of 10 mL of air, saline, or water. A 5-mL balloon will accommodate 10 mL easily and make accidental catheter distraction less likely.

7. Remove the sheath from the bladder, anterior abdominal wall, and cutaneous entry site, and then literally peel it away from the indwelling catheter.

8. Withdraw the catheter until a snug fit is ensured at the cystostomy site.

Figure 55.29 Suprapubic cystostomy using a peel-away sheath. If this equipment is not available, an analogous procedure using any catheter suitable for central venous catheterization may be performed.
Complications

A wide variety of complications have been reported, which serve as reminders that suprapubic cystostomy is not innocuous. Occasionally, the suprapubic tube or catheter cannot be positioned or maintained successfully without problems (Box 55.5).

The most serious complications involve perforation of the peritoneum or the intraperitoneal contents. Although finding the bladder using a small-gauge “scout” needle may help reduce bowel injury, even in the most apparently successful of bladder punctures, a complication may still result.

Occasionally, the clinician proceeds with suprapubic cystostomy when the bladder is not palpable and has not been located with a syringe and needle. Injury of adjacent organs is much more frequent in these circumstances. Allowing the bladder to refill over time can be an important consideration, especially if initial attempts to locate the bladder were unsuccessful. Ultrasound guidance may also be helpful for determining bladder size and location.

Conclusions

Suprapubic cystostomy can be readily performed when emergency bladder access is needed. However, the procedure may be associated with significant risks to the patient. As such, when time affords, discussion with a urologist is prudent to explore alternative methods of bladder access and urine drainage.

Box 55.5 Complications of Suprapubic Cystostomy

- Bowel perforation
- Through-and-through bladder penetration with associated rectal, vaginal, or uterine injury
- Intraperitoneal extravasation (without a previous history of surgery)
- Extraperitoneal extravasation
- Ureteral catheterization
- Obstruction of the tubing (blood, mucus, or kinking)
- Tubing comes out
- Infection
- Hematuria

LOWER GENITOURINARY TRACT IMAGING

Introduction

Although the signs of GU trauma in general can be quite subtle, lower urinary tract injury can often be quickly identified and thoroughly evaluated radiographically in the ED. Radiologic imaging of the upper urinary tract is generally a less urgent matter and can usually be done in the radiology suite or, when important for emergency operative decision-making, as a single shot intravenous pyelogram (IVP) in the operating room.

The timing of any radiologic evaluation can be challenging in the emergency setting, especially when faced with a critically ill, multiply injured patient. The trauma team of clinicians involved in each resuscitation must determine the priority and extent of such an evaluation on a case-by-case basis.

In patients with urologic trauma, the lower urinary tract should be studied before the upper urinary tract (e.g., study the urethra before the bladder, study the bladder before the kidneys). As such, specific diagnostic studies should be conducted in a retrograde fashion (Fig. 55.30).

RUG and retrograde cystography are the diagnostic procedures of choice to evaluate potential injury to the lower urinary tract. These studies must be carried out in the proper sequence and in a retrograde fashion to avoid missing potential injuries.

Figure 55.30 Diagnostic approach to genitourinary (GU) tract trauma. The GU tract is studied in a retrograde fashion, proceeding from the urethra (retrograde urethrography) to the bladder (conventional or computed tomographic [CT] cystography) to the kidneys and ureters (CT urography with delayed/excretory images). These studies must be carried out in the proper sequence and in a retrograde fashion to avoid missing potential injuries.

RUG and retrograde cystography are the diagnostic procedures of choice to evaluate potential injury to the lower urinary tract. These studies must be carried out in the proper sequence and in a retrograde fashion to avoid missing potential injuries. Retrograde refers to the technique of instilling contrast retrograde through the urethra or by gravity filling of the bladder. It must be distinguished from antegrade filling, in which intravenous contrast for IVP or abdominal computed tomography (CT) is excreted from the kidneys and allowed to fill the bladder passively over time.

Procedure Overview

Indications: RUG is indicated whenever uncertainty exists about the integrity of the urethra; retrograde cystography is indicated any time a bladder injury is suspected.

Contraindications: uncertainty about urethral integrity is a contraindication to blind urethral catheterization; ensuring urethral integrity with a RUG prior to Foley catheter placement is required for retrograde cystography; prior...
history of reaction to radiographic contrast material is a precaution. Complications: tissue irritation from contrast extravasation; progression of urethral injury from overly forceful injection of contrast material. Equipment: equipment and personnel for plain radiographs (or fluoroscopy), small Foley catheter (or Toomey syringe), contrast material for injection.

**Key Procedure Sequence**

**Retrograde Urethrogram**
1. Assemble the items and personnel needed for plain imaging (or fluoroscopy).
2. In case of pelvic fracture, keep the patient supine.
3. In other cases, orient the hips in a slightly oblique position in relation to the stretcher.
4. Gently extend and stretch the penis along the medial thigh to maximize urethral unfolding.
5. Take a plain film (kidneys, ureters, and bladder [KUB]) or fluoroscopy for a reference image prior to injecting the contrast.
6. Retract and secure foreskin/glans with a folded 4 × 4 gauze pad.
7. Hold the penis between the long and the ring fingers of the nondominant hand.
8. Insert a non–Luer-Lok syringe or a small Foley catheter snugly into the meatus.
9. Before injecting contrast, stretch the penis perpendicularly across the patient’s thigh.
10. Gently inject 10 to 15 mL of contrast; avoid an overly forceful injection.
11. Immediately following the contrast injection, obtain a radiograph (urethrogram).

**Retrograde Cystogram**
1. Assemble items and personnel needed for plain imaging (or fluoroscopy).
2. In the case of a pelvic fracture, keep the patient supine.
3. In other cases, orient the hips in a slightly oblique position in relation to the stretcher.
4. Perform a RUG to ensure urethral integrity prior to placement of the catheter needed for the cystogram.
5. Remove the central plunger from a 60-mL catheter-tip syringe and attach the tip to a Foley catheter.
6. Hold the syringe above the level of the bladder and pour contrast material into the syringe.
7. Allow the bladder to fill by gravity instillation, and fill to capacity (~400 mL in an adult).
8. Clamp the catheter, obtain both anteroposterior (AP) and oblique images (or AP and lateral if oblique not feasible).
9. Obtain an AP postevacuation film in all cases after bladder drainage.
10. If bladder injury is detected, keep the catheter in place and consult urology.

**Description**

**Background**
In the resuscitation of any trauma patient, placement of a Foley catheter is a standard method of monitoring urinary output. Blood at the urethral meatus, however, may indicate a partial or complete urethral disruption and dictates the need for a RUG to delineate urethral integrity. This study can be done by the resuscitating clinician in the ED or on the operating room table by the trauma surgeon or urologist if the patient requires immediate surgical intervention for life-threatening injuries. RUG is a quick, technically easy study to perform and should be part of every emergency clinician’s armamentarium. Similarly, retrograde cystography allows for the evaluation of bladder integrity.

**Anatomy and Physiology**
The anatomy of the male urethra is demonstrated in Fig. 55.17. The male urethra varies from 17.5 to 20 cm in length for adults and consists of anterior and posterior portions, each of which is subdivided into two parts. The pendulous and bulbous urethral segments are anterior, and the membranous and prostatic segments are posterior. The male urethra is approximately 4 cm long and extends from the bladder neck at the urethrovaginal junction to the vaginal vestibule, where it forms the external meatus between the labia minora (see Figs. 55.15 and 55.16).

**Pathophysiology**
The male posterior urethra, which includes the membranous and prostatic urethra, is injured more frequently than the anterior urethra. The urogenital diaphragm encloses and fixes the membranous urethra; the prostate and prostatic urethra are firmly attached to the posterior surface of the symphysis pubis by the puboprostatic ligaments. Blunt trauma and pelvic fractures, especially in the presence of a full bladder, may result in shearing forces that partially or completely avulse portions of the firmly attached posterior urethra. Usually, the bladder and prostate gland are sheared from the membranous urethra, resulting in a complete urethral disruption (Fig. 55.31). The female urethra, in contrast, is short and relatively mobile and generally escapes injury in blunt trauma. Occasionally, a significant pelvic fracture will result in a laceration or avulsion of the female urethra at the bladder neck. Direct injuries to the female urethra may also occur secondary to penetrating trauma to the vagina or perineum. These injuries often are disclosed by blood at the introitus or an abnormal vaginal examination in the female pelvic fracture patient.91 Contusions or lacerations of the male anterior urethra occur when the bulbous urethra is compressed against the inferior surface of the symphysis pubis. This happens most commonly as a result of straddle injuries in males but may result from any blunt perineal trauma. Penetrating injuries or urethral instrumentation may also cause injuries to the penile urethra. Anterior urethral injuries may result in extravasation of blood or urine into the penis, scrotum, or perineum or along the anterior abdominal wall, depending on whether Buck’s fascia has been violated (Fig. 55.32). This is in contrast to posterior urethral injuries, in which blood and urine extravasate into the pelvis.

**Indications**
RUG is indicated whenever uncertainty exists about the integrity of the urethra. The RUG is an anatomic rather than a physiologic examination, but is critical in the identification of a possible urethral injury prior to urinary catheter insertion to preclude a potentially devastating complication.91

The traditional physical indications to perform a RUG in the trauma setting include blood at the urethral meatus, a
Retrograde Urethrogram

A patient with an associated pelvic fracture should remain supine throughout the entire radiographic examination. In cases of suspected urethral injury not associated with pelvic fracture, it is acceptable to obtain oblique films during the nonpalpable or high-riding prostate, or evidence of urethral trauma. However, the value of these findings to the emergency clinician has been called into question, given their low sensitivity.94,95

Perform a retrograde cystogram any time a bladder injury is suspected. It assumes that the urethra is normal before passing the Foley catheter required for retrograde cystography.

Contraindications

Uncertainty about urethral integrity is a contraindication to blind urethral catheterization. As such, urethral integrity must be ensured prior to Foley catheter placement, which is essential for retrograde cystography. History of a prior reaction to radiographic contrast material is a relative contraindication to contrast use, depending on the route of administration selected. Intravenous administration for antegrade imaging carries a much greater risk of serious adverse events as compared with the negligible risks associated with retrograde injection or installation of iodinated contrast materials necessary for lower GU tract imaging.

Procedure

Several different iodinated contrast agents are suitable for retrograde studies (urethography, cystography). Common agents, dosages, and methods of administration are listed in Table 55.6.

RUG and retrograde cystography should ideally be performed under fluoroscopy to provide real-time imaging, but plain films are an acceptable and frequently utilized alternative.

Retrograde Urethrogram

A patient with an associated pelvic fracture should remain supine throughout the entire radiographic examination. In cases of suspected urethral injury not associated with pelvic fracture, it is acceptable to obtain oblique films during the
Seal the balloon portion of the catheter in the fossa navicularis of the penile urethra, and delicately inflate it with 1.0 to 1.5 mL of saline solution to ensure a snug fit. Lubrication is not recommended because it may prevent the balloon from remaining in place. Alternatively, apply steady manual pressure of the balloon against the external urethral meatus (distal to the fossa navicularis). Another option is to gently advance a catheter-tipped Toomey irrigating syringe or a regular 60-mL syringe with an attached Christmas-tree adapter or a non–Luer-Lok syringe inside the urethral meatus until a snug fit is ensured (see Fig. 55.33, step 2). Gently squeeze the meatus over the injecting device to prevent leaking.

Inject contrast through the catheter or syringe. If not done carefully, this technique often results in the spillage and deposition of contrast outside the urethra and onto the patient and the examination table, yielding a spurious result. As such, slowly inject anywhere from 10 to 60 mL of contrast material (typically less than 10–20 mL is all that is needed) under constant pressure into the urethra. Not uncommonly, a spasm of the external urethral sphincter will be encountered, preventing filling of the posterior urethra. Use slow, gentle pressure to overcome this resistance. Overly forceful injection of contrast material may cause intravasation of contrast material into the venous plexus of the urethra, simulating an injury (see Fig. 55.33, step 4). Finally, during the injection of the last 10 mL of contrast, take a film (the urethrogram) (see Fig. 55.33, step 3). Alternatively, view the urethrogram in real time using fluoroscopy.

The extravasation of contrast material from a urethral disruption usually appears as a flamelike density outside the urethral contour (see Fig. 55.33, steps 5–7). If any contrast material is seen within the bladder in conjunction with urethral extravasation, a partial rather than a complete urethral disruption is more likely. In a complete urethral disruption, urethral extravasation will be present without evidence of contrast within the bladder. The examiner needs to be certain that the lack of bladder contrast is not secondary to voluntary contraction of the external sphincter. Occasionally, as mentioned previously, intravasated contrast material is seen in the periurethral penile venous plexus (see Fig. 55.33, step 4). It is of no clinical significance and should not be mistaken for urethral extravasation. As expected, penile venous intravasation (venous plexus opacification) will clear spontaneously on any postvoid films, unlike urethral extravasation, which remains in place.

If a Foley catheter has been successfully placed into the bladder and a partial urethral injury is suspected later, such an injury can be readily demonstrated without removing the catheter. Place the lubricated end of a pediatric feeding tube into the proximal penile urethra alongside the existing Foley catheter (Fig. 55.34). Obtain a seal by compressing the glans penis with the nondominant thumb and index finger and gently inject contrast material with the dominant hand. In this way, extravasation can be demonstrated. It should be noted, however, that successful placement of the Foley catheter obviates the need for further treatment of a partial urethral tear in the emergency setting because an indwelling catheter alone is appropriate initial management for this type of injury. The finding of an associated urethral injury must be conveyed to a urologist because it will dictate the duration of definitive Foley catheter drainage.

Retrograde Cystogram
As noted, demonstrate that the urethra is normal before passing the Foley catheter required for retrograde cystography.

First, take a plain film (KUB) or fluoroscopic image for reference before injecting any contrast material. Retract and secure the penile foreskin with a folded 4 × 4 gauze sponge. Second, hold the penis between the long and the ring fingers of the nondominant hand to allow a snug fit of the contrast-filled syringe or catheter inside the urethral meatus. Stretch the penis laterally over the proximal thigh with moderate traction to prevent urethral folding (i.e., the double image of the proximal penile and bulbous urethra superimposed on one another) to ensure a high-quality RUG (Fig. 55.33, step 3).

After sterile penile preparation, get ready to inject the dye. A small Foley catheter may be used (see Fig. 55.33, step 1).
Retrograde Urethrography

Insert the balloon portion of a Foley catheter (8 Fr) into the urethra, and slowly inflate the balloon with 2 mL of air or water to create a snug fit. Then slowly inject 10 to 60 mL of a 10% solution of contrast material through the catheter lumen. Usually only 10 to 20 mL is required.

Alternatively, use a plain-tipped (non-Luer-Lok) syringe or a catheter-tipped (Toomey) syringe in the urethra. Squeeze the glans around the syringe to prevent spillage of the contrast material. Inject slowly.

Inject the contrast material slowly. Injecting too fast causes venous intravasation (arrows). This may mimic urethral extravasation but it clears immediately, as opposed to actual extravasation, which remains indefinitely. If unsure, take another film in 10 minutes. The presence of intravasation is benign but can confuse the clinician.

Penile urethral disruption. Extravasation of contrast material (arrow) is seen in the corpus cavernosum.

Bulbar urethral disruption. Extravasation of contrast material (white arrows) is seen in the scrotum.

Supramembranous urethral disruption. Extravasation of contrast material (arrow) is noted superiorly.

Obtain a preliminary KUB or fluoroscopic image to serve as the reference film for the entire examination. Next, fill the bladder under direct operator supervision by gravity instillation of contrast material. After removing the central plunger from a 60-mL catheter-tip syringe, attach the catheter-tipped end of the syringe to the Foley catheter and hold it above the level of the patient’s bladder. Pour the contrast material into the syringe and allow it to fill the bladder by gravity instillation to one of three end points: (1) 100 mL with evidence of gross extravasation on fluoroscopy or on plain film (if the examiner elects to check at this point); (2) 400 mL in an adult or any child aged 11 years or older. In children younger than 11 years, bladder capacity, and therefore appropriate contrast volumes, are estimated based on this formula \((\text{age in years} + 2) \times 30\); or (3) the point of initiating a bladder contraction (see later), then add an additional 50 mL by hand injection under gentle but steady pressure.

Obtain AP and complementary oblique projections if feasible (Fig. 55.35A1). In the presence of a pelvic fracture, obtain all films with the patient in the supine position for the same reasons that were elucidated for RUG. A lateral film may be informative when oblique films are not possible (see Fig. 55.35A2). Obtain an AP postevacuation film in all cases after bladder drainage to disclose posterior perforation, especially in cases of penetrating trauma (see Fig. 55.35A3). A dilute solution of contrast material (see Table 55.6) may be used, rather than full-strength contrast. Some authors recommend a dilute solution of contrast material (≤ 10%) because extravasation into the periurethral or perivesical tissues may cause considerable inflammatory reaction at higher concentrations. The dilute solutions do not appear to compromise the quality of the study, but this must remain a consideration. Retrograde cystography done by any technique other than hand-poured gravity instillation is subject to inadequate bladder filling or connector tubing-catheter disconnection. Both conditions will result in spurious examination results, which may adversely affect important patient management decisions.

It must be stressed that in the absence of initial gross extravasation, the bladder must be filled to 400 mL in an adult, or to an appropriate capacity in a child, and the catheter clamped with a Kelly clamp. Volumes less than 400 mL have been associated with false-negative findings, especially in penetrating bladder injuries. At times, the patient may have difficulty cooperating with bladder filling because of a head injury or associated pain; and in the case of severe injury, the patient may have involuntary bladder contractions, causing contrast material to back up into the syringe. If this occurs, refill the bladder to the point of initiating a bladder contraction, clamp the Foley catheter, remove the initial syringe and replace it with a 60-mL contrast-filled syringe, unclamp the catheter, hand-inject the additional 50 mL under pressure, and reclamp the catheter. The goal is to overdistend the bladder. Once the filled-bladder films have been obtained and reviewed, unclamp the Foley catheter and allow the contrast material to drain into a bedside drainage bag. Then obtain the AP postevacuation film to visualize any posterior extravasation that may have been hidden by the distended bladder during the AP filled-bladder film (see Fig. 55.35B2 and B3). Once again, take care to ensure that contrast material is not spilled onto the patient or the examination table during the procedure.

Extravasation from an injured bladder may be intraperitoneal, extraperitoneal, or both. Extraperitoneal extravasation is usually seen as flamelike areas of contrast material confined to the pelvis and projecting laterally to the bladder (see Fig. 55.35B1 and B4). If the contrast material extravasates intraperitoneally, it tends to fill the paracolic gutters and outline intraperitoneal structures, particularly the bowel, spleen, or liver (see Fig. 55.35B5). It is important to distinguish extraperitoneal from intraperitoneal injury because the treatment options differ significantly: surgical repair is required for all intraperitoneal injuries and for those extraperitoneal injuries that extend into or primarily involve the bladder neck, especially in women. Most other extraperitoneal injuries can be managed confidently by Foley catheter drainage alone.

Retrograde cystography may be done in conjunction with contrast-enhanced abdominal CT scanning (Fig. 55.36). The bladder must be filled just as if a conventional retrograde cystogram were being obtained. Clamp the catheter and seek evidence for contrast ascites on the CT scan. When this is encountered, bladder injury with extravasation must be further evaluated with selective images of the pelvis.

**Aftercare**

Foley catheter placement following demonstration of urethral integrity with a RUG. Consult urology to aid in decision making in any case of urethral or bladder injury.

**Complications**

Contrast extravasation into periurethral or perivesical tissues in the case of urethral or bladder injury, respectively, may cause considerable inflammatory reaction, particularly contrast materials of higher concentrations.

**Conclusions**

RUG and retrograde cystography are the diagnostic procedures of choice to evaluate potential injury to the lower urinary tract. These studies must be carried out in the proper sequence and in a retrograde fashion to avoid missing potential injuries.

**UPPER GENITOURINARY TRACT IMAGING**

Once the lower GU tract has been exonerated with RUG and retrograde cystography, the upper tract (kidneys and ureters)
**Retrograde Cystography**

**A. Normal Examination**

1. **AP view.** Fill the bladder with at least 400 mL (see text for details). The normal bladder has a smooth outer contour and no extravasation of contrast material is seen.

2. **Lateral view.** Obtain a lateral or oblique view to assess for posterior extravasation because the opacified bladder would prevent visualization of this injury on the AP view.

3. **Postvoid film.** Always perform a postvoid film to assess for subtle posterior perforation, especially in the case of penetrating trauma.

**B. Abnormal Examinations**

1. **Extraperitoneal bladder perforation.** Contrast material is seen adjacent to the bladder neck. The bladder itself is displaced rightward by an adjacent hematoma.

2. **Posterior injury.** On this AP film, the bladder appears normal, and no extravasation of contrast material is noted. However, postvoid films are required to rule out posterior injury.

3. **Postvoid film.** The postvoid film demonstrates substantial posterior extravasation of contrast material. This injury would have been missed if only the AP study were performed.

4. **Extraperitoneal bladder rupture.** Extravasated contrast material is seen surrounding the bladder (arrows) but no loops of bowel are outlined. This injury can typically be treated conservatively with a Foley catheter.

5. **Intraperitoneal bladder rupture.** Note the contrast material outlining the paravesical space, tracking up the right paracolic gutter and outlying loops of bowel (arrows). This typically requires operative repair.

6. **Pelvic hematoma.** Note the “open-book” pelvic fracture (arrows). No extravasation of contrast material is seen, however, the pear-shaped bladder indicates the presence of a pelvic hematoma.

*Figure 55.35 Retrograde cystography. AP, Anteroposterior.*
Retrograde Computed Tomographic Cystography

**Figure 55.36** Retrograde computed tomographic cystography. *CT*, computed tomography; *IV*, intravenous.

**Box 55.6** American Association for the Surgery of Trauma Classification System for Renal Injuries

**GRADE 1 INJURIES**
- Hematuria with normal imaging
- Contusions
- Subcapsular, nonexpanding hematoma without parenchymal laceration

**GRADE 2 INJURIES**
- Nonexpanding perinephric hematomas confined to the retroperitoneum
- Renal cortical lacerations less than 1 cm in depth without urinary extravasation

**GRADE 3 INJURIES**
- Renal cortical lacerations greater than 1 cm in depth without urinary extravasation

**GRADE 4 INJURIES**
- Parenchymal laceration extending through the renal cortex/medulla and into the collecting system
- Main renal artery or vein injury with contained hemorrhage
- Segmental infarction without associated laceration

**GRADE 5 INJURIES**
- Shattered or devascularized kidney
- Complete avulsion or thrombosis of the main renal artery or vein
- Avulsion of the ureteropelvic junction


should be evaluated. For adult blunt trauma patients with gross hematuria or hypotension (systolic pressure lower than 90 mm Hg), upper tract imaging is indicated. Data show that 12.5% of such patients will have a major renal injury as compared with only 0.2% of adults with microscopic hematuria and normal blood pressure. In the presence of a major mechanism of injury or other extenuating circumstances, inclusion criteria for upper tract imaging may be broadened. It should be noted that these criteria are not intended for use in pediatric populations, and opinions regarding diagnostic strategies in children remain controversial.

Contrast-enhanced CT is the gold standard for evaluating patients with suspected renal trauma. Parenchymal injuries, vascular injuries, and urinary extravasation can all be readily identified on CT (Fig. 55.37, plates 1 and 2). Traumatic renal injuries are graded on a 5-point scale (Box 55.6). For patients with a mandate for emergency laparotomy, upper tract imaging can be performed in the operating room with an excretory urogram (see Fig. 55.37, plate 3).

Ureteral injuries can also be identified on contrast-enhanced CT imaging; however, it is essential to obtain delayed images (i.e., approximately 10 minutes after the initial intravenous...
Grade 3 renal laceration. This contrast-enhanced CT scan reveals a renal laceration >1 cm in depth without urinary extravasation (arrow) and hence a grade 3 injury.

Grade 4 renal injury. This laceration extends through the renal parenchyma and into the collecting system. Extravasation of contrast material is seen (long arrow), thus making this a grade 4 injury. Note the large perinephric hematoma (short arrows).

Urethral transection. This intraoperative excretory urogram reveals extravasation (arrow) in the upper part of the right ureter consequent to a stab wound. No contrast material is seen in the ureter below the site of injury, a finding indicative of complete transection.

Normal CT urogram. A coronal projection of a CT urogram in the excretory phase demonstrates opacification of the proximal and middle section of the ureters (arrows) without evidence of extravasation of contrast material.

Distal ureter injury. This patient suffered a gunshot wound to his abdomen. After emergency laparotomy and bowel resection, abdominal pain worsened. The initial arterial phase CT demonstrated a large fluid collection in the inferior recesses of the peritoneum (arrow).

Delayed images revealed extravasation of contrast material from the distal end of the ureter (arrow) into the fluid collection. This part of the ureter was not explored during the initial operation.
contrast bolus) (see Fig. 55.37, plates 4.6). This delay allows the contrast material to become concentrated in the kidneys and excreted through the ureters. Ureteral injury can then be identified by extravasation of contrast material into adjacent tissues.

ACKNOWLEDGMENTS

The authors and editors wish to acknowledge the contributions made to previous editions by Robert E. Schneider, MD, Ivan Zbaraschuk, MD, Richard E. Berger, MD, Jerris R. Hedges, MD, Martin Schiff, Jr., MD, Morton G. Glickman, MD, and Geoffrey E. Herter, MD. In addition, the authors wish to acknowledge Dr. Robert E. Schneider, who is board certified in both emergency medicine and urology, for his mentorship.

REFERENCES ARE AVAILABLE AT www.expertconsult.com


Emergency Childbirth

Samreen Vora and Valerie A. Dobiesz

BACKGROUND

Of the more than 3.9 million births in the United States in 2013, 98.6% were delivered in a hospital setting. Although a physician was present for 91.7% of these births, the percentage of these births that occur in the emergency department (ED) setting is unknown.1 Childbirth is a relatively rare occurrence in the ED but ‘complications of pregnancy, childbirth, and puerperium’ is one of the twenty leading primary diagnoses seen in the ED.2 Emergency delivery of an infant continues to be one of the most challenging and stress-inducing procedures facing emergency physicians (EPs). The clinician needs to assess the mother and fetus, prepare for delivery, and anticipate potential difficulties or complications during and after the birthing process.

In institutions where on-site and timely obstetric services are available, the primary duty of the EP may be only to determine that labor is active and delivery imminent. However, in the case of unavailable obstetrics, precipitous delivery, or delayed arrival, the EP may be required to solely manage a patient in active labor. This role includes handling both maternal and neonatal resuscitation.

ANATOMY AND PHYSIOLOGY

Labor is the process by which the fetus is expelled from the uterus. It begins with a sequence of regular and effective uterine contractions that result in effacement and dilation of the cervix.3 Identification of true labor versus false labor is best determined in a dedicated obstetric unit, often with external uterine monitoring.

Labor can also be divided into phases. The latent phase of labor is the period between its onset and when labor becomes active, which generally requires 80% effacement and cervical dilation of greater than 4 cm.4 Active labor is normally divided into three progressive stages. The first stage begins with cervical effacement and dilation and ends when the cervix is completely dilated. In multiparous women, this stage of labor typically lasts approximately 5 to 8 hours, as opposed to 7 to 13 hours in nulliparous women, but with much individual variation.3,4 The second stage of labor begins when dilation of the cervix is complete and ends with delivery of the infant. The duration of this stage is also variable, with a median of 50 to 60 minutes in nulliparas and 15 to 20 minutes in multiparas.3,4 The third stage of labor begins after delivery of the infant and ends with delivery of the placenta. The fourth stage of labor refers to the 1 hour immediately after delivery, the period in which postpartum hemorrhage (PPH) is most likely to occur.1 It is recommended that maternal blood pressure and pulse be recorded immediately after delivery and every 15 minutes for the first hour to rapidly detect any ongoing hemorrhage.1

IDENTIFICATION OF LABOR

It is not uncommon for women to have contractions late in pregnancy, although not all are true or effective labor contractions. Brief Braxton-Hicks contractions of the uterus, usually confined to discomfort in the lower abdominal region and groin, are typically irregular in timing and strength. These pains do not cause any change in the cervix or result in descent of the fetus. Although these pains usually stop spontaneously, they can rapidly convert to true labor contractions; if they occur, a period of observation may be necessary.

True labor is characterized by a regular sequence of uterine contractions with progressively increasing intensity and decreasing intervals between contractions. The interval between contractions gradually diminishes from 10 minutes at the onset of labor to as short as 1 minute or less in the second stage of labor. This should be accompanied by effacement and dilation of the cervix, along with descent of the fetal presenting part into the pelvis. The onset of true labor can be difficult to identify given that patients are far more likely to be at home than in a hospital when labor begins.

Show or bloody show is a sign of approaching labor. The normal mucus plug sealing the cervix discharges as the cervix dilates. Show consists of a small amount of blood-tinged mucus discharged from the vagina and indicates that labor is already in progress or will probably occur during the next several hours to a few days. The mucus color can be pink or brown tinged as a result of minor bleeding. However, if more than a small amount of blood escapes with the mucus plug, an abnormal cause such as placental abruption or placenta previa should be considered. Digital vaginal examination under these circumstances is generally contraindicated.

Spontaneous rupture of membranes can occur during the course of active labor, typically evident by a sudden gush or continuous leakage of a variable amount of clear or slightly turbid vaginal fluid. Rupture of membranes before the onset of labor at any stage of gestation is referred to as premature rupture of membranes (PROM). Term PROM complicates approximately 8% of pregnancies.3,6 In the majority of cases it is followed by the onset of labor and delivery within 5 hours. The most significant maternal risk associated with term PROM is intrauterine infection (chorioamnionitis and endometritis). Fetal risks associated with PROM include umbilical cord compression and ascending infection.5 The ideal management of PROM should be deferred to the consulting obstetrician.7

Membrane rupture occurring before 37 weeks of gestation is called preterm premature rupture of membranes (pPROM). Birth within 1 week generally occurs regardless of management or the clinical findings. The most significant maternal risk related to pPROM is intrauterine infection. The most significant risks to the fetus are complications related to prematurity. Preterm delivery occurs in approximately 12% of all births in the United States and is the leading cause of perinatal morbidity and mortality.5

Accurate diagnosis of PROM is helpful in further management. Suspected PROM should be confirmed by an examination that minimizes the risk of introducing infection. Avoid digital cervical examination unless the patient is in active labor or delivery is imminent. A sterile speculum examination can be performed to look for amniotic fluid extruding from the cervical os or pooling in the posterior fornix, as well as to inspect for fetal or umbilical cord prolapse and assess cervical effacement and dilation.3,4
Amniotic fluid may be differentiated from vaginal fluid by testing the pH of the fluid with Nitrazine paper or similar swab devices (Fig. 56.1). The pH of vaginal fluid is generally 4.5 to 6.0, whereas amniotic fluid has a pH of 7.0 to 7.5. The yellow testing paper turns blue-green to deep blue in the presence of amniotic fluid. With vaginal secretions, the Nitrazine paper remains yellow. False-positive results may occur with blood, semen, bacterial vaginosis, or alkaline urine. If rupture of membranes is documented in the ED, notify the patient’s obstetrician and plan for hospital admission or possible transfer.

The biomarker fetal fibronectin (fFN), a glycoprotein produced by amniocytes and cytotrophoblast found in cervicovaginal secretions, may also be used as a screening tool for preterm labor as an adjunct to maternal/fetal monitoring and pelvic examination. The detection of increased levels of fFN between 22 and 37 weeks gestation can be considered an indicator of preterm birth. If fFN has an excellent negative predictive value (99%) in determining the likelihood of delivery within one week. The test is done with a vaginal swab but must be done before any digital exam. The test is not reliable if the patient has had sexual intercourse within 24 hours, associated bleeding, or has used a gel.

In addition, under EMTALA, a woman in labor is considered unstable for interhospital transfer unless done so at the direction of the patient or a physician who certifies that the benefits outweigh the risks of transfer. In this case, conduct a brief obstetric history, including the onset and frequency of contractions, the presence or absence of bleeding, possible loss of amniotic fluid, previous prenatal care, and due date. Monitor fetal well-being by evaluating the fetal heart rate, particularly immediately after a uterine contraction. In an ED setting, fetal assessment is increasingly done via bedside ultrasonography, which can assess fetal movement and heart rate (Fig. 56.2). The traditional approaches of using external Doppler or auscultation are also acceptable. The normal baseline fetal heart rate is 110 to 160 beats/min.

**EVALUATION OF LABOR**

When a woman is initially seen with contractions, the general condition of both the fetus and mother should be assessed quickly. Under the federal Emergency Medical Treatment and Labor Act (EMTALA), hospitals with an ED must conduct an appropriate screening examination to rule out true labor.
presenting part can be felt through the cervix on sterile vaginal examination. In longitudinal lies, the presenting part is the fetal head, the buttocks (breech), or the feet (footling breech). With a transverse lie, the presenting part is the shoulder. The presentation can be cephalic, breech, shoulder, or compound (a fetal extremity with another presenting part). All presentations except cephalic are considered malpresentations.

Cephalic presentations are classified by the bony leading landmark of the fetal skull. Ordinarily, the head is sharply flexed so that the occipital fontanelle is the presenting part. This is referred to as a vertex or occiput presentation. Less commonly, the neck is fully extended and the face is foremost in the birth canal; this is termed face presentation. A partially flexed or partially extended neck position results in sinciput and brow presentations, respectively. Sinciput and brow presentations associated with preterm infants are almost always unstable and convert to either an occiput or face presentation as labor progresses.

Breech presentations are classified as frank, complete, or incomplete (Fig. 56.4). A fetus presenting with the buttocks and hips flexed and the legs extended is termed a frank breech. Presentation of the buttocks with flexion of the fetal hips and knees results in a complete breech presentation. When one or both feet or knees are lowermost in the canal, an incomplete or footling breech results.

At or near term, 97% of fetuses will be vertex and 3% will be breech. The incidence of breech delivery is approximately 25% at 28 weeks, 17% at 30 weeks, and 11% at 32 weeks.

Position refers to the relationship of the presenting part to the maternal pelvis and may be either left or right. The occiput is the reference point in cephalic presentations, whereas the sacrum is the determining part in breech presentations. The vertex occiput anterior position is the most common and is considered normal. The other position is occiput posterior.

**LABOR MOVEMENTS: VERTEX PRESENTATION**

Full dilation of the cervix signifies the second stage of labor and heralds delivery of the infant. Typically, the patient begins to bear down, which coincides with descent of the presenting part. Uterine contractions may last 1.5 minutes and recur after a resting phase of less than 1 minute. Delivery of a vertex-presenting infant usually occurs spontaneously. The role of the clinician or attendant is principally to provide control of the birth process by preventing forceful, sudden expulsion or extraction of the infant with resultant fetal or maternal injury.

The mechanism of labor in vertex presentations consists of engagement of the presenting part, flexion, descent, internal rotation, extension, external rotation or restitution, and expulsion (Fig. 56.5). These are often referred to as cardinal movements of labor. The mechanism of labor is determined by the pelvic dimensions and configuration, the size of the fetus, and the strength of the uterine contractions. Essentially, the fetus will follow the path of least resistance by adaptation of the smallest achievable diameter of the presenting part to the most favorable dimensions and contours of the birth canal.

The sequence of movements in vertex presentations is as follows:

1. **Engagement** refers to the mechanism by which the greatest transverse diameter of the head, the biparietal diameter in occiput presentations, passes through the pelvic inlet. A fetus is engaged when the presenting part is at 0 station. In a primiparous patient, it usually occurs in the last 2 weeks.
of pregnancy; in a multiparous patient, it can occur at the onset of labor.

2. **Flexion** of the head is necessary to minimize the presenting cross-sectional diameter of the head during passage through the smallest diameter of the bony pelvis. In most cases, flexion is necessary for both engagement and descent and occurs passively.

3. **Descent** is the downward passage of the fetal presenting part. It is gradually progressive but is not necessarily continuous. Descent is affected by uterine and abdominal contractions, as well as by straightening and extension of the fetal body.

4. **Internal rotation** occurs with descent and is necessary for the head or presenting part to traverse the ischial spines. This movement essentially turns the head such that the occiput gradually moves from its original, more transverse position anteriorly toward the symphysis pubis or, less commonly, posteriorly toward the hollow of the sacrum. This is known as occiput anterior or occiput posterior, respectively.

5. **Extension** occurs as the flexed head reaches the anteriorly directed vaginal introitus. The occiput reaches the inferior aspect of the pubic symphysis. The head is born by further extension as it rotates around the pubic symphysis and the chin pass successively over the anterior margin of the perineum. Immediately after its birth, the head drops downward such that the chin lies over the maternal anal region.

6. **External rotation** or restitution is return of the head to the correct anatomic position with respect to the fetal torso. It follows delivery of the head as it rotates to the transverse position that it occupied at engagement. This is also a passive movement.

7. **Expulsion** of the remainder of the fetal body then occurs. The shoulders descend in a path similar to that traced by the head (i.e., rotating anteroposteriorly for delivery). First, the anterior shoulder is delivered beneath the symphysis pubis followed by the posterior shoulder across the perineum.

### LABOR MOVEMENTS: BREECH PRESENTATION

The mechanism of labor for breech presentations varies (see Fig. 56.4). The widest diameter that is engaged is the bitrochanteric diameter. Usually, the hips engage in one of the
oblique diameters of the pelvic inlet. As descent occurs, the anterior hip generally descends more rapidly than the posterior hip. Internal rotation occurs as the bitrochanteric diameter assumes the anteroposterior (AP) position. Lateral flexion takes place as the anterior hip catches beneath the symphysis pubis, which allows the posterior hip to be born first. The infant’s body then rotates to allow engagement of the shoulders in an oblique orientation. Gradual descent occurs, with the anterior shoulder rotating to bring the shoulders into the AP diameter of the outlet. The anterior shoulder follows lateral flexion to appear beneath the symphysis, with the posterior shoulder being delivered first as the body is supported. The head tends to engage in the same diameter as the shoulders. Subsequent flexion, descent, and rotation of the head occur to bring the posterior portion of the neck under the symphysis pubis. The head is then born in flexion.

Breech delivery is associated with a greater incidence of prematurity, prolapsed cord, and increased perinatal morbidity and mortality. Cesarean section can reduce the morbidity and mortality associated with breech delivery, but a planned vaginal delivery may be the method of choice in carefully selected cases and may be beneficial in low resource settings. The emergency clinician should be cognizant of the imminent vaginal delivery of a breech infant in any presentation: frank, complete, or footling. However, a breech presentation is always problematic for any clinician, even under the best of circumstances. It is not expected that the EP will always be able to successfully deliver a breech presentation.

Types

There are three types of vaginal breech deliveries. Spontaneous breech is a breech delivery in which the infant is delivered spontaneously without any manipulation or traction other than supporting the infant. This form of delivery is rare with term infants, and there is little associated traumatic morbidity. Partial breech extraction occurs when the infant is delivered spontaneously as far as the umbilicus and the remainder of the body is extracted. Total breech extraction occurs when the entire body of the infant is extracted by the clinician.

Similar to cephalic presentations, the role of the clinician is to assist the mother in the birthing process and allow maternal expulsive efforts to effect delivery of the infant. Premature or aggressive assistance or traction can significantly increase the risk for fetal or maternal morbidity.

To perform any vaginal breech delivery, the birth canal must be sufficiently large to allow passage of the fetus without trauma and the cervix must be completely effaced and dilated. If these conditions do not exist, cesarean section (Videos 56.1 and 56.2) is indicated. To ensure full cervical dilation in a footling or complete breech, it is important that the feet, legs, and buttocks advance through the introitus to the level of the fetal umbilicus before the clinician intervenes and further delivery is attempted. The mere appearance of the feet through the vulva is not in itself an indication to proceed with delivery. This may be a footling presentation through a cervix that is not completely dilated. In this case there may be time to transfer the patient to the labor and delivery suite, preferably in the knee-chest position to minimize the risk for cord compression. Similarly, if the breech is frank, cervical dilation and outcome are improved if the infant is allowed to deliver to the level of the umbilicus. Before this, as with complete and footling presentations, there may be time to safely transfer the mother to the labor and delivery area. Tocolytics such as subcutaneous terbutaline may be considered to inhibit labor until such patients can be safely transferred.

VAGINAL EXAMINATION

Cleanse the perineum with an antiseptic and use a sterile lubricant to decrease potential contamination. A sterile vaginal (not speculum) examination is performed to identify the fetal presentation and position and assess the progress of labor, except in cases of suspected bleeding. During the digital examination, take care to avoid the anal region and potential fecal contamination. Assess cervical effacement and dilation, as well as fetal station, presentation, and position. Do not withdraw the finger from the vagina until the examination is complete. The number of vaginal examinations during labor correlates with infectious morbidity, especially in cases of early membrane rupture.

Cervical effacement refers to the process of cervical thinning that occurs before and during the first stage of labor as the cervical canal shortens from a length of approximately 2 cm to a circular opening with almost no length remaining (Fig. 56.6). Effacement is expressed as a percentage from 0% (uneffaced and thick) to 100% (completely effaced). Assess the degree of cervical effacement by palpation and determine the palpated length of the cervical canal in comparison to that of an uneffaced, or normal, cervical canal.

Determine cervical dilation by estimating the average diameter of the internal cervical os. Sweep the examining finger from the cervical margin on one side across the cervical os to the opposite margin. Express the diameter in centimeters. Ten centimeters constitutes full cervical dilation. A diameter of less than 6 cm can be measured directly. A cervix that accommodates one index finger is 1 cm, and one that accommodates two fingers is dilated approximately 3 cm. For a diameter greater than 6 cm, it is frequently easier to determine the width of the remaining cervical rim and subtract twice that measurement from 10 cm. For example, if a 1-cm rim is felt, dilation is 8 cm.
Station refers to the level of the presenting fetal part in the birth canal relative to the ischial spines, which lie halfway between the pelvic inlet and the pelvic outlet (Fig. 56.7). Zero station is used to denote that the presenting part is at the level of the ischial spines. When the presenting part lies above the spines, the distance, estimated in centimeters ranging from 1 to 5, is stated in negative figures (−5, −4, −3, −2, −1). Below the ischial spines, the presenting fetal part passes +1, +2, +3, +4, and +5 stations to delivery. This measurement is made by simple palpation. The ischial spines can be palpated at roughly the 8 and 4 o’clock positions on the vaginal examination.

Three maneuvers are used to determine fetal presentation and position. First, introduce two fingers into the vagina and advance them to the presenting part to differentiate face, vertex, and breech presentations. In vertex presentations, move your fingers up behind the symphysis pubis and then sweep them posteriorly over the fetal head toward the maternal sacrum to identify the course of the sagittal suture. Define the positions of the two fontanelles, which are located at opposite ends of the sagittal sutures, by palpation. The anterior fontanel is diamond shaped; the posterior fontanel is triangular. In breech presentations, the fetal sacrum is the point of reference, whereas in face presentations the fetal chin is used.

FETAL WELL-BEING

Auscultation

Make the initial determination of fetal well-being by assessing the fetal heart rate with a fetoscope, bedside ultrasound, or a fetal Doppler ultrasound device. Place the device firmly on the maternal abdominal wall overlying the fetal thorax and reposition it until fetal heart tones are heard. When a Doppler device is used, apply a conducting gel to the abdominal wall to interface with the Doppler receiver. To avoid confusion of the maternal and fetal heart sounds, palpate the maternal pulse as the fetal heart rate is auscultated.

The normal baseline fetal heart rate is 110 to 160 beats/min and varies considerably from a baseline measured for a minimum of 2 minutes in a 10-minute segment of time. Rates above or below this range may indicate fetal distress. Accelerations in the fetal heart rate lasting longer than 10 seconds and less than 2 minutes commonly occur during labor and are probably a physiologic response to fetal movement. Persistent fetal tachycardia occurs most frequently in response to maternal fever or amnionitis but may also indicate fetal compromise.

As with brief accelerations in the fetal heart rate, a gradual decrease in the fetal heart rate in association with a uterine contraction with an onset to nadir of 30 seconds or more (with the nadir coinciding with the peak of contraction) is termed an early deceleration. Such decelerations are physiologic and probably the result of vagal nerve stimulation secondary to compression of the fetal head. Decelerations that occur independent of uterine contractions, are abrupt, or last between 15 seconds and 2 minutes are known as variable decelerations. Variable decelerations are relatively common, can be classified according to their severity, and may be temporarily corrected by maternal repositioning. Late decelerations are those that are delayed in timing with respect to a contraction, with the nadir of the deceleration occurring after the peak of the contraction. Late decelerations can be an ominous sign and may represent cord compression or uteroplacental insufficiency and may necessitate emergency delivery.

Changes in the fetal heart rate that indicate fetal distress are usually evident immediately after a uterine contraction, and therefore the fetal heart rate is optimally assessed at this time. Formal monitoring of labor should ideally be performed in an obstetrics unit (Fig. 56.8).

Management of Fetal Distress

Definitive evaluation of fetal distress should be performed in the obstetric unit by the delivery team. There is no role for sophisticated fetal monitoring in the ED. In the absence of a dedicated obstetric unit, transfer to another hospital is the only option, albeit a less than ideal one. An EP working in an ED without adequate obstetric backup can do little to effect a positive outcome in high-risk situations. Eclampsia, bleeding, and abnormal fetal presentation may be identified, but the EP needs to focus attention on maternal well-being while expediting transfer, referral, or both. The EP has limited options for managing fetal distress.

If fetal distress is suspected on the basis of the resting fetal heart rate or changes after contractions, change the maternal position, typically into the left lateral decubitus position, and reevaluate. Administer supplemental oxygen to the mother to optimize fetal oxygenation. In the absence of bleeding, perform a vaginal examination to rule out the possibility of umbilical cord prolapse. Cord prolapse usually occurs at the same time as rupture of the membranes and is diagnosed by palpation of the umbilical cord on vaginal examination or by visualization of the cord protruding through the introitus.
Figure 56.8 Deceleration patterns of the fetal heart rate (FHR). A, Early deceleration caused by head compression. B, Late deceleration caused by uteroplacental insufficiency. C, Variable deceleration caused by cord compression. Note: fetal monitoring is not done in the emergency department; this figure is supplied for completeness. (Modified from Lowdermilk DL, Perry SE, Bobak IM, editors: Maternity and women’s health care, ed 6, St. Louis, 1997, Mosby.)
Cord prolapse is associated with several risk factors such as malpresentation, grand multiparity, and prematurity, as well as iatrogenic causes.\(^{22-24}\)

Management of cord prolapse is directed at sustaining fetal life until delivery is accomplished. In situations with the cord prolapse and evidence of fetal distress, unless immediate delivery is feasible or the fetus is known to be dead, prepare for an emergency cesarean section. If immediate obstetric services are not available, four temporizing measures can be undertaken. (1) Place the patient in the knee-chest or deep Trendelenburg position, and keep the patient in this position until delivery.\(^{22,23,25}\) (2) Minimize compression of the umbilical cord by inserting a sterile gloved hand and exerting manual pressure in the vagina to lift and maintain the presenting part away from the prolapsed cord. (3) After manual elevation of the presenting part, instill 500 to 700 mL of saline into the bladder to raise the part and maintain cord decompression. Once the bladder is filled, remove the vaginal hand.\(^{22,25}\) (4) Tocolytic therapy can be administered to decrease uterine contractions and improve fetoplacental perfusion in cases of fetal bradycardia or pathological decelerations.\(^{23,25}\) Unfortunately, outcomes of true obstetric emergencies managed solely in the ED are often bleak and essentially out of the hands of the EP.

**Tocolytic Therapy**

Tocolytics are drugs used to suppress uterine contractions for women in preterm labor and are rarely indicated or instituted in the ED. However, under extremely limited circumstances and preferably under obstetric guidance, this intervention may be initiated in the ED. The goal of delaying delivery is to enable administration of antenatal corticosteroids for lung maturation and magnesium sulfate (MgSO\(_4\)) for neuroprotection, as well as to permit maternal transport (if indicated) to a tertiary facility.\(^{32-34}\) Preterm birth is a major contributor to perinatal mortality and morbidity globally.\(^{10}\)

Before instituting pharmacologic tocolytic therapy for either preterm labor or fetal distress, initiate basic maneuvers to improve maternal and fetal status. Because uterine hypoxia may induce uterine contractions, administer supplemental oxygen and infuse 500 mL of crystalloid intravenously. Place the mother in the left lateral decubitus position to improve uterine perfusion.\(^{32}\) Because uterine, cervical, or urinary tract infections account for 20% to 40% of cases of preterm labor, search for a specific cause and treat infections appropriately.\(^{32}\)

If the contractions persist and cervical changes are documented despite these basic interventions, consider pharmacologic therapy.\(^{1,31}\) Although tocolytic agents are commonly used and have been shown to prolong pregnancy by 2 to 7 days, data suggest that tocolysis does not improve neonatal outcomes.\(^{7,28,31}\) In addition, none of the commonly used tocolytic medications are approved for use by the U.S. Food and Drug Administration (FDA) for this indication. General contraindications to tocolytic therapy include severe preeclampsia, placental abruption, intrauterine infection, advanced cervical dilation, and evidence of fetal compromise or placental insufficiency.\(^{31}\)

There are no clear first-line tocolytic agents to manage preterm labor (Table 56.1). Clinical circumstances and preferences should dictate treatment. Agents such as calcium channel blockers (CCBs) and the prostaglandin inhibitor indomethacin have shown varying efficacy in clinical trials.\(^{2,20,11}\) The most commonly used tocolytic agents in the United States are the CCB nifedipine, MgSO\(_4\), and the \(\beta\_2\)-receptor agonist terbutaline.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>End Point</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indomethacin</td>
<td>50–100 mg PO</td>
<td>May repeat 25–50 mg PO q 6 hr for a maximum of 48 hr</td>
</tr>
<tr>
<td></td>
<td>or 50 mg PR</td>
<td></td>
</tr>
<tr>
<td>Terbutaline</td>
<td>0.25 mg SC</td>
<td>May repeat q 20–60 min</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cessation of uterine contractions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Intolerable maternal side effects</td>
</tr>
<tr>
<td>Magnesium</td>
<td>4–6 g IV over</td>
<td>Cessation of uterine contractions</td>
</tr>
<tr>
<td>sulfate</td>
<td>20 min followed</td>
<td>Signs of magnesium toxicity (e.g., respiratory depression, hypotension,</td>
</tr>
<tr>
<td></td>
<td>by an infusion</td>
<td>somnolence)</td>
</tr>
<tr>
<td></td>
<td>at 1–3 g/hr IV</td>
<td></td>
</tr>
<tr>
<td>Nifedipine(^*)</td>
<td>10 mg PO</td>
<td>May repeat q 15–20 min</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Cessation of uterine contractions</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Harmful maternal side effects, e.g., hypotension</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Maximum dose, 40 mg</td>
</tr>
</tbody>
</table>

\(^*\)Variable doses have been used.
IV, Intravenously; PO, orally; PR, per rectum; q, every; SC, subcutaneously.

**TABLE 56.1 Tocolytics for the Emergency Management/Cessation of Preterm Labor**

CCBs are the tocolytics preferred by the World Health Organization (WHO).\(^{13}\)

**\(\beta\_2\)-Receptor Agonists**

The most commonly used \(\beta\_2\)-adrenergic tocolytic agent is terbutaline. The \(\beta\_2\)-mimetic agents react with adrenergic receptors to reduce intracellular ionized calcium levels and prevent the activation of myometrial contractile proteins.\(^3\) Although terbutaline stimulates \(\beta\_2\)-receptors primarily, it has some \(\beta\_1\)-activity, and this is responsible for its cardiovascular side effects.

Terbutaline is commonly used for the treatment of preterm labor.\(^{1,10}\) Administer terbutaline subcutaneously in a 0.25-mg dose and repeat it every 20 to 60 minutes until contractions cease or intolerable maternal side effects occur.\(^{31,35,36}\) In 2011 the FDA required the addition of a *Boxed Warning and Contraindication* (“black box warning”) against the use of injectable terbutaline for the prevention or prolonged treatment of preterm labor beyond 48 to 72 hours because of adverse effects, including maternal death.\(^7\) Use terbutaline with caution in patients with cardiovascular disease, hypertension, hyperthyroidism, diabetes, and seizures and in those taking other sympathomimetic amines.\(^{31,36}\) The general clinical side effects of arrhythmias, myocardial ischemia, and pulmonary edema are related to its
inherent activity as a β-mimetic drug. Treatment of the majority of side effects is supportive; severe cardiovascular effects may be treated with β-blocking agents.35,38

MgSO₄

MgSO₄ is not approved in the United States for use as a tocolytic agent and has been shown to be ineffective at delaying birth or preventing preterm birth, has no apparent advantages as a tocolytic agent, and it may be associated with an increased risk of fetal, neonatal, or infant mortality.18 Even though the use of MgSO₄ as a tocolytic has fallen out of favor, benefits have been shown for antenatal use for its fetal neuroprotective effects.40–41 The mechanism of action is not fully understood; magnesium probably decreases myometrial contractility through its role as a calcium antagonist. Its fetoprotective effects probably result from noncompetitive antagonism of the N-methyl-D-aspartate receptor or through antiapoptosis and prevention of neuronal cell loss.42

MgSO₄ is generally administered at a loading dose of 4 g intravenously over a period of 20 to 30 minutes, followed by a maintenance intravenous infusion beginning at 1 g/hr.11,41 Infusion of MgSO₄ often produces sweating, warmth, and flushing. Rapid parenteral administration may cause transient nausea, vomiting, headache, or palpitations.11,44 The major side effect of magnesium therapy is impairment of the muscles of respiration with subsequent respiratory arrest, an effect not usually seen until the serum magnesium level exceeds 10 mEq/L. At levels of 12 mEq/L or greater, respiratory arrest may occur.18

The first sign of magnesium toxicity, a decrease in the patellar reflex, typically occurs as serum magnesium levels exceed 4 mEq/L, with loss of the reflex as levels increase further. The dosing and ongoing maintenance of magnesium therapy should be guided by the clinical status of the patient rather than by laboratory values. Monitor the patellar reflex throughout therapy. Because magnesium is almost totally excreted by the kidney, it is contraindicated in the presence of renal failure. Monitor urinary output and renal function throughout therapy. If respiratory depression develops, inject 10 mL of a 10% solution of calcium gluconate or calcium chloride over a 3-minute period as an antidote. For severe respiratory depression or arrest, prompt endotracheal intubation may be lifesaving.3

Calcium Channel Blockers

Calcium antagonists inhibit the influx of calcium ions through the muscle cell membrane and reduce uterine vascular resistance. The decrease in intracellular calcium also results in decreased myometrial activity. CCBs, mainly nifedipine, have demonstrated benefits in postponement of birth compared to β-mimetics with respect to prolongation of pregnancy, serious neonatal morbidity, and maternal adverse effects. CCBs may have some benefits over oxytocin receptor antagonists and magnesium sulfate, although oxytocin receptor antagonists had fewer maternal adverse effects. No difference has been noted in perinatal mortality.30

Although dosing regimens vary, nifedipine is frequently given as an initial loading dose of 30 mg orally and then 10 to 20 mg every 4 to 6 hours.31,33 The CCB–induced decreased vascular resistance can lead to maternal hypotension and thus decreased uteroplacental perfusion.45

Cyclo-Oxygenase Inhibitors

Cyclo-oxygenase (COX) inhibitors inhibit uterine contractions, are easily administered and appear to have few maternal side effects, although adverse effects have been reported in the fetus and newborn.46 Indomethacin is administered as an initial oral or rectal dose of 50 to 100 mg followed by 25- to 50-mg doses every 6 hours for a total of 48 hours.47 Ketorolac can be used as an alternative with a 60-mg intramuscular loading dose, followed by 30 mg intramuscularly every 6 hours for 48 hours.47 Side effects include oligohydramnios and premature closure of the ductus arteriosus.

Steroids

In addition to tocolytics, patients in preterm labor at less than 34 weeks’ gestation are candidates for antenatal steroid administration. This may reduce the incidence of neonatal respiratory distress syndrome, intraventricular hemorrhage, and necrotizing enterocolitis.31 Appropriate patients should receive betamethasone, 12 mg intramuscularly every 24 hours for a total of two doses, or dexamethasone, 6 mg intramuscularly every 12 hours for a total of four doses. Dexamethasone may have benefits compared to betamethasone such as fewer intraventricular hemorrhages and decreased lengths of stay in the neonatal intensive care unit (NICU). The intramuscular route may be more advantageous than the oral route.47

VAGINAL BLEEDING DURING THE THIRD TRIMESTER

Bleeding during the third trimester should always be considered an emergency because shock may occur within minutes. Prepare for the two most typical causes of bleeding in late gestation, placenta previa and placental abruption (Fig. 56.9). Placenta previa refers to implantation of the placenta in the lower uterine segment with varying degrees of encroachment on the cervical os. Placenta previa is classically characterized by vaginal bleeding with little or no abdominal or pelvic pain. Abruptio placentae refers to separation of the placenta from its site of implantation in the uterus before delivery of the fetus. Although the clinical signs and symptoms with placental abruption can vary considerably, abruptio placentae is typically associated with varying degrees of abdominal pain and uterine irritability or contractions.3,48

Stabilization of a patient with third-trimester bleeding should be initiated with large-bore intravenous access. Blood should be drawn for a complete blood count with platelets and a type and crossmatch. If abortion is suspected, clotting studies, including a fibrinogen level and a toxicology screen for cocaine, may be indicated because of the association of abortion with disseminated intravascular coagulation and cocaine abuse, respectively. Until the diagnosis of placenta previa is excluded, digital vaginal examination is contraindicated because of the possibility of tearing or dislodging a placenta previa, which may result in profuse, potentially fatal hemorrhage.1 The simplest and most precise method of placental localization is by transabdominal ultrasound, which has an accuracy in locating placenta previa of approximately 96%. In contrast, ultrasonography has limited sensitivity in detecting abruptio placentae, with a reported negative predictive value of between 63% and 88%.3 Therefore negative findings on
Abnormalities of Placental Implantation

Marginal placenta previa

Partial placenta previa

Total (central) placenta previa

Figure 56.9 A, Placenta previa.

Continued
Placental Abruption

External bleeding

Internal (concealed) bleeding

Obstruction of cervix by presenting part

Section through placenta in premature separation showing nodular ischemia and infarction above clots.

Figure 56.9, cont’d B, Abruptio placentae. (Netter illustrations from www.netterimages.com. © Elsevier Inc. All rights reserved.)
PROCEDURE

Technique for Uncomplicated Delivery

Although complete sterility is not always possible especially during emergency conditions, when time permits, use sterile technique and equipment. Wear sterile gloves, a gown, mask, and eye protection. Clean the perineum and vulva as for a vaginal examination and drape with sterile towels so that only the immediate area about the vulva is exposed. Be careful to avoid fecal contamination of the infant or perineum.

Ideally, place the patient on a delivery table in the dorsal lithotomy position to increase the diameter of the pelvic outlet (Fig. 56.10, step 1). Alternatively, position the patient on a stretcher with her hips and knees partially flexed, her thighs abducted, and the soles of her feet placed firmly on the stretcher. Additional personnel may assist in keeping the patient in this position. If the foot of the bed cannot be removed, enhance the delivery position by placing the underside of a bedpan under the patient's buttocks to provide additional space between the bed and the perineum.

Spontaneous Vertex Delivery

Spontaneous delivery of a vertex-presenting infant is divided into three phases: delivery of the head, delivery of the shoulders, and delivery of the body and legs.

Delivery of the Head

Anticipate delivery when the presenting part reaches the pelvic floor. With each contraction, the perineum bulges further and the vulvovaginal opening becomes more and more dilated by the fetal head. Just before delivery, crowning occurs, which is when the head is visible at the vaginal introitus and the widest portion, or the biparietal diameter of the head, distends the vulva.

Gentle, gradual, controlled delivery is desirable. Avoid explosive delivery of the head. Once the fetal head distends the vaginal introitus to 5 cm or more during a contraction, place the palm of one hand over the occipital area and provide gentle pressure to control delivery of the head (see Fig. 56.10, step 2). With the other hand preferably draped with a sterile towel to prevent contamination from the anus, exert upward pressure on the chin of the fetus through the perineum just in front of the coccyx in a modified Ritgen maneuver (Fig. 56.11). This maneuver extends the neck at the proper time such that the smallest diameter of the head passes through the introitus and over the perineum to protect the maternal perineal musculature. It is not uncommon for the vagina and perineum to tear with expulsive maternal effort during delivery of the head. If maternal expulsive efforts are insufficient to allow delivery of the head, an episiotomy may be considered at this time.

Gently support the head during subsequent delivery of the forehead, face, chin, and neck. With delivery of the neck, pass a finger around the infant's neck to determine whether it is encircled by one or more coils of the umbilical cord. If a loop of cord is felt, loosen it carefully and gently slip it over the infant's head (Fig. 56.12). If this cannot be done easily, clamp the cord doubly, cut the cord between the clamps and promptly deliver the infant.

Delivery of the Shoulders

Just before external rotation the head usually falls posteriorly, almost bringing it into contact with the mother's anus. If maternal defecation occurs, avoid fecal contamination of the fetus. As rotation takes place, the head assumes a transverse position and the transverse diameter of the thorax rotates into the AP diameter of the pelvis. In most cases the shoulders are born spontaneously. Aid delivery by grasping the sides of the head and exerting gentle downward (posterior) traction until the anterior shoulder appears beneath the symphysis pubis (see Fig. 56.10, step 3). Gently lift the head upward to aid in delivery of the posterior shoulder (see Fig. 56.10, step 4). The remainder of the body usually follows without difficulty.

If delivery of the body is delayed after the shoulders have been freed, assist by providing moderate traction on the exposed fetal body. To avoid injury to the brachial plexus, do not hook the fingers in the axilla during delivery. Always exert traction in the direction of the long axis of the infant. If traction is applied obliquely, bending of the neck and excessive stretching of the brachial plexus may occur.

Clearing the Airway

Once the head has been delivered, quickly wipe the infant's face and mouth. Routine suctioning is not indicated for vigorous newborns. Although it may be counterintuitive, current recommendations no longer advise routine oropharyngeal and nasopharyngeal suctioning of infants with meconium staining by amniotic fluid. Studies have shown that this practice offers no benefit if the infant is vigorous. A vigorous infant is one who has strong respiratory effort, good muscle tone, and a heart rate greater than 100 beats/min. Routine intubation for tracheal suction of infants with meconium staining is also no longer recommended. Per 2015 American Heart Association (AHA) guidelines, infant resuscitation should be initiated with air (21% oxygen at sea level), particularly for preterm infants less than 35 weeks, then titrated as necessary.

Clamping the Cord

The infant should be positioned at or slightly below the level of the vaginal introitus during clamping of the cord. Cut the umbilical cord with scissors between two Kelly clamps placed 4 to 5 cm from the infant's abdomen (see Fig. 56.10, step 6). Delayed clamping of the umbilical cord, no earlier than 1 minute after birth, in infants who do not require resuscitation can improve the short- and long-term hematologic and iron status of full-term infants. Later, apply an umbilical cord clamp 2 to 3 cm from the infant's abdomen. Collect blood samples from the placental end of the cord for infant serology, including Rh determination.

After cutting the umbilical cord, evaluate the infant and, if necessary, initiate resuscitation as described in the later section on The Newborn. Rapidly assess the infant by determining the following:
- Was the baby born after a full-term gestation?
- Does the baby have good muscle tone?
- Is the baby breathing or crying spontaneously?

If the answer to these questions is “yes,” the baby will probably not need resuscitation.
**Spontaneous Vertex Delivery**

1. Place the patient in the dorsal lithotomy position. Anticipate delivery when *crowning* occurs; the fetal head will be visible at the vaginal introitus.

2. Place one hand over the occiput and provide gentle pressure to control delivery of the head. Use your other hand to exert pressure on the chin of the fetus through the perineum (the modified Ritgen maneuver).

3. Apply gentle downward (posterior) traction until the anterior shoulder appears beneath the symphysis pubis.

4. Gently lift the head upward to aid in delivery of the posterior shoulder.

5. The remainder of the body usually delivers spontaneously without difficulty. Routine suctioning is not indicated for vigorous newborns.

6. Position the infant at or slightly below the level of the vaginal introitus during clamping of the cord. Cut the cord with scissors between two Kelly clamps placed 4 to 5 cm from the infant’s abdomen.

7. Deliver the placenta (placental separation usually occurs within 5 minutes after delivery). Ask the mother to bear down to aid in delivery.

---

*Figure 56.10* Spontaneous vertex delivery. (Adapted from *Ferri’s clinical advisor 2013*. St. Louis, 2013, Mosby.)
Emergency Childbirth

1199

in separation of the cord from the placenta or uterine inversion with potentially catastrophic hemodynamic consequences. Examine the placenta for completeness and save it for later evaluation by the obstetrician.3,19

Examine the vulva, vagina, and cervix for traumatic lacerations. Cervical lacerations most typically occur at the 9 or 3 o’clock position; vaginal lacerations typically occur at the point of the ischial spines. The most common areas for lacerations are the vagina, hymen, and perineum. Based on the availability of resources and the clinical setting, definitive repair of uncomplicated or minor obstetric lacerations may be performed in the ED by the EP, but this is often done by a consultant obstetrician.

After delivery of the placenta, the primary mechanism by which hemostasis is achieved at the placental site is through myometrial contraction. Agents such as oxytocin, methylergonovine, and ergonovine may be used to stimulate myometrial contraction. In an effort to prevent uterine atony and subsequent bleeding, oxytocin is often administered after delivery of the placenta.

Oxytocin is the most commonly used oxytocic drug and is usually given by continuous intravenous infusion. Add 20 units of oxytocin to 1 L of normal saline and administer the solution at a rate of 10 mL/min for several minutes until the uterus remains firmly contracted and bleeding is controlled. At this point, reduce the infusion rate to 1 to 2 mL/min.3,19 Alternatively, ergot derivatives such as methylergonovine maleate, 0.2 mg, or ergonovine maleate, 0.2 mg, may be given intramuscularly.3 Because of their vasoconstrictive properties, ergot preparations are relatively contraindicated in patients with hypertension, including pregnancy-associated hypertension or preeclampsia.3 The synthetic prostaglandin misoprostol can also be given as an 800- to 1000-µg rectal dose. In the United States, oxytocic agents are not generally administered before delivery of the placenta because of concerns that the resultant uterine contraction may entrap the placenta or trap an undiagnosed twin within the uterus.1,19 Even when oxytocics are administered, the hour after delivery of the placenta is the time during which PPH secondary to uterine atony is most

Delivery of the Placenta

Placental separation usually occurs within approximately 5 minutes after delivery of the infant, although it may take longer. It can be recognized by the following signs:
1. The uterus becomes globular and firmer as it contracts.
2. There is a sudden gush of blood as the placenta separates from the uterine wall.
3. The umbilical cord lengthens and protrudes further out of the vagina.

Ask the mother to bear down; the increased intraabdominal pressure produced by this maneuver may be enough to effect complete expulsion of the placenta (see Fig. 56.10, step 7). If maternal force alone is insufficient, aid in the delivery of the placenta. After ensuring that the uterus is firmly contracted and placental separation has occurred, use one hand to exert gentle pressure through the abdominal wall to lift the uterine fundus cephalad while keeping the umbilical cord slightly taut with the other hand (Fig. 56.13). Repeat this maneuver until the placenta reaches the introitus. At this time, stop uterine pressure and gently lift the placenta upward and out of the vagina. Such maneuvers may result in separation of the cord from the placenta or uterine inversion with potentially catastrophic hemodynamic consequences. Examine the placenta for completeness and save it for later evaluation by the obstetrician.3,19

Examine the vulva, vagina, and cervix for traumatic lacerations. Cervical lacerations most typically occur at the 9 or 3 o’clock position; vaginal lacerations typically occur at the point of the ischial spines. The most common areas for lacerations are the vagina, hymen, and perineum. Based on the availability of resources and the clinical setting, definitive repair of uncomplicated or minor obstetric lacerations may be performed in the ED by the EP, but this is often done by a consultant obstetrician.

After delivery of the placenta, the primary mechanism by which hemostasis is achieved at the placental site is through myometrial contraction. Agents such as oxytocin, methylergonovine, and ergonovine may be used to stimulate myometrial contraction. In an effort to prevent uterine atony and subsequent bleeding, oxytocin is often administered after delivery of the placenta.

Oxytocin is the most commonly used oxytocic drug and is usually given by continuous intravenous infusion. Add 20 units of oxytocin to 1 L of normal saline and administer the solution at a rate of 10 mL/min for several minutes until the uterus remains firmly contracted and bleeding is controlled. At this point, reduce the infusion rate to 1 to 2 mL/min.3,19 Alternatively, ergot derivatives such as methylergonovine maleate, 0.2 mg, or ergonovine maleate, 0.2 mg, may be given intramuscularly.3 Because of their vasoconstrictive properties, ergot preparations are relatively contraindicated in patients with hypertension, including pregnancy-associated hypertension or preeclampsia.3 The synthetic prostaglandin misoprostol can also be given as an 800- to 1000-µg rectal dose. In the United States, oxytocic agents are not generally administered before delivery of the placenta because of concerns that the resultant uterine contraction may entrap the placenta or trap an undiagnosed twin within the uterus.1,19 Even when oxytocics are administered, the hour after delivery of the placenta is the time during which PPH secondary to uterine atony is most

Figure 56.11 For controlled and gentle delivery of the head, use the modified Ritgen maneuver. (From Seils A, Noujaim S, Davis K, et al, editors: Williams obstetrics, ed 22, New York, 2005, McGraw-Hill Medical.)

Figure 56.12 As the head is delivered, use a finger to determine whether the umbilical cord is encircling the neck. If a loop of cord is felt, loosen it carefully and gently slip it over the infant’s head.

Figure 56.13 Delivery of the placenta and fetal membranes by exerting controlled traction on the cord and suprapubic pressure with the abdominal hand to prevent uterine inversion. Care should be taken to avoid avulsion of the cord. Active management with uterotonic agents such as oxytocin administered at delivery hastens delivery of the placenta and may reduce the incidence of postpartum hemorrhage and total blood loss. (From Gabbe SG, Niebyl JR, Simpson JL, editors: Obstetrics: normal and problem pregnancies, ed 5, Philadelphia, 2007, Churchill Livingstone.)
likely to occur. For this reason, palpate the uterus frequently to ensure that it is well contracted. A normally contracted uterus will feel firm with its upper margin just below the maternal umbilicus. If the uterus is flaccid or bleeding, gently massage it through the abdominal wall. Occasionally, the placenta may fail to separate completely, resulting in a retained placenta or placental fragments with persistent uterine bleeding. Support the patient with intravenous fluids and blood transfusions as indicated until definitive therapy is available. Constant firm uterine massage can lessen hemorrhage and may be lifesaving.

**COMPLICATIONS**

**Complex Deliveries**

**Shoulder Dystocia**

*Shoulder dystocia* refers to impaction of the fetal shoulders in the pelvic outlet after delivery of the head and occurs in 0.6% to 1.4% of deliveries (Fig. 56.14). Shoulder dystocia is associated with several risk factors, including fetal macrosomia, maternal diabetes, obesity, multiparity, and postterm pregnancy.

Impaction of the fetal shoulders and thorax in the maternal pelvis prohibits adequate respiration, and compression of the umbilical cord frequently compromises the fetal circulation. For these reasons, shoulder dystocia is a serious and potentially fatal complication of delivery. Fetal complications include brachial plexus injury, clavicular and humeral fractures, and rarely, death.

**Management**

The techniques used to treat shoulder dystocia frequently require an assistant, and delivery can result in fetal injury or hypoxia, as well as increased rates of PPH. Call for emergency assistance from a pediatrician, an obstetrician, and an anesthesiologist. Complications such as a fetal fracture or nerve injury are not unusual with shoulder dystocia during any delivery technique.

Although a variety of techniques have been described to free the anterior shoulder from its impacted position beneath the symphysis pubis, the initial technique of choice is the McRoberts maneuver because of its ease and potential for success (Fig. 56.15A). In one study, approximately 40% of shoulder dystocias were relieved with the McRoberts maneuver alone. To perform the McRoberts maneuver, place the mother in the extreme lithotomy position with her hips completely flexed so that her knees rest alongside her chest. This causes a flattening of the lumbar lordosis, rotates the maternal pelvis cephalad, and frequently frees the impacted anterior fetal shoulder. If the McRoberts maneuver fails to effect delivery, ask an assistant to apply moderate suprapubic, not fundal, pressure to the maternal abdomen while providing gentle downward traction on the fetal head (see Fig. 56.16B).

If these initial maneuvers fail to effect delivery, several other techniques exist, the choice of which will depend on clinician preference and experience. In the first maneuver, place two fingers in the vagina and exert pressure on the fetal scapula to rotate the *posterior* shoulder 180 degrees in a clockwise or counterclockwise fashion, depending on the orientation of the torso (reverse Wood’s screw), to free the entrapped anterior shoulder (see Fig. 56.15C). Other maneuvers include the Rubin maneuver, in which rotation is achieved by applying pressure on the posterior aspect of the anterior shoulder, or the Wood’s screw maneuver, in which pressure is placed on the anterior aspect of the posterior shoulder. This may cause release of the impacted anterior shoulder and result in progression of delivery. Alternatively, attempt to deliver the posterior arm (see Fig. 56.15D). In this maneuver, insert the hand along the hollow of the maternal sacrum to the level of the fetus’ posterior elbow. Exert pressure at the antecubital fossa, flex the posterior forearm of the fetus, and grasp the hand or forearm. Next, carefully sweep the posterior arm of the fetus across its chest to effect delivery of the posterior arm and shoulder. Rotate the shoulder girdle into one of the oblique diameters of the pelvis and subsequently deliver the anterior shoulder. No particular maneuver has a higher correlation to neonatal morbidity and mortality, but the use of multiple maneuvers does increase the chance for neonatal complications. Preliminary data suggests delivery of the posterior arm should be considered immediately following the McRoberts maneuver/subpubic pressure.

If all these strategies fail, it may be necessary to perform a controlled destructive procedure, such as fracture of the fetal clavicle, or the cephalic replacement maneuver (Zavanelli) with subsequent cesarean delivery. The Zavanelli maneuver should be a last resort. An episiotomy is not useful during the management of shoulder dystocia.

**Breech Delivery**

**Technique**

If a breech delivery becomes necessary in the ED, an episiotomy may need to be performed but is not recommended routinely. As the breech progressively distends the perineum, the posterior hip will be delivered, usually from the 6 o’clock position (Fig. 56.16, *step 1*). The anterior hip will then be delivered, followed by external rotation to the sacrum-anterior position. Ideally, two attendants should be available for a breech delivery.

**Delivery of the Presenting Part and Body.** Many breech infants deliver spontaneously with maternal expulsion efforts alone. Avoid fetal traction and apply fetal manipulation only
The McRoberts maneuver is the least invasive maneuver to disimpact the shoulders in shoulder dystocia. Position the patient in the extreme lithotomy position with the hips completely flexed (knee-chest position); this may free the anterior fetal shoulder.

Moderate suprapubic pressure will often disimpact the anterior shoulder. Desperate traction on the fetal head is not likely to facilitate delivery and might lead to trauma. Delivery of an infant with shoulder dystocia often results in fracture of the clavicle or humerus to accomplish delivery.

Rubin or reverse Wood’s screw maneuver. 1, Rotate the posterior shoulder. 2, Deliver the rotated shoulder.

Posterior shoulder delivery. Insert a hand and sweep the posterior arm across the chest and over the perineum. Take care to distribute the pressure evenly across the humerus to avoid unnecessary fracture.
As the fetus begins to emerge, the posterior hip will deliver, usually from the 6-o’clock position.

Avoid premature aggressive traction, which increases the risk for head entrapment or nuchal arm entrapment.

After spontaneous expulsion of the umbilicus, rotate the thigh externally and rotate the pelvis in the opposite direction.

As the knees flex during the rotational maneuvers, the legs can be delivered.

When the scapulae appear under the symphysis, reach over the left shoulder and sweep the arm across the chest.

Deliver the left arm.

Gently rotate the shoulder girdle to facilitate delivery of the right arm.

Rest the fetal body on your palm and forearm.

Place your index and middle fingers over the infant’s maxilla to maintain head flexion. Apply downward traction on the shoulders, and then elevate the body of the fetus to deliver the head.

Breech Delivery

Figure 56.16 Breech delivery.
after spontaneous delivery to the level of the umbilicus. Continued descent of the fetus will allow delivery of the legs, which may be aided by splinting the medial part of the thighs of the fetus with the fingers positioned parallel to the femur and exerting pressure laterally to sweep the legs away from the midline (see Fig. 56.16, steps 2 to 4). After delivery of the legs, grasp the fetal bony pelvis with both hands, with the fingers resting on the anterior superior iliac crests and the thumbs on the sacrum. Because the fetal body is slippery and difficult to hold, wrap it in a towel to assist delivery. Use the maternal expulsive efforts in conjunction with gentle downward traction. Rotate the fetal pelvis to bring the fetal sacrum into the transverse position to effect delivery of the scapulae. Two methods of shoulder delivery are commonly used (see Fig. 56.16, steps 5 to 7). In the first, with the scapulae visible, rotate the trunk so that the anterior arm and shoulder appear at the vulva and can easily be released and delivered. Next, rotate the body of the fetus in the reverse direction to deliver the other shoulder and arm beneath the symphysis pubis. In the second method, if trunk rotation is unsuccessful, deliver the posterior shoulder. Grasp the feet in one hand and draw them upward over the mother’s groin. Exert leverage on the posterior shoulder, which will slide out over the perineal margin, usually followed by the arm and hand. Deliver the anterior shoulder, arm, and hand beneath the symphysis pubis by downward traction on the fetal body.

Occasionally, spontaneous delivery of the arm and hand does not follow delivery of the shoulder. If this occurs, provide upward traction on the fetal body after delivery of the posterior shoulder. Pass two fingers along the fetal humerus until the fetal elbow is reached. Using the fingers to splint the fetal arm, sweep it downward and deliver it. Deliver the anterior arm by depression of the fetal body alone. In some cases it may be necessary to sweep the anterior arm down over the thorax by using two fingers as a splint.

Delivery of the Head. After the shoulders appear, the head usually occupies one of the oblique diameters of the pelvis, with the chin directed posteriorly. Extract the head by using the Mauriceau maneuver as follows. With the fetal body resting on the clinician’s palm and forearm, place the index and middle fingers of one hand over the infant’s maxilla, not the mandible, to maintain flexion of the fetal head (see Fig. 56.16, step 8). Hook two fingers of the other hand over the fetal neck and, while grasping the shoulders, apply downward traction until the suboccipital region appears under the symphysis pubis. Elevate the body of the fetus toward the mother’s abdomen, and the fetal mouth, nose, brow, and eventually the occiput will emerge over the perineum. Avoid excessive elevation of the fetal torso to prevent hyperextension of the neck. If available, have an assistant apply supra pubic pressure to help deliver the head. If delivery of the head is not affected by the Mauriceau maneuver, forceps delivery may be necessary but is beyond the scope of this text and most ED practitioners. Once the breech baby is delivered, further management proceeds as for a normal vertex delivery.

Rarely, breech extraction of an infant becomes necessary and is indicated only if there is a definite diagnosis of fetal distress unresponsive to routine maneuvers, if obstetric services are unavailable, and if cesarean section cannot be performed promptly. To perform the extraction, introduce the hand into the vagina and grasp both feet of the fetus, with the index finger placed between the fetal ankles. Apply gentle traction until the feet are pulled through the vulva. Continue gentle downward traction while grasping successively higher portions of both legs and thighs. When the breech appears at the vulva, apply gentle traction until the hips are delivered. As the buttocks emerge, rotate the fetal back anteriorly. Place the thumbs over the sacrum and the fingers over the hips and deliver the remainder of the breech as described earlier. At times, delivery of a frank breech may be necessary. Facilitated by an episiotomy, allow the breech to deliver spontaneously as far as possible. Place a finger on each side of the fetal groin and exert moderate traction. Once the knees appear outside the birth canal, flex the legs slowly to assist in delivery, and proceed with delivery as described earlier.

Episiotomy

Routine use of episiotomy is no longer recommended. Selected indications include breech delivery, shoulder dystocia, occiput-posterior presentations, and imminent perineal tear, but generally is not indicated and when done in these circumstances is an independent risk factor for third- or fourth-degree perineal tears. Episiotomy may be necessary to expedite delivery in situations of fetal distress. Two types of episiotomy are used: median (midline) and mediolateral (Fig. 56.17). The median approach is the easiest type to perform and repair, and results in the least amount of blood loss, heals more rapidly with minimal discomfort, and is generally more common in the United States. A major complication of median episiotomy is potential inadvertent extension of the incision into the anal sphincter or rectum, which results in third- and fourth-degree lacerations, respectively. A mediolateral episiotomy seldom results in extension into the anal sphincter, but blood loss is greater, repair is more difficult, and healing may be more painful.

Technique

With vertex presentations, perform the episiotomy during the second stage of labor, when the fetal head begins to distend the perineum and becomes visible to a diameter of 3 to 4 cm during a contraction. Anesthesia for episiotomy in the ED is usually limited to local infiltration of the perineum with 1% or 2% lidocaine.

Make the incision with Mayo scissors through the skin and subcutaneous tissue, the vaginal mucosa, the urogenital septum, and the superior fascia of the pelvic diaphragm (see Fig. 56.17). Make the incision up to half the length of the perineum, and extend it 2 to 3 cm upward into the vaginal mucosa. If the incision is in the midline, extend the incision through the lowermost fibers of the puborectalis portion of the levator ani muscles. As the head crowns, place the index and middle fingers inside the vaginal introitus to expose the mucosa, posterior fourchette, and perineal body. Use tissue scissors to incise the median raphe of the perineum halfway to the anal sphincter. For a mediolateral episiotomy, direct the incision downward and outward in the direction of the lateral margin of the anal sphincter either to the right or to the left.

Repair the episiotomy after delivery of the infant and placenta. Repairs are usually performed by the obstetric consultant in the delivery suite but can be done by an EP experienced in the procedure (see Fig. 56.17). The goals of episiotomy repair are to restore both anatomy and hemostasis with a minimal amount of suture material. Perform the closure after delivery of the placenta and following inspection and
Episiotomy And Repair

Midline Episiotomy
Make the incision through the lowermost fibers of the puborectalis portion of the levator ani muscles. Incise the median raphe of the perineum halfway to the anal sphincter.

A major complication of midline episiotomy is incision of the anal sphincter or rectum, which leads to third- and fourth-degree lacerations.

Mediolateral Episiotomy
Make the incision through the lowermost fibers of the puborectalis portion of the levator ani muscles. Direct the incision downward and outward in the direction of the lateral margin of the anal sphincter.

The mediolateral approach rarely extends into the anal sphincter but may be associated with greater blood loss and more difficult repair.

Place a taped sponge in the upper part of the vagina.

Expose the full extent of the episiotomy with the left hand. Place the first suture 1 cm cephalad to the most superior margin of the episiotomy or laceration to ensure hemostasis of the repair.

Use continuous locked absorbable suture to close the vaginal epithelium from the apex of the laceration to the hymenal ring.

Use simple interrupted (absorbable) suture to close the deep perineal fascia and underlying levator ani muscles (black arrows).

Bring the vaginal epithelial suture below the skin into the subcutaneous tissue (white arrow).

Use the vaginal epithelial suture to close the superficial fascia down to the edge of the episiotomy.

Use the same suture again as a subcuticular stitch coming back to the hymenal ring, where it is tied off.

Remove the vaginal sponge.

Figure 56.17 Episiotomy and repair. Routine episiotomy is no longer advised, but the procedure is often performed. Simple repairs may be done in the emergency department (ED), but this is often performed by the obstetrician or in the delivery room after an ED birth. (From Hacker NF, Gambone JC, Hobel CK, editors: Hacker & Moore's essentials of obstetrics and gynecology, ed 5, Philadelphia, 2009, Saunders.)
repair of the cervix and upper vaginal canal if indicated. The principles of repair are the same regardless of the type of episiotomy.

Because there is minimal tension on the closed wound, use 2-0 or 3-0 absorbable suture, such as chromic catgut or polyglycolic acid, on a large atraumatic needle. The first step is to close the vaginal mucosa with a continuous suture from just above the apex of the incision to the mucocutaneous junction to reapproximate the margins of the hymenal ring (see Fig. 56.17, step 1). Ligate large actively bleeding vessels during closure with separate absorbable sutures. Next, reapproximate the perineal musculature with three or four interrupted sutures (see Fig. 56.17, step 2). Close the superficial layers by one of two methods. In the first method, use a continuous suture to close the superficial mucosa from the mucocutaneous junction outward and then continue it upward as a subcuticular skin closure, with the suture returning to and ending at the mucocutaneous junction (see Fig. 56.17, steps 3 and 4). Alternatively, place several interrupted sutures through the skin and subcutaneous fascia and tie them loosely. This last method of skin closure avoids burying two layers of suture in the more superficial layers of the perineum.3,19

The most common complication of episiotomy is hematoma formation, which requires evacuation and drainage. Infection is an infrequent complication that usually responds to sitz baths, good hygiene, and antibiotic therapy.

Immediate PPH

PPH is the leading cause of maternal mortality globally. It is primarily seen in developing countries, but the incidence is increasing in the United States.64,65 PPH is generally defined as blood loss of more than 500 mL following vaginal delivery and more than 1000 mL following cesarean delivery. The average blood loss with a vaginal delivery is estimated to be approximately 500 mL but clinicians consistently underestimate actual blood loss.64 PPH is divided into early hemorrhage, which occurs within 24 hours of delivery, and late hemorrhage, which occurs after 24 hours and up to 6 weeks after delivery.

PPH may be sudden and massive but is frequently characterized by persistent moderate bleeding until serious hypovolemia develops. Observe carefully for blood loss, including evaluation of uterine size and consistency, during the early postpartum period. The most common cause of early PPH is uterine atony, which is involved in up to 80% of cases.38 Less common causes include lacerations of the lower genital tract, retained placenta or placental fragments, coagulation disorders, uterine rupture, uterine inversion, and placental site bleeding.4

Management

Management of PPH is similar to management of any acute blood loss. It consists of replacing intravascular volume with crystalloid and blood products, and using massive transfusion protocols as needed, and to correct the underlying cause of the hemorrhage. The diagnosis of uterine atony, the most common cause of bleeding, is made when uterine palpation reveals a soft “boggy” uterus. The diagnosis may be suspected on the basis of abdominal examination with confirmation made on bimanual examination as atony may be localized to the lower uterine segment.

Uterine atony is initially managed with firm manual massage of the uterine fundus through the abdominal wall in conjunction with the administration of oxytocic agents (Table 56.2). If bleeding persists, bimanual uterine compression is indicated. Use one hand to compress and massage the uterus through the abdominal wall while using the fist of the other hand to gently massage the anterior aspect of the uterus through the vagina (Fig. 56.18). Avoid vigorous downward massage, which can result in acute uterine inversion or can injure the blood vessels in the broad ligament.

Oxytocics

Oxytocin is the usual first-line drug for PPH secondary to uterine atony.66 Administer oxytocin as an intravenous infusion. A typical initial dose is 20 to 40 units in 1 L of crystalloid infused at a rate of 200 to 500 mL/hr. Titrate to sustain uterine contractions and control uterine hemorrhage. Slowing of hemorrhage should be observed within minutes of administration. If an intravenous line is not available, administer 10 units of intramuscular or intrauterine oxytocin.67 Do not administer oxytocin as an intravenous bolus because this can cause severe hypotension.67

| TABLE 56.2 Drugs for the Management of Immediate Postpartum Hemorrhage |
|------------------------|---------------------------|--------------------------|
| Drug                   | Dose                      | Comments                 |
| Oxytocin               | 20–40 units in 1 L of crystalloid initially infused at 200–500 mL/hr and then titrate to sustain uterine contractions and control hemorrhage | Do not administer as an IV bolus. First-line therapy. If IV access is unavailable, may use 10 units IM. |
| Methylergonovine maleate or ergonovine maleate | 0.2 mg IM | Avoid in patients with hypertensive disease, including preeclampsia |
| Carboprost tromethamine | 0.25 mg IM  
Repeat q 15 min until uterine hemorrhage is controlled or a maximum dose of 2 mg | Concurrent use of antiemetics and antidiarrheals recommended to control side effects |
| Misoprostol            | 800–1000 µg PR             | Single dose; can give PO. May cause tachycardia. |

*Note the increased concentration of oxytocin when used for the treatment of postpartum hemorrhage versus that given to stimulate uterine contractions after uncomplicated delivery.

IM, Intramuscularly; IV, intravenous; PO, orally; PR, per rectum; q, every.
If bleeding continues despite these measures, consider uterine tamponade with sterile gauze packing, insertion of an intrauterine balloon tamponade catheter (Bakri balloon [Cook Urological Inc., Spencer, IN] or BT-cath balloon tamponade catheter [Utah Medical Products, Inc., Midvale, UT]), or placing a Foley catheter into the atonic uterine cavity until surgical intervention or arterial embolization can be performed.68,73

If vaginal bleeding persists despite a firmly contracted uterus, search for additional causes of bleeding. The lower genital tract should be inspected for lacerations. Control bleeding by direct pressure or by gentle application of ring forceps to bleeding cervical lacerations. Use absorbable suture to control bleeding from accessible lacerations. Adequate visualization of the upper part of the vagina and cervix can be difficult, and repair of lacerations may require general anesthesia and obstetric intervention regardless of the location. Genital tract hematomas may also be a significant cause of hemorrhage but are not generally recognized until several hours after delivery.53 Also consider retained placental fragments and coagulopathy. Retained placental fragments may be removed by gentle manual extraction. Use one hand to follow the path of the umbilical cord and find the maternal placental plane and use the other hand to steady the uterine fundus through the maternal abdomen. The plane should feel velvety and irregular and should be gently dissected with side-to-side motion of the fingers until the placenta separates. This procedure is painful and associated with increased risk of infection, so analgesia and antibiotics should be administered.

Rarely, PPH is due to uterine inversion, with an estimated incidence of one in several thousand deliveries.74 Consider uterine inversion especially when there is severe pelvic pain, absence of a palpable fundus, brisk excessive PPH, and maternal hemodynamic instability. Shock develops in up to 40% of patients.69,75–77 The diagnosis is made by visualization or palpation of the uterine fundus in the vaginal vault or protruding through the introitus. On abdominal examination, no mass representing the uterus may be palpated, or when palpated, the uterus may have a cuplike dimpling of the fundus.68,71 Misoprostol, a synthetic prostaglandin E1 analogue, can also be given as an 800- to 1000-µg rectal dose. Clinical trials using tranexamic acid in PPH are being conducted and show promise but evidence on efficacy is insufficient to recommend use at the present time.66,72

If bleeding and poor uterine tone persist despite oxytocin therapy, consider additional therapy. Second-line therapy includes ergot derivatives such as methylergonovine or a prostaglandin. Give methylergonovine as a 0.2-mg dose intramuscularly every 2 to 4 hours, with uterine contractions occurring within 2 to 5 minutes of administration and lasting regularly for several hours.68 Though not the route of choice, methylergonovine can also be given via the oral route. Because of their tendency to cause severe hypertension, avoid administering ergot preparations to women with hypertensive disease, including preeclampsia.68,69 Alternatively, give 15-methylprostaglandin F2α (carboprost tromethamine) to stimulate uterine contractions.68–70 Administer carboprost at a dose of 0.25 mg intramuscularly, repeated at 15- to 90-minute intervals as determined by the clinical course, but not to exceed 2 mg.69,71 Misoprostol, a synthetic prostaglandin E1 analogue, can also be given as an 800- to 1000-µg rectal dose. Clinical trials using tranexamic acid in PPH are being conducted and show promise but evidence on efficacy is insufficient to recommend use at the present time.66,72

Figure 56.18 Use uterine massage to control postpartum bleeding. Insert one hand into the vagina to compress the anterior uterine wall while massaging the posterior aspect of the uterus through the abdominal wall with the other hand.

Figure 56.19 Manual replacement of an inverted uterus. Uterine inversion should be suspected with the sudden onset of brisk vaginal bleeding in association with an absent palpable fundus abdominally and maternal hemodynamic instability. It may occur before or after placental detachment. The diagnosis is made clinically with bimanual examination, during which the uterine fundus is palpated in the lower uterine segment or within the vagina. Use sonography to confirm the diagnosis if the findings on clinical examination are unclear.
Use tocolytic agents such as terbutaline or MgSO4 for uterine relaxation and repositioning, and stop all oxytocic drugs.3,6,8,69,77

To reposition the uterus, insert one hand into the vagina with the tips of the fingers at the uterocervical junction and hold the uterine fundus firmly in the palm of the hand. Gently apply pressure to the uterine fundus in the direction of the umbilicus. Do not initially exert pressure centrally on the fundus because this will cause the uterus to become compressed and force more “layers” of the uterus to lie within the relatively tight cervical ring.75,77 General anesthesia and laparotomy may be necessary for uterine positioning.

**PERIMORTEM CESAREAN SECTION**

Cardiopulmonary arrest is a rare event during pregnancy with the exact incidence unknown. Perimortem cesarean section (PCS) or resuscitative hysterotomy is a surgical procedure with the potential to save a viable fetus and improve maternal circulation in cases of maternal cardiopulmonary arrest.78 The time from when maternal circulation ceases to extraction of the neonate is the critical factor in determining maternal and fetal outcome. Delivery of the fetus increases maternal venous return and cardiac output by 25% to 30% and may lead to survival benefit for the mother.79,80 PCS should be considered in any woman who suffers a cardiac arrest after 24 weeks' gestation (uterine size exceeding the level of the umbilicus) and initiated within 4 minutes of onset of maternal cardiac arrest and the absence of spontaneous circulation.80,81 Evidence suggests the early initiation of PCS in cases of cardiac arrest may increase the restoration of hemodynamic status and likelihood of return of spontaneous circulation in the mother.80,85

**Indications**

Although the lower limit of fetal viability varies among institutions, performance of PCS before approximately 24 weeks is not generally indicated. If the duration of gestation is not known from the history, estimate fetal maturity quickly by determining the age of the fetus in weeks will correspond to the distance in centimeters from the last normal menstrual period or by measuring the height of the uterine fundus. Between 18 and 30 weeks’ gestation, the age of the fetus in weeks will correspond to the distance in centimeters from the uterine fundus to the symphysis pubis (e.g., at 28 weeks’ gestation the fundus lies approximately 28 cm above the symphysis pubis or halfway between the umbilicus and the costal margin).84

Survival of the infant is directly related to the time elapsed from maternal cardiac arrest to delivery, prompt performance of cardiopulmonary resuscitation (CPR) on the mother, the maturity of the fetus, pre-arrest health status of the mother, and in certain circumstances, the availability of neonatal intensive care facilities.85–87 In accordance with advanced cardiac life support guidelines, initiate CPR immediately on recognition of maternal arrest and continue it until the infant has been delivered. The anatomic and physiologic changes of pregnancy will inhibit the effectiveness of CPR on the mother. Aortocaval occlusion by the gravid uterus may occur after 20 weeks’ gestation. It can reduce venous return and compromise maternal cardiac output, especially in the supine position. CPR should be performed in the left lateral, head-down position or with an assistant manually displacing uterus away from the inferior vena cava.88 In addition, the decreased functional residual capacity of the lungs may impede ventilation efforts. Under these conditions, CPR generates only 30% to 40% of normal cardiac output, which severely compromises placental perfusion. The potential for infant survival decreases and the chance of neurologic damage increases as the time from maternal cardiac arrest to PCS rises. Neonatal outcome is best when PCS is performed within 4 minutes and poor if done after 20 minutes. Make every attempt to begin PCS within 4 minutes of cardiopulmonary arrest and complete the procedure within 5 minutes of arrest.84,87 Fetal prognosis is generally better in the later stages of pregnancy and after the sudden death of a previously healthy mother rather than death secondary to chronic illness.84,89 CPR should continue during and after the procedure. PCS in itself may represent the most important variable for successful maternal resuscitation.85,89 The decision to perform this procedure may be one of the most challenging that an EP will encounter. In the absence of immediate obstetric backup, it is reasonable to proceed with PCS within 4 minutes if initial maternal resuscitation efforts are unsuccessful. Prolonged attempts to resuscitate the mother are unlikely to benefit either the mother or the fetus.

**Technique**

The most experienced person present should perform the PCS, preferably an obstetrician. Under ideal circumstances a neonatologist should also be in attendance. Do not delay the procedure to allow time for arrival of consultants. Time should not be wasted searching for fetal heart tones or attempting to evaluate fetal viability with ultrasonography.

Extract the infant rapidly while avoiding fetal and maternal injury in the ED (Fig. 56.20). Hence, do not waste time preparing a sterile operating field or transporting the patient to an operating suite outside the ED. Use a large (e.g., No. 10) scalpel blade and make a midline vertical incision through the abdominal wall extending from the symphysis pubis to the umbilicus. Carry the incision through all abdominal layers into the peritoneal cavity. In most gravid women, the hyperpigmented “linea nigra” is apparent, and when present, use this as a guide for the incision. If available, place retractor in the abdominal wound and draw them laterally to expose the anterior surface of the uterus. Reflect the bladder inferiorly. If full, aspirate the bladder to evacuate it and permit better access to the uterus. While avoiding injury to fetal parts, make a small (≈5 cm) vertical incision through the uterus until amniotic fluid is obtained or the uterine cavity is clearly entered. Insert the index and long fingers into the incision and use them to lift the uterine wall away from the fetus. Use bandage scissors to extend the incision vertically to the fundus until a wide exposure is obtained. Gently deliver the infant, clamp and cut the cord, and evaluate and resuscitate the newborn as clinically indicated. Because the incision is relatively high in the uterus, the infant's head may not be readily accessible to the clinician. In this case, grasp the infant's feet and deliver the infant through maneuvers similar to those used for a breech delivery.

**THE NEWBORN**

Approximately 10% of newborns require some degree of assistance to begin breathing at birth (e.g., some sort of
stimulation to breathe), but less than 1% require extensive resuscitation. Evaluation of the newborn begins before delivery with assessment of maternal well-being and identification of risk factors for fetal distress: multiple pregnancies of less than 35 weeks, maternal infection or hypertension, oligohydramnios, preterm delivery at less than 36 weeks, breech presentation, meconium-stained amniotic fluid, non-reassuring fetal heart rate, emergency cesarean section, shoulder dystocia, and opiate use in normal labor.

Newborn resuscitation can be divided into four categories of action: (1) initial assessment and stabilization; (2) ventilation, including bag-valve-mask (BVM) or bag-tube ventilation; (3) chest compressions; and (4) administration of medications or fluids. Although most newborns require no resuscitation or only basic steps such as warming, drying, and stimulation, others will require further intervention. The most crucial action of neonatal resuscitation is establishment of adequate ventilation.

**Evaluation**

Traditionally, the Apgar scoring system, applied at 1 and 5 minutes after birth, has been the standard of newborn evaluation.
In general, the higher the score, the better the condition of the infant. Waiting for the traditional 1-minute Apgar score to indicate the need for resuscitation has been replaced by the concept of “the golden minute.” Begin basic resuscitation maneuvers, reevaluation, and positive pressure ventilation (PPV), if required, by the first 60 seconds after birth. Use the heart rate and respiratory effort to guide resuscitation efforts because skin color does not reliably predict oxygenation status in the newborn period.

**Respiration**

Normally, the newborn begins to breathe and cry almost immediately after birth. After initial respiratory efforts, the newborn should be able to establish regular respirations sufficient to improve color and should be able to maintain a heart rate higher than 100 beats/min. Generally, drying and stimulation (rubbing the back or flicking the soles of the feet) are enough to induce effective respirations in the newborn. Gasping and apnea after 30 seconds of stimulation are signs indicating the need for assisted ventilation and continuous pulse oximetry monitoring. It is important to note that even in healthy newborns, oxygen saturation does not reach extrauterine levels until 10 minutes after birth. Studies have demonstrated potential harmful effects from both excessive and insufficient supplemental oxygen versus room air during neonatal resuscitation. For term and preterm infants requiring ventilation, the current recommendation is to begin with room air and avoid the use of supplemental oxygen. It is important to note that even in healthy newborns, oxygen saturation does not normally reach extrauterine levels until 10 minutes after birth. Studies have demonstrated potential harmful effects from both excessive and insufficient supplemental oxygen versus room air during neonatal resuscitation. For term and preterm infants requiring ventilation, the current recommendation is to begin with room air and avoid the use of supplemental oxygen.

**TABLE 56.3** Apgar Scoring System

<table>
<thead>
<tr>
<th>Sign</th>
<th>0</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart rate (beats/min)</td>
<td>Absent</td>
<td>Slow (&lt; 100)</td>
<td>&gt; 100</td>
</tr>
<tr>
<td>Respiratory effort</td>
<td>Absent</td>
<td>Slow, irregular</td>
<td>Good, crying</td>
</tr>
<tr>
<td>Muscle tone</td>
<td>Flaccid</td>
<td>Some flexion of extremities</td>
<td>Active motion</td>
</tr>
<tr>
<td>Reflex irritability</td>
<td>No response</td>
<td>Grimace</td>
<td>Vigorous cry</td>
</tr>
<tr>
<td>Color</td>
<td>Blue, pale</td>
<td>Body pink, extremities blue</td>
<td>Completely pink</td>
</tr>
</tbody>
</table>

Note that oxygen administration to the newborn is not recommended after a routine delivery, even when there is early cyanosis. Excessive oxygenation may be toxic. For term and preterm infants requiring ventilation, the current recommendation is to begin with room air and avoid the use of supplemental oxygen. It is important to note that even in healthy newborns, oxygen saturation does not normally reach extrauterine levels until 10 minutes after birth.

An uncompromised newborn will be able to maintain a pink color of the mucous membranes without supplemental oxygen. Central cyanosis is determined by examining the face, trunk, and mucous membranes, but it is not a reliable method of determining oxygen saturation in the immediate neonatal period. Acrocyanosis is usually a normal finding in the newborn and not a reliable indicator of hypoxemia. It may, however, indicate other conditions such as cold stress.

**Stabilization Technique**

Following delivery of the infant and cutting the umbilical cord, place the newborn on the side in the sniffing position with the neck in a neutral or slightly extended position. Place a rolled blanket or towel under the back and shoulders of the supine infant to elevate the torso 2 to 2.5 cm because this may help maintain head position. Prevent heat loss in the newborn because cold stress can increase oxygen consumption and impede effective resuscitation. Avoid hyperthermia, however, because it is associated with perinatal respiratory depression. Place the infant under a radiant warmer, rapidly dry the skin, and wrap the infant in warmed blankets to reduce heat loss. Alternatively, use the mother’s body as a heat source for the newborn. If initial evaluation indicates that the infant is stable, dry and place the infant skin to skin on the mother’s chest or abdomen, and cover both with blankets.

Vigorous newborns should not be routinely suctioned after delivery, even those delivered through meconium-stained fluid. Reserve suctioning (including suctioning with a bulb syringe) immediately after birth for babies who exhibit obvious obstruction to spontaneous breathing or require PPV. Endotracheal intubation and suctioning is no longer recommended for nonvigorous babies born through meconium-stained amniotic fluid. If airway obstruction necessitates suctioning, first clear secretions from the mouth and nose with a bulb...
syringe or suction catheter (8- or 10-Fr). Aggressive pharyngeal suctioning can cause laryngeal spasm and vagal bradycardia, so limit the depth and duration of suctioning and do not exceed a negative pressure of 100 mm Hg.49,52

For newborns who do not quickly respond to conservative measures, PPV is indicated within 1 minute of birth. Most newborns who require PPV can be adequately ventilated with a bag and mask. Perform assisted ventilations at a rate of 40 to 60 breaths/min (30 breaths/min if mechanical compressions are being performed). As of the 2015 AHA guidelines, higher inflation pressures and longer inflation times for the first several breaths are not recommended due to insufficient data.49 Visible chest expansion is a more reliable indicator of appropriate inflation pressure than a specific manometer reading.49,52 Heart rate measurement can be used to assess adequacy of ventilation after 60 seconds of visible chest rise with PPV.49 Because BVM ventilation can produce gastric distention and impede respiration, insert an orogastric tube (8-Fr) in infants undergoing prolonged PPV.52

Endotracheal intubation may be indicated when BVM ventilation is ineffective, when the airway is obstructed, when chest compressions are performed, or when prolonged PPV is required.49,91 Alternatively, a neonatal laryngeal mask airway may provide effective airway management, especially in the case of ineffective BVM ventilation or failed endotracheal intubation.49

Monitor the heart rate during neonatal evaluation and stabilization by either direct auscultation over the chest, palpation of the pulse at the base of the umbilical cord, or a three-lead ECG. A readily discernible heartbeat of 100 beats/min or greater is acceptable. If the heart rate is less than 60 beats/min despite adequate ventilation and oxygenation for 30 seconds, institute chest compressions while continuing to ventilate.49,91 Deliver chest compressions on the lower third of the sternum and not over the xiphoid to avoid damage to the liver.49,91

There are two techniques for performing chest compressions in the newborn. In the preferred method, position two thumbs side by side over the lower third of the sternum just below the nipple line. If the infant is large or the resuscitator’s hands are too small to encircle the chest, use two-finger compressions with the ring and middle fingers.49,91 Compress to a depth that is approximately one third of the AP diameter of the newborn’s chest so that a palpable pulse is generated.49 Coordinate compressions and ventilations to avoid simultaneous delivery, which may compromise the efficacy of ventilation. Use a compression-to-ventilation ratio of 3:1 with 90 compressions and 30 breaths to achieve approximately 120 events/min.49 If the heart rate remains less than 60 beats/min despite these interventions,49 establish an umbilical or intravenous line and institute appropriate drug therapy. Alternatively, use intravenous access, but this may not be as effective in a preterm infant.52

The medications most commonly used during neonatal resuscitation are listed in Table 56.5.49,52

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Indications</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epinephrine</td>
<td>0.01–0.03 mg/kg (0.1–0.3 mL/kg of a 1:10,000 solution)</td>
<td>Bradycardia, asystole</td>
<td>May be repeated every 3–5 min as indicated</td>
</tr>
<tr>
<td>Volume expanders (normal saline or Ringer's solution)</td>
<td>10 mL/kg IV over 5–10 min</td>
<td>Suspected hypovolemia, shock, or blood loss</td>
<td>May be repeated after determination of clinical response</td>
</tr>
<tr>
<td>Bicarbonate</td>
<td>1–2 mEq/kg of a 0.5-mEq/mL solution given over at least 2 min</td>
<td>Prolonged arrests unresponsive to other therapy</td>
<td>Not indicated during brief periods of CPR</td>
</tr>
</tbody>
</table>

†The intravenous route is now strongly preferred, but one can consider administering epinephrine via an endotracheal tube in doses of 0.05 to 0.1 mg/kg while obtaining intravenous access.

CPR, Cardiopulmonary resuscitation.


ACKNOWLEDGMENTS

The authors gratefully acknowledge the work of George H. Lew and Michael S. Pulia, who contributed this chapter to the last edition.

REFERENCES ARE AVAILABLE AT www.expertconsult.com

SELECTED READINGS


REFERENCES


PELVIC EXAMINATION

In an emergency department (ED) there are several reasons to perform a pelvic examination: lower abdominal pain, vaginal discharge, vaginal bleeding, suspected sexually transmitted disease (STD), retained foreign body, and possible Bartholin abscess (BA) of the vulva. Emergency physicians do not routinely screen for cervical cancer. The evaluation of sexual assault victims is discussed in Chapter 58.

Before performing a pelvic examination, the examiner must have a basic understanding of female pelvic anatomy (Fig. 57.1). The vulva is made up of the labia majora, labia minora, clitoris, hymen, and vulvar vestibule. The labia majora extend from the mons veneris anteriorly to the perineal body and are filled with fatty tissue that varies in thickness with age—proportionately thicker in children and thinner as a woman passes through menopause. The labia minora are generally covered completely by the labia majora and are more delicate, pink, and hairless. Anteriorly the labia minora form a hood for the clitoris and posteriorly join with the labia majora to form the fourchette. The urethral meatus is just posterior to the clitoris. The hymen separates the external genitalia from the vagina and may be obscured by the labia. There are two Bartholin glands which lie below the fascia, one on each side of the lower, posterior third of the vagina. These glands secrete lubrication through ducts located at the lower pole of the labia minora. Normal Bartholin glands are rarely palpable. Healthy vaginal mucosa is reddish pink, but during pregnancy may become dusky, almost cyanotic-appearing. The cervix faces posteriorly (80%) but may face anteriorly (20%). The nonparous cervical diameter is 2 to 3 cm with a length of 2 to 4 cm. The cervix is normally pink. The cervical os changes with vaginal delivery from being smooth and circular to being fissured, oval, and slightly irregular.

Preparation for a Pelvic Examination

Before beginning an internal pelvic examination assemble the appropriate equipment. This equipment includes: a stretcher with stirrups or knee holders, various sizes of vaginal speculums, gloves, surgical lubricant, appropriate lighting, swabs, ring forceps or a Kelly clamp for the removal of foreign bodies or products of conception, transport medium for chlamydia and gonorrhea testing, pH paper, slides, coverslips, “eye dropper” bottles of 10% potassium hydroxide (KOH), and normal saline for testing vaginal discharge (Fig. 57.2). Speculums may be plastic and disposable, or may be metal (warmed to body temperature before insertion), but must be sized to the patient; a pediatric speculum for women with a narrow introitus (elderly women and young adolescents), a Smith-Pederson speculum for virginal women, a standard speculum for most, and a Graves speculum for obese women and multiparous women. Note that if stirrups or leg supports are not available, the patient can have a padded bedpan under her buttocks with the bottoms of her feet together and her knees separated. When examination in a lithotomy position is not possible, an examination may be performed with the patient in a knee-chest position.

### Culdocentesis

**Indications**

Diagnosis of acute pelvic conditions when ultrasound is not available or clinically feasible, including:

- Ruptured viscus (ectopic pregnancy or corpus luteum cyst)
- Pelvic inflammatory disease
- Other intraabdominal infections
- Splenic or liver injuries
- Ruptured aortic aneurysm

**Contraindications**

- Uncooperative patient
- Pelvic mass detected on bimanual examination
- Nonmobile retroverted uterus
- Coagulopathies

**Complications**

- Rupture of unsuspected tuboovarian abscess
- Bowel perforation
- Pelvic kidney perforation
- Bleeding

**Equipment**

- **Lidocaine with epinephrine**
- **Bivalve vaginal speculum**
- **Ring forceps**
- **18-gauge spinal needle**
- **19-gauge butterfly needle**

Review Box 57.1 Culdocentesis: indications, contraindications, complications, and equipment. Not shown is optional prepuncture topical anesthetic, such as a topical anesthetic-soaked cotton ball.
Figure 57.1 Anatomy of the female external genitalia, vagina, and rectouterine pouch (pouch of Douglas). A, Female external genitalia. B, Cross sectional view of female genitalia demonstrating the rectouterine pouch (Douglas’ pouch). C, Anatomy of cervix and posterior fornix as seen through speculum. (Modified from Drake RL: Gray’s anatomy for students, ed 2, Philadelphia, 2010, Elsevier, Fig. 5.56.)
A pelvic examination should be done with the patient in a lithotomy position. She should be gowned and should have an empty bladder. Privacy should be assured and the patient should be fully informed by the examiner. Chaperones should be fully informed by the examiner. Chaperones are commonly used during the examination. The examiner should glove and should touch the patient, reassuring her prior to each phase of the pelvic examination.

**Performance of the Pelvic Examination**

The vulva is examined visually for any lesions, and then by palpation. Women with a BA have acute, painful swelling of the posterior vulva that usually has developed over 2 to 3 days, making it difficult for them to sit or walk. The abscess appears erythematous, swollen, and forms a tender, spherical (3 to 4 cm in diameter) mass just lateral to the posterior fourchette. A Bartholin cyst is usually not tender and there is no associated erythema or swelling, differentiating it from an abscess. The treatment of a BA is similar to other abscesses in that incision and drainage is often required.

After examination of the vulva, separate the labia majora to expose the introitus and look for lesions, discharge, or blood. Next, insert a speculum with good lighting. If cervical cancer screening is anticipated, lubricate the speculum with water only. Otherwise, use surgical lubricant. Spread the labia, insert the speculum to its full length, and open the speculum. If the cervix is not seen, withdraw the speculum and palpate the location of the cervix with one finger, remove the finger, and follow with a second redirected speculum insertion. Inspect the cervix for blood, discharge and any lesions, particularly at the squamocolumnar junction (where the red columnar tissue of the endocervix changes to the reddish-pink squamous epithelium of the vagina). If there is a white lesion at the squamocolumnar junction that does not wipe off with a swab, consider dysplasia, carcinoma in situ, or condyloma acumini-tum. Collect samples for STD and vaginitis prior to withdrawal of the speculum (see later section on Vaginal Discharge and Suspected STD Sampling). Additionally, remove any foreign body or material found (see later section on Vaginal Foreign Body Removal). Withdraw the speculum slowly while inspecting the walls of the vagina. Close the speculum just prior to exiting the introitus.

A bimanual examination of a woman with a small vagina can be accomplished with one gloved, lubricated index finger. Two fingers can be used when the introitus is larger. Gently place your opposite hand on the patient’s abdomen. After palpating the labia majora, separate them, and inspect the introitus. Insert the fingers to locate the cervix. When using two fingers, place one on each side of the cervix. Pain with side to side cervical motion may indicate pelvic inflammatory disease (PID). Using the hand on the abdomen and the intravaginal fingers, evaluate the size and shape of the uterus. An asymmetric uterus may be an indication of a fibroid tumor. After moving both fingers to one side of the cervix, palpate the adnexa of that side between the intravaginal fingers and the abdominal hand. Gently palpate the pelvic adnexa since firm palpation of normal organs may cause pain and can be misleading to the examiner. If an adnexal mass is found, try to estimate a 3-D size in centimeters along with the degree of firmness, the degree of fixation to adjoining organs, and the degree of tenderness. Masses that can be detected with the bimanual examination include pedunculated fibroids, paraovarian cysts, tuboovarian abscesses (TOAs), and ectopic pregnancies.

Many examiners follow the bimanual examination with a rectal examination looking for firmness or nodules between the uterus and rectum, suggestive of tumors or uterosacral ligamentous endometriosis. Rectal material should be guaiac tested for blood, which may be falsely positive for gastrointestinal bleeding if blood is present in the vagina.

**Bartholin Abscess (BA)**

As stated previously, the BA is an acute painful swelling located in the posterior part of the vaginal introitus (Fig. 57.3). The inflammatory swelling can be small or quite large and painful. BA occurs in about 2% of women, usually in the reproductive years. The pathogenesis of the abscess is unclear, but is thought to be the result of a blocked or partially obstructed Bartholin duct leading to accumulation of material with subsequent infection. As with any abscess, the evaluation must include consideration of the patient’s comorbidities and willingness to consent for the procedure.

In most cases, a BA can be treated in the ED, but close follow-up with gynecology is needed. These abscesses can recur and the gynecologist may elect to do a marsupialization of the area which reduces the recurrence rate. If the abscess is very large, if the diagnosis is uncertain, or if there are other masses, significant cellulitis, or significantly distorted tissue, involvement of the gynecologist is warranted. In addition, if the patient has significant comorbidities, has unstable vital signs, has a bleeding dyscrasia, or is immunocompromised, the patient should be seen by the consulting gynecologist to manage the abscess.

**Incision and Drainage (Fig. 57.4)**

Incision and drainage of a BA can be painful and anxiety-provoking for many patients.

It is an invasive procedure that requires a written, witnessed, and signed consent form from the patient, parent, or guardian and should be witnessed with a notation made in the medical record documenting that the procedure was described, complications were discussed, and any alternatives such as antibiotics or warm compresses were offered when appropriate.
Once written consent is obtained, place the patient in the lithotomy position. Place the patient’s feet in stirrups. Generously premedicate with intravenous opioids, or sedatives when appropriate. Administration of nitrous oxide analgesia is also an accepted practice. Procedural sedation with propofol, etomidate, or benzodiazepines can be considered, especially for those patients who are extremely uncomfortable.

1. Use universal precautions with gown, gloves, and mask. The contents of the abscess may be under pressure and can spray when incised.
2. Clean the affected area in the usual sterile way.
3. Apply topical analgesia to the mucosal surface of the introitus where the swelling is most evident. Note that the abscess incision is made on the mucosal, not on the skin surface. Applying viscous lidocaine to this area for 10 minutes can help reduce discomfort.
4. Administer a local anesthetic using a 25- or 27-gauge needle and 2% lidocaine. Note that the abscess contents may be under pressure and may leak during this step.
5. Approach the abscess from the vaginal introitus and, using an 11-blade scalpel, make a 5-mm stab incision through the mucosal surface, not the skin, to evacuate the contents of the abscess. If the abscess is larger, the incision can be lengthened to the diameter of the abscess, but in most cases 5 mm is sufficient (Fig. 57.5).
6. After irrigating and cleaning the area, the wound can be packed or left open. Iodoform strips can be used to pack the wound. Do not pack the wound too tightly and leave just enough material to fill the base of the wound with a small tail (0.5 to 1 cm) extending from the cavity.
7. Some authors recommend inserting a small balloon catheter into the wound rather than packing. The Word catheter (Cook Medical Inc., Bloomington, IN) is often recommended (Fig. 57.6). Insert the Word catheter cautiously and fill it with 3 mL of saline. Be careful not to fill it too much because the resulting pressure in the abscess cavity can be uncomfortable. This should be sufficient to hold the catheter in place. On rare occasions, it is necessary to suture the catheter in place. The Word catheter is usually left in place for 2 to 4 weeks.
8. Close follow-up with gynecology is recommended for a repeat examination and evaluation within the next 3 days.
9. Antibiotics are usually not necessary unless there is fever, significant cellulitis, multiple comorbidities, or the patient is immunocompromised in any way. If these conditions are present, the patient may need observation and gynecology consultation prior to performing the procedure.
10. Counsel the patient that the Word catheter is usually left in place for up to 4 weeks, but close gynecology follow-up and monitoring is needed throughout that time.
11. If the patient is well enough to be discharged, prescribe analgesics, including nonsteroidal antiinflammatory medications. Advise patients that they can resume most of their normal activities pending repeat follow-up evaluation by gynecology.

**COMMON CAUSES OF VAGINAL BLEEDING IN ADULT WOMEN**

Common causes of vaginal bleeding in adult women include menstruation, oral contraceptive use, anovulatory cycles, spontaneous abortion, ectopic pregnancy, intrauterine contraceptive devices, persistent corpus luteum, gynecologic malignancy, vaginal or cervical injury, vaginal foreign bodies, gestational trophoblastic diseases, placenta abruptio, placenta previa, vasa previa, and uterine rupture. A good history along with beta human chorionic gonadotropin (β-hCG) testing and ultrasound (US) help differentiate among some of these, but pelvic examination should be considered for most of these potential conditions, except, most notably, placenta previa, where it can cause catastrophic hemorrhage. In general, avoid a pelvic examination in late third trimester bleeding.
Bartholin Abscess Drainage (Word Catheter)

Place the patient in the dorsal lithotomy position and identify the abscess, which will be located at either the 5- or 7-o’clock position, with the labia minora transecting the abscess.

Cleanse the area with antiseptic solution. While stabilizing the abscess with your thumb and index finger, inject local anesthetic through the mucosal surface (not through the skin).

Use a No. 11 scalpel blade to make an incision in the mucosal surface of the abscess. Make the incision large enough to accommodate the catheter, but small enough to prevent extrusion of the inflated balloon.

Alternatively, a hemostat can be used to puncture the abscess cavity. Stabilize the abscess with your thumb and index finger and skewer the abscess onto the hemostat (see Fig. 37.21).

Once the abscess has been entered (heralded by a palpable pop or the free flow of pus), insert the Word catheter to the hilt and inflate with 3 to 4 mL of saline. Use a 25-gauge needle to fill the balloon.

The catheter is left in place for 2 to 4 weeks to form a fistula. Antibiotics are of no proven value once drainage is performed, but practice varies.

Figure 57.4 Drainage of a Bartholin abscess (Word catheter [Cook Medical Inc., Bloomington, IN] method.) Note that the mucosal surface, not the skin, is the site of abscess drainage.
that may cause pain in the nonpregnant patient include salpingitis (PID), TOA, ovarian cyst, ovarian torsion, endometriosis, round ligament pain, uterine perforation, or uterine fibroids. Painless third trimester bleeding may be caused by placenta previa, in which a separated placenta is near or at the cervical os. Because manipulation of the uterus may further dislodge the placenta, defer rectal examination, speculum examination, and manual examination of the vagina to an obstetric professional. PID and TOA are often associated with vaginal discharge and fever. Endometriosis is often associated with dyspareunia and dysmenorrhea.

**Early Pregnancy Bleeding**

One fifth of pregnant women have vaginal bleeding during the first half of pregnancy, and 2% of women have an ectopic pregnancy, with the potential of hemorrhagic shock and maternal death. Evaluation of ectopic pregnancy is discussed in the later section on Culdocentesis.

One of the most common reasons to perform a speculum examination in early pregnancy bleeding is to evaluate a possible miscarriage, looking for cervical dilatation or the presence of products of conception. Some emergency physicians simply palpate the os intravaginally to determine the status of cervical dilatation and feel that speculum examination adds little to either the diagnosis or the plan of treatment, with the exception of early pregnancy bleeding associated with shock where the removal of obstructing products of conception is a resuscitative measure. Ultrasonography and β-hCG levels generally give the information needed to manage a patient with early pregnancy bleeding, and some practitioners feel that vaginal examination does not improve the accuracy of the diagnosis.2,3

**Pelvic and Lower Abdominal Pain**

Pelvic examination, particularly bimanual examination, is often performed for pelvic and lower abdominal pain. The pain can be caused by problems in the urinary tract, intestinal tract and the reproductive tract (in both pregnant and nonpregnant patients). Urinary tract conditions include ureteral colic, cystitis and pyelonephritis. Intestinal tract conditions include appendicitis, diverticulitis, inflammatory bowel disease, bowel obstruction or ischemia, and a perforated viscus. Reproductive tract conditions include ectopic pregnancy, threatened abortion, endometritis, endometriosis, corpus luteal cyst, salpingitis (PID and TOA), ovarian cyst, ovarian torsion, round ligament pain, uterine fibroids, and uterine perforation. Later in pregnancy these conditions of importance are ectopic pregnancy, placenta previa (which is most often painless), and placenta abruptio, all of which are suggested by vaginal bleeding. Conditions
Vaginal Foreign Body Removal

Retained vaginal foreign bodies can cause pain or bleeding and may lead to a foul-smelling vaginitis. Common vaginal foreign bodies in adults include retained tampons, broken parts of condoms, pessaries, contraceptive diaphragms, drug smuggling devices, and sexual stimulation devices. Women with psychiatric illness sometimes insert vaginal foreign bodies (vaginal polyembolokoilamania). The removal of these foreign bodies, once located, may be accomplished using a speculum and ring forceps, or a Kelly clamp. Imbedded foreign bodies may require removal in an operating room. Once the foreign body is removed, the vaginal walls should be inspected for lacerations and infection. Bleeding vaginal lacerations can be tamponaded with surgical dressing prior to repair. Deep lacerations may require operative repair by a gynecologist.

It is not uncommon for a tampon to be lodged in a horizontal position in the upper vagina, commonly following intercourse. There is frequently a foul odor. The tampon can be removed by trapping it between two gloved examining fingers. Withdraw the tampon while inverting the examining glove with the tampon inside, placing the encased tampon in the palm of the opposite glove and then inverting the second glove over the first, followed by disposal.

Vaginal Discharge and Suspected STD Sampling

History and physical exam are critical for diagnosing infectious pathology of the external genitalia, vagina, cervix, or reproductive tract. However, given the often nonspecific symptoms, laboratory evaluation is mandatory to target appropriate therapy. Diagnostic options include Gram stain, culture, nucleic acid amplification tests (NAATs), and antigen-detecting immunochromatography. Culturing is done to assess the effectiveness of antibiotics on targeted bacteria, particularly when there has been treatment failure. According to the Centers for Disease Control and Prevention, “the performance of NAATs with respect to overall sensitivity, specificity, and ease of specimen transport is better than that of any of the other tests available for the diagnosis of chlamydial and gonococcal infections.”

Bacterial Vaginosis (BV) and Candidiasis. The vaginal discharge of BV is homogeneous, thin, and white and when obtained from a speculum examination, can aid in diagnosis. pH paper applied to the vaginal fluid can suggest BV when the pH is over 4.5. This pH is elevated because of a lack of the normal acid-producing lactobacillus predominance in the vagina. Vaginal fluid generally has a pH below 4.5 and is usually in the 3.8 to 4.2 range. Conversely, mucus from the cervix generally has a pH that is in the 7.0 range and care must be taken not to sample cervical mucus when testing vaginal fluid for vaginitis. Make a wet mount of a very small smear of vaginal fluid. Take the fluid with a cotton swab and apply it to a glass slide. Mix it with a drop of 10% KOH and cover it with a coverslip. The fluid may have a fishy, amine smell (the “whiff” test). When examined microscopically (40 to 100 x power), the slide may reveal clue cells (vaginal epithelial cells studded with adherent coccobacilli that look somewhat like sugar cookies covered with poppy seeds).

The vaginal discharge of moniliasis (candidiasis) is “cheesy” and is associated with severe itching and burning of the perineum. A mixture of vaginal fluid with KOH should lyse many bacteria allowing the examiner to identify budding yeast cells, thereby helping with the diagnosis of monilial vaginitis (candidiasis).

Trichomoniasis. A wet mount of a tiny drop of vaginal fluid and a drop of normal saline, covered with a coverslip and examined microscopically, can aid in the diagnosis of trichomoniasis but such examination has poor sensitivity (51% to 65%). Trichomonads are about the size of white blood cells but are characterized by a rapid, flagella-caused movement.

The use of NAATs are highly sensitive and detect three to five times more T. vaginalis infections than wet-mount microscopy. There is also an antigen-detection test using immunochromatographic capillary flow dipstick technology (OSOM Trichomonas Rapid Test [Sekisui Diagnostics, Lexington, MA]) that can be performed at the point of care. The OSOM Trichomonas test is rapidly available and has a sensitivity of 82% to 95% with a specificity of 97% to 100%.

Cervicitis and PID. For women with a purulent cervical discharge, the diagnosis of cervicitis or PID should be considered. The cervix should be visualized during the speculum exam, assessing for mucopurulent discharge from the os. Appropriate transport culture medium should be available and a cervical swab should be obtained, placed in the medium, and sent to a bacteriologic laboratory.

Chlamydia and Gonorrhea. NAATs are the diagnostic tests of choice, when available, for both chlamydia and gonorrhea. The difference between urine, cervical, and vaginal specimen tests is not statistically significant. The sensitivity of a Gram stain of endocervical discharge for the diagnosis of chlamydia or gonorrhea is low compared to culture or NAATs and is not recommended.

CULDOCENTESIS

Culdocentesis is a procedure in which peritoneal fluid is aspirated from the retrouterine pouch by the insertion of a hollow needle through the posterior fornix of the vagina. Historically, this technique was used to diagnose ruptured ectopic pregnancies and ruptured ovarian cysts, and, rarely, to obtain cultures for diagnosis and treatment of PID. Widespread utilization of transvaginal US and β-hCG have largely replaced the use of culdocentesis. Nevertheless, culdocentesis has clinical utility to diagnose ruptured ectopic pregnancy in a patient for whom sonography is not readily available.

Anatomy for Performing a Culdocentesis

Before attempting culdocentesis, the clinician must be familiar with the anatomy of the vagina and the rectouterine pouch (pouch of Douglas) (see Fig. 57.1). In adult women, the vagina is approximately 9 cm long. From its inferior to its superior aspect, the posterior wall of the vagina is related to the anal canal by way of the perineal body, the rectum, and the peritoneum of the rectouterine pouch. The uterus lies at nearly a right angle to the vagina. The rectouterine pouch and the posterior wall of the vagina are adjacent only at the upper quarter (=2 cm) of the posterior vaginal wall. The vaginal wall in this area is less than 5 mm thick.
The blood supply to the upper part of the vagina comes from the uterine and vaginal arteries, which are branches of the internal iliac artery. This area is drained by a vaginal venous plexus that communicates with the uterine plexuses. The vagina has its greatest sensation near the introitus and little sensation in the area adjacent to the rectouterine pouch.

The rectouterine pouch is formed by reflections of the peritoneum, and it is the most dependent intraperitoneal space in both the upright and the supine positions. Blood, pus, and other free fluids in the peritoneal cavity pool in the pouch because of its dependent location. This pouch separates the upper portion of the rectum from the uterus and the upper part of the vagina. The pouch often contains small intestine and, normally, a small amount of peritoneal fluid.

Indications

Culdocentesis is indicated in any adult woman in whom aspiration of fluid from the rectouterine pouch will help confirm the clinical diagnosis. If US examination is not readily available in the ED or if the patient is too hemodynamically unstable to be transported to an off-site location for US, culdocentesis may be the fastest and most accurate diagnostic technique available to the emergency clinician.11 Analysis of peritoneal fluid is also an acceptable method of differentiating inflammatory from hemorrhagic pelvic pathologic conditions. Conditions in which culdocentesis may be of diagnostic value include a ruptured viscus (particularly an ectopic pregnancy or a corpus luteum cyst), PID and other intraabdominal infections (particularly appendicitis with rupture or diverticulitis with perforation), intraabdominal injuries to the liver or spleen, and ruptured aortic aneurysms.12

Ectopic Pregnancy

Ectopic pregnancy is often one of the most difficult gynecologic lesions to diagnose.11 The incidence of ectopic pregnancy is estimated to be 0.64% to 2.0%, although surveillance data is imprecise and limited for a variety of reasons.14 Ectopic pregnancy is the most common obstetric cause of maternal death in the first trimester.15 In a series of 300 consecutive cases of ectopic pregnancy, 50% of patients received medical evaluation at least twice before the correct diagnosis was made.

The clinical picture of ectopic pregnancy may include vascular collapse, pelvic pain, isolated rectal or back pain, amenorrhea, abnormal menses, shoulder pain, syncope, cervical or adnexal tenderness, adnexal mass, anemia, and leukocytosis. It is important to note that blood in the peritoneal cavity does not consistently correlate with peritoneal irritation, blood pressure, or pulse rate.15 In fact, bradycardia in the presence of significant intraperitoneal bleeding from a ruptured ectopic pregnancy is not unusual (Tables 57.1 and 57.2).

Risk factors for an ectopic pregnancy include a history of salpingitis, use of an intrauterine contraceptive device, or tubal ligation; however, no combination of these signs, symptoms,

### TABLE 57.1 Correlation Between the Results of Culdocentesis Performed on 77 Patients With Ectopic Gestation and Various Clinical Parameters

<table>
<thead>
<tr>
<th>Classic Triad</th>
<th>BLEEDING</th>
<th>ADNEXAL MASS</th>
<th>PERITONEAL SIGNS</th>
<th>PULSE ≥100 BEATS/MIN</th>
<th>BLOOD PRESSURE &lt; 90/40 MM Hg</th>
<th>MEAN HEMATOCRIT (%)</th>
<th>HEMOPERITONEUM ≥100 ML</th>
<th>RUPTURED TUBE</th>
<th>TOTAL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td>37</td>
<td>54</td>
<td>10</td>
<td>26</td>
<td>19</td>
<td>9</td>
<td>35</td>
<td>52</td>
<td>30</td>
</tr>
<tr>
<td>Negative</td>
<td>8</td>
<td>8</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>39</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Inadequate</td>
<td>13</td>
<td>15</td>
<td>6</td>
<td>5</td>
<td>4</td>
<td>1</td>
<td>38</td>
<td>13</td>
<td>7</td>
</tr>
<tr>
<td>Total patients</td>
<td>58</td>
<td>77</td>
<td>19</td>
<td>32</td>
<td>24</td>
<td>10</td>
<td>65</td>
<td>37</td>
<td>77</td>
</tr>
</tbody>
</table>

Note: There is a lack of correlation between positive culdocentesis and peritoneal signs and changes in vital signs. Patients are grouped by the result of culdocentesis (i.e., positive, negative, or inadequate). Note that only 10 patients were hypotensive and only 24 experienced tachycardia.


### TABLE 57.2 Correlation Between Tubal Status and Hypotension, Tachycardia, Hematocrit, Signs of Peritoneal Irritation, and Hemoperitoneum in 77 Patients With Ectopic Gestation

<table>
<thead>
<tr>
<th>CULDOCENTESIS</th>
<th>CULDOCENTESIS POSITIVE</th>
<th>PERITONEAL SIGNS</th>
<th>BLOOD PRESSURE &lt; 90/40 mm Hg</th>
<th>PULSE ≥100 BEATS/MIN</th>
<th>HEMOPERITONEUM ≥100 ML</th>
<th>AVERAGE HEMATOCRIT (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ruptured (n = 37)</td>
<td>30</td>
<td>25</td>
<td>8</td>
<td>19</td>
<td>37</td>
<td>33.6</td>
</tr>
<tr>
<td>Intact (n = 40)</td>
<td>24</td>
<td>7</td>
<td>2</td>
<td>5</td>
<td>28</td>
<td>37.3</td>
</tr>
<tr>
<td>Total patients</td>
<td>54</td>
<td>32</td>
<td>10</td>
<td>24</td>
<td>65</td>
<td>37.3</td>
</tr>
</tbody>
</table>

Note: Culdocentesis is frequently positive in the absence of rupture. Patients are grouped by the presence (i.e., “ruptured”) or absence (i.e., “intact”) of hemoperitoneum. Note that only about half the patients with “ruptured” status had tachycardia.

or historical data is diagnostic of an ectopic pregnancy. To confuse the diagnosis further, a normal menstrual history is reported in approximately 50% of patients with an ectopic pregnancy. A urine pregnancy test may occasionally be negative. Though rarely seen, the combination of a uterine decidual cast (Fig. 57.7) and a positive pregnancy test is virtually pathognomonic of an ectopic pregnancy. A uterine cast is decidua that has been hormonally stimulated by the ectopic pregnancy but is passed vaginally when the tissue can no longer be supported. The cast is an outline of the uterine cavity, but it can be mistaken for products of conception if not inspected carefully. Therefore all tissue passed vaginally should be carefully inspected before being sent to the laboratory for analysis for products of conception. An ectopic pregnancy can occasionally occur in conjunction with an intrauterine pregnancy. Patients who have undergone a therapeutic abortion may actually have had an unrecognized ectopic pregnancy, hence the need for pathologic evaluation of any tissue obtained by uterine evacuation procedures.

The greater sensitivity of the serum and urine β-hCG assay, coupled with the increased availability of emergency medicine physicians trained to perform pelvic US, has greatly increased the chance of early diagnosis of unruptured and ruptured ectopic pregnancy. Urinary β-hCG tests are sensitive at 20 mIU/mL or greater and are positive 98% of the time in the first few weeks of pregnancy. However, ectopic pregnancy is often associated with very low production of this hormone.

Quantification of the serum test adds additional information as it is sensitive to 5 mIU/mL. Therefore a negative urine β-hCG test rules out pregnancy in greater than 98% of cases, and pregnancy in any site can be ruled out in virtually all patients with a negative serum β-hCG test. A single quantitative β-hCG level is a poor predictor of the size of the pregnancy or the risk for ectopic pregnancy, but serial testing is quite helpful. It is expected that the quantitative serum β-hCG level should double approximately every 2 days in the first trimester.

To increase the accuracy of diagnosis, it is helpful to combine quantitative β-hCG testing with US examination (Fig. 57.8). An empty uterus by transvaginal or abdominal US combined with positive quantitative serum β-hCG results are strongly suggestive of ectopic pregnancy. The quantitative range in which the ultrasonographer should detect an intrauterine pregnancy varies, but an intrauterine pregnancy should be detected if the serum β-hCG level is in the range of 1200 to 1500 mIU/mL when using a transvaginal probe and greater

Figure 57.7 This decidual cast, a perfect outline of the uterine cavity, was initially thought to be a product of conception when found in the vaginal vault of a pregnant woman treated for abdominal pain and vaginal bleeding. The initial diagnosis was a spontaneous abortion, but this cast is virtually diagnostic of an ectopic pregnancy. Hypotension developed later and the woman was found to have a ruptured tubal pregnancy.

Figure 57.8 A, Pelvic ultrasound showing an empty uterus. The endometrial stripe (arrow) is clearly visible, and there is no evidence of a gestational sac. If the β human chorionic gonadotropin (β-hCG) level is below the discriminatory zone (see text), this could represent an early viable intrauterine pregnancy, a nonviable intrauterine pregnancy, completed abortion, or an ectopic pregnancy. If β-hCG exceeds the discriminatory zone, the chance of an ectopic pregnancy exceeds 85%. B, Pelvic ultrasound showing the presence of a yolk sac (arrow) within a gestational sac in the uterus. This patient was 5 weeks pregnant by dates.
peritoneum has been found in 45% to 60% of cases of a frankly ruptured ectopic pregnancy, it may be diagnostic even in a nonruptured case when bleeding has been slow or intermittent. Note that many ectopic pregnancies leak varying amounts of blood for days or weeks before rupture. Culdocentesis is an invasive procedure that requires a written, witnessed, and signed consent form from the patient, parent, or guardian when the patient’s condition permits. If verbal consent is obtained, this action should be witnessed and a notation made in the medical record documenting that the procedure was described, complications were discussed, and any alternatives (e.g., CT, sonography, or immediate laparoscopy) were offered when appropriate.

Once written or verbal consent is obtained, place the patient in a lithotomy position with the head of the table slightly elevated (reverse Trendelenburg position) so that intraperitoneal fluid gravitates toward the rectouterine pouch. Place the patient’s lower leg in a stirrup and flex the hip and knee to about 30°. Tense the patient’s abdomen to ascertain the intraabdominal pressure level below the discriminatory zone (defined as the hCG level at which a normal intrauterine pregnancy can be detected by US), is non-diagnostic and could represent an early viable normal pregnancy, a nonviable intrauterine pregnancy, a completed abortion, or an ectopic pregnancy. When no intrauterine pregnancy is detected by US and the serum β-hCG level exceeds the discriminatory zone, the chance of an ectopic pregnancy ranges from 86% to 100%.20

Culdocentesis may play an important role in some patients in the diagnosis of ectopic pregnancy. The test has an accuracy rate of 85% to 95%.11,31,32 Romero and coworkers32 reported that an ectopic pregnancy was found in 99% of patients with a positive pregnancy test and positive results on culdocentesis. Although culdocentesis is most often positive in the presence of a frankly ruptured ectopic pregnancy, it may be diagnostic even in a nonruptured case when bleeding has been slow or intermittent. Note that many ectopic pregnancies leak varying amounts of blood for days or weeks before rupture. Hemoperitoneum has been found in 45% to 60% of cases of unruptured ectopic pregnancy, as proved at surgery.12,21

Hence, culdocentesis may be helpful in a stable patient whose US examination does not demonstrate an intrauterine pregnancy despite a quantitative serum β-hCG level in the appropriate range. Although some clinicians opt for outpatient monitoring of serial β-hCG levels in this setting, patients in whom the clinician has high suspicion for an ectopic pregnancy (e.g., a patient who has or has had significant discomfort) or in whom close follow-up cannot be ensured may be candidates for culdocentesis.31 Even though a negative finding on culdocentesis does not rule out an early ectopic pregnancy, patients with a non-diagnostic US and a negative culdocentesis generally represent those at lower risk for “rupture” of an ectopic pregnancy during outpatient serum β-hCG monitoring. Patients with a non-diagnostic US examination and a serum β-hCG level below the threshold at which an intrauterine pregnancy should be visible on the US examination may need further evaluation. These patients, especially those with significant pain, an unexplained low hematocrit, or postural changes in vital signs (or near syncope), might be candidates for culdocentesis.

Blunt Abdominal Trauma

Historically, diagnostic peritoneal lavage (DPL) and computed tomography (CT) have been used to identify hemoperitoneum in blunt trauma patients. The use of culdocentesis has also been advocated to aid in this diagnosis.12,21 In the ED, two factors have largely obviated the need to perform invasive procedures for diagnosis of hemoperitoneum: (1) the increasing availability of high-resolution CT and (2) emergency clinicians trained to perform the bedside US focused assessment with sonography for trauma (FAST) examination. However, because small amounts of blood tend to collect in the rectouterine pouch, aspiration of clear peritoneal fluid is of great potential value in excluding a diagnosis of hemoperitoneum. This is especially helpful in situations in which US is unavailable or the patient is too unstable to leave the ED for a CT scan. In fact, culdocentesis may be more advantageous than DPL in some instances because there is less risk for urinary bladder perforation or bowel injury. In addition, previous abdominal surgery is not a relative contraindication to culdocentesis, as it is with DPL.21

Contraindications

Contraindications to culdocentesis are relatively few and include an uncooperative patient, a pelvic mass detected on bimanual pelvic examination, a nonmobile retroverted uterus, and coagulopathies. Pelvic masses may include TOAs, appendiceal abscesses, ovarian masses, and pelvic kidneys. It has been suggested that the only major risk with the procedure is rupture of an unsuspected TOA into the peritoneal cavity. This can be avoided by careful bimanual pelvic examination to exclude patients with large masses in the cul-de-sac.26 Although no data are available to guide the age at which culdocentesis may be performed safely, the procedure is generally limited to patients beyond puberty. This limitation is suggested on the basis of anatomy and with the consideration that the procedure is difficult to perform through a small prepubertal vagina.

Equipment

The equipment required for culdocentesis is depicted in Review Box 57.1. Either an 18-gauge spinal needle or a 19-gauge butterfly needle held by ring forceps is acceptable. It may be helpful to anesthetize the posterior vaginal wall at the site of the puncture with 1% to 2% lidocaine with epinephrine administered through a 27- or 25-gauge needle. Some physicians use a topical anesthetic (eutectic mixture of local anesthetics [EMLA], benzocaine), viscous lidocaine, or a cocaine-soaked cotton ball to anesthetize the mucosa before infiltration with a local anesthetic. Although puncture of the posterior vaginal wall at the upper fourth of the vagina is generally no more painful than a venipuncture, there is some advantage to using a local anesthetic if multiple attempts at culdocentesis are required, as is sometimes the case. In addition, the epinephrine may produce vasoconstriction and reduce bleeding associated with the needle puncture. Culdocentesis is often stressful to the patient, and all attempts should be made to render the procedure as painless as possible. Parenteral analgesia and/or sedation should be administered when the patient is uncomfortable or anxious.

Technique

Preparation

Culdocentesis is an invasive procedure that requires a written, witnessed, and signed consent form from the patient, parent, or guardian when the patient’s condition permits. If verbal consent is obtained, this action should be witnessed and a notation made in the medical record documenting that the procedure was described, complications were discussed, and any alternatives (e.g., CT, sonography, or immediate laparoscopy) were offered when appropriate.

Once written or verbal consent is obtained, place the patient in a lithotomy position with the head of the table slightly elevated (reverse Trendelenburg position) so that intraperitoneal fluid gravitates toward the rectouterine pouch. Place the patient’s
feet in stirrups. Premedicate with intravenous opioids or sedatives if appropriate. Administration of nitrous oxide analgesia is also an accepted practice. Procedural sedation with propofol, etomidate, or benzodiazepines can be considered. Although the pain associated with passage of the culdocentesis needle is generally minor, judicious use of analgesia and sedation makes the procedure easier for both the clinician and patient.

If radiographs are indicated, take them before culdocentesis to avoid confusion with procedure-induced pneumoperitoneum.

**Exposure**

Perform a bimanual pelvic examination before culdocentesis to rule out a fixed pelvic mass and to assess the position of the uterus. It is possible to palpate an adnexal mass if the mass exceeds 3 cm in diameter. Insert a bivalve vaginal speculum and open it widely by adjusting both the height and the angle thumbscREWs (Fig. 57.9, step 1). Grasp the posterior lip of the cervix with the toothed uterine cervical tenaculum and elevate the cervix. Warn the patient in advance that she may feel a sharp pain when the cervix is grasped with the tenaculum. Inform the patient also that bleeding from the tenaculum puncture site or culdocentesis site, or both, may produce postprocedural spotting.

Use the tenaculum to elevate a retroverted uterus from the pouch, to expose the puncture site, and to stabilize the posterior wall during puncture with the needle. Some clinicians prefer to use longitudinal traction on the cervix to produce the same result. The vaginal wall adjacent to the rectouterine pouch will be tightened somewhat between the inferior blade of the bivalve speculum and the elevated posterior lip of the cervix. Such tightening of the vaginal wall exposes the puncture site and keeps it from moving away from the needle when the wall is punctured.

After the tenaculum is applied and the posterior lip of the cervix is elevated or traction is applied, swab the vaginal wall in the area of the rectouterine pouch with an antiseptic, followed by a small amount of sterile water. Administer a local anesthetic (1% lidocaine with epinephrine) at this point. Anesthetic may be injected through a separate 27- or 25-gauge needle or with the spinal needle that will be used for the culdocentesis. Use a cotton ball soaked in 4% cocaine or 20% benzocaine solution or apply viscous lidocaine to the area for topical anesthesia of the posterior vaginal wall approximately 15 minutes before infiltration with a local anesthetic. This combination will make the needle puncture nearly painless. Attach the needle to a 20-mL syringe. A smaller syringe might not be long enough to allow adequate control of the needle, and the clinician's hand may block the view of the puncture site if a smaller syringe is used.

**Aspiration**

Following local anesthesia, advance the syringe and the spinal needle parallel to the lower blade of the speculum (see Fig. 57.9, step 2). Fill the syringe with 2 to 3 mL of saline (nonbacteriostatic) before puncture. After needle puncture, the free flow of fluid from the syringe will expel tissue that may have clogged the needle and will confirm that the tip of the needle is in the proper position and not lodged in the uterine wall or the intestinal wall. Use saline rather than air because if air is used, it may be difficult to interpret the presence of free peritoneal air on subsequent radiographs. To avoid the need to change the syringe during the procedure, use 1% lidocaine for both anesthesia and confirmation of proper needle placement; however, the bacteriostatic property of this agent precludes its use if the procedure is performed to obtain fluid for culture.

Penetrate the vaginal wall in the midline 1 to 1.5 cm posterior (inferior) to the point at which the vaginal wall joins the cervix (see Fig. 57.9, step 2). Pass the needle a total of 2 to 2.5 cm.27 Apply gentle suction with the syringe while slowly withdrawing the needle. Avoid aspirating any blood that has accumulated in the vagina from previous needle punctures or from cervical bleeding because this may give the false impression of a positive tap. Bleeding from the puncture site in the vaginal wall can be minimized by adding epinephrine to the local anesthetic.

Blood or fluid may be obtained immediately but may also be obtained when the needle is withdrawn from the peritoneal cavity. Therefore it is important to aspirate continuously while gradually withdrawing the needle. If no fluid is aspirated, reintroduce the needle and direct it only slightly to the left or right of the midline. Directing the needle too far laterally may result in puncture of the mesenteric or pelvic vessels. If no fluid is obtained on the first attempt, repeat the procedure.

Some physicians prefer the use of a 19-gauge butterfly needle held with ring forceps (see Fig. 57.9, step 3, and Fig. 57.10).27 This technique offers a built-in guide to needle depth and allows good control of the needle during puncture. An assistant must aspirate the tubing while the physician controls positioning and withdrawal of the needle.

Fluid that is aspirated may be old nonclotting blood, bright red blood, pus, exudate, or a straw-colored serous liquid. Any fluid that is not blood should be submitted for Gram stain, aerobic and anaerobic culture, and cell count. Blood should be observed for clotting. Blood should also be sent for determination of the hematocrit.

**Interpretation of Results**

Interpretation of the results of culdocentesis depends primarily on whether any fluid was obtained. It should be noted that in the absence of a pathologic condition, 2 to 3 mL of clear yellowish peritoneal fluid can be aspirated. When there is no return of fluid of any type (a so-called dry tap), the procedure has no diagnostic value. Because a dry tap is nondiagnostic, it should not be equated with normal peritoneal fluid. In addition, when less than 2 mL of clotting blood is obtained, this is also considered to be a nondiagnostic tap because the source of this small amount of blood may be the puncture site on the vaginal wall. Such blood will usually clot. More than 2 mL of nonclotting blood is certainly suggestive of hemoperitoneum. However, some researchers interpret as little as 0.3 mL of nonclotting blood as a positive tap.15 There is no particular significance of larger amounts of blood because absolute volume may be related to the position of the needle or the rate of bleeding. Brenner and colleagues14 reported no blood from culdocentesis in 5% of patients with proven ectopic pregnancies even when rupture had occurred. In the series of 61 patients with surgically proven ectopic pregnancy reported by Cartwright and associates,15 culdocentesis performed within 4 hours of surgery was positive in 70%, negative in 10%, and inadequate in 20%. “Positive” in their series was defined as obtaining at least 0.3 mL of nonclotting blood with a hematocrit of greater than 3%. “Negative” was defined as obtaining 0.3 mL of fluid with a hematocrit of less than 3%. An “inadequate” tap was one in which no fluid was obtained. In the 252 patients reported by Vermesh and coworkers29 who had surgically proven ectopic
A “positive tap” is one in which nonclotting blood is obtained, although the presence of nonclotted blood does not confirm a tubal pregnancy. Intrapertoneal blood from any source (ectopic pregnancy, ovarian cyst, ruptured spleen) may remain unclotted after aspiration for days as a result of the defibrination activity of the peritoneum. Return of serosanguineous fluid also suggests a ruptured ovarian cyst. The hematocrit of blood from active intraperitoneal bleeding is greater than 10%. In one series, the hematocrit of blood from a ruptured ectopic pregnancy was 15% or greater in 97% of cases.\(^{14}\)

It should be emphasized that a positive finding on culdocentesis in the presence of a positive pregnancy test does not always prove an ectopic pregnancy.\(^{29}\) A ruptured corpus

**Culdocentesis**

Place the patient in the lithotomy position with the head of the bed slightly elevated so that intraperitoneal fluid gravitates toward the rectouterine pouch. Premedicate with sedatives or intravenous opiates as clinically indicated. Perform a bimanual pelvic examination to exclude the presence of a pelvic mass.

Insert a bivalepelvic speculum and open it widely by using both the height (A) and angle (B) thumbscrews. Grasp the posterior lip of the cervix with a tenaculum and elevate the cervix.

Swab the vaginal wall in the area of the rectouterine pouch with antiseptic followed by a small amount of sterile water.

Administer a local anesthetic with a separate 25- or 27-gauge needle or with the needle that will be used for culdocentesis. The site of needle entry is 1 cm posterior to the point at which the vaginal wall joins the cervix (x). (Optional: before injecting the local anesthetic, place a cocaine [4%]- or benzocaine [20%]-soaked cotton ball on the area to be punctured for 15 minutes.)

Fill the syringe with 2 to 3 mL of saline prior to puncture. Advance the spinal needle parallel to the lower blade of the speculum. Gently depress the plunger during advancement of the needle. Free flow of fluid will confirm proper needle placement in the rectouterine pouch. Pass the needle 2 to 2.5 cm. Apply gentle suction while slowly withdrawing the syringe.

Alternatively, use a 19-gauge butterfly needle held with ring forceps. This technique offers a built-in guide to needle depth and allows good control of the needle during puncture. Use the help of an assistant to aspirate while you control needle position and withdrawal.

**Figure 57.9** Culdocentesis. (1 and 2, from Vander Salm TJ, Cutler BS, Wheeler HB: *Atlas of bedside procedures.* Boston, 1979, Little, Brown; 3, from Webb MJ: *Culdocentesis,* *JACEP* 7:452, 1978.)
A

18-gauge spinal needle

Ring forceps

19-gauge butterfly needle

B

Figure 57.10 Variations in culdocentesis technique. A, An 18-gauge spinal needle attached to a syringe. The operator applies continuous suction during withdrawal of the needle. B, A 19-gauge butterfly needle attached to a syringe and held with ring forceps. An assistant is required to aspirate the syringe while the operator controls the needle with the forceps.

BOX 57.1 Interpretation of Culdocentesis

**POSITIVE**

>0.5 mL nonclotting, bloody fluid (hematocrit >12%)

Indicates hemoperitoneum
- When β-hCG is also positive, ectopic pregnancy is found in greater than 95% of cases
- Nonspecific—can occur in intrauterine pregnancies and nonpregnant women (e.g., ruptured cyst, retrograde bleeding)

Does not necessarily indicate tubal rupture
- 50% to 62% of ectopic pregnancies with peritoneal blood may be unruptured

**NEGATIVE**

Serous fluid
- Excludes hemoperitoneum and tubal rupture
- Falsely negative in 10% to 15% of ectopic pregnancies (generally unruptured)

**NONDIAgnostIC**

Dry tap or clotting blood
- Excludes neither ectopic pregnancy nor hemoperitoneum
- 15% of procedures are nondiagnostic
- 16% of ectopic pregnancies have nondiagnostic study results

β-hCG, β subunit of human chorionic gonadotropin.

TABLE 57.3 Interpretation of Culdocentesis Fluid

<table>
<thead>
<tr>
<th>ASPIRATED FLUID</th>
<th>CONDITION AND SUGGESTED DIFFERENTIAL DIAGNOSIS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clear, serous, straw colored (usually only a few milliliters)</td>
<td>Normal peritoneal fluid</td>
</tr>
<tr>
<td>Large amount of clear fluid</td>
<td>Ruptured or large ovarian cyst (fluid may be serosanguineous); pregnancy may be coexistent</td>
</tr>
<tr>
<td>Ascites</td>
<td></td>
</tr>
<tr>
<td>Carcinoma</td>
<td></td>
</tr>
<tr>
<td>Exudate with polymorphonuclear leukocytes</td>
<td>Pelvic inflammatory disease</td>
</tr>
<tr>
<td>Gonococcal salpingitis</td>
<td></td>
</tr>
<tr>
<td>Chronic salpingitis</td>
<td></td>
</tr>
<tr>
<td>Purulent fluid</td>
<td>Bacterial infection</td>
</tr>
<tr>
<td>Tuboovarian abscess with rupture</td>
<td></td>
</tr>
<tr>
<td>Appendicitis with rupture</td>
<td></td>
</tr>
<tr>
<td>Diverticulitis with perforation</td>
<td></td>
</tr>
<tr>
<td>Bright red blood†</td>
<td>Ruptured viscus or vascular injury</td>
</tr>
<tr>
<td>Recently bleeding ectopic pregnancy† (ruptured or unruptured)</td>
<td></td>
</tr>
<tr>
<td>Bleeding corpus luteum</td>
<td></td>
</tr>
<tr>
<td>Intraabdominal injury</td>
<td></td>
</tr>
<tr>
<td>Liver</td>
<td></td>
</tr>
<tr>
<td>Spleen</td>
<td></td>
</tr>
<tr>
<td>Other organs</td>
<td></td>
</tr>
<tr>
<td>Ruptured aortic aneurysm</td>
<td></td>
</tr>
<tr>
<td>Old, brown, nonclotting blood</td>
<td>Ruptured viscus</td>
</tr>
<tr>
<td>Ectopic pregnancy with intraperitoneal bleeding over a few days or weeks</td>
<td></td>
</tr>
<tr>
<td>Old (days) intraabdominal injury (e.g., delayed splenic rupture)</td>
<td></td>
</tr>
</tbody>
</table>

†Note: The hematocrit of blood from a ruptured ectopic pregnancy is usually 15% or greater (97.5% of cases), but some authors use greater than 3% as positive.

Complications

Culdocentesis is one of the safest procedures performed in the emergency setting, and there are probably fewer complications with this technique than with peripheral venous cannulation. Complications have been reported, however, the most serious being rupture of an unsuspected TOA. Other complications include perforation of the bowel, perforation of a pelvic luteum cyst in the presence of an intrauterine pregnancy test is probably the most common cause of a false-positive scenario. Whenever possible, US can help corroborate the findings on culdocentesis.
kidney, and bleeding from the puncture site in patients with clotting disorders. Because the most common complications result from puncture of a pelvic mass, careful bimanual examination of the patient should help prevent this problem. Puncture of the bowel and the uterine wall occurs relatively frequently but does not generally result in serious morbidity. Obviously, penetration of a gravid uterus has greater potential for harm. Occasionally, one will aspirate air or fecal matter, thereby confirming inadvertent puncture of the rectum.
REFERENCES

Examination of the Sexual Assault Victim
Carolyn Joy Sachs and Malinda Wheeler

The majority of sexually assaulted individuals do not share the experience with anyone, suffering in silence. An estimated one-third of all sexual assaults are reported to law enforcement. In many cases, after contact with law enforcement, patients are taken to the emergency department (ED) for evaluation, examination, and treatment. Sexual assault patients may also present to the ED de novo, without prior law enforcement contact. In 2013, according to the Centers for Disease Control and Prevention (CDC) statistics, sexually assaulted patients accounted for slightly more than one-tenth of all assault-related visits to the ED by female patients.1

Some sexual assault victims agree to police interviews and investigations, and others do not. Federal legislation guarantees all victims the right to a forensic examination and treatment of sexual assault regardless of their cooperation with legal investigation or their desire to initially pursue prosecution.2 Some states require medical personnel treating sexual assault patients to report the assault to local law enforcement, whereas others forbid such reporting without patient consent. Clinicians must know their own state laws regarding such reports.

DEFINITIONS

Sexual assault refers to any sexual contact between one person and another without appropriate legal consent.3 Physical force may be used to overcome the victim's lack of consent, but this is not mandatory to prove assault. Coercion into sexual contact by intimidation, threats, or fear also defines sexual assault. State laws differ slightly on the definition of exact acts that constitute sexual contact and on which populations are unable to give legal consent. In general, persons under the influence of drugs or alcohol, minors, and those who are mentally incapacitated are considered unable to give consent for sexual contact.

Clinicians who treat sexual assault patients have a professional, ethical, and moral responsibility to provide the best medical and psychological care possible. At the same time, they must collect and preserve the proper medicolegal evidence that is unique to the evaluation of sexual assault cases. Often hospitals and jurisdictions affiliate with designated sexual assault examination teams to provide specialized evaluation and treatment of patients. These sexual assault response teams (SARTs) provide clear advantages outlined near the end of the chapter. However, patients may be brought to an ED that does not routinely provide specialized care for sexual assault. This chapter is designed to aid clinicians in such a general care location. Prepared emergency personnel can help attenuate the psychological and physical impact of sexual assault. Through proper care of the patient and careful acquisition of evidence, ED staff can help the patient recover from the assault and aid society in improving the prosecution and conviction rates of sexual predators.

EVALUATION AND TREATMENT OF PATIENTS SUFFERING FROM SEXUAL ASSAULT

Preparation

Most often, local jurisdictions or hospitals provide clinicians with detailed forms and instructions for examination and documentation of sexual assault. This chapter is meant to supplement such instructions and forms. Clinicians should be familiar with local documents before performing a sexual assault examination. Careful step-by-step planning and the use of written protocols ensure the best care for victims and aids in the prosecution of assailants and the exoneration of wrongly accused subjects.

ED personnel must secure patient privacy and designate a separate area for the care of sexually assaulted patients. If medically and logistically possible, interviews should be conducted in a private room separate from the examination room. EDs often have such an area, frequently called the “grieving room” or the “family room.” Law enforcement or other governmental agencies may provide examination kits for the collection of forensic evidence from victims (Fig. 58.1). These kits should be available in the ED and the staff should be familiar with them. If such kits are not provided by local sources, hospital staff may need to assemble their own kits from material found in most EDs. Alternatively, private companies assemble and sell such kits (e.g., The Lynn Peavey Company, https://www.lyn.peavey.com). Prepared kits can save a tremendous amount of nursing and clinician time. A checklist of local requirements for sexual assault examination should be included in the kits and serves as a reminder for all the necessary medicolegal procedures to be completed.

Although this chapter is devoted primarily to the evaluation of adult female sexual assault patients, guidelines for the evaluation of adult male sexual assault patients, female child patients, male child patients, and accused assailants are provided in separate sections of this chapter. The same examiners designated to perform adult female examinations may easily perform male victim and assailant examinations; however, examination of a child sexual assault patient often requires considerable expertise and training. When possible, medical staff with specialized training in the examination of child sexual assault patients should perform these examinations. If this is not possible, the section on Child Sexual Assault Examinations should provide emergency medical personnel with a basic framework to perform an initial examination.

Consent

Consent for the evaluation and treatment of a sexual assault patient is mandatory. The patient has undergone an experience in which her/his right to grant or deny consent was taken from her/him, and obtaining consent for medical treatment and gathering evidence has important psychological and legal implications. The patient has the right to decline medicolegal examination and even medical treatment. Before beginning evaluation and treatment, obtain witnessed, written, informed consent. If no local forensic examination forms are available,
use the standard ED “consent to treat” forms, but ensure that the patient is well informed and gives verbal consent to each step of the examination. Although a few states mandate that medical personnel report sexual assaults to law enforcement, patients may choose to not discuss the event with police. If the patient cannot give consent for a forensic examination because of a reversible process (e.g., intoxication, an acute psychological reaction), wait several hours for the patient’s mental status to improve to a reasonable level before consent is obtained. When patients cannot give consent because of minor status or a developmental disability, the person authorized to give medical consent for the patient may give consent for the examination unless that person is a suspect in the assault. Many states allow an adolescent patient of a certain age (e.g., > 12 to 14 years old) to consent to an examination for conditions related to sexually transmitted diseases (STDs), sexual assault, and pregnancy. State laws also differ in examiners’ requirements to make an attempt to contact the legal guardian (unless the guardian is a suspected perpetrator). Clearly, emergency personnel must be informed regarding their local laws concerning these requirements. In the rare case that a patient cannot give consent as a result of a potentially irreversible medical condition, such as severe head trauma and coma, seek the advice of institutional legal counsel before proceeding with a forensic examination. In some cases, the next of kin may provide the needed consent, whereas in other cases, it may be necessary to obtain a court order to proceed.

History
The history of the event should include only the elements necessary to complete the required forms, to perform a focused physical examination, and to collect evidence. Questions beyond this, such as the details leading up to the assault, should be left to police investigators. Avoid the urge to “help” the alleged victim by unduly embellishing or detailing uncorroborated or nonmedical information supplied during the examination. Limiting the history not only shortens the evaluation in the ED but also helps prevent discrepancies between the ED history and the official police investigation report, which could weaken the victim’s case in court. Document the pertinent medical history, including the last menstrual period, current contraception, recent anal-genital injuries or surgeries, and preexisting injuries.

The history of the event required by legal documentation or protocol usually includes the time, date, and place of the alleged assault and a description of the use of force, threats of force, and the type of assault. Elements of force may include the type of violence used (e.g., grabbing, hitting, kicking, strangling, weapon use), threats of violence, the use of restraints, the number of assailants, the use of alcohol or drugs (forcibly or willingly) by the patient, and any loss of consciousness experienced by the patient. Sexually assaultive acts may include manual or oral fondling (of breasts, genitalia, or both); vaginal, oral, or anal penetration or attempted penetration (with fingers, penis, or other objects); ejaculation on or in the body; and the use of a condom. Use of physical force or violence is partly a police matter, but from a medical standpoint it is desirable to correlate positive findings on the physical examination (e.g., abrasions, ecchymosis, and scratches) with a description of any force, restraint, or violence.

Clearly indicate the patient’s post-assault activity, which is commonly requested on the necessary documentation, including douching, bathing, urinating, defecating, gargling, and brushing teeth. These activities can alter the recovery of seminal specimens and other sexual assault evidence. However, hygiene activities should not deter the clinician from the collection of evidence because perpetrator DNA has been recovered from the patient’s skin even after multiple showers. In addition, obtain a good history from the patient about potential injuries and any body trauma that may have occurred before the assault.

Elements of the victim’s history should help in deciding which potential samples to collect. For example, sperm may be recovered from the cervix for up to 19 days after intercourse and from the vagina for up to 10 days (Table 58.1).6 Cervical samples are now recommended in all cases that will involve speculum examination because of greater forensic yield.

<table>
<thead>
<tr>
<th>BODY CAVITY</th>
<th>MOTILE SPERM</th>
<th>NONMOTILE SPERM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vagina</td>
<td>6–28 hr</td>
<td>14 hr–10 days</td>
</tr>
<tr>
<td>Cervix</td>
<td>3–7 days</td>
<td>7.5–19 days</td>
</tr>
<tr>
<td>Mouth</td>
<td>—</td>
<td>2–31 hr</td>
</tr>
<tr>
<td>Rectum</td>
<td>—</td>
<td>4–113 hr</td>
</tr>
<tr>
<td>Anus</td>
<td>—</td>
<td>2–44 hr</td>
</tr>
</tbody>
</table>

Obtain a gynecologic history in preparation for documentation of injuries and treatment plans. From a medicolegal standpoint, inquire about any recent gynecologic surgical procedures or unintentional genital trauma that might alter the expected normal genital appearance. The history should also include the use of any method of birth control before the attack (with information regarding any missed birth control pills), last normal menstrual period, last voluntary intercourse, gravidity and parity, and recent STDs. As with all assaulted patients, the medical history should include current medications, tetanus immunization status, and allergies.

While taking the history, observe the patient’s ability to understand and respond appropriately to questions. Victims of sexual assault may not possess the capacity to consent to intercourse because of a developmental disability, young age, or intoxication with drugs or alcohol. Consider obtaining blood, urine, or both and testing for drugs or alcohol when the history suggests impaired consciousness. Patients who lack consenting capacity because of a developmental disability may have sufficient prior documentation of the condition. In the rare instance in which an examiner suspects a previously undocumented developmental disability, formal examination of the patient's mental capacity can be performed at a later time by request of the district attorney.

**Physical Examination**

Physical examination of a sexual assault patient differs from most other ED examinations in that examiners are not only caring for a patient’s physical and mental well-being but also investigating a crime scene and collecting specific evidence. As always, patiently explain every step of the examination process to the patient. Remind the patient to communicate any discomfort or questions during the examination and to ask for a break from the examination if needed. In addition, remind the patient of her right to decline any portion of the examination and the ability to stop at any point. Each patient should have the opportunity to have a family member, friend, victim advocate, or any combination of such individuals in the room during all parts of the examination, if they so desire.

**Collection of Clothing**

If not already collected by law enforcement, collect the clothes that the victim wore during the assault for potential evidence. Ask the patient to disrobe by dropping her clothes onto a large piece of paper or blue disposable underpad that is protected from the floor by a sheet. Using gloved hands, place each item of clothing in a separate paper bag. Label all collected material meticulously and describe it in the chart. Labels should include date, time, contents, and name of person who collected clothing. Bundle the paper or underpad and any material that might have fallen during the patient’s disrobing and place it in a separate paper bag. Minute amounts of blood, semen, and/or saliva can produce a DNA profile. When properly dried and free of moisture DNA can persist in swabs and on clothing for years at room temperature. Therefore, be sure that clothing is dried (but not by using heat) before packaging. The DNA profile obtained by the laboratory can be uploaded into the FBI's CODIS (Combined DNA Index System) database and identify an unknown assailant. When a victim’s clothing must be collected, be sure to provide suitable clothing for the patient to wear home after release from the ED.

**General Body Examination**

After the patient disrobes and is placed in a gown, examine her body for signs of trauma and foreign material. Uncover only one part of the body at a time to examine and then carefully re-cover it. This allows the patient to retain some modesty during the examination. Important areas for evaluation are the back, thighs, breasts, wrists, and ankles (particularly if restraints were used). Even in the absence of ecchymosis, note tender areas during the examination. Evidence from the physical surroundings of the assault can occasionally be found in the hair or on the skin. Retain such material as evidence. Document areas of trauma and evaluate further (e.g., with radiographs) as indicated by the type and extent of injury. Approximately 10% to 67% of sexual assault victims display bodily injuries. Document these injuries because they correlate significantly with successful prosecution of perpetrators. Bodily evidence may range from abrasions to major blunt force and penetrating trauma. If the victim has not bathed, bodily evidence in the form of dried semen stains may be visible on the hair or the skin of the victim. In a darkened room, dried semen (and, unfortunately, many other substances) on skin may fluoresce under examination with shortwave light, such as that produced by a Woods lamp or an alternate light source, but may also be noticed equally well by its reflective appearance under regular room lighting. Use moistened swabs to collect potential dried secretions; then air-dry them thoroughly and preserve as evidence. Dried secretions may be semen or saliva stains. If the history indicates oral contact or ejaculation in specific areas, these areas should be sampled using moistened swabs even if no dried secretions are visible. Fragments of the assailant’s skin, blood, facial hair, or other foreign material from the assault site may be trapped beneath a victim’s fingernails. Obtain fingernail scrapings by cleaning under a victim’s nails with a toothpick or small swab or by cutting the nails closely over a clean piece of paper. Fingernail samples should be collected in such a way as to maximize the detection of foreign DNA and minimize the amount of the patient’s cells present, using only light pressure to avoid any injury or bleeding to the patient. Fold the toothpick and debris into the paper, place it in an envelope, and package it with the other specimens.

**Imaging**

Photographs can be a valuable addition to the documentation of bodily injury. Medical institutions may employ professional-quality photographic teams; others must rely on law enforcement for photo documentation. Most institutions require patient consent for photographs taken by hospital personnel. Optimally, institutions should have a prearranged plan to handle film or digital media according to a written “chain of custody.” Alternatively, self-developing film (Polaroid) or instant digital prints that can be permanently labeled (e.g., subject, date, details of the pictured injury) may be used but will provide inferior resolution in most cases. The photographs should be labeled immediately and may be added to the legal evidence. In some jurisdictions, photographs of physical injuries will be taken and retained by an accompanying law officer. These photographs may serve as evidence or may simply refresh the examiner’s memory at the time of the trial.

**Oral Evaluation**

If indicated by the history, inspect the oral cavity closely for signs of trauma and collect evidence if indicated. Mouth injuries from forced oral copulation include lacerations of the labial
or lingual frenulum, mucosal lacerations, and abrasions (Fig. 58.2A). Injury to the lips is often produced by the patient’s own teeth as her lips are forced inward by forced oral penetration with the perpetrator’s penis. Potential injuries to the posterior pharyngeal wall and soft palate include contusions, submucosal hemorrhage, and lacerations (see Fig. 58.2B). Document these injuries at the initial examination because mucosal injuries heal quickly and may not be present hours or days later. Collect potential evidence with swabs rubbed between the teeth and the buccal mucosa on both the upper and lower gingival surfaces bilaterally. Spermatozoa have been identified in oral smears for hours after the attack despite brushing the teeth, using mouthwash, or drinking various fluids and may provide valuable evidence up to 12 hours after examination. Collect any foreign material (e.g., hair) to include as potential evidence. During the oral inspection, local law enforcement may request that examiners collect buccal cell swabs to provide the crime laboratory with a sample for victim DNA reference.

**Genital Examination**

Once the patient is in the lithotomy position, inspect the thighs and perineum for signs of trauma and for foreign material such as seminal stains. Use an ultraviolet light again to look at suspicious dried secretions. Many jurisdictions recommend routine collection of swabs from the external genital area because of the high likelihood of evidence being present after drainage of the vaginal vault and the inconsistent fluorescence of seminal fluid with a Wood’s lamp.

**Pubic Hair Samples**

If local crime laboratories request pubic hair samples, proceed with the following protocol. Before the pelvic examination, comb the patient's pubic hair for foreign material (particularly pubic hair belonging to the assailant). Place clean paper below the victim's buttocks with the patient in the lithotomy position and comb the pubic hair onto the paper. Fold these hairs and the comb into the paper and place them directly in a large paper envelope to be given to law enforcement. Foreign pubic hairs can often provide enough cellular DNA material from the root to enable the crime laboratory to perform DNA analysis. In addition, specialized laboratories possess the capability of performing mitochondrial DNA analysis from the hair shaft in many cases.

Significant hair transfer occurs in less than 5% of assaults. For the small minority of cases in which crime labs desire to compare foreign suspect hairs with the victim’s hair, a sample pulled from the patient may be requested. Although pulling the patient’s hair from the roots may provide the best sample, this collection method is painful, considered insensitive, and not recommended by these authors during the initial evaluation. Additionally, many agencies perform DNA testing on any hair recovered initially, rendering microscopic comparison with victim hairs superfluous. In the very rare case when it is requested, a patient can provide the hairs at a later time.

Genital examination of a sexual assault patient differs considerably from most ED pelvic examinations. First, carefully explain what you are doing to the patient. Perform a careful visual evaluation of the vulva and vaginal introitus for signs of trauma. The following techniques of separation and traction move the tissues that are most likely to suffer injury into view. In performing separation, use both hands to gently separate the labia laterally in each direction and inspect the posterior fourchette and vaginal introitus. Similarly, in performing traction, use both hands to hold each labium majus and apply gentle inferior labial traction (i.e., toward the examiner); this gives a much-improved view of the hymen, especially in prepubertal females (Fig. 58.3). If these maneuvers are not performed, traumatic genital injuries may be missed.

Familiarity with female (Fig. 58.4) and male (Fig. 58.5) genital anatomy, including all terms, is important for accurate descriptions. Although most novice examiners concern themselves with detecting injuries to the hymen, the majority of sexual assault–related vaginal injuries occur to the posterior fourchette (Fig. 58.6). In fact, hymenal injuries are rare in sexually active adult women and are more commonly observed in sexually inexperienced adolescents (Fig. 58.7). More uncommon injuries to the vaginal walls and cervix may be discovered during the speculum examination.

Reported rates of genital injury in forensically examined patients range from 6% to 20% without colposcopy to 53% to 87% with colposcopy. Most importantly, examiners must be cognizant of the fact that even completely normal findings on genital examination remain consistent with forced

**Figure 58.2** Oral injuries incurred as a result of sexual assault. A, Mucosal labial injury after forced oral copulation. B, Petechial hemorrhage seen on the soft palate after a similar assault.
**Figure 58.3** The normal hymen in a prepubertal female as seen with inferior labial traction.

**Figure 58.4** Normal female anatomy.

**Figure 58.5** Normal male anatomy. A, Circumcised. B, Uncircumcised.

**Figure 58.6** The majority of sexual assault–related vaginal injuries occur to the posterior fourchette. A, Superficial epithelial injuries (*arrows*) to this region may be difficult to appreciate. B, Application of toluidine blue dye highlights the injury.
Figure 58.7 Hymenal injury at the 6-o’clock position, usually found in adolescent girls. Such injuries are uncommon in adults.

sexual assault. In fact, a study of more than 1000 sexual assault patients found that almost half of all victims with forensic evidence positive for sperm had no genital injury.15

Colposcopy

Teixeira first described the use of colposcopy for documentation of sexual assault in 1981.15 Although it is not readily available, nor a standard of care in most EDs, colposcopy has revolutionized the documentation of injury. The colposcope provides magnification, a bright light source, and usually permanent documentation of injuries in the form of still images or video (mainly in digital format but occasionally traditional film). In one small study the colposcope increased the rate of detection of genital injury from 6% to 53%.16 Colposcopes with photo or video attachments provide excellent photographic documentation for the court and allow review by expert practitioners for court testimony without subjecting the victim to reexamination (Fig. 58.8A and B). Experienced sexual assault examiner programs are increasingly using high-quality digital single-lens reflex cameras mounted on a tripod to obtain excellent images that are indistinguishable from those obtained with colposcopy (see Fig. 58.8C). ED practitioners may have access to such equipment. Colposcopically visible injuries have also been described in adolescent women after the first consensual intercourse; hence, genital injury does not always translate to nonconsensual vaginal penetration.17 Conversely, totally normal findings on genital examination by colposcopy are often found after sexual assault. Even in sexually inexperienced adolescents, forced penetration can occur without leaving discernible genital injury.17 Although previous sexual experience by the victim decreases the likelihood of finding genital injury, experts cannot fully explain the reasons why some rape victims sustain measurable genital injury whereas others do not.

Forensic Evidence Collection

Protocols for collection of evidence vary by legal jurisdiction. The following discussion draws from the model protocol suggested by the state of California and the American College of Emergency Physicians manual.1

Obtain standard specimens during inspection of the external genitalia, rectum, vagina, and cervix. Lubricate the speculum as usual. Lubricant use is no longer considered a potential threat to evidence recovery and should be used for patient comfort per examiner protocol. Generally, the specimens collected will be determined by the patient’s history and local protocol, but they may include any of the items listed in Box 58.1 and shown in Fig. 58.9. Some protocols recommend that examiners make a wet mount of one swab from the vaginal pool and look at it under the microscope for the presence of motile sperm. Because of rapid cell death, studies have shown a negligible chance of finding motile sperm from a vaginal wet mount more than 8 hours after intercourse.18 Furthermore, in complying with the Clinical Laboratory Improvement Amendments of 1988, ED practitioners in the United States rarely have sufficient access or experience with microscopy to make this step a routine recommendation. Several swabs from the vaginal pool (including the one used to make the wet mount, if done) and the external genitalia should be obtained and then applied over clean slides for a dry mount. In addition, collect cervical swabs. The crime laboratory may recover sperm from cervical specimens up to 12 days or more after coitus.18 Label all slides and then allow them to air-dry. Package swabs and slides in paper envelopes for the local crime laboratory. Some EDs maintain specific equipment (i.e., a dry box) to aid in the drying of specimens; in others, the swabs and slides must be left out until completely dried.

Crime laboratories may also request collection of a vaginal washing, although this specimen is becoming obsolete with advances in DNA technology. For this procedure, insert 5 mL of sterile (but not bacteriostatic) water or saline into the vagina and then remove it. Place the washing in a sealed container (such as those used for urine collection or a red-topped blood tube). Label each sample separately and record in the chart the area from which the specimen was collected.

Due to the extreme sensitivity of new DNA tests, exercise caution to prevent contamination of samples. Wear gloves at all times during specimen collection and packaging. Take care not to inadvertently sneeze, spit, or cough on or near samples.

<table>
<thead>
<tr>
<th>Box 58.1</th>
<th>Potential Evidence Collected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clothing (list and describe all clothing collected on the chart and checklist form)</td>
<td></td>
</tr>
<tr>
<td>Debris</td>
<td></td>
</tr>
<tr>
<td>Dried secretions, swabs, and slides</td>
<td></td>
</tr>
<tr>
<td>External genital swabs and slides</td>
<td></td>
</tr>
<tr>
<td>Pubic hair combings</td>
<td></td>
</tr>
<tr>
<td>Oral mucosal swabs and slides</td>
<td></td>
</tr>
<tr>
<td>Anal swabs and slides</td>
<td></td>
</tr>
<tr>
<td>Vaginal pool swabs and slides</td>
<td></td>
</tr>
<tr>
<td>Vaginal lavage fluid in a test tube or urine container</td>
<td></td>
</tr>
<tr>
<td>Cervical swabs and slides</td>
<td></td>
</tr>
<tr>
<td>Tampon or condom present in the vagina (dried or frozen)</td>
<td></td>
</tr>
<tr>
<td>Urine or blood toxicology sample (timed collection)</td>
<td></td>
</tr>
<tr>
<td>Reference blood, buccal mucosa, and/or hair sample</td>
<td></td>
</tr>
<tr>
<td>Collect all evidence with gloved hands to avoid DNA contamination</td>
<td></td>
</tr>
<tr>
<td>Change gloves when necessary to avoid interlocation contamination</td>
<td></td>
</tr>
</tbody>
</table>
Genital Testing for STDs

Guidelines from the CDC no longer recommend routine testing but suggest tests be ordered on an individual basis using standard methods currently employed in most EDs. The majority of SART programs in the United States do not routinely perform STD testing on adults and adolescents. STD testing during sexual assault examination can detect only infection before the assault and provides no meaningful information for the crime laboratory. In addition, routine prophylactic treatment with...
antibiotics effective against *Neisseria gonorrhoeae*, *Chlamydia trachomatis*, and incubating syphilis makes detection of these preexisting infections superfluous; however, clinicians might want to consider obtaining samples for culture or nucleic acid amplification test from child victims, in whom the presence of an STD would be indicative of previous sexual contact.

**Perineal Toluidine Blue Dye Staining**

Toluidine blue dye is a nuclear stain, also used for cancer detection and mast cell staining, that highlights areas of injury. It adheres to areas denuded by abrasions and lacerations where the epidermal layer of nonnucleated cells has been removed (Fig. 58.10). The underlying nucleated cells take up the dye. Although it is not a uniform standard of care and is unavailable in many EDs, the dye can enhance the examiner’s ability to visualize and photographically document more subtle genital injuries (Fig. 58.11; also see Fig. 58.6B). Genital lacerations may provide corroborating evidence of nonconsensual intercourse, or at least sexual activity. To outline injuries, apply a 1% aqueous solution of toluidine blue dye to the perineum and wipe the excess dye off with a cotton ball moistened with lubricating jelly. A swab containing the dye is commercially available from several distributors (http://nphi.org/products.html). After the excess dye is removed, any areas that retain the stain signify a disruption in the epidermis, most likely injury. Separate any folds of the area and carefully examine them to avoid missing injuries. Ideally, apply the dye before speculum examination to eliminate the possibility of iatrogenic injury. The procedure is described in Fig. 58.10 and Box 58.2. In one study, the use of toluidine blue dye increased the injury detection rate from 16% to 40% in women, without the use of colposcopy19; however, injuries detected with the aid of toluidine blue dye are not 100% specific for sexual assault because such injuries have also been found after consensual intercourse, especially in adolescents.20

Collect external genital samples before the application of toluidine blue to avoid washing away potential DNA evidence. The use of toluidine blue dye itself does not interfere with DNA evidence from vaginal specimens, and it has proved safe for mucosal application.21,22

**Anal Evaluation**

The anal examination follows the genital examination in most cases (Fig. 58.12). Documentation of anal penetration holds
follows: separate the anal folds as much as possible by applying gentle lateral or vertical traction, clean the external anal skin with water and gauze, and then insert swabs approximately 2 cm into the anus. Gently move them in a circular motion and then remove them. Use the swabs to make slides, air-dry them, and then include them in the evidence sent to the crime laboratory. Anoscopy is indicated to document potential internal mucosal injuries in victims who report anal penetration or who describe loss of consciousness during the assault. Perform this procedure in the same manner as diagnostic anoscopy to evaluate other anal or rectal emergencies in the ED. It is best to collect the internal rectal specimen at the end of the anoscope to prevent contamination from any external sample being dragged internally. In one retrospective observational study of male patients, the use of anoscopy and colposcopy

**Figure 58.10** A, During a sexual assault, injuries are often multiple and typically occur between the 3-, 6-, and 9-o’clock positions. B and C, Traumatic skin injuries can be highlighted by applying toluidine blue to the perineum and vaginal area and then wiping it off to show the lesions. (A, From Marx J: Rosen’s emergency medicine: concepts and clinical practice. ed 6, Philadelphia, 2006, Elsevier.)

**Figure 58.11** Posterior fourchette injury A, before and B, after application of toluidine blue dye. Toluidine dye highlights an injury that would likely go unnoticed without dye application.

**BOX 58.2 Toluidine Blue Staining of the Perineum to Detect Microabrasion**

1. Collect all external genital specimens as indicated by examination before application of the dye.¹
2. Before speculum examination or anoscopy, apply 1% toluidine blue to the entire vulva (labia majora, labia minora, posterior fourchette, perineal body, and perianal area). The anus may also be stained. Do not use dye in the vaginal vault or mucous membranes.
3. Allow it to dry for approximately 1 minute.
4. Remove excess dye with water-soluble lubricant by gently blotting the area until the excess dye is removed.
5. DO NOT rub the area.
6. Photograph the area if indicated.
7. The dye will fade in 1 to 2 days.

¹A prefilled swab (T-Blue Swab/TBS, Tri-Tech, Inc.) is available through the National Forensic Nursing Institute.
²See Fig. 58.10.
³This does not interfere with DNA or semen testing.
Modified from The National Forensic Nursing Institute [http://nfni.org/forensichighlights.html]. This testing is not usually performed in the emergency department without a special assault team.
samples may include two buccal swabs collecting mucosal epithelial cells, a blood sample in a designated tube or filter paper card, or even two nasal swabs, especially useful in pediatric cases. The need for such samples will be determined by local protocol.

**Blood Tests**

Some crime laboratories request blood samples for DNA reference, toxicology analysis, or both. Include these samples with the material sent to law enforcement. When collecting blood for toxicology testing, record the exact date and time of collection on the specimen so that the criminalist may estimate the dose and timing of substances used by perpetrators to facilitate the assault.

**Urine Tests**

Perform bedside urine \( \beta \)-human chorionic gonadotropin \( (\beta\text{-hCG}) \) testing in all female patients of childbearing age to exclude preexisting pregnancy before administering pregnancy and STD prophylaxis. Collect the patient’s first available voided urine as requested by local crime laboratories to optimize recovery of potential toxicology evidence.

**Spermatozoa, Semen, and DNA Testing**

Motile and immotile sperm may be found microscopically in wet mounts of vaginal aspirates and in vaginal, oral, and rectal swabs. If formally trained, evaluate the slide microscopically immediately after the physical examination. Examiners may find sperm in vaginal wet mount specimens. Early discovery of sperm may be helpful to law enforcement investigations. However, most ED examiners lack formal training in this process and crime laboratories possess much higher sensitivity for detection of sperm, thus making a negative initial wet mount unhelpful. For these reasons, most examiners do not routinely perform the wet mount examination. After consensual intercourse with a normal ejaculate, laboratory testing of vaginal secretions detects sperm in 50% of specimens 4 days after coitus. However, despite penile penetration during sexual assault, crime laboratories may fail to detect sperm. Reasons for failure include inadequate specimen collection, degradation of the ejaculate, azoospermia, failure of the perpetrator to ejaculate, vasectomy in the perpetrator, washing by the victim, or use of a condom. Collection of relevant clothing may prove most important for DNA recovery as laboratories have recovered spermatozoa DNA from laundered clothing as long as 8 months after deposit.

A crime laboratory analyst initially looks for semen in a given sample by searching microscopically for sperm on a concentrated specimen and by testing for other components found in semen. The presence of spermatozoa is the only confirmatory test for semen. Seminal plasma components include p30 and acid phosphatase. p30 is a glycoprotein found in the prostate and other body fluids and is no longer regarded as conclusive evidence of semen (i.e., ejaculation within 48 hours). The presence of high levels of p30 is suggestive of a vasectomized or azoospermic male. Both p30 and acid phosphatase are presumptive evidence only because they can occur in other body fluids, such as vaginal secretions, male urine, etc.

Despite negative testing for seminal plasma components, laboratories may be able to detect male-specific “Y” DNA

---

**Figure 58.12** Anal injury is best seen with separation of perianal tissues. **A,** Anal tear in a 13-year-old boy after forced penile-anal penetration. **B,** Anal contusion and tear (arrows) in an adult male after forced penile-anal penetration. **C,** Multiple superficial tears to the anus and perianal region.

provided superior documentation of injuries over colposcopy alone. The location of anoscopically detected injuries may be recorded geographically on an imaginary clock face as with vaginal injuries.

**Reference Samples**

Crime laboratories may request reference samples from various locations on the patient’s body to use in comparison testing with potential evidence from the perpetrator. These reference
evidence from persistent sperm cells or the perpetrator’s epithelial cells.\textsuperscript{28,29} As DNA testing technology rapidly changes, the ability of crime laboratories to perform a specific forensic test varies by location and over time. Most crime laboratories look for unique short tandem repeats in perpetrator DNA with polymerase chain reaction amplification testing, which requires minimal material.

**Chain of Custody**

Samples and other evidence need to be carefully and formally transferred to the police, a crime laboratory, or a forensic pathologist. Label each sample with the patient’s name, hospital number, date, time of collection, area from which the specimen was collected, and collector’s name. Package these specimens according to local crime laboratory specifications and transfer them to the next appropriate official (police officer, pathologist, or other individual) along with a written confirmation of chain of custody, including a list of the specimens, the signature of each person who collected them, and the signature of each person who received them. If this documentation is inadequate or the documentation chain is broken, important evidence may be deemed inadmissible in court.

**TREATMENT**

**STD Prophylaxis**

The approach to prophylaxis for potential diseases transmitted by sexual assault varies by region, disease prevalence, and local practice and is often influenced by the emotional state of the victim and personal and religious viewpoints. We present an overview that may serve as a general guideline, with the understanding that many issues are vague, unsettled, and not totally adopted by all clinicians (Box 58.3).

It is advisable to address the issues of STD, pregnancy, psychological distress, and follow-up in the treatment of a patient with a history of recent sexual assault. Because infection rates before the assault are not known, the risk of contracting an STD as a consequence of a sexual assault has been difficult to determine, and estimates vary widely (Table 58.2).\textsuperscript{30} Jenny and colleagues found the post-assault incidence of STDs to be 2% for chlamydia and 4% for gonorrhea.\textsuperscript{31} The reported rates of 12% for *Trichomonas* and 19% for bacterial vaginosis seem high and may reflect a preexposure infection because male transmission of these infections, especially bacterial vaginosis, is uncommon. There is a lack of specific data on risk for the development of herpes, hepatitis B, or human immunodeficiency virus (HIV) infection from sexual assault. However, HIV transmission has been noted.\textsuperscript{32}

Because some patients tend to have relatively low compliance with keeping follow-up visits, most examiners offer, at the least, treatment of gonorrhea and chlamydia at the time of the initial examination.\textsuperscript{33} Though most often termed prophylaxis, technically this antibiotic administration is considered “treatment” given so early that the disease is subclinical. The need for routine administration of medication to combat *Trichomonas* is unclear, and many clinicians do not recommend it as a routine intervention. Prophylaxis for bacterial vaginosis is rarely suggested.

With the increasing prevalence of antibiotic-resistant *N. gonorrhoeae*, ceftriaxone in an intramuscular 250-mg dose is the CDC’s most current recommended treatment of choice in targeting gonorrhea after sexual assault. Ceftriaxone also treats incubating syphilis. No single-dose regimen for gonorrhea

### BOX 58.3 Recommended Empirical Emergency Department Testing and Pharmacologic Treatment After Sexual Assault

- NO routine STD cultures of the victim unless symptomatic for an STD or the victim is a child (a positive test in a child is indicative of abuse).
- Treat empirically for GC,\textsuperscript{3} *Chlamydia, Trichomonas* (optional), and bacterial vaginosis (optional) (see Box 58.4).
- Administer a tetanus booster vaccine if indicated.
- If not immunized for hepatitis B or unsure of the victim’s vaccination status, give first dose of vaccine empirically and follow with subsequent vaccination at 1 to 2 and 4 to 6 months. If previously vaccinated, offer hepatitis surface antibody testing with follow-up vaccination if the test result is negative. In addition to vaccination, consider HBIG in nonimmune patients after high-risk exposure to a known hepatitis B–positive perpetrator, followed by a full vaccination schedule.
- If the suspect is known or suspected to be HIV positive, treat with medications recommended by local ID experts or provided in occupational exposure kits or consider either of the following: Combivir (GlaxoSmithKline, Research Triangle Park, NC) (twice a day) or Truvada (Gilead Sciences, Inc., Foster City, CA) (once a day) for 28 days.
- If the suspect is not known to be HIV positive, there is no consensus on recommendations for treatment, and clinicians must consider each patient individually.\textsuperscript{*} This is an area of uncertainty. HIV prophylaxis may be given after a discussion of the risks and benefits with the patient.
- Obtain a pregnancy test and administer pregnancy prevention if not already pregnant and physically able to conceive (see Table 58.3).

**TABLE 58.2** Risk for Sexually Transmitted Diseases After Sexual Assault

<table>
<thead>
<tr>
<th>DISEASE</th>
<th>RISK (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gonorrhea</td>
<td>6–18</td>
</tr>
<tr>
<td><em>Chlamydia</em></td>
<td>4–17</td>
</tr>
<tr>
<td>Syphilis</td>
<td>0.5–3</td>
</tr>
<tr>
<td>Human immunodeficiency virus</td>
<td>&lt;1</td>
</tr>
</tbody>
</table>

\*Assistance with postexposure prophylaxis–related decisions can be obtained by calling the National Clinician’s Post Exposure Prophylaxis Hotline (PEP Line) (telephone: 888–448–4911).

is effective against coexisting C. trachomatis infection. Therefore give patients either a single dose of azithromycin (1 g orally) or a 7-day course of doxycycline (100 mg orally twice a day) or tetracycline (500 mg orally four times a day). A negative pregnancy test is a prerequisite for using either of the latter two antibiotics. Erythromycin may be used as a second alternative for Chlamydia prophylaxis in a pregnant patient. Some examiners administer prophylaxis for Trichomonas with a single 2-g oral dose of metronidazole or tinidazole; although effective and recommended by the CDC, these medications may cause significant nausea, vomiting, and diarrhea, which can interfere with the efficacy of pregnancy prophylaxis. One option to circumvent metronidazole-induced emesis interfering with efficacy of concomitantly administered medications is to delay administration by several hours. Box 58.4 provides several options for STD prophylaxis.

**Prevention of HIV Infection**

In a non–assault-related scenario, the risk for transmission of HIV from one episode of unprotected consensual receptive vaginal intercourse with an infected individual is approximately 1 in 1000. The incidence with unprotected receptive anal intercourse is significantly higher at 8 to 32 per 1000. However, sexual assault victims often sustain tissue injury because of the violent nature of the act, which may increase the rate of transmission of the virus. Although postexposure prophylaxis (PEP) for parenteral occupational exposure to infected body fluids (i.e., needlestick) is believed to be effective based on case-control studies, there is no proof that PEP for human sexual exposure prevents transmission of the virus. Furthermore, victims of sexual assault frequently arrive for treatment much later than those who have occupational exposures.

However, 40% of sexual assault victims fear contracting HIV after assault and should, at a minimum, receive counseling and, many argue, the option of taking anti-HIV medications because they may be effective. Unfortunately, immediate testing of the perpetrator remains a remote option. The majority of cases lack a perpetrator in custody for testing, and few state laws provide for mandated legal preconception HIV testing of alleged perpetrators. The CDC recommends administering PEP to sexual assault victims only in cases in which the perpetrator is known to be HIV positive. This recommendation specifies a 28-day medication course for patients with a history of sexual assault, who arrive for care less than 72 hours after the event, with an HIV-positive perpetrator, and when that exposure represents a substantial risk for transmission (i.e., sexual contact with genital secretions). As with occupational exposure, antiretroviral medications should be initiated as soon as possible after exposure. HIV seroconversion due to failures of PEP following sexual exposure have been reported (http://cid.oxfordjournals.org/content/41/10/1507.long). For sexual assault exposure with a perpetrator of unknown HIV status, the CDC states PEP must be addressed by practitioners on an individual case-by-case basis. Given the extreme negative outcome, the relative safety of treatment, and the lack of conclusive scientific evidence, at least two states have written policies legislating examiners to offer HIV PEP to all sexual assault victims, and practitioners must be aware of local mandates. The CDC guidelines provide a useful framework to approach individual decisions in prescribing HIV PEP (http://www.cdc.gov/std/tg2015/sexual-assault.htm).

**Pregnancy Prophylaxis**

Pregnancy occurs in up to 4.7% of sexual assault victims. An estimated 22,000 annual rape-related pregnancies could be avoided if all victims received pregnancy prophylaxis. As an initial step, perform a urine pregnancy test before administering postcoital contraception (PcC). Modern urine pregnancy tests possess a β-hCG detection threshold approaching 20 mIU/mL to 25 mIU/mL and will usually be positive 1 to 2 weeks after conception, often before a menstrual period is missed. Offer pregnancy prevention with available oral PcC to all patients with a history of sexual assault. Several methods can
be used to prevent pregnancy after forced, nonconsenting intercourse (Table 58.3). The drug of choice for PcC is levonorgestrel, which is available without a prescription in the United States in commercial kits called Plan B (Teva Women’s Health, Inc., Frazer, PA), Plan B One-Step (Teva Women’s Health, Inc.), Next Choice One-Dose (Watson Pharma, Inc., Parsippany, NJ), Take Action (Teva Pharmaceuticals, Israel), My Way (GAVIS Pharmaceuticals, Somerset, NJ), or other generics. In 2013, the US Food and Drug Administration (FDA) extended the approved sale of Plan B without a prescription to all women of childbearing status, from a more restrictive policy which limited its sale to individuals 17 years or older.31 Although the original Plan B kit included two pills containing 0.75 mg of levonorgestrel, each approved for administration 12 hours apart, a large World Health Organization (WHO) trial demonstrated that both pills may be taken at once with the same efficacy as the divided dose, with the potential for increased compliance.44 The newer Plan B One-Step is a single 1.5-mg pill to be taken as outlined in the WHO study. In the rare instance in which levonorgestrel is unavailable, there are several combined oral contraceptive pills, known as the Yuzpe regimen, that may be used for PcC (see Table 58.3).46 The Yuzpe method is somewhat less effective but prevents approximately 75% of pregnancies that would otherwise have occurred.43 Levonorgestrel prevents 89% of pregnancies that would otherwise have occurred and causes fewer side effects. Potential adverse side effects of both methods include nausea, vomiting, and breast tenderness. If the patient vomits within 1 hour of taking a dose, repeat the dose. Some practitioners routinely offer prophylactic antiemetic therapy, and such a strategy seems appropriate. Others reserve such treatment for patients who vomit. In 2010, the FDA approved a third option for pharmacologic emergency contraception, ulipristal acetate (brand name Ella [Afaxys, Inc, Charleston, SC]). This selective progesterone receptor modulator has been demonstrated to be as effective as levonorgestrel for prevention of pregnancy 72 hours after intercourse and more effective for longer postcoital use. Ulipristal requires a prescription and is approved for use up to 120 hours after intercourse.47 Despite taking PcC, patients who meet criteria for obesity may be two to four times more likely to become pregnant than those who do not meet obesity criteria, though this failure is somewhat less with ulipristal.46,47 All available evidence demonstrates no untoward effects on the fetus should pregnancy occur despite oral PcC.48 The common practice of obtaining written patient consent for these medications seems unwarranted. Insertion of a copper intrauterine device is another form of PcC and remains the most effective emergency contraception available, preventing more than 99% of pregnancies, but is generally outside the purview of the emergency physician.49

Unfortunately, religious preferences may deter some hospital EDs from providing PcC.50 In these instances, the website and number listed at the end of this paragraph provide practitioners information to give referral for easy access to PcC for patients who lack funds to pay for the medication. In addition, given the ever-increasing availability of new methods and drugs for PcC, examiners may want to obtain up-to-date information from this site (http://www.not-2-late.com; telephone: 1-800 -not-2-late). The site also provides links for patients to order levonorgestrel and ulipristal via courier, including methods for concomitantly obtaining a prescription for ulipristal.

**Psychological Support**

Sexual assault precipitates a psychological crisis for the patient, and psychological care should begin as soon as possible, ideally initiated when the patient first arrives in the ED.43,51 Reassure the victim that she will be in control of the examination, that she may ask questions at any point, and that she should notify the examiner if anything hurts or if she needs a break. Ensure that the patient has a complete understanding that in the ED she has full control over her body and the examination is the first step toward psychological support and healing. Unfortunately, if this is not made a priority, full recovery may be impaired. Posttraumatic stress disorder (PTSD), manifested as numbed responsiveness to the external world, disturbances in sleep, feelings of guilt, memory impairment, avoidance of activities, and other symptoms, often develops in sexual assault victims. Rape trauma syndrome is the specific label for PTSD in this population.51 Rape trauma syndrome stems from the following characteristics of sexual assault: (1) it is sudden and the victim is unable to develop adequate defenses, (2) it involves intentional cruelty or inhumanity, (3) it makes the victim feel trapped and unable to fight back, and (4) it often involves physical injury. Attention to the initial psychological care of a rape victim in the ED is fundamental and can reduce distress during forensic examination.52

Many areas have a local sexual assault crisis agency that can dispatch an advocate that will accompany patients with a history of sexual assault during the interview and examination. This same agency may help to provide the follow-up psychological support that must be offered to all victims. It is critical that all examiners maintain current contact information with these agencies and use their services. The importance of this contact is emphasized in some areas by the fact that state law dictates that medical personnel contact a local sexual assault crisis agency and use their services. The importance of this contact is emphasized in some areas by the fact that state law dictates that medical personnel contact a local sexual assault crisis agency when a victim arrives for examination (California penal code 264.2, Notification of a Counseling Center [Amended by Stats. 2014, Ch. 136, Sec. 1. Effective January 1, 2015]). In the absence of immediate local crisis services, a hospital social worker may fill this role.

---

**TABLE 58.3 Emergency Contraception**

<table>
<thead>
<tr>
<th>BRAND</th>
<th>MANUFACTURER</th>
<th>PILLS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Progestin-Only Emergency Contraception Oral Therapy: Recommended</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plan B One-Step</td>
<td>WCC</td>
<td>1 pill (levonorgestrel, 1.5 mg) immediately</td>
</tr>
<tr>
<td>Or</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Progesterone Receptor Agonist/Antagonist</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ulipristal (Ella)</td>
<td>Watson Pharma, Inc. Morristown, NJ</td>
<td>One 30-mg pill immediately</td>
</tr>
</tbody>
</table>

*For the most up-to-date and alternative recommendations and general information, call 1-800-not-2-late or view [http://www.not-2-late.com](http://www.not-2-late.com). Some regimens cause nausea and an antiemetic may be used. If vomiting occurs, repeat the dose of antiemetic. Alternatively, the older Yuzpe method may be used with any combination oral contraceptive containing levonorgestrel, ethinylestradiol 0.03 mg/0.3 mg dosed as two pills immediately and two pills after 12 hours.
Postexamination Follow-Up

Medical and psychological follow-up of sexual assault victims is essential. Unfortunately, less than one-third of victims complete follow-up medical care. Many protocols recommend a 2-week follow-up to reexamine any injuries and to repeat testing for STDs and pregnancy. The timing of this follow-up seems less important given the widespread use of prophylactic medication to prevent STDs and pregnancy. However, because of the measurable failure rate of PeC, repeated pregnancy testing is critical for a victim who does not experience an expected menses. Further follow-up evaluations may be performed at 4 or 6 weeks and 3 and 6 months to repeat serologic tests for HIV, hepatitis B, hepatitis C, and syphilis. In addition, local volunteer support groups can be of immense assistance to a sexual assault victim; contact with such a group should be offered to each victim.

SPECIFIC POPULATIONS

Male Evidentiary Examinations

Male evidentiary examinations include all the same forensic evidence collection as for female victims except vaginal and cervical specimens. As with all patients who have been sexually assaulted, the forensic examination is guided by the history of events related by the patient. Most male victims suffer from anal penetration, or sodomy, by a perpetrator. In addition, male victims may also suffer from rape trauma syndrome but may be less likely to get the psychological support they require. The increased risk for transmission of HIV with anal intercourse has been noted (see earlier section on Prevention of HIV Infection). Because of the extreme emotional reaction that men often feel after a sexual assault, they report the crime even more sporadically than female victims do. Male patients with a history of sexual assault deserve the same unhurried, nonjudgmental, compassionate care that female victims do. Penile samples from the shaft, glans, corona, and scrotum may be obtained if there is oral or anal contact with the perpetrator.

Child Sexual Assault Examinations

In general, the care and treatment of a pediatric sexual assault patient requires expert knowledge and experience. Frequently, ED practitioners are the first professionals to examine a child victim, and based on the history provided, the presence of obvious genital injury and trauma should be an adequate prompt to manage the forensic process. However, in less obvious cases, the subtle variations in developmental changes and congenital anomalies may leave many clinicians ill equipped to render an opinion concerning findings indicative of sexual assault. The lives of children and families may be disrupted or severely affected, depending on the practitioner’s opinion of the presence of genital penetration-type findings. The history in these cases can be challenging to obtain given the age of the patient, their developmental stage, and psychological state. Emergency providers must remain vigilant for any clues, no matter how insignificant they may appear initially.

A well-known study by Adams and associates demonstrated that the majority of children reporting sexual abuse have normal or nonspecific genital findings. As these authors succinctly stated regarding child sexual assault, it is “normal to be normal.” Despite expert physical examination, the vast majority of sexually abused children cannot be differentiated from nonabused children. Discovery of one of the rare examination markers of injury should be confirmed by experts, and a discussion of these findings, though beyond the scope of this chapter, is covered extensively in other resources. The potential sexual assault history provided by the child or caretaker should therefore remain the primary indicator that inappropriate genital contact has occurred. Any concerning elements of the history warrant an investigation of the possibility of sexual abuse.

It cannot be emphasized enough that the examiner’s responsibility in the care of a child victim of sexual abuse remains within the realm of experts. However, in EDs lacking timely availability of local experts, inspect the genitalia carefully in an unhurried, child-friendly manner, and if indicated, collect forensic specimens. For a very young child with small genital orifices, the aid of a magnification source may be extremely helpful. Ask a parent (unless a suspect) to assist in the calming, reassurance, and positioning of the child for careful inspection. Whereas the basic lithotomy position may be used for an older, more mature child or an adolescent patient, use of alternative positioning of a pediatric female patient is essential for inspection. The frog leg position (the feet together and the knees spread widely apart) with the use of labial or gluteal (or both) separation and traction is often beneficial in children (Fig. 58.13). Take care to gently separate the labia to avoid superficial examiner-induced injuries. In addition, to get a better look at the hymenal perimeter in prepubertal girls and the anus in girls and boys, ask them to turn over into the knee-chest position (Fig. 58.14). Genital findings that are deemed definitive of

![Figure 58.13](image1.png) “Frog leg” position to examine children.

![Figure 58.14](image2.png) Child in the knee-chest position to facilitate examination of the hymen.
ACUTE TRAUMA TO EXTERNAL GENITAL/ANAL TISSUES, WHICH COULD BE ACCIDENTAL OR INFlicted

- Acute laceration(s) or bruising of labia, penis, scrotum, perianal tissues, or perineum.
- Acute laceration of the posterior fourchette or vestibule, not involving the hymen.
- Residual (healing) injuries to external genital/anal tissues. (These rare findings are difficult to diagnose unless an acute injury was previously documented at the same location.)
- Perianal scar.
- Scar of posterior fourchette or fossa.

INJURIES INDICATIVE OF ACUTE OR HEALED TRAUMA TO THE GENITAL/ANAL TISSUES

- Bruising, petechiae, or abrasions on the hymen.
- Acute laceration of the hymen, of any depth; partial or complete.
- Vaginal laceration.
- Perianal laceration with exposure of tissues below the dermis.
- Healed hymenal transection/complete hymen cleft (a defect in the hymen between 4 o’clock and 8 o’clock that extends to the base of the hymen, with no hymenal tissue discernable at that location.)
- A defect in the posterior (inferior) half of the hymen, wider than a transection, with an absence of hymenal tissue extending to the base of the hymen.

INFECTIONS TRANSMITTED BY SEXUAL CONTACT, UNLESS THERE IS EVIDENCE OF PERINATAL TRANSMISSION OR CLEARLY, REASONABLY, AND INDEPENDENTLY DOCUMENTED BUT RARE NONSEXUAL TRANSMISSION

- Genital, rectal, or pharyngeal Neisseria gonorrhoea infection.
- Syphilis.
- Genital or rectal Chlamydia trachomatis infection.
- Trichomonas vaginalis infection.
- HIV, if transmission by blood transfusion has been ruled out.

DIAGNOSTIC OF SEXUAL CONTACT

- Pregnancy.
- Semen identified in forensic specimens taken directly from a child's body.

These findings support a disclosure of sexual abuse and are highly suggestive of abuse even in the absence of a disclosure, unless a timely and plausible description of accidental injury is provided by the child and/or caretaker. Physical findings should be confirmed using additional examination positions and/or techniques. Diagnosis of sexually transmitted infections must be confirmed by additional testing to avoid assigning significance to possible false positive screening test results. Photographs or video recordings of these findings should be evaluated and confirmed by an expert in sexual abuse evaluation to ensure accurate interpretation.


Suspect Examinations

As forensic evidence collection in the form of DNA retrieval continues to evolve, EDs may see more requests from local law enforcement for collection of evidence from suspects. EDs should be familiar with local and state protocols, especially regarding consent. Some jurisdictions permit examination of suspects without consent, given the imminent degradation of potential biologic evidence. Other jurisdictions require that suspects give consent or, at the very least, that police obtain a search warrant from the court. The sooner that a suspect is

sexual abuse or penetration or are nonspecific are included in Box 58.5. However, many normal hymenal differences exist from one child to the next, and the definitive diagnosis of “abnormal” is often difficult even for experts. When any doubt exists in the ED, completely describe the findings and refer the child for later examination by experts. The availability of a colposcope or alternative photographic equipment with magnification clearly aids in the documentation of any injuries that may heal before examination by an expert can be performed.

When disclosure or genital injuries confirm possible penetration of the child, collect specimens for potential evidence. On all conscious prepubertal children, collect the specimens without inserting a pediatric speculum. If there is no bleeding or significant trauma, procedural sedation is rarely indicated. If a child proves to be too uncooperative for an ED examination, refer the patient to a child sexual assault expert for examination in a controlled setting or make the appropriate referral to a specialist. For the rare cases involving severe vaginal trauma or suspected internal genital injury (active bleeding) that will possibly require surgical repair, conduct the examination under deep procedural sedation or refer for examination by a consulting gynecologist under general anesthesia. External anal and vulvar swabs are usually collected quite easily; however, lack of estrogen in prepubertal children may increase hymenal sensitivity making vaginal samples difficult to obtain. Therefore many local protocols do not require collection of internal samples. For extraordinary circumstances, internal samples should remain the very last evidence collected. Make every effort possible to avoid swab contact with hymenal tissue during collection. Vaginal aspirates can be obtained with a feeding tube or plastic angiocatheter and may provide an alternative to vaginal swabs.

Collecting genital specimens to screen for STDs remains a controversial issue in the realm of child abuse experts. For N. gonorrhoeae, at least, the literature supports the notion that all infected children will display an abnormal discharge. With very young children, the practitioner may have only one opportunity to collect vaginal specimens without causing agitation and prohibiting further examination. Forcing specimen collection under physical restraint is considered a second assault on the child. Because the child is being evaluated for possible sexual abuse, the primary specimens collected should be for forensic DNA analysis. STD detection and treatment can be performed at a later time. Clinicians should consult local child abuse centers and state agencies for protocols regarding immediate STD specimen collection, referral, and follow-up services.
apprehended and brought in for a medical-forensic examination, the better the quality of forensic biologic evidence.

Performing a medical-forensic examination on a suspect can give important corroborating information for the investigation of a crime. It can also help exonerate the innocent. Law enforcement should be in attendance during the examination of any suspect to ensure the safety of the examiner, the witness, and the cooperation of the suspect. Be sure that the suspect and victim do not encounter one another in the hospital setting during the examination period. Examine the victim and suspect in separate locations within the ED or clinical setting. It is extremely beneficial to conduct the examination and history of the suspect before the victim’s examination, to search for physical findings on the suspect that were indicated by the victim’s history. For example, if during the victim’s history she relates that she scratched the suspect’s left shoulder in defense, the examiner should examine, document, and preferably photograph the presence (or absence) of an injury on the suspect’s left shoulder.

The physical and evidentiary examination of the suspect is similar to that of the victim. The primary differences lie in history taking, reference samples, and more “blind” samples. During the examination of a suspect, law enforcement officers, rather than the suspect, provide the history of the event. Collect reference samples of blood and buccal swabs if possible. Previously recommended, head and pubic hair reference samples are no longer required in most areas and practitioners should refer to local protocols for guidance on this. Apply special attention not only to nail scrapings but also to swabbing all the fingers for possible vaginal epithelial cells from digital penetration. Female DNA persists in the male fingernail scrapings for up to 6 hours or more. The shaft and corona of the glans penis should be swabbed along with separate swabs of the scrotum for vaginal secretions. Criminalists often recover a female victim’s DNA from the scrotal swab even if the perpetrator wore a condom. With an unwashed penis, swabs almost uniformly show evidence of female cells up to 24 hours after coitus.

Examinin suspect requires the same amount of professional sensitivity and respect that any patient receives within the ED. It is not within the realm of the clinician’s expertise to determine whether the suspect is guilty or innocent.

### The Unconscious Victim and “Drug-Facilitated Sexual Assault”

Alcohol and other drugs play an important role in many sexual assaults. Half of all sexual assaults involve drug or alcohol ingestion. In many cases it is unclear whether a drug was taken voluntarily or whether it was surreptitiously given to the assaulted victim.

Popular media has raised public awareness of drugs used to facilitate sexual assault under the term date-rape drugs (Box 58.6). Although date-rape drugs are of significant concern, extensive forensic testing in the United States shows that only a minority of sexual assault cases involve the suspect covertly spiking a victim’s drink with a tablet, capsule, powder, or liquid mind-altering drug. A routine hospital urine drug screen is inadequate to evaluate a sexual assault victim. Forensic laboratories usually offer an evaluation for multiple drugs in a specific test designed for the sexual assault victim. However, testing may not be adequately sensitive to test for all substances used during drug-facilitated sexual assault. The drugs most commonly associated with drug-facilitated sexual assault are ethanol, marijuana, cocaine, and benzodiazepines. Zolpidem, a common sleeping medication, has been used, but it is difficult to detect and only transiently positive in urine samples. Frequently, more than one drug is found. Although any type of sedative or hypnotic drug, or a combination of both, may be used to facilitate sexual assault, the most publicized drugs include flunitrazepam (Rohypnol), γ-hydroxybutyrate (GHB), and most recently, beverages containing high amounts of alcohol and caffeine (e.g., Four Loko [2017 Phusion Projects, Chicago IL]). In previous decades, laboratory testing implicated flunitrazepam and GHB in drug-facilitated sexual assault in approximately 3% to 5% of cases. Flunitrazepam is a benzodiazepine unavailable in the United States but available in Mexico. It can be detected in urine up to 3 weeks after ingestion. In the United States, GHB is a schedule 1, federally banned central nervous system depressant. Legally, it is available only by prescription as the drug Xyrem (Jazz Pharmaceuticals plc, Dublin, Ireland) for narcolepsy with a schedule 3 exception, but it can easily be manufactured illegally by users. It can be detected in drinking material residue by crime laboratories, as well as in the victim’s urine, up to 4 hours after a sufficiently large ingestion. Drugs similar to GHB are 1,4 butanediol and γ-butyrolactone.

Often, the victim’s last memory is of using drugs or alcohol and then passing out. A common scenario is for the victim to have one glass of wine (or another usual drink), suddenly feel nauseated, and then wake up hours later in a different location and lacking intervening memory. Some remember short segments of activity that may indicate some type of sexual acts. Victims may lack any memory at all but desire to be “checked” for intercourse. A comprehensive medical-forensic examination should be conducted on these individuals. Without

### BOX 58.6 Possible Date Rape Drugs

<table>
<thead>
<tr>
<th>Drug</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alprazolam</td>
</tr>
<tr>
<td>Ketamine</td>
</tr>
<tr>
<td>Amphetamines</td>
</tr>
<tr>
<td>Lorazepam</td>
</tr>
<tr>
<td>Barbiturates</td>
</tr>
<tr>
<td>Meprobamate</td>
</tr>
<tr>
<td>1,4-Butanediol (BD)</td>
</tr>
<tr>
<td>Methamphetamine</td>
</tr>
<tr>
<td>γ-Butyrolactone (GBL)</td>
</tr>
<tr>
<td>Midazolam (Versed)</td>
</tr>
<tr>
<td>Cannabis</td>
</tr>
<tr>
<td>Diazepam</td>
</tr>
<tr>
<td>phenylethylalanine (PCP)</td>
</tr>
<tr>
<td>Cocaine</td>
</tr>
<tr>
<td>Chloral hydrate</td>
</tr>
<tr>
<td>clonazepam</td>
</tr>
<tr>
<td>Clonidine</td>
</tr>
<tr>
<td>Diazepam</td>
</tr>
<tr>
<td>clonazepam</td>
</tr>
<tr>
<td>Diazepam</td>
</tr>
<tr>
<td>Ethanol</td>
</tr>
<tr>
<td>Flunitrazepam (Rohypnol)</td>
</tr>
<tr>
<td>γ-Hydroxybutyrate (GHB)</td>
</tr>
<tr>
<td>Flunitrazepam (Rohypnol)</td>
</tr>
<tr>
<td>Phencyclidine (PCP)</td>
</tr>
<tr>
<td>Flunitrazepam (Rohypnol)</td>
</tr>
<tr>
<td>Temazepam</td>
</tr>
<tr>
<td>Flunitrazepam (Rohypnol)</td>
</tr>
<tr>
<td>Triazolam</td>
</tr>
<tr>
<td>Flunitrazepam (Rohypnol)</td>
</tr>
<tr>
<td>Zolpidem</td>
</tr>
</tbody>
</table>

*Frequently, more than one drug is found. Most common are alcohol, marijuana, cocaine, and benzodiazepines; others account for less than 5% of positive tests.

Will not be detected on a routine immunoassay drug screen. A more detailed analysis will be required.

a history from the victim, examiners should collect samples from every potential oral or genital contact, including the neck, breasts, vulva, and from all orifices (oral, vaginal, and anal). Obtain samples of both blood and urine, if possible, for toxicology (including ethanol), with exact times of collection documented. Remember to collect the first voided urine to optimize potential recovery. Many of the drugs used to facilitate sexual assault are not found on routine hospital laboratory testing. Some forensic laboratories offer a “date-rape panel” that tests for a variety of commonly used substances. Obviously, a positive drug test does not prove date rape, and it may be impossible to distinguish self-administration from clandestine ingestion.

Extreme sensitivity must be used when discussing positive genital findings with a victim who has no memory of any sexual activity. Many times, the imagined sexual acts can create just as severe a traumatic response as an actual remembered sexual assault. For the unconscious victim, there is no memory of events to fill in the blanks, only his or her terrifying imagination of what could have happened.

LEGAL ISSUES

When the local government decides to proceed with a sex crime case against an alleged perpetrator, the district attorney will commonly contact the examiner to give legal testimony. A concise and complete well-documented chart often negates the need for a clinician to appear in court. When required for this task, it is best for the examiner to work with the prosecuting attorney to prepare testimony. As is the case for all ED patients, chart notes should be written objectively and carefully in the expectation that the ED evaluation and evidence collection may be presented in court. In some jurisdictions it is possible to minimize the time spent away from work by arranging to be called to the courtroom just before the time of testimony or by giving a deposition before the court date. Once on the witness stand, the examining clinician is most often considered a percipient witness and not necessarily an expert in the area of sexual assault. The law requires that one testify only to one’s best recollection and to what is indicated in the chart. Factual information in answer to questions should be given only if one knows the facts; assumptions should be avoided. One should not be afraid to acknowledge the limits of one’s knowledge or expertise. Statements such as “there were marks on the body that were consistent with bite marks” are preferable to statements such as “there were bite marks.” The court decides if a person was sexually assaulted, and the clinician is there to give information about the patient’s findings and statements, what was found, and what was done for treatment.

SARTS

Before the 1990s, sexual assault examinations were mostly the responsibility of emergency clinicians. However, since the early 1990s, nurses or nurse clinicians have been performing an increasing number of sexual assault examinations. Called sexual assault nurse examiners (SANEs), these nurses are the core members of SARTs. Other members of SARTs include law enforcement individuals, victim advocates, prosecutors, and forensic laboratory personnel.

Most examinations still take place in the ED but may be done in a space near the ED or an affiliated clinic. To establish SARTs, extra funding by government or a charitable organization is often needed because many local police jurisdictions often do not have the necessary funding to adequately reimburse for the evidentiary examination to support a program. However, law enforcement is increasingly willing to pay more for a SANE-performed forensic examination because they believe that it provides superior documentation for legal proceedings. Nurse examiners have formed the International Association of Forensic Nurses. This group has drafted standards of practice for sexual assault examiners’ education and the examinations themselves. Advantages of SARTs using SANEs include the following:

1. The practitioner performing the examination is specifically dedicated to treating the patient with a history of sexual assault, not tending to multiple patients in a busy ED.
2. The clinician has usually completed more extensive training on sexual assault examination (mean of 80 hours) and evidence collection and, accordingly, may perform a more comprehensive examination with better collection of evidence.
3. Many involved feel that designated clinicians understand and consider the emotional needs of the victim more fully because of their extra specialty training.

Useful guidelines and resources for establishing SANE programs are currently available.1,66,67

ACKNOWLEDGMENT

The editors and author wish to acknowledge the contributions of G. Richard Braen to this chapter in previous editions; Beverly Kerr, Supervising Criminalist at Los Angeles County Sheriff’s Department; Mary Hong, Sr. Forensic Scientist-DNA Supervisor at Orange County Crime Laboratory; and Patricia A. Huck, Criminalist II of the LAPD Crime Laboratory for their technical forensic advice.


This page intentionally left blank
Headache and head injury are encountered commonly in the emergency department (ED). If either is accompanied by vomiting, decreased level of consciousness, or abnormal vital signs, the possibility of increased intracranial pressure (ICP) should be considered. Acutely increased ICP is a neurologic emergency that must be managed rapidly to prevent further brain damage and death. In some cases the accompanying clinical symptoms may be vague or subtle and make diagnosis difficult. Familiarity with the pathophysiology of increased ICP facilitates its diagnosis and management.

**PATHOPHYSIOLOGY OF ICP**

Alexander Monro, an anatomist in the 18th century, described the intracranial contents as containing a fixed volume. The fixed-volume theory was supported by George Kellie a few years later and became known as the Monro-Kellie doctrine. This doctrine has since guided our understanding of intracranial dynamics and the principles of autoregulation.

The components of the calvaria are the brain parenchyma, cerebrospinal fluid (CSF), the venous blood supply, and the arterial blood supply (Fig. 59.1). CSF and the venous blood supply have the greatest ability to change their volume to compensate for increases in pressure. These dynamic changes in the relative proportion of the cranial content may not affect the patient if ICP is not excessive. However, if a pathologic process overwhelms the compensatory mechanisms, the result will be a nearly exponential increase in ICP (Fig. 59.2).

Normal supine ICP ranges from 5 to 15 mm Hg. Transient increases in ICP as high as 80 to 100 mm Hg occur with coughing or straining. Other factors that can transiently increase ICP are movement, pain, and fever. A space-occupying lesion such as a tumor, hematoma, abscess, or foreign body can also raise ICP. Fig. 59.3 demonstrates that the area and cause of the increased ICP will determine where shifts occur to result in brain herniation.

**Brain**

Brain volume can be increased by edema, idiopathic intracranial hypertension (IIH), tumor, or bleeding. The three types of edema are vasogenic, cytotoxic, and interstitial. Vasogenic edema results from increased permeability of the capillaries, which leads to passage of excess fluid into the extracellular space. Cytotoxic edema is due to accumulation of intracellular fluid in brain tissue (neurons and glia) secondary to dysfunction of the adenosine triphosphatase pump. Interstitial edema occurs when fluid accumulates as a result of blockage of CSF absorption.

IIH, formerly known as pseudotumor cerebri or benign intracranial hypertension, is a chronic condition characterized by increased CSF pressure not caused by a tumor, edema, hydrocephalus, or change in CSF composition. It occurs most frequently in obese women. Symptoms may include headache, nausea, and blurry vision. The headache is typically worse on waking or with exertion. In general, patients with IIH have normal findings on neurologic examination except for the frequent presence of papilledema. The precise pathophysiology behind IIH remains unclear, with several proposed etiologies, including: cerebral venous outflow abnormalities (e.g., venous stenosis); increased CSF outflow resistance at either the level of the arachnoid granulations or CSF lymphatic drainage sites; obesity-related increased abdominal and intracranial venous pressure and obesity-related chronic inflammation; altered sodium and water retention mechanisms; abnormalities of vitamin A metabolism; and sex hormone dysfunction.1-4 Severe and untreated cases of IIH can lead to permanent vision loss. There appears to be some crossover with the more serious and life-threatening diagnosis of cerebral venous thrombosis (CVT), and therefore magnetic resonance (MR) (ideally with venography) or computed tomography (CT) venography of the cerebral venous system is recommended in patients with presumed IIH to rule out CVT.4-5 Other commonly associated conditions are listed in Box 59.1.

Brain tumors encompass neoplasms that originate in the brain itself (primary brain tumors) or involve the brain as a metastatic site (secondary brain tumors). Primary brain tumors include tumors of the brain parenchyma, meninges, cranial nerves, and other intracranial structures (the pituitary and pineal glands) (Fig. 59.4). Primary central nervous system lymphoma refers to high-grade B-cell non-Hodgkin’s lymphoma confined to the central nervous system. Secondary brain tumors, the most common type, originate elsewhere in the body and metastasize to the intracranial compartment.
CSF

CSF is produced by the choroid plexus at a daily rate of 400 to 600 mL (~ 20 mL per hour) with complete turnover of total volume occurring four to five times per day in young adults (Fig. 59.5). It flows from the ventricles into the cisternae of the subarachnoid space and is drained by the arachnoid villi of the
dural sinuses to maintain a constant volume of 100 to 150 mL. Obstructive hydrocephalus occurs when flow is blocked at any point in the ventricular system by clotted blood, tumor, colloid cyst, edema, or primary stenosis. Communicating hydrocephalus is due to impedance of flow beyond the ventricular system at the level of the basal cisternae or lack of absorption by the arachnoid villi. Communicating hydrocephalus can occur with both infection and subarachnoid hemorrhage (Fig. 59.6).

**Blood**

Up to a certain range, cerebral blood flow (CBF) is maintained by an autoregulatory mechanism despite fluctuations in cerebral perfusion pressure (CPP) (Fig. 59.7). Constant CBF can typically be maintained at any CPP between 60 and 160 mm Hg. Once CPP is out of the autoregulatory zone, CBF is linearly related to CPP. CPP lower than 60 mm Hg can lead to ischemia, whereas CPP higher than 160 mm Hg can result in hypertensive encephalopathy.

**SIGNS AND SYMPTOMS**

Findings on neurologic examination can be normal in a patient with a mild increase in ICP because of the brain’s compensatory mechanisms. Patients with a complaint of headache or head
injury may not initially manifest the more dramatic and worrisome symptoms of increased ICP such as vomiting, syncope, altered mentation, or Cushing’s reflex (bradycardia, increased blood pressure, and irregular respirations). ICP correlates poorly with clinical symptomatology. One of the earliest clinical signs is decreased venous pulsation on funduscopic examination, but this may be difficult to appreciate in an acutely ill patient in a busy ED. Moreover, the initial findings on head CT might not reveal the true extent of injury, especially with early stroke or when DAI is involved. However, as compensatory mechanisms fail, CT findings, as well as clinical symptoms, will become more obvious.

Signs and symptoms of severely increased ICP include a decreasing level of consciousness, papilledema, cranial nerve palsies, and lateralizing neurologic deficits. When any of these are noted, particularly when CT confirms the presence of a mass effect, such as hydrocephalus or a midline shift, urgent intervention is necessary (Fig. 59.8). Neurosurgical consultation for possible invasive measures to reduce ICP is indicated. Medical management of increased ICP should also proceed without delay.

**MEDICAL TREATMENT OF INCREASED ICP**

**Oxygenation**

Management of the airway and breathing is paramount in patients with brain injury. If the patient is hypoxic, supplemental oxygen is critical in preventing further ischemia. Patients with a Glasgow Coma Scale (GCS) score of 8 or lower or with impending signs of inadequate respiratory status should undergo rapid-sequence intubation (RSI) to protect their airway and better control blood levels of oxygen and carbon dioxide (partial pressure of oxygen [PO2] and partial pressure of carbon dioxide [PCO2]). Apneic patients should receive bag-valve-mask ventilation while preparing for intubation, with specific avoidance of hyperventilation, and apneic oxygenation via a high-flow nasal cannula (rate of 15 L/min or higher) during airway securement.

**Sedation and Paralytics**

A rapid neurologic examination that assesses the patient’s pupillary reflexes, motor response to voice and painful stimuli, and the ability to interact with his/her surroundings should be performed before sedation and RSI. To blunt the hemodynamic response to intubation, fentanyl, at an intravenous (IV) dose of 2 to 5 µg/kg, can be given over 30 to 60 seconds while preoxygenating the patient. Fentanyl is an excellent drug for control of pain, which if untreated, can lead to increased ICP. Fentanyl can also be given during the pretreatment phase of RSI (3 minutes before administration of the paralytic agent) if time allows. Caution should be exercised, however, to avoid precipitous drops in blood pressure, which can threaten CBF and exacerbate brain injury. Therefore it should be avoided in hypotensive patients and hypovolemic patients. We also recommend caution in brain-injured patients with concomitant trauma, who may have exaggerated episodes of hypotension with narcotics and sedatives. Pretreatment lidocaine, which was traditionally believed to decrease the sympathomimetic response to intubation, is no longer recommended as multiple studies have shown conflicting and insufficient evidence to support its use. A defasciculating dose (typically 1/10 the intubating dose) of a nondepolarizing neuromuscular blocking agent given prior to intubation has also fallen out of favor as no sufficient evidence has supported its use.

In patients who are normotensive or hypotensive, etomidate (0.3 mg/kg IV) is the induction agent of choice due to its minimal effect on systemic blood pressure and its lowering of ICP. Ketamine, at a dose of 1 to 2 mg/kg IV, can also be used as it does not appear to increase ICP, as previously believed, and should be strongly considered in severely hypotensive patients due to its ability to increase blood pressure and its analgesic effects that minimize the adverse sympathetic stimulation of laryngoscopy. In hypertensive patients, thiopental (50 to 100 mg IV or 3 to 5 mg/kg IV) or propofol (1 mg/kg IV) can be used for induction. Propofol has gained acceptance as a sedative in patients with increased ICP because of its short duration of action and depression of cerebral metabolism and oxygen consumption. This may have a neuroprotective effect. However, with higher doses propofol can cause profound decreases in systemic blood pressure and extended periods of use can be associated with significant morbidity.

Succinylcholine (1.5 mg/kg in adults and up to 2.5 mg/kg in pediatric patients) is the ideal paralytic agent in patients with increased ICP due to its rapid onset, short duration of
action, and consistent and reliable effects.\textsuperscript{15} Nondepolarizing agents (e.g., rocuronium) are appropriate when contraindications to succinylcholine exist (see Chapter 5). Sedatives and paralytics should be short-acting to facilitate close monitoring of the patient’s neurologic status.

**Oxygenation and Hyperventilation**

As the airway is secured, adequate supplemental oxygen should be provided, with titration down rapidly from the initial fraction of inspired oxygen of 1.0 used for RSI to ensure oxygen saturation greater than 90%.\textsuperscript{2}

In general, avoid hyperventilation in patients with brain injury as low P\textsubscript{CO\textsubscript{2}} levels cause cerebral vasoconstriction, resulting in decreased CBF in the critical hours following injury.\textsuperscript{6,8} Hyperventilation is also associated with poor survival and neurologic outcomes.\textsuperscript{16,17} For the majority of brain-injured patients, the target is thus eucapnia with a P\textsubscript{CO\textsubscript{2}} of 35 to 40 mm Hg.\textsuperscript{16} Nevertheless, if a patient displays evolving signs of brain herniation (e.g., anisocoria, hemiparesis, asymmetric posturing, Cushing’s reflex, or rapid deterioration in GCS score), hyperventilation may be necessary to arrest the process. Current recommendations target a P\textsubscript{CO\textsubscript{2}} of 28 to 35 mm Hg in these scenarios as a temporizing measure until surgical or other interventions to lower ICP can occur.\textsuperscript{16}

**Head Position**

If not in shock, routinely elevate the patient’s head to decrease ICP, risk of aspiration, and in those who are intubated, rates of ventilator-associated pneumonia. This position allows drainage of cerebral veins. Feldman,\textsuperscript{18} Ng,\textsuperscript{19} and their colleagues demonstrated that elevation of the head to 30 degrees significantly reduces ICP in most patients without impairing CBF, CPP, or cerebral metabolism. Raising the head of a patient with hypotension, however, exacerbates any decrease in the patient’s mean arterial pressure (MAP) and hence lowers CPP. If head elevation is to be used, MAP must be maintained above 90 mm Hg to facilitate a CPP of approximately 60 mm Hg.\textsuperscript{20} It is also important to avoid neck rotation and flexion or any other intervention that could result in compression of the jugular vein. If the jugular veins are compressed, venous outflow from the head can be further compromised, increasing ICP. Therefore the head and neck should be kept in a neutral position.

**Fluid Management**

The goal in fluid management is to maintain euvolement; head-injured patients with hypotension have double the mortality of normotensive patients. Patients with increased ICP may be hypovolemic from profuse vomiting, decreased fluid intake, or hemorrhage. Distributive shock may also be present and occurs as a result of loss of vasomotor sympathetic tone, especially in patients with concomitant cervical spine injury. Only isotonic or hypertonic fluids should be used (normal or hypertonic saline), as hypotonic fluids may lead to worsening brain edema. Vasopressors may also be necessary. A target CPP of 60 mm Hg and MAP of greater than 90 mm Hg are recommended.\textsuperscript{21}

**Hyperosmolar Therapy**

Mannitol effectively and rapidly reduces ICP. Mannitol can be given at doses of 0.25 to 1 g/kg (with the highest dose used in emergency situations) infused every 2 to 4 or more hours.\textsuperscript{22} Current recommendations favor bolus therapy over continuous infusion. Mannitol can cause a precipitous drop in blood pressure (and hence CPP) in patients who are hypovolemic. Thus, mannitol is contraindicated in those with preexisting hypotension (typically defined as systolic blood pressure lower than 90 mm Hg). The effect of mannitol on ICP is transient; repeated doses lose their ability to decrease ICP over time. It may also cause renal damage as serum osmolarity increases. Similar to hyperventilation, mannitol should be viewed as a bridge to definitive neurosurgical intervention and its use reserved for patients with signs of impending herniation.

Mannitol has two properties: it initially acts as a volume expander and then serves as an osmotic agent. On administration of mannitol, intravascular volume expands and blood viscosity decreases, resulting in augmentation of CBF. Once volume expansion has occurred, osmotic movement of fluid from the brain parenchyma to the intravascular compartment begins and results in a decrease in ICP. The osmotic effect usually occurs within 15 minutes. The half-life of mannitol ranges from 90 minutes to 6 hours.\textsuperscript{21} Its effect is most pronounced in patients with CPP lower than 70 mm Hg.\textsuperscript{21}

Hypertonic saline is a good alternative to mannitol in patients who are found to be hypotensive. Hypertonic saline is an ideal resuscitation fluid for patients with concomitant head injury and hemorrhagic shock as it effectively expands intravascular volume while causing osmotic diuresis of the brain. Hypertonic saline is typically used as a 3% solution in 150-mL boluses, as a 7.5% solution in 75-mL boluses, or as a 23.4% solution in 30-mL boluses. There is no consensus on the preferred concentration or volume of administration.\textsuperscript{21–23} Hypertonic saline can be redosed if needed and its onset of action occurs rapidly.

Although the use of mannitol and hypertonic saline have both been endorsed by the Brain Trauma Foundation and a large consortium of pediatric societies, no clear evidence favors the use of one or the other in normotensive or hypertensive patients.\textsuperscript{22,23–25} However, in patients with hypotension, hypertonic saline is clearly preferred. A serum osmolarity of 320 mOsm/L has previously been stated to be the upper limit of safety with both mannitol and hypertonic saline; however, this has been safely exceeded in practice without deleterious effects.\textsuperscript{22,25}

**Seizure Prophylaxis**

Controlling seizure activity is paramount in the early stages of traumatic brain injury to avoid hypoxia and aspiration, as well as elevation of ICP and possible herniation. In the absence of a seizure, it is currently controversial whether prophylactic anticonvulsant medication should be routinely administered after spontaneous intracranial hemorrhage. If a seizure occurs, anticonvulsant therapy is warranted. Following traumatic brain injury, there is low-quality evidence supporting the use of prophylactic phenytoin during the first week following injury to prevent early posttraumatic seizures, with the recognition that this has no effect on the occurrence of late seizures or mortality.\textsuperscript{16} Rapidly acting benzodiazepines such as lorazepam and diazepam are first-line therapy for actively seizing patients, followed by anticonvulsant agents such as pentobarbital or phenytoin. Phenytoin can be given more rapidly (up to 150 mg/min IV by infusion) than phenytoin, which can be administered no faster than 50 mg/min IV. Phenytoin can induce hypotension and lower CPP, even at rates below 50 mg/
min. The use of levetiracetam in cases of severe head injury and stroke has been increasing. Although beneficial effects of prophylactic anticonvulsants on outcomes have not been established, the risks associated with administration of anticonvulsants are small in comparison to the potentially devastating secondary injury that results from uncontrolled seizure activity.

Patients who are paralyzed, either chemically or by their neurologic disease, are difficult to monitor for seizure activity without the use of electroencephalography (EEG). These patients may manifest seizures by subtle rhythmic movements of the extremities, tonic gaze deviation, an elevation in heart rate or blood pressure, or spikes on an ICP monitor (if one is placed by the neurosurgeon). Whenever possible, avoid giving paralytic agents so that seizure activity can be monitored. If seizures are not controlled despite upward titration of benzodiazepines, phenytoin, or other agents, endotracheal intubation and induction of a barbiturate or propofol coma may be necessary. Pentobarbital has been used to treat uncontrolled elevations in ICP when other medical and surgical treatments have failed. It decreases CBF, metabolism, oxygen consumption, and cerebral edema. It also scavenges free radicals. Pentobarbital is given at a loading dose of 10 mg/kg over a 30-minute period followed by infusion at 1 to 3 mg/kg per hour with an EEG monitor available for burst suppression. Intensive monitoring of the patient’s hemodynamic status is necessary because of pentobarbital’s hypotensive effect.

**Steroids**

Steroids have been shown to be beneficial in patients with vasogenic edema, that is, edema associated with brain tumors. Steroids decrease CSF production, stabilize membranes, and restore normal membrane permeability. Dexamethasone is usually administered at 10 to 20 mg IV as a loading dose, followed by 4 to 10 mg every 6 hours. No studies support the use of steroids in head-injured patients; in fact, steroid use in this population has been found to increase mortality. Dexamethasone decreases CBF, metabolism, oxygen consumption, and cerebral edema. It also scavenges free radicals. Pentobarbital is given at a loading dose of 10 mg/kg over a 30-minute period followed by infusion at 1 to 3 mg/kg per hour with an EEG monitor available for burst suppression. Intensive monitoring of the patient’s hemodynamic status is necessary because of pentobarbital’s hypotensive effect.

**Glucose Control**

Head-injured patients tend to be hyperglycemic in response to stress or steroid administration. Optimization of blood glucose levels is currently undertaken in the management of these patients to avoid cellular edema in brain tissue. Hypoglycemic states can exacerbate brain injury and therefore should be scrupulously avoided. However, the long-term value of tight glycemic control in the management of acute brain injury with increased ICP is unclear.

**Hypothermia**

Lower body temperatures have been associated with a decrease in CBF, ICP, and metabolism. The previous method of keeping core temperatures lower than 30°C has been abandoned due to increased incidence of cardiac arrhythmias, severe coagulopathy, and the difficulty of maintaining such profound hypothermia. More recently, mild therapeutic hypothermia (32°C to 34°C) has become popular in the management of post–cardiac arrest victims who do not regain consciousness. Although one study has shown a benefit in survival and decreased ICP with mild hypothermia (32°C to 34°C), in the setting of traumatic brain injury and stroke it is not currently recommended and should only be considered if all other treatment modalities have been ineffective in lowering ICP. Hyperthermia increases ICP and should be aggressively treated with antipyretics and cooling measures.

**Skull Trephination**

Skull trephination, also known as trepanning, trepanation, or burr hole placement, is a technique that has been traced back to the Neolithic age. Though rarely performed by non-surgeons, it can be a lifesaving procedure for severely head-injured patients. In a patient with an acute epidural hematoma and deteriorating findings on neurologic examination (decreasing level of consciousness, anisocoria, or hemiparesis), skull trephination has been shown to improve both short- and long-term neurologic outcomes. Trephination is an appropriate treatment of acute subdural hematoma as well, but this requires incision through the dura mater and is less realistic for non-neurosurgeons (Figs. 59.9 and 59.10). Trephination may be lifesaving if performed before transport in selected patients from remote-practice environments without neurosurgical capability. This heroic procedure is not standard nor is it an expected skill for most emergency clinicians; nonetheless, it may be attempted when the situation is dire.

Trephination in the ED should be performed only in the temporal regions. Trephination in the parietal and occipital regions is associated with a much higher risk for hemorrhage and air embolism because of their proximity to the dural venous sinuses. These approaches should generally be performed only after the results of CT are available for guidance.

In practice environments without access to CT, it is not unreasonable to attempt this procedure before transport of the patient when all other measures to lower ICP have failed and signs of evolving brain herniation are present. The blind approach for temporal trephination is depicted in Fig. 59.11.
**Emergency Skull Trephination**

**Indications**
Acute epidural hematoma and both of the following:
1. Deteriorating findings on neurologic examination (e.g., decreasing level of consciousness, anisocoria, hemiparesis)
2. Delay in obtaining definitive neurosurgical care (e.g., remote-practice environments)

**Contraindications**
Parietal or occipital epidural hematomas (high risk for hemorrhage and air embolism because of dural venous sinuses)
Subdural hematomas (requires an incision through the dura mater and is less realistic for the non-neurosurgeon)

**Complications**
Bleeding
Infection
Brain parenchymal injury

**Equipment**
- A. Self-retaining scalp retractor
- B. 1/2-inch Galt trepine
- C. Raney clips and applier
- D. Suction
- E. Periosteal elevator
- F. Scalpel
- G. Gauze 4- × 4-in pads
- H. Kelly clamp
- I. Scissors
- J. Adson (toothed) forceps

**Review Box 59.1** Emergency skull trephination: indications, contraindications, complications, and equipment.

**Figure 59.10** Epidural and subdural hematomas. **A**, Acute epidural hematoma. The hyperdense (white) appearance of this lesion indicates that it is acute. The inner margins of epiurals (arrows) have a convex appearance, which results in a lens-shaped hematoma. Note that the hematoma does not cross the suture lines (temporal-sphenoid anteriorly, temporal-parietal posteriorly.) **B**, Acute-on-chronic subdural hematoma. The heterogeneous density of this lesion (hypodense anteriorly [double asterisk], hyperdense posteriorly [asterisk]) suggests an acute-on-chronic hemorrhage. The inner margins of subdurs (arrows) are concave, which results in a crescent-shaped hematoma. Subdural hematomas may cross suture lines (note that this hematoma extends from the frontal to the parietal regions). Both these examples demonstrate effacement of the lateral ventricles and midline shift, which are indicators of increased intracranial pressure.
Shave, prepare, and drape the entry site. Use CT to guide the location of the burr hole placement. If CT is not available, choose a site 2 cm anterior and 2 cm superior to the tragus.

Insert a self-retaining scalp retractor to provide exposure. The exposed area should be about 2.5 cm in diameter.

Adjust the centering point of the trephine so that it protrudes 1/8 inch. With the trephine at a 90-degree angle to the skull, apply gentle pressure with a clockwise-counterclockwise rotating motion.

Make a 4-cm vertical incision extending down to the periosteum. Optionally apply Raney clips to the skin edges to control bleeding.

Stop sawing into the skull when you feel a slight give, which indicates that you have penetrated the full thickness of the skull. Remove the trephine.

If the bone fragment does not come out in the trephine, use toothed forceps or a hooked instrument to remove it. Preserve the bone fragment in sterile saline.

Allow the clot and blood to extrude from the burr hole. Apply gentle suction as needed to assist evacuation of the clot. A small suction catheter can be left in place during transport if needed.

**Figure 59.11** Emergency skull trephination. This technique is for patients with epidural hematomas in the temporal region. It should not be used for parietal or occipital hematomas. If a subdural hematoma is present, the dura must be opened. If no blood is encountered on a blind procedure done on the same side as a blown pupil, repeat it on the opposite side. *CT,* Computed tomography.
and involves locating an area 2 cm anterior and 2 cm superior to the tragus on the side of the suspected hematoma. In the majority of cases, the appropriate side for trephination is ipsilateral to pupillary dilation and contralateral to motor paresis. The temporal area (between the ear and the orbit) should be shaved and prepared with chlorhexidine or povidone-iodine via sterile technique. Local anesthetic can be used to infiltrate the area of the planned incision; however, this is not necessary in an unconscious patient and adds time to the procedure.

Once the proper location is identified, either with a blind approach or by using the results of CT, make a 4-cm vertical incision through the skin and temporalis fascia. Divide the temporalis muscle. Once the periosteum is identified, incise it. Expose the skull by elevating the periosteum (with a periosteal elevator if available). Insert a trephine or hand drill into the area underneath the periosteum. While holding the drill perpendicular to the skull, apply continuous gentle downward pressure and rotate the drill in an alternating clockwise-counterclockwise motion. As progress is made with the hand drill, gradually reduce pressure to avoid inadvertent “plunging” into the brain parenchyma. The operator will know when penetration through both the outer and inner tables of the skull has been accomplished once resistance against the drill is no longer felt. After skull penetration has been accomplished, remove the round piece of bone that has been cored out (with the diameter of the drill) and place it in saline.

In many cases, epidural blood and clots under pressure will extrude from the site on full penetration of the skull. However, insertion of a suction catheter into the trephinated space may be necessary for full evacuation of clotted material. If easily identified, the bleeding artery (usually the middle meningeal artery) may be clamped. After successful trephination, the patient’s status should immediately be reassessed for signs of improvement. In a significant minority of patients, false localizing signs may lead the clinician to suspect a hematoma on the wrong side. Thus, if no improvement is noted with trephination on the side of the suspected hematoma, the procedure may be repeated on the opposite side. However, in all cases the delay in definitive neurosurgical care caused by attempts at trephination must be weighed against the possible benefits of the procedure. Moreover, trephination should ideally be performed after consultation with the accepting neurosurgeon.

Complications of the procedure include bleeding, infection, and injury to the brain parenchyma.

**Operative Management**

Operative management is the definitive treatment of increased ICP secondary to a space-occupying lesion. This entails either craniotomy or ventriculostomy with drainage of blood and CSF.

**INTRACRANIAL SHUNTS**

Intracranial shunts are used for the long-term management of increased ICP. An estimated 30,000 intracranial shunts are placed in the United States every year. Intracranial shunts have a high rate of failure and represent a disproportionately high number of hospital readmissions. Approximately 80% of patients will require at least one revision in 20 years and more than half of these occur within the first 2 years after shunt placement. Shunt failure tends to be more common in the pediatric population. A recurrent elevation in ICP because of shunt malfunction can occur days to years after shunt placement.

The basic principle behind an intracranial shunt is to divert CSF into a body cavity from which it can readily be eliminated and drained. The essential elements of the shunt system include a proximal and a distal catheter, a valve, and a reservoir. The valve allows unidirectional flow, incorporates a pumping chamber, and regulates the pressure at which flow will occur across it. The reservoir is usually located between two valves. The proximal valve allows flow from the ventricles to the reservoir, whereas the distal valve allows flow from the reservoir to the distal catheter (Fig. 59.12). Many different types of shunt systems, incorporating a variety of designs, are available (Fig. 59.13). Some have unique characteristics, such as a double dome, whereas in others, valves are absent altogether.

In most cases the reservoir allows for measurement of pressure, testing for patency, fluid sampling, and injection of medication or contrast material. In rare cases, other equipment is incorporated into the shunt system for specific purposes, including an on-off switch, a telemetric pressure sensor, and an anti-siphon device.

Ventriculoperitoneal (VP) shunts are at present the mainstay for treatment of hydrocephalus in infants and children because of the ease of insertion of the catheter and the capacity of the peritoneal cavity to absorb CSF. Other types of shunts include ventriculovenous, ventriculoatrial, ventriculopleural, and lumboperitoneal. These are usually reserved for circumstances in which VP shunting has failed or when the patient has a history of multiple abdominal surgeries or peritoneal infections.

In the VP shunt system, a ventricular catheter is placed in the right frontal horn of the lateral ventricle (nondominant hemisphere) and connected to a subcutaneous valve traversing the temporal aspect (Fig. 59.14). This valve is then connected
to a distal catheter threaded subcutaneously into the neck and finally into the peritoneum (Fig. 59.15). Ventricular catheters can be either straight or angled, with the latter having the option of a reservoir component attachment. Valves come in four different types (ball, diaphragm, miter, slit), each with unique flow characteristics. The distal catheter has either an open or closed end. Identifying the type of shunt in place is often difficult unless the patient or caretaker has the information available. Moreover, the skin overlying the subcutaneous component of the shunt in the temporal regions can scar, thus rendering palpation of the shunt type impossible.

Programmable VP shunts have recently become more popular and several different brands are now available. Unlike a traditional shunt, which has a fixed pressure threshold for

![Shunt components](image1)

*Figure 59.13* Shunt components. (From Rengachary SS, Wilkins RH, editors: *Principles of neurosurgery*. Philadelphia, 1994, Mosby.)

![Cross-section of a Pudenz flushing valve](image2)

*Figure 59.14* Cross-section of a Pudenz flushing valve (American Heyer-Schulte, Santa Barbara, CA) illustrating the diaphragm valve. The proximal inlet tube and silicone base are placed in the bur hole so that only the reservoir (silicon dome) protrudes above the skull. (Courtesy PS Medical, Goleta, CA.)

![Ventriculoperitoneal shunt and alternative occipital placement](image3)

CSF drainage, the opening pressure of the valve in a programmable shunt can be adjusted transcutaneously using a magnetic coding device.

Shunt Failure

Shunt failure can occur because of mechanical obstruction, fracture or migration of the shunt, excessive CSF drainage or infection.44

Mechanical Obstruction

The most common cause of shunt malfunction is catheter obstruction. This usually occurs at the ventricular catheter tip or at the valve.39 The proximal blockage may be due to tissue debris or choroid plexus within the ventricular catheter, whereas the distal obstruction of venous shunts may result from thrombus or venous occlusion (such as in a ventriculovenous shunt). Peritoneal shunts may be associated with infection (peritonitis) or mechanical obstruction (e.g., omental blockage). Some evidence suggests that delayed hypersensitivity to the shunt material is also an occasional cause of obstruction.

Postoperatively, following shunt placement, the ventricles usually begin to diminish in size within a week in patients with high-pressure hydrocephalus, so continued enlargement suggests (but does not confirm) shunt malfunction. With normal-pressure hydrocephalus, however, the ventricles may remain large despite adequate drainage of the CSF.

Fracture and migration of the catheter usually present with subtle signs of elevated ICP and clinically are similar to shunt obstruction.44

Excessive CSF Drainage

Extra-axial fluid collection may occur in response to rapid decompression of enlarged ventricles. This is a relatively rare cause of shunt failure and is estimated to occur in approximately 3.4% of cases. Although most commonly due to a benign collection of CSF, at times a subdural hematoma may form in response to the sudden decompression of the ventricles, which leads to tearing of the bridging veins as the brain pulls away from the dura. Slit ventricle syndrome (SVS) can also occur in response to chronic overdrainage in children and is characterized by small ventricles demonstrated on CT scan with symptoms suggestive of shunt malfunction. In contrast to shunt obstruction, symptoms of SVS are typically influenced by postural changes and improve after resting supine.45

Shunt Infection: Treatment and Prevention

Of all the potential complications associated with shunting procedures, infection is the most notorious and occurs in 2% to 10% of cases.46 Two studies have shown that approximately 8% of neurosurgical patients with implanted shunts acquire infections.47,48 Infection can be due to skin organisms overlying the valve that may ulcerate the skin and colonize the shunt; however, the most common cause of shunt infection is colonization at the time of shunt placement.50 Typical organisms include *Staphylococcus epidermidis* and *Staphylococcus aureus*, as well as gram-negative bacilli, which are thought to be introduced into the system during manipulation of the site.51 Approximately 70% of these infections are seen within 2 months of shunt placement. Other risk factors associated with shunt infection include perioperative infection and dental or urologic instrumentation.

Most shunt infections are caused by otherwise nonvirulent bacteria such as *S. epidermidis*. In the presence of the shunt, these organisms exhibit unusual virulence. This reflects a variety of factors, including the ability to adhere to shunt surfaces, as well as production of mucoid substances that protect the bacteria from host defenses. The Silastic shunt material itself has an adverse effect on the immune system. Specifically, leukocytes cannot adhere to such surfaces as well as bacteria can.

The diagnosis may be obvious in patients with variable systematic signs, wound infection, meningitis, peritonitis, or septicemia. However, the shunt may harbor an indolent infection without symptoms or signs. Culture of CSF obtained from a shunt tap may be negative even when the shunt is ultimately shown to be infected.52,53

A 2014 systematic review recommends removal of the shunt “with a moderate degree of clinical certainty” in the presence of shunt infection; however, controversy still exists over the need for complete or partial shunt removal.54 Nevertheless, many neurosurgeons still recommend replacement of the entire shunt after the infection is eradicated.

If a shunt infection is suspected, empiric systemic antibiotics should be given as soon as possible in the ED. Consultation with a neurosurgeon and infectious disease specialist is recommended before administering antibiotics directly into the CSF, as the available evidence is insufficient to support intrathecal administration of antibiotics in addition to those administered systemically.54

Clinical Presentation

Patients with intracranial shunt malfunctions frequently present to the ED. In many cases the patient’s significant other or caretaker provides the subtle clues that lead to the diagnosis. Depending on the age of the patient, symptoms of increased ICP may include changes in behavior and level of consciousness. Headache, nausea, vomiting, and visual disturbances are also common. In children, lethargy, poor feeding, vomiting, ataxia, decreased or increased activity level, fussiness, fever and diaphoresis, as well as bulging fontanelles can all be indications of shunt malfunction. Bradycardia not associated with hypertension is also suggestive of increased ICP and shunt malfunction. Fever, particularly within the first 6 months after surgery, is strongly associated with shunt infection.43 In children, lethargy and shunt site swelling are the most predictive signs of shunt malfunction.31 Failure of upward gaze is another sign (“sunset eyes” or Parinaud’s syndrome).55 Cranial nerve, visual field, and fundoscopic examination should be routine in the assessment of older, cooperative patients with intracranial shunts.

Symptoms of shunt malfunction may be difficult to interpret, particularly if the symptoms are atypical or nonspecific or if they occur in young children. Likewise, asymptomatic shunt obstruction can occur in children in whom shunt independence has developed. Clinical evaluation of a child may not always be diagnostic of a shunt malfunction.

Shunt Assessment

The characteristics of the valve reservoir on compression or “pumping” the shunt may be useful in diagnosing shunt malfunction in some cases, although the utility of this technique has been challenged by some experts.56,57 Partial shunt valve compression should be done because full depression and release may lead to suction of the choroid plexus. If the CT scan
shows narrow (slit) ventricles, indicative of overdrainage. Further valve compression may similarly cause blockage of the shunt. When compressing the shunt, a normal refill time of 15 to 30 seconds should be observed. If the valve fills more slowly than this but can be compressed easily, the obstruction is proximal to the valve. If the valve is not compressible, the blockage is either at the valve or distal to it. Proximal obstruction is more common than distal obstruction. Other sources of proximal blockage include blood clots or debris related to the surgical procedure. Sources of distal obstruction include malposition, infection, shunt disconnection, and pseudocyst formation. The entire shunt tract and all surgical incisions should be examined for signs of wound infection, disruption of the tubing, or CSF leakage around the tract. Definitive treatment usually requires shunt revision.

Radiographic imaging is the cornerstone of shunt evaluation when dysfunction is suspected in the ED. A non–contrast-enhanced head CT is the initial imaging modality of choice. Radiographic findings indicative of hydrocephalus include enlarged ventricles and evidence of transependymal flow of CSF. It should be noted, however, that the finding of enlarged ventricles on CT reveals little about shunt function unless comparison with previous images or serial scans show progressive ventricular expansion. Head CT has a sensitivity of 83% in detecting shunt obstruction and a negative predictive value of 93%.

Although shunt series radiographs have long been a mainstay in the evaluation of VP shunts, evidence challenges the role of these radiographs as a routine part of the diagnostic workup for shunt malfunction in the setting of a normal CT or radionuclide imaging. Findings of an abnormal shunt series alone changed management only in 0.3% of cases in a recently published (2011) ED retrospective review.41

Patients with intracranial shunts are subjected to repeat radiological evaluation over the course of their lifetime and excessive exposure to ionizing radiation is a very real concern. Therefore sparing additional unnecessary exposure to radiation has been advocated by certain experts.41 Other techniques, such as low-dose CT scan obtained with decreased tube current and a lower number of CT slices (as low as three), have significantly decreased radiation exposure (up to 87%) while still accurately diagnosing shunt malfunction.58 Modified magnetic resonance imaging (MRI) protocols have also decreased the use of ionizing radiation in the evaluation of shunt function. One note of caution: MRI studies may interfere with the settings of programmable VP shunts; consultation with the patient’s neurosurgeon is advised before proceeding with MRI in these patients.59

Shunt series radiographs consist of anteroposterior (AP) and lateral skull, AP chest, and AP abdominal radiographs (Figs. 59.16 and 59.17). They still have an essential role in the setting of an abnormal CT as these films may reveal kinking, breakage, or disconnection of the catheter (Fig. 59.18) and help localize the site of shunt malfunction. Shunt series radiographs have a sensitivity of 20% and a negative predictive value of 22%.

![Figure 59.16 Computed tomography (CT) evaluation of a ventriculoperitoneal shunt. Axial (top row) and coronal (middle row) CT images reveal the intracranial portion of the shunt. Mild ventriculomegaly is noted; however, this reveals little about shunt function without comparison to previous scans. The shunt (arrows) can also be visualized on the CT scout images (bottom row, left and middle). A companion chest radiograph (bottom row, right) demonstrates continuity of the shunt (arrows) through to the peritoneal cavity.](image-url)
accessing the VP shunt and strict sterile technique should be maintained throughout the procedure.

Other invasive techniques include injection of either radio-nuclide or contrast material into the shunt as a marker of flow. If ventricular fluid pressure is low, there may be little evidence of flow, thus giving a false indication of shunt malfunction.

Various noninvasive techniques have been devised for assessing shunt function. Changes in visual evoked potentials associated with elevated ICP have been suggested as a means of determining function. Thermographic and Doppler detection of shunt flow is also possible. MRI techniques can likewise assess CSF flow and shunt patency.

Shunt Tapping

Percutaneous access of the VP shunt is generally performed by a neurosurgeon; however, if neurosurgical consultation is

If the distal catheter is in the peritoneum and a distal obstruction is suspected or if the patient complains of abdominal pain, abdominal ultrasound should be obtained. Ultrasound may reveal a pseudocyst at the distal portion of the catheter or an abnormal fluid collection. Ultrasound has also been used to rapidly identify the cause of catheter malfunction at the bedside, in the case of catheter discontinuity, and it is likely that its use in this role will continue to increase, particularly if radiographic studies are not readily available.

Tapping the reservoir proximal to the distal valve allows percutaneous testing of proximal shunt patency. Although this is useful in most instances, it can be difficult to assess the rate of inflow or runoff adequately with a small-bore needle, particularly if the patient is a child or is unable to cooperate. CSF should be sent for analysis, including cell count, glucose, protein, Gram stain and culture, when infection is suspected. The potential for introducing infection is a concern when accessing the VP shunt and strict sterile technique should be maintained throughout the procedure.

Other invasive techniques include injection of either radio-nuclide or contrast material into the shunt as a marker of flow. If ventricular fluid pressure is low, there may be little evidence of flow, thus giving a false indication of shunt malfunction.

Various noninvasive techniques have been devised for assessing shunt function. Changes in visual evoked potentials associated with elevated ICP have been suggested as a means of determining function. Thermographic and Doppler detection of shunt flow is also possible. MRI techniques can likewise assess CSF flow and shunt patency.
allow it to fully dry. After appropriate draping, infiltrate the skin with 1% plain lidocaine to a level of adequate local anesthesia.

Enter the reservoir percutaneously at a 20- to 30-degree angle (Fig. 59.19). Lack of CSF flow from the reservoir indicates a

not available, the emergency physician may need to perform the procedure as described.

Clip the scalp hair over and around the reservoir, prepare the skin with a surgical scrub brush for 10 minutes, followed by the application of a povidone-iodine solution, and then

Figure 59.18 Before and after series of a shunt malfunction. The left column (A–C) demonstrates a malfunctioning shunt. The right column (D–F) is the same patient after shunt revision. A, The shunt has become disconnected at the valve; no efferent catheter tubing is seen coming from the inferior portion of the valve (arrow). B, The catheter has migrated inferiorly and has coiled in the pelvis (arrow). C, Head computed tomography (CT) scan showing ventriculomegaly (arrow), a direct result of the shunt malfunction. D, After shunt revision, the lateral skull radiograph shows that a catheter is now connected to the valve (arrows, difficult to appreciate on this film). E, The catheter is seen coursing through the abdomen (arrows). F, Repeated head CT shows resolution of the ventriculomegaly.
proximal obstruction unless the ventricles are completely deflated (SVS). Even so, a small amount of CSF should be obtained within the tubing, thus ensuring that entry into the lumen of the reservoir has been accomplished. If the ventricles are deflated and only a small amount of CSF is aspirated, hold the end of the tubing 5 to 10 cm below the level of the reservoir to see whether CSF will fill the tubing and eventually begin to drip at the rate of 2 to 3 drops/min. If CSF is readily aspirated from the reservoir, hold the tubing vertically to obtain an indication of intraventricular pressure. This pressure reading, as well as the ease with which CSF is aspirated, will give some indication of proximal obstruction. To assess runoff or patency of the distal end, apply pressure to the tubing proximal to the reservoir and then deflate the reservoir without any resistance. If any resistance is encountered when deflating the reservoir, suspect a distal obstruction.

It is essential to send any aspirated CSF for laboratory analysis, including a cell count, protein, glucose, and most importantly, Gram stain and culture to evaluate the system for infection.

**Future Directions**

The diagnosis of shunt malfunction remains challenging and delays in diagnosis can cause permanent neurologic sequelae and death. Proposed solutions include developing new techniques to drain CSF that are not reliant on intracranial shunts with their high failure rates and frequent need for revision. Endoscopic third ventriculostomy (EVT) has been considered a valuable alternative to VP shunt placement in selected cases. Although there are no trials directly comparing the two techniques, a 2014 systematic review of the evidence has found that ETV is a viable treatment option for pediatric hydrocephalus, with equivalent outcomes in determined clinical scenarios.61

Another possible solution would rely on improved evaluation of the implanted shunts. An indwelling device that monitors ICP would obviate the need for repeated exposure to ionizing radiation and the risk of infection of repeated shunt taps. One such device, currently still in the experimental phase, shows clinical promises. The baric probe is an implantable, long-term ICP monitor based on a compressible gas design that can be interrogated by handheld ultrasound and provide continuous data on the status of the shunt.62

**ACKNOWLEDGMENTS**

The authors thank Jessica L. Osterman, MD and Megan L. Rischall, MD as well as Frederick K. Korley, MD, Khosrow Tabassi, MD, and Cecile G. Silvestre, MD, for their contributions to this chapter in previous editions, as well as special thanks to Stuart P. Swadron, MD.

REFERENCES ARE AVAILABLE AT [www.expertconsult.com](http://www.expertconsult.com)
REFERENCES


Cerebrospinal fluid (CSF) examination is performed in the emergency department (ED) to obtain information relevant to the diagnosis and treatment of specific disease entities. Many urgent and life-threatening conditions require immediate and accurate knowledge of the nature of the CSF. However, on rare occasions, certain harmful consequences may result from a spinal puncture. Perform a careful neurologic examination before the procedure, and give special thought to the risks and merits of the procedure in each situation.

HISTORICAL PERSPECTIVE

In 1885, Corning punctured the subarachnoid space of a living patient to induce cocaine anesthesia.1 In 1891, Quincke first removed CSF in a diagnostic study and introduced the use of a stylet.2 He studied the cellular contents and measured protein and glucose levels. Quincke was also the first to record CSF pressure with a manometer. Subsequently, increasingly sophisticated bacteriologic, biochemical, cytologic, and serologic techniques were introduced. In 1919, Dandy reported on replacing CSF with air to determine normal brain anatomy and to identify pathologic changes.3 Water-soluble contrast media have since been used to delineate the spinal subarachnoid space and cerebral cisterns. Other uses of spinal dural puncture include drainage of fluids and injection of anesthetic agents, chemotherapeutic agents, and antibiotics.

ANATOMY AND PHYSIOLOGY

In adults, CSF occupies approximately 140 mL of the spinal and cranial cavities, with approximately 30 mL in the spinal canal. This volume is the result of a balance between continuous secretion (primarily by the ventricular choroid plexus) and absorption into the venous system (mainly by way of the arachnoid villi). After formation, the fluid passes out of the ventricles via the midline dorsal foramen of Luschka and the lateral ventral foramina of Magendie. The fluid then flows into the spinal subarachnoid space, the basilar cisterns, and the cerebral subarachnoid space. The rate of production is approximately 0.35 mL/min, and ventricular production of CSF is such that there is a net flow out of the ventricles of 50 to 100 mL/day. The usual volume of CSF (15 to 20 mL) removed at lumbar puncture is commonly regenerated in approximately 1 hour.

CSF may have an embryologic nutritive function; at maturity, CSF most likely acts as a mechanical barrier between the soft brain and the rigid fibro-osseous dura, skull, and vertebral column. It also appears to support the weight of the brain.4 When buoyed by CSF, the functional weight of the brain is...
INDICATIONS FOR SPINAL PUNCTURE

General Indications

The indications for spinal puncture have been reduced with the introduction of noninvasive diagnostic procedures, that is, magnetic resonance imaging (MRI) and computed tomography (CT). A few clinical situations require early or even emergency spinal puncture. The primary indication for an emergency spinal tap is the possibility of central nervous system (CNS) infection (meningitis), with the exception of a suspected brain abscess or a parameningeal process. The need for early detection of meningitis results in the performance of many more lumbar punctures than ultimate diagnoses of infection. No other method can be used to completely exclude meningitis.

The mere presence of a fever does not mandate lumbar puncture. However, CSF should generally be examined for evidence of infection in patients with a fever of unknown origin, especially if consciousness is altered or the immune system is impaired, even in the absence of meningeal irritation. Meningeal signs may not be present in patients who are old, debilitated, or immunosuppressed; are receiving antiinflammatory drugs; or have had partial treatment with antibiotics. In a newborn, even a fever is not a dependable sign because temperatures may be normal or even subnormal. For infants younger than 1 year, a high index of suspicion is required to make the diagnosis of meningitis.

Approximately 25% of infants with meningitis will not have nuchal rigidity, but many appear toxic or moribund. A tense and bulging fontanelle is somewhat more reliable, although this sign may be absent in a dehydrated child. Neonatal meningitis occurs in 25% of sepsis cases. In addition, 15% to 20% of infants with meningitis have negative blood cultures. In a child between the ages of 1 month and 3 years, fever, irritability, and vomiting are the most common signs of meningitis. Typically, handling is painful for the child, and the child cannot be comforted. In addition, an older child may complain of a headache. At all ages, patients generally appear ill and drowsy with a dulled sensorium. Physical signs become more useful in diagnosing meningitis in children older than 3 years and include nuchal rigidity, Kernig’s sign (effort to extend the knee is resisted), and Brudzinski’s sign (passive flexion of one hip causes the other leg to rise, and effort to flex the neck makes the knees come up). The jolt accentuation test is a more sensitive sign of meningeal irritation. This test is considered positive when the patient’s pain is exacerbated by lateral rotation of the head to either side. A petechial rash in a febrile patient should also raise suspicion for Neisseria meningitidis. Previous use of antimicrobial agents may modify the clinical and CSF findings; partially treated children are less likely to be febrile or exhibit an altered mental status. In addition, patients in the early stages of meningitis may lack the classic features associated with advanced disease.

The second indication for emergency spinal puncture is suspected spontaneous subarachnoid hemorrhage (SAH). The diagnosis is usually made by imaging (head CT) or by the direct finding of blood in CSF obtained by spinal puncture. Before a major hemorrhage, 20% to 60% of patients with aneurysmal SAH have had a “sentinel thunderclap” or “warning leak” headache (i.e., an unusual sudden headache caused by a “minor” leak of blood from the aneurysm). The headache may precede a major rupture by hours to months. The goal of early clinical recognition is surgical or other interventional therapy before a recurrent major hemorrhage. After a warning leak, head CT is more likely to be falsely negative, thus giving added importance to the performance of a lumbar puncture.

The usual clinical picture of SAH is an instantaneous excruciating headache. Patients generally recall the exact moment that the headache occurred. The location of the headache is variable and does not necessarily indicate the site of hemorrhage. Nausea, vomiting, and a decreased level of consciousness are common symptoms, with approximately one third of patients becoming unconscious at the onset. Examination usually reveals an acutely ill patient with irritability or overtly altered mental status. Meningeal signs are commonly present at the time of initial examination and usually develop within 2 to 3 days. The meningeal signs may become more severe during the first week after hemorrhage and correspond to the breakdown of blood in CSF. During the first week, many patients are febrile, a reflection of chemical hemic (blood-related) meningitis. Failure to detect blood radiographically in an awake patient may indicate a small hemorrhage or a predominant basal accumulation of blood. If a patient is initially seen several days after the hemorrhage, the blood may have become isodense with brain tissue and may no longer be visible on CT. The proper diagnosis would then require spinal puncture.

Modern CT scanners detect recent SAH quite accurately. However, because acute SAH is not detected by the initial CT scan in all patients, lumbar puncture has been used to rule out the diagnosis with certainty. A 2011 study conducted at academic medical centers found that a non-contrast head CT done with a third-generation scanner within 6 hours after onset of the headache and read by a qualified radiologist was 100% sensitive for the detection of SAH. Another study performed at non-academic hospitals determined the negative predictive value for the detection of SAH of a head CT done within 6 hours after the onset of headache to be 99.9%. If the neurologic picture demonstrates localizing findings, the presence of a large intracranial hematoa should be suspected, and spinal puncture is contraindicated until imaging studies delineate the nature of the lesion.

Other non-emergent reasons for examination of CSF include evaluation for CNS syphilis, unexplained seizures, instillation of chemotherapy and contrast agents, assessment for a suspected demyelinating or inflammatory CNS process, and treatment of headache from SAH. Carcinomatous meningitis and suspected spinal cord compression from metastatic disease may require spinal puncture for myelography and cytologic examination. MRI is a suitable alternative for identifying compressive myelopathy and has largely replaced contrast-enhanced myelography in developed countries.

Idiopathic Intracranial Hypertension (Pseudotumor Cerebri)

Idiopathic intracranial hypertension (IIH) is a rare condition of unclear etiology. Patients with IIH have a marked elevation in intracranial pressure (ICP; usually to 250 to 450 cm H2O) without hydrocephalus or mass lesions in the setting of normal CSF composition. Findings on neuroimaging studies are typically normal, and most cases are idiopathic. Because
of the nonspecific signs and symptoms, many cases initially escape detection by clinicians. IIH has been associated with hypervitaminosis A, tetracycline, estrogen therapy, and a plethora of other conditions such as sarcoidosis, tuberculosis, and carcinomatosis. IIH is most common in obese adolescent girls and young women, but it can also occur in children and men. Patients suffer from chronic headaches, typically worse with maneuvers that increase ICP (e.g., Valsalva maneuver, squatting, bending, coughing). Papilledema is frequently present. Cranial nerve palsies, particularly of the sixth cranial nerve, are caused by increased ICP alone and may be falsely localizing.

IIH may regress spontaneously after a few months. The major concern is visual loss, which may be permanent. The diagnosis can be made only by lumbar puncture performed after neuroimaging. This condition underscores the need to measure opening pressure during lumbar puncture whenever possible. Many cases can be controlled with medication, but occasionally, lumbar puncture is required to lower ICP. One way to lower ICP in the ED is to drain CSF via lumbar puncture by removing 5- to 10-mL aliquots of CSF and checking ICP after each removal until a pressure lower than 200 mm H₂O is achieved. Herniation does not occur despite the elevated pressure in IIH, probably because ICP is uniform in all CNS compartments. As spinal fluid is regenerated rapidly, the procedure may need to be repeated every few days to keep CSF pressure at this level. Analysis of CSF should demonstrate all parameters to be normal. Drug therapy, in the form of acetazolamide, glycerol, diuretics, or corticosteroids, has been advocated and may negate the need for repeated lumbar puncture. Resistant cases may require shunting procedures.

**CONTRAINDICATIONS TO SPINAL PUNCTURE**

Spinal puncture is absolutely contraindicated in patients with infection in the tissues near the puncture site. Spinal puncture is relatively contraindicated in the presence of increased ICP caused by a space-occupying lesion. Caution is particularly advised when lateralizing signs (hemiparesis) or signs of uncal herniation (unilateral third-nerve palsy with an altered level of consciousness) are present. In such cases, a tentorial or cerebellar pressure cone may be precipitated or aggravated by the spinal puncture. Cardiorespiratory collapse, stupor, seizures, and sudden death may occur when pressure is reduced in the spinal canal.

The risk for herniation seems to be particularly pronounced in patients with brain abscesses. Brain abscesses are frequently manifested as expanding intracranial lesions that induce headache, mental disturbances, and focal neurologic signs rather than as an infectious process with signs of meningeal irritation. Fever is often absent. In 75% of cases, a primary source of chronic suppuration is present. Common predisposing factors for brain abscess include craniofacial trauma, cranioencephalic trauma, penetrating injuries that push bone fragments into the brain, large animal bites of infants’ skulls, neurosurgical procedures, cardiovascular disorders treated with right-to-left shunts, bacterial endocarditis, gram-negative sepsis in neonates, dental infections, chronic sinuitis, otitis, mastoiditis, chronic abdominal, pulmonary, or pelvic infection, bacterial meningitis, and immunosuppression. Abscesses may also develop in infarcted brain tissue in septic patients if the blood–brain barrier is compromised. Although the CSF is usually abnormal (elevated pressure, elevated white blood cell [WBC] count, and elevated protein concentration), spinal puncture in patients with a known or suspected abscess is contraindicated. Brain herniation markedly reduces the patient’s likelihood of survival. If the history suggests abscess, CT can rapidly diagnose and localize the lesion (Fig. 60.1). Because the appearance of brain abscesses on CT is similar to that of neoplastic and vascular lesions, false-positive reports of brain abscess are possible.

Spinal epidural hematomas are very rare but may occur in certain subpopulations of patients undergoing lumbar puncture. Those most at risk are individuals with a bleeding diathesis, including those treated with anticoagulants either before or immediately after lumbar puncture and those with abnormal clotting mechanisms, especially thrombocytopenia. The condition occurs, but rarely, even with an atraumatic tap and in the absence of a coagulation defect. Spinal subdural hematomas after lumbar puncture are even rarer. Lumbar puncture can injure the dural or arachnoid vessels, which may result in minor hemorrhage into the CSF. This is generally of little consequence. However, the number of patients with hemophilia and human immunodeficiency virus (HIV) infection who require lumbar puncture has increased since the late 1990s. In a coagulopathic patient, attempt to correct the clotting deficiency if clinically feasible and time permits. The procedure should be performed by experienced clinicians, who are less likely to traumatize the dura. After the procedure, monitor the patient carefully for progressive back pain, lower extremity motor and sensory deficits, and sphincter impairment. Thoroughly investigate any complaints of motor weakness, sensory loss, or incontinence after lumbar puncture. Lumbar puncture may be performed in patients with a coagulation defect if the procedure is expected to provide essential information, such as in the diagnosis of meningitis and the pathogen responsible. Correct warfarin-induced coagulopathy with fresh frozen plasma (FFP) or prothrombin complex concentrate together with...
vitamin K. The vitamin K–FFP protocol might need 12 to 24 hours to totally reverse the effect of warfarin. Correct the coagulopathy before a puncture if the clinical situation permits such delay. Butler has described the off-label use of low-dose prothrombin complex concentrate for the correction of warfarin-induced coagulopathy prior to lumbar puncture.12 Lumbar puncture can be performed safely in patients with hemophilia A or B whose deficient clotting factor is replenished before the procedure.13 Additional replacement of the clotting factor after the procedure is of unknown value.

Performance of lumbar puncture in patients with leukemia and low platelet counts has also been studied. Howard and coworkers reported on 5223 lumbar punctures performed on 958 children with newly diagnosed acute lymphoblastic leukemia.34 The platelet count was 10 × 109/L, acknowledging that the position was having elective diagnostic lumbar puncture with a platelet count less than 50 × 109/L, opting for fluoroscopic guidance is prudent. The 2015 clinical guidelines from the AABB (formerly the American Association of Blood Banks) include a weak recommendation for prophylactic platelet transfusion for lumbar puncture that is not required if the platelet count is higher than 10 × 109/L. The number of patients with platelet counts lower than 10 × 109/L was too small to allow any conclusion about this group. A more recent review (2010) concluded that in the absence of additional risk factors, a platelet count of 40 × 109/L is “safe” for lumbar puncture. Lower counts are probably safe as well, but there was insufficient evidence to make recommendations.33 The 2015 clinical guidelines from the AABB (formerly the American Association of Blood Banks) include a weak recommendation for prophylactic platelet transfusion for patients having elective diagnostic lumbar puncture with a platelet count less than 50 × 109/L, acknowledging that the position was based on very-low-quality evidence.16

Aspirin and nonsteroidal antiinflammatory agents have not been shown to increase the risk for bleeding following lumbar puncture. Subcutaneous heparin administration is not believed to pose a substantial risk for bleeding after lumbar puncture if the total daily dose is less than 10,000 units. The risk for bleeding in patients taking clopidogrel, ticlopidine, or a glycoprotein (GP) IIb/IIIa receptor antagonist is not known but is probably small; however, use of these agents should cause the clinician to proceed with caution. One scenario is the need to diagnose meningitis because of an unusual or difficult-to-treat pathogen. It is necessary to withhold clopidogrel and ticlopidine 1 to 2 weeks before their anticoagulant effect has fully dissipated, making the decision for emergency lumbar puncture in such patients a risk-benefit scenario, with no firm guidelines. Pharmacologic data suggest that cessation of the GP IIb/IIIa receptor antagonist tirofiban may allow lumbar puncture after 8 hours and cessation of abciximab after 24 to 48 hours. Spinal hematoma may develop in 1% to 2% of patients who receive full anticoagulant therapy after undergoing lumbar puncture.

Proceed with caution in a patient with prior lumbar fusion or laminectomy. It is technically difficult to enter the subarachnoid space in the presence of significant postoperative changes. It would be acceptable to perform the puncture in an applicable space above or below the surgical scar; otherwise, opting for fluoroscopic guidance is prudent.

Attempts have been made to link bacterial meningitis to a lumbar puncture procedure in a bacteremic patient. Presumably, the procedure introduces bacteria into the CSF. This phenomenon is rare, difficult to substantiate, and not supported by clinical data. Though of theoretical concern, bacteremia should not be used as a criterion to forego lumbar puncture if meningitis is suspected.

If the history and physical examination suggest a treatable illness such as meningitis or SAH, the clinician may perform a spinal puncture after careful consideration of the entire clinical picture. In all cases, undertake the study after careful thought regarding how the results will contribute to the evaluation and treatment of the patient. It is unlikely that spinal puncture will beneficially alter management in patients with neoplastic disease, intracranial hematoma, abscess, a completed nonembolic infarction, or cranial trauma. Moreover, even when acute life-threatening emergencies such as SAH or meningitis are suspected, delaying CSF examination temporarily while empirical treatment proceeds is an important management option when there are concerns about the safety of the procedure.

**EQUIPMENT**

Assemble the standard equipment for a spinal puncture before starting the procedure. Place it where the operator can easily access it. A standard-point Quincke cutting needle is most often used and supplied with the kit. Some operators prefer to use a Sprotte needle (Havel’s, Inc., Cincinnati, OH) or a Whitacre needle (Becton Dickinson and Company, Rutherford, NJ) to minimize any dural injury associated with passage of the needle (Fig. 60.2). These styletted needles have a side port for withdrawal of fluid and, theoretically, are more likely to separate than cut the dural tissue. Although commercial kits provide most of the items needed for lumbar puncture, the operator should bring additional supplies, including supplemental spinal needles, specimen tubes, gauze, antiseptic solution, additional local anesthetic, needles and syringes, and extra sterile gloves of the appropriate size (see Review Box 60.1).

**PROCEDURE**

Lumbar puncture (Videos 60.1 and 60.2) is commonly carried out with the patient in the lateral recumbent position (Fig. 60.3). A line connecting the posterior superior iliac crests intersects the midline at approximately the L4 spinous process (Fig. 60.4). Spinal needles entering the subarachnoid space at this point are well below the termination of the spinal cord, and the only important neurologic structure is the cauda equina. Generally, the needle pushes isolated nerves to the side during advancement. The adjacent interspace above or below may be used, depending on which area appears to be most accessible to palpation. The space between the lumbar vertebrae is relatively wide. In the thoracic region, the spinous processes overlap and are directed caudally; therefore there is no midline area free of overlying bone. In adults, the spinal cord extends to the lower level of L1 or the body of L2 in 31% of persons, thus eliminating higher levels as sites for puncture. Puncture in adults and older children may be performed from the L2–L3 interspace to the L3–S1 interspace.

Developmentally, the spinal canal and the spinal cord are of equal length in the fetus. Growth of the cord does not keep pace with the longitudinal growth of the spinal canal. At birth, the cord ends at the level of the L3 vertebra. Consequently, in infants the needle should be placed at the L4-L5 or L5-S1 interval.
is critically ill or eliminate it when the patient is mentally incapacitated and no guardian is present. Many patients greatly fear lumbar puncture, and hence some clinicians provide routine preprocedure sedation or analgesia if not clinically contraindicated. Intravenous midazolam and fentanyl are useful adjuncts, but practices vary and there is no consensus on standards with regard to the use or nonuse of preprocedure medications. If the patient is anxious, it is reasonable to give a benzodiazepine agent parenterally (e.g., midazolam, 0.1 to 2.5 mg intravenously for a healthy adult younger than 60 years of age) to facilitate the procedure. Fentanyl at a dose of 0.5 to 1.5 µg/kg (adults) is a reasonable alternative.

The next important step is positioning the patient. Generally, place older children and adults in the lateral decubitus position for the procedure. Give the patient a pillow to keep the head in line with the vertebral axis. Position the shoulders and hips perpendicular to the stretcher or table. Use a firm table or bed when available. Because flexion of the neck does not facilitate the procedure to any great extent and severe flexion may add to the patient’s discomfort, this step may be omitted. Severe flexion of the neck in an infant may cause airway compromise. Arch the lower part of the patient’s back toward the clinician by having the patient’s knees drawn toward the chest.

Some clinicians place the patient in an upright sitting position because the midline is more easily identified. This position can be used in both adults and infants (Fig. 60.5). The higher CSF hydrostatic pressure while sitting may aid flow of CSF in a dehydrated patient. Observe caution regarding orthostatic changes in blood pressure and airway maintenance. Generally, allow a sitting patient to lean onto a bedside stand and use a pillow to rest the head and arms. Have an assistant support the patient during the procedure. Radiographic studies by Fisher and colleagues have demonstrated the advantages of hip flexion when the sitting position is used. To accomplish hip flexion, use a stool to support the patient’s feet, which pulls the knees up toward the chest. This increases lumbar interspinous width, which may increase the success and ease of needle passage.

Iatrogenic infection after lumbar puncture is extremely rare. Use sterile gloves during the procedure; the need for face masks is debatable. Some authors have suggested applying the same guidelines for control of central line infection (caps, gowns, gloves, and masks) to lumbar puncture. Wash the patient’s back with an antiseptic solution applied in a circular motion and increase the circumference of the cleansed area with each motion. Place a sterile towel or drape between the patient’s hip and the bed. Commercial trays have a second sterile drape with a hole that may be centered over the site selected for the procedure.

Infiltrate the skin and deeper subcutaneous tissue generously with local anesthetic. Buffered or warmed 1% lidocaine is preferred. Warn the patient about transient discomfort from the anesthetic. Anesthetizing the deeper subcutaneous tissue significantly reduces procedural discomfort. Merely raising a skin wheal is insufficient anesthesia. Some operators not only anesthetize the interspinous ligament but also apply local anesthetic in a vertically fanning distribution on both sides of the spinous processes near the lamina. Such a field block on each side of the spinous processes anesthetizes the recurrent spinal nerves that innervate the interspinous ligaments and muscles.

While waiting for the anesthetic to take effect, connect the stopcock and manometer and ensure that the valve is working. Commonly, a 3.5-inch, 20-gauge needle is used in adults, and a 2.5-inch, 22-gauge needle is used in children (a 1.5-inch,
Spinal Puncture

1. Position the patient in the bed. Generally, the lateral decubitus position is preferred. Arch the patient's back towards you. Consider mild sedation or analgesia when clinically appropriate.

2. Identify and mark anatomic landmarks. The L4 spinous process is at the level of the posterior-superior iliac crests.

3. Prepare the skin with antiseptic solution. Apply in a circular motion with a gradually increasing circumference.

4. Apply a sterile drape.

5. Create a wheal with anesthetic in the skin overlying the entry site. Then, infiltrate and anesthetize the deeper tissues.

6. Insert the needle in the midline. Hold the needle parallel to the bed, and advance it toward the umbilicus. Remove the stylet periodically to check for CSF.

7. CSF will flow from the needle hub when the subarachnoid space has been penetrated.

8. Attach the manometer and measure the opening pressure.

9. Collect the CSF sample in sequential, numbered vials.

10. Replace the stylet before removing the needle.

Figure 60.3 Spinal puncture. CSF, Cerebrospinal fluid.
A 22-gauge needle is available for infants. Needles of these sizes have enough rigidity to allow the procedure to be accomplished easily but make less of a dural tear than larger needles do. Patients should be told to report any pain and should be informed that they will feel some pressure.

With the patient in the lateral decubitus position, place the needle into the skin in the midline, parallel to the bed. Hold the needle between both thumbs and index fingers (Fig. 60.6). After the subcutaneous tissue has been penetrated, angle the needle toward the umbilicus. The bevel of the needle should be facing straight up toward the ceiling. The supraspinous ligament connects the spinous processes, and the interspinal ligaments join the inferior and superior borders of adjacent spinous processes. The ligamentum flavum is a strong, elastic membrane that may reach a thickness of 1 cm in the lumbar region. The ligamentum flavum covers the interlaminar space between the vertebrae and assists the paraspinous muscles in maintaining an upright posture (Fig. 60.7). The ligaments are stretched in a flexed position and are more easily crossed by the needle. The ligaments offer resistance to the needle, and a “pop” is often felt as they are penetrated. Some clinicians choose to advance the needle in small increments, removing the stylet to check for CSF flow. If no fluid flows, the stylet is replaced and the needle is advanced. Hold the stylet in place during advancement, until the subarachnoid space has been reached. The pop may not be felt with the very sharp needles contained in disposable trays.

If bone is encountered, partially withdraw the needle to subcutaneous tissue. Repalpate the back and ascertain that the needle is in the midline. Directing the tip of the needle toward the navel often enhances navigation of the interspinal space. If bone is encountered again, slightly withdraw and reangle the needle so that the point is placed at a more sharply cephalad angle (Fig. 60.8). This approach should avoid the inferior spinous process.

Normal CSF is a clear fluid and will flow from the needle when the subarachnoid space has been penetrated and the stylet is removed. In normal patients, the dura will be penetrated when the needle is advanced approximately one-half to three-fourths of its length. In obese patients, the entire length of the needle may be required to reach the subdural space (Fig. 60.9).

If feasible, attach the manometer and record the opening pressure (Fig. 60.10). This step is commonly omitted in critically ill patients. Pressure readings from a struggling patient may be inaccurate. Readings are valid only if taken with the patient relaxed and in the lateral decubitus position (not the
A three-way stopcock, supplied in disposable trays, allows both collection of CSF and measurement of opening pressure with a single needle. Positioning of the manometer is often more convenient if an extension tube (provided with most disposable trays) connects the needle hub to the stopcock, which in turn is attached to the manometer. Position the manometer so that the “zero” mark is at the level of the spinal needle. Then ask the patient to relax. Changing position from a straight to a fully extended position increases CSF pressure minimally, by 2 to 8 cm of water. Extending the legs after needle placement does not meaningfully decrease opening CSF pressure. CSF flow is enhanced if the patient straightens or extends the legs following confirmation of proper needle position. The observation of phasic changes in the fluid column with respirations and arterial pulsations confirms needle placement in the subarachnoid space. If the needle is against a nerve root or is only partially within the dura, the pressure may be falsely low, and respiratory excursions will not be reflected in the manometer. Minor rotation of the needle may solve these problems. Hyperventilation will reduce the pressure readings because of hypocapnia and the resultant cerebral vasoconstriction.

After measuring the pressure, turn the stopcock and collect enough fluid to perform all the studies desired. It is generally advised that CSF should be collected by a free flow from the needle, and not by aspiration. The first sample of fluid exits from the manometer if pressure has been measured, and then additional fluid flows from the spinal canal. Even if the pressure is elevated, remove sufficient fluid for performance of all indicated studies because the risk associated with the procedure involves the dural rent, not only the amount of fluid initially removed. Presumably, more fluid will subsequently be lost through the hole in the dura. Replace the stylet into the needle before withdrawing it.

Commercial trays generally supply four specimen tubes. One tube is commonly used for determining protein and glucose levels and for electrophoretic studies, another is used for microbiologic and cytologic studies, and a third is used for serologic tests. Cell counts should be performed in the first
Lumbar Puncture in Infants

Lumbar puncture in infants is usually performed to exclude meningitis or encephalitis. The sitting position may allow the midline to be identified more easily. Use of a needle without a stylet has been suggested for small infants because this device allows the pressure to be estimated as the needle punctures the dura. However, failure to use a stylet may cause the subsequent development of an intraspinal epidermoid tumor. Additional tests may be indicated. Use special stains, such as India ink for suspected Cryptococcus infection in patients with acquired immunodeficiency syndrome (AIDS), acid-fast stain in patients with possible tuberculous meningitis, and viral studies in patients with suspected encephalitis. The fourth tube can be stored under refrigeration in the laboratory for any additional studies that may be required after the initial assessment.

Traumatic taps are common and usually clinically inconsequential. Nevertheless, they can be minimized by proper patient and needle positioning. A traumatic tap most commonly occurs when the subarachnoid space is transfixed at the entrance of the ventral epidural space, where the venous plexus is heavier. A plexus of veins forms a ring around the cord, and these veins may be entered if the needle is advanced too far ventrally or is directed laterally (Fig. 60.11). If blood is encountered and the fluid does not clear, repeat the procedure at a higher interspace with a fresh needle. A traumatic tap, per se, is not a particularly dangerous problem in a patient with normal coagulation, and no specific precautions are needed if blood-tinged fluid is obtained. However, observe for signs of cord or spinal nerve compression from a hematoma developing within the first several hours in patients with a coagulopathy.

Lateral Approach for Lumbar Puncture

The supraspinal ligament might be calcified in older persons, making a midline perforation difficult. A calcified ligament may deflect the needle. In this case, use a slightly lateral approach. Because the lower lamina rises upward from the midline, direct the needle slightly cephalad to miss the lamina and slightly medially to compensate for the lateral approach. The needle passes through the skin, superficial fascia, fat, the dense posterior layer of the thoracolumbar fascia, and the erector spinae muscles. The needle then penetrates the ligamentum flavum (bypassing the supraspinal and interspinal ligaments), the epidural space, and the dura before CSF is obtained (Fig. 60.12). Lateral cervical puncture is an alternative approach that can be performed utilizing an insertion site 1 cm inferior and 1 cm dorsal to the mastoid process.

Lumbar Puncture in Infants

Lumbar puncture in infants is usually performed to exclude meningitis or encephalitis. The sitting position may allow the midline to be identified more easily. Use of a needle without a stylet has been suggested for small infants because this device allows the pressure to be estimated as the needle punctures the dura. However, failure to use a stylet may cause the subsequent development of an intraspinal epidermoid tumor.
Use of a butterfly infusion set needle simplifies the procedure, which is helpful when managing a squirming or hyperactive patient. In general, a stylet is recommended primarily at the time of skin penetration and on needle withdrawal, although many operators use the stylet during any advancement of the needle.

If the child’s neck is very tightly flexed, CSF might not be obtained. However, if the head is held in midflexion, CSF usually flows briskly. If CSF fails to flow, gently suction with a 1.0-mL syringe to exclude a low-pressure syndrome. Because pressure readings are inaccurate in a struggling child, measurement of pressure is not commonly attempted in infants and young children.

Avoid prolonged severe flexion of the neck in an infant because it may produce dangerous airway obstruction. If the infant suddenly stops crying, check the airway immediately. Proper positioning is best accomplished by an assistant, who maintains the spine maximally flexed by partially overlying the child and using the chest and body weight to immobilize the thorax and hips while holding the child behind the shoulders and knees. Infants have poor neck control; therefore the assistant must also ensure that the child maintains an open airway. Pay particular attention to avoiding marked neck and trunk flexion. Incorrect positioning usually results in multiple punctures and a bloody tap.

Newborn and preterm infants may experience significant hypoxia and clinical deterioration during lumbar puncture; a sitting position appears to be preferable. Lumbar puncture in infants with respiratory distress syndrome may pose greater risk than benefit. This is a problem primarily in neonates but may also apply to younger infants with sepsis. Closely monitor all infants with serious cardiopulmonary disease during the procedure. Preoxygenation with or without monitoring oxygen saturation may be used as a precaution.

Although local anesthesia or sedation has not been a routine practice during lumbar puncture in an infant or child, it is being reconsidered. Neonates perceive pain, and local anesthesia neither produces physiologic instability nor makes the procedure more difficult. The use of topically applied eutectic mixture of local anesthetics (EMLA) reduces the pain associated with needle insertion in newborns. Sedation of an anxious child may be considered, but sedatives are relatively contraindicated in an obtunded patient without a protected airway and in the setting of hemodynamic instability.

The Difficult Lumbar Puncture

The traditional approach to lumbar puncture depends on palpation of bony landmarks to determine the correct location...
Section X
Neurologic Procedures

Comlications

Headache After Lumbar Puncture

A number of complications from lumbar puncture have been reported. One of the most common is headache, which occurs after 1% to 70% of spinal taps. In general, the development of post-puncture headache can be neither prognosticated nor prevented. The syndrome most commonly starts within the first 48 hours after the procedure (although a case occurring 12 days after the procedure has been reported) and usually lasts for 1 to 2 days (occasionally as long as 14 days). Cases lasting months have been described. The headache usually

Procedure and Technique

To perform ultrasound-guided lumbar puncture, position the patient in the usual manner. Place the ultrasound probe over the spine in a transverse orientation at the level of the iliac crests such that the shadow caused by the spinous process is centered on the screen. Use a pen to mark the skin on each side of the transducer, exactly at the midpoints (Fig. 60.US3). These two marks can then be connected to mark the midline of the spine. Next, rotate the transducer 90 degrees to obtain a midline longitudinal view and position it until the gap between two spinal processes is in the center of the screen. Mark the skin at the midpoint of each side of the transducer and then connect the marks to form a single line (Fig. 60.US4). The intersection of the two lines is the site for entry of the needle (Fig. 60.US5). Cleanse the skin with antiseptic solution and perform the remainder of the procedure in the usual fashion.

Ultrasound Box 60.1: Lumbar Puncture—cont’d

Figure 60.US3 First use the ultrasound (US) probe in the transverse orientation at the level of the iliac crests, and obtain an image with the shadow of the spinous process centered on the screen (as in Fig. 60.US1). Mark the skin at the exact midpoint of the transducer. This represents the anatomic midline.

Figure 60.US4 Rotate the ultrasound (US) probe 90 degrees and obtain a midline longitudinal view. Position the probe so that the interspace is centered on the screen (as in Fig. 60.US2). Mark the skin at the midpoint of the transducer. This represents the level of needle entry.

Figure 60.US5 The intersection of the anatomic midline and the level of the interspinous space is the site of needle entry.

for insertion of the needle. However, landmarks are difficult to palpate in overweight and obese patients. When lumbar puncture fails, the usual alternative has been fluoroscopic guidance, which necessitates movement of the patient out of the ED, as well as the availability of an appropriately trained radiologist. Brook found CT guidance to be comparable to fluoroscopy in the facilitation of lumbar puncture in obese patients with a previously unsuccessful attempt. Bedside ultrasound is being used increasingly in emergency and critical care medicine, and its scope of practice has expanded to include guidance for a number of procedures, including lumbar puncture (see Ultrasound Box). Ultrasound can be used in adults as well as neonates and infants.
begins within minutes after the patient arises and characteristically ceases as soon as the patient lies down. The pain is mild to incapacitating and is generally cervical and suboccipital in location but might involve the shoulders and the entire cranium. Exceptional cases include nausea, vomiting, vertigo, blurred vision, ear pressure, tinnitus, and stiff neck. The headache may change to a positional backache or neck ache.

The technique of spinal puncture probably has little to do with the development of a postprocedure headache. The syndrome is widely thought to be caused by leakage of fluid through the dural puncture site. This results in a reduction in CSF volume below the cisterna magna and downward movement of brain tissue, along with displacement and stretching of pain-sensitive structures such as the meninges and vessels, and causes a traction headache. The recumbent position brings relief because the weight of the brain is shifted cephalad. Another proposed mechanism is cerebral vasodilation. Another hypothesis suggests that the headache is caused by an altered distribution of craniospinal elasticity and acute intracranial venous dilation. Some authors have commented on the incidence and severity of post–dural puncture headache as being related to the orientation of the spinal needle bevel and its type and size. It is clear that the incidence of post–dural puncture headache is lower when the bevel of the needle is oriented parallel to the longitudinal axis of the spine. The traditional explanation for this was that the parallel bevel would separate rather than cut the longitudinal dural fibers and thus produce a smaller dural hole and less CSF leakage. However, the dural fibers are oriented randomly, not longitudinally. Even with random fiber orientation, back flexion is more likely to close a dural hole in the longitudinal plane than a hole in the horizontal plane.

The two basic types of spinal needles are cutting (Quincke) and noncutting or pencil point (Sprotte and Whitacre) (see Fig. 60.2). Noncutting needles cause a lower incidence of headache after dural puncture, perhaps because the tip of the needle tends to separate rather than cut the dural fibers. It has traditionally been thought that the dural hole resulting from puncture with a noncutting needle is smaller than that created by puncture with a cutting needle; however, this may not be true. The important difference may be that the cutting needle causes a clean-cut opening in the dura whereas the noncutting needle produces a jagged opening with rough edges. The jagged opening may produce a more intense inflammatory response that results in edema and more complete closure of the hole. The use of atraumatic spinal needles is not routine. As more spinal kit manufacturers include them in kits, their use might increase. A single-institution study suggested that routine use of noncutting needles yields a potential cost saving. However, in thick-skinned individuals, passage of a thin, noncutting needle may be technically difficult. Making the initial pass with a thicker cutting needle to the level of the interspinous ligament, followed by removal and advancement of a noncutting needle in the same soft tissue tract, can be helpful.

Use of a smaller-diameter needle is one intervention that will probably cause a lower incidence of post–puncture headache because it creates a smaller dural hole. If diameter were the only consideration, as small a needle as possible would be used. However, a needle must provide adequate CSF flow rates to allow timely CSF collection and pressure measurement. With a very small needle, a syringe may be needed to withdraw fluid, and pressure cannot be recorded easily. In addition, technically, a small needle such as a 26-gauge is difficult to place and manipulate into a position in which it does not become intermittently obstructed by nerve roots. A 2014 randomized crossover trial compared the use of 22-gauge and 25-gauge needles in children and found no difference in the incidence of post–puncture headache, but the 25-gauge needles were found to be flimsy and difficult to use, with fluid collection taking much longer. When these characteristics are considered, a 20- to 22-gauge atraumatic needle seems to be the best overall choice for diagnostic lumbar puncture.

Despite common beliefs about techniques and post-procedure interventions, lumbar puncture headaches may not be completely preventable. Studies of the influence of directives such as strict bed rest on post-puncture headache have yielded contradictory results, with no consensus on any specific preventive intervention forthcoming concerning the worsening of, improvement in, or effect on the incidence. Brocker reported a reduction in the incidence of headache from 36.5% to 0.5% when patients lie prone instead of supine for 3 hours after puncture with an 18-gauge needle. He postulated that the prone position caused hyperextension of the spine and disrupted alignment of the holes in the dura and the arachnoid, making leakage less likely. Theonnissen and coworkers concluded that there was no evidence that longer bed rest after lumbar puncture was better than immediate mobilization or short bed rest in reducing the incidence of headache. There appears to be no reason to enforce bed rest following lumbar puncture.

Other factors that might influence the incidence of a post–spinal puncture headache were reviewed by Fishman and by Lin and Giederman. The incidence is higher in young patients than in older patients and is also increased in females and individuals with a history of headache. Many medications have been advocated for the treatment of headache following lumbar puncture: barbiturates, codeine, neostigmine, ergots, diphenhydramine (Benadryl [Johnson & Johnson, New Brunswick, NJ]), dimenhydrinate (Dramamine [APP Pharmaceuticals, Schaumburg, IL]), caffeine, amphetamine sulfate (Benzedrine [GlaxoSmithKline plc, London, United Kingdom]), ephedrine, intravenous fluids (normal saline, lactated Ringer's solution), magnesium sulfate, and vitamins. Niraj and colleagues tested the use of greater occipital nerve block in patients with post-procedural headache, a less invasive alternative to epidural blood patch, and achieved modest success in a small trial.

Traditionally, caffeine has often been used as the initial intervention for post–puncture headache, and it appears to be of benefit. Although caffeine may decrease post lumbar puncture headache, there is a high recurrence rate. Caffeine can be administered in the ED, bringing pain relief within a few hours. It is thought to result from caffeine's vasoconstrictive effect on the cerebral vasculature. One suggested regimen is 500 mg of caffeine (sodium benzoate) diluted in 1000 mL of normal saline infused over a 1- to 2-hour period. A second dose can be given if the headache is not relieved. If successful, consumption of caffeinated beverages may be continued as an outpatient. A cup of coffee contains 50 to 100 mg of caffeine.

Most post–puncture headaches can be managed by bed rest with the head in the horizontal position. Avoid dehydration because it lowers CSF pressure and might aggravate the headache. Although dehydration should be avoided, the role of fluid supplementation in the prevention of post–dural puncture headache remains uncertain.
Simple analgesics are commonly prescribed, but they have no apparent advantage over bed rest and fluid intake. A patient with a prolonged headache after spinal puncture should be reassessed to rule out structural causes. Because a spinal headache has classic signs and symptoms, if the headache is not postural, consider other causes.

In summary, because most post–lumbar puncture headaches are mild and self-limited, conservative therapy for the first 24 to 48 hours is recommended. Bed rest and oral analgesics, including opioids, are usually effective, as well as, perhaps, oral caffeine drinks, but for those refractory to conservative measures, an epidural blood patch is recommended. For patients with a prolonged low-pressure headache, placement of an epidural blood patch by experienced operators is highly successful and often provides dramatic relief. Consider a blood patch for all patients with seriously symptomatic spinal headaches. Perform an epidural tap at the level of the previous lumbar puncture. Use the loss-of-resistance technique with sterile saline to locate the epidural space. Draw 10 to 20 mL of autologous blood into a syringe aseptically and slowly inject it (1 to 2 mL every 10 seconds) into the epidural space at the site of the durapuncture. Slow or discontinue the injection if back pain or paresthesia develops. Keep the patient supine for 1 hour while hydration is administered intravenously. Relief usually occurs within 20 to 30 minutes after the procedure. Epidural patches are less likely to be effective if symptoms have been present for more than 2 weeks. Pain is relieved when the blood patch forms a gelatinous tamponade that stops the CSF leak and immediately elevates CSF pressure. Patch failures (15% to 20%) are believed to be caused by improper needle placement, injection of an inadequate quantity of blood (after which a second patch is usually successful), or an incorrect diagnosis.

Complications reported after placement of an epidural patch include back stiffness, paresthesia, radicular pain, subdural hematoma, adhesive arachnoiditis, and bacterial meningitis. The procedure should be used in patients with refractory headaches that do not respond to conservative therapy, and it should be performed by clinicians trained in its use.

**Infection**

Spinal puncture is contraindicated in the presence of local infection at the puncture site (cellulitis, suspected epidural abscess, or furunculosis) because of the danger of inducing meningitis. A large concentration of bacteria in the bloodstream at the time of CSF examination is associated with meningitis. The meningitis could be coincidental (“spontaneous”) or could result from leakage of blood containing bacteria into the subarachnoid space after lumbar puncture (“induced”). It is likely that many cases of puncture-induced meningitis emerge after a cautious clinician performs a lumbar puncture early in the course of meningitis, before the infection has had time to be reflected in CSF. A 2006 review concluded that the majority of cases of post-puncture meningitis are probably caused by contamination of the site with aerosolized bacteria from medical personnel, contamination from skin flora, or least commonly, direct or hematogenous spread from an endogenous infectious site. Suspected bacteremia is not a contraindication to lumbar puncture. Delay in diagnosis because of concern regarding the risks associated with lumbar puncture is probably more serious than the risk of causing meningitis with the procedure.

**Herniation Syndromes After Lumbar Puncture**

Lumbar puncture is of value in confirming a diagnosis of meningitis, encephalitis, or SAH. Generally, when the patient has symptoms consistent with bacterial meningitis but not increased ICP, it is safe to perform lumbar puncture before a head CT scan. Lumbar puncture may also be the best initial procedure to diagnose SAH, thus reducing the need for routine CT scanning in certain low-risk patients with acute sudden headache. However, in patients with severely altered mentation, focal neurologic findings, papilledema, or a suspected intracranial mass lesion, one should perform a CT scan before lumbar puncture to rule out increased ICP or mass effect. Others have considered a recent seizure an indication for CT scan prior to LP. When meningitis remains in the differential diagnosis, antibiotics are best administered after blood is obtained for culture and before the CT scan. In patients with suspected elevated ICP based on clinical features, head elevation, hyperventilation, and intravenous mannitol have been recommended.

Particularly in patients with supratentorial masses, there may be a large pressure gradient between the cranial and lumbar compartments. When brain volume is increased because of a mass or edema, rostrocaudal displacement may occur after lumbar puncture if the skull is intact.

The question of whether lumbar puncture precipitates brain herniation in cases in which herniation would not have occurred spontaneously cannot be answered with certainty. Controversy still exists regarding the risk for brain herniation from lumbar puncture in patients with acute bacterial meningitis, and data continue to be sparse. Herniation may occur in the absence of lumbar puncture in the setting of acute bacterial meningitis and has been temporally related to the procedure. Brain herniation is usually fatal, and occurs in approximately 5% of patients with acute bacterial meningitis and accounts for approximately 30% of the deaths associated with the disease. Although CT imaging may reveal contraindications to lumbar puncture, normal findings on CT do not eliminate the risk for herniation and do not necessarily mean that a lumbar puncture is safe.

Lowering pressure in the lumbar spinal canal by removing CSF can increase the gradient between the cranial and lumbar compartments and thereby theoretically promote both transventricular and foramen magnum (cerebellar) herniation. The frequency with which lumbar puncture causes or accelerates transtentorial herniation is unknown because herniation might have developed spontaneously in a seriously ill patient without the procedure. With the current use of small-caliber spinal needles, herniation appears to be extremely rare; nonetheless, it is not fully predictable by CT or the opening CSF pressure readings.

A careful neurologic examination should precede all spinal punctures. When the patient has a history of headache and fever with progressive deterioration in mental status and localizing neurologic signs, spinal puncture should not be performed as the initial diagnostic procedure. Gopal and colleagues identified three statistically significant predictors of new intracranial masses: papilledema, focal abnormalities on neurologic examination, and altered mental status. Greig and Goroszeniuk made similar recommendations. Hasbun and associates demonstrated that in adults with suspected meningitis, the presence of any of 13 clinical features is predictive of abnormal findings on CT (Box 60.1). Theoretically,
the presence of abnormal findings on CT portends potential herniation, with or without lumbar puncture. The absence of abnormal findings suggests the patient is a good candidate for immediate lumbar puncture because the risk for brain herniation as a result of the procedure is low (see Box 60.1). Joffe argues that clinical signs of impending herniation are the most appropriate criteria on which to base decisions regarding the timing of lumbar puncture in patients with acute bacterial meningitis. Such signs include a significantly decreased level of consciousness (Glasgow Coma Scale score \( \leq 11 \)), brain stem findings (pupillary changes, posturing, irregular respirations), and a very recent seizure. It is recommended that these patients undergo neuroimaging (CT or MRI) and receive antibiotics empirically while awaiting the results of imaging, which will help in assessment of the safety of a subsequent lumbar puncture. Measures to lower ICP before spinal puncture may also be considered. Appropriate cultures of blood and more easily accessible body fluids should be obtained before the administration of antibiotics. Blood cultures are positive in 80% of infants with meningitis.

Critically ill patients with acute bacterial meningitis often deteriorate rapidly and experience fatal brain herniation, both shortly after lumbar puncture and in the absence of the procedure. A direct cause-and-effect relationship between lumbar puncture and brain herniation in the setting of suspected meningitis is obscure and probably cannot be defined prospectively in the ED when clinical decisions must be made. A 2007 review identified 22 case reports of rapid deterioration and herniation after lumbar puncture in adults and children with acute bacterial meningitis. The general consensus is that less ill patients (clearly a subjective clinical judgment) with a clinical scenario that includes meningitis as a possibility can be evaluated safely with lumbar puncture. In critically ill patients, especially those with localizing neurologic signs, severely depressed level of consciousness, or papilledema, diagnostic lumbar puncture may be delayed until the risk for herniation is lower, and aggressive and empirical treatment with meningitis doses of antibiotics should proceed.

Head CT can identify hemorrhagic lesions and most neoplasms. Its results should contribute to the decision regarding the need for and the risk involved with spinal puncture. Head CT can identify patients with unequal pressure between intracranial compartments, who are at greater risk for cerebral herniation. Findings that suggest unequal pressure include (1) lateral shift of midline structures, (2) loss of the suprachiasmatic and perimesencephalic cisterns, (3) shift or obliteration of the fourth ventricle, and (4) failure to visualize the superior cerebellar and quadrigeminal plate cisterns with sparing of the ambient cisterns. The presence of a mass in the posterior fossa is a strong contraindication to lumbar puncture. Unfortunately, because of bone and motion artifact, the posterior fossa may be a difficult area to visualize on CT.

Although there is no clear standard with respect to the use of CT before lumbar puncture, a reasonable and logical approach is to avoid initial lumbar puncture and first perform head CT when a mass lesion is suspected or if the patient has signs and symptoms of increased ICP (see Box 60.1). This approach correlates with the clinical policy promulgated by the American College of Emergency Physicians in 2002.

### Epidermoid Tumor

An epidermoid tumor or cyst is a mass of desquamated cells containing keratin within a capsule of well-differentiated stratified squamous epithelium. Congenital lesions arise from epithelial tissue that becomes sequestered at the time of closure of the neural groove between the third and the fifth weeks of embryonic life, but such lesions are rare. Acquired intraspinal epidermoid tumors result from the implantation of epidermoid tissue into the spinal canal at the time of lumbar puncture performed with needles without stylets or with ill-fitting stylets. The clinical syndrome consists of pain in the back and lower extremities developing years after spinal puncture. Failure to use a stylet on needle withdrawal might also result in aspiration of a nerve root into the epidural space.

### Backache and Radicular Symptoms

Minor backache from the trauma of the spinal needle occurs in 90% of patients. Frank disk herniation has been reported from passage of the needle beyond the subarachnoid space into the anulus fibrosus. Transient sensory symptoms from irritation of the cauda equina are also common.

Other reported complications include transient unilateral or bilateral sixth-nerve palsies caused by stretching or displacement of the abducens nerve as it crosses the petrous ridge of the temporal bone, SAH, subdural and epidural hematoma, epidural CSF collection, cauda equina syndrome, anaphylactoid reactions to local anesthetics, settling of cord tumors, and retroperitoneal abscess produced by laceration of the dura in patients with meningitis. Most of these complications are rare and seldom encountered; however, they should be considered if a patient returns after a lumbar puncture with worsening back pain or abnormal findings on neurologic examination.

The complications associated with lateral cervical and cisternal puncture are similar to those encountered with lumbar

---

**BOX 60.1 Clinical Characteristics Associated With Abnormal Findings on Head CT in Adults With Suspected Meningitis**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age 60 years or older</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Immunocompromised state</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>History of central nervous system disease</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Seizure within 1 week before initial evaluation</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Abnormal level of consciousness</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Inability to answer two questions correctly</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Inability to follow two commands correctly</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Gaze palsy</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Abnormal visual fields</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Facial palsy</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Arm drift</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Leg drift</td>
<td>Increased risk for herniation.</td>
</tr>
<tr>
<td>Abnormal language</td>
<td>Increased risk for herniation.</td>
</tr>
</tbody>
</table>

---

*Includes patients with human immunodeficiency virus infection or acquired immunodeficiency syndrome, those receiving immunosuppressive therapy, and those who have undergone transplantation.

*Mass, stroke, or focal infection.

*Aphasia, dyssartria, or extinction.

CT, Computed tomography.

puncture. Perforation of a large vessel with resultant cisterna magna hematoma or obstruction of vertebral artery flow has been described. Puncture of the medulla oblongata may cause vomiting or apnea, and puncture of the cord may be associated with pain. Long-lasting side effects of cord puncture seem to be rare. A traumatic tap and post-puncture headache may occur.

**Spinal Epidural Hemorrhage**

Rarely, serious bleeding leading to spinal hematoma after lumbar puncture can produce spinal cord compromise and permanent significant neurologic deficits, such as cauda equina syndrome. This may occur more frequently in coagulopathic patients but can also arise in those with a normal coagulation profile and a seemingly atraumatic tap. Aspirin and nonsteroidal medications are not associated with an increased bleeding risk after lumbar puncture. The bleeding is concealed and might be suspected only by persistent severe backache or neurologic findings. Weakness, numbness, and incontinence after spinal puncture must be investigated, usually with MRI, for a possible spinal hematoma. Surgical intervention, including laminectomy and evacuation of blood, may be required and must occur in a timely manner to avoid permanent loss of neurologic function. Those with mild symptoms and progressive recovery may be managed conservatively with close monitoring.

**INTERPRETATION**

**Pressure**

CSF pressure is clinically important. Measure it accurately whenever feasible. Unfortunately, in some cases it will be logistically impossible to obtain. If the lumbar puncture is being performed with the patient in a seated position, place the patient in the lateral decubitus position before a measurement is obtained. This may be done initially, before fluid is collected, or after fluid collection, in which case a closing pressure is obtained. By repositioning the patient and measuring the pressure after fluid is collected, the likelihood of the needle being displaced as a result of the repositioning is minimized. Accurate measurement depends on cooperation of the patient. Measurements from struggling or agitated patients will probably be inaccurate; in such cases, sedation may allow more accurate readings to be obtained.

Normal CSF pressure is between 7 and 20 cm H2O. Obese patients may have a CSF pressure of up to 25 cm H2O. Elevated pressure is abnormal. Opening pressure is taken promptly, thereby avoiding falsely low values caused by leakage through and around the needle. Herniating cerebellar tonsils may occlude the foramen magnum and prevent increased ICP from being reflected in the lumbar pressure reading. Increased ICP can result from expansion of the brain (edema, hemorrhage, or neoplasm), overproduction of CSF (choroid plexus papilloma), a defect in absorption, or obstruction of flow of CSF through the ventricles. Cerebral edema may be associated with meningitis, CO2 retention, SAH, anoxia, congestive heart failure, or superior vena cava obstruction. Pressure may be falsely elevated in a tense patient when the head is elevated above the plane of the needle and, possibly, in markedly obese patients and those experiencing muscle contraction. Pressure is not usually measured in neonates because a struggling or crying child will have a falsely elevated pressure. Avery and colleagues concluded that for most children (1 to 18 years of age), an opening pressure above 28 cm H2O should be considered elevated.

Low pressure suggests obstruction of the needle by the meninges. Low pressure can also be seen with a spinal block. Rarely, a primary low-pressure syndrome occurs in a setting of trauma, after neurosurgical procedures, secondary to subdural hematoma in elderly patients, with barbiturate intoxication, and in cases of CSF leakage through holes in the arachnoid.

The Queckenstedt test is useful for demonstrating obstruction in the spinal subarachnoid space, but it is seldom performed today because myelographic techniques have been refined and the availability of MRI has reduced the number of myelograms and associated lumbar puncture studies. However, because situations might arise when this simple and reliable test is important diagnostically, the technique is described here.

With the patient in the lateral recumbent position, compression of the jugular vein causes decreased venous return to the heart. This distends the cerebral veins and causes a rise in ICP, which is transmitted throughout the system and measured in the manometer. After 10 seconds of bilateral compression, CSF pressure usually rises to 15 cm H2O over the initial reading and returns to baseline 10 to 20 seconds after release. If there is no change in lumbar pressure or if the rise and fall are delayed, one may conclude that the spinal subarachnoid space does not communicate with the cranial subarachnoid space. In this situation, to facilitate subsequent myelography, consider injecting Pantopaque before removing the needle. This is necessary because the lumbar dural sac may collapse, making it impossible to reenter the canal. If cervical cord disease is suspected, repeat the test with the neck in the neutral position, hyperextended, and flexed. When lateral sinus obstruction is suspected, use unilateral jugular venous compression (Tobey-Ayer test).

**Appearance**

If the CSF is not crystal clear and colorless, a pathologic condition of the CNS should be suspected. The examiner should compare the fluid with water and view down the long axis of the tube or by holding both tubes against a white background. A glass tube is preferred because plastic tubes are frequently not clear. CSF will appear grossly bloody if more than 6000 red blood cells (RBCs)/µL are present. Note that the fluid may appear clear with as many as 400 RBCs/µL and 200 WBCs/µL. Xanathochromia, a yellow-orange discoloration of the supernate of centrifuged CSF, is generally considered to be the result of SAH of at least a few hours’ duration and has been used to differentiate prior bleeding from a traumatic tap. A traumatic tap does not usually exhibit xanthochromia (see the later section on The Traumatic Tap). Xanthochromia is produced by red cell lysis and is caused by one or more of the following pigments: oxyhemoglobin, bilirubin, or methemoglobin.

Traditionally, CSF samples have been assessed for xanthochromia by visual inspection by a laboratory technician after centrifugation of a sample of CSF. Most hospitals still use this method. Recently, spectrophotometry, designed to demonstrate both oxyhemoglobin and bilirubin, has been advocated as a more precise way of determining xanthochromia. Oxyhemoglobin alone without bilirubin in a CSF sample is thought
to be artifactual (traumatic). Visually, bilirubin and oxyhemoglobin cannot be differentiated. Therefore detection of bilirubin by spectrophotometry should define xanthochromia and prompt additional investigation for SAH. Relying on spectrophotometry to identify xanthochromia without pigment differentiation will cause a high false-positive interpretation. The absence of both oxyhemoglobin and bilirubin by spectrophotometry does not support the diagnosis of SAH. Oxyhemoglobin causes red coloration; bilirubin, yellow; and methemoglobin, brown. Oxyhemoglobin is seen within 2 hours after subarachnoid bleeding and red cell lysis, but it may be detected immediately if the bleeding is profuse. Formation of oxyhemoglobin peaks 24 to 48 hours after hemorrhage, and the discoloration disappears in 3 to 30 days.4

Blood must be present in CSF (in vivo) for a number of hours for bilirubin to appear; it will not appear spontaneously once CSF is in the collection tube. The appearance of bilirubin in CSF involves the conversion of oxyhemoglobin by the enzyme heme oxygenase. The enzyme is found in the choroid plexus, the arachnoid, and the meninges. Enzyme activity appears approximately 12 hours after the hemorrhage.4 Bilirubin may persist in CSF for 2 to 4 weeks. Moreover, bilirubin in CSF caused by hepatic or hemolytic disease does not appear until a serum level of 10 to 15 mg of total bilirubin per 100 mL is reached, unless underlying disease associated with high CSF protein levels is present. Xanthochromia may also be seen with CSF protein values above 150 mg/dL. CSF may clot in patients with a complete spinal block and very high CSF protein.

Graves and Sidman noted that an RBC concentration of 5000/µL, created by adding RBCs to acellular CSF, will produce xanthochromia evident on spectrophotometry by 2 hours.129 The addition of RBCs to achieve RBC concentrations of 20,000/µL and 30,000/µL will produce xanthochromia by 1 hour or immediately, respectively. Therefore xanthochromia may occur with a traumatic tap and does not always indicate SAH.

Methemoglobin is a reduction product of oxyhemoglobin that is characteristically found in encapsulated subdural hematomas and in old intracerebral hematomas.

**Cells**

In adults, WBC counts higher than 5 cells/µL indicate the presence of a pathologic condition. A large study by Kestenaum and colleagues in 2010 determined 95th percentile WBC reference values in normal infants: 19 cells/µL for infants 28 days of age or younger and 9 cells/µL for those between the ages of 29 and 56 days.130 A median CSF WBC count of 271 WBCs/µL has been reported in infants with group B Streptococcus in the era of intrapartum antibiotic prophylaxis. The CSF WBC count in infants is higher with gram-negative meningitis than with gram-positive CNS infections. In general, polymorphonuclear leukocytes are never seen in normal adults. However, with use of the cytocentrifuge, an occasional specimen from an otherwise normal individual may show one to two neutrophils.131 More than three neutrophils is always abnormal in an adult. Moreover, as many as 30% of patients exhibit CSF pleocytosis after a generalized or focal seizure. Nonetheless, such a finding should prompt culture of CSF because the presence of neutrophilic pleocytosis is commonly associated with bacterial meningitis or the early stages of viral or tuberculous meningitis. Small lymphocytes may be seen in normal individuals. Small and large immunocompetent cells are found with a variety of bacterial, fungal, viral, granulomatous, and spirochetal diseases.

Eosinophils always indicate an abnormal condition, most commonly a parasitic infestation of the CNS. They may also be seen after myelography and pneumoencephalography and, to a minor degree, with other inflammatory diseases, including tuberculous meningitis and neurosyphilis. Normal CSF RBC counts are lower than 10 cells/µL. Herpes simplex virus (HSV) encephalitis may elevate the RBC count. Finally, myeloid and RBC precursors may contaminate CSF with bone marrow cells from an adjacent vertebral body.132

**Glucose**

Glucose enters CSF by way of the choroid plexus, as well as by transcapillary movement into the extracellular space of the brain and the cord via carrier-mediated transport. It then equilibrates freely within the CSF subarachnoid space. Once in the CSF, glucose undergoes glycolysis and there is an invariable rise in CSF lactate levels. Glucose levels remain subnormal for 1 to 2 weeks after effective treatment of bacterial meningitis.

The normal range of CSF glucose is 50 to 80 mg/dL, which is 60% to 70% of the glucose concentration in blood. Ventricular fluid glucose levels are 6 to 8 mg/dL higher than in lumbar fluid. A ratio of CSF glucose to blood glucose of less than 0.5 or a CSF glucose level below 40 mg/dL is invariably abnormal. The ratio is higher in infants, in whom a ratio of less than 0.6 is considered abnormal. Hyperglycemia may mask a depressed CSF glucose level; when present, the ratio of CSF glucose to blood glucose should be measured routinely. With extreme hyperglycemia, a ratio of less than 0.3 is abnormal.133 Between 90 and 120 minutes is required before CSF glucose reaches a steady state with changes in blood glucose (e.g., after an intravenous injection of glucose). When CSF glucose is of diagnostic importance, obtain CSF and blood samples, ideally after a 4-hour fast.

Low CSF glucose levels may be associated with several diseases of the nervous system (Box 60.2). Only low concentrations of glucose are of diagnostic value; elevated CSF glucose levels generally have little significance and usually reflect hyperglycemia. A rapid estimate of the CSF glucose level can be obtained by using bedside reagent strip testing with a

---

**BOX 60.2 Low CSF Glucose Syndromes**

- Bacterial meningitis
- Tuberculous meningitis
- Fungal meningitis
- Sarcoidosis
- Meningeal carcinomatosis
- Amebic meningitis
- Cysticercosis
- Trichinosis
- Syphilis
- Chemical meningitis
- Subarachnoid hemorrhage
- Mumps meningitis
- Herpes simplex encephalitis
- Hypoglycemia
commercial autoanalyzer. Formal laboratory testing is recommended for confirmation of the bedside levels.

Protein

The normal range of lumbar CSF protein is 15 to 45 mg/dL. Infants normally have a lower level than adults, and protein levels may drop after lumbar puncture. The concentration is lower in the ventricles (5 to 15 mg/dL) and the basilar cisterns (10 to 25 mg/dL) as a result of a gradient in the permeability of capillary endothelial cells to proteins in blood. Levels of CSF protein in premature infants and full-term neonates are higher than in adults, with a mean of 90 mg/dL; protein levels decline by the age of 8 weeks because of maturation of the blood–brain barrier.

Most of the proteins in CSF come from blood, which normally has a protein concentration of up to 8000 mg/dL. Protein entry is determined by its molecular size and the relative impermeability of the blood–CSF barrier. A full range of serum proteins is found in CSF at a several hundred-fold dilution.

An increase in the total CSF protein level is a nonspecific abnormality associated with many disease states. Levels higher than 500 mg/dL are uncommon and seen mainly with meningitis, SAH, and spinal tumors. The high levels that occur with cord tumors result from an increase in local capillary permeability. With high levels (generally 1000 mg/dL), CSF may clot (Froin's syndrome).

Hemorrhage into CSF or the introduction of blood by a traumatic tap increases CSF protein levels. If the serum protein concentration is normal, the CSF protein level should theoretically rise by 1 mg/dL for every 1000 RBCs, but this relationship varies. The inflammatory effect of hemolyzed RBCs may also significantly increase CSF protein.

Selective measurement of immunoglobulin fractions in CSF has proved to be of diagnostic value in suspected cases of multiple sclerosis. Elevated CSF immunoglobulin levels may reflect disruption of the blood–brain barrier or a local antibody response to a CNS immune response.134 Stimuli may be infectious or antigenic and produce an inflammatory response. Elevated immunoglobulin levels have been found in many conditions, including syphilis, viral encephalitis, subacute sclerosing panencephalitis, progressive rubella encephalitis, tuberculous meningitis, sarcoidosis, cysticercosis, andacute inflammatory demyelinating polyneuropathy (Guillian-Barré syndrome).

The Traumatic Tap

The incidence of a traumatic tap is 10% to 30%, depending on the criteria used to define the condition.135–137 Currently, there is no consensus on what constitutes a traumatic tap. Traditionally, the number of RBCs in CSF, the rate of clearance of RBCs from tube 1 to tube 3 or 4, and the presence or absence of xanthochromia have guided clinicians in attempts to define the need for further investigation for SAH when blood is detected during lumbar puncture.

Absolute Number of RBCs

The current literature makes no firm recommendations regarding the absolute CSF RBC count that can be used as a cutoff to differentiate SAH from a traumatic tap. However, SAH consistently produces more RBCs in CSF than a traumatic tap does. In tube 3 or 4, an absolute RBC value of 400 to 500 RBCs/µL or less is very suggestive of a traumatic tap, and this value has been traditionally used by clinicians.11 In a retrospective study of 300 patients, Gorczynski and coworkers reported a 100% negative predictive value for SAH with an RBC count in tube 4 of 500 RBCs/µL or less, with a sensitivity of 100% for SAH.138 No radiographically normal subject had an RBC count higher than 10,000 RBCs/µL, suggesting that results above this number are suspicious for radiographically detectable SAH. When the RBC count in tube 4 ranged between 500 and 10,000 RBCs/µL, SAH could not be ruled out without further study. Perry and colleagues concluded that the combination of an RBC count of fewer than 2000 × 10⁶ in the final tube and no xanthochromia reasonably excluded the diagnosis of aneurysmal SAH with a sensitivity of 100% and 95% confidence interval of 74.7% to 100%.139 They added that patients with a high pretest probability might need further investigation.

Any attempt to define precisely how low an RBC count must be to eliminate the possibility of SAH would result in an arbitrary threshold. As the foregoing discussion suggests, SAH is exceedingly unlikely with RBC counts lower than 500 RBCs/µL and becomes progressively less likely as this number decreases.

RBC Clearance From First to Last Tubes

In traumatic punctures, the fluid generally clears of RBCs between tubes 1 and tubes 3 or 4 as the needle is washed by CSF. In fact, a decrease in the RBC count between the first and last tubes of at least 25% to 30% has traditionally been considered strong evidence of a traumatic tap.140 However, it must be remembered that a traumatic tap may also occur in a patient with true SAH. Thus, regardless of the change in RBC count from the first to last tubes, SAH cannot be excluded unless the RBC count approaches zero in either tube. If this is not the case and xanthochromia is not present (see later), it may be most prudent to repeat the puncture at a different interspace or site. To help avoid this situation, when a clinician encounters what is suspected to be a traumatic tap (e.g., a streak of blood flowing into an otherwise clear-appearing stream of CSF), wasting the first 2 or 3 mL of CSF while positioning the needle to obtain the clearest possible sample will increase the odds of an RBC count approaching zero.

Xanthochromia

RBCs undergo hemolysis in CSF after a few hours and xanthochromia is produced. Xanthochromia persists for up to 4 weeks, depending on the number of RBCs originally present. Xanthochromia may be suggestive of but not pathognomonic for SAH.141 An early CSF examination may show clear fluid before the development of hemolysis, even after spontaneous subarachnoid bleeding. However, xanthochromia may be detected immediately after a traumatic tap if the RBC count exceeds 30,000/µL.138 The presence of a clot in one of the tubes strongly favors a traumatic tap. With SAH, clotting does not occur because blood is defibrinated at the site of the hemorrhage. Lumbar puncture performed several days after a traumatic tap may also yield stained fluid. Collection of clear CSF from an immediately repeated puncture at a higher interspace also indicates a traumatic tap. The fluid from a traumatic tap should contain approximately 1 WBC per 700 RBCs if the complete blood cell count is normal, but this ratio is highly variable. All blood-contaminated CSF should be cultured, especially samples from uncooperative infants and children being evaluated for sepsis.
A D-dimer test on CSF can be used to determine SAH by identifying local fibrinolysis. Other conditions such as disseminated intravascular coagulation, previous traumatic taps, or prior thrombolytic therapy may produce false-positive results.

Fluoroscopically guided lumbar puncture in patients with suspected SAH and negative findings on CT is another option that may reduce the frequency of traumatic punctures, but it requires specific expertise and is frequently performed in the radiology suite.

**CSF Analysis With Infections**

**Bacterial Infections**

The CSF findings are essential to establish a provisional diagnosis of acute bacterial meningitis. CSF analysis establishes not only the diagnosis but also the causative organism and therefore the choice of antibiotics (Table 60.1). CSF must be transported to the laboratory immediately and examined at once. CSF cells begin to lyse within 1 hour after collection; this process can be slowed by refrigeration. In cases of meningococcal infection, a delay in processing may cause the diagnosis to be missed because the organism tends to autolyze rapidly. For other organisms, speed is somewhat less important but still warranted.

Gram stain is of great importance because the results direct antibiotic therapy. Gram-negative intracellular or extracellular diplococci are indicative of *Neisseria meningitidis*. Small gram-negative bacilli may indicate *Haemophilus influenzae*, especially in children. The presence of gram-positive cocci indicates *Streptococcus pneumoniae*, other *Streptococcus* species, or *Staphylococcus*. Twenty percent of Gram stains are falsely negative because too few organisms are present. The Gram stain smear is more likely to be positive in patients who have not received prior antibiotic therapy. Acidine orange stain may improve the yield with gram-negative organisms.

For culture, blood and chocolate agar are required. *N. meningitidis* and *H. influenzae* grow best on chocolate agar. The plates are incubated under 10% CO₂. Thioglycolate medium is used for possible anaerobic organisms. Cultures are examined at 24 and 48 hours, but the plates should be kept for at least 7 days. Large volumes of CSF may improve yields.

While the culture results are pending, bacterial infection should be suspected in patients with an elevated opening pressure and marked pleocytosis ranging between 500 and 20,000 WBCs/µL. The differential count with bacterial infections is usually neutrophil predominant. A count higher than 1000 cells/µL seldom occurs with viral infections. Occasionally, acellular fluid may be collected from a severely immunosuppressed patient. Moreover, repeated lumbar puncture may be required in febrile patients whose clinical features remain compatible with meningitis. In such scenarios, broad-spectrum empirical antibiotics should be continued until the results of repeated testing or bacterial culture are available.

**CSF Analysis in Bacterial and Viral Meningitis**

<table>
<thead>
<tr>
<th>TABLE 60.1 CSF Analysis in Bacterial and Viral Meningitis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BACTERIAL MENINGITIS</strong></td>
</tr>
<tr>
<td><strong>VIRAL MENINGITIS</strong></td>
</tr>
<tr>
<td>Opening pressure</td>
</tr>
<tr>
<td>White blood cell count (per mm³)</td>
</tr>
<tr>
<td>Differential count</td>
</tr>
<tr>
<td>Glucose level</td>
</tr>
<tr>
<td>Protein level</td>
</tr>
</tbody>
</table>

*This is only a guide. Care must be taken when interpreting these parameters, especially early in the clinical course.

CSF, Cerebrospinal fluid.

In addition, blood and urine can be examined for antigen. Agglutination tests can detect levels of antigen that are approximately 10 times lower than CIE is capable of detecting. False-positive results can occur in the presence of rheumatoid factor, serum complement components, and possibly other serum proteins. The technique may be used for infections with *H. influenzae*, *S. pneumoniae*, *N. meningitidis*, and group B streptococci.

Another technique that has some potential use for identification of bacterial meningitis is the enzyme-linked immunosorbent assay. This technique may detect levels of antigen 100 to 1000 times lower than agglutination tests can but is technically more difficult and requires 4 hours to perform.

A positive CSF antigen test may be expected in 70% to 90% of patients with *Neisseria* meningitis. This compares with a positive Gram stain in approximately 70% of patients. Positive latex antigen tests have been reported in approximately 60% of *S. pneumoniae* meningitis cases, with a positive Gram stain in 80%. A positive latex test and Gram stain are reported in approximately 85% of *H. influenzae* meningitis cases. Group B streptococci can be detected with 60% to 90% sensitivity.

Bacterial antigens may persist in CSF for several days after antibiotic therapy. With appropriate antimicrobial therapy, 25% to 33% of positive tests convert to negative per day. A negative test, however, does not rule out bacterial meningitis.

In addition, blood and urine can be examined for antigen. Frequently, antigen is found only in urine. Urine needs to be concentrated and may have the disadvantage of reflecting urinary tract infections. Moreover, the particle agglutination test for *H. influenzae* type B may be positive up to 10 days after children have received *H. influenzae* polysaccharide vaccine. Antigen tests are not useful in diagnosing gram-negative bacillary, staphylococcal, and *Listeria* meningitis. In addition, although antigen tests may identify the bacterial pathogen, they do not provide information about the antibiotic susceptibility of the organism.

Polymerase chain reaction (PCR) may aid in the rapid diagnosis of CNS infection when results of the aforementioned common techniques are suboptimal. PCR amplifies target nucleic acid in CSF by the use of repeated cycles of DNA synthesis. PCR requires the use of flanking DNA sequences at the opposite ends of the target DNA. Synthetic primers anneal to their respective recognition sequences at the opposite end of the target sequence; they serve as primers for new DNA synthesis. PCR allows detection and quantification of organisms whose genetic material is DNA or messenger RNA. PCR permits the diagnosis of infectious disease with a high degree of sensitivity and specificity and allows rapid reliable detection of microbes present in small numbers. Nonetheless, false-negative and false-positive laboratory results may occur.

**Empirical Antibiotic Use Before Lumbar Puncture**

Many patients are transported within a facility or to a referral center for a CT to rule out an intracranial mass after clinical concern for meningitis is raised. In such instances, CSF examination might not be performed before transport because of technical problems (an uncooperative or a large patient) or concerns regarding the safety of lumbar puncture in an obtunded patient with possible increased ICP. The initial clinician may have to decide whether to initiate empirical antibiotic therapy. Antibiotic administration could obscure the bacterial source, whereas a delay in initiating therapy increases morbidity and mortality. It may be difficult to identify individuals at risk for a fulminant course, and bacterial meningitis cannot always be diagnosed with confidence; some cases might be misdiagnosed as SAH or metabolic encephalopathy.

After administration of parenteral antibiotics, CSF cultures are not adversely affected for 2 to 3 hours. Twenty-four hours after treatment, as many as 38% of patients with meningitis could still have positive CSF cultures. Kanegaye and associates demonstrated that CSF sterilization may depend on the infecting organism. They reported CSF sterilization after antibiotic administration within the following time frames: meningococcal, less than 2 hours; pneumococcal, less than 4.3 hours; and group B streptococcal, longer than 8 hours.

Whenever possible, blood should be obtained for culture immediately before the administration of antibiotics. When CSF is cultured more than a few hours after parenteral antibiotics are administered, antigen tests may be helpful. Occasionally, lymphocytic pleocytosis may develop in response to antibiotic therapy, but in most cases, the cell count, differential, glucose and protein concentrations are unchanged in the first 2 to 3 days of antibiotic therapy. It is thus reasonable to initiate therapy on the premise that a delay might be deleterious. If a lumbar puncture cannot be performed, consultation with clinicians at a referral center seems appropriate. If a lumbar puncture is performed before transfer, a portion of the CSF (chilled on ice) should be sent with the patient or held in the referring hospital laboratory.

Bacterial meningitis in children younger than 10 years has historically been caused by *H. influenzae*. Fortunately, this organism is easy to grow in early postantibiotic cultures and is likely to be associated with positive blood cultures and antigen tests. In the pediatric population, a single dose of an antibiotic before transport is unlikely to prevent identification of bacteria. In neonates, adults, and immunosuppressed patients, the sensitivity of blood cultures and immunologic tests is less reliable. CSF examination as early as possible in the course of treatment is preferred.

For suspected or confirmed cases of acute bacterial meningitis, antimicrobial therapy can be started and based on the most likely causative organism with respect to the age of the subject, associated diseases, and renal function. Tables 60.2 through 60.4 offer guidelines for emergency antibiotic therapy. For immunocompromised patients and after neurosurgery, a third-generation cephalosporin (cefotaxime, ceftizoxime, cefazidime, or ceftriaxone) plus ampicillin and vancomycin should be used for coverage against staphylococci, *L. monocytogenes*, and gram-negative organisms. The third-generation cephalosporins are efficacious in many empirical regimens or situations in which the organism is known. Reliance on third-generation cephalosporins alone for all cases of bacterial meningitis would result in treatment failure for all *Listeria* species and increasing numbers of *Enterobacter*, *Serratia*, and *Pseudomonas* groups.

**Dexamethasone Therapy for Bacterial Meningitis**

In acute bacterial meningitis, bacterial cell wall components, including lipopolysaccharides and teichoic acid, initiate and exacerbate the host response. These substances stimulate the production of cytokines, including interleukin-1 and tumor necrosis factor, from macrophages and monocytes. Cytokines may injure vessels, diminish cerebral perfusion, and stimulate cerebral swelling. The outcome of acute bacterial meningitis has been related to the severity of the inflammatory process in the subarachnoid space.
### TABLE 60.2 Empirical Therapy for Purulent Meningitis

<table>
<thead>
<tr>
<th>PREDISPOSING FACTOR</th>
<th>ANTIMICROBIAL THERAPY</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td></td>
</tr>
<tr>
<td>0–4 wk</td>
<td>Ampicillin plus cefotaxime or ampicillin plus an aminoglycoside</td>
</tr>
<tr>
<td>4–12 wk</td>
<td>Ampicillin plus a third-generation cefalosporin</td>
</tr>
<tr>
<td>3 mo–50 yr</td>
<td>Vancomycin plus a third-generation cefalosporin</td>
</tr>
<tr>
<td>&gt;50 yr</td>
<td>Vancomycin plus ampicillin plus a third-generation cefalosporin</td>
</tr>
<tr>
<td>Immunocompromised state</td>
<td>Vancomycin plus ampicillin plus cefazidime</td>
</tr>
<tr>
<td>Basilar skull fracture</td>
<td>Vancomycin plus a third-generation cefalosporin</td>
</tr>
<tr>
<td>Head trauma; post neurosurgery</td>
<td>Vancomycin plus cefazidime</td>
</tr>
<tr>
<td>Cerebrospinal fluid shunt</td>
<td>Vancomycin plus cefazidime</td>
</tr>
</tbody>
</table>

*Consider corticosteroids before administration of antibiotics (see text).

*Vancomycin should be added to all empirical therapeutic regimens when strains of *Streptococcus pneumoniae* that are highly resistant to penicillin or cefalosporin are suspected.

*Cefotaxime or ceftriaxone.

*Add ampicillin if meningitis caused by *Listeria monocytogenes* is suspected.

### TABLE 60.3 Recommended Total Daily Dose (With Divided Dosing Intervals in Hours) of Antimicrobial Agents for Meningitis in Adults With Normal Renal and Hepatic Function

<table>
<thead>
<tr>
<th>ANTIMICROBIAL AGENT</th>
<th>TOTAL DAILY DOSE (INTRAVENOUS)</th>
<th>DOSING INTERVAL (hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ampicillin</td>
<td>8–12 g/day</td>
<td>4</td>
</tr>
<tr>
<td>Cefotaxime</td>
<td>8–12 g/day</td>
<td>4–6</td>
</tr>
<tr>
<td>Ceftazidime</td>
<td>6 g/day</td>
<td>8</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>4 g/day</td>
<td>12</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>4 g/day</td>
<td>6</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>2 mg/kg IV load, then 1.7 mg/kg q 8 hr</td>
<td>8</td>
</tr>
<tr>
<td>Tobramycin</td>
<td>2 mg/kg IV load, then 1.7 mg/kg IV q 8 hr</td>
<td>8</td>
</tr>
<tr>
<td>Vancomycin</td>
<td>30–60 mg/kg per day</td>
<td>8–12</td>
</tr>
</tbody>
</table>

*Consider corticosteroids before the administration of antibiotics (see text).

*High dose recommended for pneumococcal meningitis.

*Peak and trough serum concentrations must be monitored.

### TABLE 60.4 Recommended Total Daily Doses With Divided Dosing Intervals in Hours of Antimicrobial Agents for Meningitis in Neonates, Infants, and Children With Normal Renal and Hepatic Function

<table>
<thead>
<tr>
<th>ANTIMICROBIAL AGENT</th>
<th>NEONATES (0–7 DAYS)</th>
<th>NEONATES (8–28 DAYS)</th>
<th>INFANTS AND CHILDREN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amikacin†</td>
<td>15–20 mg/kg per day IV divided q 12 hr</td>
<td>30 mg/kg per day IV divided q 8 hr</td>
<td>15–22.5 mg/kg per day IV divided q 8 hr (max dose 1.5 g/day)</td>
</tr>
<tr>
<td>Ampicillin</td>
<td>200–300 mg/kg per day IV divided q 8 hr</td>
<td>400 mg/kg per day IV divided q 6 hr</td>
<td>300–400 mg/kg per day IV divided q 3–4 hr (max dose 10–12 g/day)</td>
</tr>
<tr>
<td>Cefotaxime</td>
<td>100–150 mg/kg per day IV divided q 8–12 hr</td>
<td>200 mg/kg per day IV divided q 6 hr</td>
<td>200–300 mg/kg per day IV divided q 6–8 hr (max 12 g/day)</td>
</tr>
<tr>
<td>Ceftazidime</td>
<td>100–150 mg/kg per day IV divided q 8–12 hr</td>
<td>100 mg/kg per day IV divided q 12 hr</td>
<td>150 mg/kg per day IV divided q 8 hr (max 6 g/day)</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>—</td>
<td>—</td>
<td>100 mg/kg per day IV divided q 12 hr</td>
</tr>
<tr>
<td>Chloramphenicol</td>
<td>25 mg/kg per day IV q 24 hr (initial loading 20 mg/kg, then first maintenance dose 12 hr later)</td>
<td>25–50 mg/kg per day IV divided q 12–24 hr (initial loading 20 mg/kg, then first maintenance dose 12 hr later)</td>
<td>75–100 mg/kg per day IV divided q 6 hr (max 2–4 g/day)</td>
</tr>
<tr>
<td>Gentamicin†</td>
<td>2.5 mg/kg per day IV divided q 8 hr†</td>
<td>4–5 mg/kg per day IV divided q 12–48 hr†</td>
<td>7.5 mg/kg per day IV divided q 8 hr†</td>
</tr>
<tr>
<td>Tobramycin†</td>
<td>4–5 mg/kg per day IV q 24–48 hr†</td>
<td>4–5 mg/kg per day IV divided q 12–48 hr†</td>
<td>5–7.5 mg/kg per day IV divided q 8 hr†</td>
</tr>
<tr>
<td>Vancomycin†</td>
<td>10–15 mg/kg per day IV divided q 8–12 hr</td>
<td>45–60 mg/kg per day IV divided q 8–12 hr</td>
<td>60 mg/kg per day IV divided q 8 hr</td>
</tr>
</tbody>
</table>

*Consider corticosteroids before the administration of antibiotics (see text).

*Smaller dosages and longer intervals of administration may be advisable for very-low-birth-weight neonates (< 2000 g).

*Peak and trough serum concentrations must be monitored.
Studies have suggested benefit from adjunctive dexamethasone therapy in reducing neurologic sequelae, especially hearing loss, in children with *H. influenzae* meningoencephalitis and in lowering the mortality rate and providing a better overall outcome in adults with community-acquired acute bacterial meningitis caused by *S. pneumoniae*. There appear to be few adverse sequelae from steroid therapy. The adjunctive benefit of corticosteroids during treatment of meningitis caused by other viral, fungal, or parasitic organisms is unknown. A Cochrane review concluded that “the corticosteroid dexamethasone leads to a major reduction in hearing loss and death in both children and adults with bacterial meningitis, without major adverse effects.” Interestingly, in children in low-income countries, the use of corticosteroids was associated with neither benefit nor harmful effects.

The dosage of dexamethasone is 0.15 mg/kg (10 mg intravenously in adults) every 6 hours for 4 days. It is recommended that the corticosteroid be given before or simultaneously with antibiotic use, but the exact timing and specific benefits are unclear. It has been suggested that caution be exercised when administering corticosteroids to immunocompromised and leukopenic patients with bacterial meningitis, but no specific recommendations have been published.

A moderate inflammatory response in the meninges is required for penetration of the CNS by many antibiotics. Reducing meningeal inflammation reduces the concentration of antibiotics in CSF. Corticosteroids should be discontinued after approximately 4 days of treatment, at which time the meningeal inflammation should be reduced.

**Neurosyphilis**

The true incidence of neurosyphilis is unknown. Approximately 5000 new cases of this disease occur in the United States each year. Its natural history and clinical manifestations have been modified in the antibiotic era. The widespread use of oral antibiotics has changed neurosyphilis into chronic, partially treated meningitis. Partial therapy might clear peripheral infection and attenuate the immune response. Therapy may be sufficient to minimize symptoms but insufficient to eradicate organisms in the CNS and eye, which could then multiply.

CSF findings suggestive of neurosyphilis include more than 5 WBCs/µL, an elevated protein concentration, an elevated γ-globulin concentration, and a positive serologic test for syphilis. The glucose concentration is usually normal. Cell and protein values are higher in early neurosyphilis than in late neurosyphilis.

Diagnostic certainty remains difficult. The diagnostic criterion standard is darkfield microscopy to identify the morphology and flexing “corkscrew” motility of spirochetes. Serologic tests for syphilis are either treponemal or nontreponemal. Nontreponemal tests detect a nonspecific globulin complex called reagin. Reagin tests, such as the Venereal Disease Research Laboratory (VDRL) flocculation test, lack sensitivity and should not be used to exclude the diagnosis of neurosyphilis. One third to one half of patients with neurosyphilis have a negative VDRL test on serum, and more than one-third have negative VDRL tests on CSF. CSF VDRL is quite specific, with false-positive results seen primarily after traumatic taps.

Treponemal tests provide evidence of a specific immune response to *Treponema pallidum*. These tests include serum fluorescent treponemal antibody absorption (FTA-ABS), microhemagglutination tests for *T. pallidum*, and the *T. pallidum* hemagglutination assay. A positive serum treponemal test indicates past infection with syphilis and may be reactive indefinitely, even after treatment. Therefore CSF is used as a guide to the presence and activity of neurosyphilis. The VDRL test is commonly used on CSF and, when positive, is strong evidence of neurosyphilis. False-positive CSF VDRL tests are rare. The FTA-ABS test can be used on CSF; the false-positive rate is between 4% and 6% and is believed to represent antibodies that have entered passively from serum. The FTA-ABS test measures immunoglobulin G (IgG) antibody and cannot differentiate active from past infection. The CSF VDRL may be reactive as a result of contamination with seropositive blood (traumatic tap, SAH) or because of entry of serum reagin into CSF during meningitis. In summary, CSF VDRL and CSF FTA-ABS are complementary tests in the diagnosis of neurosyphilis: CSF VDRL is highly specific but not sensitive, whereas CSF FTA-ABS is less specific but more sensitive.

There is some concern that many patients with parenchymal neurosyphilis have normal CSF. This finding has led to the recommendation that a patient with signs of progressive neurosyphilis and a positive treponemal serologic test be treated with antibiotics regardless of the CSF findings. CSF pleocytosis may be provoked after 1 week of therapy and may provide supportive evidence for a diagnosis of neurosyphilis. PCR may have future applications for the diagnosis of neurosyphilis (see later section on Neurosyphilis in Patients Infected with HIV).

**Viral Meningitis**

The organisms most commonly isolated in viral meningitis are the enteroviruses (coxackieviruses, echoviruses) and mumps virus. Enteroviruses are most commonly seen in the summer and fall, and mumps appears most frequently in the winter and spring. Viral cultures in most hospitals are not available and play little role in acute decisions regarding diagnosis and treatment. A serial rise in CSF antibody titers may be helpful but are difficult to obtain in patients who have recovered clinically. Intrathecal production of organ-specific antibodies (IgM, IgG, and IgA isotopes) may be diagnostic of neurologic infection if there is no history of infection. Serum and CSF antibody titers must be measured in a specialized laboratory. Viral meningitis is diagnosed when bacterial culture and Gram stain are negative. A tentative diagnosis may be based on analysis of CSF.

The WBC count in viral meningitis and encephalitis is characteristically 10 to 1000 cells/µL. The differential cell count is predominantly lymphocytic and mononuclear in type. In the early stages of meningoencephalitis, however, polymorphonuclear cells may predominate, making the distinction between viral and bacterial infection difficult. In such cases, a tap repeated in 12 to 24 hours will assist in clarifying the diagnosis. Protein levels are usually mildly elevated, but normal levels may be seen. Antibiotic coverage pending the results of culture may be reasonably initiated if the diagnosis of viral meningitis is in doubt. The CSF glucose concentration is characteristically normal; notable exceptions include some cases of mumps meningoencephalitis and HSV encephalitis. CSF pleocytosis and elevated protein levels have also been found in asymptomatic HIV-seropositive individuals.

If CSF cannot be delivered to the viral laboratory in 24 to 48 hours, it should be refrigerated at 4°C. Members of the enterovirus group are occasionally isolated from CSF. Herpesviruses and arboviruses are also found in CSF. PCR is the diagnostic test of choice for HSV meningoencephalitis and
will be used increasingly for the diagnosis of other CNS viral infections. Although the majority of cases of viral meningitis are self-limited and have a good outcome, HSV encephalitis is a rapidly progressive and life-threatening emergency that responds to treatment with intravenous acyclovir. Thus, in an acutely ill patient in whom the diagnosis of acute meningitis-encephalitis is being considered, a negative Gram stain and pleocytosis are sufficient grounds to initiate empirical acyclovir therapy until the results of more definitive diagnostic data from PCR are available.162

**CSF Analysis in Immunocompromised Patients**

The number of immunocompromised individuals is increasing because of the HIV epidemic and the increased survival of patients with cancer and autoimmune disorders. The nervous system is a major target of the HIV virus: neurologic disease develops in 40% to 60% of infected individuals during their lifetime. One third of HIV-infected patients have neurologic complaints as the initial manifestations of AIDS, and an even higher incidence of nervous system involvement is found at autopsy.161 The risk for CNS infection depends on the underlying disease, treatment, duration, and type of immune abnormality. Abnormalities include defects in T-lymphocyte and macrophage cellular immune function, defects in humoral immunity, defects in the number and function of neutrophils, and loss of splenic function with an inability to remove encapsulated bacteria.164

Patients with defective cell-mediated immunity include those with lymphoma or organ transplants, those taking corticosteroids daily, and patients with AIDS. These individuals are vulnerable to infections with microorganisms that are intracellular parasites. A common source of acute bacterial meningitis in such patients is *L. monocytogenes*. Clinical findings include fever, headache, seizures, focal neurologic deficits, and brain stem encephalitis.165

Patients with defective humoral immunity include those with chronic lymphocytic leukemia, multiple myeloma, and Hodgkin’s disease after radiotherapy or chemotherapy. These patients have difficulty controlling infection by encapsulated bacteria. A fulminating meningitis caused by *S. pneumoniae*, *H. influenzae* type B, or *N. meningitidis* may develop. After splenectomy, patients are at risk for the development of meningitis for the same reason. Neutropenic patients are at risk for meningitis caused by *Pseudomonas aeruginosa* and the *Enterobacteriaceae*.166

Establishing a specific diagnosis in HIV and organ transplant patients may be difficult or impossible because of overlapping clinical and radiographic findings, the presence of simultaneous infections with more than one organism, and changes in CSF that may be nonspecific. The immune response can be altered, with absence of the usual signs of meningeal irritation. Patients might have diffuse encephalopathy or focal neurologic deficits. CSF is abnormal in 60% of asymptomatic HIV-infected individuals, complicating correlation of the CSF and clinical findings.163

**Toxoplasmosis**

*Toxoplasma gondii*, an intracellular protozoan, is associated with CNS infection in up to 30% of AIDS patients who have antibodies to this organism. Most adults have antibodies against this organism; infection is believed to represent reactivation of latent primary infection. *Toxoplasma* encephalitis usually develops within the first 2 years after the diagnosis of AIDS. Cerebral toxoplasmosis is usually accompanied by the acute or subacute development of focal disease, including seizures. MRI or CT scanning often reveals multiple abscesses that might represent multiple pathogens. Toxoplasmosis and lymphoma can be difficult to distinguish clinically. Less commonly, the manifestation is one of chronic meningitis with confusion, memory loss, and lethargy similar to the AIDS-dementia complex. CSF in these patients is nonspecific with increased protein, mononuclear pleocytosis (<100 cells/µL), and, rarely, a reduced glucose concentration. Serum and CSF serology may be either positive or negative and does not help make a diagnosis, although most patients with encephalitis...
have detectable IgG antibodies. The diagnosis is usually based on clinical and imaging responses to antibiotics (pyrimethamine or sulfadiazine) or on brain biopsy. Treatment failures with relapse occur in 50% of AIDS patients and in 15% to 25% of non-AIDS patients and necessitate life-long treatment.

**Mycobacterial Tuberculosis**

CNS mycobacterial infection is almost always the result of infection with *Mycobacterium tuberculosis*. Infection typically occurs in the setting of disseminated tuberculosis. Atypical *Mycobacterium* infection in HIV-infected individuals uncommonly affects the CNS.

The clinical manifestation is meningitis (particularly meningitis involving the basal cisterns), encephalitis, or abscess formation. If tuberculosis is suspected, a large volume of CSF (10 mL) is required for adequate culture. The cell count varies from 100 to 400 cells/µL, with a lymphocytic predominance; 30% may show predominantly neutrophils early in the course of infection. Protein levels are elevated (100 to 500 mg/dL), and the CSF glucose level may be depressed. Acid-fast stains should be examined by experienced technicians. Fluid is inoculated onto Löwenstein-Jensen medium, and the absence of visible growth on the medium should not be considered negative until 8 weeks has elapsed. CSF cultures are more sensitive than stains.

**Primary CNS Lymphoma**

CNS lymphoma develops in 2% of AIDS patients. The signs and symptoms resemble those of diffuse encephalopathy, although focal neurologic deficits occur occasionally. Leptomeningeal spread of malignancy is reflected in a modest lymphocytic pleocytosis with slightly elevated protein and decreased glucose levels. Cytologic yield is improved by repeated lumbar punctures and submitting large quantities of CSF or fluid obtained by cisternal puncture.

**Progressive Multifocal Leukoencephalopathy**

Progressive multifocal leukoencephalopathy, an uncommon disorder in individuals with impaired cell-mediated immunity, is caused by reactivation of JC papovavirus in the kidney. Progressive demyelination is manifested as dementia, blindness, aphasia, hemiparesis, and seizures, which progress until death. MRI and CT demonstrate nonenhancing white matter lesions without a mass effect. Definitive diagnosis is made by brain biopsy, but CSF may show the presence of myelin basic protein, increased IgG, and an acellular or a mild CSF pleocytosis (<50 WBCs/µL). Average survival is 4 months.

**Cytomegalovirus Infection**

Cytomegalovirus is detected in the brains of 30% of HIV-infected persons at autopsy. A distinct CNS disorder has not been defined. CSF pleocytosis may be minimal. Retinitis and painful polyradiculopathies are recognized, with a prominent CSF pleocytosis found in the latter condition.

**ACKNOWLEDGMENTS**

The editors and author acknowledge the contributions of Jon Kooiker to this chapter in previous editions. The author would like to thank Linda J. Kesselring, MS, ELS, for copyediting the manuscript and incorporating revisions into the final document.

References are available at www.expertconsult.com
REFERENCES


EVALUATION OF THE COMATOSE PATIENT

Coma is defined as a state of deeply reduced consciousness from which a patient cannot be aroused by verbal, tactile, or noxious external stimuli. This subsection will review procedures that have important diagnostic and prognostic significance in the comatose patient: caloric testing and the evaluation of brain death.

Caloric Testing

Background

In a comatose individual with normal brain stem and cranial nerve function, stimulation of the vestibular labyrinth results in well-described and reproducible extraocular movements through the vestibulo-ocular reflex (VOR). Pathologic conditions involving the labyrinth, vestibulocochlear nerve, or the oculomotor reflex pathways in the brain stem will alter or abolish the VOR.

Physiology and Functional Anatomy

The anatomic pathways underlying the VOR begin in the lateral semicircular canal of the inner ear. Movement of endolymphatic fluid within the canal results in changes in polarization in the underlying hair cells, which in turn are relayed via cranial nerve VIII (vestibular) to the brain stem (Fig. 61.1).

There are two main pathways between the vestibular and oculomotor nuclei in the brain stem. The direct pathway travels from the vestibular complex to the nuclei of the ipsilateral third cranial nerve (oculomotor) and contralateral sixth cranial nerve (abducens), and ultimately to the ipsilateral medial rectus and contralateral lateral rectus muscles. The indirect pathway occurs over multisynaptic circuits in the tegmental reticular formation.

Rotation of the head generates flow of endolymphatic fluid within the semicircular canals; the direction of flow increases or decreases stimulation of the vestibular neuron, which results in conjugate deviation of the eyes away from or towards the affected side, respectively. This principle forms the physiologic basis of caloric testing. When the lateral canal is in the vertical position (head elevated 30 degrees above horizontal) and ice water is infused into the ear, the endolymph nearest the canal cools and sinks, which results in decreased stimulation of the vestibular neuron, and the eyes deviate conjugately toward the side of irrigation. When warm water is used in the same position or when the canal is inverted 180 degrees, the opposite occurs.

Indications and Contraindications

Caloric testing is infrequently used in emergency department (ED) settings in the United States where advanced neuroimaging is readily available. However, in resource-limited settings, caloric testing at the bedside may assist in the evaluation of comatose patients by differentiating structural, metabolic, and psychogenic causes of unresponsiveness.

Few contraindications exist to caloric testing of an unresponsive patient. An absolute contraindication is the presence of a basilar skull fracture, either documented radiologically or suspected by clinical signs, because of the risk of introducing infection into the central nervous system (CNS) through an associated dural tear. If active bleeding or cerebrospinal fluid otorrhea or rhinorrhea is present in a trauma patient, defer caloric testing and evaluate the patient for a potential basilar skull fracture. Tympanic rupture, hemotympanum, or step-off deformities of the auditory canal may indicate fracture of the temporal bone, which is a contraindication to caloric testing. Signs of active ear infection and perforation of the tympanic membrane are relative contraindications to caloric testing.

Equipment

A 30- or 50-mL plastic syringe is ideal for irrigation. The syringe may be used as is, or a short length of soft plastic tubing (e.g., a butterfly catheter with the needle cut off) may be attached. At least 100 mL of ice water should be available, although larger quantities of cool tap water (<25°C) can be used with similar results if ice is unavailable. Saline or tap water may be used. A small, curved plastic emesis basin is useful to collect water as it drains from the ear canal. Other equipment required includes an otoscope, ear specula, and equipment for removal of cerumen. Towels and a thermometer may be helpful.

Procedure

Inspect the ears before inserting the otoscope. Remove excess cerumen and foreign material. Clearly visualize the tympanic membrane. Leave the ear speculum in the canal as a guide for irrigation. Perform the test with the patient supine and the head and upper part of the body raised to 30 degrees, if possible. Drape the patient with a towel and position a small emesis basin below the ear to collect the outflow of water from the ear. Fill a container with ice water and place it near the bedside.

Fill a syringe and catheter system (without the needle) with 10 mL of ice water and direct the irrigation stream toward the tympanic membrane. Because the goal of qualitative caloric testing is to induce a maximum response, the amount and rate of infusion are not critical. As a general guide, infuse 5 to 10 mL of ice water initially over a period of 5 to 10 seconds. Amounts less than 5 mL may be advisable in suspected cases of light coma or psychogenic unresponsiveness. If no response is noted at first, infuse at least 100 mL before declaring that there is no response. Ask an assistant to hold the patient’s eyelids open, which makes it easier to observe for eye deviation. Movement usually occurs after a latency of 10 to 40 seconds, with the response persisting for as long as 4 to 5 minutes. Focusing on a scleral vessel makes small deviations easier to detect. Wait at least 5 to 10 minutes after the eyes have returned to their original position before testing the contralateral ear.

Complications

Carefully selecting equipment can avoid the few complications that are possible with caloric testing. Using plastic syringes and soft catheter tubing instead of needles to irrigate the ear will reduce the risk of perforation of the tympanic membrane or canal wall injury if the patient moves unexpectedly.
As mentioned in the contraindications section, caloric testing should not be performed in patients with possible basilar skull fractures, due to the theoretical risk of meningitis. Other potential complications include otitis media and induction of vomiting with subsequent aspiration. Although ice water irrigation might produce nausea and emesis in awake patients, vomiting with aspiration has not been reported as a complication of caloric testing in comatose patients. Nevertheless, some may prefer to delay testing until the patient’s airway is protected.

**Interpretation**

After irrigation, ocular movements should be observed for any response to the stimulus. Typically, there is a latency of response of 10 to 40 seconds. Reactions to ice water irrigation may be divided into four categories: (1) caloric nystagmus, (2) conjugate deviation, (3) dysconjugate deviation, and (4) absent responses (Fig. 61.2).

The first reaction, caloric nystagmus, is seen in normal, alert individuals. Ice water infusion induces a rhythmic jerking of the eyes that includes a slow deviation toward the irrigated side followed by a quick compensatory saccade toward the midline. In an apparently comatose individual, caloric nystagmus usually signifies psychogenic unresponsiveness due to catatonia, conversion reactions, schizophrenia, or feigned coma, but can also be present in patients with very mild organic disturbances in consciousness. The response is present in more than 90% of children by 6 months of age and declines in magnitude only after the seventh decade of life.

In the second type of response to cold caloric stimulation, conjugate deviation, the eyes deviate conjugately toward the
CHAPTER 61  Special Neurologic Tests and Procedures

1. Normal nystagmus

2. Conjugate deviation

3. Dysconjugate deviation (with MLF lesion)

4. Absent responses

<table>
<thead>
<tr>
<th>Oculocerephalic Testing</th>
<th>Vertical head rotation</th>
<th>Oculovestibular Testing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Horizontal head rotation</td>
<td>Cold H₂O</td>
<td>Cold H₂O</td>
</tr>
<tr>
<td>Vertical head rotation</td>
<td>Cold H₂O</td>
<td>Cold H₂O</td>
</tr>
</tbody>
</table>

- **Doll's eyes movements do not occur in alert persons.**
- **Conjugate downward eye movement when head is moved backward.**
- **No consistent movement.**
- **Dysconjugate movement. Left eye does not move medially.**
- **No horizontal movement.**
- **Eyes move downward by third cranial nerve action.**

**Note:** Illustrations by Charles H. Boyer.
side of ice water stimulation (they “look” toward the source of irritation). When present, this reaction indicates intact brain stem function, as well as intact afferent and efferent limbs of the VOR. This is seen during general anesthesia, in supratentorial lesions without brain stem compression, and with many metabolic and drug-induced comas.

Dysconjugate reactions constitute the third type of caloric response to ice water stimuli. The most common dysconjugate reaction is internuclear ophthalmoplegia, in which a lesion of the medial longitudinal fasciculus causes weakness or paralysis of the adducting eye after caloric irrigation. Internuclear ophthalmoplegia can be due to acute damage to the rostral pons or as a manifestation of multiple sclerosis or stroke. With acute supratentorial lesions, the development of dysconjugate caloric responses is a significant sign that may indicate compression of the brain stem and impending herniation. Caloric responses of this type are less common with metabolic and drug-induced coma. Reversible internuclear ophthalmoplegia has been reported in patients with hepatic coma and may occur during toxic responses to phenytoin, barbiturates, or amitriptyline.

Absent caloric response is the fourth category of reactions to ice water stimuli. Loss of caloric responses in comatose patients with structural lesions is usually a sign of brain stem damage. The VOR may also be transiently absent or decreased on the side opposite massive supratentorial damage during the first hours after injury.4 Whereas the initial absence of the VOR is generally a poor prognostic indicator, it does not uniformly predict a poor outcome when performed in the first 24 hours after presentation.5 This has particular relevance in cases of traumatic coma, where testing in the first 24 hours after injury may generate variable responses and is of less prognostic value.6 Absent caloric responses may occur with any subtentorial lesion that affects the vestibular reflex pathways, including pontine hemorrhage, basilar artery occlusion, cerebellar hemorrhage, or infarction with encroachment on the brain stem, and with any expanding mass lesion within the posterior fossa. Caloric responses may disappear in patients with deep coma resulting from subarachnoid hemorrhage, perhaps because of pressure on the brain stem.

The VOR is usually retained until the late stages of metabolic coma. Nevertheless, caloric responses may be transiently absent in certain types of drug-induced coma, with eventual complete recovery of the patient. The VOR seems to be particularly sensitive to the effects of sedative-hypnotic drugs, antidepressants (e.g., amitriptyline, doxepine), and anticonvulsants (e.g., phenytoin, carbamazepine).7,8 As one would expect, neuromuscular blocking agents (e.g., succinylcholine) will abolish caloric-induced ocular movements.

Finally, the caloric response may be absent for reasons other than the neurologic causes responsible for the coma. Inadequate irrigation because of excessive cerumen or poor technique and unilateral or bilateral dysfunction of the peripheral vestibular apparatus must be considered. Bilateral loss of the caloric response (areflexia vestibularis) is uncommon in conscious patients, constituting 1.7% and 0.2% in two large series of patients.9,10

**Summary**

Caloric testing is a simple, easily performed bedside procedure that may enhance the neurologic assessment of comatose patients. In the ED this test should be reserved for stable patients undergoing secondary assessment. The examination requires minimal equipment and can be particularly useful in settings where access to advanced neuroimaging is limited or delayed. Complications are few if patients are properly selected and correct technique is used.

**Brain Death Testing**

**Background**

Brain death is defined as irreversible and complete loss of cerebral and brain stem function with preserved cardiac function.11 Prior to declaring brain death, the cause of the brain injury should be definitively identified, as some toxicologic syndromes may mimic brain death. The most common scenario where emergency clinicians may be asked to perform a brain death examination is to identify potential organ and tissue donors. It should also be noted that a determination of brain death is not necessary for withdrawal of life-supporting measures. If a cause of coma or severe neurologic injury is identified and a poor prognosis is shared with the family, withholding or withdrawing life-sustaining treatment may be appropriate and formal determination of brain death need not be performed.

**Indications and Contraindications**

Evaluation for brain death implies that severe CNS dysfunction has been identified, that the cause of the CNS dysfunction is known, and that reversible causes of coma have been confidently excluded.12,13 Complex medical issues that may confound the assessment should be considered and ruled out, including severe electrolyte disturbances, hypothermia (defined as a core temperature <32°C), hypotension, drug intoxication or poisoning, and pharmacologic neuromuscular blockade.14 Neuroimaging studies should be carefully reviewed.

**Procedure**

The clinical neurologic examination remains the standard for determination of brain death.15 It typically involves assessment for cerebral function, brain stem reflexes, and respiratory drive. The emergency clinician should be familiar with local practices and policies, which may also require an electroencephalogram, documentation of absent cerebral blood flow by angiography or radioisotope brain scan, or other techniques. The following components of the clinical examination are generally utilized to establish brain death.

**Coma Assessment**

While holding the patient’s eyes open, give loud verbal commands such as “Look up!” and assess for voluntary eye movements; this assessment is particularly important to identify patients with locked-in syndrome. Additionally, deliver a strong painful stimulus by forcefully pressing on the brow, sternum, or nail bed. Any purposeful response to stimuli in any extremity indicates the patient is not brain-dead. The absence of responses documents the presence of coma.

**Brain Stem Reflex Testing**

**Pupillary Response.** The pupils in brain-dead patients are unreactive and midposition to dilated. Shine a bright light into the pupil and observe for a reaction; none will be seen in a brain-dead patient.

**Auditory Reflex.** Deliver a loud handclap into each ear. Observe for eye blink or other reaction. Any reaction establishes that some brain stem function remains and excludes brain death.
Oculocephalic Reflex. Hold eyelids open and rotate the head abruptly from side to side in the horizontal plane. If the eyes do not turn with the head and appear to maintain visual fixation on a point in space, the oculocephalic reflex is present and brain death is excluded.

Caloric Testing. Perform cold water irrigation of the external auditory canals with large volumes (≥100 mL) to elicit any eye movements through the VOR. In a brain-dead patient, there will be no movement of the eyes in response to irrigation.

Corneal Reflex. Stimulate the cornea with a cotton wisp or applicator. Any eye closure indicates an intact cranial nerve V to VII reflex and excludes the diagnosis of brain death.

Cough Reflex. Stimulate the trachea or main stem bronchi by deep suctioning and observe for coughing. A cough excludes brain death.

Apneic Oxygenation Test. If brain stem reflexes are absent, apnea is formally tested to evaluate the function of the medulla, where respiratory drive originates. As hypercapnia (and not hypoxia) triggers respiratory effort, simply disconnecting the ventilator to allow hypercapnia to develop may lead to hypoxia and should therefore be avoided.14 The most commonly described technique to avoid hypoxia while generating hypercapnia is to: (1) disconnect the patient from the ventilator, (2) deliver oxygen at 10 to 15 L/min through a catheter inserted into the trachea, and (3) observe for respiratory effort for 8 minutes. An alternative technique is to set the ventilator rate to zero while allowing continuous oxygen flow to continue and maintaining any necessary continuous positive pressure through the ventilator.15 Any observed excursion of the abdomen or chest sufficient to produce a tidal volume suggests that brain death is not present. If no respiratory excursions are observed, an arterial blood gas analysis should be obtained and artificial ventilation resumed pending results. An arterial partial carbon dioxide pressure (PCO₂) of 60 mm Hg or higher and the absence of respiratory excursions are the criteria for a positive apnea test (i.e., that the patient lacks spontaneous respirations).16

Declaration of Death
If the criteria for brain death are satisfied, the family and all clinicians involved in patient care should be informed to allow further management decisions. At some institutions, the patient is declared dead at the time that the criteria are met, and further care is assumed by the transplant services if that is the anticipated course. Some institutions require evaluation by two independent clinicians with specific specialty training, or repeated examinations several hours apart; however, the need for a second brain death determination has been questioned.17,18 Families are generally given the option of being present at the bedside when mechanical ventilation is discontinued, although some advise against this policy because spontaneous reflex movements may occur and disturb the family.19

Brain death in the pediatric population is more complex, with varying recommendations for repeated examinations and ancillary tests; such discussion is outside the scope of this chapter but is summarized in other resources.15,20,25 A model for direct family conversation in this sensitive interaction has been described and includes a sample script and procedure.22

Complications
A substantial proportion of brain-dead patients (33% to 75%) may exhibit movements originating in the spinal cord or peripheral nerves.22,23 These movements can be spontaneous or provoked by tactile stimuli, and may falsely be interpreted as evidence of cerebral or brain stem function. Among the most frequently observed movements are finger jerks, the undulating toe flexion response (slow and repetitive flexion/extension of the toes), and the triple flexion reflex (flexion of thigh, leg, and foot); however, many other movements have been described.23,25–28

Summary
The emergency clinician may become involved in the assessment of patients for brain death. The cause of irreversible coma should be definitely identified prior to initiating a systematic brain-death examination, which includes an assessment of cerebral function, brain stem reflexes, and apnea. The clinician should be familiar with local practices and policies in this sometimes complex medicolegal process.

PROCEDURES TO EVALUATE AND TREAT VERTIGO
Vertigo is the illusory sensation of movement, sometimes experienced by patients as self-motion, other times as motion of the environment. There are a wide variety of diseases that can cause vertigo, including damage to or dysfunction of the labyrinth, vestibular nerve, or central vestibular structures in the brain stem. In this subsection, we focus on two distinct disorders, each with characteristic clinical features that present with vertigo: benign paroxysmal positional vertigo (BPPV) and acute vestibular syndrome (AVS).

BPPV is a common mechanical disorder of the inner ear in which certain head movements precipitate vertigo; nystagmus, nausea, and vomiting commonly accompany the vertigo. Although patients may have quite dramatic findings with severe symptoms, the actual episodes of vertigo are extremely brief and typically last less than 1 minute. In contrast, AVS refers to a separate group of disorders characterized by an acute onset of vertigo, nausea, vomiting, and gait instability that lasts for days to weeks.

Dix-Hallpike Test
Background
In BPPV, it is thought that small crystals of calcium carbonate, or canaliths, are displaced into the endolymph of the posterior semicircular canal. Head movement shifts the canaliths, which induce bidirectional forces in the fluid that create the sensation of spinning and trigger the attack of BPPV.11–14 The posterior semicircular canal is most commonly affected,13–15 but at times the lateral canal is thought to be involved, thereby leading to variants of typical BPPV.16–18

Indications and Contraindications
The Dix-Hallpike test may be used to confirm the diagnosis of BPPV by provoking a specific type of nystagmus. The maneuver should not be performed on patients with severe cervical spine disease, unstable spinal injury, high-grade carotid stenosis, or unstable heart disease.11 Patient discomfort and physical infirmity are relative contraindications; some elderly
The Dix-Hallpike test may be useful in confirming the diagnosis of BPPV and in localizing the affected ear. Accurate identification of BPPV is of interest because canalith-repositioning maneuvers may be offered as a therapeutic option.

Canalith-Repositioning Maneuvers

**Background**

If the clinical evaluation of a patient with vertigo is consistent with the diagnosis of BPPV, the patient may be a candidate for canalith-repositioning maneuvers. Based on the theory that stray material in the posterior semicircular canal causes the symptoms of BPPV, maneuvers were designed that involved sequential head movements to reposition the debris.34,35,40,41 Manipulation of head position theoretically allows the debris (canaliths) to sequentially fall from the problematic location in the semicircular canal into the vestibule where they can adhere. The most commonly recommended maneuvers are the Epley and Semont maneuvers.34,41 Several randomized studies have compared the Epley maneuver with a sham maneuver, reporting success rates of 50% to 90%.42–45 One large study of 965 patients with BPPV found a single canalith-repositioning maneuver improved symptoms in 85% of patients.46 Both the Epley and Semont maneuver appear to have similar efficacy (reported in the range of 90% in one small study),47 although the Semont is less well studied.48

**Indications and Contraindications**

The indication for canalith-repositioning maneuvers is the clinical diagnosis of BPPV as confirmed by the history and physical examination, including the Dix-Hallpike maneuver. Careful patient selection is key to the success of these maneuvers, as well as correct identification of the impaired ear, which

**Procedure**

The Dix-Hallpike maneuver is illustrated in Fig. 61.3 and Video 61.1. Place the patient initially in the upright seated position on the stretcher with the head turned 45 degrees to one side. Instruct the patient to keep the eyes open and focused on the examiner, then quickly lay the patient down flat so the head hangs over the edge of the bed and observe the eyes for nystagmus. Repeat the entire maneuver with the head turned 45 degrees toward the opposite side.35

**Interpretation**

The typical positive response in patients with BPPV is vertigo that develops after a brief delay (1 to 10 seconds), lasting less than a minute, and with rotatory nystagmus (where the eyes rotate about the visual axis). Nausea or other systemic symptoms are often present.35,39 The eye movements are mixed rotatory and vertical nystagmus of both eyes with the upper pole of the eye beating toward the dependent ear and the vertical nystagmus beating toward the forehead. The side with the ear in the downward position during the Dix-Hallpike test that elicits greater nystagmus usually identifies the affected ear. If the positional nystagmus is atypical, or if the maneuver fails to elicit nystagmus in a patient with ongoing symptoms of vertigo, another diagnosis should be considered.35

**Complications**

The maneuver may precipitate brief, intense vertigo and nausea in patients with BPPV.

**Summary**

The Dix-Hallpike test may be useful in confirming the diagnosis of BPPV and in localizing the affected ear. Accurate identification of BPPV is of interest because canalith-repositioning maneuvers may be offered as a therapeutic option.
will dictate the initial position and movement of the patient. Contraindications are the same as for the Dix-Hallpike maneuver, including severe cervical spine disease, unstable spinal injury, and high-grade carotid stenosis.

**Procedure**

**Epley Maneuver**

The Epley procedure is illustrated in Fig. 61.4 and Video 61.2. Place the patient initially in the seated position on the stretcher with the head turned 45 degrees toward the affected side. Lay the patient down flat with the head hanging over the edge of the bed. After 20 seconds or after the symptoms subside, rotate the patient's head so that it faces the opposite shoulder while maintaining the head-hanging orientation. Wait another 20 seconds or for symptom resolution and then roll the patient further onto the side and rotate the head further into a face-down position. Again, after 20 seconds or after any symptoms subside, return the patient to a seated position.34

Some authors suggest keeping the patient's head in each position long enough for any provoked symptoms of nystagmus or vertigo to resolve.35 Others suggest a period of 3 to 4 minutes in each position.31,47,49 The maneuver may be repeated a few times until some improvement in symptoms occurs.47,50 After a successful procedure, advise the patient to remain in a head-upright position for 24 hours.

**Semont Maneuver**

The Semont maneuver involves larger and more abrupt body movements (Fig. 61.5). With the patient seated on the side of an examination table or bed, turn the patient's head 45 degrees toward the right ear and the neck slightly extended with the chin pointed slightly upward. This position results in the patient's head hanging to the right.

First, a Dix-Hallpike test is performed with the patient's head rotated 45 degrees toward the right ear and the neck slightly extended with the chin pointed slightly upward. This position results in the patient's head hanging to the right.

Then the head and body are further rotated until the head is face down. The vertex of the head is kept tilted downward throughout rotation. The maneuver usually provokes brief vertigo. The patient should be kept in the final, face-down position for about 10 to 15 seconds.

Once the vertigo and nystagmus provoked by the Dix-Hallpike test cease, the patient's head is rotated about the rostral-caudal body axis until the left ear is down.

With the head kept turned toward the left shoulder, the patient is brought into the seated position. Once the patient is upright, the head is tilted so that the chin is pointed slightly downward.

**Figure 61.4** The Epley procedure, a bedside maneuver for the treatment of a patient with benign paroxysmal positional vertigo affecting the right ear. The presumed position of debris within the labyrinth during the maneuver is shown in each panel. (Adapted from Furman C: Benign paroxysmal positional vertigo, *N Engl J Med* 341:1590, 1999.)
toward the unaffected side; the head will maintain this position throughout the entire maneuver. Quickly lay the patient down into a side-lying position onto the affected side, keeping the head turned 45 degrees so the patient’s head is directed partially towards the ceiling. Once the symptoms subside, move the patient abruptly through the sitting position to the opposite side-lying position, maintaining the head position so that now the head faces down partially towards the floor. Keep the patient there until the symptoms subside. Return the patient to the upright position.47,49 Advise the patient to remain in a head-upright position for 24 hours after a successful procedure.

**Complications**

Nausea is reported in 16% to 32% of patients during the Epley maneuver.48 Exacerbation of vertigo occurs occasionally and is thought to result from displacement or dislodgment of canal debris. Repeating the procedure is recommended for relief. Symptoms of vertigo may recur in as many as 50% of patients, including 20% during the first 2 weeks.31,48

**Summary**

In patients with correctly identified BPPV, canalith repositioning may bring immediate relief or improvement of symptoms. Evidence of success in the ED setting is limited but growing.51

**HINTS Examination**

**Background**

Whereas the majority of AVS conditions are the result of a peripheral neuropathy (such as acute labyrinthitis or vestibular neuritis), some AVS presentations (as many as 25% in some series) are due to a central pathology such as a brain stem or cerebellar stroke.52 It can be very difficult to distinguish a peripheral from a central process in patients with AVS using the standard neurologic examination. The “HINTS” examination was created for this very purpose and is intended to assist providers in distinguishing central from peripheral causes of AVS. The HINTS exam is composed of three separate tests: the horizontal Head Impulse test, a test of Nystagmus, and a Test of Skew (HINTS).53

**Indications and Contraindications**

Each of the three components of the HINTS examination has separate contraindications. The horizontal head impulse test should not be performed in patients with severe cervical spine disease, unstable spinal injury, high-grade carotid stenosis, or unstable heart disease. Patient discomfort is also a relative contraindication. If vertical or bidirectional nystagmus is present at rest, or if vertical skew is present at baseline, a central cause of vertigo should be suspected and a full HINTS examination is not indicated.

**Procedure**

**Horizontal Head Impulse Test (Video 61.3)**

Have the patient seated in a comfortable position and stand in front of the patient. Ask the patient to gaze directly on the examiner’s eyes at all times and relax the neck as much as possible. Place hands on either side of the patient’s head and rapidly rotate the head by approximately 20 degrees to each side and back to midline several times while carefully observing the eyes for loss of fixation on the central target of the examiner’s eyes. If the eyes fall off target when the head is rotated laterally to the affected side, a corrective saccade to resume
Interpretation

Horizontal Head Impulse Test

The terminology used to describe the results of the horizontal head impulse test can be confusing, as the test was initially designed to detect peripheral vestibular disease. Therefore, a positive (abnormal) test is suggestive of a peripheral lesion, whereas a negative (normal) test is consistent with a more concerning central lesion. In the setting of AVS symptoms, the inability to maintain fixation during the horizontal head impulse test suggests a peripheral vestibular lesion. This is presumed to occur because there is no afferent input from the affected nerve when the head is rotated to the affected side. If a patient with AVS symptoms maintains fixation on the examiner during the horizontal head impulse test, it suggests a central pathology. In rare instances, patients with lateral pontine and cerebellar stroke syndromes can have a positive horizontal head impulse test that erroneously suggests a peripheral lesion.

Visual fixation on the central target occurs; this represents an abnormal horizontal head impulse test and is suggestive of a peripheral lesion. Randomly alter the speed and direction of rotation to generate an unpredictable sequence so the patient cannot anticipate the direction of rotation. A normal and abnormal horizontal head impulse test maneuver is simulated in Fig. 61.6.

Test of Nystagmus

Ask the patient to fixate on the examiner’s finger and slowly track the finger laterally in both directions. Observe for nystagmus at the extremes of gaze.

Test of Skew

Ask the patient to fixate on an object directly ahead. Alternately cover each eye in rapid succession. Look closely for subtle vertical refixation of the eyes (i.e., the eye moves up or down as it is uncovered to compensate for misalignment).
Test of Nystagmus
The nystagmus that results from a peripheral lesion is unidirectional; its fast or corrective phase beats away from the affected side when looking away from the affected side. With gaze toward the affected side, the nystagmus may decrease in intensity or disappear, but the direction of the fast beats will not change. If the direction of the nystagmus differs with horizontal gaze to either side (e.g., left beating when looking left and right beating when looking right), a central cause should be suspected.36

Test of Skew
Skew deviation refers to a vertical misalignment of the eyes (e.g., one eye is looking straight ahead, while the other eye is deviated downward or upward); it is usually associated with a central lesion. Although skew deviation can be subtle, it can be unmasked by alternately covering each eye in rapid succession while the patient fixes her gaze on the examiner.37

HINTS Exam
If any one of the components of the HINTS examination suggests a central cause, the composite HINTS examination is suggestive of a central lesion. Several studies have examined the diagnostic accuracy of the HINTS exam for identifying central causes of AVS.53,58–60 In one study of 101 patients with chronic risk factors for cerebrovascular disease and new onset AVS, the HINTS examination was 100% sensitive (95% confidence interval [CI] 95%–100%) and 96% specific (95% CI 80%–99%) for a central cause. A subsequent study by the same group and two other smaller studies have reported sensitivities of 88% to 100% and specificities of 85% to 99%.58,59,61 Of note, the studies supporting the diagnostic accuracy of the HINTS examination have been conducted in patients with a moderate to high risk of stroke, where the examination was performed by experienced neuro-otologists53,58–60; test performance in the ED setting by emergency clinicians is not known.

Complications
The horizontal head impulse test may exacerbate injury in improperly selected patients with unstable cervical spinal injury or, theoretically, with severe carotid atherosclerotic disease, although no reports of such complications have been reported.

Summary
The HINTS examination may be useful in identifying patients with a central cause of vertigo. Although early investigations appear to confirm its role in differentiating central from peripheral vertigo, it should be used with caution and in conjunction with other diagnostic data. Further studies are needed to confirm the reliability of these tests in the ED setting performed by emergency clinicians.

EVALUATION OF WEAKNESS

Background
Myasthenia gravis (MG) is the most common disease of neuromuscular transmission, with an annual incidence of seven to nine new cases per million.62 Patients with MG may be grouped into two major categories. The first group shows weakness in the proximal muscles that increases with activity and improves with rest. The second group has ocular complaints of diplopia or ptosis. Patients in the ocular weakness group may or may not have generalized symptoms as well. Fatigue is the hallmark of the disease; the symptoms typically wax and wane.

Under normal circumstances, when the nerve terminal is stimulated, acetylcholine (Ach) is released and diffuses across the synaptic cleft to the postsynaptic receptors. Acetylcholinesterase, located deep within the synaptic folds, hydrolyzes acetylcholine. In myasthenia gravis there are a reduced number of postsynaptic acetylcholine receptors, which can lead to failed neuromuscular transmission and weakness.

Figure 61.7 Neuromuscular junctions. Acetylcholine is released from presynaptic vesicles and diffuses across the synaptic cleft to the postsynaptic receptors. Acetylcholinesterase, located deep within the synaptic folds, hydrolyzes acetylcholine. In myasthenia gravis there are a reduced number of postsynaptic acetylcholine receptors, which can lead to failed neuromuscular transmission and weakness.
proprietary preparation), due to the drug’s rapid onset and short duration of action, and the ice-pack test, because it is safe and noninvasive.66,70,71

**Edrophonium (Tensilon) Test**

**Background**

Edrophonium chloride is an acetylcholinesterase inhibitor that has been used for the diagnosis of MG since the 1960s. The short duration of action of edrophonium that made it unsatisfactory as a therapeutic agent for MG makes it useful as a diagnostic agent. The drug’s onset of action is rapid (30–45 seconds), and the duration of maximal effect is short (usually less than 2 minutes). Any effect resolves within 5 to 10 minutes.72

**Indications and Contraindications**

The bedside Tensilon test is indicated for the diagnosis of patients with suspected MG when there is a clinical need to make this diagnosis in the ED. If other bedside tests, such as the ice-pack test (discussed later), are conclusive, or if eyelid fatigue on prolonged upgaze can be demonstrated, there is no need to administer edrophonium.73 If the patient will be admitted to an inpatient setting where timely electromyelograms or serologic testing can be performed, it is reasonable to defer the Tensilon test. Although the Tensilon test was once touted as a way to distinguish myasthenic crisis from cholinergic crisis, it is not reliable and is not recommended.74

A muscle that is clearly weak must be identified to monitor during Tensilon testing; some advocate only performing the test in patients with ptosis or ophthalmoparesis, as quantifying strength in other muscle groups independent of effort makes the test too unreliable.75,76

Administration of edrophonium potentiates the muscarinic effects of ACh and is not entirely benign, particularly in patients with cardiac disease or asthma, which are relative contraindications to the test. Patients often have increased salivation and mild gastrointestinal cramping, but serious complications (e.g., symptomatic bradycardia and syncope) are rare.75–77 One study reported the complication rate in office settings by neuro-ophthalmologists to be 0.2%.77 There is one report of transient asystole after the administration of edrophonium in a patient with suspected MG; however, the patient was critically ill and had been receiving intravenous labetalol, which may have exacerbated the effect of the edrophonium.77 Hence, caution is advised in administering edrophonium to patients receiving β-blocking agents, digoxin, or other drugs with atrioventricular-blocking properties.79–82

**Equipment**

Intravenous access should be secured with a saline lock. Ten milligrams of edrophonium chloride (Tensilon, ICN Pharmaceuticals) should be drawn up in a tuberculin syringe. Edrophonium chloride is supplied in 1-mL and 10-mL vials at a concentration of 10 mg/mL. A second syringe of normal saline should be available to administer as a double-blind placebo, although some clinicians have recommended nicotine, calcium chloride, or atropine for this purpose.83 Atropine and other cardiovascular drugs and resuscitative equipment should be readily available. It may be useful to prepare a syringe with 0.5 mg of atropine prior to the test should bradycardia or bronchospasm occur. Cardiac monitoring is recommended.75 Photographic recording equipment may help objectively document any improvement in motor function.

**Procedure (Video 61.4)**

Identify a muscle that is clearly weak. Obvious ptosis or ophthalmoparesis allows direct observation of a single weak muscle becoming stronger in response to the drug. Ideally, one person is available to administer the edrophonium or placebo and a second person is free to observe the effect of medication on the patient. Neither the observer nor the patient should know which syringe contains edrophonium and which contains placebo, thus creating a double-blind testing situation.

Again, ptosis is a reliable sign to test and is generally used if present (Fig. 61.8). The principles involved in assessing the effect of edrophonium on ptosis may be extended to testing other muscles. Ask the patient to look upward for several moments to fatigue the levator muscles. Note the degree of ptosis and document it by measurements or photographs. After a moment’s rest, ask the patient to look straight ahead. Inject 0.2 mL of the test substance in one syringe (2 mg of edrophonium or saline). If no response is seen within 1 minute, inject further increments of 2 mg up to a maximum of 10 mg. Note any increase in strength as reflected by an increase in size of the distance between the upper and lower lids (the palpebral fissure).

Incremental administration of edrophonium has several benefits. Many patients may respond to doses less than 10 mg and will be at less risk for muscarinic side effects at lower doses. Additionally, some patients may have improved strength.

![Figure 61.8](image-url) **Figure 61.8** Tensilon test. A, Bilateral ptosis (right worse than left) and defective upgaze (right worse than left). B, Improvement of ptosis and left upgaze following the intravenous injection of edrophonium chloride (Tensilon). (From Kanski JJ: Clinical diagnosis in ophthalmology. St. Louis, 2006, Mosby.)
of extraocular muscles at lower doses that paradoxically worsens at higher doses.\textsuperscript{73} Giving edrophonium in 2-mg aliquots will allow these patients to be properly identified as having a response. Repeat the procedure with the other test substance. Any improvement should occur within 30 seconds and disappear after 5 minutes.

**Complications**

A small percentage of individuals are hypersensitive to the initial small dose of edrophonium and exhibit the cholinergic side effects of salivation, lacrimation, and miosis. These effects are transient. Atropine, 0.5 mg, may be given intravenously, if necessary, to counteract these symptoms. A smaller number of patients may experience symptomatic bradycardia that responds to atropine. As described earlier, rare cardiac arrhythmias and death have been reported in patients taking digoxin or β-blockers.

**Interpretation**

Objective signs of improvement in the strength of an identified paretic muscle within 30 seconds of administration of edrophonium and fading of that improvement over the next 5 minutes are the criteria for a confirmatory test result. Subjective increases in general strength or relief of fatigue do not constitute a positive test. Normal individuals should have no change in muscle strength, but may transiently experience the side effects of salivation, lacrimation, and diaphoresis. Perioral, periocular, or lingual fasciculations are almost always noted in normal patients after edrophonium administration.\textsuperscript{72} The Tensilon test may be repeated in 30 minutes if desired.

The sensitivity of the edrophonium test for MG is estimated to be approximately 80% to 90%.\textsuperscript{75,83–85} The specificity is difficult to estimate, but false-positive tests have been reported in patients with Eaton-Lambert syndrome, intracranial lesions, amyotrophic lateral sclerosis, botulism, and Guillan-Barré syndrome.\textsuperscript{75,86–88}

**Ice-Pack Test**

**Background**

It has been observed clinically that myasthenic patients have exacerbations of weakness with environmental heat and improvement in strength with cold temperatures, possibly because lower temperatures inhibit acetylcholinesterase function.\textsuperscript{71} A simple bedside test uses these observations to evaluate ptosis.\textsuperscript{66,70,71} An ice pack is placed lightly over the closed eyelid for 2 minutes. Ptosis has been noted to improve in 80% or more of patients with ocular MG and may be more sensitive than the edrophonium test.\textsuperscript{71,89} Although the reported number of patients evaluated by this method remains small, the test is included here because its safety, speed, and ease of administration make it an attractive ED application.

**Indications**

Unilateral or bilateral ptosis of uncertain etiology in patients being evaluated for MG is the sole indication for this test.

**Procedure**

Ice and a bag (or a surgical glove) is the only equipment required. A camera to record a response is optional. Measure or photograph the degree of the patient's ptosis prior to the test. Studies have used the vertical distance between the upper and lower lid, or palpebral fissure, as an objective measure of the degree of ptosis.\textsuperscript{89} If bilateral ptosis is present, test the more affected eye. Fill a bag (or surgical glove) with ice and place gently on the patient's closed eyelid for 2 minutes or until patient discomfort limits application. After removing the ice, immediately assess the width of the palpebral fissure and compare with the pretest width (Fig. 61.9).

![Image of ice-pack test](image-url)
**Complications**
Patient discomfort from ice pack application may limit cold exposure time to less than 2 minutes, but may still allow a test to be successfully performed.

**Interpretation**
A clear improvement in ptosis in the cooled eye constitutes a positive test. Studies have defined 2 mm or more of improvement in the width of the palpebral fissure as a positive test. The effect should be reproducible. In small clinical studies the ice-pack test was found to be at least as sensitive as administration of edrophonium in improving ptosis in patients with ocular MG. False-negative results do occur, more commonly in patients with complete ptosis, and probably at approximately the same frequency as with Tensilon testing. Normal individuals show no change in width of the palpebral fissure after cold exposure. False-positive results are rare.

**Summary**
MG is typically diagnosed by serologic testing for autoantibodies and electrodiagnostic studies in the ambulatory setting. However, when diagnosis in the ED is desired, the Tensilon test and ice-pack test may provide valuable information at the bedside. False-negative results do occur, and additional testing should be performed if clinical suspicion for MG persists.

**ACKNOWLEDGMENTS**
The author would like to thank J. Stephen Huff, MD and Paul B. Baker, MD for their significant contributions to this chapter in prior editions.
The following discussion focuses on procedures performed by emergency clinicians during the evaluation and treatment of injuries and diseases of the eye. Emphasis is placed on practical application of the techniques; cautions to be heeded by the emergency clinician are included. Not all procedures are mandated to be performed by the emergency clinician, and if beyond the expertise available in the ED, they may be referred to a specialist.

**ASSESSMENT OF VISUAL ACUITY**

Evaluation of visual acuity may initially be deferred with simple, obvious, or straightforward cases, such as a stye, periorbital laceration, or minor eye irritation; however, assessment of visual acuity should be the first procedure performed in the majority of patients seen in the emergency department (ED) with an eye complaint (Videos 62.1 and 62.2). Even though it may initially be deferred in the triage or trauma room setting or under other relevant scenarios, the emergency clinician must ensure that visual acuity or function is assessed adequately as part of the initial evaluation.

**Indications**

Visual acuity should be assessed as soon as practicable and before the patient is examined with bright lights. In the event of blepharospasm from an injury (e.g., abrasion, chemical exposure), a topical anesthetic may facilitate the examination. Patients with eye complaints often say that they “can’t see.” In these instances, emergency visual acuity assessment should be performed first, beginning with evaluation of light perception, then hand motion, and finally counting fingers at 3 ft (Fig. 62.1). If the patient succeeds in performing these assessments, a near vision card may then be used or distant visual acuity assessed. In emergency circumstances, detailed formal vision testing is not essential; however, some form of visual acuity assessment is needed. In this situation, the ability to count fingers or read newsprint gives some indication of gross visual function. Formal visual acuity testing should never delay critical therapeutic interventions such as eye irrigation in the case of eye exposures.1

**Distant Visual Acuity Procedure**

For formal vision testing, ask the patient to face a well-lit standard Snellen or similar eye chart from a premeasured distance of 20 ft. Use a card or the palm of the hand to occlude one eye at a time. If possible, examine all patients while wearing their current lens correction to obtain the best corrected distant visual acuity. If not available, measure visual acuity first without correction and then with a pinhole device, and note any improvement in visual acuity. The pinhole device functions as a corrective lens by eliminating divergent light rays and allowing light only through the center of the lens, thus reducing corneal refractive error. In general, visual acuity is improved with the pinhole device. Decreased visual acuity that is not improved with this device suggests that corneal refractive error is not the cause. Construct a pinhole device by punching one to several holes in the center of a card (3 × 5 inch index card) with an 18-gauge needle. Devices with one or more pinholes drilled into an eye cover are available commercially (Fig. 62.2A and B) or can be constructed from readily available material in the ED, such as the index card with holes just mentioned (Fig. 62.3). Fig. 62.2C presents a chart for testing visual acuity while the patient is on a stretcher or in a chair. The chart can be used directly from this text if held 14 inches from the eye. Begin by testing the affected eye or the one presumed to have the worst visual acuity. First, instruct the patient to read the smallest letters on the chart that can easily be seen. Then ask the patient to read letters that can just barely be made out (i.e., they do not have to be clear). If the patient is unable to read the largest letter on the chart, move the patient to half the distance from the chart (10 ft), or if using the figure in this text, move it 7 inches closer to the eye and repeat the procedure. Record the results reflecting the change in distance (e.g., 10/200). The numerator in the vision ratio is the distance of the patient from the chart and the denominator is the distance at which a patient with normal vision can read the line of letters. For patients who still cannot read the letters on the chart, test vision progressively as follows: the ability to count fingers, detect hand motion, and perceive light (with or without projection; i.e., the ability to perceive the direction of light), and finally, the inability to perceive light.2

**Near Visual Acuity Procedure**

Perform near visual acuity assessment in the ED at the bedside or at triage. Hold a pocket near vision card (see Fig. 62.2C)
or any printed material in good light at a distance of approximately 14 inches in front of the patient and occlude each eye alternately as described earlier. Alternately, an “app” may be used from a smart phone device, but directions vary and not all apps measure near distance.1 When using available printed material in lieu of a near vision card, measure the size of the letters that are discerned by the patient. At a later time compare these letters with the size of the letters on the near vision card to deduce the patient’s actual visual acuity. When near vision is decreased, it is usually caused by either loss of visual function or poor accommodation as a result of advancing age (presbyopia). Less commonly, it is caused by traumatic mydriasis. Thus, examine patients with presbyopia with their reading correction in place to obtain the best corrected near visual acuity.

For patients who cannot communicate or in whom factitious blindness or malingering is suspected, check for optokinetic nystagmus (OKN) to determine whether the visual pathway is intact. To test for OKN, pass a regularly sequenced pattern in front of the eyes. If an optokinetic drum is available, rotate the drum in front of the patient (Fig. 62.4). This is not available in many EDs, however multiple apps are available for use with an iPhone or smart phone which show a moving stripe which elicits nystagmus. Some apps have the ability to record the nystagmus in a video format (Optokinetic Drum Provision 1.0, Bluestone Publishing, Inc, updated Sept 18, 2015). In place of the drum, substitute a printed piece of paper such as newsprint (without photographs or large areas with no print) or a standard tape measure. Pass it in front of the patient’s eyes at reading distance while instructing the patient to look at it as it moves rapidly by. Evaluate for tracking as demonstrated by involuntary nystagmus-like eye movements seen when the test object is moved from side to side in front of the patient. Such movement indicates an intact visual pathway. Finally, another effective method is to hold a mirror in front of the patient and slowly rotate the mirror to either side of the patient. Patients with an intact visual pathway will maintain eye contact with themselves as demonstrated by eye movement as the mirror is moved. A large mirror that reflects the patient’s entire face is most effective for this purpose. All patients with decreased visual acuity from baseline require routine referral for further ophthalmologic follow-up; however, patients with moderately or severely decreased visual acuity not explained by refractive error require ophthalmologic consultation in the ED.

**USE OF OPHTHALMIC ANESTHETIC AND ANALGESIC AGENTS**

Topical anesthetic agents are widely used in the ED setting. Their use is essential in achieving adequate pain control and to evaluate and treat painful eye conditions. Two agents most commonly used are tetracaine and proparacaine. Comparative studies suggest that proparacaine is less painful on instillation, whereas tetracaine is longer lasting, and thus possibly more effective.4,5 Topical anesthetic agents have been used in the outpatient setting following ophthalmic surgery and are considered safe and effective for this condition. Although still controversial, there is a growing body of evidence that supports their use for brief outpatient treatment (a few days only) of corneal abrasions.6,7

Ophthalmic nonsteroidal antiinflammatory drugs (NSAIDs) have been evaluated for their effectiveness and in the treatment of traumatic corneal abrasions. Examples include ketorolac tromethamine, diclofenac, and flurbiprofen. These agents are thought to be safe to use and effective for the relief of pain associated with corneal abrasions.8-11 Topical ophthalmic NSAIDs have also been shown to be safe and effective when used for the treatment of corneal abrasions in conjunction with bandage contact lenses.12 Despite similar potential complications (delayed corneal healing) topical NSAIDs have gained widespread acceptance compared to topical anesthetic agents in their use for treatment of corneal abrasions.

**Indications and Contraindications**

Application of topical anesthetic agents can be both diagnostic and therapeutic. Relief of discomfort with a topical anesthetic strongly suggests, but does not ensure, a conjunctival or corneal injury. An ocular irritant may also be masked by the use of these agents. Classic teaching is that patients should not self-administer anesthetic preparations. It is thought that they delay wound healing by disrupting surface microvilli and causing a decrease in the tear film layer and tear break-up time.13 Although self-administered topical anesthetic agents are now routinely used after photorefractive keratectomy for the first 3 or 4 postoperative days, this likely safe technique has not yet become a common ED practice.14

As evident from Table 62.1, the anesthetic solutions commonly used have a duration of action of less than 20 minutes, though tetracaine may have a slightly longer clinical effect.4 Patients with a large corneal lesion may need a more extended period of pain relief and may require bed rest, opioid analgesics, and/or appropriate sedatives.

A final word of caution should be added regarding the use of ophthalmic solutions. Guaiac solutions are commonly supplied in dropper bottles similar in size and appearance to those containing ophthalmic solutions. Well-intentioned ED personnel may store the guaiac reagent bottles with the ophthalmic bottles. One should encourage both color coding of the bottles and examination of them and their labels before each use to avoid corneal injury from inadvertently instilling guaiac reagent into the eye.

**Procedure**

Instillation of anesthetic or analgesic agents is similar to the administration of other eye solutions as described later. Forewarn the patient that the medication will sting transiently.
Figure 62.2 A commercial pinhole device reveals refractive error caused by corneal aberration (excessive tearing or nearsightedness). The pinhole device functions as a corrective lens by eliminating divergent light rays and allowing light only through the center of the lens, thus reducing corneal refractive error. In general, visual acuity is improved with the pinhole device. Decreased visual acuity that is not improved with this device suggests that corneal refractive error is not the cause. A, First measure visual acuity without the device. B, Measure again with the patient looking through the pinhole. Document the acuity with and without the pinhole device. C, If the patient cannot stand or a formal eye chart is not available, ask the patient to read this “distance equivalent” chart by holding the card 14 inches away from the patient.
on instillation but will quickly offer relief. Repeat instillation in one minute to ensure maximum effect. Instruct the patient not to rub the insensate eyes to avoid causing or worsening an abrasion. Provide the patient material to blot the eyes gently, if needed, as tearing occurs.

Complications

Open bottles can harbor bacteria and introduce infection. Out-of-date medication may not be effective and result in inadequate anesthesia. The absence of protective reflexes while the patient is under the effect of the medication may encourage use of the eye and result in further corneal injury from the foreign body (FB) or corneal infection.

A systemic review of the safety of topical anesthetics in the treatment of corneal abrasions found corneal erosions, edema, or iritis reported when agents were used for prolonged periods of time (up to 10 weeks), too often (up to every 15 minutes), and with higher concentrations. Delayed corneal healing has been long taught as a significant complication of topical anesthetic and analgesic agents, but several well-designed randomized clinical trials in the ED-based and ophthalmology (PRK surgery) literature report no adverse events from topical anesthetic use and corneal re-epithelialization occurring by 72 hours, indicating that any delay in healing does not appear to be clinically significant. Thus, though not currently used for routine outpatient treatment, topical anesthetic drops can be considered for use safely for 2 to 3 days without documented adverse effects in selected patients.

DILATING THE EYE

Dilating the eye is useful for both diagnostic and therapeutic purposes. Be advised, however, that an attack of narrow-angle (angle-closure) glaucoma may be precipitated by dilating the pupil. The most common form of glaucoma, however, is open-angle glaucoma, and this type is not precipitated by dilating the pupil. Some patients may have a “mixed mechanism” glaucoma with both open-angle and narrow-angle components. Systemic reactions, such as bradycardia or even heart block from β-blocker eye drops, can be induced by mucosal absorption of dilating medications.

![Figure 62.3](image1.png) A bedside pinhole device can be easily fashioned if commercial devices are not available. In this image, a single hole was crafted from a needle and an index card. Multiple holes work effectively as well. When assessing visual acuity with a handmade pinhole device, have the patient hold the device in front of the eye tested, while covering the other eye. (Courtesy Mary Jo Chandler, PA-C.)

![Figure 62.4](image2.png) Optokinetic nystagmus (OKN) testing will determine whether the visual pathway is intact. It will be normal in the patient with factitious blindness or malingering. Induce OKN by passing a regularly sequenced pattern in front of the eye such as this commercially available drum. Hold the drum in front of the patient. Direct the patient to look at the drum as you rotate it slowly. Alternatively, draw a tape measure across the line of sight while asking the patient to look directly at it as it passes.

### TABLE 62.1 Ophthalmic Anesthetic Agents

<table>
<thead>
<tr>
<th>GENERIC NAME</th>
<th>TRADE NAME</th>
<th>CONCENTRATION (%)</th>
<th>ONSET OF ANESTHESIA</th>
<th>DURATION OF ANESTHESIA (min)</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tetracaine</td>
<td>Pontocaine</td>
<td>0.5–1.0</td>
<td>&lt; 1 min</td>
<td>15–20</td>
<td>Marked stinging; ointment and preservative free unit dose available</td>
</tr>
<tr>
<td>Proparacaine</td>
<td>Ophthaine, Ophthetic</td>
<td>0.5</td>
<td>&lt; 20 sec</td>
<td>10–15</td>
<td>Least irritating; no cross-sensitization with other agents</td>
</tr>
<tr>
<td>Benoxinate</td>
<td>Dorsacaine</td>
<td>0.4</td>
<td>1–2 min</td>
<td>10–15</td>
<td>Contains fluorescein in solution</td>
</tr>
</tbody>
</table>
There are two types of dilators: sympathomimetic agents, which stimulate the dilator muscle of the iris, and cycloplegic agents, which block the parasympathetic stimulus that constricts the iris sphincter. Cycloplegic agents also block contraction of the ciliary muscles, which control focusing of the lens of the eye. This second effect of cycloplegic agents is beneficial in the therapeutic use of dilators for iritis.

Cycloplegic agents were used cosmetically as early as Galen’s time. Beginning in the early 1800s, extracts from the plants Hyoscyamus and belladonna were used in ophthalmology. Atropine was first isolated in 1833. Epinephrine was used on eyes in 1900 as the first sympathomimetic agent.

**Indications and Contraindications**

There are several diagnostic and therapeutic indications for dilating the pupil. Dilation is indicated for diagnosis when the fundus cannot be examined adequately through an undilated pupil. An elderly patient with miotic pupils and cataracts is an example of a patient in whom dilation may facilitate funduscopic examination. Dilation is therapeutically useful for many ophthalmic conditions, including inflammation of the eye. In the emergency setting, corneal injury with secondary traumatic iritis is a common example. Dilation helps the inflamed eye in two ways. First, it may help to prevent adhesions (synechiae) from forming between the iris and other ocular structures. Such adhesions eventually limit movement of the pupil and may precipitate glaucoma. Second, cycloplegic dilating agents relax the ciliary muscle spasm that often accompanies an inflamed eye and thus may reduce the pain associated with inflammation. Though traditionally used for these purposes, both benefits are largely theoretical with little formal evidence to support or refute their use in the ED.

Dilation is discouraged in patients with head injury who are at risk for herniation when it is necessary to monitor pupil findings. Dilation is contraindicated in the presence of narrow anterior chamber angles. Patients predisposed to having narrow angles may be unaware of this condition. Evaluate the depth of the anterior chamber before this procedure, and do not dilate the eye if there is any question of a narrow angle.

To estimate the depth of the anterior chamber, shine a penlight tangentially from the lateral side of the eye. When the depth of the anterior chamber is normal, uniform illumination of the iris is seen. However, when the iris has a forward convexity as in the case of a narrow anterior chamber, only a sector of iris is illuminated and there will be a shadow on the medial (nasal) side of the iris (Fig. 62.5). With a slit lamp, the depth of the anterior chamber angle can be assessed directly. The definitive test for assessing the anterior chamber angle is gonioscopy, in which the anterior chamber angle structures are viewed directly by means of a special mirrored contact lens and a slit lamp. Gonioscopy is not a technique normally performed by emergency clinicians.

Systemic effects can develop after the application of eye drops. Review the following sections on agents and complications before using these drugs in patients with compromised cardiovascular function.

**Agents**

Only two dilating agents are really needed in the ED. Phenylephrine (Neo-Synephrine, Hospira, Inc., Lake Forest, IL) 2.5%, a potent sympathomimetic, is used for diagnostic dilation of the pupil and visualization of the fundus. The drug is short acting, and because accommodation is not affected, the patient’s vision is not altered. Phenylephrine 10% should not be used routinely because it can be absorbed systemically and, in rare cases, has caused hypertensive crisis, myocardial infarction, and death.

For therapeutic cycloplegia in patients with iritis, 5% homatropine works well. Even though Table 62.2 indicates a maximum duration of 3 days, 24 hours is more common. Therefore 5% homatropine is a useful therapeutic agent for traumatic iritis. Atropine itself should not be used for traumatic

**TABLE 62.2 Mydriatic Agents**

<table>
<thead>
<tr>
<th>AGENT</th>
<th>MAXIMUM MYDRIASIS</th>
<th>DURATION OF MYDRIASIS</th>
<th>COMMON TRADE NAME</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sympathomimetics</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phenylephrine, 2.5%</td>
<td>20 min</td>
<td>3 hr</td>
<td>Neo-Synephrine, Hospira, Inc., Lake Forest, IL</td>
</tr>
<tr>
<td>Cocaine, 5% or 4%</td>
<td>20 min</td>
<td>2 hr</td>
<td>—</td>
</tr>
<tr>
<td><strong>Parasympatholytics (Cycloplegics)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atropine, 1%</td>
<td>40 min</td>
<td>12 days</td>
<td>—</td>
</tr>
<tr>
<td>Scopolamine, 0.25%</td>
<td>30 min</td>
<td>7 days</td>
<td>—</td>
</tr>
<tr>
<td>Homatropine, 5%</td>
<td>30 min</td>
<td>1–3 days</td>
<td>—</td>
</tr>
<tr>
<td>Cyclopentolate, 1%</td>
<td>30 min</td>
<td>6–24 hr</td>
<td>Cyclogyl, Alcon Laboratories, Inc., Fort Worth, TX</td>
</tr>
<tr>
<td>Tropicamide, 1%</td>
<td>30 min</td>
<td>4 hr</td>
<td>Mydriacyl, Alcon</td>
</tr>
</tbody>
</table>

*bottle caps of all these agents except cocaine are usually red.

*The duration of effect shows considerable individual variation. These are general estimates.

*Preferred for funduscopic examination.

*A 10% solution may produce cardiovascular reaction and hence should not be used.

*Preferred for iritis or corneal abrasion therapy.
Obstructed flow leads to increased posterior chamber pressure. Pupillary block (iris bombé).

NORMAL ANGLE- CLOSURE GLAUCOMA

**Figure 62.5** A, Left: The normal eye. Aqueous humor, produced in the posterior chamber, flows through the pupil into the anterior chamber. The major pathway for the egress of aqueous humor is through the trabecular meshwork into Schlemm’s canal. Right: Primary angle-closure glaucoma. In anatomically predisposed eyes, transient apposition of the iris at the pupillary margin to the lens during pupil dilation blocks the passage of aqueous humor from the posterior chamber to the anterior chamber. Buildup of pressure in the posterior chamber bows the iris forward and occludes the trabecular meshwork. B, Left: Normal anterior chamber with a negative transillumination test. Note that the entire iris is illuminated. Right: Shallow anterior chamber with a positive transillumination test. Note the shadow on the outer half of the iris. C, Left: Clinical use of the penlight examination to assess the depth of the anterior chamber of the right eye. The examiner sits face to face with the patient to ensure that the light source is perfectly perpendicular to the line of vision. Right: The axial anterior chamber depth of the right eye is less than normal. This can be demonstrated by shining a light across the eye from the temporal side. Because of the convexity of the iris-lens diaphragm, the nasal iris manifests a crescentic shadow (eclipse sign). (A, From Kumar V, Abbas AK, Fausto N, et al, editors: *Robbins and Cotran pathologic basis of disease, professional edition*, ed 8, Philadelphia, 2009, Saunders; B, from Swartz MH: *Textbook of physical diagnosis*, ed 6, Philadelphia, 2009, Saunders; C, from Kanski JJ: *Clinical diagnosis in ophthalmology*, St. Louis, 2006, Mosby.)
iritis because the undesirable effects of pupillary dilation and blurred vision persist for a week or longer after the associated corneal abrasions have healed. Atropine drops may be prescribed as part of the therapy for nontraumatic iritis after appropriate ophthalmologic consultation. Individuals with lightly pigmented irides tend to have greater sensitivity to cycloplegic agents than do individuals with greater pigmentation; the cycloplegic effect might therefore be more prolonged in people with light eyes. It might be difficult to dilate some patients with deeply pigmented irides, however, and numerous applications of drops might be required.

Malingers may use mydriatic agents to dilate a pupil unilaterally for the purpose of feigning neurologic disease. Normally, the pupillary dilation caused by intracranial compression of the third cranial nerve will constrict with 2% pilocarpine eye drops. A mydriatic-treated eye can be identified by full motor function of the third cranial nerve and absence of miosis after the instillation of pilocarpine. A fixed and dilated pupil in an awake and alert patient cannot be secondary to brain herniation. Although other neurologic problems may be present, in a normal-appearing patient with a fixed and dilated pupil, a pharmacologic cause is highly likely. It should be noted that legitimate patients may not recall the name of an eye medicine that they used but will usually recall whether the bottle had a red cap, as is found on all cycloplegic solutions marketed in the United States, though these agents are marketed in some countries (e.g., Greece, Turkey, India) with different colored caps. An unexpected mydriasis in a trusted patient may be the result of such an agent. Medications that constrict the pupil, such as pilocarpine, generally have a green cap. Pressure-lowering drops for glaucoma may be yellow or blue topped (β-blockers), purple topped (adrenergic agents), or orange topped (topical carbonic anhydrase inhibitors).

A fixed and dilated pupil from a pharmacologic cause may be encountered after both nasotracheal and orotracheal intubation (Fig. 62.6). In such ill or injured patients, cerebral herniation must be considered. When phenylephrine is used to constrict the nasal mucosa before nasal intubation (endotracheal tube, nasogastric tube), inadvertent spread to the eye can result in a fixed and dilated pupil. The same scenario may occur during resuscitation when endotracheal epinephrine has been instilled into the lungs and cardiopulmonary resuscitation has expelled epinephrine into the eye. In such scenarios, the affected pupil will not constrict after intraocular pilocarpine administration. Finally, a fixed and dilated pupil might occur as a result of inadvertent contamination of the eye with scopolamine after the application of a scopolamine patch.

Procedure

Instillation of mydriatic agents is similar to the administration of other eye solutions. For medicolegal purposes, note the patient’s visual acuity before instillation of the medicine. This documents that any decreased vision is not the result of the mydriatic agent. Whenever dilation is performed, note on the patient’s chart the dose and time that agents have been given to avoid confusion during subsequent neurologic evaluation.

Place the patient in a supine or a comfortable semirecumbent position. Instruct the patient to gaze at an object in the upper visual field, such as a fixture on the ceiling. Gently depress the lower lid with a finger on the epidermis. Instill a single drop of the solution into the lower lid fornix, and ask the patient to blink to spread the medication. Do not use more than a single drop because it produces reflex tearing and reduces the concentration in contact with the conjunctiva. Forewarn the patient that the medication is uncomfortable when it goes into the eyes. After the medication has been instilled, the patient may blot the eye when it is closed but should not rub it with a tissue. If the desired effect is not noted in 15 to 20 minutes, repeat the dose, but this is seldom required.

Complications

Any dilator can precipitate an attack of angle-closure glaucoma in susceptible patients. In the case of angle-closure glaucoma,
it may take several hours before symptoms become evident. The patient often complains of smoky vision with “halos” around lights, as well as an aching pain that at times can be quite severe. Nausea and vomiting may occur. If the affected eye becomes infected in association with a hazy cornea, elevated pressure on tonometry, and an oval, fixed pupil, consult an ophthalmologist immediately. Place the patient supine and administer analgesics and antiemetics. Treatment usually includes the agents suggested in Box 62.1, and later, definitive laser or surgical procedures.

Be aware that if contaminated, an eye medication can introduce infection. Most solutions contain bactericidal ingredients, but contamination of the tips of droppers can still occur (Fig. 62.7A).27 Use only newly opened bottles of eye medication or single-use vials, particularly if the patient has a deep corneal injury or has recently undergone eye surgery. Promptly discard out-of-date drops and those in which crust or other material is found around the nozzle.

Forewarn the patient that any cycloplegic (in contrast to a sympathomimetic) will blur a patient’s near vision. Vision will be less blurred in adults older than 45 years, who generally have a reduced ability to focus for near vision. If administering large amounts of eye drops that have systemic effects, such as β-blocker drops, place the operator’s index finger under the inferior eyelid along the nasal borders of the eye and firmly compress the nasolacrimal duct against the globe for a few minutes, thereby preventing migration of the drops into the nose and reducing systemic absorption.

**THE FLUORESCEIN EXAMINATION**

Perform fluorescein staining of the eye as part of the evaluation of patients with eye trauma and infection. It is a quick and easy technique that is crucial for the proper diagnosis and management of common eye emergencies. View the fluorescein-stained cornea and conjunctiva under a “blue” light and ideally in conjunction with slit lamp magnification (see later section on Slit Lamp Examination).

Sodium fluorescein is a water-soluble chemical that fluoresces. It absorbs light in the blue wavelengths and emits the energy in the longer green wavelengths. It fluoresces in an

---

**Figure 62.7** A, Administration of eye drops. Position the patient with the head tilted back. Direct the patient’s gaze upward. Pull the lower lid downward and instill a single drop of medicine in the lower conjunctival fornix. Ensure that the tip of the dropper does not touch the lid or lashes. Instruct the patient to close the eyelids for 1 minute to increase contact of the medicine with the globe and to decrease outflow of medication down the tear duct and over the lid margin. B, If administering large amounts of eye drops that have systemic effects, such as β-blocker drops, place the operator’s index finger under the inferior eyelid along the nasal borders of the eye and firmly compress the nasolacrimal duct against the globe for a few minutes, thereby preventing migration of the drops into the nose and reducing systemic absorption.

---

**BOX 62.1 Treatment Options for Acute Angle-Closure Glaucoma**

Place patient supine; administer analgesics (such as IV morphine titrated to effect) and antiemetics (such as ondansetron 8 mg IV):

1. β-Blocker*: Timolol 0.5% (Timoptic, Aton Pharma, Lawrenceville, NJ), 1 drop to the affected eye, wait 1 minute, and then
2. α-Agonist: Apraclonidine 1% (Iopidine, Alcon Laboratories, Inc., Fort Worth, TX), 1 drop to the affected eye, wait 1 minute, and then
3. Miotic agent: Pilocarpine 2% (Isoprop Carpine, Alcon), 1 drop to the affected eye every 15 minutes for 2 total doses, wait 1 minute after first dose, and then
4. Prednisolone acetate 1%, 1 drop to the affected eye every 15 minutes for 4 total doses
5. Acetazolamide 500 mg IV (may give by mouth, two 250-mg tablets, if IV medication not available)

*Empiric therapy, no controlled trials, based on clinical experience.

aRelative contraindications: severe bronchospasm, second- to third-degree heart block, uncompensated congestive heart failure.

alkaline environment (such as in Bowman’s membrane, which is located below the corneal epithelium) but not in an acidic environment (such as in the tear film over intact corneal epithelium). Thus it is useful in revealing even minute abrasions on the cornea.

Fluorescein was initially used in ophthalmology in the 1880s. It was first used as a drop, but when the danger of contamination by bacteria (especially Pseudomonas) was recognized in the 1950s, paper strips impregnated with fluorescein were developed. These strips are now supplied in individual sterile wrappers and should be used instead of the premixed solution.

Indications and Contraindications
Fluorescein staining is indicated for the evaluation of suspected abrasions, FBs, and infections of the eye, including “simple” cases of conjunctivitis, which may actually be herpetic keratitis. Exposure of the face to pepper spray has been associated with corneal abrasions, and such patients should undergo fluorescein staining and be evaluated with a slit lamp or Wood’s lamp. Conical defects may be seen after even a few seconds of unprotected viewing of a welder’s arc flame.

Fluorescein permanently stains soft contact lenses. Therefore when fluorescein is used, remove soft contact lenses before instilling the fluorescein and caution the patient to not put the lenses back into the eye for several hours. Topically administered fluorescein is considered nontoxic, although reactions to a fluorescein-containing solution (not impregnated strips) have been described. These reports, which consist of vage reactions and generalized convulsions, are rare, not rigorously supported, and believed to be caused by agents other than fluorescein in the solution. If using one of these fluorescein-containing solutions rather than the fluorescein-impregnated strips, beware of these potential, yet scientifically suspect idiosyncratic reactions.

In addition, beware that fluorescein dye may enter the anterior chamber of the eye in patients with deep corneal defects. This form of intracocular fluorescein accumulation is nontoxic. When the anterior chamber is viewed under the blue filter of the slit lamp, a fluorescein “flare” is visible and should not be confused with the flare reaction noted with iritis.

Procedure
Theoretically, one should not use topical anesthetics before fluorescein staining because a superficial punctate keratitis may develop in some patients from the anesthetic, which can confuse the diagnosis. However, in patients who are tearing profusely and squeezing their eyes shut from an abrasion or an FB, the examination is often impossible if a topical anesthetic is not first used. Thus, it is practical to apply a local anesthetic before instilling fluorescein.

Grasp the fluorescein strip by the non-orange end and wet the orange end with 1 drop of saline. Several convenient forms are available, including a small bottle of artificial tears or a 5-mL “bullet” or “fish” of normal saline commonly used for nebulizer treatments. Alternatively, wet the strip with tap water or the recently used local anesthetic drops. Once the strip is moistened, place it gently on the inside of the patient’s lower lid. Withdraw the strip and ask the patient to blink, which spreads the fluorescein over the surface of the eye. The key to a good examination is to have a thin layer of fluorescein over the corneal and conjunctival surfaces. If the strip is too heavily moistened before placing it in the lower fornix, the eye may become flooded with the solution, which makes evaluation difficult. If too much dye accumulates, the patient can remove the excess dye by blotting the closed eye with a tissue. Conversely, placing a dry strip in an unanesthetized eye may be irritating. Next, use a Wood’s lamp (4× magnification), the blue filter of a slit lamp, or simply a penlight with a blue filter to examine the eye in a darkened room (see Fig. 62.8, step 2). Check for areas of bright green fluorescence on the corneal and conjunctival surfaces. The naked eye may not be able to see small defects. Ideally, use a slit lamp with 10× or 25× magnification to examine the stained cornea before ruling out a pathologic process. A new handheld magnification device, the Eidolon Bluminator ophthalmic illuminator (Eidolon Optical, LLC of Natick, MA), produces an intense blue light from a light-emitting diode with 7× magnification (see Fig. 62.8, step 3). After completion of the fluorescein examination, irrigate excess dye from the eye to minimize damage to the patient’s clothing from dye-stained tears.

The Seidel test uses fluorescein to detect perforation of the eye. To perform this test, install a large amount of fluorescein onto the eye by profusely wetting the strip. Examine the eye for a small stream of fluid leaking from the globe (see Fig. 62.8, plate 4). This stream will fluoresce blue or green, in contrast to the orange appearance of the rest of the globe flooded with fluorescein.

Interpretation
Fluorescein is used mainly for evaluation of corneal injuries. Although conjunctival abrasions pick up the stain, most of the staining on the conjunctiva represents patches of mucus rather than a real pathologic condition. Corneal staining is more specific for injury, and the pattern of injury often reflects the original insult.

Corneal staining patterns are illustrated in Figs. 62.8 and 62.9. Abrasions usually occur in the central part of the cornea because of the limited protection of closure of the patient’s eyelids. The margins of the abrasions are usually sharp and linear if seen in the first 24 hours (see Fig. 62.8, plate 6). Circular defects are seen about embedded FBs and may persist for up to 48 hours after removal of a superficial foreign object. Deeply embedded objects may be associated with defects persisting for longer than 48 hours. Objects under the upper lid (including some chalazia) often produce vertical linear lesions on the upper surface of the cornea (see Fig. 62.8, plate 7). When vertical lesions are noted, search diligently for a retained FB under the upper lid. Overuse of hard contact lenses diminishes the nutrient supply to the cornea. The central part of the cornea sustains the most injury and thus fluoresces brightly when stained. Excessive exposure to ultraviolet light as a result of sunlamp abuse, snow blindness, or welding flashes produces a superficial punctate keratitis, which in its mildest form may not be visible without a slit lamp (see Fig. 62.8, plate 8). The central part of the cornea is the least protected by the lids, and a central, horizontal band–like keratitis can result. Herpetic lesions may develop anywhere on the cornea. Classically, these lesions are dendritic, although ulcers may also be punctate or stellate (see Fig. 62.8, plate 9). Any area of corneal staining with an infiltrate or opacification beneath or around the lesion should alert the practitioner to the possibility of a viral, bacterial, or fungal keratitis.
Fluorescein staining is a quick and easy diagnostic procedure that should be part of every eye evaluation. The extra time that the examination takes provides a wealth of diagnostic information on patients with eye trauma or infection. No complications are associated with the procedure with the exception of some dryness or irritation.

Obtain urgent ophthalmologic consultation so that cultures of the possible etiologic agents can be procured and appropriate treatment initiated.

Many Pseudomonas organisms fluoresce when exposed to ultraviolet light; therefore the presence of fluorescence before the instillation of fluorescein in a red eye should suggest the possibility of infection with this organism.

**Summary**

Fluorescein staining is a quick and easy diagnostic procedure that should be part of every eye evaluation. The extra time that the examination takes provides a wealth of diagnostic information on patients with eye trauma or infection. No complications are associated with the procedure with the exception of some dryness or irritation.
The crucial first step in the treatment of chemical injuries to the eye is irrigation. Irrigate as clinically appropriate for the severity, length of exposure, and causative agent. Serious chemical injury to the eye requires irrigation as soon as possible, even at the site where the exposure occurred, before the patient is brought to the ED. Corneal injury can occur within seconds even at the site where the exposure occurred, before the patient presents to the ED. Irrigation in the ED is indicated for all acute chemical injuries involving the eye.

**Indications and Contraindications**

Irrigation is indicated for all acute chemical injuries involving the eyes. Irrigation may also be therapeutic in patients with an FB sensation but no visible FB. Small, unseen foreign material in the conjunctival tissues may be flushed out with irrigation. There is no contraindication to eye irrigation, but in patients with a possible perforating injury, perform the irrigation especially gently and carefully.

**Equipment**

The following equipment is necessary for eye irrigation:

- Topical anesthetic, such as 0.5% proparacaine
- Sterile irrigating solution (warmed intravenous saline or lactated Ringer's [LR] solution in a bag with tubing)*
- A basin to catch the fluid
- Cotton-tipped applicators
- Gauze pads to help hold the patient’s lids open
- Lid retractors
- Irrigating device (e.g., Morgan Therapeutic Lens [MorTan Inc., Missoula, MT], modified central venous catheter, or Eye Irrigator) for prolonged irrigation
- Optimal: 10 mL of 1% lidocaine added to a liter of irrigating fluid

**Procedure**

**Basic Technique**

First, instill a topical anesthetic into the eye (Fig. 62.10, step 1). Evert the eyelid and sweep out any particulate matter in the conjunctival fornices with a moistened, cotton-tipped applicator (see later section on Ocular FB Removal and Fig. 62.10, step 2). Hold the eyelids open during irrigation (see Fig. 62.10, step 3). The easiest method is to use gauze pads to grasp the wet, slippery lids and hold them open. If the patient has severe blepharospasm, consider using lid retractors (Desmarres [Sklar Surgical Instruments, West Chester, PA] or paper clip retractors; Fig. 62.11). When lid retractors are used, be certain that the eye is well anesthetized, that the retractors do not injure the globe or the lids, and that chemicals are not harbored under the retractors. Be aware that simple retractors fashioned from metal paper clips (especially those that are nickel plated and shiny) may have surface chipping, which can create an ocular FB. Exercise caution to avoid ocular injury when using such a makeshift retractor.

Deutsch and Feller recommended an ipsilateral facial nerve block for severe blepharospasm (Fig. 62.12). To avoid swelling of periorbital tissue, block the facial nerve just anterior to the condylar process of the ipsilateral mandible. Place a subcutaneous line of anesthetic (2% lidocaine) to temporarily paralyze the orbicularis muscle.

Irrigate with normal saline or LR solution directed over the globe and into the upper and lower fornices (see Fig. 62.10, step 4). The choice of fluid initially is less important than initiating irrigation as rapidly as possible. If tap water is available at the site of the injury, begin irrigation immediately with copious amounts of fluid before transporting the patient to the hospital. Teach out-of-hospital care providers to irrigate all acid injuries of the eye for at least 5 minutes at the scene and to irrigate all alkali injuries for at least 15 minutes. LR or normal saline solution is preferred over tap water or 5%

*A balanced salt solution designed for eye irrigation is preferred by some (when available) and may produce less corneal edema with chemical injuries. Readily available normal saline and lactated Ringer's solution are equally well tolerated.
result in the development of a superficial punctate epithelial keratopathy.

Before irrigation, instill anesthetic eye drops, such as 0.5% tetracaine. Adding 10 mL of 1% lidocaine to a liter of irrigating fluid can decrease the patient's discomfort during prolonged irrigation.

**Duration of Irrigation**

Although Deutsch and Feller\(^3\) recommended that a full liter of irrigating solution be used in every case of caustic injury, the duration of irrigation is best determined by the extent of the exposure and the causative agent. Acids are quickly neutralized by proteins in the surface tissues of the eye and, once irrigated out, cause no further damage. The only exceptions are hydrofluoric and heavy metal acids, which can penetrate through the cornea. Alkalis can penetrate rapidly and, if not
In addition to litmus paper, the pH indicator on urine multi-indicator sticks can be used. The pH indicator on urine dipsticks is conveniently closest to the handle; all the distal indicator squares can be cut off with scissors. Normal tear film pH is 7.4. Use the noninjured eye as a control if the results are equivocal. If the pH measured in the conjunctival fornices is still abnormal after the initial irrigation, continue to irrigate. If the pH is normal after irrigation, wait 20 minutes and check it again to make sure that it remains normal, especially if alkaline contamination has occurred. Delayed changes in pH are usually the result of incomplete irrigation and inadequate swabbing of the fornices. In anticipation of this deficiency, measure the pH deep in the fornices. Consider double-lid eversion with a lid elevator to expose the upper fornix for swabbing, irrigation, and pH testing.

Prolonged Irrigation

Alkaline burns may require prolonged irrigation, and it is essential to consult ophthalmology in such cases. The Morgan Therapeutic Lens is a contact lens–type irrigation device that can provide slow, continuous irrigation once the more vigorous initial irrigation has been completed (Fig. 62.13 and Video 62.3). First, anesthetize the eye with topical anesthetic drops. Then place the device carefully on the surface of the eye with the lids closed around the intravenous tubing adaptor. Attach the intravenous tubing to the adaptor and provide continuous flow through the device onto the cornea and into the fornices. As the local anesthetic agents wash out during the irrigation process, the device can become uncomfortable, so reapply the anesthetic drops frequently during irrigation for patient comfort. Such short-term use of local anesthetics will not inhibit healing of the cornea.

removed, will continue to produce damage for days because of slow dissociation of the cation from combination with proteins. Therefore prolonged irrigation is indicated and at least 2 L of solution should be used. Although rapid flushing with the first 500 mL is prudent, slow continuous irrigation, as discussed later, at a rate sufficient to generate a continuous trickle is often more effective and better tolerated than continued high-volume flushing. If the nature of the offending agent is unknown or in question, use prolonged irrigation.

Consult ophthalmology for all alkaline, hydrofluoric acid, and heavy metal acid injuries. Irrigation on an inpatient basis may be required for a period of 24 hours or longer, especially when the cornea is hazy or obviously thickened. Note that the magnesium contained in sparklers combines with water from tears to produce magnesium hydroxide. Treat such fireworks injuries as alkaline injuries rather than as thermal injuries.

The pH of the conjunctival fornices with a pH paper strip to check the effectiveness of irrigation (see Fig. 62.10, step 6). In addition to litmus paper, the pH indicator on urine multi-indicator sticks can be used. The pH indicator on urine dipsticks is conveniently closest to the handle; all the distal indicator squares can be cut off with scissors. Normal tear film pH is 7.4. Use the noninjured eye as a control if the results are equivocal. If the pH measured in the conjunctival fornices is still abnormal after the initial irrigation, continue to irrigate. If the pH is normal after irrigation, wait 20 minutes and check it again to make sure that it remains normal, especially if alkaline contamination has occurred. Delayed changes in pH are usually the result of incomplete irrigation and inadequate swabbing of the fornices. In anticipation of this deficiency, measure the pH deep in the fornices. Consider double-lid eversion with a lid elevator to expose the upper fornix for swabbing, irrigation, and pH testing.

Prolonged Irrigation

Alkaline burns may require prolonged irrigation, and it is essential to consult ophthalmology in such cases. The Morgan Therapeutic Lens is a contact lens–type irrigation device that can provide slow, continuous irrigation once the more vigorous initial irrigation has been completed (Fig. 62.13 and Video 62.3). First, anesthetize the eye with topical anesthetic drops. Then place the device carefully on the surface of the eye with the lids closed around the intravenous tubing adaptor. Attach the intravenous tubing to the adaptor and provide continuous flow through the device onto the cornea and into the fornices. As the local anesthetic agents wash out during the irrigation process, the device can become uncomfortable, so reapply the anesthetic drops frequently during irrigation for patient comfort. Such short-term use of local anesthetics will not inhibit healing of the cornea.
experimental evidence suggests that massive parenteral or oral ascorbic acid supplementation may prevent the development of deep corneal injury, but such treatment has not gained universal acceptance.

**Summary**

Eye irrigation is easy, and complications associated with the technique are usually minimal. At times the clinician may be unsure whether a chemical injury is toxic enough to warrant irrigation. If any doubt exists, err on the side of irrigating the eye rather than omitting this vital procedure and risking progression of the eye injury.
In the evaluation of a patient in whom a penetrating injury to the globe is suspected, perform a careful expeditious examination of the eye, preferably with a slit lamp. Avoid any pressure on the eye or rapid eye movements. If perforation is obvious (e.g., teardrop pupil, flaccid globe, flat anterior chamber, prolapsed iris) or confirmed by slit lamp (positive Seidel test; see Fig. 62.8, plate 4), do not perform any procedures (except perhaps irrigation or ultrasound; see later) and consult ophthalmology early for definitive diagnosis and care (Fig. 62.14). Until the ophthalmologist arrives, protect the eye from further harm by keeping the patient quiet, elevating the head of the bed, and placing a protective shield over the eye. Commercial shields are available for this purpose. When a metal shield is not available, construct a makeshift protective shield with the material available (e.g., paper, plastic, or Styrofoam cups; Fig. 62.15). The protective shield helps avoid pressure on the globe and overlying tissue and assists in preventing extrusion of vitreous and other ocular contents. Extend the edges of the shield up to or beyond the bony orbital rim for this purpose. Apply adhesive tape over the shield from the forehead to the cheek to secure the shield in position.

If a patient has a globe perforation, treat with systemic antibiotics (a combination of cefazolin and gentamicin is a good initial choice), tetanus toxoid, and antiemetics in doses aggressive enough to halt vomiting.

**Figure 62.14** Serious eye injuries. A, Corneal laceration with prolapse of the iris. The extruded iris is dark and mimics a corneal foreign body (arrow). Frequently, the only clue is an abnormal pupil, and the extruded iris may not be appreciated as intraocular tissue. The pupil is irregular (often pear or teardrop shaped) and points toward the laceration. B, A pear-shaped pupil without protrusion of the lens is a more subtle, yet characteristic indication of a perforated globe (arrow). C, Another indication of a penetrating globe injury is periorbital fat protruding from an upper eyelid laceration (arrow). This patient was stabbed with a knife. D, This patient has an obviously cloudy lens soon after trauma. A projectile entered the temporal portion of the globe and produced a seemingly minor scleral hemorrhage. Patients with penetrating injuries to the globe should be treated with systemic antibiotics (such as a combination of cefazolin and gentamicin), tetanus toxoid if indicated, and antiemetics to control vomiting (which raises intraocular pressure). (A, Courtesy Lawrence B. Stack, MD.)

**Figure 62.15** When a penetrating globe injury is suspected and a metal shield is not available in the emergency department or prehospital setting, a makeshift shield can be fashioned with available material. A paper cup was used to fashion this shield.
BEDSIDE ED OCULAR ULTRASOUND

ED ocular ultrasonography has markedly advanced the emergency physician’s (EPs) ability to quickly address important ocular conditions. In 2000, Blaivas and colleagues published the first series of patients who were successfully evaluated with bedside ocular ultrasound by EPs. Many subsequent studies have demonstrated that with proper training and equipment, EPs are competent and care is improved by using this noninvasive technology at the bedside. 54–56

Indications and Contraindications

Bedside ED ocular ultrasonography may be appropriately used as an adjunct in the evaluation of altered vision, suspected FB, ocular pain, headache, eye trauma, head injury, or altered mental status. Conditions that are reliably diagnosed include globe rupture, intraocular FB, vitreous hemorrhage, lens dislocation, retinal detachment, and elevated intraocular pressure.

There are no absolute contraindications for bedside ED ocular ultrasound, given proper equipment, an experienced sonographer, and a cooperative patient. The evaluation of a potential open globe deserves special mention. Whereas it is generally true that the eye should be protected from further manipulation after suspecting an open globe, evaluation with bedside ED ocular ultrasound is the exception. It is imperative that the EP performing the bedside ultrasound be mindful of the possibility of an open globe and utilizes the no pressure technique described later. An immediate benefit is gained when the EP carefully uses this technique to avoid a delay in diagnosis and treatment. Any delay incurs an increased risk of adverse outcome. In general, the benefit obtained from the use of bedside ED ocular ultrasound outweighs the risk of further examination of the eye in this setting. That said, if the physical exam obviously suggests open globe pathology, the clinician is discouraged from this procedure, as it carries the risk of vitreous extrusion (even with minimal globe compression) with little benefit to the diagnostic evaluation.

Equipment

Appropriate ultrasound machine with linear array transducer, 7.5 to 10 mhz or higher.

Procedure

Ensure ultrasound equipment is clean and disinfected. Adjust setting on machine for ocular ultrasound. Place the patient in a recumbent or semi-reclined position to minimize runoff of gel. Instruct the patient to keep eyes closed, but not clenched, to look straight ahead, and not to move the eyes. It is suggested that the closed eye be covered with a large piece of clear Tegaderm dressing (3M, St. Paul, MN) to facilitate the ultrasound examination without contaminating it with gel or the probe. Place a copious amount of gel over the Tegaderm dressing (Fig. 62.16). The large amount of gel placed over the closed eyelid forms an acoustic window for the ultrasound probe. This allows an image to be obtained without additional pressure applied to the eye.

Instruct the patient to keep eyes closed, but not clenched, to look straight ahead, and not to move the eyes. It is suggested that the closed eye be covered with a large piece of clear Tegaderm dressing (3M, St. Paul, MN) to facilitate the ultrasound examination without contaminating it with gel or the probe. Place a copious amount of gel over the Tegaderm dressing (3M, St. Paul, MN) to facilitate the ultrasound examination without contaminating it with gel or the probe. Place a copious amount of gel over the closed eyelid to be examined, enough to completely fill the sulcus formed by the eye so that the transducer can touch the gel in a transverse plane without touching the eyelid and without breaking contact of the transducer with the gel (Fig. 62.16). Hold the transducer in such a way so that the examiner’s hand can stabilize the probe against movement by resting the hand on the patient’s forehead, bridge of the nose, or maxilla (Fig. 62.17). Barely touch the probe to the gel without applying pressure to the eyelid (no-pressure technique) (Fig. 62.18). Maintain this contact throughout the examination. As soon as the probe touches the gel, the ocular structures should immediately come into view. Adjust probe position to obtain a clear cross section of the eye showing anatomic structures (anterior chamber, iris, lens, vitreous, retina, nerve sheath). Adjust gain for a clear image, starting with maximal settings and reducing it until structures come into view again. Position the optic nerve in the middle of the screen and adjust the depth to view approximately 1 cm of the optic nerve. Identify the optic structures (Fig. 62.19). Identify abnormal findings (see Figs. 62.19B and C), and describe abnormal findings in the caption. Use multiple planes to clarify normal and abnormal findings. A direct and consensual pupillary light reflex can be observed by viewing the pupillary response to light via ultrasound.
Probe stabilization is accomplished by resting the sonographer’s hand on the patient’s nose and face while the probe is gently touched to the surface of the gel. The acoustic window can be seen underneath the probe. (Courtesy Mary Jo Chandler, PA-C.)

To measure optic nerve sheath diameter, ask the patient to adjust the gaze of the examined eye, approximately 10 degrees laterally. When viewed at this angle, the optic nerve is aligned with the ultrasound beam and decreases the possibility of obtaining a falsely widened measurement. This can occur if the optic nerve is imaged at an angle. Measure the width of the optic nerve sheath 3-mm deep to the retina (Fig. 62.20). Obtain at least two measurements and average for a final measurement. A diameter of greater than 5 mm is considered to be widened and suggests increased intracranial pressure.

**Complications**

Inadvertent pressure applied to an eye with a ruptured globe could result in further damage to the globe. Excessive scan times could exceed exposure limit guidelines.

**Findings**

Globe rupture, intraocular FB, vitreous hemorrhage, retinal detachment, dislocated lens, and elevated intraocular pressure may be seen. These findings will be obscured if periorbital or orbital emphysema is present or if there is excessive air in the gel.

![Figure 62.18](image1)

![Figure 62.19A](image2) Normal anatomy is easily identified. Findings shown include: B, Retinal detachment, C, Vitreous hemorrhage. (Courtesy Lauren Oliveira, MD.)
Summary
ED bedside ocular ultrasound is a valuable technique that can be used by the EP to discover important findings in the ED setting. It offers significant advantages for evaluation of ocular and central nervous system complaints, and it is particularly applicable to a possible open globe injury. It can be performed reliably at the bedside by EPs trained in its use.

OCULAR FB REMOVAL
Patients with an external FB in the eye are frequently seen in EDs. They are often in pain and desperate for help. Maintain a high degree of suspicion for FB injuries and perforation of the eye because such injuries may be occult and not readily detected. Not all FB injuries are associated with pain. Glass embedded in the cornea may be particularly difficult to detect. This section reviews procedures for locating and removing extraocular FBs and for appropriate postprocedural care. Finally, a brief discussion covering evaluation of the eye for potential perforation of the globe and detection of the presence of an intraocular FB is provided.

Indications and Contraindications
Extraocular FBs must always be removed. The timing of removal and the technique required vary according to the patient’s clinical status and the type of injury. For the most part, the emergency clinician can proceed directly to removal of the object via the techniques described in this section. When the patient is extremely uncooperative (e.g., an intoxicated patient, a mentally deficient patient, or a young child) or when the injury is complicated (e.g., deeply embedded object, multiple foreign objects from a blast injury, or possible globe penetration), consult ophthalmology immediately. A patient with a suspected FB or abrasion after exposure to a projectile (e.g., grinding wheel, hammering, metal objects colliding) should be rigorously evaluated for the presence of a deep intraocular FB. See further discussion later in this chapter. A penetrating injury of the cornea is of particular concern because the iris tissue may prolapse and have an appearance similar to a corneal FB (see Fig. 62.14). Hence, in addition to the history of projectile exposure, an irregular pupil, especially a pear-shaped pupil, should alert the clinician that a penetrating injury might have occurred.

Equipment
The following equipment is necessary for removal of an extraocular FB:
- Topical anesthetic, such as 0.5% proparacaine
- Sterile cotton-tipped applicators
- Fluorescein strips
- Magnification: loupes plus a Wood’s lamp, Eidolon Blumina-tor ophthalmic illuminator, or slit lamp
- Eye spud or 25-gauge needle attached to a 1- or 3-mL syringe or to the tip of a cotton-tipped applicator
- Dilator drops, such as 5% homatropine
- Antibiotic ointment, such as erythromycin

Consideration of an Intraocular FB
When examining a patient with an ocular FB sensation, always remain cognizant of the potential for an intraorbital or intraocular FB. Penetrating injuries represent a greater threat to visual loss than an extraocular FB does and can be disastrous if overlooked. Note that an intraocular FB can be deceptively subtle on initial evaluation. Bedside ED ultrasound is a useful adjunct for identification of intraocular FBs (Fig. 62.21).

The clinical findings are most helpful in determining which patients are at risk for a penetrating injury to the globe. An individual who complains of an FB sensation in the absence of trauma or one whose history is simply that something “fell” or “blew” into the eye is at low risk for perforation of the globe. Conversely, there is a greater probability of globe penetration in an individual who has sustained a high-velocity wound to the eye (e.g., drilling, hammering, grading metal, blasting rock). The presence of any of the following findings on physical examination should alert the clinician to a probable intraocular FB: irregular pupil, shallow anterior chamber on
slit lamp examination, prolapsed iris, positive Seidel test (see earlier section on The Fluorescein Examination), focal conjunctival swelling, hemorrhage, hyphema, lens opacification, and reduced intraocular pressure (IOP). Do not perform tonometry if penetrating injury to the globe is suspected. Be aware that a penetrating injury may not be associated with eye pain. If there is strong historical evidence and physical findings to support a diagnosis of globe penetration, obtain emergency ophthalmologic consultation.

An intraocular FB is often not visible on direct ophthalmoscopy. Although orbital radiography for radiopaque objects and ultrasonography of the globe have been used for indirect FB localization, computed tomography of the orbit is considered the most useful technique (Fig. 62.22). When plain orbital radiography is performed to look for an intraocular FB, be aware that an eyelid FB may mimic an intraocular FB. Patients with a suspected metallic FB should not undergo computed tomography and ophthalmologic consultation. The role of the emergency clinician is to suspect the diagnosis, protect the eye from further harm, and obtain ophthalmologic consultation. The remainder of this section addresses the problem of extraocular FBs.

**Procedure**

**FB Location**

The first step is to locate the FB. Apply a drop of topical anesthetic to the inside of the lower lid as described previously (see Fig. 62.7A). The presence of vertical corneal abrasions from FBs under the lids is helpful in localizing these hidden foreign objects (see Fig. 62.8, plate 7). Use a penlight and loups or a slit lamp to examine the bulbar conjunctiva by having the patient look in all directions. Examine the inside of the lower lid by pulling it down with the thumb while asking the patient to look up. Evert the upper lid by asking the patient to look down as the end of an applicator stick is pressed against the superior edge of the tarsal plate of the upper lid. Meanwhile, grasp the lashes and pull down, out, and then up to flip the eyelid over (Fig. 62.23).

Minute FBs under the lid may be missed with simple visual inspection. Ideally, examine the everted lid under magnification with loupes or a slit lamp. With simple lid eversion, it is still not possible to see the far recesses of the upper conjunctival fornix. Although double eversion of the upper lid is helpful, the best way to rule out an FB in the upper fornix is to sweep the anesthetized fornix with a moistened applicator as the upper lid is held everted. Examine the tip of the applicator for minute FBs. Small conjunctival FBs not hidden by the lids are often best removed with a moistened nasopharyngeal swab (e.g., nasopharyngeal Calgiswab [Puritan, Guilford, ME]).

Reexamine the cornea. Most corneal FBs have an area of fluorescein staining around them. Use a slit lamp or other magnification device such as the Bluminator (Eidolon) to make the examination easier. If the clinician is limited to loups and a penlight, shine the light diagonally on the cornea to locate the FB. With a history of a high-speed projectile hitting the eye, rule out an intraocular FB. In the case of a blast injury, multiple FBs may penetrate the eye. If an FB cannot be found on the surface despite a suggestive history, examine the eye for physical evidence of penetration, as discussed earlier. Dilate the pupil and examine the fundus. Though not foolproof, bedside ultrasonography may identify the presence of a metallic FB (sensitivity of 87.5%, specificity of 95.8%, and positive and negative predictive values of 96.5% and 85.2%, respectively). If in doubt regarding an intraocular FB, consider computed tomography and ophthalmologic consultation.

**FB Removal**

Once an extraocular FB is located, the technique for removal depends on whether it is embedded. Reapplication of topical anesthetic drops may be needed. If the FB is lying on the surface, eject a stream of water from a syringe through a plastic catheter, which will usually wash the object onto the bulbar conjunctiva. Once the FB is on the lower lid or bulbar conjunctiva, gently touch a wetted cotton-tipped applicator to the conjunctiva and the object will adhere to the tip of the applicator. Be aware that overzealous use of an applicator for removing corneal FBs can lead to extensive corneal epithelial injury. A spud device is required for removal of objects that cannot be irrigated off the cornea.

To remove embedded corneal FBs (Fig. 62.24), use a commercial spud device, a bur drill, a short 25- or 27-gauge needle on a small-diameter syringe (e.g., insulin or tuberculin syringe), or a cotton-tipped applicator. Use the applicator or syringe as a handle for the attached needle. Contrary to what one might expect, it is difficult to penetrate the sclera or the cornea with a needle, especially when it is applied tangentially to the cornea.

As with removal of conjunctival FBs, anesthetize the eye. Position the patient so that the head is well secured (preferably in a slit lamp frame). At this point, provide a simple explanation of the procedure, which usually ensures excellent compliance on the part of the patient. Rest your hand on the patient’s cheek so that unexpected movements by the patient will not result in large movements of the removal device. Instruct the patient to gaze at an object in the distance (e.g., the practitioner’s ear when a slit lamp is used) to further stabilize the eye. Bring the removal device close to the eye under direct vision; then while it is in focus, manipulate it under the magnification device (e.g., Wood’s lamp, Eidolon Bluminator ophthalmic illuminator [see Fig. 62.8], or slit lamp) to remove the FB. Hold the device tangential to the globe, and pick up or scoop out the foreign object using the magnification device.
To evert the upper eyelid, place a cotton-tipped applicator against the superior edge of the tarsal plate.

Grasp the eyelashes and pull them down, then out, and then up and over the applicator.

Hold the everted lid in place with the applicator. Use a second applicator to sweep the interior surface of the lid.

This patient complained of an FB in the eye despite irrigation. Lid eversion revealed a small speck (arrow) under the upper lid. This could cause a cornea abrasion characterized by vertical striations (see Fig. 62.7, plate 7).

The foreign body was easily removed (arrow) by touching it with a moistened cotton-tipped applicator.

This patient had a swollen and tender upper eyelid thought to be secondary to a stye. With lid eversion, a small pustule (arrow) was found under the upper eyelid.

With a 27-gauge needle, the pustule was incised and a drop of pus was expressed; she made a rapid and uneventful recovery.

Figure 62.23 Lid eversion and foreign body (FB) removal.
Corneal Foreign Body Removal

This embedded corneal FB is readily seen under slit lamp examination. A removal device (needle, spud, or bur drill) should be used for careful removal. A rust ring will remain if the FB has been there for only a few hours.

Rust rings (arrow) are retained FBs and are removed in a similar manner. Most rust rings should be removed, but there is no urgency. Small ones out of the line of sight may remain. A bur drill can be used for attempted removal, which if unsuccessful, can be reattempted in 24 hours. Alternatively, a small needle can be used to loosen the edges and then the ring scooped out. Both procedures will leave a corneal abrasion.

Under direct vision (not looking through the slit lamp), bring the syringe close to the eye while resting the hand on the patient's cheek. Be sure that the patient's forehead maintains continual contact with the crossbar on the slit lamp.

While looking through the slit lamp, bring the needle to the cornea and remove the FB.

Hold the side of the instrument (drill bit or beveled edge of the needle) tangential to the cornea.

A variety of instruments may be used for FB removal, including an eye spud (A), a cotton-tipped applicator (B), and a 25- or 27-gauge needle on a tuberculin syringe (C).

Figure 62.24 Corneal foreign body (FB) removal.
object. If a bur drill is used, press the side of the drill against the FB until removal is accomplished.

During removal, rest your hand against the patient’s face. It may also be helpful to brace the elbow with a pad or half-full tissue box to provide further support for the arm while removing the FB. If right-handed, place the lower part of your hand against the left maxillary bone when removing a foreign object from the patient’s left eye and against the bridge of the patient’s nose or infranasal area when removing an object from the right eye. If left-handed, reverse these positions. Using loupes or a slit lamp for magnification is highly recommended to monitor the entire process. In particular, corneal contact with the spud device is more readily discerned when magnification is used. Only topical anesthesia is required to remove FBs from the cornea. Although patients may feel pressure during removal of FBs, pain should not be felt after the eye is anesthetized.

Rust Rings
A common problem with metallic FBs is that rust rings can develop (see Fig. 62.24, plate 2). They can develop within hours because of oxidation of the iron in the FB. There are two preferred techniques for removal of a rust ring. The most direct technique is to remove the ring at the same time as the FB, either with repeated picking away with a needle, spud device, or with a rotating bur. The second approach is to let the iron of the rust ring oxidize and kill the surrounding epithelial cells during a 24- to 48-hour period. After that, the rust ring will be soft and often comes out in one solid plug. Generally, a small rust ring produces little visual difficulty unless it is directly in the line of sight. If large, a rust ring may delay corneal healing. Close follow-up is important to ensure healing of the cornea and total removal of the rust ring and FB.

Multiple FBs
If multiple FBs are present in the eye, such as from an explosion, refer the patient to an ophthalmologist. One technique that may be chosen by the ophthalmologist is to denude the entire epithelium with alcohol and remove the superficial FBs. The deeper ones gradually work their way to the surface, sometimes years later.

Aftercare
After removing the FB, an antibiotic ointment is frequently instilled. Though commonly used, the value of the ointment for superficial corneal defects after removal of an FB is unproved, and no specific standard of care is supported by scientific evidence. Contrary to previous techniques, conjunctival and corneal abrasions do not need patching. Data suggest that eye patching offers no benefit in healing corneal abrasions secondary to FBs. If a superficial injury is sustained from the FB, instruct the patient to return only if the eye does not feel completely normal or if there is any blurred vision. The majority of superficial injuries heal without difficulty. The patient should be warned that the FB sensation might return temporarily when the anesthetic agent wears off. One animal study involving direct ocular exposure to Clostridium tetani organisms suggested that nonpenetrating ocular injuries are unlikely to lead to tetanus. Tetanus prophylaxis after corneal FB removal is not standard, but it should be considered. However, tetanus prophylaxis appears to be essential for injuries that penetrate through the cornea or sclera.

Complications
Complications associated with ocular FB removal are rare. The most frequent problem is incomplete removal of the FB. In such cases the epithelium has difficulty healing over the affected area, and thus the eye stays inflamed. Eventually, the diseased epithelium either sloughs off and heals or heals over the FB remnants, which are gradually absorbed. In either case, adverse effects on the eye are minimal; a minute scar on the cornea, even directly in the center, will rarely affect vision. Nonetheless, incomplete removal of a corneal foreign object warrants ophthalmologic follow-up.

Conjunctivitis may develop after removal of an extraocular FB. In most cases the bacteria producing the infection are introduced by the patient rubbing the irritated eye.

Although perforation of the globe by the clinician’s spud device is theoretically possible, this complication is exceedingly rare. Treatment of this type of corneal puncture wound consists of antibiotics, placement of an eye shield, and ophthalmologic consultation. In the absence of resultant endophthalmitis, permanent sequelae are unlikely to develop.

Epithelial injury can occur when cotton-tipped applicators are used to vigorously remove corneal FBs. Indeed, the use of cotton-tipped applicators for embedded corneal FB removal is condemned.

Summary
Ocular FBs are one of the most common eye emergencies. Searching for and removing the FB is usually straightforward. The only real trap is missing an intraocular FB. This must be ruled out if there is a history of a high-speed projectile hitting the eye or if the findings on physical examination suggest globe penetration.

EYE PATCHING
Patching the lids shut has traditionally been the last step in the treatment of a number of common eye emergencies; however, multiple studies have shown that eye patching offers no benefit in pain relief or healing rates in patients with conjunctival or corneal abrasions. A meta-analysis of studies on eye patching and corneal abrasions or ulcers showed that patching might actually slow healing rates and patients might actually have worsening of their pain. This was found to be true in children as well. Patching is also contraindicated in contact lens wearers and in situations in which the abrasion or ulcer may be infected. In summary, eye patching is no longer indicated and might actually worsen the ophthalmologic process that it once was thought to help. Application of a therapeutic bandage contact lens directly to the cornea has been recommended as a possible treatment of corneal epithelial defects. Evidence from several small studies suggests that a bandage contact lens is safe, effective, and well tolerated and allows a significant number of patients to immediately resume their regular activities while maintaining baseline visual acuity. Further study on the application of this modality in the ED setting is needed.
CONTACT LENS PROCEDURES

An estimated 24 million Americans wear a form of contact lenses. Removal of these lenses in the ED may be required to permit further evaluation of the eye or to prevent injury from prolonged wear. Emergency clinicians also evaluate patients for “lost” contact lenses, which may be trapped under the upper lid. At times, patients may request that the clinician remove a lens that they have failed to extract from the cornea. Corneal ulcers can occur in patients who wear contact lenses and may require prompt treatment. This section on contact lens procedures addresses these concerns and discusses injuries associated with attempts at removal, the mechanism of injury from prolonged wear, and instructions to be given to patients at discharge. Furthermore, this section introduces the use of bandage contact lenses for the treatment of acute corneal abrasions.

The first contact lenses were scleral lenses made of glass. These lenses, which covered the cornea as well as much of the surrounding sclera, are reported to have been in use from 1888 to 1948. Glass corneal lenses (sitting entirely on the cornea) made by the Carl Zeiss Optical Works of Jena were first described in 1912. A practical scleral lens using methyl methacrylate rather than glass was discussed by Obrig and Mullin in 1938. In 1947, Tuohy redeveloped the corneal lens with methylmethacrylate. This was the forerunner of the current hard contact lens. The development of lenses made of soft gas-permeable polymers was reported in Czechoslovakia in 1960. These hydrogel (hydrophilic gelatinous–like) lenses have evolved into today’s soft contact lenses. Soft contact lenses now come in a variety of types, including extended and daily wear. The majority of soft contact lenses in use are now disposable. Soft contact lenses have been used therapeutically by ophthalmologists for decades.

Corneal Injury From Contact Lens Wear
Hard Contact Lenses

Oxygenation of the cornea is dependent on the movement of oxygen–rich tears under hard contact lenses during blinking. During the “adaptation” phase of early wear, a wearer of hard contact lenses produces hypotonic tears as a result of mechanical irritation from the lens. This results in corneal edema, which reduces subsequent tear flow under the lens during blinking. Overwearing a lens at this time leads to corneal ischemia, with superficial epithelial defects found predominantly in the central corneal area (see Fig. 62.9), where the least tear flow occurs. With adaptation, the tears become isotonic and the blinking rate normalizes, thus permitting increased wearing time. During early adaptation, blinking is more rapid than normal and then slows to a subnormal rate during late adaptation. Mucous delivery to the cornea in the tear film may also play an important role in maintaining corneal lubrication. Tight-fitting contact lenses may never permit good tear flow despite an adaptation phase; individuals with tightly fitted lenses may never be able to wear their original contact lenses for longer than 6 to 8 hours. Lenses that are excessively loose can also cause irritation by moving during blinking. Rough or cracked edges can cause corneal abrasions.

In the ED, a patient with eye irritation caused by prolonged contact lens wear may be either a new or an adapted wearer. An adapted wearer may have been exposed to chemical irritants (e.g., smoke), which reduces the tonicity of tears and leads to corneal edema and decreased tear flow. Alternatively, an adapted wearer with irritation may have ingested sedatives (e.g., alcohol) or may have fallen asleep wearing the contact lenses, thus decreasing blinking and tear flow. Another possibility is that the patient may actually be wearing tight-fitting contact lenses that have never allowed true adaptation despite many months of wear.

A patient with overwear syndrome usually awakens a few hours after removing the lenses. The patient experiences intense pain and tearing similar to that caused by an FB. The delay in onset of the symptoms until after removal of the lenses is caused by a temporary corneal anesthesia produced by the anoxic metabolic by-products that build up during extended wear. A second factor is the slow passage of microcysts of edema, which are pushed up to the corneal surface by mitosis of the underlying cells. When the cysts break open on the surface, the corneal nerve endings are exposed.

Most patients with overwear syndrome can be managed with reassurance, frequent administration of artificial tears, oral analgesics, and advice to “wait it out” in a darkened room. In the past some patients required patching for comfort. A patient who has experienced no problems with contact lenses before an overwear episode can return to using the lenses after 2 or 3 days of wearing glasses but should be advised to build up wearing time gradually. A patient who was having chronic problems with lens comfort before the episode should check with an ophthalmologist before using the contact lenses again.

Soft Contact Lenses

Although the cornea is also oxygenated by way of the tear film with soft contact lenses, only approximately one-tenth of the flow behind the lens that occurs with a hard lens is present during soft contact wear. The high degree of lens gas permeability permits the majority of oxygenation to occur directly through the lens. A hydrogel lens is more comfortable than a hard contact lens because lid motion over the lens is smooth. The minimization of lid and corneal irritation allows a more rapid adaptation phase because the initial reflex-induced changes in tearing and blinking are reduced. Nonetheless, the lenses may still lead to corneal edema and secondary hypoxic epithelial changes if worn for an excessive period when blinking is inhibited. Some individuals can tolerate the lenses for extended periods and may on occasion sleep with the contact lenses in place, although this practice is not encouraged. Newer extended-wear hydrogel lenses permit wear for up to 1 week without injury. These lenses are not discernible from standard soft lenses on examination.

Although the acute overwear syndrome that occurs with hard contact lenses can also occur with soft lenses, it is infrequent. More commonly, ocular damage from soft contact lenses falls into one of three categories:
1. Corneal neovascularization (Fig. 62.25A). Frequently, the patient is asymptomatic, but on slit lamp examination, fine vessels are seen invading the periphery of the cornea. Refer the patient to an ophthalmologist for refitting with looser or thinner lenses or with contact lenses that are more gas permeable.
2. Giant papillary conjunctivitis (see Fig. 62.25B). The patient notes decreased lens tolerance and increased mucous production. On examination of the tarsal conjunctiva (best seen with eversion of the upper lid), large papillae are seen. These papillae appear grossly as a cobblestoned surface. Instruct the patient to discontinue wearing the lenses until
Indications for Removal

Remove a contact lens in the following situations:

1. Contact lens wearer with an altered state of consciousness. The emergency clinician should always be aware that a patient with a depressed or acutely agitated sensorium might be unable to express the need to have the contact lenses removed. Furthermore, it is likely that patients with a depressed sensorium will have decreased lid motion. During the secondary survey of these patients, identify the presence of the lenses and arrange for their removal and storage to prevent harm from excessive wear or possible accidental dislodgment at a later time. Without magnification, soft contact lenses may be difficult to see. Examine the eye with an obliquely directed penlight to reveal the edge of the soft lens 1 to 2 mm from the limbus on the bulbar conjunctiva.

2. Eye trauma with lenses in place. After measurement of visual acuity with the patient's lenses in place, remove them and perform a more detailed examination of the cornea. Fluorescein may discolor hydrogel lenses; when possible, remove extended-wear lenses before using this chemical. After the dye is instilled, flush the eyes with normal saline. Advise the patient to wait at least 1 hour before reinserting the lenses. The availability of single-use droppers of 0.35% fluorexon (Fluoresoft, Alden Optical, Lancaster, NY) has permitted safe staining of eyes when soft lenses are to be worn immediately after the examination. Limited eye irrigation after the use of fluorexon drops is still recommended before the reinsertion of soft contact lenses.

3. Inability of the patient to remove the contact lens. A patient may have a hard contact lens that cannot be removed because of corneal edema from prolonged wear. Alternatively, the patient may have a “lost” contact lens believed to be located behind the upper lid. Because there is no urgency for removal of the contact lens in the out-of-hospital setting, it can wait until the patient has been evaluated by a clinician.

Contraindication to Removal

The only major problem with contact lens removal occurs when the cornea may be perforated. In this case the suction cup technique of removal, described later, is preferred.

Procedure

Hard Contact Lens Removal

A number of maneuvers have been devised for removal of a corneal lens. One technique is to first lean the patient's face over a table or a collecting cloth. Pull the lids temporally from the lateral palpebral margin to lock the lids against the edges of the contact lens. Ask the patient to look toward the nose and then downward toward the chin. This movement works...
the lower eyelid under the lower edge of the lens and flips the lens off the eye. The technique requires a cooperative patient because the clinician must pull the patient's lids tightly against the edge of the contact lens. The movement of the patient's eye then flips the contact free.

In an unresponsive patient in the supine position, modify the technique. Take a more active role in lid movement by using the following procedure (Fig. 62.26A). Place one thumb on the upper eyelid and the other on the lower eyelid near the margin of each lid. With the lens centered over the cornea, open the eyelids until the margins of the lid are beyond the edges of the lens. Then press both eyelids gently but firmly on the globe of the eye and move the lids so that they are barely touching the edges of the lens. Press slightly harder on the lower lid to move it under the bottom edge of the lens. As the lower edge of the lens begins to tip away from the eye, move the lids together, which allows the lens to slide out sufficiently so that it can be grasped. Remember to use clean hands (and preferably wear examination gloves that have been rinsed in tap water or saline) when removing the lens.

Alternatively, move the lens gently off the cornea with a cotton-tipped applicator to guide the lens onto the sclera. Force the tip of the applicator under an edge of the lens and flip the contact loose. Apply topical anesthetic when using an applicator if the patient is awake. Take care with this technique to avoid contact of the applicator with the cornea when the lens is moved off the eye. Perhaps the easiest technique is to use a moistened suction-tipped device and simply lift the lens off the cornea (see Fig. 62.26B).

Scleral lenses (hard contact lenses that cover both the cornea and an amount of the sclera) can be removed by an exaggeration of the manual technique described earlier (see Fig. 62.26C). Elevation of the lens with a cotton-tipped applicator or a suction-tipped device is also an effective technique. Soft contact lenses should not be removed with a suction-tipped device because tearing or splitting of the lenses might occur.

**Soft Contact Lens Removal**

With clean hands (preferably using gloves rinsed in saline or tap water), pull down the lower eyelid with the middle finger. Place the tip of the index finger on the lower edge of the lens. Slide the lens down onto the sclera and compress it slightly between the thumb and index finger. This pinching motion folds the lens so it can be removed from the eye (see Fig. 62.26D). Alternatively, use a cotton-tipped applicator (e.g., Q-Tip) instead of a gloved hand. Occasionally, a tight-fitting lens will be difficult to remove. One potential method is the use of topical anesthetic drops, lubricating eye drops (e.g., Refresh Celluvisc [Allergan plc., Dublin, Ireland] lubricant eye drops), and a cotton-tipped applicator to lift the edge of the lens from the limbus. This breaks the seal of the lens on the cornea and allows removal.

**Lens Storage**

After a contact lens is removed, store it in sterile normal saline solution. Use the patient's own storage container and lens solution if available. A variety of alternative sterile containers are available for use in the ED. Be certain to keep the right and left lenses separate and in appropriately labeled containers. The containers should be kept with the patient until a friend or family member can procure them, or they should be locked up with the patient's valuables.

---

**Evaluation of a “Lost” Contact Lens**

A patient may request examination for a “lost” contact lens. The patient may be unsure whether the lens is hidden under a lid, remains on the cornea, or is truly outside the eye. Evaluation of a patient with a lost contact should begin, as should all eye examinations, with measurement of visual acuity. Measure visual acuity preferably with a 20-ft eye chart. Diminished visual acuity in the eye with the lost contact is convincing evidence that the lens is missing. Though transparent, soft contact lenses in proper position are usually seen when viewed closely with loupes or a slit lamp. The lens forms a fine line at the point where it ends on the sclera several millimeters peripheral to the limbus. Hard contact lenses are even more evident as they change in position on the cornea.

If the contact lens is not evident on initial inspection, evert the lids as discussed in the earlier section on Ocular FB Removal (double eversion of the upper lid). If the lens is still not visible, place a drop of topical anesthetic in the eye. Gently sweep the upper fornix with a moistened cotton-tipped applicator while the patient looks toward the chin. If the lens is still not evident even though the patient remains insistent that it is in the eye, perform a fluorescein examination after explaining that the dye will color the lens (permanently). Evert the upper lid again and examine with an ultraviolet light source.

If the lens remains elusive, reassure the patient that a thorough examination was performed and that no object was located under the eyelids or on the cornea. Next, examine the cornea for defects that warrant antibiotic ointment for treatment of corneal abrasion. Follow-up with the patient's eye specialist for a replacement lens and to provide further reassurance is advised. Ask the patient to retrace his movements at the time that the contact began to give trouble or was missed. Check the clothing being worn at that time and look for the lens there. A final possibility is that the patient may have accidentally placed the two lenses together in the same side of the carrying case, thereby causing them to stick together. Take a methodical approach, as outlined earlier, to ensure that no lens remains hidden in the eye.

**Complications of Lens Removal**

A corneal abrasion can occur during lens removal. It is difficult at times to determine whether the injury was produced by the patient or was a result of removal by the clinician. Fortunately, the corneal injury is generally of a superficial nature and responds well to symptomatic care.

**Summary**

Contact lens removal is usually simple. Challenging situations include identifying patients at risk for corneal injury from overuse, helping patients who have lost a contact lens in the eye, and providing aftercare instructions for patients with contact lens–related problems.

**BANDAGE CONTACT LENSES FOR TREATMENT OF CORNEAL ABRASIONS**

Acute corneal abrasions can cause significant pain, limit function, and result in lost days of work. Several studies have shown that bandage contact lenses are very effective in reducing pain
Contact Lens Removal

A. Manual Removal of Hard Contact Lens
1. Separate the eyelids.
2. Entrap the lens edges with the eyelids.
3. Expel the lens by forcing the lower lid under the inferior edge of the lens.

B. Suction Cup Removal of Hard Contact Lens
Use a moistened suction cup to remove a hard contact lens.

C. Removal of a Hard Scleral Lens
1. Separate the eyelids.
2. Force the lower lid beneath the edge of the scleral lens by temporal traction on the lower lid.
3. Lift the lens off the eye.

D. Removal of a Soft Contact Lens
1. Separate the eyelids and then move the contact onto the sclera with the index finger.
2. Pinch the lens between the thumb and index finger.

without a requirement for narcotic analgesia, in decreasing
time away from work, and in returning to baseline functioning
when used for uncomplicated acute corneal abrasions. Yet
bandage contact lenses are seldom used in the ED. An immedi-
ate benefit can be gained when the emergency practitioner
selectively uses this treatment modality.

**Indications and Contraindications**

Bandage contact lenses may be used appropriately as an adjunct
for the treatment of acute corneal abrasions resulting from
minor trauma when symptomatic relief or rapid return of
functionality is desired. Obtain a clear history of an acute
traumatic abrasion and a fluorescein stain pattern consistent
with this diagnosis.

Bandage contact lenses are contraindicated when the cause
of the corneal epithelial defect is suspected to be due to infec-
tion. Ensure that a corneal ulcer is not present. The hallmark
fluorescein stain pattern consistent with a corneal ulcer is a
round stain with blurred margins, commonly in the central
visual axis (midpupil region) (see Fig. 62.25C). Pain is dispro-
portionate to the findings. An intense ciliary flush and anterior
chamber reaction (cells and flare) may also be present. Avoid
the use of bandage contact lenses in patients with a history of
soft contact lens use and a nontraumatic abrasion in the central
part of the cornea. Follow-up for removal of the bandage
contact lenses and reevaluation of the eye are required to
confirm improvement and identify complications. Unreliable
patients who are not likely to be compliant with follow-up are
at increased risk for infection and thus not good candidates
for this modality.

**Equipment**

Hydrophilic bandage contact lenses are available and include,
but are not limited to, Biomedics 55 (Ocular Sciences, Inc., San
Francisco), −0.50 dioptic power, 8.6 posterior curvature, and
14.2 diameter of the lens, and Acuvue Oasys BC (Johnson &
Johnson Vision Care, Inc., Jacksonville, FL), zero power (plano)
or −0.50 diopter, 8.4 posterior curvature, and 14.0 diameter.
Acuvue Oasys has been approved by the U.S. Food and Drug
Administration (FDA) for up to 1 week of continuous wear.

**Procedure**

Assess and document the pain. Instill 1 drop of a fluoroqui-
nolone ophthalmic solution in the affected eye as prophylaxis
against infection at the time of lens placement. Use of topical
anesthetics is not required, but they have often been used in
the evaluation of abrasions. Use a gloved hand that has been
rinsed with water to remove the t alc. Orient the contact lens
on the tip of the examiner's index finger. Ensure that the lens is
not inverted. The normal configuration of the lens resembles a
cup, whereas an inverted lens resembles a saucer with its edges
flaring outward. Another method to determine orientation
of the lens is to try to close the edges or have them meet
together with your fingers. If the edges curve away from each
other, the lens is inverted. Direct the patient's gaze upward
and then pull the lower lid slightly downward while inserting
the lens over the cornea. Ask the patient to gently blink and
then assess for placement. After a few minutes, reassess and
again document whether pain is present. Instruct the patient
to return for follow-up in 1 day. On return, remove the soft lens
as described earlier, assess vision, document pain, and reevaluate
the eye.

**Complications**

Because of the hypoxic microenvironment of the cornea,
covering the cornea with a bandage contact lens could increase
the opportunity for an infection to occur. Patients who do not
disclose the use of soft contact lenses are at increased risk for
corneal ulcers. Some patients may experience “tight lens
syndrome” because of a lens that does not fit well or from
drying out as the lens is exposed to the elements. Bandage
contact lenses are usually of a generic size and thus do not
offer a precise fit, but they are in place only for a short duration.
Signs and symptoms of a tight lens may include redness, eye
irritation, burning, a dry sensation, blurry vision, halos around
objects, and lack of mobility of the contact lens over the cornea.
Lack of mobility can reduce oxygen tension and thereby cause
corneal edema and an even tighter lens. Rewetting solutions
can improve hydration. Removal of a tight-fitting lens was
described earlier.

**Summary**

Bandage contact lenses have been shown to be safe, effective,
and comfortable in the treatment of corneal abrasions. Careful
patient selection and proper technique will optimize this
treatment modality when appropriately indicated for routine
ED use. Patients will experience significantly decreased pain
without the use of narcotic analgesics, significantly increased
functionality, and decreased time away from work.

**INFECTIOUS KERATITIS**

Infectious keratitis with corneal ulceration can have a variety
of causes, and commonly includes overwear of contact lenses.
Diagnosis of a corneal ulcer requires the use of a slit lamp and
accurate determination of the patient's history. Infectious
keratitis is a frequent problem in ophthalmic practice. Herpes
simplex is a common corneal pathogen. *Acanthamoeba* is another
pathogen particularly associated with contact lens use and
exposure to organism-tainted environments. When a patient
is seen in the ED with a corneal ulcer, promptly refer the
patient to an ophthalmologist. When immediate referral is
not possible, obtain telephone guidance from an ophthalmologist
for initiation of therapy, and arrange for ophthalmology follow-
up within 24 hours.

Patients with herpes simplex keratitis often give a history
of previous episodes of the disease. Patients who undergo almost
any form of corneal stress may sustain an activation of preexist-
ing corneal disease. Herpes simplex keratitis is classically
recognized by its dendritic pattern on fluorescein staining (see
Fig. 62.8, plate 9).

*Acanthamoeba* keratitis is a disease with potentially devastating
consequences. Its frequency seems to be increasing, particularly
in contact lens wearers, and its pathophysiology is not com-
pletely understood. Patients often have a red eye in which the
initial bacterial culture results are negative.

Bacterial keratitis occurs in a variety of settings. Organisms
range from the relatively common *Staphylococcus* (including
methicillin-resistant *Staphylococcus aureus*) or *Streptococcus*
to *Mycobacterium*, which can be difficult to identify. A variety of
antibiotics are used against bacterial agents. Ciprofloxacin is a quinolone that has demonstrated efficacy against most of the common causative agents. Bacterial organisms in the cornea can develop resistance to any antibiotic, and resistance to fluoroquinolones has also been observed. Ideally, treatment follows acquisition of material from the ulcer for culture.

In instances in which a cellular infiltrate is seen on slit lamp examination and there will be a delay of hours before an ophthalmologic consultant can perform the culture, it is prudent to initiate therapy with topical fluoroquinolones such as ciprofloxacin. In such circumstances, obtain corneal samples for culture under the telephone guidance of the consultant before starting the antibiotic. One approach is to lightly touch a culture medium–moistened cotton-tipped swab against the ulcer and then streak standard culture media. If the ulcer is chronic or the patient is immunocompromised, a fungal organism may be the causative agent. Finally, a saline-moistened cotton-tipped swab may be used to obtain a Gram stain of the ulcer. Initiation of therapy before obtaining specimens for culture makes subsequent identification of an organism difficult. For this reason, consider the circumstances of the individual case before initiating treatment.

**TONOMETRY**

Tonometry (Video 62.4) is the estimation of IOP. It is obtained by measuring the resistance of the eyeball to indentation by an applied force. Prolonged elevated IOP is associated with visual field loss and blindness. A sudden elevation in IOP can result from trauma or primary angle-closure glaucoma. Patients with primary angle-closure glaucoma are often seen in the ED with systemic complaints, including nausea, vomiting, and headache. Occasionally, these patients are surprisingly free of pain in or about the eye. The emergency clinician must determine the IOP and its relationship to the systemic symptoms.

Ophthalmologists depended on tactile estimation of eye pressure until the 1860s, when von Graefe developed the first mechanical tonometer. Applanation tonometry was introduced in 1885 by Maklakoff but was not popularized until Goldmann improved the instrument. Schiøtz developed an impression tonometer in 1905 and modified it in the 1920s; this form is still in use today. Aside from modifications in configuration, current tonometers closely resemble the devices popularized by Schiøtz and Goldmann. Pocket-sized tonometers using the principle of the MacKay-Marg tonometer are available. One such device is the Tono-Pen XL (Reichert Inc, Depew, NY). These devices are portable, lightweight, and relatively accurate, with built-in provisions for calibration. They have the advantage of a one-time-use replaceable cover that eliminates concern about the possible transmission of an infectious agent. The Icare (Tiolat Oy, Helsinki, Finland) is a rebound tonometer that has recently appeared for use in clinical practice. This device uses a newer technology that calculates IOP from the rebound of a lightweight disposable plastic covered magnetized metal rod that is bounced off the cornea. Its advantages include portability and lack of need for corneal anesthesia, and correlates closely with the Tonometer-XL. Whereas the Schiotz and Tono-Pen XL tonometers are the most commonly used devices for measuring IOP in the ED, the Icare shows promise to be useful in the ED and primary care settings.

**Tonometric Techniques**

Four tonometric techniques are reliable and clinically useful for estimating IOP:

1. The impression method uses a plunger (3 mm in diameter) to deform the cornea, and the “indentation” is then measured. This technique was popularized by Schiøtz and commonly bears his name.
2. The MacKay–Marg method is a refined version of the impression technique in which smaller amounts of cornea are indented.
3. In the applanation method, a planar surface is pressed against the cornea.
4. Rebound tonometry is an induction-based impact method in which a very lightweight magnetized probe is bounced off the cornea.

The Schiøtz tonometer actually measures total IOP (initial pressure plus the pressure added by the weight of the tonometer and the plunger). Friedenwald empirically found that a “rigidity coefficient” could be introduced to allow an estimation of the true IOP. One must be aware, however, that calculated conversion tables for Schiøtz tonometers use an average estimate of the rigidity coefficient and hence are not accurate when eye rigidity is altered (e.g., after scleral buckle procedures for retinal detachment or with extreme myopia).

Measurement of IOP in the ED by tonometry is a technique available to most emergency clinicians. Tonometry is not a standard procedure for many eye-related complaints, but in the following special situations, tonometry may be particularly helpful:

- Confirmation of a clinical diagnosis of acute angle-closure glaucoma. For example, a middle-aged or elderly patient with acute aching pain in one eye, blurred vision (including halos around lights), and a red eye with a smoky cornea and a fixed midposition pupil needs a pressure reading (Fig. 62.27A). Sometimes the findings are less dramatic, and sometimes the patient complains mostly of nausea and vomiting, which suggests “flu” rather than an eye disorder.
- Determination of a baseline ocular pressure after blunt ocular injury. Patients with hyphema often have acute rises in IOP because of blood obstructing the trabecular meshwork (see Fig. 62.27B). Later, angle recession can cause a permanent form of open-angle glaucoma. Arts and colleagues suggested that an IOP greater than 22 mm Hg or a difference of 3 mm Hg or greater between the eyes is a good marker of “ocular injury” in the setting of an orbital fracture.

Tonometry may also be considered in the following scenarios:

- Determination of a baseline ocular pressure in a patient with iritis. Both open- and closed-angle glaucoma, as well as corticosteroid-induced glaucoma, can develop in patients with iritis. Because most cases of iritis are referred, tonometry may also be deferred unless signs of increased IOP are present.
- Documentation of ocular pressure in a patient at risk for open-angle glaucoma. All patients older than 40 years with a family history of open-angle glaucoma, optic disc changes, visual field defects, and pressure of 21 mm Hg or higher should be referred to an ophthalmologist for further workup. Referral should also be made for patients with suspiciously cupped discs who have normal pressure; some of these patients may have “low-pressure” glaucoma associated with...
Contraindications to Tonometry

Tonometry is relatively contraindicated in eyes that are infected unless one is using a device such as the Tono-Pen XL, which uses a sterilized cover. Sterilize a tonometer before and after applying it to a potentially infected eye. Measure infected eyes with either a noncontact tonometer or a device with a covered tip (e.g., Tono-Pen). Swab the contact portions of any device with alcohol and allow it to dry before use on another eye. Not all viruses are destroyed by cleansing with alcohol. Hydrogen peroxide is effective in deactivating the human immunodeficiency virus (HIV) responsible for acquired immunodeficiency syndrome (AIDS). Ultraviolet sterilization, cold sterilizer bathing of the footplate and plunger, and ethylene oxide sterilization have all been advocated as alternatives to sterilizing the tip of the Schiøtz tonometer. The Schiøtz tonometer may also be used with sterile disposable coverings (marketed as Tonofilm). Nonetheless, defer measurement of IOP in an obviously infected eye until a subsequent visit to the ED or private clinician unless the red eye demands an immediate determination of IOP.

Examples of indications for immediate tonometry in the setting of a red eye are suspected angle-closure glaucoma (acute onset of redness and pain in the eye with smoky vision, a cloudy cornea, and a fixed pupil in mid-dilation, often with headache and nausea) and iritis (ciliary injection with photophobia), in which secondary angle-closure glaucoma or corticosteroid-induced changes in pressure may occur. Reported cases of conjunctivitis spread by tonometry predominantly tend to be viral infections. Particular effort should be made to avoid use of the instrument on patients with active facial or ocular herpetic lesions or those who may have AIDS.

The presence of corneal defects also represents a relative contraindication to tonometry. Use of a tonometer on an abraded cornea may lead to further injury and is commonly deferred until a subsequent visit. Patients who cannot maintain a relaxed position (e.g., because of significant apprehension, blepharospasm, uncontrolled coughing, nystagmus, or uncontrolled hiccup) are unlikely to permit an adequate examination and can sustain a corneal injury when sudden movements occur during an examination. Furthermore, tonometric examination, with the exception of the palpation technique (through the lids) and the noncontact method, should not be performed on a cornea without complete anesthesia.

Tonometry should not be performed on a patient with a suspected penetrating ocular injury. Globe perforation may be exacerbated by pressure on the globe with resultant extrusion of intraocular contents. Slit lamp examination can be used for detection of a possible perforation.

Procedure

Palpation Technique

All forms of tonometry are essentially ways of determining the ease of deforming the eye; an eye that is easily deformed has low pressure. The most direct way to do this is simply to press on the sclera through the lids and grossly compare one eye with the other. One can easily distinguish the rock-hard eye of acute glaucoma from the normal opposite eye by this method. Direct the patient to look down without closing the lids. Rest both hands on the patient’s forehead and apply just enough digital pressure on the involved eye to indent it slightly with one index finger. With the other index finger, alternately feel and compare the compliance of the other eye (Fig. 62.28, plate 1). An experienced examiner is able to estimate IOP within 3 to 5 mm Hg of the actual IOP with the palpation technique, but most emergency clinicians do not have enough experience to trust this method.

Another method is to anesthetize the eyes topically and press a wetted applicator on the sclera of each eye. Again, eye deformation is inversely related to ocular pressure. Rigidity of the globe is also a factor in this crude method of tonometry.

Impression (Schiøtz) Technique

Use of the Schiøtz tonometer requires relaxation on the part of the patient and steadiness on the part of the clinician. After placing the patient in either a supine or a semi-recumbent position, instruct the patient to gaze at a spot directly above the eyes. A spot on the ceiling should suffice; alternatively, patients can stretch their arm up over their head and gaze at
Tonometry: Palpation And Schiøtz Techniques

A relatively unskilled examiner can detect the very high intraocular pressure of acute angle-closure glaucoma with tactile tonometry. The examiner rests both hands on the patient’s forehead and alternately applies just enough digital pressure on the globe to indent it slightly with one index finger while feeling the compliance of the globe with the other.

Before using the Schiøtz tonometer, test it on a flat surface to ensure smooth motion of the device and that the zero line is achieved.

Apply lid separation pressure to the bony orbital rims. An assistant may separate the lids while you concentrate on proper placement of the tonometer. Hold the tonometer vertically during use, and rest your hand against the patient’s facial bones. After anesthetic drops are instilled, the patient will not experience any pain from this procedure. It is important to have a relaxed patient because squinting and blepharospasm may interfere with the reading. Note: Gloves should be worn.

*The table provides estimates of intraocular pressure to the nearest mm Hg for the different weight of the Schiøtz tonometer. Accuracy is most dependable with scale readings greater than 5. If the scale reading is less than 5, use the next highest weight that will give a reading of 5 or more.

Use the above chart to determine the converted reading based on the reading and the amount of weight on the scale.

Figure 62.28 Tonometry: palpation and Schiøtz techniques.

their thumb. Place a drop of topical anesthetic in each eye. After the irritation of the drop passes, allow the patient to blink while blotting the tears away with a tissue. Rubbing the eyes lowers IOP. Reassure the patient that further discomfort will not occur during the procedure.

Ask the patient to keep both eyes wide open and fixed on an object. Separate the eyelids on the side where you are standing. Test the tonometer on a flat surface to confirm smooth movement of the device (see Fig. 62.28, plate 2). Take care to direct pressure onto the orbital rims rather than the orbit.
because pressure directed into the orbit falsely raises the reading. Hold the tonometer momentarily over the open eye, and inform the patient that the instrument will block vision in that eye. Instruct the patient to continue to gaze at the fixation point as though the instrument were not there. After the patient relaxes the involuntary muscle contraction that occurs when the instrument is first placed in the line of sight, gently lower the instrument onto the middle portion of the cornea (see Fig. 62.28, plate 3). This is a painless experience for patients with an anesthetized cornea. Vertically align the instrument with the footplate resting on the cornea; the reading should be in midscale. Should the reading be on the low end of the scale (<5 units), place additional weight on the plunger after the instrument has been removed. Repeat the process as before with the additional weight.

Measure the opposite eye in the same fashion. Use the chart provided to determine the converted reading based on the reading and amount of weight on the scale. Refer to ophthalmology if the converted scale reading is higher than 21 mm Hg (see Fig. 62.28, plate 4). Patients with elevations in IOP of 30 mm Hg or greater require more urgent consultation and initiation of therapy. Associated symptoms or signs of angle-closure glaucoma (primary or secondary) represent an ophthalmologic emergency.

**Errors With Impression Tonometry**

Inaccurate readings can occur with the Schiotz tonometer for a variety of reasons. Falsely low readings may occur if the plunger is sticky. Check plunger motion and the zero point of the tonometer on a firm test button before use. If the plunger is sticky, clean it with isopropyl alcohol and dry it with a tissue. Inadvertently directing pressure onto the orbit when the lids are held open may elevate IOP and provide a falsely elevated reading. The following eye movements have been found to elevate IOP: closing the lids (increases by 5 mm Hg), blinking (increases by 5 to 10 mm Hg), accommodation (increases by 2 mm Hg), and looking toward the nose (increases by 5 to 10 mm Hg). Repeated or prolonged measurements have been found to lower IOP approximately 2 mm Hg. Patients with elevations of angle-closure glaucoma (primary or secondary) represent an ophthalmologic emergency.

**Calibration**

To perform calibration of the Tono-Pen XL (required before use at least once each day), depress and release the activation switch momentarily. The liquid crystal display (LCD) should show a single dashed line “—”. If the device beeps and double dashed lines “— — — ” appear on the LCD, push the activation switch again so that “— — ” reappears. If the previous calibration shows “bAd” on the LCD, a long beep sounds, followed by “CAL” on the LCD. A short beep follows and then the desired “—” is displayed. Once “—” is displayed, hold the probe vertically with the tip pointing straight down. Press and release the activation switch twice in rapid succession. Two beeps will then sound and “CAL” will appear on the LCD. Hold the probe in this position (up to 20 seconds) until a beep sounds and “—UP—” appears on the LCD. Immediately turn the probe 180 degrees so that the tip points straight up. In a few seconds, another beep occurs and the LCD changes. If the LCD reads “Good”, the calibration was successful. If the LCD reads “bAd”, the calibration was unsuccessful.

With an unsuccessful calibration, repeat the calibration steps described earlier until two consecutive “Good” readings are obtained. If further attempts are unsuccessful, loosen the Ocu-Film tip cover and repeat the calibration process. If attempts are still unsuccessful, press the reset button and repeat the process. If still unsuccessful, use compressed air to clean the mechanism and ensure free movement. Place an Ocu-Film (latex; Reichert) cover snugly (but without tension) over the probe tip.

**Measurement**

To proceed to measurement, prepare the patient as outlined earlier. Depress and release the activation switch to obtain “— — — ” on the LCD. A beep will occur when ready. If the switch is not depressed long enough, the LCD will be blank. If a blank screen is seen, press and release the activation switch again to obtain “— — — ” on the LCD. Hold the probe like a pen and touch it to the cornea briefly and lightly. Touch the cornea four times. A click will sound and a reading will appear on the LCD each time that a valid reading is obtained. In the Tono-Pen XL, after four valid readings, a final beep will sound that the devices can be used in patients with irregular or high corneal astigmatism.

Ideally, the complete instructions provided with the device should be consulted before each use; however, the following synopsis is provided to help in circumstances in which instructions are unavailable.

First, spray the tip of the probe with compressed gas to clean the mechanism and ensure free movement. Place an Ocu-Film (latex; Reichert) cover snugly (but without tension) over the probe tip.
and the averaged measurement will appear on the LCD. The
taken bar reflects statistical reliability (a reading > 20% reflects
an unreliable measurement and should be repeated). In the
Tono-Pen AVIA, after depressing the activation button, a green
light and beep signal that the unit is ready for a measurement.
The dual sided LCD display shows “= = =” and the unit can
be operated as described earlier. When 10 readings are obtained
within 15 seconds, the LCD displays an IOP with a statistical
confidence indicator. (Fig. 62.31)

If single dashes (“—”) appear on the LCD after the final
beep, too few valid readings were obtained. In such a case,
reactivate the probe (without recalibration) and repeat the
measurement procedure. If the probe is not reactivated within
20 seconds, the LCD will clear, but the device can be activated
as noted previously without recalibration.

The values are interpreted as outlined earlier for the Schiotz
device. Readings may be affected by the same features noted
as causes of error with impression tonometry via the Schiotz
device. Store the device with an unused Ocu-Film cover protec-
ing the tip of the probe. As with any device, be familiar with
the manufacturer’s instructions for use.

Impression/Rebound (Icare) Technique
When using the Icare tonometer, the preparations for testing
are similar to the Tono-Pen described earlier, except that this
device can be used without topical anesthetic drops. Review
the user’s manual to become familiar with the instructions
before use. The following brief synopsis is provided if instruc-
tions are not available. Various parts of the tonometer are
shown in Fig. 62.32 and referred to in these instructions.

Use of the wrist strap is recommended. Press the measuring
button to turn on the device. When the display shows “LoAd”,
load the single use probe into the probe base, being careful
not to drop the probe out of the tonometer. (Fig. 62.31)

If single dashes (“—”) appear on the LCD after the final
beep, too few valid readings were obtained. In such a case,
reactivate the probe (without recalibration) and repeat the
measurement procedure. If the probe is not reactivated within
20 seconds, the LCD will clear, but the device can be activated
as noted previously without recalibration.

The values are interpreted as outlined earlier for the Schiotz
device. Readings may be affected by the same features noted
as causes of error with impression tonometry via the Schiotz
device. Store the device with an unused Ocu-Film cover protect-

Figure 62.29 Tonometry: Tono-Pen (Reichert Inc, Depew, NY) technique. IOP, intraocular pressure.
Tonometry: Tono-Pen Technique

Wait a few seconds. If the calibration was successful, a beep will sound and “Good” will appear on the LCD screen.

Instill a drop of topical anesthetic (e.g., tetracaine 0.5%) into both eyes.

Hold the Tono-Pen as you would hold a pencil, and brace your hand against the patient’s cheek for stability.

Activate the Tono-Pen by pressing the switch. A beep will sound and “==” will appear on the LCD screen.

Touch the Tono-Pen against the cornea lightly and briefly. Repeat several times; a click will sound every time that a pressure is measured. After four valid readings, a final beep will sound, and the averaged measurement will appear on the LCD.

Check the reading on the screen. The number represents IOP in mm Hg. The bar below the number represents the statistical reliability. (A reading > 20% reflects an unreliable reading and should be repeated.)

Figure 62.30 The Tono-pen AVIA (Reichert Inc, Depew, NY) is designed to fit comfortably in the hand, and has a large dual-sided LCD display.

Figure 62.31 The Tono-pen AVIA (Reichert Inc, Depew, NY) LCD display indicates that the device is ready to measure with double dashed lines (top). During the measuring mode, the display indicates the number of successful measurements (middle). Once 10 measurements are completed successfully, the display will show the IOP (large number) and a statistical confidence indicator (bottom).
Complications

When tonometric instruments are used properly and reasonable precautions are taken, complications are unusual. An eye with preexisting corneal injury should be spared the additional trauma of tonometer placement. Corneal abrasions can be produced or worsened by ocular movement during testing. In particular, patients with uncontrollable nystagmus, hiccups, coughing, or those who are extremely apprehensive should not be subjected to tonometry. Infection can be transmitted by use of the instrument. Careful cleansing of the device and avoidance of tonometry in patients with obvious conjunctivitis, corneal ulcers, or active herpetic lesions should minimize the risk of spreading the infection to the unaffected eye or to subsequent patients. Although protective coverings can be placed over the tonometer contact, tonometry can usually be postponed in the aforementioned individuals until the risk for infection is minimal. Extrusion of ocular contents with penetrating injuries is a potential, but rare complication.

SLIT LAMP EXAMINATION

The slit lamp is an extremely useful instrument for examination of the anterior segment of the eye. The instrument can reveal pathologic conditions that would otherwise be invisible, such as minor corneal defects, anterior chamber hemorrhage, and inflammation.

Indications and Contraindications

The slit lamp can be used in the majority of eye examinations (Video 62.5). It is especially useful in the ED for the diagnosis of corneal abrasions, FBs, and iritis. The slit lamp facilitates
FB removal and is also used in conjunction with most applation tonometers. Portable slit lamp instruments are readily available (Fig. 62.35) but seldom used in the ED; thus, emergency practitioners generally have access only to a stationary, upright device. Therefore in the absence of a portable device, a slit lamp examination is contraindicated in patients who cannot tolerate an upright sitting position (e.g., those with orthostatic syncope).

**Equipment**

The slit lamp has three essential components: a binocular microscope mounted horizontally, a light source that can create a beam of variable width, and a mechanical assembly to immobilize the patient’s head and manipulate the microscope and light source. The location and arrangement of the knobs that control these components vary in devices made by different manufacturers. Usually, by simply turning each knob and watching the results, one can quickly master a new machine. Fig. 62.36 illustrates the location of the functional controls on one particular instrument.

First, locate the on/off switch for the instrument. Frequently, this switch incorporates or is adjacent to a rheostat that provides two or three different power settings. The lowest setting is adequate for routine examination and will preserve bulb life. One can use a high-intensity setting when examining the anterior chamber with a narrow slit beam. Often, these controls are located on a transformer placed beneath the table to which the slit lamp has been attached. The second knob that one should find is the locking nut for the mechanical assembly. Loosen the nut so that the assembly can be moved relative to the stand.

Make adjustments so that the patient is comfortable while sitting with the head in the device. Ask the patient to press the forehead firmly against the headrest with the chin in the chin rest. By varying the height of the table and height of the chin rest, one should be able to maximize comfort of the patient’s neck and back. Adjust the chin rest to align the patient’s eye level with the mark on the headrest support rods.

The binocular microscope has a control for varying the magnification. Usually, low powers such as 10× or 16× are the most useful. Use a higher power to examine the anterior chamber for cells and flare and when the cornea is examined in minute detail. Adjust the binocular interpupillary distance to match that of the examiner. Focus the eyepieces by moving the instrument forward and backward until the narrowed vertical beam is sharpest on the patient’s cornea when viewed with the unaided eye. Then, while viewing through each eyepiece individually, adjust the focus of each to produce a sharp image of the anterior surface of the cornea.

Notice that the light source is mounted on a swinging arm. Find the knobs that adjust the width and the height of the light beam. Click various filters in as needed, usually white and blue filters for standard examination. Alter the angle of the slit lamp beam to vertical from horizontal. The vertical alignment is preferred for routine examinations in the ED.

Both the microscope and the light source are mounted on swivel arms linked at their base to the movable table. Change the position of this table by pushing on any part of it. Use the joystick on the table for finer movements. Vary the height of the microscope and the light source by twisting either the joystick or a separate knob at the base, depending on the design of the instrument.

**Procedure**

There are three setups that every slit lamp operator must know. The first is for an overall screening of the anterior segment of the eye. For examination of the patient’s right eye, swing the light source to your left at a 45-degree angle while the microscope is directly in front of the patient’s eye. Set the slit beam to the maximum height and the minimum width using the white light. To scan across the patient’s cornea, first focus the beam on the cornea by moving the entire base of the slit lamp forward and backward. Then move the whole base of the mechanical apparatus left and right to scan across the cornea. The 45-degree angle between the microscope and the light source is the default position. The most common mistake is to try to scan by swinging the arm of the light source in an arc; this does not work because the light beam will remain centered on the same point of the patient’s eye. Scan across at the level of the conjunctiva and the cornea and then push slightly forward on the base or joystick and scan at the level of the iris. The depth of the anterior chamber is easily appreciated with this low-magnification setup (Fig. 62.37). When the depth of the anterior chamber is reduced, suspect a corneal perforation or a predisposition to angle-closure glaucoma.

Use this basic setup to examine the conjunctiva for traumatic lesions, inflammation, and FBs. Examine the eyelids for hordeolum, blepharitis, or trichiasis. Completely evert the lids (as described in the earlier section on Ocular FB Removal) in conjunction with the slit lamp examination to permit evaluation of the undersurface of the upper lid for FB retention.

Corneal FB removal can be enhanced by use of the slit lamp. In particular, the instrument allows stabilization of the patient’s head. Magnification also minimizes corneal injury during FB or rust ring removal. The upper eyelid may be immobilized with a cotton-tipped applicator, as discussed previously. The clinician’s hand can be steadied against the patient’s nose, cheek, or forehead or against the support rods of the headrest. The patient should be instructed to stare straight ahead at a fixed light or at the clinician’s ear during removal of the FB.

The second setup is essentially the same as the first but uses the blue filter. The purpose is to identify any areas of
Figure 62.36 Slit lamp (BM 900, Haag-Streit, Mason, OH).
**Figure 62.37** Slit lamp photograph of a normal left eye under low power. The curved slit of light on the right is reflected off the cornea and the slit on the left is reflected off the iris. The depth of the anterior chamber can easily be appreciated under this low-magnification setup. Note that the light source is on the patient’s left side to examine the left eye, with the path of the light going in a temporal-to-nasal direction. (Courtesy D. Price.)

Fluorescein staining. After fluorescein is applied, “click” the blue filter into position and widen the beam to 3 or 4 mm. A patient can tolerate a wider beam without photophobia if it is blue. Search for corneal defects (as discussed in the earlier section on The Fluorescein Examination) with this setup. The blue filter may also be used with applanation tonometry, as discussed in the earlier section on Tonometry.

The purpose of the third setup is to search for cells in the anterior chamber, either the white cells of iritis or the red cells of a microscopic hyphema. Shorten the height of the beam to 3 or 4 mm and make it as narrow as possible. Switch the microscope to high power. Focus the beam on the center of the cornea and then push forward slightly so that it is focused on the anterior surface of the lens. Pull the joystick back again to a focus point midway between the cornea and the lens, where it will be focused on the anterior chamber (Fig. 62.38). Keep the beam centered over the pupil so that there is a black background. Normally, the aqueous humor of the anterior chamber is totally clear. Small particles seen floating up or down through the beam are usually circulating cells. If the beam lights up the aqueous like a searchlight in the fog, the examiner has found the protein flare that accompanies iritis. Note that fluorescein can penetrate an abraded cornea and produce a fluorescein flare on slit lamp evaluation. To avoid confusion, some clinicians prefer to examine for anterior chamber flare before the stain is used. A variety of conditions evaluated by the slit lamp are pictured in Fig. 62.39.

**UNILATERAL LOSS OF VISION**

There are a variety of reasons why an individual may sustain complete loss of vision in one eye, but most commonly, such loss is caused by occlusion of the central retinal vein, occlusion of the central retinal artery, or optic nerve damage. Less commonly, pressure on the orbit from a retroorbital hemorrhage may compromise the ophthalmic artery.

Although discussion of all the potential causes of unilateral loss of vision is beyond the scope of this text, amaurosis fugax deserves special mention. Amaurosis fugax is a transient loss of vision that is most commonly due to cholesterol or platelet emboli from atherosclerotic carotid occlusive disease. When plaques are visualized in the retinal vasculature, auscultate the neck for carotid bruits and consider referring the patient for a more thorough evaluation including ultrasound examination of the carotid artery.

**Central Renal Artery Occlusion**

A patient with central retinal artery occlusion (CRAO) has generally experienced a recent sudden painless (complete or nearly complete) unilateral loss of vision. On examination, there is an afferent pupillary defect (APD) (i.e., sluggish or nonreactive pupil in the affected eye with direct illumination and a normal consensual response) and reduced visual acuity. Immediately after the event the fundus may appear nearly normal; however, it soon becomes pale and a classic “cherry-red spot” in the macula may be evident as a result of patent choroidal vessels showing through the transparent fovea (Fig. 62.40).

**Therapy**

Visual recovery has been noted to occur up to 3 days after CRAO. Start treatment if the patient is seen within 24 hours after the onset of symptoms. Consult ophthalmology while initiating therapy. In many centers, CRAO is considered an ischemic event and neurology as well as ophthalmology are often involved in acute management. In some stroke centers, patients with CRAO are admitted to neurology on confirmation of the diagnosis.

Most of the emergency techniques suggested for treating vascular insults to the eye in the ED are theoretically sound but are not supported or refuted by rigorous scientific data. No specific standard of care has been promulgated for these interventions by emergency clinicians. Techniques discussed later are probably safe and possibly useful and may be attempted in an emergency situation. It is unknown whether these interventions will be vision saving.

Slow rebreathing into a paper bag is believed to increase the arterial carbon dioxide level, thus aiding vasodilation, permitting the occlusion to move more peripherally, and possibly...
A more aggressive therapy, generally performed only by ophthalmologists, is anterior chamber paracentesis. In the absence of available consultation, consider this technique when CRAO is recent and unresponsive to the previously described therapeutic approaches. For this procedure, keep the patient supine with the head and eyelids secured. Anesthetize the cornea with topical anesthetic drops (e.g., 0.5% proparacaine drops). Inject the conjunctiva adjacent to the limbus with a 27- or 30-gauge needle until the entire perilimbal area is infiltrated and has the appearance of chemosis in all quadrants. During the remainder of the procedure, ask an assistant to firmly grasp the conjunctiva with toothless forceps at the 3 and 9 o’clock...
symptoms improved or all steps were complete. Of the 11 patients in this arm of the study, eight had improved visual acuity. In those who improved, all had their symptoms improved in 12 hours or less. The presumed cause was either a platelet-derived or cholesterol embolus from atheroma or glaucoma. Although this study is small, it supports emergency ophthalmology consultation and aggressive treatment of patients seen within 12 to 24 hours of the onset of symptoms.

Complications
Overzealous globe massage has the potential to produce intraocular trauma, including retinal detachment and intraocular hemorrhage. Anterior chamber paracentesis may produce hemorrhage, infection, or mechanical injury to the cornea, iris, or lens. Although these complications are rare, ophthalmologic consultation for assistance with the underlying CRAO and surveillance for these potential complications should be initiated on an emergency basis.

Orbital Compartment Syndrome
Acute facial trauma or recent retrobulbar anesthesia may produce retrobulbar hemorrhage with sufficient pressure to compromise the ophthalmic artery and result in an orbital compartment syndrome (Fig. 62.43). A form of posttraumatic glaucoma may also occur when the retrobulbar hematoma forces the globe against the eyelids. In this case, IOP rises precipitously because the globe is in a relatively closed space as a result of the firm attachment of the eyelids to the orbital rim by the medial and lateral canthal ligaments. The optic
The retina will not retain function if it is deprived of blood for a long period.

**Technique: Lateral Canthotomy and Cantholysis** *(Fig. 62.44 and Video 62.6)*

The goals of the procedure are to release pressure on the globe and decrease IOP sufficiently to reinstitute retinal artery blood flow. Because retinal recovery is unlikely to occur if rapid relief of ischemia is not accomplished, taking extra time to clean the eye beyond simple saline cleansing of the lids and lateral canthus must be carefully considered. Stabilize the patient's head and lids and anesthetize the lateral canthus by injecting 1% to 2% lidocaine with epinephrine. Before incising, crush the lateral canthus with a small hemostat for 1 to 2 minutes to minimize bleeding. Incise the canthus with iris or Steven's scissors. Take precautions to avoid injury to the protruding globe. Begin the incision at the lateral canthus and extend it toward the orbital rim. Find the superior and inferior crus of the lateral canthal tendon and release them from the orbital rim. Some operators prefer to release the inferior crus and reassess IOP before considering release of the superior crus.

Although hemorrhage, infection, and mechanical injury might result from the procedure, these complications generally respond to therapy better than retinal injury from prolonged ischemia. Emergency ophthalmologic consultation should be obtained, although when the procedure is indicated, it may be considered by the emergency clinician. Lateral canthotomy incisions generally heal without suturing or significant scarring.

**Complications**

Although hemorrhage, infection, and mechanical injury might result from the procedure, these complications generally respond to therapy better than retinal injury from prolonged ischemia. Emergency ophthalmologic consultation should be obtained, although when the procedure is indicated, it may be considered by the emergency clinician. Lateral canthotomy incisions generally heal without suturing or significant scarring.

**REDUCTION OF GLOBE LUXATION**

Although luxation of the globe is uncommon, the emergency clinician should be aware of the condition and its mechanisms, know how to reduce the globe, and know when to prioritize ophthalmologic consultation. With luxation of the globe there is extreme proptosis, which permits the lids to slip behind the globe equator (Fig. 62.45). Subsequent spasm of the orbicularis oculi muscles sustains the luxation and limits extraocular movement. Traction on the optic nerve and retinal vessels may produce direct or indirect injury to the optic nerve and retina.

Luxation may be spontaneous, voluntary, or traumatic. A variety of conditions (e.g., orbital neoplasms, Graves' disease, histiocytosis X, cerebral gumma, and craniofacial dysostoses) may predispose the patient to luxation. Triggering events include maneuvers that increase IOP (e.g., the Valsalva maneuver), trauma to the orbit or forehead, or eyelid manipulation.

**Indications and Contraindications**

Early globe reduction is indicated to relieve symptoms and minimize visual impairment. Attempts at reduction in the ED are relatively contraindicated when there is obvious rupture of the globe.

**Technique**

Before globe reduction, perform a rapid eye examination to document visual acuity, range of eye motion, pupillary reactivity,
Lateral Canthotomy And Cantholysis

Identify the lateral canthus (arrow). Cleanse the area with antiseptic and anesthetize with 1% lidocaine with epinephrine. (The left eye is depicted in this image sequence.)

Crush the lateral canthus with a hemostat for 1 to 2 minutes to reduce incisional bleeding (not shown). Then, cut through the crushed tissue with iris scissors (as depicted above) to perform the canthotomy.

Pull the lower eyelid away from the globe with toothed forceps (arrow).

“Strum” the tissue under the canthotomy with the scissors to identify the inferior crus of the lateral canthal ligament. Cut through this ligament with scissors to perform the inferior cantholysis. Note that the scissors are directed inferiorly during this step, perpendicular to the canthotomy incision.

The eye after canthotomy and cantholysis. This procedure relieves increased intraocular pressure by allowing the globe and orbital contents to move forward.

NOTE:
If intraocular pressure remains elevated after inferior cantholysis, the superior crus of the lateral canthal ligament may be released in a similar fashion.

Figure 62.44 Lateral canthotomy and cantholysis. (From Eisele OW, Smith RV, editors: Complications in head and neck surgery, ed 2, St. Louis, 2008, Mosby.)
STYE

A stye, or hordeolum, is an acute purulent inflammation (bacterial infection) of the eyelid characterized by pain, swelling, and redness. It can be quite annoying and painful to the touch. A small nodule or abscess first develops in an eyelid hair follicle or a modified sebaceous gland at the margin of the eyelid. This may be external (pointing at the lid margin) or internal (pointing under the conjunctival lid; Fig. 62.46). An obvious pustule may be seen, and if so, incising it with a small needle and expressing pus produces a faster cure. The lid may be inverted to find a small pustule on the inner surface of the lid that can be nicked with a 25-gauge needle (see Fig. 62.23, plates 6 and 7). S. aureus is the organism most frequently isolated from the infection.1 Treat with warm compresses on the eyelid as frequently as possible. One method is to fill a sink with very hot water and alternate wet washcloths for 15 to 20 minutes. Topical ophthalmic antibiotics (drops every 2 hours or ointment five to six times a day) are usually prescribed. Erythromycin ointment is often suggested. Topical treatment is generally sufficient, but antistaphylococcal oral antibiotics (dicloxacillin, cephalosporins) might occasionally be needed, especially if in patients with significant surrounding lid cellulitis. More formal incision and drainage may be necessary if the infection is unresponsive to conservative therapy.118 Spread of the infection can lead to preseptal cellulitis.

APD OR MARCUS GUNN PUPIL

An APD, or Marcus Gunn pupil, is caused by a variety of diseases of the afferent, or “in-going,” pathways of the eye. It is produced by a unilateral lesion of the retina or optic nerve. Causes include optic neuritis (as seen with multiple sclerosis), ischemic optic neuropathy, optic tumor, and retrobulbar hematoma (an indication for lateral canthotomy). To evaluate for an APD, the swinging flashlight test may be used (Fig. 62.47). Normally, shining a light in either eye causes bilateral

and any evidence of globe rupture (see earlier discussion).117 Place the patient in a recumbent position and administer a topical ocular anesthetic agent (e.g., 0.5% proparacaine). When the lashes are visible, ask an assistant to apply steady outward and upward traction while the globe is gently pushed behind the lids. Use gloved fingers to apply steady scleral pressure and to manipulate the globe back into the orbit. When the lashes cannot be grasped, introduce a lid retractor behind the lid to provide countertraction. Others recommend placing a suture through the anesthetized skin of each lid to provide countertraction.

After the procedure, repeat the eye examination to document visual acuity and extracocular movement. It is not uncommon for return of full visual function to be delayed for several days or occasionally longer.

Complications

It is common with this procedure for lashes to be retained in the conjunctival fornices. Evaluate for and remove any free lashes to prevent corneal injury. Edema, retrobulbar hemorrhage, or orbital deformity may prevent outpatient reduction. When reduction is not possible in the ED, saline drops should be applied to the globe and a noncontact eye shield used.

Aftercare

Patients with spontaneous luxation and no visual impairment in whom the globe is easily reduced warrant follow-up within 24 to 48 hours. Instructions to avoid potential triggering maneuvers should be given. Recurrent luxation may warrant lateral tarsorrhaphy. Further evaluation of potential precipitating factors can be pursued on an outpatient basis.

Patients with traumatic luxation are at greater risk for underlying ophthalmic injury and warrant emergency consultation. A computed tomography scan of the orbit is helpful for evaluating both the soft tissue and the bony structures about the globe.

**Figure 62.45** Appearance of a luxated globe. A, This globe protruded when the lower eyelid was retracted downward to examine the conjunctiva for anemia. B, It was easily reduced with slight manual pressure on the closed upper lid. C, Afterward, the patient had no eye or vision complaints. This is a normal variant in some people.
To evaluate for an aPD (no light reaching the brain via the optic nerve on the affected side), record the pupil size at baseline. Shine a light into the affected eye. Record the direct response (constriction of the illuminated pupil in response to light) and the indirect or consensual response (constriction of the opposite pupil in response to light). Next, shine the light into the other eye and record the direct and indirect responses. Repeat this procedure back and forth until the pattern of response to light is identified. With an aPD, there is a decreased direct response to light along the afferent or in-going pathways, whereas the efferent or “out-going” pathways to the opposite eye are preserved. Thus, light shined into the affected eye will cause neither a direct nor a consensual response, but light shined into the unaffected eye will cause bilateral pupillary constriction.

SUBCONJUNCTIVAL HEMORRHAGE

Subconjunctival hemorrhage may occur spontaneously (often noticed on awakening) or after straining, vomiting, or severe coughing. The patient notices a painless bright red hemorrhage of the sclera. The cornea and anterior chamber are unaffected, and there are no visual symptoms.

ACKNOWLEDGMENT

The authors recognize the many contributions by Jerris R. Hedges, MD, to this chapter through the first five editions of the textbook.
The procedures presented in this chapter are most effectively performed with special equipment and techniques (Videos 63.1–63.10). Some are within the realm of general emergency medicine clinical expertise; others are not. They are reviewed from the perspective of the emergency clinician who must decide whether the patient needs treatment acutely in the emergency department (ED), can be managed with timely referral, or requires urgent consultant expertise.

PHARYNX AND LARYNX

Examination of the Larynx

Visualization of the larynx and pharynx is a critical part of the complete evaluation of patients with complaints of sore throat, hoarseness, foreign body (FB), or stridor.

Anatomy

The pharynx is the part of the throat located posterior to the mouth and the nasal cavity and superior to the esophagus and larynx. It is commonly divided into three sections, the nasopharynx, the oropharynx, and the laryngopharynx (hypopharynx). The nasopharynx is the most cephalad portion and extends from the base of the skull to the upper surface of the soft palate. The oropharynx is behind the oral cavity and extends from the uvula to the level of the hyoid bone. Its anterior wall consists of the base of the tongue and the vallecula; its lateral wall consists of the tonsils, tonsilar fossa, and pillars; and its superior wall is formed by the inferior surface of the soft palate and the uvula. Pharyngeal branches of the ascending pharyngeal artery, ascending and descending palatine arteries, and pharyngeal branches of the inferior thyroid artery supply blood to the pharynx. The pharyngeal plexus and the maxillary and mandibular nerves innervate the pharynx.

The larynx in adults is located in the anterior part of the neck at the level of the C3–C6 vertebrae. It connects the inferior portion of the pharynx (hypopharynx) with the trachea. The larynx extends vertically from the tip of the epiglottis to the inferior border of the cricoid cartilage. It consists of nine cartilages. Branches of the vagus nerve, the superior laryngeal nerve, and the recurrent laryngeal nerve innervate the larynx. The relevant anatomy is depicted in Fig. 63.1.

Laryngoscopy Indications and Contraindications

Laryngoscopy is indicated for the evaluation of patients with complaints of dysphagia or odynophagia. More specifically, it should be performed in patients complaining of dysphagia, hoarseness, FB ingestion or sensation in the throat, angioedema, and in patients who require assessment of their airway status. In general, laryngoscopy can be used to evaluate a problem, to exclude airway compromise, and to diagnose several other diseases such as gastroesophageal reflux, cancer, and allergy.

Laryngoscopy has traditionally been discouraged in patients with impending airway compromise; however, it may be performed carefully in stridulous patients as long as a predesignated team (usually consisting of an anesthesiologist, otolaryngologist, or another physician skilled in the management of a difficult airway) is readily available and able to intervene if necessary. Care should be taken to avoid accidental trauma to the laryngopharynx, which may exacerbate swelling and further compromise the airway.

Equipment

The equipment required depends on the type of laryngoscopy procedure performed. For flexible laryngoscopy a standard flexible nasopharyngolaryngoscope, a light source, gloves, a nasal speculum, surgical lubricant, antifogging solution, decongestant spray, anesthetic spray, and a wall suction setup with a Frazier suction-tip catheter are needed (Fig. 63.2, plate 1). Many choices of decongestant are available; however, 0.05% oxymetazoline (Afrin, Bayer, Whippany, NJ) or 0.1% to 1% phenylephrine is commonly used. Lidocaine (4%) is typically used as the anesthetic. A 5% cocaine solution serves as both an anesthetic and decongestant. If nasal spray formulations are not available, medication-soaked cotton pledgets, an atomizer bottle, or a syringe atomizer may be used.

Mirror laryngoscopy requires a curved dental mirror, an external light source (preferably a headlamp), 4 × 4-inch gauze, and an antifogging solution. If antifogging solution is not available, hot water can be used to prevent fogging of the mirror. Anesthetic solution may be required if the patient cannot tolerate the procedure.

Procedure

Flexible Laryngoscopy

Attach the nasopharyngolaryngoscope to its light source and the suction tubing to its port (if available). Ensure that both are functioning properly before beginning. Before inserting the scope, adjust the eyepiece to your visual acuity; it is helpful to check the focus on newsprint or a small object. Review the scope’s directional controls. Examine both nares and choose the more patent one to enter. Anesthetize and vasoconstrict the naris (see Fig. 63.2, plate 2). Because this procedure is irritating, allow enough time for these medications to become effective. You may also anesthetize the pharynx to minimize gagging (see Fig. 63.2, plate 3). Warm the end of the scope in warm water to help prevent fogging. Place the patient in the seated position with the head placed against a headrest in the “sniffing position.” Insert the tip of the lubricated scope just inside the naris. (Some authors recommend a series of soft nasal trumpets to gradually dilate the nasal cavity and allow easier passage of the scope.) Movement of the scope against the inside of the nasal passage may be irritating to the patient. Minimize this sensation by resting your fourth and fifth fingers on the bridge of the patient’s nose while stabilizing and guiding the scope between your thumb and index finger (see Fig. 63.2, plate 4).
While looking through the eyepiece, slowly advance the endoscope past the middle turbinate into the nasopharynx or through the lumen of a nasal trumpet. To clear fogging or mucus off the lens, ask the patient to swallow, wipe the lens against the pharyngeal mucosa, or use the suction. Once the scope is in the nasopharynx, direct the tip inferiorly by using the thumb control near the eyepiece. Use the thumb control to accomplish up and down movements of the scope. Rotate the scope about its axis and then apply thumb control to provide lateral movement and visualization. At this point the base of the tongue and the tonsils will come into view. Slide the scope farther caudad to bring the larynx into focus. Once again, systematically view the patient’s anatomy and function during both respiration and phonation (see Fig. 63.2, plate 6).

If the nasopharyngeal scope will not pass through either naris, pass it through the oropharynx. Properly anesthetize the oropharynx and avoid contacting the posterior portion of the tongue to prevent gagging. A plastic bite block can be used. Alternatively, cut a 10-mL syringe (without the plunger) in half and ask the patient to hold it in the mouth between the incisors (see Fig. 63.2, plate 5). Pass the fragile endoscope through this tube into the oropharynx to prevent accidental biting of the scope.

**Mirror Laryngoscopy**

Otolaryngologists most commonly use mirror laryngoscopy, but it can be used in the ED setting if the necessary equipment is readily available. Clinicians unfamiliar with this method should practice frequently because significant eye-to-hand coordination is required to reflect the light beam off the angulated mirror onto the larynx. When this procedure is properly performed, most patients are able to tolerate it without anesthesia of the oropharynx. Fiberoptic nasopharyngoscopy has largely replaced mirror laryngoscopy in the ED when the equipment is available.

Establish rapport with the patient by explaining how the examination will be performed. Have the patient sit erect in the sniffing position, with the feet flat on the floor and leaning slightly forward. Attach your headlamp and adjust the beam of light (Fig. 63.3). Warm the mirror with warm water to prevent fogging, but check the temperature of the mirror with your hand before placing it into the oropharynx so that the

![Figure 63.1 A, Anatomy of the oropharynx.](image)
Figure 63.1, cont’d  B, Sagittal section of the neck. (Netter illustration from www.netterimages.com. Elsevier, Inc. All rights reserved.)
### Flexible Laryngoscopy

1. **Flexible nasolaryngoscope**
   - Basic equipment required for flexible laryngoscopy. Additional items include anesthetic spray, a nasal speculum, surgical lubricant, and antifogging solution.

2. **Oxymetazoline spray**
   - Anesthetize and vasoconstrict the naris before inserting the scope. Oxymetazoline (e.g., Afrin) spray is an ideal vasoconstrictor. Atomized 4% lidocaine can be used as an anesthetic.

3. **Benzocaine spray**
   - Anesthetize the oropharynx with benzocaine spray or atomized 4% lidocaine to prevent gagging.

4. **Rest your fingers on the bridge of the patient’s nose while stabilizing and guiding the scope between the thumb and middle finger.**

5. **10-mL syringe with end cut off**
   - To pass the scope orally, first anesthetize the oropharynx (with atomized 4% lidocaine or benzocaine spray). Use a plungerless 10-mL syringe with the end cut off as a bite block to prevent accidental biting of the scope.

6. **Laryngeal anatomy**
   - Laryngeal anatomy as seen through the nasopharyngoscope.

---

**Figure 63.2** Flexible laryngoscopy.
Benzocaine (Hurricane spray [Beutlich Pharmaceuticals, LLC., Bunnell, FL] or Cetacaine gargle [Cetylite Industries, Inc., Pennsauken, NJ]) or aerosolized tetracaine or lidocaine may be used. One or two quick sprays of benzocaine into the posterior aspect of the oropharynx is sufficient. Though rare, prolonged or repeated spraying of benzocaine can result in methemoglobinemia. Reassure the patient beforehand that although this may make the throat feel as though it is swelling or paralyzed, in actuality, it is just the numbness that accounts for the sensation. The tendency to gag can also be minimized by having the patient concentrate on breathing efforts and keep the eyes open and fixed on an object in the distance.

Once the patient is anesthetized, repeat the steps described earlier and position the mirror against the soft palate. Rotate the angle of the mirror and systematically inspect the base of the tongue, valleculae, epiglottis, piriform recess, arytenoids, false and true vocal cords, and if possible, the superior aspect of the trachea (see Fig. 63.2, plate 6). Observe for masses, evidence of infection, asymmetry, or FBs. Further evaluate the anterior structure of the larynx and function of the vocal cords by having the patient say “eeee” in a high-pitched voice. This should move the epiglottis away from blocking the view of the larynx and bring the true cords together at the midline (Fig. 63.5).

Complications

There are very few complications with laryngoscopy. Occasionally, it may not be possible to complete the procedure because of a prominent gag reflex or patient apprehension and discomfort. Complications include traumatic abrasions and bleeding anywhere along the path of the laryngoscope or on the soft palate or pharynx if a mirror is used. Epistaxis and hemoptysis are uncommon. In patients with head injury, there is always a slight risk of passing the scope intracranially if a basilar skull fracture has occurred; use of a soft nasal trumpet significantly reduces this risk. Laryngospasm and acute airway compromise can be induced in patients with paraglottic infections.
Peritonsillar Abscess Drainage

**Indications**
Clinical suspicion of peritonsillar abscess (PTA)
- Signs and symptoms of PTA include
  - Asymmetric tonsillar bulging
  - Uvular deviation (away from the bulging)
  - "Hot potato" voice

**Contraindications**
- Severe trismus
- Coagulopathy
- Uncooperative patient

**Complications**
- Failure to completely drain the abscess
- Aspiration of pus and/or blood
- Hemorrhage
- Carotid artery puncture

**Equipment**
- 10-mL syringe with 18-gauge needle and covered with a cut cap
- Scalpel with a No. 11 blade and tape covering all but 1 cm of the blade
- Tongue depressor
- Lidocaine with epinephrine
- Topical anesthetic spray
- Frazier-tip suction
- Kelly clamp

**Review Box 63.1** Peritonsillar abscess drainage: indications, contraindications, complications, and equipment. Use a long 25- to 27-gauge needle and a 3- to 5-mL syringe for local anesthesia and a 10-mL syringe for aspiration. Larger syringes obscure the operator's view of the anatomy.

**Figure 63.6** Anatomy of a peritonsillar abscess. The palatine tonsil and peritonsillar space are identified on the patient's right. A peritonsillar abscess is shown on the patient's left. Note that the abscess can extend medially and displace the uvula. The carotid artery and jugular vein are posterior and lateral to the abscess. Avoid lateral angulation of the aspirating needle and use a needle guard to limit depth of penetration (see Fig. 63.10).

**TONSIL: PERITONSILLAR ABSCESs**

Peritonsillar abscess (PTA), also known as quinsy, is most common during the second and third decades. It is rarely seen in children younger than 6 years and is the most common head and neck abscess in anyone older than 6 years. Treatment of PTA has undergone significant change in the past 100 years and continues to do so at this writing. A myriad of opinions exist on the appropriate treatment method, although most agree that some form of drainage procedure is usually required and should be performed in conjunction with the administration of antibiotics and pain control medication. Three options for surgical drainage include needle aspiration (most common), incision and drainage, and immediate (quinsy) tonsillectomy.

**Anatomy**

Understanding the relative anatomy before attempting to treat PTA is important (Fig. 63.6). The palatine tonsils are located between the anterior and posterior pillars of the throat, bound in a capsule, and covered by mucosa. The superior pharyngeal constrictor muscle defines the lateral wall of the tonsil. Of great importance is the internal carotid artery, which lies approximately 2.5 cm posterolateral to the tonsil.
The abscess is defined as a collection of pus between the tonsillar capsule, the superior constrictor muscle, and the palatopharyngeus muscle (Fig. 63.7). The abscess is not within the tonsil itself. PTA is believed to arise from spread of infection from the tonsil or from the mucous glands of Weber located in the superior tonsillar pole. The abscess is most commonly initiated from the upper pole of the tonsil. However, it can also spread from the middle or inferior poles.

**Pathophysiology and Clinical Findings**

PTAs can occur in patients with inadequately treated tonsillitis and in those with recurrent tonsillitis, but in some patients they arise de novo. There are no data proving that antibiotics, even the correct ones in proper doses, invariably prevent the progression of tonsillitis to abscess formation. The abscess is generally unilateral, and bilateral involvement is rare.

Patients with PTA have a sore throat, odynophagia, low-grade fever, and a variable degree of trismus. The trismus develops secondary to irritation of the pterygoid muscle. The patient may also complain of ipsilateral otalgia. As the abscess expands, the patient may experience dysphagia with drooling. Patients may be dehydrated secondary to poor oral intake. Changes in voice are common (“hot-potato voice”) and caused by transient velopharyngeal insufficiency and muffled oral resonance. Rancid breath is also common. Tender ipsilateral anterior cervical lymphadenopathy is usually present. Although leukocytosis is often present, a complete blood count and other laboratory tests are nonspecific.

The differential diagnosis for this acute process includes unilateral tonsillitis, peritonsillar cellulitis, retropharyngeal abscess, infectious mononucleosis, epiglottitis, herpes simplex tonsillitis, retromolar abscess, neoplasm, FB, and internal carotid artery aneurysm. Chronic conditions include leukemia, carcinoma, and tumor in the parapharyngeal space. Differentiation of PTA from peritonsillar cellulitis may be difficult, especially in the early stages of an abscess. The history and time course of the two disease processes are quite similar. Trismus and uvular deviation are uncommon with peritonsillar cellulitis.

Complications of PTA may include pharyngeal obstruction or extension into the closely approximated neurovascular bundles and parapharyngeal space. Specific complications include airway obstruction, rupture of the abscess with aspiration pneumonia, septicemia, internal jugular vein thrombosis, supplicative thrombophlebitis of the jugular vein (Lemierre’s syndrome), carotid artery rupture, mediastinitis, necrotizing fasciitis, and cardiac and renal sequelae of group A streptococcal infection.

**Indications and Contraindications**

The indication for PTA drainage is usually straightforward, but some advocate that initial medical options exist, especially in children. If obviously present in an adult, the PTA may be drained. However, the difficulty can be in the diagnosis of PTA. As an option to immediate surgical intervention, patients with suspected PTA and no airway symptoms, who do not appear to be in a toxic state clinically, may be admitted to the hospital without imaging for 24 hours of hydration; for empirical intravenous antibiotics to cover group A streptococci, *Staphylococcus aureus*, and respiratory anaerobes (clindamycin is often used); and for analgesia. If a rapid response to medical therapy is not seen, drainage or tonsillectomy should be performed. Approximately 50% of patients, predominately children rather than adults, will respond to medical therapy alone.

It may be difficult to differentiate cellulitis from abscess. To diagnose PTA, in addition to visualization, place a gloved index finger into the mouth to feel for hardness or fluctuance in the peritonsillar region (Fig. 63.8).

Intraoral sonography may augment diagnostic accuracy and direct localization for drainage (Fig. 63.9; see Chapter 66). It is performed with an intracavitary probe. Blaivas and coworkers found that ED ultrasound was effective in diagnosing and aiding drainage in five cases of PTA. Ultrasound excluded the diagnosis in one. If there is still a question regarding the diagnosis or actual location of the abscess, computed tomography (CT) may be helpful but is not regularly performed and is not standard before a drainage procedure in straightforward cases.

There are very few, if any, contraindications to draining a PTA. One contraindication is the absence of a PTA; however, needle aspiration may be performed to confirm the presence of an abscess. Other contraindications can include severe trismus, coagulopathy, and inability of the patient to cooperate with the procedure.
For either procedure, the patient should be given intravenous pain medication and may require mild sedation or even procedural sedation. One should administer parenteral narcotic analgesia, mild sedation, or both, before attempting aspiration. Fentanyl, 1 to 3 µg/kg administered intravenously a few minutes before the procedure, is often ideal. Midazolam may be used judiciously, but the patient should not be overly sedated. The combination of midazolam, ketamine, and glycopyrrolate is reported to be safe and effective for outpatient peritonsillar drainage in children.4

Procedure
Before describing the procedures for drainage of a PTA, one needs to know the clinical circumstances under which to perform the different procedures. A myriad of opinions exist on the appropriate treatment method, although most agree that some form of drainage procedure should be performed in conjunction with the administration of antibiotics and pain control. Three options for surgical drainage include needle aspiration (most common), incision and drainage, and immediate (quinsy) tonsillectomy. Each method is discussed.

Needle aspiration is relatively simple, can be performed by emergency clinicians, does not require special equipment, and is relatively inexpensive. Other benefits of needle aspiration over incision and drainage include decreased pain and trauma. Many believe that this should be the initial surgical drainage procedure for adults and children. The recurrence rate after aspiration is 10%,3 and its cure rate is 93% to 95%.5 Approximately 4% to 10% of patients require repeated aspiration.3,5,6 One drawback is that needle aspiration may miss the PTA and therefore allow misdiagnosis as peritonsillar cellulitis. For this reason, some authors propose admission of patients with negative aspirations and the presumed diagnosis of peritonsillar cellulitis for intravenous antibiotics and observation to prevent further morbidity. Although most studies involved hospitalization and intravenous antibiotics, selected outpatient treatment with oral antibiotics has also been successful and is usually the option chosen unless the patient appears to be in a septic state.6

The incision and drainage procedure is commonly done on an outpatient basis under local anesthesia. It is usually performed after pus is obtained by needle aspiration, but occasionally it is the primary procedure. It seems most logical to first attempt needle aspiration and follow with incision and drainage only if additional pus is suspected or other extenuating circumstances are present. The success rate for incision and drainage is high, with a recurrence rate similar to or lower than that with aspiration alone.5

Treatment guidelines based on a review of the literature3,5–7 suggest that patients with PTA should initially be treated by needle aspiration. Incision and drainage and immediate tonsillectomy should be reserved for treatment failures or recurrences. These procedures can be performed in conjunction with hospital admission and administration of intravenous antibiotics or as outpatient treatment with oral antibiotics. One evidence-based review analyzed 42 articles, 5 of which were clinical studies on surgical technique.5 All three techniques were found to be effective in treating PTA, and the recurrence rate was low (grade C recommendation). The approach depends on the patient’s clinical status and medical history. The emergency clinician makes decisions about the treatment of PTA in the ED, but as local protocols dictate, consultation with an otolaryngologist may be appropriate.
The two procedures described here include needle aspiration and incision and drainage. They should be performed only in a cooperative patient without severe trismus. As the carotid artery is located 2.5 cm behind and lateral to the tonsil, there is minimal room for error, patient movement, or poor anesthesia.

**Needle Aspiration**

Have the patient sit upright with a support behind the head. This is best done as a two-person procedure. Ask an assistant to retract the cheek laterally to maximize visibility. A headlamp provides optimal lighting; a double–tongue blade setup aids in visualization of the operative area (see Fig. 63.3). Administer a parenteral narcotic analgesic, a mild sedative, or both before attempting aspiration.

Use digital palpation to locate the fluctuant area of the abscess. Anesthetize the area topically or with local infiltration. Infiltrate with 2 to 3 mL of 1% lidocaine with epinephrine via a 25- to 27-gauge needle. Use a 3- to 5-mL syringe with a long needle so it is possible to visualize the area to be injected (Fig. 63.10, step 1). A large syringe can block the operator’s view. Displacing the tongue with a finger rather than a tongue blade may provide a better view. Infiltrate the lidocaine intramuscularly for the best results, but be careful not to increase the size of the abscess by direct injection into the abscess cavity. The area should blanch. With proper local infiltration, the patient will not feel the penetration of the aspirating needle. If the trismus is so pronounced that it prevents adequate anesthesia, it will probably be too difficult to aspirate or incise the abscess properly.

Novel techniques to assist in the drainage procedure have recently been described. Afarian and Lin described the use of a laryngoscope with a curved blade (Fig. 63.11A and B). The blade is inserted into the patient’s mouth as far posteriorly as the patient can tolerate. The blade shines light from inside the mouth onto the posterior aspect of the pharynx. In addition, the laryngoscope provides better exposure of the area because the handle is below the patient’s mouth and the holding hand does not obscure the view. Moreover, the curved blade sweeps the tongue out of the way and the weight of the handle will help overcome trismus. Braude and Shalit described a similar process involving the use of a disassembled disposable vaginal speculum with a fiberoptic light (see Fig. 63.11C). One advantage of both techniques is that an assistant can hold either light source without getting in the way of the operator. Finally, Chang and Hamilton reported that performing the procedure with the patient in the Trendelenburg position and the operator seated behind the patient’s head provides comparable success rates and patient comfort.

For abscess aspiration, attach a long 18- to 20-gauge needle to a 10-mL syringe. Fashion a needle guard by cutting off the distal 1 cm of the plastic needle cover, replace the cover on the needle, and securely attach this guard to the needle and syringe with tape to prevent inadvertent displacement (see Fig. 63.10, step 2). Ensure that the needle protrudes only 1 cm beyond the cover to limit the depth of needle penetration and lessen the risk of entering any major vascular structures. If pus is not obtained at a 1-cm depth, deeper penetration is discouraged. Insert the needle into the most fluctuant (or prominent) area as previously determined, which is most commonly the superior pole of the tonsil (see Fig. 63.10, step 3). Importantly, advance the syringe and needle in the sagittal plane only; do not angle to the side toward the carotid artery. Do not aspirate the tonsil itself because the abscess develops in the peritonsillar space surrounding the tonsil. Continually aspirate while advancing the needle in the sagittal plane and do not direct it laterally, where it could injure the carotid artery. If the aspirate is positive for pus, remove as much purulent material as possible. If the aspirate is negative, attempt aspiration again in the middle pole of the peritonsillar space, approximately 1 cm caudal to the first aspiration. If still negative, perform a third and final attempt at the inferior pole (see Fig. 63.10, step 4). Up to 30% of abscesses will be missed if only the superior pole is aspirated. It must be stressed that a negative aspirate does not rule out a PTA.

Usually, 2 to 6 mL of pus is obtained. It is unusual to recover more than 8 to 10 mL (see Fig. 63.10, step 5). Although culture is recommended, the results rarely alter subsequent therapy. When significant amounts of pus are aspirated, the patient usually feels immediate improvement in pain and dysphagia. After the needle is removed, some bleeding will be noted (see Fig. 63.10, step 6). Slight oozing may occur for a few hours, especially if warm water rinses are used. Drainage of pus may continue and is often sensed as a foul taste by the patient. Significant additional drainage of pus may be an indication for repeated aspiration, incision and drainage, or hospital admission.

Some clinicians advise a formal incision and drainage procedure if frank pus is obtained, whereas others now accept needle aspiration (with close follow-up) as the definitive initial treatment. Combined aspiration and formal drainage in the same visit may be indicated if large amounts of pus are obtained (> 5 to 6 mL) or if pus continues to drain from the aspiration site. There are no agreed standards regarding the best practice for this issue.

A new device, the Reciprocating Procedure Device (RPD; AVANCA Medical Devices, Inc., Albuquerque, NM), has been developed that allows one-handed aspiration of the abscess (Fig. 63.12). The RPD consists of two syringe barrels and plungers. The plungers are linked by a pulley system in opposing fashion, which results in a set of reciprocating plungers. When one plunger is pressed with the thumb, the syringe injects; when the accessory plunger is depressed, the syringe aspirates. The RPD allows stable finger positioning and finer control of the needle and syringe. To use the RPD, attach a needle to the RPD. Press the injection plunger with the thumb while advancing the RPD simultaneously in the oral cavity until the needle penetrates the abscess. Once the mucosal surface has been penetrated, depress the aspiration plunger to provide a vacuum for aspiration without moving the needle tip. Studies have shown that the RPD allows enhanced needle control, safer and more accurate aspiration procedures, and decreased complications by 35% to 60%.

**Incision and Drainage**

To incise a PTA, anesthetize the area as described earlier. Prepare a No. 11 or 15 scalpel blade by tapering over all but the distal 0.5 cm of the blade to prevent deeper penetration (Fig. 63.13, step 1). Incise the area of maximal fluctuance or the area where a preceding aspiration (if one was performed) located pus. Do not incise the tonsil itself; instead, incise the peritonsillar area where pus accumulates. Incise the mucosa in an area 0.5 cm long in a posterior-to-anterior direction. A stab incision with a No. 11 blade usually suffices (see Fig. 63.13, step 2). Warn the patient that the pus will flow posteriorly and must be expectorated. Expect bleeding because this is a
Use a 25- to 27-gauge needle to inject the mucosa with 1–2 mL of lidocaine with epinephrine; observe for blanching. Often, the tongue is best displaced with the finger rather than a tongue blade. Topical anesthetic spray (e.g., Hurricaine) may be used prior to injection.

Use a long 18- to 20-gauge needle for aspiration. As a safeguard to prevent overpenetration of the needle, remove the plastic needle guard and cut off the distal 1 cm. Then, replace the cut-off guard on the needle (arrow) and secure it to the hub.

While applying continuous suction, direct the needle in the sagittal plane only (directly anterior to posterior), not to the side toward the carotid artery. Aspirate the superior pole initially. If pus is not obtained from the superior pole, next try the middle pole. The inferior pole may be aspirated last if needed. Note that the tonsil itself is not aspirated.

Usually, 2–6 mL of pus is obtained. It is unusual to recover more than 8–10 mL. If substantial amounts of pus are obtained, the patient usually notices immediate improvement in symptoms.

After needle removal, some bleeding will be noted (arrow). Slight oozing of blood and pus may continue for several hours. Significant additional drainage may be an indication for repeated aspiration, incision and drainage, or hospital admission.

Figure 63.10 Drainage of a peritonsillar abscess: needle aspiration technique.
discharged with 24-hour follow-up. Toxic patients, those with excessive volumes of aspirate, those with persistent bleeding, and those unable to take oral antibiotics are candidates for admission or more prolonged observation. Frequent rinses with warm saline are quite helpful in relieving postaspiration symptoms.

Complications of Surgical Drainage

Needle aspiration is an accepted, safe, and effective technique for treatment of PTA in the ED. There is an approximate 10% failure rate and need for subsequent drainage. Complications can include aspiration of pus or blood and hemorrhage. If the patient has cellulitis, the aspiration will be of no help, but it will not worsen morbidity. Failure to obtain pus should prompt high-dose antibiotics and reevaluation in 24 hours. Many clinicians will opt for admission in such instances. Though often feared, injury to the carotid artery has not been reported as a complication of needle aspiration of PTA. Catastrophic hemorrhage may result from the extremely rare and largely theoretical aspiration of a pseudoaneurysm mimicking a PTA or similarly rare necrosis of the carotid artery. In addition, incisions that are too large or too small may lead to poor healing or an inability to completely evacuate the abscess, respectively.

Antibiotic Therapy

After drainage, empirical oral antibiotics are used in outpatients and intravenous antibiotics are administered to inpatients. PTAs are often polymicrobial, and hence culture of aspirated pus is prudent, although the results rarely affect subsequent treatment. The predominant bacterial species are *Streptococcus pyogenes* (group A streptococcus), *S. aureus* (including in rare cases methicillin-resistant *S. aureus* [MRSA]), and various respiratory anaerobes (including *Bacteroides*, *Fusobacterium*, *Prevotella*, and *Veillonella* species). *Haemophilus* species are found occasionally. Aerobes and anaerobes may be recovered simultaneously if appropriate culture techniques are used. Appropriate oral antibiotics for outpatient therapy in areas where *S. aureus* remains susceptible to methicillin include amoxicillin-clavulanate or clindamycin. For inpatients, administer ampicillin-sulbactam or clindamycin. Penicillin plus metronidazole is also a recommended antibiotic regimen. In septic patients, it is prudent to also cover for MRSA. Although clindamycin may be adequate for MRSA, some clinicians add vancomycin or linezolid to the aforementioned initial antibiotic regimens if MRSA is suspected or predominant in the population. It is prudent to initially cover for MRSA in septic or very toxic patients.

Glucocorticoid Therapy

It is common practice for clinicians to administer short-term corticosteroids, such as intravenous dexamethasone or methylprednisolone, as an adjunct to other therapies in an attempt to provide relief from the symptoms of PTA. Evidence of the benefits of glucocorticoids in the management of PTA is inconsistent, but such intervention may hasten improvement of symptoms in adolescent and adult patients treated with needle aspiration and intravenous antimicrobial therapy. The benefits may be marginal, but as there are no significant complications, it is reasonable to empirically treat adults with short-term corticosteroids. A single intramuscular dose of dexamethasone (10 mg) is a common protocol. Additional data are necessary before the routine use of glucocorticoids can be recommended in the management of PTA in children.

EAR

Anatomy

The ear consists of three sections, the outer, middle, and inner ear. The outer ear includes the pinna (auricle), the external
EAC extends from the head to the external auditory meatus in the skull and measures approximately 2.5 cm in adults. It is relatively short and straight in early infancy but begins to take on its adult S-shape and overall anterocaudal orientation at 2 years of age. Initially, the EAC is almost entirely cartilaginous, but by adulthood its medial two thirds is composed of bony support with an overlying thin, stratified squamous epithelium. The lateral third has a less sensitive, thicker hairy epithelium that produces cerumen and retains its cartilage as support. The arterial supply to the EAC originates from the external carotid artery via the posterior auricular, maxillary, and superficial temporal branches. The mandibular branch of the fifth cranial nerve (V3) and the vagus nerve innervate the ear.

Other important anatomic considerations include two natural narrowings in the EAC, which are important when considering FBs. One is located at the junction of bone and cartilage and the other lies just lateral to the TM. A blind spot may occur in the tympanic sulcus (inferior and anterior to the TM) because of the oblique orientation of the TM. An examiner using a simple otoscope may not visualize an FB in this sulcus.

Anesthesia of the External Ear

Auricle

Indications for local anesthesia of the auricle include closure of extensive lacerations or performance of other painful procedures such as incision and drainage of hematomas (Fig. 63.14). Four nerve branches supply the external ear; knowledge of their anatomy is required to understand the location for injection of anesthetic (see Fig. 63.14B). The great auricular nerve (branch of the cervical plexus) innervates most of the posteroomedial, posterolateral, and inferior aspect of the auricle. A few branches of the lesser occipital nerve may contribute to this area. The auricular branch of the vagus supplies the concha and most of the area around the auditory meatus. The auriculotemporal nerve (from the mandibular branch of the trigeminal nerve) supplies the anterosuperior and antero- medial aspects of the auricle.
Procedure

Fill a 10-mL syringe with either 1% lidocaine or 0.25% bupivacaine. Mix with epinephrine if a regional block is planned in an area without evidence of traumatized vascularity. Attach the syringe to a 25- or 27-gauge needle (5 to 7 cm in length). One of several methods may be used to induce partial or complete anesthesia, depending on the area of concern. To anesthetize the nerve branches of the great auricular and lesser occipital nerve branches, inject between 3 and 4 mL of anesthetic into the posterior sulcus (Fig. 63.16, plate 1). Insert the needle behind the inferior pole of the auricle and gradually aspirate and inject toward the superior pole along the crescent-shaped contour of the posterior aspect of the auricle. Anesthetize the auriculotemporal nerve anteriorly by placing 3 to 4 mL of anesthetic just superior and anterior to the cartilaginous tragus. Provide anesthesia to the auricular branch of the vagus nerve and the more central areas of the auricle by using the technique shown in Fig. 63.16, plates 3 and 4.

Figure 63.15 Closure of extensive lacerations of the ear such as this require a nerve block of the auricle.

Anesthesia of the Ear

Auricular Field Blocks

1. Inject anesthetic along the course of the posterior sulcus.

One method of auricular block uses approximately 3–4 mL of anesthetic injected both at the posterior sulcus (red arrow) and at a point just anterior to the tragus (blue circle).

External Auditory Canal Blocks

3. Four-quadrant field block. Inject anesthetic subcutaneously in the four quadrants of the lateral portion of the ear canal. Use the largest speculum that will fit to guide the injections. Withdraw the speculum, tilt it toward each of the four quadrants, and insert the needle subcutaneously (x). Inject 0.25–0.50 mL of anesthetic to produce a slight bulge in the soft tissue. A total of 1.5–2.0 mL of anesthetic is usually sufficient to anesthetize the ear canal and permit painless removal of a foreign body. Ketamine procedural sedation may facilitate the procedure.

Figure 63.16 Anesthesia of the ear.
Another and possibly more effective option is the regional block shown in Fig. 63.16, plate 2. Insert the needle subcutaneously approximately 1 cm above the superior pole of the auricle and direct it to a point just anterior to the tragus. Be sure to inject the skin of the scalp while avoiding the auricular cartilage. Aspirate and then slowly withdraw the needle while injecting anesthetic until the needle is almost to the puncture site. Redirect the needle posteriorly and repeat the process while aiming at the skin just behind the mid-auricular area. Remove the needle and perform the same procedure, but insert the needle just inferior to the insertion of the ear lobe and anesthetize it in a superior direction. Again, block the auricular branch of the vagus as described in Fig. 63.16, plates 3 and 4, if additional anesthesia of the concha is required.

Use caution if adding epinephrine to the anesthetic solution when performing regional nerve blocks of the ear, especially if the blood supply has already been traumatically reduced. Do not include epinephrine when directly infiltrating wounds of the auricle because restriction of blood flow through the end-arteries may result in tissue necrosis. Other complications related to local anesthesia and regional blocks of the head and neck are reviewed elsewhere in this text.

EAC and TM

The EAC is innervated by the auricular branch of the vagus nerve (inferiorly and posteriorly) and by the auriculotemporal nerve (superiorly, anteriorly, and inferiorly). The primary indication for local anesthesia of the auditory canal is for removal of FBs, including débridement of otitis externa or removal of significant impacted cerumen. It is nearly impossible for even the most cooperative patient to be able to tolerate even minimal manipulation of the auditory canal or TM. It is very difficult to achieve adequate anesthesia of the inner ear and TM for painful procedures. Simply stated, no easy and completely effective procedure consistently works well. If total anesthesia is required, general anesthesia, especially in children, is often the only alternative. Ketamine is an ideal agent for short procedures, especially for children with foreign objects in the ear. Topical anesthetics are inadequate because of their poor absorption through the rather impermeable and keratinized epithelial surface of the EAC. The editors advise instilling 4% cocaine solution to fully fill the ear canal of the supine patient (affected ear facing upward), and waiting a full 20 minutes for anesthesia of the TM to occur. Some anesthesia of the canal is also achieved with topical cocaine. Though effective for some procedures, injection of local anesthetics into and around the auditory meatus is quite painful and often difficult to perform in a struggling and uncooperative patient. Certain instances warrant adjunctive use of procedural sedation. Auralgan, a combination of benzocaine and other ingredients, may provide analgesia for painful earaches secondary to otitis, but it does little to benefit painful procedures.

Procedure

For local anesthesia use a 25- or 27-gauge needle (3 to 5 cm in length) attached to a syringe containing 1% lidocaine with epinephrine. A 1:10 mixture of 8.4% sodium bicarbonate to lidocaine helps reduce pain during injection in this sensitive area. Place a speculum just inside the auditory meatus, inject 0.3 to 0.5 mL of the anesthetic into subcutaneous tissue, and stop after a small bulge is raised in the skin. Inject all four quadrants in this manner by moving the speculum after each injection (see Fig. 63.16, plate 3). If additional anesthesia is necessary, give two more small injections. Inject the same amount slightly farther into the canal, once along the anterior wall and again at the posterior wall at the bone-cartilage junction.

Another similar technique involves depositing the anesthetic just lateral, or exterior, to the external auditory meatus. Using the same size of needle and type of anesthetic solution as just described, inject approximately 0.5 to 1.0 mL into each of five points around the auditory meatus and tragus (see Fig. 63.16, plate 4).

Examination

Several methods can be used to examine the EAC and TM. In all methods, grasp the superior aspect of the pinna and pull cephalad and posterior to straighten the slightly tortuous EAC. Examination is most commonly done with a fiberoptic otoscope (Fig. 63.17). Place a plastic or metal speculum into the auditory meatus for examination and use a headlamp or head mirror/lightbulb as a light source. After inspection, the operating hand can be used to pass instruments into the EAC and to maneuver them more easily. Although this technique provides excellent illumination, the use of magnifying loupes can improve visualization during procedures. The ideal setup for removal of cerumen or an FB consists of an operating microscope and a speculum. This provides binocular vision and frees the examiner’s hands for instrumentation. Unfortunately, this equipment is seldom found outside the otolaryngology clinic setting. If using a standard otoscope, stabilize the hand holding the otoscope against the temporal part of the patient’s skull to prevent inadvertent injury to the canal if the patient moves unexpectedly.

Removal of Impacted Cerumen

Excretions from the ceruminous or apocrine and sebaceous glands, together with cells exfoliated from the EAC, combine

---

**Figure 63.17** Examination of the ear canal. A, Retract the pinna in a superior and posterior direction to straighten out the ear canal (arrow). Hold the scope in the other hand and stabilize it against the patient’s head. This prevents inadvertent injury if the patient moves unexpectedly. B, The pinna can be retracted and the scope held with a single hand that is also resting against the patient’s head (arrow). This technique is useful if instruments are to be passed through the otoscope.
# Cerumen Impaction Removal

## Indications
Symptomatic cerumen impaction
Sudden hearing loss in the setting of cerumen impaction
Evaluation of otitis media in the setting of cerumen impaction

## Contraindications
Few if any absolute contraindications
Contraindications to irrigation:
- Patient aversion to irrigation
- History of middle ear disease
- History of ear surgery
- Perforated tympanic membrane
- Severe otitis externa
- Narrow ear canals

## Complications
- Otitis externa
- Tympanic membrane perforation
- Middle ear injury
- External auditory canal trauma

## Equipment
- Ear scoop
- Alligator forceps
- Frazier-tip suction
- Syringe and Teflon catheter for irrigation
- Otoscope
- Ceruminolytic

## Procedure
Removal of cerumen can be accomplished by irrigation, manual extraction, or a combination of both. Generally, the procedures used to remove cerumen are safe; however, otologic injury has

---

**Review Box 63.2** Cerumen impaction removal: indications, contraindications, complications, and equipment.

---

*Figure 63.18* A, Extensive contact dermatitis from the use of neomycin-containing ear drops. B, Malignant otitis externa in a diabetic patient developed after the patient manually removed cerumen with a cotton swab and a pencil, but it may develop without manipulation of the canal.

- Narrow ear canals.
- FBs, especially sharp objects and vegetable matter.
- Uncooperative patient.
- Occluding aural exostoses.
- Known inner ear disturbance, especially if the patient has severe vertigo.
- History of radiation therapy encompassing the external or middle ear, base of the skull, or mastoid.

---

Cerumen is composed of lipids, complex proteins, and simple sugars. Cerumen repels water, has documented antimicrobial activity, and forms a protective barrier against infection. Cerumen often becomes impacted, which results in complaints of a “blocked” ear, sudden-onset impaired hearing, or dizziness.

**Indications and Contraindications**
Symptomatic impaction is an indication for removal, although symptoms are rare until complete obstruction is present. Sudden loss of hearing is a common complaint in patients with totally occluding, impacted cerumen. Cerumen obstructs visualization of the TM and can be evacuated as a part of the evaluation of a febrile child or a patient complaining of ear pain. However, removal of cerumen in a child is rarely indicated in the ED simply to visualize the TM.

There are few, if any, true contraindications to removal of impacted cerumen. Cerumen is usually impacted for prolonged periods, and vigorous attempts to remove it may precipitate otitis externa. It is reasonable to instill antiseptics (acetate acid otic solution) (VoSol [ECR Pharmaceuticals Co., Inc., Richmond, VA] and others) or antibiotic ear drops for a few days after removal of the cerumen to prevent otitis externa. Neomycin-containing ear drops are best avoided because of precipitation of a contact dermatitis (Fig. 63.18A). Caution should be used in removing impacted cerumen in diabetic patients. Diabetics commonly experience otitis externa after seemingly minor manipulation of the ear canal (see Fig. 63.18B).

Contraindications to irrigation include the following:

- Patient aversion to or a history of injury from previous syringe irrigation.
- History of middle ear disease.
- History of ear surgery.
- Known or suspected perforated TM.
- Severe otitis externa.
occurred after this “minor” procedure and has even resulted in litigation.\(^\text{17}\)

Irrigation is an effective approach for removal of cerumen and has the advantage of being painless and simple to perform. It is usually most successful after the instillation of a ceruminolytic (see later). Because the patient does not have to remain completely still, it is ideal for the pediatric population. It is estimated that 150,000 ears are irrigated in the United States each week.\(^\text{17}\) Though usually more time-consuming and messy than manual extraction, irrigation is an appropriate initial method to attempt and can be performed by technicians with guidance from the clinician.

A 2004 evidence-based review concluded that the current evidence suggests little difference in the efficacy of water-based and oil-based preparations for removing cerumen.\(^\text{18}\) Non–water-, non–oil-based preparations appear to be most effective in clearing cerumen and improving syringing, but further research is needed.\(^\text{18}\) Whichever of the following techniques are used, some tips for successful removal of cerumen include proper lighting, attention to patient comfort, and abrupt cessation when the patient's comfort level is breached.

**Ceruminolytics**

These products may soften hardened or impacted cerumen. They are used as adjuncts to other procedures; simply instilling ceruminolytics into the canal will not remove enough cerumen to aid the emergency clinician. If irrigation fails, continued outpatient use of ceruminolytics is often prescribed, usually combined with home irrigation via a bulb syringe or a repeated visit in a few days. Although many products are available as ceruminolytics, a 5% or 10% solution of sodium bicarbonate disintegrates cerumen much more quickly and efficiently than commercially prepared ceruminolytics and other products.\(^\text{19}\)

Cerumenex (Purdue Frederick Company, Stamford, CT), Cerumol (Thornton and Ross Ltd., Huddersfield, United Kingdom), Auralgan, Buro-Sol, alcohol, and oils were all tested and took more than 18 hours to disintegrate cerumen versus approximately 90 minutes for the sodium bicarbonate solutions.\(^\text{19}\) Cerumenex and Cerumol have been since removed from the market due to otic cell toxicity. Hydrogen peroxide is another commonly used ceruminolytic, but its use has not been systematically studied. One study found that the liquid preparation of the stool softener docusate sodium (Colace, Purdue, Stamford, CT) was much more effective than Cerumenex as a ceruminolytic.\(^\text{19}\) An evidence-based review of agents found that docusate sodium administered 15 minutes before irrigation was most effective in facilitating removal of cerumen. Triethanolamine (Cerumenex) and olive oil were the next most effective treatments.\(^\text{21}\)

Place the patient in the supine position with the affected ear up, instill the solution, and wait at least 15 to 30 minutes before attempts at removal (Fig. 63.19, step 1). Repeat the instillation between attempts at manual extraction or irrigation.

**Irrigation (Ear Syringing)**

After a ceruminolytic has been instilled and left in the canal for 15 to 30 minutes, irrigation of the canal is often effective in flushing out impacted cerumen. Ask the patient to sit upright and hold an emesis or ear irrigation basin flush tightly against the skin just below the earlobe. Insert the irrigation tip into the EAC only as far as the cartilage-bone junction, and direct the stream of water superiorly to wash the impacted cerumen away from the TM (see Fig. 63.19, step 2). It is important to warm the water to body temperature to prevent caloric stimulation. Multiple attempts may be necessary, and intermittent attempts at manual removal of loosened cerumen may help hasten the process. During the irrigation, ask an assistant to apply traction to the pinna to straighten the canal for more efficient irrigation. Patients usually feel some discomfort with forceful irrigation, but not severe pain.

Attach a 30- to 60-mL syringe to a 19-gauge or larger butterfly device, cut off the needle and wings, and use the resultant tubing for irrigation. A large plastic or Teflon intravenous catheter (16- or 18-gauge with the needle removed) can similarly be affixed to a syringe.

The most common way to irrigate an ear is with a syringe and catheter. The use of oral jet irrigators (Waterpik, Fort Collins, CO) is another accepted method, but a syringe-catheter setup is readily found in the ED and unlikely to generate enough pressure to cause injury. After irrigating the EAC, apply several drops of isopropanol to the EAC to facilitate evaporation of residual moisture. Do not use isopropanol if the TM is ruptured. Furthermore, topical steroid-containing suspension drops (ciprofloxacin/hydrocortisone) may be soothing after prolonged irrigation. Because severe otitis externa can develop in diabetics after irrigation, some clinicians routinely prescribe antibiotic ear drops (e.g., fluoroquinolones) for a few days after irrigation in high-risk patients.

**Manual Instrumentation**

Manual instrumentation is more advantageous because it is usually quicker and the examiner may more easily remove hardened or larger concretions of cerumen under direct visualization. However, it is difficult to manually remove cerumen without causing significant pain, so irrigation is preferred. Manual removal may be the initial procedure in some cases, followed by irrigation when the cerumen is partially disrupted. Place the diagnostic or operating head of the fiberoptic otoscope or a speculum as a protective port through which instruments are passed and manipulated (see Fig. 63.19, step 3). An operating microscope works best in this situation but, again, is not usually available. To prevent startling or agitating an already anxious patient, allow the patient to experience the sensation of an instrument in the canal by first placing it softly against the wall of the ear canal.

Instruments used for removal of cerumen include flexible plastic or wire loops, right-angle hooks, suction-tipped catheters, or plastic scoops (see Review Box 63.2 and Fig. 63.19, steps 4 to 6). The spoonlike instruments and irrigation are both more effective in removing softer cerumen. Firm cerumen is ordinarily more easily withdrawn with loops or right-angle hooks. Gently tease the cerumen off the canal wall with loops and then pass hooks or loops around the cerumen and withdraw it slowly. Take care to keep both hands in contact with the patient’s head because any sudden movement may cause trauma to the canal or the TM.

**Complications**

Complications from removal of cerumen are rare but can have serious consequences. Although complications from ear syringing are more common with jet irrigators, they may occur with any method of ear irrigation and include otitis externa, TM perforation, or middle ear injury from a preexisting defect in the TM. If the patient is experiencing sudden pain, tinnitus, hearing loss, nausea, or vertigo, stop the irrigation and examine the TM. If the membrane is ruptured, give prophylactic oral
Cerumen Impaction Removal

1. Instill a ceruminolytic solution into the ear at least 15 minutes before attempts at removal (see text for discussion of various agents). Repeat the instillation between attempts.

2. Irrigate with body-temperature water via a syringe with a plastic IV catheter (with the needle removed). Apply traction to the pinna during irrigation to straighten the ear canal. Periodic attempts at manual removal of loosened cerumen may be beneficial.

3. For manual removal, use the otoscope (with the view piece retracted) as a port for passage of the instrument.

4. Gently tease the cerumen off the canal wall with a loop or plastic ear scoop. This should be performed only under direct visualization.

5. Remove dislodged pieces of cerumen from the canal with an instrument such as alligator forceps.

6. A suction-tip catheter may also be used. However, this should be done under direct visualization only to prevent iatrogenic injury to the fragile ear canal.

Figure 63.19 Cerumen impaction removal. Place liquid ducolax or other ceruminolytic into the canal for 20 to 30 minutes prior to wax removal. Most wax is removed with irrigation alone, but multiple irrigations may be required. When removal of cerumen is unsuccessful in the emergency department, discharge the patient with a ceruminolytic and attempt irrigation again in 24 to 48 hours. IV, Intravenous.
antibiotics for otitis media, keep the ear canal perfectly dry with cotton, and refer the patient to an otolaryngologist. This complication is usually benign.

Complications from manual extraction most commonly occur when inadvertent contact is made with the thin, friable skin of the bony canal. Trauma may cause EAC laceration, otitis externa, or perforation of the TM.

**Ear Canal Débridement and Wick Placement**

Débridement of the ear canal and wick placement are essential components in the management and treatment of otitis externa or “swimmer’s ear,” an acute inflammation of the skin of the EAC. This is essentially a cellulitis of the ear canal. Precipitants of otitis externa include water exposure and trauma. Excessive moisture in the canal raises the pH and removes the cerumen. Keratin cannot absorb water, thereby creating a medium for bacterial growth. Trauma, especially self-manipulation with FBs (e.g., cotton swabs, fingernails), causes abrasions in the ear canal and introduces infection. Removal of cerumen by water irrigation is a well-recognized risk factor for the development of otitis externa.

Diabetics and other immunocompromised patients, especially human immunodeficiency virus–positive patients, are susceptible to malignant (necrotizing) otitis externa, a life-threatening form of otitis externa caused by *Pseudomonas* (see Fig. 63.18B). Deep tissue necrosis, osteomyelitis, intracranial extension, and systemic toxicity are hallmark features. Malignant otitis externa is difficult to treat and the mortality rate can be as high as 53%. The diagnosis of malignant otitis externa should be considered in a diabetic or immunocompromised patient with significant symptoms who fails to respond to initial outpatient treatment.

**Indications and Contraindications**

In patients with suspected otitis externa, attempts should be made to clean debris from the canal to aid in healing. In addition, wick placement will be helpful in delivering antibiotic medications to the swollen canal. It has been touted that the key to successful treatment is adequate removal of canal debris. However, vigorous attempts to remove debris on the first visit are frequently painful, of unknown value, and often eschewed.

Removal of debris is contraindicated in cases of suspected malignant otitis externa and otolaryngology consultation is required.

**Procedure**

Because of patient discomfort and canal swelling, use small swabs (e.g., urethral swabs) to gently remove debris. Irrigate the canal gently, but realize that many patients will be cured without extensive débridement. Perform irrigation only in the canal gently, but realize that many patients will be cured.

**Complications**

There are very few complications with canal débridement and ear wick placement. Usually, the wick will fall out as edema in the canal subsides; if not, it may become an FB of the EAC. Care should be taken to not injure the already friable skin of the EAC, which can lead to bleeding and possible infection or cellulitis of the EAC when placing or removing a wick.

**FBs in the Ear Canal**

Despite its small size, the EAC may play host to numerous types of FBs. Living insects account for most FBs found in adults. Children frequently place food (e.g., peas, beans), organic matter (e.g., grass, leaves, flowers), and inorganic objects (e.g., beads, rocks, dirt) into their ear canals during play, and they often fail to admit this to parents. Button batteries may cause significant tissue destruction in a matter of hours, and it is vital to immediately obtain otolaryngology consultation for removal if the button battery is not easily extracted. Symptoms of FB retention usually consist of ear pain, fullness, or impaired hearing in adults; pediatric patients may not be encountered until an associated otitis externa with a purulent discharge has developed.

As described previously, the anatomy of the EAC predisposes to entrapment of FBs in either a lateral or a deeper position. Removal of more medial objects can be much more painful, and anesthesia is usually required. Even the most cooperative patient may become difficult after feeling pain during manipulation of the ear canal. It is probably impossible to adequately immobilize the head of an uncooperative awake child and to delicately extract an FB.
SECTION XI  OPHTHALMOLOGIC, OTOLARYNGOLOGIC, AND DENTAL PROCEDURES

Ear Wick Placement

Cotton or 0.25-inch Nu-Gauze packing strip can be coated with Cortisporin Otic cream and placed into the external auditory canal with alligator forceps.

An easier solution is to use a Merocel ear wick. Place the dehydrated wick into the canal and apply Cortisporin Otic drops to it. Instruct the patient to place the drops directly on the wick until the follow-up visit.

Figure 63.20 Placement of an ear wick. Ear wicks are used for the treatment of otitis externa and function to deliver antibiotic medications to the swollen external auditory canal.

Some authorities claim that injecting local anesthetics makes extracting FBs even more difficult because of soft tissue distortion, although swelling should be minimal if proper amounts of anesthetic are used. Anesthesia of the EAC may be difficult to achieve. Topical anesthetics have a partial effect, and a four-quadrant technique may not produce complete anesthesia, especially of the TM (see Fig. 63.16, step 3). The editors advise filling the ear canal with 4% cocaine and waiting 20 minutes for topical anesthesia of the canal and TM. Procedural sedation (preferably an analgesic-sedative combination or dissociative anesthetic) can aid in the removal of FBs in a distraught child by preventing further struggling and potential canal trauma. Ketamine is an excellent anesthetic for simple FB removal in the outpatient setting. The care provider must weigh the inherent risks related to procedural sedation against those of general anesthesia and the cost of hospital admission.

Adequate visualization of the object is needed for successful removal. One study showed that canal lacerations occurred in 48% of patients in whom removal was attempted without a microscope and in 4% when it was used. The otoscope is the traditional ED instrument for viewing FBs in the ear canal. It is less likely to be useful for retrieval because it is difficult to insert the instrument through or around the end of the speculum. A specialized ear, nose, and throat (ENT) speculum allows more space for instrumentation. A headlamp provides a good light source and leaves both hands free. Magnifying loupes also provide hands-free magnification.

Indications and Contraindications

In the majority of cases, removal of an FB should be attempted in the ED. Before initiating removal, the clinician should set realistic limits on the number of attempts to be made. Even the best clinician can become too aggressive as frustration builds with failed attempts to extract the object. Early consultation with an otolaryngologist should not be considered a failure with difficult FBs. Indeed, with the proper equipment and experience, most objects can be removed atraumatically.

Severely impacted objects and concomitant EAC infection are relative contraindications to removal, and ENT consultation should be obtained. In addition, patient cooperation is essential for successful removal. The only other contraindication to removal of FBs from the EAC involves certain circumstances when irrigation will be the technique used. Irrigation should not be performed in cases of known or suspected TM rupture (tinnitus, vertigo, significant hearing loss, or bleeding from behind the object). In addition, irrigation should not be performed if the object is soft or if it is a seed or other vegetable or organic matter because the water may cause the object to swell.

Procedures

Make a judicious effort to remove an ear FB in the ED setting. Avoid prolonged traumatic attempts because this often terrifies the patient, complicates subsequent attempts, and can cause bleeding and swelling, thus making subsequent efforts more difficult. Most approaches to removal of FBs are anecdotal and found in the literature as case reports or case series rather than as prospective clinical trials. Familiarize yourself with several techniques because the most appropriate choice depends on the size, shape, consistency, and depth of impaction of the object.

Irrigation is the least invasive method; the techniques were explained in detail earlier in this chapter (see section on Removal of Impacted Cerumen and Fig. 63.19). Irrigation works particularly well with small rocks, dirt, or sand that lie deep in the canal next to the TM.

Suction-Tipped Catheters

This technique works well with objects that are round and difficult to grasp. Suction is readily available in the ED but should provide 100 to 140 mm Hg of negative pressure to be
useful. To avoid iatrogenic injury, inform the patient of the impending noise to prevent sudden movements caused by a startle reflex. Place either the blunt or the soft plastic tip against the object and withdraw it slowly. If using a suction instrument with a thumb-controlled release valve (as with the Frazier suction tip), remember to cover the port to activate the suction.

The Hognose (IQDr, Inc., Manitou Springs CO), a commercially available device designed by an emergency clinician, aids in the removal of FBs in the auditory canal. It is used in combination with an otoscope and suction setup. It is essentially an otoscope speculum with suction attachment and a soft self-molding tip that can attach to objects. The flange comes in three color-coded sizes: 4, 5, and 6 mm. To use, first attach the Hognose to the otoscope and set the standard wall suction at a low to medium vacuum setting (Fig. 63.21). Next, under direct visualization, approach the FB with the otoscope. Finally, engage suction by applying finger pressure on the open insufflation port and withdraw.

**Manual Instrumentation**

This approach can be attempted with various instruments (Fig. 63.22, plate 1). Use the diagnostic or operating head of a fiberoptic otoscope for illumination and magnification. Ask an assistant to hold the pinna back and out so that you can hold the otoscope with one hand and manipulate the instrument with the other. A speculum and either a headlamp or a head mirror/flat plate lamp will also provide adequate light. Magnifying loupes are usually required for adequate visualization. Use small alligator forceps to remove objects with edges that can be grasped, but avoid trying to encircle an impacted round FB because this may cause trauma to the canal wall. A small right-angle hook is another choice. Place the tip past the object, rotate it 90 degrees, and then pull the object from the canal. Fine tissue or Adson forceps, curets, and skin hooks are other instruments used occasionally. Use of these instruments is commonly associated with abrasions and bleeding of the ear canal.24,25 Instruments should be used only on compliant, cooperative patients. Direct visualization of the object is essential.

**Fogarty Catheters**

Small Fogarty catheters (biliary or vascular) may be used in a manner similar to that described later in this chapter, in the section on Nasal FB Removal. Attach the tip of the catheter to a 3-mL syringe. Pass the catheter beyond the FB. Once the tip is past the object, gradually inflate the balloon and drag the FB out along with the balloon. Immediately deflate the balloon if pain suddenly occurs because rupture of the TM is a potential complication.

**Cyanoacrylate (Superglue)**

The use of glue to remove FBs was first reported in India in 1977.24,25 Glue is most effective in removing smooth, round objects that are difficult to grasp (see Fig. 63.22, plate 2). The FB should be dry and easily visualized. Apply a small amount of glue to the tip of a thin paintbrush, a straightened paper clip, or the blunt end of a wooden cotton-tipped applicator. Allow the glue to become tacky. Place the tip against the object, allow it to dry, and then carefully withdraw the FB. Minor complications are possible if the tip dries against the canal wall (abrasion, excoriation) or if the glue spills or drips onto the wall (thereby creating a new FB). This technique may be more useful in adults because cooperation is required.26

**Removal of Insects**

Cockroaches are the most commonly found live insects in the ear (see Fig. 63.22, plate 3). A suggested treatment is to instill various substances into the ear canal to immobilize or kill the insect before removing it. This helps in retrieval by allowing a stationary target and also halts the disturbing and painful movement of the insect. Controversy exists about which agent can most effectively accomplish this task. Mineral oil has traditionally been used, but lidocaine has been reported to paralyze insects and allow easier extraction than with the more viscous mineral oil. An in vitro comparative study showed that immersion in microscope oil versus 2% or 4% lidocaine solution killed roaches in less than 60 seconds (=27 and 41 seconds, respectively).27 The roaches struggled less in the viscous oil than in the lidocaine, which did not appear to cause paralysis. Other substances (isopropanol, water, succinylcholine, hydrogen peroxide) were shown to be ineffective in killing the roaches in a reasonable amount of time. Once disabled, insects are removed by mechanical extraction as described previously; pieces can be suctioned out if fragmentation occurs.

If an insect cannot easily be removed or if parts remain, it is quite acceptable to prescribe outpatient antibiotic drops for 24 to 48 hours and try irrigation again or refer to a specialist. As with any FB in the ear canal, prolonged attempts at removal in the ED are counterproductive.

**Complications**

Hearing should be evaluated before and after removal of an FB, especially in patients with suspected TM or middle ear injuries. Also examine the opposite ear and the nose of children to search for the rare but possible second FB. Minor lacerations or excoriations of the canal usually heal quickly with or without antibiotic ear drops, as long as the canal is kept clean and dry. Document preexisting canal trauma or suspected TM rupture before attempts at removal; otherwise, this may be falsely attributed to iatrogenic causes at a later date. Indications for otolaryngology referral include failed removal of the object in the ED, existent injury to the EAC or TM, TM rupture, EAC infection, object wedged in the medial part of the EAC or up against the TM, glass or other sharp-edged FBs, and special circumstances (disk batteries and putty). Generally, no routine follow-up is necessary except in
Indications and Contraindications

Diagnosis of an auricular hematoma is based on the history and physical examination (Fig. 63.23). The presence of tender, anterior auricular swelling following trauma that deforms the anatomy of the pinna should prompt drainage of the hematoma. The goal is to prevent cartilage damage and deformation of the pinna.

If the patient is initially seen more than 7 days after injury, drainage may be difficult because of the formation of granulation tissue, and these patients should be referred to an otolaryngologist. In addition, patients with concomitant cellulitis, perichondritis, or recurrent and chronic hematomas should be referred to a specialist.

Ear Canal Foreign Body Removal


2. Putting superglue on a stick and allowing it to attach to an FB may be successful. This technique is most effective in removing smooth, round objects that are difficult to grasp. The FB should be dry and easily visualized. A high degree of patient cooperation is required.

3. Removal of an insect can be a disaster or relatively easy. An easy way is to use ketamine anesthesia with careful direct extraction. No patient can cooperate with more than minimal manipulation of the inner ear canal. Instilling mineral oil or lidocaine into the canal to kill the insect before extraction is recommended.

Figure 63.22 Removal of a foreign body from the ear canal. Avoid excessive and prolonged attempts in the emergency department to remove difficult FBs. For difficult-to-extract insects, it is appropriate to prescribe outpatient antibiotic drops for 24 to 48 hours and again try to irrigate the insect or refer to a specialist. Adequate anesthesia is essential because no patient can fully cooperate with painful deep ear canal manipulation. FB, Foreign body.
Auricular Hematoma Evacuation

Auricular hematomas manifest after trauma with anterior auricular swelling. The hematoma forms between the cartilage and perichondrium and will lead to permanent deformity if not treated.

1. Anesthetize the pinna via local infiltration of 1% lidocaine without epinephrine or via an auricular block.

2. Use a scalpel to incise the skin along the natural skin folds at the edge of the hematoma. Follow the natural curve of the pinna. Gently peel the skin and perichondrium off the hematoma and underlying cartilage. Completely evacuate the hematoma and irrigate the pocket with normal saline.

3. A compression dressing must be placed to prevent reaccumulation of the hematoma. Place dry cotton into the EAC and fill all external auricular crevices with saline-soaked or Vaseline gauze.

4. Carefully conform the material into all the convolutions of the auricle. Vaseline gauze, saline-soaked 1/4-inch packing gauze, or saline-soaked cotton may be used. Place gauze behind the ear.

5. Place three to four layers of gauze behind the ear as a posterior pack, and then cover the entire ear with multiple layers of fluffed gauze.

6. Secure the dressing to the head with Kerlix or an elastic wrap.

Figure 63.23 Evacuation of an auricular hematoma. EAC, External auditory canal.
**Procedure**

Treatment of an auricular hematoma is complete evacuation of the subperichondrial hematoma and approximation of the perichondrium to the cartilage.

**Needle Aspiration**

Though used widely, this technique is no longer recommended by most authorities because of the high risk for reaccumulation of the hematoma. Aspiration is often inadequate and other management is required. Other sources recommend needle aspiration followed by incision and drainage if reaccumulation occurs.28

Aspiration of an auricular hematoma is performed by perforating the hematoma with a 20-gauge needle. “Milk” the hematoma between the thumb and forefinger until the entire hematoma is evacuated. Apply a pressure dressing. Reexamine the ear frequently for reaccumulation of the hematoma. Reaccumulation of blood requires respiration. For small hematomas that are acute, needle aspiration alone with a bolster dressing is adequate therapy.28

A newer variation of needle aspiration has been described by Brickman and coworkers.29 In their method, after preparation and local anesthesia, needle aspiration was performed using an 18-gauge angiocath and 10-cc syringe. After the collection was aspirated, the syringe and needle were removed, leaving the plastic catheter in place. A compressive dressing was then applied and patients were closely followed up. The catheter was removed in 5 days. Of the 53 patients enrolled, 94.3% initially had adequate hematoma drainage with excellent long-term results. In the initial 72 hours, six patients had catheter dislodgement, with hematoma reaccumulation in three patients (5.6%). These were successfully drained by respiration and there were no further recurrences.

**Incision** (see Fig. 63.23)

An auricular hematoma may be incised along the natural skin folds. Anesthetize the pinna by local infiltration of 1% lidocaine (without epinephrine) or with an auricular block (described earlier). Incise the skin with a No. 15 blade at the edge of the hematoma and follow the curvature of the pinna. Gently peel the skin and perichondrium off the hematoma and underlying cartilage. Completely evacuate the hematoma and irrigate the remaining pocket with normal saline.

After removing the hematoma, apply antibiotic ointment and reapproximate the perichondrium to the cartilage with a pressure dressing. A compression dressing, noninvasive or surgical, must be applied because a simple dressing will allow the hematoma to reaccumulate. First, dry cotton should be placed into the EAC. Next, fill all external auricular crevices with mineral oil, saline-soaked gauze, or Vaseline gauze. Then place three to four layers of gauze behind the ear as a posterior pack. Cut a V-shape out of the gauze first to allow a snug fit. Cover the packed external ear with multiple layers of fluffed gauze. Bandage the fluffed gauze in place with Kerlix or an elastic wrap.

A surgical compressive dressing involves sutureing dental rolls over the area. To accomplish this, pass a 4-0 nylon suture through the entire thickness of the ear and over the hematoma. Wrap the suture around a dental roll on the posterior aspect of the ear and then pass the needle back through the pinna. Wrap and tie the suture around a second dental roll on the anterior aspect of the pinna. A second suture may be placed to secure a third dental roll. The dressing should firmly reapproximate the perichondrium to the cartilage without compromising the vasculature. Remove the dressing in 1 week. The procedures are compared in Table 63.1.30

Prescribe antistaphylococcal antibiotics and instruct the patient to inspect the wound frequently for evidence of vascular compromise, infection, or both. Reevaluate the wound in 24 hours for recurrence of the hematoma. Treat infection by removal of the bandage, surgical drainage, and intravenous antibiotics.

**Complications**

There are a few complications associated with management of auricular hematomas. The first is incomplete evacuation of the subperichondrial hematoma...
of the hematoma, which can lead to destruction of cartilage (cauliflower ear). The hematoma can reaccumulate. Local site infection can develop and result in the development of chondritis. Finally, scar formation leading to deformity is possible.

**NOSE**

**Anatomy**

The nose consists of the vestibule, nasal septum, lateral wall, and nasopharynx. The vestibule is the anterior-most portion of the nares and is composed of skin and hair follicles. The nasal septum is the midline structure and is composed of cartilage anteriorly and bone posteriorly. The lateral wall of the nose contains the superior, middle, and inferior turbinates, as well as the auditory tube opening.

Three major arteries supply the nose and conjoin via anastomoses. The sphenopalatine artery emerges from the sphenopalatine foramen, which is located at the posterior aspect of the middle turbinate (Fig. 63.24). This is the most common source of posterior epistaxis. This artery supplies the lateral turbinates and the posterior septum. The anterior and posterior ethmoidal arteries branch off the ophthalmic artery and penetrate the cribriform plate to supply the superior nasal mucosa. The superior labial branch of the facial artery completes the triad and supplies the nasal septum and vestibule. The watershed area on the anterior septum, also known as Kieselbach’s plexus, is the most common source of anterior epistaxis (Fig. 63.25).

**Anesthesia of the Nose**

Most nasal anesthesia can be accomplished topically. Numerous preparations are available. Cocaine (4% solution) is the preferred agent for both vasoconstriction and anesthesia. Unfortunately, cocaine is not routinely stocked. Alternatively, 2% lidocaine with epinephrine (local anesthetic solution) may be used but is less effective. Be aware of the total amount of lidocaine being administered and stay within recommendations for the maximum safe dosage. This is an issue primarily with elderly patients who have cardiovascular disease. Lidocaine 4% is also quite effective for anesthesia of the nose. A solution of 1% tetracaine and 0.05% oxymetazoline (Afrin, Bayer) is an effective topical anesthetic and vasoconstrictor.

Apply the local anesthetic and vasoconstrictor to cotton swabs (Fig. 63.26). If a larger area of anesthesia is needed, use cotton pledgets. Fig. 63.27 describes the procedure of making pledgets. Soak each pledget in an anesthetic or vasoconstrictor and then squeeze the excess fluid out of the pledget. Place each pledget horizontally on the floor of the nasal cavity and stack the next pledget on top. Three pledgets are usually required to pack the nasal cavity. They can be replaced with new pledgets in 5 minutes if the desired anesthetic effect is not achieved. Benzocaine (Hurricane, Beutlich Pharmaceuticals) spray may also be used as a topical anesthetic. Remind the patient that excess anesthetic may numb the throat but will not inhibit swallowing.

![Figure 63.24 Vascular supply to the lateral wall of the nose. The most common site of posterior epistaxis is the sphenopalatine artery as it emerges posterior to the middle turbinate. (From Maceri DR: Epistaxis and nasal trauma. In Cummings CW, editor: Otolaryngology—head and neck surgery, ed 2, St. Louis, 1993, Mosby–Year Book, p 728.)](image)

![Figure 63.25 Vascular supply to the nasal septum. The most common site of anterior epistaxis is within the area labeled Kieselbach’s plexus. (From Maceri DR: Epistaxis and nasal trauma. In Cummings CW, editor: Otolaryngology—head and neck surgery, ed 2, St. Louis, 1993, Mosby–Year Book, p 728.)](image)

![Figure 63.26 Placement of local anesthetic in the nose to block the anterior ethmoidal nerve superiorly and the sphenopalatine ganglion at the posterior end of the middle turbinate before reducing a nasal fracture. Cocaine is the preferred agent. (From Schuller DE, Schleuning AJ, DeMaria TF, et al, editors: DeWeese and Saunders otolaryngology: head and neck surgery, ed 8, St. Louis, 1994, Mosby, p 152.)](image)
A local nasal block can be performed for more painful and prolonged procedures. The sensory innervation of the nose is illustrated in Fig. 63.28. It begins with topical anesthetic as just described. Next, using a thin 25- or 27-gauge needle, inject 1% lidocaine with 1:100,000 epinephrine along the septum, lateral walls, and nasopalatine nerves. Depending on the procedure, it may be necessary to anesthetize both sides. First, inject along and beneath the soft tissue of the nasal dorsum (infratrochlear nerve). Next, inject in the area of the infraorbital foramen to anesthetize the infraorbital nerve. Finally, inject at the base of the columella (base of the nose between the inferior and the middle turbinates, and the third in the roof of the nasal cavity and the anterior nasal vestibule. Interestingly, studies have shown that the use of EMLA (eutectic mixture of local anesthetics) cream applied over the nose 1 hour before reduction, in combination with topical intranasal anesthetic, provides similar nasal anesthesia for reduction of nasal fractures with less discomfort than occurs with needle infiltration.32

**Examination**

Examination of the nares is relatively straightforward, albeit often quite stressful to the patient. When using a nasal speculum, insert it into the naris with the handle parallel to the floor and slowly open the blades in a superior-to-inferior direction. It is a common error to attempt to spread the nares horizontally. Stabilize your hand on the patient’s nose to prevent damage to the mucosa from unexpected movement (Fig. 63.29). When attempting to visualize the nasal passageway, remember to have the patient keep the floor of the nose parallel to the ground. Tilting the head allows a view of only the anterosuperior area. A nasopharyngoscope may be used to view the nasal passageways as well, and its use is described in the previous section on examination of the pharynx.
large-bore intravenous line for administration of fluid boluses. Hematologic testing is rarely useful and not required for most patients, but in extenuating circumstances, obtain a complete blood count and consider a type and screen. Coagulation studies are not routinely indicated but should be undertaken in patients taking anticoagulants, those with underlying hematologic abnormalities, or individuals with recurrent or prolonged epistaxis.34

Many patients with epistaxis are hypertensive as well, often transiently secondary to stress and anxiety. No direct causal correlation between hypertension and overt epistaxis has been proved. The hypertension that is so often seen with epistaxis is probably a stress response instead of an inciting event. In a hypertensive patient, one can administer intravenous opioids (such as morphine) to relieve stress and anxiety, thereby usually also lowering blood pressure. In most cases, hypertension does not require treatment until evaluation after the bleeding is controlled and the anxiety of the situation has resolved. However, any patient exhibiting other signs of a true hypertensive emergency needs immediate antihypertensive treatment in addition to control of the epistaxis.

Epistaxis is rarely a manifestation of a nasopharyngeal neoplasm, unknown blood dyscrasia, nasal FB, nasal cocaine use, chronic use of nasal corticosteroid sprays, or aneurysm of the carotid artery.

Anticoagulated Patients With Epistaxis
Anticoagulated patients are at high risk for nosebleeds. However, the need for cessation and reversal of oral anticoagulants (i.e., warfarin) is controversial and not well studied. It is reasonable to continue warfarin when hemostasis is achieved and the international normalized ratio (INR) is in the intended therapeutic range. An excessive INR calls for temporary cessation of warfarin and reversal if markedly abnormal. Methods to reverse newer, non–vitamin K oral anticoagulants (i.e., dabigatran, rivaroxaban) are also elsewhere in the text. Specific factor replacement will be required in patients with hemophilia.

Additional Testing for Epistaxis
Although an evaluation of coagulation parameters (prothrombin time, INR, platelet count) is not standard for patients

Management of Epistaxis
Patients with nasal hemorrhage are commonly seen in the ED and account for approximately 1 in 200 visits.33 Epistaxis occurs more frequently in the young (<10 years) and old (70 to 79 years). Most cases are traumatic and occur in the winter months. Approximately 6% require hospitalization.34 Identification of the source of bleeding and subsequent control are paramount to the treatment of epistaxis. Although this can be frightening to both the clinician and patient, a systematic approach with the proper equipment will lessen the anxiety associated with the situation (Fig. 63.30). The goal of the procedure is to tamponade or cauterize the bleeding site. If the source is anterior, this may be the final treatment. For posterior bleeding, temporizing maneuvers are generally used until the process stops or a consultant can complete a definitive hemostatic procedure. The procedures can be performed in the ED with proper lighting and the equipment listed later in this section. Controlling epistaxis may be a time-consuming process without proper equipment or patient cooperation.

In preparation for any procedure to treat epistaxis, evaluate the patient's hemodynamic status by assessing vital signs and orthostatic symptoms and by quantifying the amount of blood lost. If the patient is symptomatic in any of these areas or if the blood loss is deemed significant, consider starting a
Epistaxis Management

**Indications**
Persistent epistaxis

**Contraindications**
Massive facial trauma with the possibility of a basilar skull fracture

**Complications**
Cautery
Nasal septal injury/perforation
Rebleeding

Anterior packing
Sinusitis
Nasolacrimal bleeding
Nasal mucosa pressure necrosis

Posterior packing
Infection
Dysphagia
Eustachian tube dysfunction
Tissue necrosis
Dislodgment

**Equipment**
- chair with a headrest or gurney with an inclinable back
- headlight with a light source and head mirror
- wall suction with multiple suction catheters
- gloves, mask, and gown for the clinician
- gown or drapes for the patient
- topical anesthetic
- topical vasoconstrictor
- nasal speculum
- tongue depressors
- small red rubber catheters
- bayonet forceps
- scissors
- kidney basin
- gauze (4 × 4 inch, 2 × 2 inch), dental rolls or cotton, No. 2 surgical silk ties
- pediatric Foley catheters (12 Fr)
- nasal tampons
- dual-balloon pack

**Review Box 63.3** Epistaxis management: indications, contraindications, complications, and equipment.

With epistaxis, these studies should be ordered routinely for anticoagulated patients. In addition, if the patient has chronic persistent bleeding, an evaluation of coagulation is prudent. A complete blood count is suggested for those with prolonged or recurrent bleeding to assess for significant blood loss. Though a rare consideration in the ED, epistaxis is a significant problem for patients with hereditary telangiectasia, von Willebrand’s disease, and hemophilia. Generally, these conditions are already known. CT or magnetic resonance imaging of the nasopharynx may be considered for elderly patients with recurrent epistaxis to evaluate for neoplasm.

**Indications and Contraindications to Treatment of Epistaxis**
Any continuing episode of epistaxis can be treated with the techniques listed in the section on Procedure. Massive facial trauma with the possibility of a basilar skull fracture would preclude the use of an intranasal balloon or packing because it may travel into the skull cavity.

**Procedure**
Because most patients are frightened by continued epistaxis and because nasal instrumentation can be annoying or painful, provide reassurance to the patient that the bleeding can be controlled with minimal discomfort. Judicious use of parenteral sedation or narcotic analgesia is well supported to make the entire interaction more palatable to the patient and ultimately more successful. Drape the patient with a gown to protect clothing from the bleeding. Have the patient hold an emesis basin to collect any continued bleeding and as a precaution to emesis of swallowed blood. Minor anterior bleeding is usually easily controlled with minimal techniques.

Ask the patient to sit upright in the sniffing position with the neck flexed and the head extended (Fig. 63.31). The base of the nose should remain parallel to the floor. After putting on a face shield, gown, and protective gloves, position yourself in front of the patient, level with the patient’s nose. Have the
Epistaxis Management: Initial Steps

1. Position the patient sitting upright in the sniffing position with the base of the nose parallel to the floor. Proper lighting with a headlamp or mirror is essential.

2. Ask the patient to blow his nose to remove all blood and clots from the nasal passage.

3. Alternatively, gently suction the nasal cavity. Suction from front to back along the nasal septum and then laterally.

4. If bleeding is minimal, attempt to locate the specific bleeding source. Use a nasal speculum to maximize visibility.

5. If bleeding is too profuse for visualization, apply a topical vasoconstrictor such as oxymetazoline.

6. Alternatively, cotton pledgets soaked in cocaine may be inserted into the nasal cavity with bayonet forceps.

Figure 63.31 Management of epistaxis: initial steps.

Patient blow the nose to remove clots or suction the nasal passageway carefully. This may temporarily increase bleeding but can aid in identifying the bleeding site. Suction from front to back along the nasal septum and then laterally. If the bleeding is minimal, attempt to locate the specific bleeding source. If the bleeding is too profuse for visualization of the source, administer a topical anesthetic and vasoconstrictor. One method is to insert lidocaine with epinephrine-soaked cotton swabs into each nare. Ask the patient to clamp the nostrils to limit bleeding and promote contact with the mucosa. If a discrete bleeding site is initially identified, an effective way to provide hemostasis and anesthesia is to inject the mucosa at the base of the bleeder with 2% lidocaine with epinephrine via a tuberculin needle and syringe device.

Insert the nasal speculum into the naris and use the suction catheter to evacuate any blood. Reapply anesthetic or vasoconstrictor if necessary. Because most cases of anterior epistaxis occur in Kiesselbach’s plexus, inspect this area closely for areas of bleeding, ulceration, or erosion. After vasoconstriction, the only evidence of the former bleeding site may be a small prominent vessel. Many clinicians will gently stroke the septum with a cotton swab to initiate bleeding so that an exact area of pathology can be identified and then cauterized. If no bleeding source is found and the bleeding has ceased, pack the nose only if the epistaxis is recurrent. Wait 15 to 20 minutes before deciding on further intervention. If no anterior source is found and bleeding continues down the posterior aspect of the pharynx, assume a posterior source and consider packing the nose with anterior and posterior packs.

Cautery

After an anterior source of bleeding is identified, cautery may be used to achieve hemostasis (Fig. 63.32). Silver nitrate sticks may be used to cauterize but will not work on an actively bleeding source; hemostasis must be achieved first. For brisk bleeding inject lidocaine with epinephrine at the base of the vessel with a tuberculin syringe. Silver nitrate works well for a small, circumscribed area of bleeding. To apply, hold the tip of the silver nitrate stick against the site for 4 to 5 seconds. Apply again if necessary. Wipe away any excess silver nitrate to prevent inadvertent cautery of other areas of the nose. Most patients will sneeze after the application of silver nitrate, so be careful of blood splatter. If bleeding restarts, the initial cautery was insufficient and should be applied again. The cauterized area immediately turns white or gray. Electrocautery works in the same manner but will penetrate more quickly than silver nitrate does. With either cautery technique, be careful to not cause septal perforation with overaggressive or repeated cautery. If cautery has not been successful after two attempts, use another technique.

Multiple attempts at cautery can significantly injure the nasal septum, and bilateral cautery should not be performed.
If this is the initial bleeding and hemostasis is achieved, no packing is necessary. If it is recurrent bleeding within 72 hours of another or if cautery does not provide hemostasis, pack the anterior cavity. If hemostasis is accomplished, apply petroleum jelly or antibiotic ointment to the area to prevent desiccation. Loughran and coworkers found antimicrobial ointment to be better than petroleum jelly in preventing bleeding. Do not administer aspirin or nonsteroidal antiinflammatory drugs for 4 days after epistaxis. If bleeding recurs at home, instruct the patient to pinch the nostrils closed for 20 minutes. Instruct the patient to return to the ED if this maneuver is unsuccessful or if the bleeding is profuse.

**Anterior Nasal Packing**

Anterior packing achieves hemostasis, prevents desiccation, and protects the area from trauma. However, improperly placed packing may further abrade the area, dislodge prematurely, or migrate into the posterior pharyngeal area. Anterior packing must be placed with adequate analgesia, proper visualization, and deliberate movements. Coating any packing material with antibiotic ointment (if not contraindicated by the manufacturer) aids in placement and theoretically helps prevent infections and toxic shock syndrome (TSS) secondary to nasal packing. Areas that continue to ooze after cautery can be treated with an anterior pack.

Easier-to-use commercial devices have largely supplanted traditional petrolatum gauze. Packing is applied in an “accordion” fashion so that each layer extends the entire length of the nasal cavity (Fig. 63.33A). Place the speculum properly to allow visualization of the floor of the nasal cavity. Lay a strip of petrolatum gauze 1.2 cm across the nasal floor, with the starting end of the gauze at the naris. Gently pack the gauze strip into the floor of the nose. Measure the gauze so that it is twice the length of the nasal cavity. Grasp the gauze at the midpoint and insert this point all the way back to the posterior aspect of the nasal cavity. Attempt to place this layer of gauze without movement of the underlying layer. Continue this pattern, with replacement of the speculum after each layer, until the cavity is filled.

When compared with gauze packing, compression devices are easier to place, better tolerated, and very successful. These are preferred over gauze packing by the editors. Preformed nasal packing products are convenient alternatives to anterior nasal packing (Figs. 63.34 and 63.35). The Merocel packing consists of compressed polyvinyl acetate with or without a drawstring that markedly expands on contact with fluid and thereby exerts pressure on the bleeding site. The pack can be trimmed with scissors or a scalpel before insertion. The Merocel Doyle nasal pack (Medtronic) has an airway tube in the center of the compressed material and a more anatomic shape. Each is available in various sizes, but usually an 8-× 1.5-× 2-cm standard Merocel or 8-× 1.5-× 3-cm Doyle will suffice. The Rapid Rhino Stat Pac (ArthroCare ENT, Austin, TX; see Fig. 63.34) is a high-volume, low-pressure balloon device with an open lumen air passage, a pilot cuff to check pressure, and a specialized Gel-Knit (carboxymethyl cellulose) covering designed to promote platelet aggregation. Numerous variations for anterior, posterior, and combination packs are available.

The easily applied nasal tampon is a reasonable first choice for most anterior bleeding (see Fig. 63.33B). Lubricate the tampon generously with antibiotic ointment and trim the length and width carefully to minimize trauma to the nose. Using bayonet forceps, advance the packing carefully along the floor of the nose. Remember to direct it parallel to the floor, not upward toward the top of the nose. Insertion may be painful, so use a single rapid movement. Once the packing is in the nasal cavity, expand it with 5 to 10 mL of saline, although contact with the moisture of the nose will often cause it to swell spontaneously. It is sometimes necessary to place two tampons side by side before inserting them, to fill the nasal cavity and provide better pressure on the areas of bleeding. Observe for 10 minutes after anterior packing to identify continued bleeding either anteriorly from the naris or running down the posterior aspect of the pharynx. Advantages of the Merocel tampon include rapid insertion, little discomfort, ease of use even by inexperienced personnel, and possible inhibition of bacterial growth.
Corbridge and colleagues found no significant difference in efficacy, patient tolerance, or complications between commercial products and gauze packing. Singer and associates found that the Rapid Rhino nasal tampon is less painful to insert and easier to remove than the Rhino Rocket (Shippert Medical Technologies, Centennial, CO) and that both were similarly effective in stopping nosebleeds.

Anterior packs are usually left in place for 3 to 5 days. Premature removal may result in rebleeding. A pack or device

The Rapid Rhino device is first soaked in sterile water for 30 seconds (see Fig. 63.33C). Do not use saline because it can inhibit its gelling characteristics. In addition, lubricants or antibacterial ointments are not needed. Insert the device along the nasal septal floor parallel to the hard palate until the plastic ring is well within the nasal cavity. Use a 20-mL syringe to inflate the device with air only. Stop inflating when the pilot cuff becomes rounded and feels firm to the touch.

Corbridge and colleagues found no significant difference in efficacy, patient tolerance, or complications between commercial products and gauze packing. Singer and associates found that the Rapid Rhino nasal tampon is less painful to insert and easier to remove than the Rhino Rocket (Shippert Medical Technologies, Centennial, CO) and that both were similarly effective in stopping nosebleeds.

Anterior packs are usually left in place for 3 to 5 days. Premature removal may result in rebleeding. A pack or device
may stimulate mucous production and act as an impetus for infection. Oral antibiotics (e.g., cephalexin, amoxicillin, or trimethoprim-sulfamethoxazole) may be prescribed with any nasal packing after emergency treatment because of the minimal risk for sinusitis and TSS. The necessity of antibiotics for short-term anterior packing is unproved. Decongestants are also prescribed to decrease secretions. Practices vary and no common standards exist.

During use and before removal, keep the nasal tampon hydrated with saline. If it contains an airway tube, first remove the tube, irrigate the space once occupied by the tube, and then remove it. To remove the Rapid Rhino, first remove the air and then slide the device out.

**Complications**

Minor oozing of blood can be expected. Any packing in the anterior nasal cavity may obstruct drainage of the paranasal sinuses or block the nasolacrimal ducts and lead to sinusitis. Occasionally, blood will exit the nasolacrimal duct and be noted in the eye (Fig. 63.36). Other complications can include nasal mucosal pressure necrosis from the packing, balloon migration, and aspiration of the packing. Hollis reported massive pneumocephalus after insertion of a Merocel nasal tampon in an elderly woman, presumably from fracture of the ethmoid plate. There have been case reports of ethmoid fracture after anterior nasal gauze packing and with the use of an intranasal balloon.

**Posterior Nasal Packing**

If no bleeding source is found anteriorly and the patient continues to hemorrhage down the posterior aspect of the pharynx, the patient most likely has a posterior source of epistaxis (Fig. 63.37). Posterior epistaxis may respond to topical vasoconstrictors. However, anterior nasal packing will not provide hemostasis for posterior bleeding because it will not cover the source of the bleeding. A posterior pack directly compresses the sphenopalatine artery and prevents the passage of blood or anterior packing into the nasopharynx.

**Posterior Gauze Pack**

A posterior nasal gauze pack is the classic method of treating posterior epistaxis (Fig. 63.38). However, because balloon devices are easier to use and less distressing to the patient, formal posterior nasal packing is less commonly used. To place a formal traditional posterior nasal gauze pack, anesthetize the patient’s nares and posterior pharynx with topical anesthetic. Prepare a roll of gauze with two silk ties (2-0) secured around
the silk ties exiting the nostril to maintain the position of the cheek to aid in removal or rescue of the posterior pack. Use the silk tie that remains in the oropharynx to the patient’s cheek to aid in removal or rescue of the posterior pack. When it is seen in the posterior of the pharynx, grasp it with forceps and guide it out of the mouth. Attach it to one set of the ends of silk ties secured to the gauze pack. Retract the red rubber catheter, thus carrying the No. 2 silk tie through the nasopharynx and out of the nose. Grasp the suture and pull the pack into the nasopharynx. Guide the pack swiftly into the oral cavity and nasopharynx with the other hand. Attach the silk tie that remains in the oropharynx to the patient’s cheek to aid in removal or rescue of the posterior pack. Use the silk ties exiting the nostril to maintain the position of the posterior pack. Pack the anterior passage as described for anterior epistaxis. Secure the silk ties over a gauze pad or dental roll. In the past, patients with traditional posterior packing were often admitted to the hospital for the duration of the posterior packing. Administering humidified air or oxygen often makes this pack more comfortable.

Inflatable Balloon Packs

Inflatable balloons come in two varieties. A Foley catheter is often used as a posterior pack because of its availability, ease of use, and successful tamponading effect (Fig. 63.39f). Insert a 12-Fr Foley catheter through the bleeding naris into the posterior aspect of the pharynx. Inflate the balloon halfway with approximately 5 to 7 mL of normal saline or water. Slowly pull the Foley catheter into the posterior part of the nasopharynx and secure it against the posterior aspect of the middle turbinate. Finish inflating the balloon with another 5 to 7 mL of normal saline or water. If pain or inferior displacement of the soft palate occurs, deflate the balloon until the pain resolves. Ensure proper placement before completely inflating the balloon because the balloon will remain too posterior in the nasopharynx and fail to achieve hemostasis. While maintaining constant gentle anterior tension on the Foley catheter, place anterior nasal packing of layered petrolatum gauze. Pack the opposite nasal cavity to counteract septal deviation. Finally, place a short section of plastic tubing over the catheter and secure it with a nasogastric tube clamp or umbilical clamp. Be careful to not exert undue pressure on the nasal alae to avoid causing necrosis.

The second type of inflatable balloon pack is the premade dual-balloon tamponading system (see Fig. 63.39B). These devices have been a significant advance in the treatment of epistaxis. Several balloon devices are available (Goitschach Nasostat [Sparta Surgical Corp, Hayward, CA], Xomed Epistat [Medtronic Xomed], and Epi-Max Balloon Catheter [Shippert Medical]). The dual-balloon pack has a posterior balloon that inflates with approximately 10 mL of air and an anterior balloon that inflates with approximately 30 mL of air. Each device may vary slightly. After appropriately anesthetizing the naris, place the lubricated pack along the floor of the affected naris as far back as possible. Inflate the posterior balloon approximately halfway with air; and then, with traction, pull the balloon into place up against the posterior aspect of the middle turbinate. Complete the inflation of the posterior balloon with air. Some clinicians prefer to inflate all balloons with saline instead of air because air may deflate slowly. Inflate slowly, and stop if pain is felt. This is usually an uncomfortable sensation to the patient. If the patient complains of pain or if the posterior soft palate deviates downward, deflate the balloon until the symptoms are relieved. Maintain the position of the balloon and inflate the anterior balloon with up to 30 mL of air. Again, halt inflation if the patient experiences increasing pain or deviation of the nasal septum. Some authors suggest packing the opposite naris to prevent such lateral deviation. Place a small piece of gauze between the nose and the external catheter hub to decrease skin irritation.

Most patients with a posterior pack, especially the elderly and those with pulmonary and cardiovascular diseases, should be admitted to the hospital for sedation and monitoring. This recommendation was common for formal posterior packs, but the ease and safety of balloon devices now allow selected patients to be treated as outpatients despite the presence of posterior packing.
After applying topical anesthetic, pass a red rubber catheter through the nose, carefully grasp it in the oropharynx with ringed forceps, and bring it out through the mouth.

Alternatively, fold a gauze pad, roll it into a cylinder, and tie it with two strings. Use two of the long strings to tie the pack to the tip of the catheter, and use the other two to remove the pack.

Remove the optional "retraction" catheter after the pack is in the proper position. Digitally guide the pack into the nasopharynx.

Make a posterior nasal pack by wrapping a cotton ball in a 4 x 4-inch gauze pad and tying two long silk sutures or umbilical tape around the neck of the pack. Leave one tie long so that it can be taped to the cheek until needed for removal of the pack.

As an option, use a second catheter that has been passed through the nonbleeding side and brought out the mouth to retract the palate forward to aid in placement of the pack.

Use a gauze roll to secure the pack to the nose, and tape the rescue ties to the cheek.

Figure 63.38 Management of epistaxis: traditional posterior packing.
Epistaxis Management: Posterior Packing With Inflatable Devices

A  Foley Catheter Technique

1. Insert a 12-Fr Foley catheter through the naris and into the posterior pharynx.

2. Look into the mouth to confirm that the catheter is properly positioned.

3. Inflate the balloon halfway with about 5–7 mL of water.

4. Slowly pull the catheter into the posterior nasopharynx up against the posterior aspect of the middle turbinate.

5. Foley catheter in proper position in the posterior nasopharynx. Inflate the balloon with another 5–7 mL of water.

6. While maintaining traction, place anterior packing with layered gauze. Packing of the opposite side may be required to prevent septal deviation. Place a piece of gauze on the exposed catheter and secure with an umbilical clamp.

B  Dual-Balloon Tamponade Catheter

1. Double-balloon epistaxis catheters have both an anterior and posterior balloon, and some have an integral airway tube. These devices serve as both an anterior and a posterior pack. They are easily inserted and are often successful in temporary control of posterior epistaxis in the ED.

2. Insert the lubricated device along the nasal floor as far back as possible. Inflate the posterior balloon halfway with air, apply traction to pull the balloon up against the middle turbinate, and then complete the inflation. Maintain the position of the balloon and then inflate the anterior balloon with 30 mL of air.

3. This patient with posterior epistaxis was successfully treated in the ED and discharged. Historically, most patients with posterior packs were admitted to the hospital; however, the ease and safety of balloon devices allow selected patients to be treated as outpatients. Consider admission for the elderly, and those with pulmonary or cardiovascular disease.

Figure 63.39 Management of epistaxis: posterior packing with inflatable devices. ED, Emergency department.
Other Techniques

ENT consultation may be required for posterior nosebleeds that do not respond to the posterior packing techniques. Other treatment options that consultants may consider include ligation or embolization of the internal maxillary artery and posterior endoscopic cautery.

Complications

Posterior nasal packing is uncomfortable and often painful, but serious medical complications from posterior packing are rare. Complications associated with posterior packing include infection, dysphagia, dysfunction of the Eustachian tube, tissue necrosis, and dislodgment. Other serious complications rarely and often anecdotally associated with posterior packing are hypoxia, hypercapnia, aspiration, hypertension, bradycardia, arrhythmias, myocardial infarction (MI), and death.41 It is questionable whether packing alone is the cause of these associated complications. Rebleeding may also be seen with early pack removal; one series found that removal of the packing within 48 hours increased the risk for rebleeding.42 Most posterior packs are left in place for 72 to 96 hours.

Dysphagia from the packing can lead to poor oral intake and possibly necessitate intravenous fluid hydration. A decrease in the arterial partial pressure of oxygen (PaO2; 7.5 to 11 mm Hg) and an increase in the arterial partial pressure of carbon dioxide (PaCO2; 7 to 13 mm Hg) can be seen in patients with nasal packing who are treated with sedation.41 However, despite traditional theoretical concerns, such altered pulmonary physiology cannot be attributed to posterior packing alone. Although it has been hypothesized that a posterior pack will cause vagal stimulation and thereby result in varying degrees of bradycardia and bronchoconstriction because of the so-called nasopalatine reflex, studies have failed to substantiate substantial physiologic changes attributed to posterior packing. Observe patients with significant cardiopulmonary disease and posterior nasal packs in a monitored setting.

Tissue necrosis of the nasal ala, nasal mucosa, and soft palate secondary to improper placement or padding has been described. Protect the skin with gauze placed under the device to reduce skin maceration. The risk for necrosis increases with the duration of the packing, so all packing should be removed in 3 to 5 days.

If the posterior pack becomes dislodged, it will fall into the oropharynx and place the patient at risk for asphyxiation, vomiting, and aspiration. The patient and nursing personnel need to be familiar with the technique for removing the pack. To remove the pack, cut the anterior sutures that exit the naris from the gauze roll if they have not already broken. Grasp the sutures exiting the mouth and guide the packing out of the nasopharynx. It may be necessary to extract the packing with forceps or digits.

Antibiotics Following Nasal Packing

Posterior packing is associated with a risk for infection, including nasopharyngitis, sinusitis, and rarely, TSS.17,41 Packing blocks the sinus ostia, which prevents proper drainage of the sinuses. TSS has been described rarely with nasal packing97 and is estimated to occur in approximately 16 per 100,000 packings. The syndrome is caused by a toxin released by S. aureus infection of the packing. A sudden onset of vomiting and diarrhea with high fever, as well as the development of an erythematous rash, heralds the onset of the disease. A systematic review did not find any methodologically rigorous studies evaluating the incidence of TSS following nasal packing; simply small, retrospective, observational, and nonrandomized prospective studies. None of the studies reported TSS as a complication of nasal packing. There is no strong evidence to support or refute the practice of prescribing antibiotics. The choice is left up to the provider.39 Awareness of the condition is paramount. Importantly, antibiotics will not eradicate the carrier state of MRSA or other nasopharyngeal flora.

Likewise, no data support or refute the routine use of prophylactic antibiotics after nasal packing to reduce the incidence of bacterial sinus infections. Short-term anterior packing is not likely to be a cause of bacterial infection of the nasopharynx and does not call for antibiotic prophylaxis. As with all forms of antibiotic prophylaxis, routine antibiotic use is associated with risks and side effects, including the selection of resistant bacterial strains. Despite the lack of proven efficacy, many clinicians provide prophylaxis against TSS and bacterial infection for patients with posterior packing for the duration of the packing. There are, however, no universally accepted standards or mandates regarding this issue. It may be reasonable to provide prophylaxis to patients at greater risk for infection, such as those with diabetes, advanced age, or an immunosuppressed state. If used, an antibiotic with staphylococcal coverage should be selected, such as amoxicillin-clavulanate. Topical mupirocin may also be used on various packing material, but the efficacy of this is unknown.

Patient Disposition Following Nasal Packing

After successful cautery or a simple anterior pack, patients can be discharged with follow-up in 48 to 72 hours. Aspirin is avoided. Packs should be left in place. Merocel packs should be moistened by the patient three times a day with saline or water. If no packing is used, the patient can coat the cauterized area four times a day with antibiotic ointment or Vaseline and should avoid the urge to pick the nose or remove any debris. If minimal bleeding recurs, oxytetracycline spray is usually effective. After 48 to 72 hours, the packing is removed and the condition reassessed.

Healthy, stable patients with various forms of packing or commercial packs can usually be discharged if the bleeding is controlled in the ED for 1 to 2 hours. Minor bleeding may be experienced, but return of significant bleeding requires reevaluation and generally hospitalization. Hospitalization is often appropriate for those with posterior packs, especially the elderly or those with concerning underlying medical conditions. Clinical judgment is the best arbiter for admission decisions.

Tranexamic Acid

In cases of poorly controlled epistaxis, tranexamic acid (TXA) may be considered. TXA is an antifibrinolytic agent that is 10 times more potent than aminocaproic acid, with a longer half-life. TXA reversibly binds to plasminogen and inhibits binding to fibrin. In turn, this prevents plasminogen activation and transformation to plasmin. Currently, TXA is U.S. Food and Drug Administration (FDA)-approved for short-term use in hemophilia to prevent hemorrhage and reduce the need for factor replacement during and following dental extraction and for heavy menstrual bleeding. Despite these limited and specific indications, TXA has been evaluated in cardiac surgery, orthopedic surgery, trauma, spinal surgery, postpartum hemorrhage, gastrointestinal bleeding, hyphema, dental extractions, and epistaxis.41
Two studies have evaluated TXA for the management of epistaxis. In a randomized, double-blind, parallel-group study of 68 patients,\(^4\) 60% of patients treated with TXA topical gel had control of epistaxis within 30 minutes compared to 76% receiving placebo. Rebleeding rates at 8 and 30 days were 11% and 44% for TXA, and 31% and 66% for placebo. A 2013 randomized control trial by Zahed\(^6\) and colleagues used injectable TXA applied topically to the nares. They compared a 15-cm cotton pledget soaked in injectable TXA (500 mg in 5 mL) to anterior nasal packing. The authors found that within 10 minutes of treatment, bleeding stopped in 71% of the TXA group compared to 51% of the anterior nasal packing group (odds ratio 2.27; 95% confidence interval 1.68–3.06, \(p < 0.001\)). In addition, more patients in the TXA group were discharged in 2 hours or less. Rebleeding rates were less in the TXA group. Finally, a 2015 case report demonstrated that TXA in 2 hours or less. Rebleeding rates were less in the TXA group.

Given these results, TXA seems to be an effective agent to treat epistaxis. It should be used when conventional therapies fail to halt bleeding and should also be given to prevent rebleeding in patients with hereditary hemorrhagic telangiectasia. The injectable form can be applied topically to the nares in soaked pledgets. TXA also shows promise in controlling epistaxis in patients being treated with novel oral anticoagulants, though its use cannot routinely be recommended for this indication at the time of this writing.

To date, TXA has not been studied in patients with posterior epistaxis. Due to its antifibrinolytic effect, venous thromboembolic events (VTE), MI, and stroke can be seen following its administration. Importantly, a Cochrane review of topical TXA found no adverse events related to VTE, MI, or stroke. It was noted that TXA decreased the need for blood transfusion by 45%.\(^9\)

**Septal Hematoma**

Trauma to the anterior portion of the nasal septum may cause a hematoma to form. A buckling stress tears the submucosal blood vessels. If the mucosa remains intact, the blood will accumulate between the mucoperichondrium and the septal cartilage. Stagnant blood is an excellent medium for bacterial growth and the formation of an abscess. Common bacteria include *S. aureus*, *Streptococcus pneumoniae*, and group A β-hemolytic streptococci. Other complications of an untreated hematoma include septal perforation and cartilage destruction with a resultant saddle nose deformity. Septal hematomas may occur immediately after the trauma or, more commonly, in the first 24 to 72 hours after the injury.\(^5\) The hematoma can cause significant destruction of the nasal cartilage and result in a cosmetic deformity.

**Indications and Contraindications**

The presence of a nasal septal hematoma requires drainage to prevent a cosmetic defect, septal necrosis, and other complications. The most common symptoms of a septal hematoma are nasal obstruction, pain, rhinorrhea, and fever. Most patients will complain of an inability to breathe through the affected side, but the absence of nasal obstruction does not rule out a septal hematoma. It is usually possible to diagnose a septal hematoma by inspecting the nasal septum with a speculum for swelling, and a fluctuant area (Fig. 63.40). The presence of septal asymmetry with a bluish or reddish hue of the mucosa is suggestive of a septal hematoma. To some it looks like a grape. Inspect both sides because bilateral hematomas are possible. Direct palpation with the littlest finger may be necessary because newly formed hematomas may not yet be ecchymotic. Palpation can further differentiate septal hematoma from septal deviation, which may appear to be similar because of asymmetry. The best way to palpate for a septal hematoma is to insert the gloved small fingers in each side of the nose and palpate the entire septum to feel for swelling, fluctuance, or widening of the septal space.

There are no absolute contraindications to drainage of a nasal septal hematoma. Caution should be used in those with known bleeding diathesis or those who are taking anticoagulants.

**Equipment**

Very little equipment is required to drain a nasal septal hematoma. A topical or injectable anesthetic should be used.
For the procedure itself, a light source, a nasal speculum, a No. 11 scalpel blade, suction apparatus, scissors, nasal saline, small Penrose drain, and some form of nasal packing material are all that is needed.

**Procedure**

Treatment of a septal hematoma consists of evacuation of the clot with subsequent reapproximation of the perichondrium to the cartilage (Fig. 63.41). To drain the hematoma, incise the mucosa over the hematoma horizontally after adequate anesthesia is achieved. Suction out all the clot and then irrigate with normal saline. Excise a small amount of mucosa to prevent premature closure of the incision and place a section of a sterile rubber band to act as a drain. Pack the nostril, as for anterior epistaxis, to reapproximate the perichondrium to the cartilage.

Prescribe broad-spectrum antibiotic therapy. Inspect the septum daily for signs of infection, recurrent hematoma, or necrosis. Evacuate recurrent hematomas. When there is no further hematoma formation over a 24-hour period, remove the drain. Pack the affected naris for one more day to complete the apposition of perichondrium to cartilage where the drain had been. If any evidence of infection is present, admit for intravenous antibiotics and surgical débridement.

**Complications**

Though rare, nasal septal abscess formation is the most common complication of septal hematomas. The infection can spread to the sinus cavities and lead to meningitis, cavernous sinus thrombosis, intracranial abscess, and orbital cellulitis.

A large or rapidly expanding hematoma may cause pressure on the septum and lead to avascular necrosis of the septal cartilage. The nasal septum can collapse and lose its shape, which causes a noticeable cosmetic defect.

After drainage, the hematoma may recur and should be treated by repeated drainage to prevent cartilage damage. Reaccumulation can be prevented by incising a piece of mucosa before packing the nasal cavity.

**Reduction of Nasal Fractures**

A nasal fracture is the most common facial fracture. Nasal fractures are accompanied by a broad range of symptoms, including mild swelling, epistaxis, and periorbital ecchymosis with obvious deformity. As with any trauma involving the head, evaluate for coexistent intracranial injury or neck injury. In the evaluation of nasal trauma, rule out the existence of a septal hematoma or cerebrospinal fluid rhinorrhea. In most cases the swelling and soft tissue deformity prevent adequate evaluation, treatment, or both. Evaluation of a patient with a suspected nasal fracture includes a thorough history, external nasal examination, and internal nasal examination using a nasal speculum with or without the use of a rigid nasal endoscope. Nasal radiographs are not routinely needed because they will not alter the course of treatment or injury.

Ask the patient to apply ice to the area and keep the head elevated to reduce soft tissue swelling. Refer the patient to an otolaryngologist or plastic surgeon for reexamination and definitive treatment in 3 to 5 days. Stress the importance of reevaluation within 10 days so that the bones do not set in a misaligned state.

**Indications and Contraindications**

The indications for reduction of a nasal fracture are first based on the type of deformity and degree of swelling present (timing of reduction). Only simple nasal bone or nasal-septal complex fractures, nasal obstruction or airway compromise from a deviated septum, and fracture of the nasal-septal complex with nasal deviation less than half the nasal bridge should be reduced in the ED. Other indications include less than 3 hours after injury in adults and children if minimal edema is present, reduction 6 to 10 days after injury in adults once the edema has resolved but before setting of the fracture fragments, and reduction 3 to 7 days after injury in children once the edema has resolved. The presence of nasal obstruction or airway compromise from a deviated septum should prompt reduction in the ED.
Contraindications to reduction of nasal fractures in the ED include severe comminution of the nasal bones and septum, associated fractures of the orbital wall or ethmoid bone, deviation of the nasal pyramid greater than half the width of the nasal bridge, caudal septal fracture-dislocation, open septal fractures, and any fracture older than 3 weeks.

**Equipment**

The standard equipment of nasal decongestant and anesthetic are needed; in addition, nasal speculum, bayonet forceps, Frazier suction tip, anterior nasal packing material, a good light source, and some specialized equipment are needed, including elevators (Goldman, Boies, Salinger, Ballenger), Walsham forceps for grasping the nasal bones, and Asch forceps for reduction of the septum. An external nasal splint is also needed.

**Procedure**

Most fractures and patients with significant soft tissue swelling should be seen in follow-up for definitive evaluation and possible reduction of the fracture. Complicated fractures and septal injuries are usually referred for follow-up. Simple fractures with minimal local swelling can be treated by closed reduction (Fig. 63.42). Some patients prefer immediate correction, are not concerned with aesthetics, or are unable to comply with

---

**Nasal Fracture Reduction**

1. Unilateral fracture of the nasal pyramid
2. Dislocation of the septum
3. Asch forceps can be used to reduce a displaced nasal septum. Insert each arm of the instrument on either side of the nasal septum.
4. Closed nasal reduction. Minor deformities with minimal swelling may be reduced in the ED and mitigate further therapy. After marking the distance of the intercanthal line on the elevator with a thumb, the tip of the instrument is used to reduce the medialized fragment by elevating it. The opposite thumb may simultaneously reduce a contralateral outfractured nasal bone (pyramid). Use the handle of a scalpel if an elevator is unavailable.

**Figure 63.42** Reduction of a nasal fracture. *ED*, Emergency department. *(Steps 3 and 4, from Flint PW, Haughey BH, Lund VJ, et al, editors: Cummings otolaryngology: head and neck surgery, ed 5, Philadelphia, 2010, Mosby Elsevier.)*
the effects of the anesthetic wear off. Finally, if nasal packing may result in nerve damage, dysesthesias, or paresthesias after lead to cartilage destruction and deformity. Reduction can also may produce a nasal septal hematoma, which if untreated, can
reduction may result in a poor cosmetic outcome. Reduction
(ENT, plastic, or oral and maxillofacial surgeon). Incomplete
be referred for open reduction by a cosmetic facial surgeon
Not all fractures will be reducible and such fractures should
Condensation
Complications
Not all fractures will be reducible and such fractures should
be referred for open reduction by a cosmetic facial surgeon
(ENT, plastic, or oral and maxillofacial surgeon). Incomplete
reduction may result in a poor cosmetic outcome. Reduction
may produce a nasal septal hematoma, which if untreated, can
lead to cartilage destruction and deformity. Reduction can also
cause excessive bleeding. This can be managed with direct
pressure and nasal packing. Direct infiltration of anesthetic
may result in nerve damage, dysesthesias, or paresthesias after
the effects of the anesthetic wear off. Finally, if nasal packing
is placed, there is a risk for sinus infection.

Nasal FB Removal

Nasal FBs most frequently occur in pediatric patients, but it
is not uncommon to find them in psychiatric or cognitively
impaired patients as well. Usually, a family member has wit-
tnessed the event or the patient complains of discomfort from
the FB. Patients may also have unilateral purulent or bloody
nasal discharge, unilateral sinusitis, or recurrent unilateral
epistaxis. Retained FBs, especially plastic ones, often initially
fail to cause pain or other symptoms. The lack of a history
of FB insertion is of little value in children because many will
not admit it. Therefore emergency clinicians need to maintain
a high level of suspicion for nasal FBs.

Types of nasal FBs vary widely and include food (e.g., meat,
nuts, beans), rubber erasers, paper wads, pebbles, marbles,
sponges, beads, jewelry, hardware (e.g., nuts, screws), and even
certain living larvae or worms.

Alkaline button batteries pose a unique problem because
they may cause significant nasal injury within hours to days.
They are composed of heavy metals such as mercury, zinc,
silver, nickel, cadmium, and lithium. Injuries can occur and
include mucosal burns, ulcerations, liquefaction necrosis, septal
perforation, synechiae, and stenosis of the nasal cavity. It is
imperative that these batteries be removed promptly before
tissue damage occurs as a result of leakage of the battery con-
tensts, electrical currents, or direct pressure. A relatively new
and interesting nasal FB is the magnetic nose ring. These small,
commercially available earth magnets are usually worn on either
side of the alar cartilage and give the appearance of a pierced
nadal stud. The magnets can be displaced and become polarized
across the nasal septum. The magnetic attraction can be quite
strong and may lead to pressure necrosis of the nasal mucosa
and possibly septal perforation. This attraction can also make
removal difficult, as well as painful for the patient. Suggested
techniques include using polarized or nonferromagnetic tools.

Many nasal FBs come to rest on the floor of the anterior
or middle third of the nose. Metallic or calcified objects may
show up on radiographs, but physical examination remains the
most reliable means for diagnosis. Maxillary, ethmoid, or
sphenoid sinusitis may also accompany FB retention. Plain
radiographs or facial CT scanning may be of value in detecting
sinusitis, although these studies are not usually necessary with
an acutely retained object.

The inability to remove a nasal FB should necessitate ENT
consultation and may result in admission for removal under
anesthesia; therefore it behooves emergency clinicians to be
skilled in this procedure. Although admission incurs increased
cost, has inherent procedural risks, and causes psychological
stress in parents and patients, providers should be prepared to
call a consultant after several attempts. As with the removal
of auricular FBs, removal of nasal FBs can be both frustrating
and time-consuming.

Indications and Contraindications

Attempt removal of a nasal FB only if it is likely to be retrieved.
ENT consultation is needed if there is any doubt about the
ability to successfully retrieve the object. Repeated attempts
may result in trauma and displacement of the object into a
less favorable location.

Do not attempt mechanical removal if the object appears
to be out of range of the instruments. In addition, do not
attempt removal in an uncooperative patient unless sedation
is provided. Use sedation cautiously because it may increase
the risk for aspiration, especially when using agents that blunt
the protective airway reflexes.

Equipment

The majority of the equipment used for removing FBs in the
ear canal can also be used for nasal FB removal. Topical nasal
anesthetic and vasoconstrictor solutions, such as oxymetazoline, can greatly aid in the removal of nasal FBs.

Procedure
A cooperative patient is essential. Weigh the risks and benefits of using simple or procedural sedation. Ketamine anesthesia works well for children. Before attempting removal, anesthetize and vasoconstrict the mucosa of the affected naris topically. Ask an assistant to stabilize the patient’s head, and immobilize a younger patient if necessary. Place more cooperative patients in the sniffing position and use a headlamp for proper illumination. Several of the techniques previously mentioned for removing an FB from the EAC can also be used in the management of a nasal FB, including the use of cyanoacrylate glue.

Manual Instrumentation
Use alligator forceps or bayonet forceps to retrieve anteriorly lodged FBs that have edges amenable to grasping (Fig. 63.43, plate 1). For harder or larger objects, carefully pass a wire loop (one can be made using 24-gauge wire loop grasped by a hemostat), a right-angle hook (if unavailable, one can be made out of an 18-gauge needle), or even a properly bent paper clip. Carefully insert it beyond the object and rotate it to allow the FB to be pulled from the naris. Direct mucosal trauma and epistaxis may occur with any of these methods. For a nasal FB with smooth, round edges that are difficult to grasp or get behind, attempt to extract it with a suction-tipped catheter in a manner similar to that described earlier for FBs in the EAC. The Hognose catheter described earlier also works well for nasal FBs (see Fig. 63.21). Local vasoconstriction and anesthesia are helpful, and 2% lidocaine with epinephrine (local anesthetic) may be used.

Balloon Catheter
For an object that cannot be removed with anterior instrumentation, one consideration is to attempt removal with a balloon catheter. A Fogarty catheter can be highly effective in removing a nasal FB (see Fig. 63.43, plates 2 and 3). A No. 4 or 5 vascular Fogarty catheter, a 12-Fr Foley catheter, and a No. 6 biliary

---

**NASAL FOREIGN BODY REMOVAL**

1. Following vasoconstriction and anesthesia (2% lidocaine with epinephrine was dripped into the nose), some FBs may be easily extracted with forceps.

2. A Fogarty catheter may be used to extract a nasal FB. Insert the lubricated catheter tip above the FB and then gradually inflate the balloon.

3. Slowly withdraw until resistance is met and then pull the object out of the naris.

4. Balloon tip

5. Bag-valve-mask technique to blow an FB out of the naris. Ensure that the face mask forms a tight seal around the patient’s mouth and that the unaffected nostril is completely occluded. Attempt to firmly compress the bag as the patient exhales (an assistant is helpful to hold the mask snugly and to occlude the other nostril). This technique works best with objects that completely occlude the nostril.

6. To blow an FB out of the nose, have the parent blow into the mouth while occluding the unaffected side with the thumb.

*Figure 63.43 Removal of a nasal foreign body. FB, Foreign body.*
Two other techniques using positive pressure have been described and use simple equipment and oxygen or medical air. In the first technique, wall oxygen or medical air, suction tubing, and a suction-tubing adapter are needed (Fig. 63.44 A).

Have the patient sit upright and lean forward with the head tilted down. Attach one end of standard suction tubing to the wall oxygen or compressed air unit. The plastic suction-tubing adapter should be firmly placed into the opposite end of the suction tubing. Insert the tubing with the adapter into the Fogarty catheter have all been described in the literature for this use. A biliary catheter is reportedly less apt to rupture. Place the patient in the supine position and apply a vasoconstrictor and anesthetic to the nasal mucosa. With a 5-mL syringe attached and the catheter lubricated with lidocaine gel, pass the tip above the object and into the nasopharynx. Inflate the balloon with air or water (~2 mL in small children and 3 mL in older children) and control the syringe plunger and balloon size with your thumb. Withdraw the catheter until resistance is felt, and then slowly pull the object out. The Katz Extractor otorhinologic FB remover (InHealth Technologies, Carpinteria, CA; see Fig. 63.43, plate 4) is a disposable, single unit composed of a flexible catheter with a balloon tip attached to a syringe. The procedure is similar to that for the Fogarty catheter. Complications mentioned in the literature include mild post-traumatic bleeding, as well as the theoretical risk for airway obstruction by the balloon or aspiration from further displacement of the object.

Positive Pressure

Another approach to a posteriorly placed nasal FB is to blow the object out with positive air pressure. The simplest way is to ask patients to blow their nose while occluding the unaffected nostril. This is only really effective in older children and adults. Alternatively, place a bag-valve-mask device over the child's mouth to provide positive pressure (see Fig. 63.43, plate 5). Occlude the opposite nostril and apply the Sellick maneuver to prevent passage of air into the esophagus. This technique often requires restraint and can also be threatening to a young child.

The “parent's kiss” technique may be used in children. First gain the child's cooperation by saying that parent is going to “give them a big kiss.” The technique is performed by having the child lie supine. Ask the parent to occlude the child's unaffected nostril with the thumb (see Fig. 63.43, plate 6). Next, as in mouth-to-mouth resuscitation, ask the parent to make a firm seal with his or her mouth over the child's open mouth and then give a short, sharp puff of air briskly into the mouth to produce outward pressure behind the object. Keep the opposite nostril occluded throughout the procedure. If successful, the object will move within grasping reach of an instrument or pop completely out of the naris. If it fails, the technique can be repeated. Studies of the parent's kiss technique have found it to be highly effective, nontraumatic, and preferable to restraint or instrumentation.

A modification of the technique that may be easier for some parents to master uses a drinking straw placed between the parent's mouth and the child's mouth. Instruct the child to make a tight seal, as though drinking, and ask the parent to deliver a quick puff.

Another similar technique may be successful. After a nasal decongestant is instilled into the affected side, place a 6- to 10-inch section of 1/4- or 1/8-inch wide rubber or soft vinyl tubing into or at the contralateral nostril and hold the tubing in place with the fingers of one hand. Place the free hand gently over the child's mouth, take a good-sized breath, and blow forcefully through the tubing. This step may be repeated up to four times. In one study, success was reported in 40 of 41 cases. Navitsky and colleagues modified this technique to reduce the risk for transmission of disease. Turn a wall oxygen source to a flow of 10 to 15 L/min. Use a male-to-male adapter at the end of oxygen tubing to direct the flow of oxygen into the contralateral (nonobstructed) nostril.

Two other techniques using positive pressure have been described and use simple equipment and oxygen or medical air. In the first technique, wall oxygen or medical air, suction tubing, and a suction-tubing adapter are needed (Fig. 63.44 A). Have the patient sit upright and lean forward with the head tilted down. Attach one end of standard suction tubing to the wall oxygen or compressed air unit. The plastic suction-tubing adapter should be firmly placed into the opposite end of the suction tubing. Insert the tubing with the adapter into the

Figure 63.44 Additional techniques for positive pressure removal of nasal foreign bodies. A, Attach suction tubing to wall oxygen or medical air, and use a suction adapter inserted in the contralateral naris. B, An alternative technique uses oxygen tubing, an earpiece from a disposable stethoscope, and an 8-Fr suction catheter. C, Equipment assembled and positioned for use.
with dystonic reactions to medications. Patients with a previous history of TMJ dislocation are more prone to repeated dislocations. The condition can be unilateral or bilateral (Fig. 63.45).

TMJ dislocation occurs when the mandibular condyle moves anteriorly along the articular eminence and becomes locked in the anterosuperior aspect of the eminence. Spasm of the masseter, internal pterygoid, and temporalis muscles occurs during an attempt to close the mandible. Trismus then results and the condyle cannot return to its normal position. Predisposing factors include anatomic disharmony between the mandibular fossa and the articular eminence and weakness of the capsule and the temporomandibular ligaments.

**Indications and Contraindications**

The presence of mandibular dislocation is the indication for reduction. The diagnosis is usually straightforward, but the condition can be misinterpreted as an acute dystonic reaction. With dislocation, patients cannot close their mouth, and speech is affected. Pain varies, and patients are often very anxious. With unilateral dislocation, the jaw will deviate to the opposite side. More commonly, bilateral dislocation occurs. With traumatic dislocation, radiographic evaluation should be performed to exclude a fracture. A mandibular series, Panorex, TMJ radiographs, and facial CT are acceptable. In the absence of a fracture, there are no real contraindications to reduction.

**Equipment**

Minimal equipment is required. An ENT or dental chair is helpful. Gloves, gauze, and a bite block should be available. Items to protect the operator’s fingers such as tongue blades or plastic finger splints can be used.

**Complications**

Most complications are due to the FB itself and include pain, obstruction, rhinorrhea, ulceration of the nasal mucosa, septal perforation, infection, and nasal or choanal stenosis. Bleeding and mucosal laceration are the most commonly reported complications of removal. The bleeding is usually minor and generally resolves with simple pressure. It may occur as a result of the object itself or from trauma during removal. Inadvertent posterior dislodgement of the object may occur and result in ingestion or aspiration of the object. Theoretically, barotrauma to the lungs and TMJs may occur with positive pressure techniques.

**Dislocation of the Mandible**

Mandibular dislocation is more properly known as temporomandibular joint (TMJ) dislocation. It is actually the mandibular condyles that dislocate. It may result from trauma but more commonly follows extreme opening of the mouth such as may occur while eating, laughing, or yawning. It may also be seen with dystonic reactions to medications. Patients with a previous history of TMJ dislocation are more prone to repeated dislocations. The condition can be unilateral or bilateral (Fig. 63.45).

Figure 63.45 Mandible dislocations. A and B, This patient yawned and then could not close his mouth. This was a recurrent bilateral mandible dislocation. C, This patient was suspected of having a dystonic reaction because she could not speak and her mandible was misaligned. It was a unilateral mandibular dislocation.
Mandible Dislocation Reduction

A Anatomy

In the normally closed position, the mandibular condyle rests in the mandibular fossa behind the articular eminence. In the maximally open position, the condyle is just under and slightly behind the eminence. In the dislocated position, the condyle moves forward and upward slightly above the eminence; muscle spasm then occurs.

B Reduction Maneuver

1 Sedate the patient or locally inject the TMJ with lidocaine. Wrap your thumbs with gauze and place them in the mouth on the back molars. Apply downward pressure on the lower molar ridge near the angle of the jaw.

2 Continue to apply downward pressure and push the mandible posteriorly. A rocking motion may be helpful.

C Alternative Techniques

1 Recumbent approach with the operator at the head of the patient.

2 Ipsilateral extraoral approach.

3 Wrist pivot method.

directly into the lateral pterygoid muscle. For TMJ injection, prepare the skin anterior to the ear, and introduce the needle into the TMJ space at the palpable depression caused by the dislocated condyle. Direct the needle anteriorly and superiorly onto the inferior surface of the glenoid fossa, and inject 2 mL of local anesthetic. For the pterygoid muscle, inject 2 to 3 mL of anesthetic into the muscle posterior to the maxillary tuberosity.

**Classic Technique (see Fig. 63.46B)**

With the patient seated and the head stabilized against the headrest, place a bite block in the mouth to prevent injury to the clinician. Wear gloves to protect your thumbs. Stand and face the patient and place your thumbs on the patient’s lower molar teeth. The level of the mandible should not be higher than the operator’s elbow. Once the thumbs are positioned on the molars as far posteriorly as possible, curve the fingers and hand around the angle and body of the mandible along the jaw and chin. Then exert steady, constant downward pressure on the lower molar region, and direct the mandible inferiorly and posteriorly back into the temporal fossa. Be careful because once reduced, reflex spasm of the jaw muscles can suddenly snap the mandible shut and injure your thumbs. Alternatively, place the thumbs on the mandibular ridge instead of the molars. For bilateral dislocations, both sides can be reduced simultaneously, but it is easier to do one side at a time.

**Recumbent Approach (see Fig. 63.46C)**

With the patient lying recumbent, stand in front of the patient. Apply caudal and posterior force on the mandible as for the classic technique. Alternatively, stand behind the patient’s head and place your thumbs on the molars. Apply downward and backward pressure for reduction.

**Posterior Approach**

With the patient seated, stand behind the patient. Place your thumbs posterior to the last molar on the retromolar gum and along the ramus of the mandible. Exert downward force in this position. This procedure carries less risk of the fingers being bitten, but proper positioning of the thumbs may be difficult to accomplish.

**Ipsilateral Approach (see Fig. 63.46C)**

The ipsilateral approach involves three routes, extraoral, intraoral, and combined, which are conducted in sequential fashion. With this technique one side at a time is reduced. Stand at the patient’s side. Attempt the extraoral route first. Use the thumb of your dominant hand to apply downward pressure on the displaced condyle just inferior to the zygomatic arch. Use the other hand to stabilize the patient’s head.

If unsuccessful, attempt the intraoral approach. Exert downward pressure intraorally on the ipsilateral molar teeth. If unsuccessful, use the combined approach. Apply extraoral pressure with one thumb and intraoral pressure with the other thumb on the ipsilateral side.

**Alternative Manual Method**

This is similar to the extraoral technique described earlier. Place your fingers directly over the preauricular prominence of the dislocated condyle. Massage the condyle in a posterior and inferior direction to induce relaxation of the muscles and guide the condylar head back into the fossa. This pressure on the condyle may be uncomfortable for patients.

**Wrist Pivot Method (see Fig. 63.46C)**

Face the patient and place your thumbs at the apex of the meniscus of the mandible. Wrap the other fingers laterally around the mandible and onto the occlusal surface of the inferior molars. Apply an upward force on the chin with your thumbs, and apply a downward force on the mandible with your fingers. Move the wrists in the direction of ulnar deviation. With these maneuvers, rotate the condyles inferiorly and posteriorly into the fossa.

**Gag Reflex Method**

The gag reflex method uses a component of jaw relaxation and transient descent of the mandible inferiorly. Induction of the gag reflex has been described as a successful method for reduction of mandibular dislocations. Provide tactile stimulation of the soft palate with a dental mirror or tongue blade in an awake patient. Muscle relaxation occurs and the mandible descends caudally so that the condyle moves inferiorly and relocates back into place.

**Syringe Technique (Fig. 63.47)**

The syringe technique is a novel, hands-free method for reduction of nontraumatic mandible dislocations that is fast, safe, effective and does not require procedural sedation. In one study, 97% of patients were successfully reduced, with the majority occurring in less than 1 minute. In this method, with the patient seated, place a 5-mL or 10-mL syringe between the posterior upper and lower molars or the gums on the affected side. Instruct the patient to bite down and grasp the syringe between the teeth or gums. Next, ask the patient to roll the syringe back and forth, resulting in reduction of the dislocated TMJ joint. Syringe size varies with each patient and is determined by the distance between the upper and lower molars or gums and the ability to open the mouth on the affected side to allow syringe placement. The syringe acts as a rolling fulcrum. The mandible and maxilla apply pressure on the syringe. As the molars or gums move over the syringe, the gliding action moves the mandible posteriorly. The displaced condyle moves posteriorly to allow the condyle to slide back into its normal position. In bilateral dislocations, reducing one side will spontaneously reduce the other.

After relocation and on discharge, advise the patient to avoid extreme mouth opening and to eat soft food for 1 week. Warm compresses, nonsteroidal antiinflammatory agents, and muscle relaxants may be helpful. The patient may be referred to an ENT or oral and maxillofacial surgeon for further care. Patients with chronic dislocation may require surgical fixation.

**Complications**

Complications include injuries to both the patient and operator. Patient injuries include redislocation, mandibular fracture (rare), joint cartilage injuries, torn ligaments or muscles, and dental or gum injuries as a result of trauma from the attempt at reduction. There may be complications associated with procedural sedation as well.

Operator injuries are prominent in techniques that require placement of the operator’s fingers in the mouth or on the teeth. These are usually caused by sudden masseter spasm after reduction. This results in the patient involuntarily biting down on the operator’s thumbs. Take measures such as gauze and plastic finger splints to prevent such injuries to the thumbs.
Uvulitis/Angioedema of the Uvula

Most cases of acute angioedema of the uvula (uvula hydrops), also known as Quincke’s disease, are spontaneous and no cause can be found. Affected patients wake up with a lump in their throat or a fullness when swallowing, look in the mirror, and see an enlarged uvula (Fig. 63.48). It appears as an edematous, pale, and watery-filled structure.67 It is not painful. Most patients are young and otherwise healthy.

Although most of the time the cause remains unidentified, there are numerous known precipitants, including smoking of marijuana and cocaine (crack), trauma to the uvula by endotracheal tubes or suction catheters during general anesthesia, and sticking a finger down the throat to induce vomiting. Do not confuse this condition with hereditary angioedema (hereditary angioneurotic edema), which is due to C1 esterase inhibitor deficiency, a recurrent and potentially fatal condition. Rarely, uvulitis may be caused by bacterial infection, particularly *Haemophilus influenzae* B, and can coexist with epiglottitis. Angiotensin-converting enzyme inhibitor use does not seem to be related, but nonspecific allergic reactions (food, environmental) have been postulated. This condition, though annoying, is usually benign, self-limited, and resolves in 24 to 48 hours. Airway compromise is a theoretical concern but occurs only rarely. Evaluating and maintaining the airway are
hemostasis. Up to 50% of patients who have major postoperative bleeding require control in the operating room. It is difficult and occasionally impossible to even initially control severe hemorrhage in the ED. Do not hesitate to intubate a massively bleeding patient to protect the airway and more easily perform local hemostatic maneuvers. Until the surgeon arrives, apply pressure directly on the bleeding area with a sponge on a long clamp (Fig. 63.50). The sponge may be dipped in epinephrine or thrombin powder if available. The bleeding area can also be infiltrated with lidocaine with epinephrine.

If a patient is seen in the ED with a complaint of posttonsillectomy bleeding but no active bleeding is found on examination and a blood clot is present, do not remove the clot. It may be tempting to discharge stable patients but definitive treatment and disposition are best made by the surgeon. If bleeding is present, or has temporarily stopped, it is prudent to admit the patient for observation. Obtain a sample of blood for a coagulation profile and a complete blood count. Blood transfusions may be required.

**Posttonsillectomy Bleeding**

Hemorrhage is the most serious complication of adenotonsillectomy, with reported rates of 0.5% to 10%, depending on the technique (Fig. 63.49). Bleeding is categorized as intraoperative, primary (within 24 hours), and secondary (between 24 hours and 10 days). Bleeding is serious at all times and must be evaluated and controlled quickly if active. The most common time for patients to be seen in the ED with delayed bleeding is between the fifth and seventh postoperative days. Although some bleeding is minor, sudden severe hemorrhage can be fatal if not managed appropriately. If active bleeding is confirmed by physical examination, consult otolaryngology. Minor bleeding 5 to 10 days postoperatively secondary to eschar separation may be evaluated and treated in the ED. However, multiple bleeding episodes are common in those who bleed, and minor bleeding may herald more significant subsequent hemorrhage. A severely bleeding patient should be taken to the operating room immediately for hemostasis. Up to 50% of patients who have major postoperative bleeding require control in the operating room. It is difficult and occasionally impossible to even initially control severe hemorrhage in the ED. Do not hesitate to intubate a massively bleeding patient to protect the airway and more easily perform local hemostatic maneuvers. Until the surgeon arrives, apply pressure directly on the bleeding area with a sponge on a long clamp (Fig. 63.50). The sponge may be dipped in epinephrine or thrombin powder if available. The bleeding area can also be infiltrated with lidocaine with epinephrine.

If a patient is seen in the ED with a complaint of posttonsillectomy bleeding but no active bleeding is found on examination and a blood clot is present, do not remove the clot. It may be tempting to discharge stable patients but definitive treatment and disposition are best made by the surgeon. If bleeding is present, or has temporarily stopped, it is prudent to admit the patient for observation. Obtain a sample of blood for a coagulation profile and a complete blood count. Blood transfusions may be required.

**References are available at www.expertconsult.com**
REFERENCES


Complaints pertaining to the teeth and supporting maxillofacial structures are common, and patients frequently go to the emergency department (ED) for evaluation. Complaints may range in scope from a simple chipped tooth to an odontogenic deep space infection or a maxillofacial injury. Treating these patients can be challenging and frustrating for busy emergency clinicians. Many emergency clinicians and other acute care providers do not receive specific training in dental emergencies during their training, yet it is important for them to be able to recognize and treat a wide range of dental and related maxillofacial problems. Some dental emergencies can lead to significant morbidity such as loss of teeth, chronic pain, infection, and craniofacial abnormality, whereas others can lead to life-threatening airway compromise.

Management of specific dental emergencies requires a thorough understanding of adult and pediatric dentition. The relevant anatomy of both populations is outlined. The techniques described for management of the various traumatic and infectious problems are, in most cases, temporizing until definitive dental or maxillofacial surgery referral can be obtained. Conditions that require emergency consultation are discussed. Topical, local, and regional anesthesia are of particular importance and utility in the management of odontogenic emergencies, so the clinician should be very familiar with these techniques.

Although this chapter describes the diagnosis and treatment of dental injuries that may confront emergency clinicians, no standard of care mandates that complex dental problems (e.g., replacement of avulsed teeth, drainage of infection) be definitively handled in the ED setting. Advances in ED equipment and clinician training, as well as the introduction of dental skills laboratories into the resident curriculum, are gradually raising the existing standard of care. The initial stabilization of fractured, subluxed, luxated, and avulsed teeth, as well as bleeding and dry sockets, is now within the treatment realm of the emergency clinician. Likewise, abscesses and infections should be properly identified and drained if necessary. It is appropriate to refer all significant dental pathology to a dentist or oral surgeon.

**TEETH**

The adult dentition normally consists of 32 teeth: 8 incisors, 4 canines, 8 premolars, and 12 molars. From the midline to the back of the mouth on each side, there is a central incisor, a lateral incisor, a canine (cuspid), two premolars (bicuspids), and three molars, the last of which is called the wisdom tooth (Fig. 64.1). The 20 primary or deciduous (baby) teeth include 8 incisors, 4 canines, and 8 molars. From the midline to the back of the mouth, there is a central incisor, a lateral incisor, a canine, and two molars (Fig. 64.2). There are no premolars in the primary dentition. Agenesis, or lack of proper formation of a tooth or teeth, is not uncommon, especially in the maxilla. Likewise, supernumerary, or extra, teeth may also occur. The adult teeth are numbered from 1 to 32, with the first tooth being the right upper third molar and the 16th tooth being the left upper third molar. The left lower third molar is the 17th, and the 32nd tooth is the right lower third molar. Numerous classification and numbering systems of the teeth exist; however, it is best for clinicians to simply describe the location and type of tooth in question (e.g., upper left second premolar, lower right canine). This removes any question when discussing a case with a consultant.

A tooth consists of the central pulp, the dentin, and the enamel (Fig. 64.3). The pulp contains the neurovascular supply of the tooth, which is responsible for carrying nutrients to the dentin, a microporous substance that consists of a system of microtubules. The dentin makes up the majority of the tooth, is a primary determinant of tooth color, and cushions the tooth during mastication. The enamel is the relatively translucent, outermost portion of the tooth and the hardest part of the body. The tooth may also be described in terms of the crown (coronal portion) and the root (apex portion). The crown is the portion covered in enamel; the root is the part that serves to anchor the tooth in the alveolar bone.

The following descriptive terminology is used for the different anatomic surfaces of the tooth. These terms are useful when describing the specific tooth injury to a consultant or colleague:

- **Facial**: the part of the tooth that faces the opening of the mouth. This is the part that you see when somebody smiles. It is a general term applicable to all teeth.
- **Labial**: the facial surface of the incisors and canines.
- **Buccal**: the facial surface of the premolars and molars.
- **Oral**: the part of the tooth that faces the tongue or the palate. This is a general term applicable to all teeth.
- **Lingual**: toward the tongue; the oral surface of the mandibular teeth.
- **Palatal**: toward the palate; the oral surface of the maxillary teeth.
- **Approximal/interproximal**: the contacting surfaces between two adjacent teeth.
- **Mesial**: the interproximal surface facing anteriorly or closest to the midline.
- **Distal**: the interproximal surface facing posteriorly or away from the midline.
- **Occlusal**: biting or chewing surface of the premolars and molars.
- **Incisal**: biting or chewing surface of the incisors and canines.
- **Apical**: toward the tip of the root of the tooth.
- **Coronal**: toward the crown or the biting surface of the tooth.

**THE PERIODONTIUM**

The periodontium, also known as the attachment apparatus, consists of two major subunits and is necessary for maintaining the integrity of the normal dentoalveolar unit. The gingival subunit consists of the junctional epithelium and gingival tissue. Gingival tissue is composed of keratinized, stratified squamous epithelium; it can be divided into the free gingival margin and the attached gingiva. The free gingiva is the cuff of tissue formed around the neck of the tooth. The
The gingival subunit is primarily responsible for maintaining the integrity of the periodontal subunit. Certain disease states such as gingivitis weaken the attachment apparatus and can result in loss of a tooth.

**ACUTE TOOTHACHE IN THE ED**

Patients with an acute toothache (odontalgia) often come to the ED for dental evaluation and relief of symptoms. Although multiple problems can initially cause pain in the area of the teeth, the cause is usually pulpitis, abscess, or dental trauma. Pulpitis can be either acute or chronic and the physiology,
Pain, depending on the scenario. Hile and Linklater\(^2\) reported significant pain relief in a fractured tooth by applying 2-octylcyanoacrylate tissue adhesive (Super Glue) directly to the tooth. This intervention is currently anecdotal but may also provide temporary pain relief for patients with open decay when air and temperature exacerbate the pain. Dry the tooth thoroughly with gauze, and generously apply a few layers of the product to the affected area. This intervention lasts less than 24 hours before enzymatic breakdown occurs. Note that the use of skin adhesives has not been approved for intraoral use; they tend to break down quickly in the oral cavity.

For years, clove oil (containing eugenol) has been a popular and reasonably effective short-term home remedy for an acute toothache or inflamed gingiva. For a cavity or gingiva pain, saturate a piece of cotton in clove oil and place the cotton directly in the cavity or along the gum. This will provide relief for a few hours. Clove oil should not be used more than a few times because of irritation and possible nerve damage. Another simple method of pain management is to apply viscous lidocaine directly onto the tooth or saturate a small cotton ball with the gel and place it in a cavity. This treatment is temporary, and caution should be exercised if reapplication is considered. Do not exceed the total recommended dosage of lidocaine.

Acute dental pain may also be referred pain, so a complete evaluation should be conducted if the anatomy in question appears to be normal (Fig. 64.4). For example, acute sinusitis can cause tooth pain and vice versa. Obvious dental infection should be treated with antibiotics (e.g., penicillin, clindamycin, erythromycin, metronidazole, amoxicillin-clavulanate) while awaiting dental evaluation. Chronic acetaminophen overdose is a known complication of overaggressive use of analgesics by patients unable to obtain dental care for an acute toothache.\(^3\)

Antibiotics provide no benefit for pain from a simple toothache, dental cavity, or pulpitis, although some clinicians prescribe them because follow-up dental care may be delayed or difficult to obtain.\(^3\) The emergency clinician, likewise, doesn’t

---

**Figure 64.2** Identification of teeth, adult and child. Each tooth has a number assigned to it. By 14 years of age, all primary teeth should normally be lost. Do not reimplant an avulsed primary tooth; rather, refer to a dentist to prevent future misalignment of the permanent teeth.

**Figure 64.3** The dental anatomic unit.
Dental Fractures

Dentoalveolar trauma is common reason for ED visits. Injury to the maxillary central incisors accounts for between 70% and 80% of all fractured teeth. Acute trauma to the teeth related to a dental problem. Two days after onset of the pain, the rash of herpes zoster appeared, which was the cause of the pain.

Individual teeth (except the posterior molars) can be temporarily anesthetized with total pain relief by simply giving a supraperiosteal injection of a local anesthetic (see Chapter 30). Bupivacaine has a long duration of action (4 to 12 hours) and has been shown to decrease the narcotic requirement of postoperative oral surgery patients even after the anesthetic properties of the medication have worn off. Molars, which are more difficult to anesthetize with supraperiosteal injections, can be anesthetized via different nerve block techniques, such as the inferior alveolar nerve block.

Dentoalveolar Trauma

Dental Fractures

Dentoalveolar trauma is a common reason for ED visits. Injury to the maxillary central incisors accounts for between 70% and 80% of all fractured teeth. Acute trauma to the teeth is not usually life-threatening; however, the morbidity associated with poorly managed dental fractures can be significant and includes failure to complete eruption, change in color of the tooth, abscess, loss of space in the dental arch, ankylosis, abnormal exfoliation, and root resorption. Dental injuries are often associated with intraoral lacerations. When a tooth is chipped or missing and there is a concomitant intraoral laceration, there is a risk that the missing portion of the tooth is embedded in the depths of the laceration (Fig. 64.5).

Figure 64.4 This physician had severe pain in his jaw thought to be related to a dental problem. Two days after onset of the pain, the rash of herpes zoster appeared, which was the cause of the pain.

Some general principles apply to the evaluation and management of dental trauma. First, identify all fracture fragments and mobile teeth. Percuss each tooth surface for mobility and sensitivity. If a tooth is missing, it cannot always be assumed that it has been avulsed. Teeth can be aspirated into the respiratory tract, swallowed into the gastrointestinal tract, or fully intruded into the maxillary sinus, alveolar bone, or nasal cavity. Take radiographs if there is any suspicion of aspiration of tooth fragments or intrusion of fragments into the gingiva or alveolar bone. Second, the dentition is much more easily manipulated if the patient is not in significant discomfort. Tooth injection and common dental blocks should be part of the emergency clinician’s armamentarium. Third, topical tooth remedies and analgesics, both over-the-counter and prescribed, should be discouraged because their long-term use can lead to the development of sterile abscesses and soft tissue irritation. Fourth, administer tetanus vaccine if needed.

Management of fractured teeth depends on the extent of fracture with regard to the pulp, the degree of development of the apex of the tooth, and the age of the patient. Dentoalveolar injuries and, in particular, tooth fractures can be classified in many ways. The Ellis classification is one system often cited in the emergency medicine literature; however, many dentists and maxillofacial surgeons do not use this nomenclature, thus making it less than ideal when discussing these types of injuries with specialists (Fig. 64.6). The most easily understood method of classification is one based on a description of the injury.

Crown fractures may be divided into uncomplicated and complicated categories. Uncomplicated crown fractures result from injuries to the enamel alone or to a combination of the enamel and dentin. Complicated crown fractures extend into the pulp.

Ellis Class I Fractures

Uncomplicated crown fractures through only the enamel are known as Ellis class I fractures (see Fig. 64.6B). They are not usually sensitive to either temperature or forced air. These fractures generally pose minimal threat to the health of the dental pulp. They may feel sharp to the patient’s tongue, lips, or buccal mucosa. Immediate treatment is not necessary but may consist of smoothing the sharp edge of the tooth with an emery board or rotary disk sander. The patient should be reassured that a dentist can restore the tooth to its normal appearance with composite resins and bonding material. Follow-up is important with these injuries because pulp necrosis and color change can occur in some cases (< 3%).

Ellis Class II Fractures

Uncomplicated fractures through the enamel and dentin are called Ellis class II fractures. Fractures that extend into the dentin are at higher risk for pulp necrosis and therefore need more
that may be applied to the surface of the tooth include CaOH, zinc oxide, skin adhesives, and glass ionomer composites. Some literature suggests that glass ionomer may be superior to other coverings; however, the difference is probably slight, and the increased cost of glass ionomer is not justified for routine use in the ED at this time. Bone wax and skin glue such as the cyanoacrylates are not recommended as dressings. Many dressings come as a base and a catalyst which require mixing, but are often available in premixed formulations as well. This is easily accomplished with a dental spatula and a mixing pad, which can be obtained from any dental supply house. A commonly used ED dressing is CaOH (Dycal [Dentsply Sirona, York, PA], Preline [Henry Schein, Melville, NY], or other similar products). Mix the catalyst and the base in equal portions, and place a small amount on the exposed area with an applicator such as a dental spatula or another appropriate instrument (Fig. 64.7). The fracture site must be dry before application of the CaOH to ensure adherence.

To dry the surface of the tooth, have the patient bite into gauze pads or mouth breathe. Likewise, connecting a nasal cannula to high-pressure wall O₂ and aiming it at the tooth will facilitate drying. CaOH will dry within minutes after being exposed to the moist environment of the mouth. Although aggressive treatment by the emergency clinician (see Fig. 64.6C).

The risk for pulp necrosis in these patients is less than 10%, but it increases as treatment time extends beyond 24 hours. These patients often complain of sensitivity to heat, cold, or forced air. Physical examination reveals the yellow tint of the dentin in contrast to the white hue of the enamel. With fractures closer to the pulp cavity, the dentin will have a pink tinge. The tooth is usually sensitive to percussion with a tongue blade. The porous nature of dentin allows passage of bacteria from the oral cavity to the pulp, which may result in inflammation and infection of the pulp chamber. This is more likely to occur after 24 hours of dentin exposure but occurs sooner if the fracture site is closer to the pulp. Likewise, patients younger than 12 years have a pulp-to-dentin ratio larger than that in mature adults and are at increased risk for pulp contamination. For this reason, younger patients should be treated aggressively and be seen by a dentist within 24 hours.

The goal of treating dentin fractures is twofold: to cover the exposed dentin and thus prevent secondary contamination or infection and to provide relief of the pain. After the tooth is covered, the dentist, using modern composites, can often rebuild the tooth directly over the calcium hydroxide (CaOH) cap that was placed in the ED. Perform supraperiosteal injection or a regional tooth block before any manipulation of the tooth. This will make application of the dressing easier because manipulation of the tooth will not cause discomfort. Dressings

---

**Figure 64.6** A, The Ellis classification for fractured alveolar teeth. The easiest method to classify fractured teeth is by description (e.g., fracture through the dentin of the first upper right molar). B, Ellis class I: only the enamel is fractured. These fractures pose no threat to the dental pulp and are not sensitive to temperature or forced air. C, Ellis class II: the crown fracture demonstrates involvement of the enamel and dentin, without exposure of the pulp. Immediate dental referral is necessary to prevent contamination of the pulp through the dentinal tubules. D, This crown fracture involves enamel, dentin, and the soft tissue of the pulp as well. Immediate dental referral is mandatory to save the tooth. (C and D, from Zitelli BJ McIntire SC, Nowalk AJ, editors: Zitelli and Davis' atlas of pediatric physical diagnosis, ed 6, St. Louis, 2012, Saunders.)
teeth, pulpotomy (partial removal of the pulp) as definitive treatment.5,9 The longer the pulp is exposed, the greater the likelihood of contamination and abscess formation. If a dentist cannot see the patient immediately, attempt to relieve the pain and cover the exposed pulp (Fig. 64.8). If significant pain is present, perform a supraperiosteal injection. Subsequently, placing dental foil over the CaOH dressing is recommended in some texts, it is not necessary if the patient plans to follow up with a dentist within 24 to 36 hours. It is also not technically easy to perform. To prevent dislodgment of the CaOH dressing, instruct the patient to eat only soft foods until seen by a dentist. Although there is no evidence to support antibiotic therapy for fractured teeth in the ED, many consultants recommend initial treatment with penicillin or clindamycin until definitive dental care can be obtained.13

Many patients who sustain a fracture through the dentin will require a root canal or other definitive endodontic treatment. Timely application of an appropriate dressing in the ED, however, may prevent contamination of the pulp and make root canal therapy unnecessary. As with any trauma to the anterior teeth, explain to the patient that disruption of the neurovascular supply is possible and that long-term complications such as pulp necrosis, color change, and resorption of the root might occur.

Ellis Class III Fractures
Complicated fractures involving the pulp are also known as Ellis class III fractures (see Fig. 64.6D). Complicated fractures of the crown that extend into the pulp of the tooth are true dental emergencies. These fractures result in pulp necrosis in 10% to 30% of cases even with appropriate treatment, so the vast majority of these injuries end up with root canal therapy.6 They may be distinguished from fractures of the dentin by the pink color of the pulp. Wipe the fractured surface of the tooth with gauze and observe for frank bleeding or a pink blush, which indicates exposure of the pulp. Fractures through the pulp are often excruciatingly painful, but occasionally there is lack of sensitivity secondary to disruption or concussion of the neurovascular supply of the tooth.

Immediate management includes referral to a dentist, oral surgeon, or endodontist. The patient usually requires pulpectomy (complete removal of the pulp) or, in the case of primary teeth, pulpotomy (partial removal of the pulp) as definitive treatment.5,9 The longer the pulp is exposed, the greater the likelihood of contamination and abscess formation. If a dentist cannot see the patient immediately, attempt to relieve the pain and cover the exposed pulp (Fig. 64.8). If significant pain is present, perform a supraperiosteal injection. Subsequently,
cover the tooth with one of the dressings described earlier. The bleed can be brisk at times. Control such bleeding by having the patient bite onto a gauze pad that has been soaked with a topical anesthetic containing a vasoconstrictor such as epinephrine. Alternatively, inject a small amount of anesthetic/vasoconstrictor into the pulp to control bleeding. After the covering is applied, instruct the patient to follow up as soon as possible with a dentist. Antibiotics with coverage directed at oral flora (e.g., penicillin, metronidazole, or clindamycin) should be considered, with recommendations to eat only soft food and to avoid mastication on the affected side. Removal of the pulp with specialized instruments by the emergency clinician is not recommended, although some authors have advocated this in the past. This procedure is the realm of the dental professional and is likely to result in complications if not done properly.

**Luxation, Subluxation, Intrusion, and Avulsion**

**Luxation and Subluxation**

*Subluxation* refers to teeth that are mobile but not displaced. *Luxation* refers to teeth that are displaced, either partially or completely, from their sockets. Luxation injuries are divided into four types (Fig. 64.9):

1. **Extrusive luxation** is an injury in which the tooth is forced partially out of the socket in an axial direction (see Fig. 64.9A).

2. **Intrusive luxation**, or intrusion, occurs when the tooth is forced apically. It may be accompanied by crushing or fracture of the tooth apex (see Fig. 64.9B).

3. **Lateral luxation** occurs when the tooth is displaced either facially, lingually, mesially, or distally (see Fig. 64.9C). This injury is often associated with injuries to the alveolar wall.

4. **Complete luxation**, also known as complete avulsion, results in loss of the entire tooth from the socket.

Even minor trauma to the oral cavity requires meticulous examination for loose or missing teeth. Examine each tooth for mobility by applying a back-and-forth motion on each side of the tooth surface with either the fingertips or two tongue blades. Any blood in the gingival crevice (area where the gingiva touches the tooth) suggests a traumatized tooth.

Teeth that are minimally mobile and are not displaced do very well with just conservative treatment. The tooth will tighten up in the socket if not retraumatized or reinfused in any way. Instruct patients to eat only a soft diet for 1 to 2 weeks and see their dentist as soon as possible. Note that a seemingly lost (avulsed) tooth may actually be deeply intruded into the soft tissue (Fig. 64.10).

Grossly mobile teeth require some form of stabilization as soon as possible. It is important to note that in certain patients with poor gingival health, luxated teeth may not be salvageable because of preexisting disease of the attachment apparatus. Stabilization is best performed by a dental specialist with enamel bonding material or wire ligation. Although many different “home remedies” exist for splinting loosened teeth in the ED, the clinician must be aware of the possibility of aspiration if the splint fails. Avoid the use of unapproved medications in the mouth. Splinting techniques are suitable for the emergency clinician to perform as temporizing measures until definitive care can be arranged. One simple technique for emergency use is to apply periodontal paste, commercially available as Coe-Pak (GC America Inc., Alsip, IL) or other similar commercial products (Fig. 64.11). These products usually consist of a base and a catalyst that when mixed, form a moderately sticky claylike dressing that becomes firm after application. It is applied over the enamel and gingiva, as well as the adjacent teeth, to splint the subluxed tooth into place. Although the splint performs best if placed on the facial and buccal surfaces of the teeth, it is usually sufficient to apply the paste only to the front (facial) surface of the teeth. Make sure that the gingiva and enamel are completely dry. Lubricate gloves with water or lubricating jelly before applying the dressing. Apply the dressing into the grooves between the teeth, as well as on the adjacent teeth. Remind the patient to eat a soft diet until seen in follow-up within 24 hours. Periodontal pastes are usually

---

**Figure 64.9** Classification of tooth trauma. A, Extrusive luxation occurs when the tooth is forced partially out of the socket in an axial direction. B, Intrusive luxation of a tooth compresses the periodontal ligament and vascular supply of the pulp. It may even crush the apical bone. C, Lateral luxation occurs when the tooth is displaced in a lingual, mesial, distal, or facial direction. Fractures of the alveolus frequently accompany lateral luxation injuries. (A–C, Adapted from King R: Orofacial infections. In Montgomery MT, Redding SW, editors: Oral-facial emergencies—diagnosis and management. Portland, OR, 1994, JBK Publishing.)
Intrusion of an upper tooth into the maxillary sinus can cause recurrent sinusitis. Teeth can also intrude into the nasal cavity and cause infection or bleeding, or they can be aspirated into the airway. The incisors are the teeth most commonly intruded.

Intruded teeth create the potential for infection or cosmetic deformities. Although certain cell culture media have been developed to stimulate cells of the periodontal ligament to proliferate and remain viable, milk and the commercially available Save-A-Tooth and EMT Toothsaver (Smart Practice, Inc., Phoenix, AZ) are the best and easiest for both prehospital care and ED storage (Fig. 64.12). Milk will preserve the periodontal ligament for 4 to 8 hours; the commercial products will preserve the ligament for 12 to 24 hours. However, reimplantation (Video 64.1) should take place at the earliest possible opportunity after the socket has been adequately prepared. The key is to get the tooth into the transport medium immediately because even 10 minutes outside some type of storage medium can cause desiccation and death of the periodontal ligament cells. Use saliva at the scene if milk, EMT Toothsaver, or

**Figure 64.10** Intruded tooth secondary to trauma. A, On superficial examination it appears that the tooth was simply knocked out. This missing tooth could be simply lost, fully intruded, aspirated, or swallowed. B, In some cases a dental radiograph or computed tomography scan is necessary to determine intrusion or avulsion. Intruded teeth create the potential for infection or cosmetic deformities. Intrusion of an upper tooth into the maxillary sinus can cause recurrent sinusitis. Teeth can also intrude into the nasal cavity and cause infection or bleeding, or they can be aspirated into the airway. The incisors are the teeth most commonly intruded.

Teeth that are luxated in either the horizontal or the axial plane or are slightly extruded can also be splinted with the techniques described earlier. It is important that the loosened tooth be in as perfect alignment as possible when the final adjustments are made at the dentist’s office. However, the alignment does not need to be as precise when the tooth is splinted in the ED. The important point is that the tooth be splinted adequately and follow-up ensured.

**Intrusion and Avulsion**

Intruded teeth are those that have been forced apically into the alveolar bone. This often results in disruption of the attachment apparatus or fracture of the supporting alveolar bone, especially in permanent teeth with mature roots. These teeth are usually immobile and do not require stabilization in the ED. Intruded teeth frequently require endodontic treatment because of pulp necrosis. It is important to consider the possibility of an intruded tooth anytime there is a space in the dentition (see Fig. 64.10). Undiagnosed intrusion of the teeth can lead to infection and craniofacial abnormalities. Obtain radiographs when it is uncertain whether a tooth is intruded or simply avulsed. Intruded teeth are best managed by a dentist or dental specialist; referral should take place within 24 hours. Permanent teeth often require repositioning and immobilization, but primary teeth are usually given a trial period to erupt on their own before any intervention is taken.

Primary teeth are not replaced after avulsion because they can fuse to the alveolar bone and potentially cause craniofacial abnormalities or infection. Reimplanted primary teeth may also interfere with eruption of the secondary teeth. The parents of these patients need to be reassured that a prosthetic replacement for the avulsed teeth can easily be made and worn until the permanent teeth erupt, if desired. See Figs. 64.1 and 64.2 as a guide to identifying permanent versus primary teeth.

Avulsed permanent teeth are those that have been completely removed from their ligamentous attachments. These are true dental emergencies. The majority of patients seen in the ED with an avulsed tooth will lose that tooth, so patient and physician expectations should not be overly optimistic. Under ideal circumstances, such as arrival at a dentist’s office with a properly stored tooth avulsed less than 60 minutes previously, this situation may result in successful reimplantation 80% to 90% of the time but often requires specialized procedures and endodontics (root canal). Emergency clinicians are not expected to save an avulsed tooth, but prompt action may give that replanted tooth some chance for survival.

The first consideration in treating dental avulsions is to ask, “Where is the tooth?” Missing teeth may have been intruded, fractured, aspirated, swallowed, or embedded into the soft tissues of the oral mucosa. Therefore radiographs should be considered anytime that an avulsed tooth cannot be located. Management of an avulsed tooth in the ED depends on a number of factors, including the age of the patient, the amount of time that has elapsed since the tooth was avulsed, associated trauma to the oral cavity such as alveolar ridge fractures, and the overall health of the periodontium. Time is the other important consideration when deciding whether to replace an avulsed tooth. In general, the longer the tooth is out of the socket, the higher the incidence of necrosis of the periodontal ligament and subsequent failure of reimplantation. Periodontal ligament cells generally die within 60 minutes outside the oral cavity if they are not placed in an appropriate transport medium. A significant amount of research has been conducted on different media used to keep cells of the periodontal ligament alive. Various transport media have been studied, including milk, Hank’s Balanced Salt Solution (Thermo Fisher Scientific, Waltham, MA), Save-A-Tooth (Phoenix Lazarus, Portland OR), saliva, cell culture media, and water. Although certain cell culture media have been developed to stimulate cells of the periodontal ligament to proliferate and remain viable, milk and the commercially available Save-A-Tooth and EMT Toothsaver (Smart Practice, Inc., Phoenix, AZ) are the best and easiest for both prehospital care and ED storage (Fig. 64.12). Milk will preserve the periodontal ligament for 4 to 8 hours; the commercial products will preserve the ligament for 12 to 24 hours. However, reimplantation (Video 64.1) should take place at the earliest possible opportunity after the socket has been adequately prepared. The key is to get the tooth into the transport medium immediately because even 10 minutes outside some type of storage medium can cause desiccation and death of the periodontal ligament cells. Use saliva at the scene if milk, EMT Toothsaver, or
• If the tooth cannot be reimplanted successfully in the field, place it in a transport medium as described earlier. Do not transport the tooth in the oral cavity such as inside the cheek because of the risk for aspiration. This location is also not ideal for keeping the periodontal ligament alive because of the bacterial flora and low osmolality of saliva.

Once the patient arrives in the ED, confirm proper placement and alignment. It is not important that the tooth be in perfect position because the dentist can make final adjustments. Splinting the repositioned tooth with periodontal paste or composite as outlined earlier may be necessary if mobility is present.

Should reimplantation not be successful in the prehospital setting, it must be done in the ED according to the following guidelines:

• Determine whether this is a permanent tooth. By 14 years of age, all primary teeth should have been replaced by permanent teeth.
• Handle the tooth by the crown only because handling the tooth by the root can damage the periodontal ligament.
• Do not replace the tooth if it is fractured or if significant maxillofacial trauma has occurred such as an alveolar ridge fracture.
• If the tooth can be replaced in the prehospital setting, gently rinse off the root first to remove any debris. Do not wipe off the root because this removes the periodontal ligament.

Save-A-Tooth is not available. The patient should reimplant the tooth in the prehospital setting if possible. Do not have the patient place the tooth in the cheek or lip as this risks aspiration. The principles cited here should be followed when providing instructions to prehospital providers or to a patient who calls for advice. It is always preferable to refer such patients directly to a dentist rather than the ED if this is practical. Patients should be requested to do the following:

Dental Splint (Coe-Pak) Application

Coe-Pak is a periodontal paste used to splint loosened or avulsed teeth. Like calcium hydroxide, it is supplied as a catalyst and a base.

Mix equal parts of the catalyst and base on the mixing pad supplied with the product.

Roll the claylike mixture between your fingers into an elongated roll. Apply lubricating jelly to your gloves prior to handling the product to prevent it from sticking.

Dry the teeth and gingiva with gauze prior to application. Press the Coe-Pak into the grooves between the teeth, as well as across the adjacent teeth and gingiva. This type of splint works best when applied to both the front and back surfaces of the teeth, although it is usually sufficient to apply it only to the front.

Figure 64.11 Dental splint (Coe-Pak, GC America Inc., Alsip, IL) application.
Ideally, the patient is immediately referred to a dentist, and the reimplanted tooth is held in place by biting on gauze. The tooth may be temporarily held in place with periodontal paste or another similar product (see Fig. 64.11) or the technique described in Fig. 64.13. This system prolongs the time to successful reimplantation but does not ensure success.

- Perform supraperiosteal injection with a local anesthetic before manipulating or replacing teeth to make the procedure more comfortable for the patient and easier to perform.
- Check the oral cavity for trauma. If an alveolar ridge fracture is present or the socket is significantly damaged, do not reimplant the tooth.
- Gently suction the socket first with a Frasier suction tip to remove any accumulated clot. Be careful to not damage the walls of the socket because this can further damage periodontal ligament fibers. Irrigate gently after suctioning. If the clot is not removed, reimplantation and realignment will be difficult. Rinse off any debris on the tooth with saline but do not scrub it. Implant the tooth into the socket with firm, but gentle pressure. Remember to handle the tooth only by the crown.
- Ask the patient to bite down gently on gauze to help align the tooth. The tooth may require splinting after replacement. This can be performed in the ED or urgent care clinic with periodontal pastes.
- Update the patient’s tetanus status as necessary.
- Prescribe a liquid diet until the patient is seen in follow-up. Antibiotics are controversial in the treatment of fractured and avulsed teeth. Although the American Association of Endodontics does not recommend the routine use of antibiotics for fractures or avulsions, other authors recommend the use of antibiotics effective against mouth flora (e.g., penicillin, clindamycin) to decrease inflammatory resorption of the root. It is probably reasonable to use antibiotics if the root or socket is heavily soiled; otherwise, treatment should be tailored to the individual patient and discussed with a knowledgeable and experienced consultant.

Prognosis

The prognosis of a reimplanted tooth depends on many things. As discussed earlier, the time until reimplantation is critical. Likewise, the age of the patient, the stage of development of the root (younger is better), and the overall health of the gingiva are also very important. An individual with gingival disease is more likely to have an unsuccessful reimplantation.

Figure 64.12 With the A, Save-A-Tooth (Phoenix Lazarus, Portland OR) or B, EMT Toothsaver (SmartHealth, Phoenix, AZ) systems, an avulsed permanent tooth is placed into the container and closed. Avulsed primary teeth are not replanted. The preservative will increase the lifespan of traumatized periodontal ligament cells. Unpreserved teeth replanted after 60 minutes rarely survive; those that do require root canal procedures and close follow-up. This system prolongs the time to successful reimplantation but does not ensure success.

Figure 64.13 Temporary suturing to hold a replanted tooth in place. A, Use a silk suture. Start by puncturing the gingiva at the border of the replaced tooth (1). Bring the suture behind the tooth and then cross over the front of the tooth to the other side. Penetrate the gingiva (2), go behind the tooth, cross over the front again, and tie the suture (A). B and C, Multiple teeth can be reimplanted.

Ideally, the patient is immediately referred to a dentist, and the reimplanted tooth is held in place by biting on gauze. The tooth may be temporarily held in place with periodontal paste or another similar product (see Fig. 64.11) or the technique described in Fig. 64.13.
The goal in any tooth avulsion or fracture is to keep the native tooth if at all possible. A tooth that has been avulsed and reimplanted usually loses the majority of its neurovascular supply and undergoes pulp necrosis, necessitating root canal therapy. However, if the periodontal ligament remains intact, there is a greater chance of a functional tooth. It is important that the patient be aware that some root resorption is always going to occur after reimplantation and that loss of the tooth might occur.

**Alveolar Bone Fractures**

Trauma involving the anterior teeth may be associated with fracture of the alveolus, which is the tooth-bearing portion of the maxilla or mandible. Alveolus or alveolar ridge fractures often occur in multitooth segments and will vary in the number of teeth involved, the amount of displacement, and the mobility of the affected segment. The patient generally complains of pain, as well as malocclusion. The diagnosis is usually clinically apparent and is notable for a section of teeth that are misaligned and variable in mobility (Fig. 64.14). Avulsed teeth, fractured teeth, or displaced teeth may be present within the alveolar segment itself. Dental bite-wing radiographs confirm the diagnosis. In the ED, Panorex or facial x-ray films may show the fracture line just apical to the root of the involved teeth; however, these films are often inconclusive or have normal findings. Facial computed tomography (CT) may provide additional information (Fig. 64.15).

Treatment of alveolar ridge fractures involves rigid splinting after repositioning the involved segment. The displaced alveolar ridge should not be repositioned by the emergency clinician as this often converts a nonmobile displaced segment into a mobile displaced segment. Repositioning is usually beyond the scope of the emergency clinician, and urgent consultation with an oral surgeon or dentist is necessary. The role of the emergency clinician is to identify the injury, as well as any avulsed or fractured teeth, and preserve as much of the alveolar bone and surrounding mucosa as possible. Alveolar bone that is lost, debrided, or missing is difficult to restore properly. Alveolar ridge fractures, if not mobile or bleeding, may be discharged if urgent follow-up is arranged for within 24 hours. If bleeding or movement of the segment is a risk, specialty consultation will need to take place within the hospital.

**Lacerations and Dentoalveolar Soft Tissue Trauma**

Trauma to the face and perioral region is often associated with soft tissue injuries such as abrasions or lacerations. Before any repair can take place, thoroughly inspect all wounds and abrasions to determine the extent of the wound and whether foreign bodies are present. Through-and-through lacerations are easily overlooked, as are small foreign bodies and debris such as tooth fragments. Obtain radiographs if there is any question of tooth fragments. Evaluate the patient for potential airway compromise.

As a general rule, repair injured teeth before undertaking soft tissue repair because manipulating the soft tissue while repairing teeth may damage sutures already in place in the soft tissues. Begin repair in the perioral region with standard wound care. After appropriate local or regional anesthesia, débride devitalized, crushed, or macerated tissue. Irrigate profusely. The role of antibiotics in mucosal trauma has not been conclusively established, and there is no definitive standard of care. Several studies suggest a minimal benefit; however, this remains to be completely proved. A reasonable guideline to follow is to use antibiotics if a significant amount of devitalized or crushed tissue is present, if there is significant wound contamination, or if the wound is a through-and-through laceration. Coverage of oral flora (e.g., penicillin, metronidazole, or clindamycin) is fine for mouth lacerations, and additional skin coverage (e.g., clindamycin, or dicloxacillin) should be considered for through-and-through lacerations. Dentoalveolar trauma may present the emergency clinician with several different situations that should generally be approached as follows.

**Buccal Mucosa**

Most small lacerations and abrasions of the buccal mucosa heal quickly and rapidly without repair, but large lacerations (> 1 to 2 cm) should be repaired. Use any absorbable suture such as chromic gut or Vicryl (Ethicon US LLC, Somerville, New Jersey).
NJ) in the mouth, but place the sutures so that the knots are buried. Silk is an alternative but has higher reactivity and is nonabsorbable. Avoid using nylon because it is sharp and irritating to tissues.

Through-and-through lacerations of the oral cavity present a special situation. Evaluate for damage to the salivary ducts (Wharton's duct and Stensen's duct) and to the facial nerve. If they are intact, proceed with the repair. Guidelines for closure are controversial, but larger lacerations (> 1 to 2 cm) should generally be closed. Close the mucosa with absorbable sutures as noted earlier, and close the skin aesthetically with 6-0 nylon, Prolene (Ethicon), or a rapidly absorbable suture. Close the mucosa first so that the skin repair is not disturbed. If the mucosal wound is small or is a puncture wound, it is reasonable to close only the skin layer. Refer very large, gaping, or complicated lacerations to an oral surgeon.

Recheck large or through-and-through lacerations of the oral cavity in 2 to 3 days. Remove nonabsorbable sutures in 7 to 10 days. Advise the patient to rinse four to six times a day with saline solution. Prescribe a soft diet. Apply a topical skin antibiotic for 24 to 48 hours.

**Gingiva**

Small lacerations of the hard gingiva overlying the maxillary or mandibular alveolus usually heal uneventfully without repair. If the laceration is large, if a flap is present, or if bone is exposed, approximate the gingiva with a 4-0 or 5-0 Vicryl (Ethicon) or Dexon (Medtronic, Minneapolis, MN) suture. As mentioned earlier, silk is another option. It is difficult to suture gingiva because there is little supporting soft tissue underneath. A helpful technique is to wrap the suture around the teeth circumferentially and use the teeth as anchors (Fig. 64.16). Large lacerations should be repaired to approximate the gingiva and cover the base of the teeth (Figs. 64.17 and 64.18). Nongapping, nonbleeding gingival lacerations may be covered with periodontal paste as described earlier in lieu of suturing, as long as dental or maxillofacial follow-up is assured.

**Frenulum**

The maxillary frenulum rarely requires sutures for simple lacerations. If the laceration is extensive or extends significantly into the surrounding mucosa or gingiva, approximate it with chromic, Vicryl (Ethicon), or Dexon (Medtronic) suture. These wounds are often significantly painful. Prescribe analgesic medications even if the wound does not require suturing. The lingual frenulum is very vascular in nature and will often need a suture or two to control hemostasis. Use a local anesthetic with a vasoconstrictor to aid in hemostasis while the wound is repaired.

**The Tongue**

Tongue lacerations are challenging (Video 64.2). Although they may be tempting to suture, most large lacerations of the body of the tongue, such as those that occur from a seizure, will heal well without suturing (Fig. 64.19). Tongue lacerations that have the wound edges approximated do not need to be sutured. Repair larger lacerations that gape because the cleft left by the wound will epithelialize and leave a grooved, bifid, or lateral flap appearance. Approximate wounds that are bleeding profusely, are flap shaped, involve muscle, or are on the edge of the tongue. Small avulsions (divots) or those in the center of the tongue usually heal without intervention.
Bleeding from the oral cavity is not unusual and is most commonly associated with dental procedures, such as extracted teeth. It is important to ascertain whether any recent dental work has been performed and what was done. Spontaneous bleeding of the gingiva or oral cavity not associated with dental manipulation or trauma is suggestive of advanced periodontal disease or an underlying systemic process. Ask the patient about other medical conditions that predispose to bleeding (e.g., liver disease, platelet abnormalities), as well as historical factors that may suggest a bleeding abnormality or clotting factor deficiency. Find out whether the patient is taking aspirin or other anticoagulants. Consider laboratory testing if pathologic coagulopathy is a significant concern, but not routinely in a patient seen after dental manipulation.

Control gingival bleeding after scaling or minor dental procedures with direct pressure and saline or hydrogen peroxide rinses. Persistent bleeding from the gingival areas despite pressure and rinses raises suspicion of a bleeding abnormality. A much more common cause of oral hemorrhage seen in the ED is postextraction bleeding. Minor oozing after dental extractions, such as wisdom tooth extraction, is normal for 2 to 4 days after surgery, but many patients get concerned when the bleeding persists despite warnings from the oral surgeon. These patients usually go to the ED when their dentist cannot explain the procedure in detail to the patient before repairing these wounds. Ask an assistant to secure the tongue by holding it with gauze. If the tongue cannot be secured in this manner, apply a towel clip to the end of the anesthetized tongue. Children with tongue lacerations that need repair generally require sedation or repair by a specialist in the surgical suite, but many of these lacerations are small and heal uneventfully on their own.

Begin the repair with either local injection of an anesthetic or a lingual block (see Chapter 30). To promote hemostasis, infiltrate locally with lidocaine with epinephrine. Use absorbable sutures such as 4-0 chromic, Vicryl (Ethicon), or Dexon (Medtronic). Silk can be used, but it must be removed in 7 to 10 days. Do not use nylon because it is very irritating to the surrounding tissues. For lacerations extending through muscle, close with one deep stitch penetrating both the mucosa and the muscle. When possible, bury the knots of absorbable suture because they will often work their way loose. Full-thickness lacerations can be closed in a number of ways. Place a suture through all three layers or close the top mucosa and muscle together and do the same thing on the underside of the tongue. Bleeding from large lacerations is almost always controlled with primary repair. In some instances, hemostasis can be achieved without the use of sutures by using Gelfoam (Pharmacia and Upjohn Company, Kalamazoo, MI) impregnated with topical thrombin (Fig. 64.20).
If the bleeding persists, insert a coagulation sponge, such as Gelfoam (Pharmacia and Upjohn Company), into the socket and then loosely close the gingiva surrounding the socket with a 3-0 absorbable figure-of-eight suture. Instruct the patient to bite down on gauze placed over the sutures. Soaking Gelfoam with topical thrombin before placement is a good way to halt minor persistent bleeding (see Fig. 64.20). Newer hemostatic dressings such as the Hemcon dental dressing (Tricoll Biomedical Inc., Portland, OR) are specifically designed for postextraction and oral bleeding. This shrimp-based bandage forms a sticky matrix when it contacts blood, and it quickly forms a seal that stops the bleeding. Other hemostatic agents that should be considered include cellulose or collagen (Surgicel [Ethicon], Avitene [Bard Davol, Warwick, RI]). Tranexamic acid used as a soaked pledget (500 mg/5 cc) has also been recently described as being very useful for oral and nasal bleeding. If these measures fail to control the bleeding, consult a specialist. It is also reasonable to check blood counts and coagulation profiles at this time.

Patients whose bleeding is controlled can be discharged and instructed not to take anything by mouth for 4 hours and then only liquids and soft foods. If silk sutures are used, remove them in 7 days.

**ALVEOLAR OSTEITIS (DRY SOCKET)**

The pain associated with an extracted tooth is significant but usually manageable with current pharmacologic modalities.
The pain associated with a dry socket, however, can be very severe and often requires more definitive treatment. Alveolar osteitis, or dry socket (Video 64.3), is a localized inflammation that occurs when the alveolar bone becomes inflamed. This condition usually occurs when the clot that is normally present in the socket after a tooth extraction becomes dislodged or dissolves. It is most common in the 2- to 4-day period after a tooth extraction. The examination is essentially unremarkable with the exception of a missing clot where the tooth was extracted. Signs and symptoms of dry socket include the following:
1. Moderate to severe pain localized to the area or frequently radiating to the ear.
2. A foul odor or taste in the absence of purulence or suppuration.
3. Symptoms that occur 3 to 5 days after tooth extraction.
4. Absence of swelling, purulence, or lymphadenitis.
5. Duration of 5 to 40 days.

Anything that increases negative intraoral pressure in the mouth (e.g., smoking, excessive rinsing, spitting, drinking from a straw), as well as hormone replacement and periodontal disease, will predispose a patient to a dry socket. In only a small percentage of patients (2% to 5%) will a dry socket develop; however, this number increases with traumatic extractions or impacted third molars.13,19

The following contribute to the development of a dry socket:
1. Excessive trauma during extraction.
2. Inadequate blood supply to the extraction site.
3. Preexisting localized infection.
4. Loss of clot from sucking, straw use, rinsing, or smoking.
5. Foreign bodies remaining in the socket.
6. Use of oral contraceptives.
7. Use of corticosteroids.
8. Pericoronalitis.

The pain associated with a dry socket is extremely severe, and if a patient is seen several days after an extraction with relatively normal findings on examination and severe pain, it is probably a dry socket. It must be distinguished from osteomyelitis, which is characterized by fever, leukocytosis, malaise, and nausea. The pain related to a dry socket will not be relieved with traditional pain medications, but a dental block usually provides instant relief. Once the block is performed, the alveolar osteitis can be treated. Irrigate the socket and gently suction out any accumulated debris. Next, fill the socket to prevent recurrence of pain and allow healing to begin. A variety of materials are suitable to fill the socket again.

Gauze (3/4 inch) impregnated with eugenol (oil of cloves) or a local anesthetic may be used. The gauze should be packed into the socket to the level of the gingival cusps. Replace the gauze in 24 to 36 hours because it tends to dry out and loosen. The patient should be seen by a dentist the next day if at all possible. The socket may also be packed with a slurry of Gelfoam (Pharmacia and Upjohn Company) and eugenol. The Gelfoam acts as a matrix to hold the eugenol in the socket. A commercial product, such as Dry Socket Paste (Sultan Healthcare, York, PA) or Dressol-X (Rainbow Speciality and Health Products Inc., Niagara Falls, NY) (Fig. 64.21), can also be applied by itself into the socket or mixed with Gelfoam and placed into the socket. Dry Socket Paste is a very sticky thick paste containing eugenol. It may stay in place longer than gauze and does not dry out. Whichever packing material is used for a dry socket, one or more packings might be necessary before healing is complete; therefore, the patient must be referred back to see the dentist who performed the extraction if possible.

Although antibiotics may be given to prevent alveolar osteitis, they are not usually necessary once the socket has been packed and should be prescribed at the discretion of the patient’s oral surgeon or dentist.11-13 NSAIDs should also be prescribed because they seem to work better than narcotics for dry socket.20

**DENTALVEOLAR INFECTIONS**

Infections of the oral cavity run the spectrum from minor, easily managed abscesses to severe, life-threatening, deep space infections that require airway management and operative drainage. Although dental infections of all severity are encountered in the ED, the most common are those related to pulp disease. Others are associated with the attachment structures of the teeth such as the gingiva, periodontal ligament, and alveolar bone. These infections are often chronic conditions, but they can progress to the point where periodontal abscesses form and emergency treatment is required.

Emergency clinicians will be called on to drain abscesses of dental origin that do not extend into the deep spaces and that have well-defined boundaries easily accessible by intraoral or external drainage.

**Disease of the Pulp**

Disease of the pulp can occur as a result of trauma, operations, or other unknown causes, but the most frequent cause is invasion of microorganisms after carious destruction of the enamel. As the enamel is destroyed, caries progresses more rapidly through the dentin and into the pulp chamber and causes an inflammatory response referred to as pulpitis. If the path of carious destruction through the tooth is adequate for drainage of the developing inflammation, the patient may be only mildly symptomatic or even asymptomatic for a long time. If drainage is blocked, however, the process may progress to rapidly involve the entire pulp cavity and the periapical space. The tooth is usually exquisitely tender at this point. Abscesses in the periapical region are generally picked up on dental x-rays and less commonly on a Panorex film. However, unless extension through the cortex exists, it is not important for the emergency clinician to make the distinction between pulpitis and a periapical abscess. Examination may reveal gross decay of one or
Disease of the Periodontium

Periodontal disease is also very common and affects practically all adults to some degree. Periodontal disease refers to infection of the attachment apparatus of the teeth: the gingiva, periodontal ligament, and alveolar bone. Unlike pulpal disease, periodontal disease is not usually symptomatic and is therefore rarely a primary reason to seek treatment in the ED. Gingivitis is an inflammation of the gingiva caused by bacterial plaque. In advanced disease, the gingiva becomes red and inflamed and tends to bleed easily. With chronic periodontal disease, an abscess can form when organisms become trapped in the periodontal pocket. The purulent material usually escapes through the gingival sulcus; however, it occasionally invades the supporting tissues, the alveolar bone, and the periodontal ligament (periodontitis). Periodontal abscesses that are not draining spontaneously through the sulcus can be drained in the ED. Saline rinses are encouraged to promote drainage. Antibiotics should be reserved for severe cases or for abscesses that cannot be drained. If it is uncertain whether the abscess originates from the pulp or the periodontium, prescribe antibiotics even if the abscess has been drained. Pericoronitis is a localized inflammation that occurs when the gingiva overlying erupting teeth becomes traumatized and inflamed. Third molars are especially susceptible; however, any tooth can be affected. The gingiva overlying the crown may entrap bacteria and debris, and infection may subsequently develop. Typical signs of inflammation and infection may develop, including erythema, edema, pus, and foul breath. Examination of the overlying gingiva with a tongue blade or finger will elicit tenderness and may produce drainage from the infection underlying the tissue flap. The pain may be moderate to severe, and referral of the pain to an ear region is common. The localized infection occasionally spreads to deeper areas such as the pterygomandibular or submasseteric spaces. Clinically, patients with significant spread of a periapical infection will have trismus secondary to irritation of the masseter and pterygoid muscles.

ED treatment of pericoronitis is directed at detecting regional spread to the deeper spaces. Trismus or other systemic signs of advanced infection require intravenous antibiotics and urgent consultation for drainage procedures, which usually requires extraction of the offending tooth. If pericoronal infection is localized, local or nerve block anesthesia is followed by removal of submucosal debris. Saline rinses and oral antibiotics are prescribed along with dental follow-up in 24 to 48 hours.

Drainage of Dentoalveolar Infections

The important determination for the emergency clinician to make is whether the odontogenic infection is localized, confined, and easily accessible or whether it is complex and involves several potential spaces. Likewise, determine whether the patient appears to be in a toxic state, has trismus, or exhibits any signs of airway compromise. Patients not meeting these criteria require specialist referral. Dental infections are not always obvious and can be mistaken for sinusitis, or vice versa (Figs. 64.22 and 64.23).

Several anesthetic techniques can make the drainage process more comfortable. Nerve blocks and local injections work best. Benzocaine 20% gel, 5% lidocaine gel, or a combination of lidocaine, prilocaine, and tetracaine generally provide good topical anesthesia before injection (Fig. 64.24). Application of these medications to the dry mucosa before injection of the local anesthetic decreases the pain associated with injection. After applying the topical anesthetic, slowly infiltrate local anesthetic with a vasoconstrictor until the tissue blanches. Either a short-acting anesthetic (2% lidocaine) or a longer-acting anesthetic (0.5% bupivacaine) may be used, depending on the clinical circumstances. Regional or dental blocks may be performed instead of local infiltration if needle placement would track already infected tissues into healthy areas. Otherwise, consider local infiltration over the site of the abscess. Instruments necessary for drainage of dentoalveolar abscesses are those usually found on a standard incision and drainage tray and include hemostats, scalpel (No. 15 or 11 blade), packing material (¼-inch gauze), and a fenestrated Penrose drain.

Intraoral Technique

Intraoral abscesses do not routinely require any antiseptic mucosa preparation before drainage. After anesthetizing the region, make a small incision (0.5 to 1.0 cm) over the area of fluctuance while keeping the point of the blade directed toward the alveolar bone. Use a hemostat to bluntly dissect the abscess and break up any loculations. Cultures are not necessary unless the patient is immunocompromised. Irrigate the wound profusely with normal saline. If the wound is large enough to place a drain or gauze inside, tack one end to the mucosa with a silk suture to prevent aspiration. Advise the patient to perform saltwater rinses hourly and arrange follow-up in 24 to 48 hours with a dentist or oral surgeon to remove the drain and provide continued management. Because the source of the abscess is not always known to the emergency clinician, prescribe antibiotics.

Extraoral Technique

Most simple dental infections can be drained intraorally, but occasionally, an abscess spreads to the face and requires drainage through the skin. It is important to realize that most dental infections should be drained through the mouth, if possible, because any extraoral drainage will cause some scarring. Never make any incisions on the face in direct proximity to important structures such as the facial nerve or the parotid gland and...
speech or swallowing, and trismus. Although infections of certain teeth usually spread to particular contiguous spaces, the rapid spread of these infections often makes localizing the exact space difficult. Any space, including the buccal, temporal, submasseteric, sublingual, submandibular, parapharyngeal, and others, may be involved (Fig. 64.26).

Maxillary extension of periapical abscesses can spread into the infraorbital space and, subsequently, to the cavernous sinus through the ophthalmic veins and result in cavernous sinus thrombosis. Cavernous sinus involvement is associated with periorbital cellulitis, as well as meningeal signs or a decreased level of consciousness. Periapical infections of the anterior

**Deep Space Infections of the Head and Neck**

It is not unusual for odontogenic infections to spread into the various potential spaces of the face and neck (Fig. 64.25). The signs and symptoms consist of fever, chills, pain, difficulty with
mandibular teeth often spread to the buccinator space or the sublingual space, whereas those of the mandibular molars spread into the submandibular space.

The submandibular space connects with the sublingual space. Infection involving both these spaces is known as Ludwig’s angina, which can be life-threatening. The source is usually a decayed lower tooth, with the tooth itself being relatively asymptomatic (Fig. 64.27). Initially, this infection may be subtle, but it can progress rapidly. As the infection progresses, the submandibular, submental, and sublingual spaces all become edematous, and there may be elevation of the tongue and the soft tissues of the mouth (Fig. 64.28). This is not a simple abscess that can be readily drained. It is considered a true oral emergency. Interventions include definitive airway management, intravenous antibiotics, and emergency ear, nose, and throat (ENT) or oral surgery consultation for emergency operative intervention. Patients with established cases of Ludwig’s angina are admitted to the hospital and undergo operative intervention often involving multiple drains. Minor early cases can be treated and observed for 6 to 8 hours in the ED with close follow-up if the infection is deemed benign, but care must be taken as Ludwig’s angina sometimes presents rather unimpressively. The soft tissues of the posterior aspect of the pharynx can also become involved. Securing the airway becomes of paramount importance. The suprahyoid region of the neck appears tense and indurated, and landmarks may be obscured. A CT scan is usually used to clarify deep space involvement, but emergency oral and maxillofacial (OMF) or ENT consultation, initiation of antibiotic therapy, and definitive airway management should be instituted immediately. This rapidly progressive disease can cause unexpected or rapid-onset airway compromise, and surgical intervention is warranted, as is consultation with the team that handles difficult airways.

Treatment of complicated odontogenic head and neck infections, including Ludwig’s angina, centers on airway management, surgical drainage, and antibiotics. If it is uncertain whether the deep spaces are involved, obtain a CT scan to delineate any extension of the infectious process, but be sure to involve the appropriate consulting services as soon as possible. Perform airway interventions early if there is any question of compromise, and consider tracheostomy. Consult an OMF or ENT surgeon because drainage and removal of necrotic tissue might be necessary. Administer antibiotics to slow spread of the infection and decrease hematogenous dissemination. The bacteria involved are typically polymicrobial, involving both aerobic and anaerobic bacteria from the primary source. There has been an emergence of β-lactamase–producing organisms in upward of 40% of isolates from odontogenic neck abscesses. Antimicrobials of choice for complicated odontogenic infections usually include the penicillins. The expanded-spectrum penicillins (ampicillin-sulbactam, ticarcillin–clavulanic acid, piperacillin-tazobactam) are effective against β-lactamase–producing bacteria and also

Figure 64.24 Apply a dollop of 20% benzocaine gel to dry mucosa before injection of local anesthetic to decrease the pain of infiltration.

Figure 64.25 A, Location of temporal space abscesses. B, Route of infection into the buccal space, vestibular spaces, submandibular space, sublingual space, and palatal space. (From Eisele D, McQuone S, editors: Emergencies of the head and neck. St. Louis, 2000, Mosby.)
Advanced dental infection with characteristic facial swelling, probably involving the masseter space. Local anesthetic (lidocaine with epinephrine) is injected with a 27-gauge needle into an area of obvious fluctuance (shown by the probe). A mandibular nerve block is an alternative to local injection. After the site is punctured with a No. 11 blade, a hemostat is inserted into the cavity and spread. Copious pus is drained with a suction catheter. Initial intravenous antibiotics (clindamycin), oral antibiotics, and outpatient follow-up yielded good results. The offending tooth was extracted when the infection was controlled.

Ludwig's angina may initially appear to be benign. This patient's main complaint was dental pain. Tooth No. 27 (arrow) was noted to be decayed and a probable source of the infection. Physical examination revealed erythema and edema of the submandibular space, consistent with early Ludwig's angina. She was admitted to the hospital for intravenous antibiotics (clindamycin). A high index of suspicion and early treatment are mandated in such patients because this infection can progress rapidly. This patient had the beginnings of an infection as manifested by elevation of the tongue and swelling of the floor of the mouth, which characteristically raises the openings of the submandibular (Wharton's) ducts (arrow). The cause was a badly decayed lower tooth (arrow). Tenderness and induration can be palpated in the floor of the mouth.
Regional dental supply houses are good sources for these items. The Dental Box is a commercially available kit that contains many of these items and is designed for ED use (Fig. 64.29). It is available at https://www.thedentalbox.com. Dentrauma is another dental emergency kit available to EDs.

**DENTAL MATERIAL**

As a general rule, EDs should have a well-stocked supply of basic dental material. Many commercially available products can be used interchangeably with many of the items listed here. These products can often be kept in a cart or in another appropriate location. The following is a basic list:

1. Packing gauze.
2. Dental roll gauze.
3. CaOH paste, glass ionomer cement, or zinc oxide cement.
4. Dry socket medicament or eugenol.
5. Topical anesthetic gel (20% benzocaine or 5% lidocaine).
6. Periodontal paste (Coe-Pak [GC America Inc.], others) or self-cure composite.
7. Articaine (Septocaine, SEPTODONT, Louisville, CO) cartridges with epinephrine.
10. Ringed injection syringe.
12. Oral surgery tray with arch bars and ligature wires.
13. Tongue blades and cotton-tipped applicators.
14. Gelfoam (Pharmacia and Upjohn Company), HemCon (Tricol Biomedical), Surgicel (Ethicon), or topical thrombin.

Regional dental supply houses are good sources for these items. The Dental Box is a commercially available kit that contains many of these items and is designed for ED use (Fig. 64.29). It is available at https://www.thedentalbox.com. Dentrauma is another dental emergency kit available to EDs.

**INTRAORAL PIERCING**

Dating back to antiquity, body piercing has been practiced in many countries and cultures as part of ceremonial and religious rites. In recent years, body and intraoral piercing has gained tremendous popularity throughout the world as a means of self-expression. In the ED, clinicians may see patients with piercings of the lips, tongue, and even uvula. Complications of oral piercing on the gums and teeth include pain, infection, bleeding, increased salivary flow, difficulty swallowing, gingival recession, gingival trauma, and chipped or fractured teeth. Repeated trauma from a metal tongue ball can be quite detrimental to teeth and cause microfractures and a sudden shattered tooth (Fig. 64.30). One study of 400 consecutive patients at a military dental office found that 20% of their
patients had at least one type of oral piercing, with the tongue being the most common location. Of these patients, 14% had fractured teeth and 27% had recession of the gums. Another study assessed the impact of time and found that tooth chipping was found on molars and premolars in 47% of patients who had a tongue piercing for longer than 4 years.

**ACKNOWLEDGMENTS**

The contributions of Robert H. Benko, DDS, to this material are greatly appreciated. The editors and author also wish to acknowledge the contributions of James T. Amsterdam to this chapter in previous editions and would like to thank Michael B. Pavel, DMD, for his review of this manuscript.

**REFERENCES ARE AVAILABLE AT** [www.expertconsult.com](http://www.expertconsult.com)
REFERENCES

Special Procedures

Procedures Pertaining to Hypothermia and Hyperthermia
Heather M. Prendergast and Timothy B. Erickson

PROCEDURES PERTAINING TO HYPOTHERMIA

With an increase in outdoor activities, changing weather patterns, and the growing epidemic of homelessness in our country, issues pertaining to hypothermia remain in the forefront. Hypothermia is not only a common diagnosis in rural areas but has also become more commonplace in urban centers across the nation secondary to inadequate housing or lack of preparation for cold weather changes. It is also important to note that numerous cases of accidental hypothermia (AH) are reported each year in areas typically considered warm weather locales such as Florida, Texas, California, Alabama, and even the Sahara desert. Every year, many recreational and elite athletes participate in outdoor sporting events. The higher the environmental stress, the greater the potential for failure in performance and the development of hypothermia. The high-altitude expeditions on Mt. Denali in 2003, Mt. Hood in 2006, and Mt. Everest in 1996, 2014 and 2015, are reminders that even well-protected, acclimatized individuals can succumb to cold-related fatalities.

Neonates are at particular risk of hypothermia due to large surface area-to-mass ratio, deficient subcutaneous tissue and inefficient shivering. Neonates also lack behavioral defense mechanisms. Acute neonatal hypothermia is common after emergency delivery or resuscitation and has been reported in cases of infant abandonment. Optimal treatment of hypothermia remains controversial. It is a well-accepted practice to carry out resuscitation of these individuals for extended periods. The medical literature contains numerous anecdotal reports of profoundly hypothermic individuals who are successfully resuscitated and discharged neurologically intact, the longest being in cardiac arrest for 8 hours and 40 minutes. Despite these spectacular reports of survival, both morbidity and mortality from hypothermia are common. Between 1972 and 2002, 16,555 deaths in the United States were attributed to hypothermia, which equates to 689 deaths per year. The United States Centers for Disease Control and Prevention (CDC) reported that in the United States during the years of 1999–2011, there were an average of 1301 deaths from environmental cold exposure per year. The actual number of patients seen in emergency departments (EDs) with hypothermia is unknown. Poverty, homelessness, alcoholism, and psychiatric illnesses are commonly associated conditions. Ethanol intoxication is the most common cause of excessive heat loss in urban settings. This chapter critically reviews approaches and procedures appropriate to the management of several categories of hypothermic patients. The recommendations combine treatment efficacy with safety. Before describing procedures and making recommendations, essential terms are defined and the pathophysiology of hypothermia is briefly reviewed.

Definitions

Accidental hypothermia has been defined as an unintentional decrease in core (vital organ) temperature to below 35°C (< 95°F). Victims of hypothermia can be separated into the following categories: mild hypothermia, 35°C to 32°C (95°F to 90°F); moderate hypothermia, lower than 32°C to 30°C (< 90°F to 86°F); and severe hypothermia, colder than 30°C (< 86°F). Other factors that may be useful in separating groups of patients with AH include the presence of underlying illness, altered neurologic state on arrival at the ED, hypotension, and the need for prehospital cardiopulmonary resuscitation (CPR). Shivering has been observed down to 31°C and may not cease until 30°C. Additionally, there are case reports of patients being verbally responsive at 25°C and having signs of life at temperatures less than 24°C. A hypothermia outcome score has been developed that incorporates some of these factors and may permit comparison of outcomes in patient groups treated with different modalities.

Risk factors for the development of AH include burn injuries, extremes of age, ethanol intoxication, dehydration, major psychiatric illness, trauma, use of intoxicants, significant blood loss, sleep deprivation, malnutrition, and concomitant medical illnesses. Risk factors for the development of hypothermia indoors include advanced age, coexisting medical conditions, being alone at the time of illness, being found on the floor, and abnormal perception or regulation of temperature. Unlike healthy exposed outdoor enthusiasts, such as skiers or mountaineers, hypothermia in urban populations is most often associated with conditions that impair either thermoregulation or the ability to seek shelter. In the majority of studies of urban hypothermia, death has been attributed to the severity of the underlying disease.
Because signs and symptoms may be vague and nonspecific, mild to moderate hypothermia may easily be overlooked in the ED. A common error is failure to routinely obtain an accurate core temperature in all patients at risk. The diagnosis is frequently delayed because of false reliance on standard oral temperatures. Symptoms such as confusion in the elderly and combativeness in intoxicated patients might not initially be recognized as symptoms of hypothermia. Hypothermic patients frequently will not feel cold or shiver, particularly the elderly population, who have impaired thermoregulatory responses because of their advanced age. Paradoxical undressing, a cold-induced psychiatric dysfunction, has been described in confused patients in whom a sensation of heat develops at lowered body temperatures. It occurs as a result of constricted blood vessels near the surface of the body that suddenly dilate. In many cases these patients are mislabeled as psychotic, thereby leading to further delays in appropriate treatment.

Measurement of Core Temperature

Because of the nonspecific nature of the symptoms of hypothermia, accurate assessment of temperature is a necessity when considering this diagnosis. It is of paramount importance not only for confirmation of the diagnosis but also for guidance in further diagnostic and therapeutic decisions. Any thermometer that does not record temperatures in the hypothermic range is inappropriate for evaluating significant hypothermia. Standard glass/mercury thermometers generally cannot record temperatures lower than 34°C (< 93.2°F), although some models are available that record temperatures as low as 24°C (75.2°F) (Dynamed, Inc., Carlsbad, CA). An electronic probe with accompanying calibrated thermometer is recommended when monitoring this vital sign. Examples of thermometers with accompanying accuracy at various temperature ranges are shown in Fig. 65.1.

<table>
<thead>
<tr>
<th>°F</th>
<th>0–2 years</th>
<th>3–10 years</th>
<th>11–65 years</th>
<th>&gt;65 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oral</td>
<td>—</td>
<td>95.9</td>
<td>99.5</td>
<td>97.6</td>
</tr>
<tr>
<td>Rectal</td>
<td>97.9</td>
<td>100.4</td>
<td>97.9</td>
<td>100.4</td>
</tr>
<tr>
<td>Axillary</td>
<td>94.5</td>
<td>99.1</td>
<td>96.6</td>
<td>98.2</td>
</tr>
<tr>
<td>Ear</td>
<td>97.5</td>
<td>100.4</td>
<td>97.0</td>
<td>100.2</td>
</tr>
<tr>
<td>Core</td>
<td>97.5</td>
<td>100.0</td>
<td>97.5</td>
<td>100.0</td>
</tr>
</tbody>
</table>

**Figure 65.1** Electronic thermometers provide accurate temperatures over various ranges. **A,** The Welch Allyn Model 692/690 SureTemp Plus (Skaneateles Falls, NY) (oral, axillary, rectal) is accurate from 80°F to 110°F. Note the various ranges for normal temperatures from various sites by age. It is generally accepted that a rectal temperature higher than 100.3°F represents a fever and that oral readings can be misleading. **B,** The IVAC electronic thermometer (Alaris CNA Medical, Royce City, TX) is also commonly used in the emergency department (accurate from 88°F to 108°F). For severe hypothermia or hyperthermia, it is important to know the accuracy of the thermometer being used. **C,** The graph demonstrates variations in normal body temperature by site and age. (A and B, Images courtesy WelchAllyn, Inc.)
Core temperature is traditionally estimated with a rectal probe, but due to large gradients within the body, rectal temperature often lags behind core temperature by up to 1 hour, reading higher than esophageal temperature during cooling and lower than esophageal temperature during rewarming. Esophageal probes may be used, although they may be affected by warm humidified air therapy. Other possible sites for measurement of temperature include the tympanic membrane, nasopharyngeal tract, and urinary bladder. Fresh urine temperature can closely approximate core temperature. “Deep forehead” temperatures measured with a Coretemp thermometer (Teramo, Tokyo) have also demonstrated excellent accuracy and approximation of core temperatures. For continuous monitoring purposes, rectal or bladder probes are preferred. Infrared tympanic temperatures have demonstrated excellent correlation with core temperatures. However, studies show that although easier to use and faster, infrared tympanic temperatures can be inaccurate at extremes of temperature by underestimating higher temperatures and overestimating lower temperatures. When a rectal probe is used, insert it at least 15 cm beyond the anal sphincter and verify its position frequently. One should remember that temperature gradients exist in the human body and therefore consistency of monitoring at one or more sites is mandatory. A chart and formula that convert centigrade to Fahrenheit temperatures will assist the clinician in assessing the severity of hypothermia (see Fig. 65.1C).

Pathophysiology

AH results from failure of the body’s thermoregulatory responses to generate enough heat to compensate for heat losses. These thermoregulatory responses include shivering, tachycardia, tachypnea, increased gluconeogenesis, peripheral vasoconstriction, and shunting of blood to central organs (Fig. 65.2). As core temperature drops despite these compensatory mechanisms, the patient becomes poikilothermic and cools to ambient temperature.

Four methods of heat loss affect the body: radiation, convection, and evaporation. Radiation involves transfer of heat from a warmer body to a cooler environment and accounts for approximately 60% of heat loss in a normothermic individual. Convection refers to loss of heat from direct contact with a cooler surface. These losses are most profound with immersion hypothermia. Evaporation refers to loss of heat by evaporation of water. Other possible sites for measurement of temperature include the tympanic membrane, nasopharyngeal tract, and urinary bladder. Evaporation loss, especially with a wind chill factor. Evaporation refers to loss of heat through sweating and insensible water loss.

With hypothermia, the enzymatic rate of metabolism decreases twofold to threefold with each 10°C (18°F) drop, and cerebral blood flow decreases 6% to 7% per 1°C (1.8°F) drop. Signs and symptoms of hypothermia vary according to the core temperature. The overall functioning of all organ systems is impaired by the cold. The greatest effects are seen in the cardiovascular, neurologic, and respiratory systems (Table 65.1). As core body temperature drops below 33°C (< 91.4°F), the patient becomes confused and ataxic. The initiation of involuntary motor activity (shivering) prevents the reduction in core temperature. Shivering thermogenesis in skeletal muscle operates on acute cold stress. In a malnourished patient, the mechanism may be rendered ineffective secondary to reduced muscle mass. Shivering stops at approximately 32°C (89.6°F), and shivering artifact on an electrocardiogram has been associated with increased survival of individuals with severe hypothermia. Atrial fibrillation occurs frequently as the temperature continues to drop and the patient loses consciousness. A J wave on the electrocardiogram often appears before ventricular fibrillation (Fig. 65.3). Though classically considered pathognomonic for hypothermia, the J or “Osborne” wave has no prognostic or predictive value in cases of hypothermia. Studies have found that Osborne waves are present in 36% of AH survivors and in 38% of nonsurvivors. Ventricular fibrillation may occur below 29°C (< 84.2°F) and becomes common as the core temperature drops to 25°C (77°F). The electroencephalogram flattens at 19°C to 20°C (65.2°F to 68°F). Asystole commonly develops at 18°C (64.4°F) but has been seen at higher temperatures. Initial core temperature does not necessarily correlate with patient outcome. The lowest recorded temperature in a survivor of AH is 9°C (43.7°F).

### TABLE 65.1 Signs and Symptoms of Hypothermia

<table>
<thead>
<tr>
<th>CORE TEMPERATURE (°C)</th>
<th>CARDIOVASCULAR SYSTEM</th>
<th>RESPIRATORY SYSTEM</th>
<th>CENTRAL NERVOUS SYSTEM</th>
</tr>
</thead>
<tbody>
<tr>
<td>30–34</td>
<td>Tachycardia</td>
<td>Tachypnea</td>
<td>Lethargy</td>
</tr>
<tr>
<td></td>
<td>Increased afterload</td>
<td>Increased minute ventilation</td>
<td>Mild confusion</td>
</tr>
<tr>
<td></td>
<td>Increased systemic blood pressure</td>
<td></td>
<td>Loss of fine motor coordination</td>
</tr>
<tr>
<td>30–34</td>
<td>Progressive bradycardia</td>
<td>Increased bronchial secretions</td>
<td>Delirium</td>
</tr>
<tr>
<td></td>
<td>Decreased cardiac output</td>
<td>Diminished gag reflex</td>
<td>Slowed reflexes</td>
</tr>
<tr>
<td></td>
<td>Hypotension</td>
<td>Depressed cough response</td>
<td>Muscle rigidity</td>
</tr>
<tr>
<td></td>
<td>Lengthening of cardiac conduction</td>
<td></td>
<td>Abnormal EEG findings</td>
</tr>
<tr>
<td></td>
<td>Atrial/ventricular dysrhythmias</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 30</td>
<td>Spontaneous ventricular fibrillation</td>
<td>Respiratory rate decreased to 5 breaths/min</td>
<td>Areflexia</td>
</tr>
<tr>
<td></td>
<td>Osborne waves at 25°C</td>
<td></td>
<td>Coma</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fixed pupils</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Rigidity</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>EEG silent at 19°C</td>
</tr>
</tbody>
</table>

EEG, Electroencephalogram.
Cold-induced injuries such as hypothermia and frostbite lead to a thermoregulatory response (e.g., shivering and increased sympathetic activity), cellular and tissue effects (e.g., membrane damage, electrolyte imbalance, endothelial injury, and thrombosis), and systemic effects (e.g., shock, arrhythmia, and neuromuscular dysfunction). (From Coughlin MJ, Zumwalt R, Fallico F, editors: Surgery of the foot and ankle, ed 8, St. Louis, 2006, Mosby.)
heating augments thermal comfort, increases core temperature by approximately 0.8°C/hr (33.4°F/hr), and reduces patient pain and anxiety during transport.51 In one study, resistive heating more than doubled the rewarming rate when compared with passive insulation and did not produce an afterdrop.52 With longer transport times, use active rewarming methods limited to heated inhalation and truncal heat application. Place insulated hot water bottles near the patient’s axilla or groin. The Res-Q-Air device (CF Electronics, Inc., Commack, NY) is lightweight, portable, and delivers heated humidified air or oxygen at temperatures ranging from 42°C to 44°C (107.6°F to 111.2°F) and down to ambient conditions of −20°C (−4°F).

In more remote settings, another option is to use a modified forced-air warming system in the field. The Portable Rigid Forced-Air Cover is heated with a Bair Hugger heater/blower (Augustine Medical, Inc., Eden Prairie, MN). It covers the patient’s trunk and thighs and can adapt to various transport vehicle power sources.8 Immobilize patients with potential traumatic injuries to the spine or extremities before transport. Pay continuous attention to airway maintenance. Initiate fluid resuscitation with intravenous (IV) crystalloid, preferably 5% dextrose in normal saline (D5NS). Alternatively, give warmed oral glucose-containing drinks to a patient who is awake and alert. Most hypothermic patients are dehydrated because fluid intake is reduced and cold causes diuresis. Avoid using lactated Ringer’s solution because it can theoretically decrease the metabolism of lactate by cold-induced hepatic dysfunction. If possible, use warmed

### Initial Evaluation and Stabilization of Hypothermic Patients

Several guidelines for the treatment of AH exist. The State of Alaska Cold Injuries Guidelines were last updated in 2014 and include a section on AH.47 Additionally, in 2014, the Wilderness Medical Society developed practice guidelines for out-of-hospital care of AH.48 Treatment of hypothermia can be divided intoprehospital care and ED management.

#### Prehospital Care

In the prehospital setting, focus primarily on removing the patient from the current environment to prevent further decreases in core temperature. Studies have shown that oral temperatures are sufficiently accurate for field use; however, infrared tympanic thermometers may not be reliable in the prehospital setting. Handle these patients with special care and anticipate the presence of an irritable myocardium because aggressive measures can inadvertently trigger cardiac dysrhythmias. Hypovolemia and a large temperature gradient often exist between the periphery and the core in a hypothermic patient.8 Avoid aggressive field management and prolonged transport times.49,50 After removing the patient’s wet clothing, wrap the patient in dry blankets or sleeping bags. Field rewarming is a misnomer because adding significant heat to a hypothermic patient in the field is extremely difficult. Studies have shown that for mild hypothermia, resistive heating (e.g., warming blankets) can be used safely in the prehospital setting. Resistive

---

**Figure 65.3** In severe hypothermia, the electrocardiogram (ECG) exhibits marked elevation of the J deflection, so-called Osborne waves (arrowheads). The height of the J wave is proportional to the degree of hypothermia, and this finding is usually most marked in the midprecordial leads. This ECG is from a patient with sinus bradycardia, but in approximately half the patients with a temperature below 32°C (89.6°F), slow atrial (arrow) fibrillation develops, a rhythm that usually converts spontaneously with rewarming. (Adapted from Krantz MJ, Lowery CM: Giant Osborne waves in hypothermia, *N Engl J Med* 352:184, 2005. Used with permission.)
IV fluids because they are generally well tolerated. If available, use a flameless heater, which is currently being used by military medical units and provides an easy and expedient means of warming fluids in the prehospital setting.

Intubate unresponsive patients, but recognize that there is no universal agreement on when to intubate a hypothermic patient who has detectable vital signs. Pulse oximetry is not usually helpful because vasoconstriction limits blood flow to the periphery and readings may be inaccurate or not possible. Prehospital providers should assess for pulses and initiate CPR at a standard rate of 100 compressions per minute if no pulses are palpable after 1 minute. Delayed or intermittent CPR can be associated with good outcomes with neurologically intact survivors after prolonged chest compressions, but should be used only if evacuation efforts necessitate. Some authors suggest that pulseless victims with core temperatures below 32°C (<89.6°F) should be transported with continuous CPR. Other authors believe that it is unnecessary to perform CPR on a patient who has any perfusing cardiac rhythm because it may precipitate ventricular fibrillation. There is no universally accepted standard for intubation or CPR in hypothermic patients with detectable vital signs.

Definitive prehospital determination of cardiac activity requires a cardiac monitor. Cardiac arrest is a common misdiagnosis because peripheral pulses are difficult to palpate when extreme bradycardia is present along with peripheral vasoconstriction. Some authors report that asystole is a more common rhythm than ventricular fibrillation. In the field, differentiating between ventricular fibrillation and asystole may be impractical. Successful defibrillation has been reported at 20°C but attempted defibrillation is often unsuccessful until the core temperature is above 30°C. If the defibrillation attempt is unsuccessful, initiate active rewarming while continuing CPR defibrillation. These attempts can be given occasionally during the rewarming process.

Transport cold, stiff, cyanotic patients with fixed and dilated pupils because the treatment dictum for prehospital personnel remains, “No one is dead until warm and dead.” Some patients are actually cold and dead. It is useful and humane if they can be safely identified. Because human physiologic responses are variable, it is difficult to predict outcome. A succinct summary of the prehospital care of a hypothermic patient is rescue, examine, insulate, and transport. In the majority of cases, evacuation, resuscitation, and transport should be initiated, provided that doing so does not endanger prehospital responders.

**ED Management**

There are no universally established standards of care regarding the use of specific techniques for rewarming a hypothermic patient or cooling a hyperthermic patient. This chapter describes all potentially useful modalities. Many are not applicable for general use in the ED, whereas others are safe, beneficial, and easily accomplished in the general ED setting. Some invasive procedures, however, such as cardiopulmonary bypass and irrigation of the peritoneal or thoracic cavity, may be overly aggressive or of anecdotal or theoretical benefit only. Exactly when to institute any given intervention is best determined by the resources available, the initial scenario, and clinical judgement individualized for each patient.

Treatment priorities in the ED setting are to prevent further decreases in core body temperature; establish a steady, safe rewarming rate; maintain stability of the cardiopulmonary system; and provide sufficient physiologic support. Adjust the rate of rewarming and the techniques used according to the degree of hypothermia and the severity of the patient’s clinical condition (Table 65.2). Anticipate and prevent complications.

In a pulseless, apneic patient, initiate CPR and continue until the core temperature is above 34°C (>93.2°F). Profound hypothermia results in coma, hyporeflexia, fixed and dilated pupils, severe bradycardia, and often an unobtainable blood pressure. With severe hypothermia, a pulse might not be palpable and measurement of blood pressure might require the use of a Doppler device. If available, use ultrasound to detect the presence of cardiac wall motion. Follow the heart rate and rhythm with electrocardiographic monitoring. In patients who have anything more than minimal impairment, perform arterial blood gas analysis frequently to determine oxygenation, ventilation, and acid-base status. If feasible, establish large-bore IV lines. Avoid central lines if possible because insertion of such lines may exacerbate the myocardial irritation. Give maintenance IV fluids. Warm all IV fluids to 40°C to 42°C (104°F to 107.6°F), but be aware that the usual volumes administered will not contribute significant heat calories. With long standard IV tubing, the heated IV fluids may actually cool to room temperature before entering the patient’s IV site.

With a mild to moderate reduction in core temperature, the level of mentation correlates with the severity of the AH, associated illnesses, or both. Noteworthy exceptions are alcoholics and diabetics, who can be in a coma at higher core temperatures because of concomitant hypoglycemia. Perform bedside glucose measurements on patients when they arrive.

**TABLE 65.2 Rewarming Techniques**

<table>
<thead>
<tr>
<th>CORE TEMPERATURE (°C)</th>
<th>METHOD AND TECHNIQUES</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt; 32</td>
<td>PER</td>
</tr>
<tr>
<td></td>
<td>Dry blankets, clothing</td>
</tr>
<tr>
<td></td>
<td>Heated intravenous solutions (43°C), D5NS</td>
</tr>
<tr>
<td></td>
<td>Warm fluids if fully alert</td>
</tr>
<tr>
<td>≤ 32</td>
<td>AER</td>
</tr>
<tr>
<td></td>
<td>Heated blankets, heating pads, warm air convection</td>
</tr>
<tr>
<td></td>
<td>Radiant heat sources</td>
</tr>
<tr>
<td></td>
<td>Alcohol-circulating blankets</td>
</tr>
<tr>
<td></td>
<td>ACR</td>
</tr>
<tr>
<td></td>
<td>Peritoneal dialysis</td>
</tr>
<tr>
<td></td>
<td>Bladder, gastric, or colonic lavage with warm fluids (43°C)</td>
</tr>
<tr>
<td></td>
<td>Heated intravenous fluids</td>
</tr>
<tr>
<td></td>
<td>Heated humidified oxygen</td>
</tr>
<tr>
<td></td>
<td>Thoracic cavity lavage (43°C)</td>
</tr>
<tr>
<td></td>
<td>Extracorporeal blood rewarming</td>
</tr>
<tr>
<td></td>
<td>Hemodialysis</td>
</tr>
<tr>
<td></td>
<td>Ultrasonic and low-frequency microwave diathermy</td>
</tr>
<tr>
<td></td>
<td>Arteriovenous anastomoses rewarming</td>
</tr>
</tbody>
</table>

**ACR.** Active core rewarming; **AER.** active external rewarming; **PER.** passive external rewarming.
in the ED. A high correlation exists between alcohol consumption and the development of hypothermia, especially in colder climates. A review of 68 cases of hypothermic deaths in Jefferson County, Alabama, found that a significant number of cases involved middle-aged men who had consumed alcohol. In other urban cases of AH reviewed, the majority were alcoholics. This study noted glycosuria in patients, even when low serum glucose values were evident, and described a renal tubular glycosuria in patients with AH. Such glycosuria may worsen or cause hypoglycemia. Glycosuria in AH is no guarantee of an adequate serum glucose concentration. This supports the routine use of supplemental IV glucose unless a normal serum glucose value can be quickly ensured. Consider administering IV thiamine (100 mg) and a trial dose of 0.4 to 2 mg of IV naloxone (Narcan, Adapt Pharma, Inc. Radnor, PA) in a comatose patient. Although failure to rewarm spontaneously has been noted in victims with hypothyroidism and other endocrine deficiencies, reserve the use of thyroid hormones and corticosteroids for patients with suspected thyroid and adrenal insufficiency, respectively.

The thermoregulatory vasoconstriction caused by hypothermia significantly decreases subcutaneous oxygen tension. Good correlation exists between the incidence of wound infection and subcutaneous oxygen tension. As core temperatures decrease from 41°C to 26°C (105.8°F to 78.8°F), neutrophil function is significantly impaired. In animal models, hypothermia appears to decrease leukocyte sequestration within the brain parenchyma, thus offering some resistance to meningitis. Although antibiotics are not routinely indicated for each case. Standard rewarming rates are a 0.5°C/hr to 2.0°C/hr (0.9°F/hr to 3.6°F/hr) rise in temperature in an otherwise stable patient (Table 65.3). Carefully consider and individualize invasive therapy to the severity of the hypothermia and the condition of the patient. Avoid overtreating and oversudging invasive techniques in an otherwise stable hypothermic patient. In patients with severe underlying problems such as hypoglycemia, hyperglycemia, sepsis, adrenal crisis, drug overdose, or hypothyroidism, treat these conditions appropriately in addition to treating the hypothermia. Long-term outcome may depend more on treatment of the underlying illness than on treating the hypothermia.

### Active External Rewarming

The cornerstone of the effectiveness of passive external rewarming relies on the body's ability to restore normal body temperature through its own mechanisms for heat production. Stop further heat loss with insulation and manipulation of the environment. Give warm fluids containing glucose to patients who are fully alert. For patients with mild AH, remove wet clothing and then provide passive external rewarming with blankets. The technique is simple, but the patient must be capable of generating enough body heat for this method to be successful. Give warmed IV fluids to counteract the cold-induced diuresis. Internal heat generation is required for rewarming, and this effect will be relatively slow. In an otherwise stable patient, aggressive intervention with drugs and invasive monitoring might be more harmful than beneficial. Patients who cannot shiver, those who are hypotensive, or those who are intoxicated or malnourished may not have this capability. Survival rates with passive external rewarming have ranged from 55% to 100%. For patients in the moderate or severe category of hypothermia, a more aggressive approach may be warranted. The options available are active external rewarming and active core rewarming. Active core rewarming techniques can be further divided into less invasive and more invasive techniques. The aggressiveness of therapy depends more on the patient's underlying health, hemodynamic status, and response to initial therapy than on the initial temperature.

### Active External Rewarming

The application of heat to the skin of a hypothermic patient has been termed active external rewarming.

#### Indications

Although there is some suggestion that active external rewarming of profoundly hypothermic patients by immersion may be associated with an increase in mortality over other treatments, other studies suggest that this technique is highly effective for mild hypothermia. Use it selectively and limit it to the
trunk. Other forms of active external rewarming are increasingly being used in the ED as adjunctive care of moderately hypothermic, otherwise healthy individuals. Vasodilatation limits the ability to increase core temperature with techniques that primarily warm the skin.

Active external rewarming is most beneficial when the heat supplied by the external source is greater than the loss of rewarming heat incurred by the cessation of shivering. In more remote wilderness settings where more aggressive warming techniques are precluded because of the lack of equipment or personnel, active external rewarming with body-to-body contact may be the only option. The rewarming contribution of body-to-body contact appears to be limited, however.

**Equipment**

Traditionally, immersion therapy has used a heated (40°C to 42°C [104°F to 107.6°F]) water tank of the type present in most burn units. Generally, immerse a hypothermic patient entirely except for the extremities and head, but immersion of the extremities may hasten rewarming. A major drawback is the inability to closely monitor patients undergoing immersion. Alternatively, use a warm water–filled heat exchange blanket (e.g., Blanketrol, Cincinnati Sub-Zero Products, Cincinnati, OH) for conduction warming. Intraoperative studies have demonstrated excellent results. A forced warm air convection system (Bair Hugger, Augustine Medical, Eden Prairie, MN; Snuggle Warm Convective Warming System, Sims Level 1, Inc., Rockland, MA) has been used for postsurgical rewarming. This approach has also been used successfully for ED-based AH therapy (Fig. 65.4). Rewarming by warm air convection permits continued monitoring in the ED and is better tolerated than immersion because of the less rapid development of vasodilatation in peripheral tissues. As an alternative for noninvasive active external rewarming, there are case reports describing rewarming hypothermic patients with a device called Arctic Sun (Medivance, Inc., Artic Sun, Louisville, CO), a system also used to induce therapeutic hypothermia. The system circulates warm water through gel pads placed on areas of the body with large surface areas (e.g., the trunk and thighs). Case reports exist showing rewarming rates from 0.63°C to 2.5°C per hour, with one patient rewarmed from an initial temperature of 22°C.

**Technique**

Because profound fluid shifts can occur with conduction warming, give the patient supplemental IV fluid warmed to 40°C (104.0°F; Hotline Fluid Warmer, Sims Level 1, Inc., Rockland, MA) at a rate sufficient to generate a urinary output of 0.5 to 1.0 mL/kg per hr. Give an initial fluid bolus of 500 mL of D,NS. Note that blood pressure is not an accurate means of gauging fluid resuscitation because serious hypothermia is always accompanied by “physiologic” hypotension. Because patients requiring mechanical ventilation have rarely been subjected to tank immersion, it cannot be recommended for hypothermic patients who require intubation. Rewarming rates ranging from 0.9°C to 8.8°C (1.6°F to 15.8°F) per hour have been reported with immersion therapy.

A heat exchange blanket allows the patient to receive other treatments that may be difficult or impossible to carry out in a tub, such as defibrillation, CPR, or more invasive warming techniques. Place the heating blanket and overlying cloth sheet underneath the patient. Set the blanket temperature to 40°C to 42°C (104.0°F to 107.6°F), and initiate the measures described in the section on Passive Rewarming Techniques. Forced-air rewarming (convection) uses a blanket cradle to create an environment through which heated air is blown. Access to the patient is quite good with this system because the overlying blankets can be raised temporarily to evaluate the patient or perform procedures. Experience with mild immersion-induced hypothermia in volunteers suggests that the forced-air technique warms at a rate comparable to that of vigorous shivering, but with less metabolic stress and less afterdrop.

**Arteriovenous Anastomoses Rewarming**

Arteriovenous anastomoses rewarming (AVR) involves immersion of the distal end of the extremities (hands, forearms, feet, and lower part of the legs). Advantages include rapid rewarming rates. A study of AVR immersion at temperatures of 45°C and 42°C (113.0°F and 107.6°F) in healthy volunteers demonstrated rewarming rates of 9.9°C/hr (±3.2°C/hr) for the former and 6.1°C/hr (±1.2°C/hr) (43.0°F ± 34.2°F/hr) for the latter. There was also a decrease in postcooling afterdrop. AVR is well tolerated by patients because of the rapid rise in core temperature and the shortened period of shivering.

**Complications**

There is concern that surface warming with accompanying vasodilatation may produce relative hypovolemia in a hypothermic patient. Other complications described with the active external rewarming method include core temperature afterdrop and rewarming acidosis. In core temperature afterdrop, colder peripheral blood is transported to the warmer core organs, thereby further reducing core temperature. In rewarming acidosis, colder blood and lactic acid return to the core organs and worsen the acidosis. To limit these complications in patients with moderate hypothermia, some authors advocate using active external warming only after active internal techniques have been initiated. Others suggest that core afterdrop is inevitable regardless of the rewarming method, as temperature will temporarily decrease in any object with a warm core and cool periphery, due to conductive properties.
CPR and other advanced cardiac therapy and monitoring are impossible with immersion rewarming. Until studied further, active external rewarming should be considered only in a clinically monitored setting for mildly hypothermic patients who can protect their airways. When using a heating device, also monitor the potential for burns in areas that have the greatest contact with the heating source.

**Active Core Rewarming**

There is evidence that active core rewarming may decrease mortality from severe hypothermic exposure when compared with other techniques. In the face of circulatory failure, often the best chance of survival is treatment with extracorporeal circulation (ECC) and warming of the blood. Several methods have been described, including the use of warm humidified air through an endotracheal (ET) tube or mask, peritoneal lavage, gastric or bladder lavage with warm fluid, thoracic tube lavage, cardiopulmonary bypass, AVR, peripheral vascular extracorporeal warming, hemodialysis, and thoracotomy with mediastinal lavage. These techniques transfer heat actively to the body core and achieve varying rewarming rates. The specific techniques and some of the advantages and disadvantages for each procedure follow.

**Emergency Warming of Saline in a Microwave**

Under ideal circumstances, keep saline in a standard warming device. When large amounts of saline are required for such procedures as peritoneal lavage, warm 1-L saline bags rapidly in a standard microwave oven (Fig. 65.5). Although devices vary, a 650 W microwave oven has been demonstrated to warm 1 L of room-temperature non–dextrose-containing saline from 21.1°C to 38.3°C (70°F to 101°F) in 120 seconds on the high setting. At midcycle (i.e., after 60 seconds), interrupt the heating with agitation, and repeat the agitation at the end of the cycle before infusion. Fluids containing dextrose should not be warmed in this manner, as glucose caramelizes at 60°C. Fluids in glass bottles and blood products are also not safe to be warmed with this method.

**Inhalation of Heated Humidified Oxygen or Air**

The use of warm humidified oxygen to treat hypothermia has been well established. Average rates of rewarming of 1°C/hr (33.8°F/hr) via mask and 1.5°C/hr to 2.0°C/hr (34.7°F/hr to 35.6°F/hr) via ET tube with heated aerosol at 40°C (104.0°F) can be obtained. Faster rewarming rates may be accomplished with a maximum safe aerosol temperature of 45°C (113°F). Core rewarming with this technique occurs through the following mechanisms. The warmed alveolar blood returns to the heart and warms the myocardium. The heated, humidified air delivered to the alveoli also warms contiguous structures in the mediastinum by conduction. Warming the inhaled air or oxygen eliminates a major source of heat loss.

**Indications and Contraindications.** The use of heated humidified air or oxygen is a simple technique that should be used routinely in all patients with hypothermia, regardless of severity. If the correct equipment is available, it can be used in the field and in the hospital. One must address the risk for burns during the inhalation of warm air in the field environment. Mouth-to-tube ventilation in an intubated hypothermic prehospital patient has the theoretical advantage of providing warm humidified air without special equipment. A ventilating rescuer can inhale oxygen and then expire it into the patient's ET tube to provide air with increased oxygen content. There are no contraindications to or reported complications from the use of warm humidified air for hypothermia, and there is no afterdrop.

**Technique.** Use a heated cascade nebulizer with a mask for patients with spontaneous respirations. Use a volume ventilator for intubated patients. Monitor the inspired air to maintain a temperature of approximately 45°C (113.0°F). Temperatures higher than 50°C (122°F) may burn the mucosa, and temperatures lower than 45°C (<113°F) do not deliver the maximum heat. Humidify the air or oxygen and note that the heater module may need modification because many units have feedback mechanisms that shut off at a given temperature. It may be difficult to deliver oxygen at the recommended temperature because of equipment limitations. In many cases the air temperature is only 30°C (86°F).

**Summary.** Inhalation of warm humidified air or oxygen results in gradual rewarming of the core and should be the
mainstay of all rewarming therapy. Studies have suggested that the rewarming rate of inhalation therapy is inferior to that of peritoneal lavage, thoracic lavage, and bath rewarming. Inhalation therapy can be combined with any and all other methods of rewarming and is relatively noninvasive and inexpensive. This therapy should be considered as the initial treatment of choice for hypothermic patients.

**Peritoneal Dialysis (Lavage)**

Peritoneal dialysis (lavage) is an attractive treatment for severe hypothermia because it is available in most hospitals and does not require any unusual equipment or training. Rewarming rates of 2°C to 3°C (3.6°F to 5.4°F) per hour, depending on the dialysis rate, can be achieved without sophisticated equipment that may delay therapy or require transfer of the patient to a tertiary care facility. This technique can also be used to help correct electrolyte imbalances. Peritoneal lavage is useful primarily in severe cases in combination with other rewarming techniques for patients without spontaneous perfusion, but has also been used alone in patients undergoing CPR in whom ECC was felt to be contraindicated or not available.

Rewarming by peritoneal dialysis was first used successfully in a patient in ventricular fibrillation with a temperature of 21°C (69.8°F). Since that time, there have been reports of successful rewarming with peritoneal lavage in stable, severely hypothermic patients and unstable hypothermic patients in cardiac arrest. Peritoneal lavage works via transfer of heat from lavage fluid to the peritoneal cavity. The peritoneal great vessels and abdominal organs provide a large surface area for exchange of heat. The use of warmed peritoneal lavage fluid is an effective approach to rewarming. There have been reports in the literature of success with rapid high-volume peritoneal lavage in pediatric patients. The technique involves the use of an infraumbilical "mini-laparotomy" incision followed by placement of a large silicone peritoneal dialysis catheter. The catheter is connected to a rapid infusion device with delivery of 1 L of warmed normal saline every 90 seconds.

**Indications and Contraindications.** Peritoneal dialysis is appropriate therapy in a severely hypothermic patient. In practice, it is often omitted if other measures appear to be successful. There are no universally established criteria for performing peritoneal lavage in hypothermic patients who have detectable vital signs. Though theoretically less effective than other techniques that directly warm the thorax in the setting of cardiac arrest, it has been used successfully in that situation. It is theoretically useful in hypothermic patients who have overdosed with a dialyzable toxin. Other less invasive methods, such as gastric or bladder lavage or warm nebulized air or oxygen inhalation, may be preferred in stable patients with temperatures higher than 26°C to 28°C (>78.7°F to 82.4°F). Peritoneal dialysis should not be performed on patients with previous abdominal surgery. It should be used with extreme caution in patients with a coagulopathy.

**Equipment.** We recommend using the Seldinger technique with a commercially available disposable kit (e.g., Arrow Peritoneal Lavage Kit, product no. AK-09000, Arrow International, Inc., Reading, PA) because of the ease of performance and minimal morbidity associated with this procedure.

**Technique.** In a noncritical patient, obtain a coagulation profile before the procedure, but in life-threatening situations, initiate the procedure immediately before laboratory studies. Place the patient in the supine position with a Foley catheter and nasogastric tube in place. After infiltrating with lidocaine, make an infraumbilical stab incision with a No. 11 scalpel blade, and place an 18-gauge needle into the peritoneal cavity directed toward the pelvis at a 45-degree angle. Insert a standard flexible J wire through the needle, and then remove the needle. Pass the 8-Fr dialysis catheter over the wire with a twisting motion, and then remove the wire.

Lavage rates of 4 to 12 L/hr can be achieved with two catheters. Warm the fluid with a standard blood warmer to 40°C to 45°C (104.0°F to 113.0°F). Use a standard 1.5% dextrose dialysate solution. Add potassium (4 mmol/L) if the patient becomes hypokalemic. Saline has also been used successfully. The rate should be at least 6 L/hr and preferably 10 L/hr.

**Complications.** The Seldinger method has a complication rate of less than 1%. A “mini-lap” performed via direct dissection may also be used but might have a higher complication rate. Further discussion of potential complications is provided in Chapter 43.

**Summary.** Peritoneal dialysis is a useful method because it entails readily available fluid and can be done with a self-contained disposable kit. If a hospital also treats trauma victims, the lavage kit can be the same as that used for evaluation of abdominal trauma. If this technique is combined with warm nebulized inhalation, warming rates of 4°C/hr (7.2°F/hr) can be achieved. Peritoneal lavage rewarms the liver and restores its synthetic and metabolic properties.

**Gastrointestinal and Bladder Rewarming**

Gastric or bladder irrigation offers some of the same advantages as peritoneal dialysis without invading the peritoneal cavity. Heat is delivered to structures in close proximity to the core. In the Multicenter Hypothermia Study, gastric/bladder/colon lavage had a first-hour rewarming rate of 1.0°C to 1.5°C/hr (33.8°F/hr to 34.7°F/hr) and a second-hour rewarming rate of 1.5°C/hr to 2.0°C/hr (34.7°F/hr to 35.6°F/hr) for severe hypothermia. In a multifactorial analysis of the Multicenter Hypothermia Study there was a trend toward improved survival in patients treated in this manner.

Although the amount of heat delivered with gastric lavage appears to be less than that delivered with peritoneal dialysis, it is somewhat easier to use and less invasive. When combined with other methods, gastric or bladder lavage provides significant warming. Serum electrolyte levels should be monitored if large volumes of tap water are used because dilutional electrolyte disturbances may occur. Children and geriatric patients might be more susceptible to electrolyte changes with tap water irrigation.

**Indications and Contraindications.** Warmed gastric or bladder lavage may be used as adjunctive therapy for moderate or severe hypothermia. It can be combined with other warming techniques when rapid rewarming is needed. Patients who are obtunded and lack protective airway reflexes should undergo ET intubation before gastric lavage to prevent aspiration of gastric contents. Refer to the appropriate chapters concerning nasogastric tube placement (see Chapter 40), gastric lavage (see Chapter 42), and urethral catheterization for specific contraindications to these procedures.
Equipment. Use a large-diameter 32- to 40-Fr lavage tube with normal saline solution warmed to 40°C to 45°C (104.0°F to 113.0°F) in a microwave or blood warmer with verification of temperature before use. Although smaller tubes are easily passed nasally, use oral placement of the large lavage tubes. A modified Sengstaken tube with gastric and esophageal balloons may also be used.

Technique. Instill 200- to 300-mL aliquots of fluid into the stomach before removal by gravity drainage. For bladder irrigation the optimal volume is not known, but avoid distention of the bladder (100- to 200-mL aliquots should be sufficient). The amount of time that the irrigant should be left in place before removal is not known, but use of rapid exchanges with a dwell time of 1 to 2 minutes is suggested.

Complications. Complications of lavage include trauma to the nasal turbinates, gastric and esophageal perforation, dilutional hyponatremia, inadvertent placement of the tube in the lungs, and pulmonary aspiration, all of which can be minimized by careful, proper technique. Fluid overload and electrolyte disturbances when using tap water are potential complications in pediatric and geriatric patients.

Summary. Gastrointestinal and bladder lavage with heated fluids is easily performed with equipment and solutions available in any hospital. The stomach, colon, and bladder are poor sites for body cavity lavage as a result of the small surface area for heat exchange.79 Because of its ease and availability, it can be started early in the resuscitation and be combined with any other rewarming method to significantly add heat,72 although its specific effect on morbidity and mortality is not known.

Thoracic Cavity Lavage
Thoracic cavity lavage can be performed either by closed means, through chest tubes placed in one hemithorax, or in open fashion, after resuscitative thoracotomy.84 The former approach offers the advantages of being less invasive and is an effective form of treatment in hospitals not equipped for cardiopulmonary bypass. Furthermore, closed-chest CPR can be continued while this technique is used. The open thorax approach offers the theoretical advantage of direct warming of the heart and the option of open-chest cardiac massage. Rapid warming rates of 6°C to 7°C (42.8°F to 44.6°F) in 20 minutes have been described. Pleural irrigation results in cardiac rewarming and might be the method of choice, particularly in patients with an arrhythmia.82

Indications and Contraindications. Thoracic cavity lavage should be considered for patients requiring rapid core rewarming in the setting of cardiac arrest or inadequate perfusion (e.g., shock, during CPR) when cardiac bypass is not available.85 Open thoracic lavage should be considered in patients who will receive open-chest massage or thoracotomy for other reasons (e.g., hypothermic arrest with penetrating trauma). Thoracic lavage is not necessary for patients with mild or moderate hypothermia who can be warmed by other less invasive methods. Avoid the technique in patients with a coagulopathy unless required as a lifesaving measure.

Closed Thoracic Lavage. An alternative that is more practical in the ED is pleural rewarming by repeatedly using warmed saline placed intermittently and then withdrawn through a chest tube. Place two large-bore thoracostomy tubes (e.g., 36- to 38-Fr in 70-kg adults) in one hemithorax. Infuse one chest tube with 3-L bags of heated normal saline (40°C to 41°C [104°F to 105.8°F]) via a high-flow fluid infuser (e.g., Level-1 Fluid Warmer, Technologies, Inc., Marshfield, MA). Collect the effluent with an autotransfusion thoracostomy drainage set (e.g., Pleur-evac, Deknatel A-5000-ATS, Fall River, MA). Empty the removable reservoir as needed. Alternatively, use a single-chest tube system with a Y-connector arrangement similar to that used for gastric lavage. Place aliquots of 200 to 300 mL with a 2-minute dwell time followed by suction drainage (at 20 cm H₂O).

Provide closed-chest massage until adequate spontaneous perfusion occurs. Perform closed-chest defibrillation if the patient is warmed to 30°C (86°F) and has persistent ventricular fibrillation. Continue thoracic lavage until the patient's temperature approaches 35°C (95°F).

Open Thoracic Lavage. Perform a left thoracotomy and pour saline warmed to 40°C to 41°C (104.0°F to 105.8°F) continuously into the thoracic cavity to bathe the heart while an assistant suction the excess fluid from the lateral edge of the thoracotomy. Alternatively, add fluid to the thorax and mediastinum intermittently and suction after several minutes. Follow this with more warmed saline and repeat. This technique also allows direct monitoring of myocardial temperature. Perform direct cardiac massage until adequate spontaneous perfusion occurs. Perform direct cardiac defibrillation in a patient warmed to 30°C (86°F) with persistent ventricular fibrillation. When defibrillation is successful, continue direct myocardial warming until the patient's temperature approaches 35°C (95°F). If defibrillation is unsuccessful at a core temperature of 30°C (86°F), continue warming while oxygenation, perfusion, and other physiologic parameters are optimized before further attempts at defibrillation.

Summary. Thoracic lavage is an effective form of active core rewarming that is usually reserved for hypothermic arrest patients.84 Thoracic lavage may be considered when vital signs are inadequate or unstable enough to severely limit perfusion. Precise indications have not been clarified beyond patients in cardiac arrest.

Cardiac Bypass
The use of cardiac bypass or an extracorporeal shunt through either the femoral artery–femoral vein or the aortocaval procedure can result in rapid rewarming but requires surgical expertise, the availability of appropriate equipment, and technical support.766 ECC usually refers to cardiopulmonary bypass or extracorporeal membrane oxygenation (ECMO).84 ECMO appears to reduce the risk of intractable cardiorespiratory failure or severe pulmonary edema after rewarming.84 This procedure has not been compared with other rewarming methods in a controlled fashion, and few centers have this modality available in a time frame that would affect survival rates. Its main advantages appear to be the rapid rate of warming that it produces and optimal patient oxygenation and perfusion. Femoral flow rates of 2 to 3 L/min with the warmer set at 38°C to 40°C (100.4°F to 104°F) will raise the core temperature 1°C to 2°C (33.8°F to 35.6°F) every 3 to 5 minutes.7 Drawbacks include potential delays in assembling the appropriate team and equipment, delays because of the time necessary to complete...
the operation, complications from the operation, the expense of the procedure and bypass equipment, and the potential for infection. Its use in extreme situations that may include cardiac arrest should be based on individual characteristics of the patient, clinician team, and hospital resources. If readily available, it should be strongly considered in hypothermic patients with asystole or ventricular fibrillation. If oxygenation is not a consideration, venovenous rewarming with an extracorporeal venovenous rewamer can achieve rapid rewarming rates (2°C/hr to 3°C/hr [3.6°F/hr to 5.4°F/hr]), although they are slower than rates with cardiopulmonary bypass. Such a device is relatively easy to use, involves readily available technology, and probably does not require heparin. This equipment needs to be assembled before patients with hypothermia arrive. In severely hypothermic patients, extracorporeal rewarming using venovenous hemofiltration has also been reported to be successful. When compared with adults, children, especially smaller ones, require special consideration with regard to IV cannulation because drainage can be inadequate with femoral-femoral cannulation. In smaller hypothermic children, some sources recommend a more aggressive emergency median sternotomy for cardiopulmonary bypass.

Cardiopulmonary bypass is indicated in the following situations: (1) cardiac arrest or hemodynamic instability with a temperature lower than 32°C (<89.6°F), (2) no response to less invasive techniques, (3) completely frozen extremities, or (4) rhabdomyolysis with severe hyperkalemia. A 47% long-term survival rate was obtained in a Swiss study of 32 young, otherwise healthy individuals, including mountain climbers, hikers, and victims of suicide attempts. Cardiopulmonary bypass is unlikely to confer similar benefit in older, poorly conditioned populations with underlying chronic diseases.

Hemodialysis
Hemodialysis was first described for the management of AH in 1965. It is a rapid and efficient modality for rapid internal rewarming of patients with moderate to severe AH, but it is uncommonly used in clinical practice. Temporary dialysis catheters are readily available and relatively easy to place. One study reported that 26 patients with AH combined with circulatory arrest or severe circulatory failure were rewarmed to normothermia with the use of ECC. Core rewarming by hemodialysis has been achieved after placement of a dialysis catheter or with the use of an existing shunt. Some of the potential advantages and drawbacks of cardiac bypass also apply to this procedure, although slower warming rates have been reported. A range from 0.6°C/hr (33.1°F/hr) to rates as high as 4.5°C/hr (40.1°F/hr) have been achieved with fluid warmed to 40°C (104.0°F). For patients who have ingested a dialyzable toxin (such as barbiturates and toxic alcohols), hemodialysis can be used to both remove the toxin and rewarm the blood. In such cases its use may be appropriate.

Experimental Techniques
Ultrasound, radioave, and low-frequency microwave diathermy rewarming appears to be a rapid, safe, noninvasive technique that has shown promise in animal studies. Frequencies of 13.6 to 40.7 MHz are typically used. In a volunteer study the technique seemed to be less effective than immersion therapy and equivalent to passive rewarming techniques. Total liquid ventilation with warmed oxygenated perfluorcarbon is currently being studied in animals as a method of rapidly rewarming the core. Benefits include shorter rewarming times than with warm humidified oxygen (1.98 ± 0.5 hours vs. 8.61 ± 1.6 hours; P < 0.0001), no afterdrop phenomenon, and no increase in lactate dehydrogenase and aspartate transaminase.

Very hot IV fluids (65°C [149°F]) have been used in animals with little vascular damage or hemolysis. Trials in humans undergoing burn débridement have been successful in preventing hypothermia during operative procedures. Saline heated to 60°C (140°F) with modified fluid warmers was infused through central venous access. There was no evidence of intravascular hemolysis or coagulopathy after the infusions. The role of hot IV fluids in the management of AH is currently undefined.

Special Situations
Cardiac Arrest
Cardiac arrest secondary to AH requires immediate treatment for the best chance of a successful outcome. Rapid rewarming and restoration of cardiac rhythm are essential for patients in cardiopulmonary arrest and can best be achieved with a combination of passive and multiple active core rewarming techniques. Because of numerous cases of survival from hypothermic cardiac arrest with prolonged external cardiac compression, thoracotomy is not mandatory. Thoracotomy does offer some theoretical advantages, however, such as increased cardiac output with open-chest massage, direct observation of cardiac activity, and direct warming of cardiac tissue with thoracic cavity lavage of warm fluid. Cardiopulmonary bypass is an effective technique for rapid rewarming. Blunt trauma and head trauma victims were previously not ideal candidates for cardiac bypass because of the anticoagulation requirement, but some authors have advocated this technique with heparin-bonded tubing even in the setting of known traumatic injury. A review of outcomes after hypothermic cardiac arrest from one institution found that the average time from thoracotomy to the development of a perfusing rhythm was 38 minutes (range, 10 to 90 minutes). The optimal rate of cardiac compressions in hypothermic patients is not known. Because of decreased oxygen consumption by vital organs, the rate required in hypothermic cardiac arrest is less than that recommended for normothermic cardiac arrest. Cardiac compressions should be initiated at half the normal rate in profoundly hypothermic patients. Guidelines developed by the American Heart Association and the Wilderness Medical Society recommend that CPR be initiated in patients with AH unless any of the following conditions exist: a “do-not-resuscitate” status is documented and verified, obvious lethal injuries are present, chest wall depression is impossible, no signs of life are present, or rescuers are endangered by delays in evacuation and altered triage conditions.

The duration of CPR depends on the time required to raise the core temperature to a level at which defibrillation should be successful (i.e., >30°C [>86°F]). Previously, it was recommended that patients not receive a set of three countershocks until a core temperature above 30°C (>86°F) could be attained. There have been reports of successful defibrillation in patients with profound hypothermia with core temperatures of 25.6°C (78.1°F). The decision to terminate resuscitative efforts remains a clinical one, but there are certain poor prognostic factors. Certainly, survival is unlikely in patients who persist in asystole or go from ventricular fibrillation to asystole as they are warmed past 32°C (>89.6°F). Some authors describe prognostic markers for patients with severe hypothermia and
cardiac arrest which have been proposed as contraindications to ED thoracotomy and cardiac bypass. Such markers include potassium levels elevated to above 10 mmol/L (mEq/L) and pH levels below 6.5. Nonetheless, there are reports of survival in patients with higher potassium levels and a pH as low as 6.29. Presenting rhythm of asystole, advanced age, underlying illness (including infection) and elevated serum lactate may also be poor prognostic indicators. Multiple reviews have shown that asphyxia prior to hypothermia arrest (as seen in avalanche and drowning victims) is a poor prognostic factor. Ultimately, the decision to continue resuscitative efforts should not be based solely on specific laboratory values or the initial core temperature.

Isolated reports of survival of hypothermic patients with prolonged CPR make extended efforts to resuscitate such patients reasonable. Children may be the best candidates for heroic measures. Under ideal conditions, hypothermic cardiac arrest patients may reasonably be admitted to an intensive care unit for a 4- to 5-hour trial of rewarming with CPR in progress. Manual CPR should be replaced by mechanical methods if the equipment is available. The oxygen-powered “thumper” has been successful during prolonged hypothermic resuscitation. Absence of responsiveness to treatment, in conjunction with a highly elevated potassium level, is an indication for termination of resuscitative efforts.

**Airway Management**

Maintain a secure functioning airway for hypothermic patients, just as in any critically ill patient. With mild hypothermia, deliver heated, humidified oxygen by face mask. Recognize that a hypothermic patient can be combative and uncooperative and may require arm restraints if a mask is used. Intubate patients with decreased sensorium who cannot reliably maintain their airway or hypothermic patients who may be hypoxic. ET intubation may be performed safely without the added risk of ventricular dysrhythmias. The technique for ET intubation depends on the specific circumstances and the expertise of the operator. Once an ET tube has been placed and secured, use it to provide warm humidified oxygen. There is no evidence that tracheal intubation is detrimental in severely hypothermic patients, and it should be considered if indicated for ventilation, oxygenation, or airway protection.

**Acid-Base Disturbances**

Acid-base disturbances are variable and can lead to metabolic acidosis from carbon dioxide retention and to lactic acidosis or metabolic alkalosis from decreased carbon dioxide production or hyperventilation. Interpretation of arterial blood gases in a hypothermic patient has been the cause of some confusion. Previously, it was suggested that all blood gases be corrected for temperature with correlation factors. With a decrease in temperature of 1°C (33.8°F), pH rises 0.015, carbon dioxide pressure (PCO₂) drops by 4.4%, and oxygen pressure (PO₂) drops 7.2% relative to values that would be obtained with blood analyzed under normal conditions. Despite the conversion guide, optimal or normal values in hypothermia have not been well documented. Other literature supports the use of uncorrected arterial blood gas values to guide therapy with bicarbonate or hyperventilation. This approach appears appropriate to support optimal enzymatic function. Gradual correction of acid-base imbalance will allow increased efficiency of the bicarbonate buffering system as the body warms. Arterial pH did not correlate with patient death in the Multicenter Hypothermia Study and should not be used as a prognostic guide to resuscitation.

**Coagulopathies**

Abnormal clotting occurs frequently in hypothermic patients, probably because cold inhibits the enzymatic coagulation cascade. Hypothermia-induced coagulopathy does not result from excessive clot lysis, but rather from impaired clot formation. Platelet function is also impaired during hypothermia because production of thromboxane B₂ is inhibited. Hypothermia-induced platelet aggregation with or without neutrophil involvement has been associated with neurologic dysfunction in patients undergoing surgical procedures. Hypercoagulability with a risk for thromboembolism may also occur, but the importance of cold-induced coagulopathy mainly involves patients with coincidental trauma. Such victims often have bleeding that is difficult to control. Replace appropriate clotting factors and use warm blood to limit further blood loss and worsening of the hypothermia.

**Trauma and Hypothermia**

Mortality is increased in trauma patients with temperatures below 32°C (< 89.6°F). It is not clear whether this increased mortality is actually a result of the hypothermia or whether the hypothermia is merely an indicator of severe injury and response to a massive transfusion of cold fluid. Patients with severe trauma are prone to hypothermia because their injuries often expose them to environmental heat loss. Patients with hemorrhagic shock are often acidotic and coagulopathic, leading to the “lethal triad” of acidosis, coagulopathy, and hypothermia. These patients are at extremely high risk for multisystem organ failure and death. Concurrent alcohol intoxication may add to the heat loss as a result of its vasodilatory effects on cutaneous vasculature and the prolonged cold exposure secondary to altered mental status. Victims of severe injury also lose heat because of exposure during resuscitation and rapid administration of cold fluids.

It is unknown to what degree correcting the hypothermia improves outcome. Nevertheless, devices to rapidly infuse warm fluids such as the Level 1 fluid warmer (Level 1 Technologies, Rockland, MA) and the Thermostat 900 (Arrow International, Reading, PA) are frequently used to warm large-volume fluid transfusions. Use of these devices seems reasonable to prevent the hypothermia associated with massive transfusions (see Chapter 28). Their use for hypothermia not associated with severe trauma is limited by the relatively low fluid requirements of patients with environmental exposure. Another Thermostat device (Aquarius Medical Corp., Phoenix, AZ) is used to accelerate recovery from hypothermia by mechanically distending blood vessels in the hand, thereby increasing transfer of exogenous heat to the body core.

**Pharmacotherapy and Monitoring**

Hypothermia alters the pharmacodynamics of various drugs. It markedly alters drug kinetics, but not enough is known about this phenomenon to define specific therapeutic guidelines. Administer drugs with caution to hypothermic patients. Because of the negative effects of hypothermia on both hepatic and renal metabolism, toxic levels of medications can accumulate rapidly after repeated use. Avoid certain drugs, such as digitalis. Sinus bradycardia and most atrial arrhythmias do not require pharmacologic treatment because the majority
resolve with rewarmin'. Transient ventricular dysrhythmias also do not require treatment. Bretylium is the preferred agent for patients requiring medication for ventricular dysrhythmias, but lidocaine, magnesium, isoproterenol, and amiodarone have also been used. For severe acidosis (pH < 7.1), IV sodium bicarbonate can be used with extreme caution. Vasopressors should be used with care, perhaps in much smaller doses than usual, because of the arrhythmogenic potential and the delayed metabolism of these agents. A review of intensive care unit admission of hypothermic patients found that treatment with vasoactive drugs was an independent risk factor for mortality, but this phenomenon remains poorly understood. In animal studies, use of epinephrine impaired myocardial efficiency in cases of moderate hypothermia. There was no advantage to repeated doses of epinephrine or high-dose epinephrine in hypothermic cardiac arrest animal models. The use of inamrinone, formerly known as amrinone, has been investigated in cases of deliberate mild hypothermia. Initial results indicate that inamrinone accelerates the cooling rate of the core temperature, thereby potentially limiting its usefulness in the management of AH.

Administer IV fluids slowly to prevent fluid overload potentiated by the decreased cardiac output. Fluids should be started early because intravascular volume is depleted in most hypothermic patients. D5W has been advocated as the ideal initial resuscitation fluid. Avoid potassium until electrolytes are measured and normal renal function is confirmed. Check serum levels of creatine phosphokinase in hypothermic patients, which may indicate rhabdomyolysis. If elevated, carefully monitor renal function. Replace fluids aggressively because this may help prevent the development of renal failure. In severely hypothermic patients, consider placing a Swan-Ganz catheter and closely monitor urinary output to assist in fluid management. The risk of precipitating ventricular fibrillation should be weighed against the potential benefits of the Swan-Ganz catheter.

It should be emphasized that hypothermic patients exhibit a "classic physiologic response" that may be somewhat protective. This response depends on the severity of the decrease in core temperature and classically consists of hypotension, hypoventilation, depressed mental status, and bradycardia. This prohibits a precise recommendation of the indications and use of medications, intubation, CPR, and other resuscitative interventions, which are better defined in normothermic patients. Hypothermic patients with a blood pressure, respiratory rate, or mental status that would prognosticate certain morbidity in normothermic patients may recover with minimal intervention on their normal pre-hypothermic state. Avoid aggressive therapies or medications aimed at providing hypothermic patients with vital signs that would be desirable in normothermic patients but may be supraphysiologic in hypothermic patients.

**Frostbite**

Hypothermic patients frequently suffer other forms of cold-related injuries in addition to their systemic hypothermia. The mildest form of frostbite is termed *frostnip*, a condition that involves only the skin and spares subcutaneous tissue. The skin is blanched and numb, but the injury is immediately reversible with no permanent sequelae if the area is quickly rewarmed. Rewarm rapidly in a water bath at 40°C to 42°C (104.0°F to 107.6°F). Frostnip occurs most frequently on the distal ends of the extremities, the nose, and the ears. Nonfreezing temperatures also produce *trench foot*, an intermediate step in the progression to true frostbite. Trench foot is the result of prolonged immersion in cold water. Rewarm patients and apply dry dressings.

In frostbite, the body parts most susceptible are those farthest away from the body’s core: the hands, feet, earlobes, and nose. Exposure of the fingers to severe cold leads to cold-induced vasodilation. Apical structures rich in arteriovenous anastomoses can shunt blood flow away from tissues. Freezing of the corneas has been reported to occur in individuals who keep their eyes open in highwind chill situations without protective goggles (e.g., snowmobilers and skiers).

The pathophysiology of frostbite includes three pathways of tissue freezing: (1) through the extracellular formation of ice crystals, (2) hypoxia as a result of cold-induced local vasoconstriction, and (3) release of inflammatory mediators. These pathways often occur simultaneously and intensify the tissue damage. At the early stages of frostbite the “hunting reaction” is observed whereby the body alternates between periods of vasoconstriction and periods of vasodilatation. As the temperature continues to decrease, the reaction stops and vasodilation persists. Cold also increases blood viscosity, promotes vasospasm, and precipitates the formation of microthrombi. Release of the inflammatory mediators, prostaglandin F2 and thromboxane A2, causes further vasoconstriction leading to cell death. Release of these mediators peaks during rewarming, and cycles of recurrent freezing and rewarming only increase their tissue levels. Avoid rewarming until refreezing can be prevented.

The clinical signs and symptoms of frostbite vary according to the degree of injury. Though useful clinically, the degree classification does not predict the extent of further tissue damage. The appearance of the affected extremity depends on the extent of the frostbite. With superficial frostbite, the affected extremity appears pale, waxy, and numb. The limb has poor capillary refill and is very painful on rewarming. With deeper frostbite, the affected extremity is hard, solid, and blanched. Hemorrhagic blisters may be present (Fig. 65.6). Initially, there is no pain or feeling in the frostbitten extremity. After rewarming, severe edema and blistering develop in the affected areas, and victims eventually exhibit dry gangrene, mummification, and ultimately tissue sloughing.

---

**TABLE 65.4 Commonly Used Medications for Hypothermia**

<table>
<thead>
<tr>
<th>Clinical Situation</th>
<th>Medication</th>
<th>Dosage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypoglycemia</td>
<td>D50W</td>
<td>1 mg/kg IV</td>
</tr>
<tr>
<td>Alcoholic/ malnourished</td>
<td>Thiamine</td>
<td>100 mg IV</td>
</tr>
<tr>
<td>Altered mental status</td>
<td>Naloxone</td>
<td>0.4–2 mg IV</td>
</tr>
<tr>
<td>Ventricular fibrillation</td>
<td>Bretylium*</td>
<td>5 mg/kg IV</td>
</tr>
<tr>
<td></td>
<td>Magnesium sulfate</td>
<td>100 mg/kg IV</td>
</tr>
</tbody>
</table>

*The role of more available antidysrhythmics such as amiodarone in patients with hypothermia remains to be determined. D5W: 50% dextrose in water; IV: intravenously.
in the frozen state. The most practical way to rewarm an extremity is to totally immerse the area in warm water at 40°C to 42°C (104.0°F to 107.6°F) for 15 to 30 minutes. Carefully protect the affected area to ensure that the tissue is not additionally injured by contact with the sides or rim of the container. After thawing, meticulously protect the area from injury. Elevate the extremity and place cotton or gauze between the toes or fingers to limit maceration. At some point, necrotic tissue should be débrided, most often after the ED encounter has allowed identification of viable tissue; however, the ideal timing and best method or intervention have not been elucidated. A conservative approach is advocated. One method is to débride white or clear blisters. Leave hemorrhagic or dark blisters intact because disruption may theoretically cause damage to the vascular supply and viable tissue.

Use topical aloe vera, a thromboxane inhibitor, and administer systemic antiprostaglandins such as ibuprofen. Aspirin and ibuprofen inhibit the arachidonic and acid cascade, although there is no evidence of efficacy for either agent. Some prefer ibuprofen because it also produces fibrinolysis. The use of semiocclusive dressings has shown promising results in the management of deep frostbite injuries of the fingertips. Provide tetanus prophylaxis. Adjuvant therapies involving the use of heparin or low-molecular-weight heparin, warfarin, vasodilators, corticosteroids, or immediate surgical sympathectomy have failed to improve outcomes.

The ideal intervention to ameliorate or limit tissue injury has not been proved, and it is uncertain if any protocol will prove effective. Mixed success has been achieved with the use of hyperbaric oxygen and thrombolitics. In a small study of frostbite victims, Twomey and coworkers suggested the following treatment algorithm for severe frostbite: (1) rapid rewarming; (2) assessment of the patient’s clinical appearance; (3) early-phase 99mTc scintiscan to assess the distal circulation; (4) administration of tissue plasminogen activator (t-PA), 0.15 mg/kg by IV bolus, followed by 0.15 mg/kg per hr to a maximum dose of 100 mg over a 4- to 6-hour period, for patients with digits or limbs showing no flow and an absence of contraindications; (5) therapeutic heparin for 3 to 5 days; (6) administration of warfarin to an international normalized ratio two times control for 4 weeks; (7) pain management as needed; (8) ibuprofen, 400 to 600 mg orally four times daily; (9) light dressings with topical antimicrobials; and (10) no ambulation on frostbitten feet.

Figure 65.6 Frostbite. Rapid rewarming is the treatment of choice; immersion in warm water (40°C to 42°C) for 15 to 30 minutes is a practical method. A conservative approach to frostbite débridement is suggested, with many alternatives acceptable. One approach is to débride white or clear blisters and leave hemorrhagic or dark blisters intact.

Favorable prognostic signs for frostbite include intact sensation, normal color, warm tissues, early appearance of clear blisters, and edema. Early intervention is critical in terms of the ultimate outcome. Delay in seeking medical care for more than 24 hours is associated with an 85% likelihood that surgical intervention will be required. Patients seen within the first 24 hours require surgery less than 30% of the time. The predictive value of the initial physical examination is limited, but the presence of nonblanching cyanosis, hemorrhagic blisters, and impaired sensation appears to indicate a poor prognosis.

Based on early bone scans and retrospective studies, researchers from France proposed a new classification for predicting frostbite outcomes on day 0. Four degrees of severity are defined. With first degree, there is complete recovery. Second degree often leads to soft tissue amputation. With third degree, bone amputation is needed, and with fourth degree, systemic effects occur.

It is critical to stabilize hypothermia and other life-threatening conditions before warming frostbitten extremities.

Rapid rewarming is the treatment of choice for frostbite. The aim is to limit the length of time that the tissue remains in the frozen state. The most practical way to rewarm an extremity is to totally immerse the area in warm water at 40°C to 42°C (104.0°F to 107.6°F) for 15 to 30 minutes. Carefully protect the affected area to ensure that the tissue is not additionally injured by contact with the sides or rim of the container. After thawing, meticulously protect the area from injury. Elevate the extremity and place cotton or gauze between the toes or fingers to limit maceration. At some point, necrotic tissue should be débrided, most often after the ED encounter has allowed identification of viable tissue; however, the ideal timing and best method or intervention have not been elucidated. A conservative approach is advocated. One method is to débride white or clear blisters. Leave hemorrhagic or dark blisters intact because disruption may theoretically cause damage to the vascular supply and viable tissue.

Use topical aloe vera, a thromboxane inhibitor, and administer systemic antiprostaglandins such as ibuprofen. Aspirin and ibuprofen inhibit the arachidonic and acid cascade, although there is no evidence of efficacy for either agent. Some prefer ibuprofen because it also produces fibrinolysis. The use of semiocclusive dressings has shown promising results in the management of deep frostbite injuries of the fingertips. Provide tetanus prophylaxis. Adjuvant therapies involving the use of heparin or low-molecular-weight heparin, warfarin, vasodilators, corticosteroids, or immediate surgical sympathectomy have failed to improve outcomes.

The ideal intervention to ameliorate or limit tissue injury has not been proved, and it is uncertain if any protocol will prove effective. Mixed success has been achieved with the use of hyperbaric oxygen and thrombolitics. In a small study of frostbite victims, Twomey and coworkers suggested the following treatment algorithm for severe frostbite: (1) rapid rewarming; (2) assessment of the patient’s clinical appearance; (3) early-phase 99mTc scintiscan to assess the distal circulation; (4) administration of tissue plasminogen activator (t-PA), 0.15 mg/kg by IV bolus, followed by 0.15 mg/kg per hr to a maximum dose of 100 mg over a 4- to 6-hour period, for patients with digits or limbs showing no flow and an absence of contraindications; (5) therapeutic heparin for 3 to 5 days; (6) administration of warfarin to an international normalized ratio two times control for 4 weeks; (7) pain management as needed; (8) ibuprofen, 400 to 600 mg orally four times daily; (9) light dressings with topical antimicrobials; and (10) no ambulation on frostbitten feet.

Bruen and colleagues in a small retrospective study, reported that administration of t-PA within 24 hours of frostbite injury improved tissue perfusion and reduced amputations. The protocol included t-PA administered at an initial rate of 0.5 to 1.0 mg/hr into the extremity via a femoral or brachial arterial catheter sheath. Heparin was also administered at 500 Units/hr into the intraarterial catheter. Thrombolytic therapy for frostbite is encouraging, but the exact parameters for its use are still being investigated. Iloprost, a prostacyclin analogue, has vasodilatory properties that mimic a chemical sympathectomy. The risk of amputation was significantly lower in a controlled trial of patients with severe frostbite who received IV iloprost plus aspirin after thawing. Agents that can inhibit the formation of free radicals are also promising. Such agents include superoxide dismutase, prostaglandin E1 analogues, and drugs containing antiprostaglandin activity such as pentoxifylline. The use of antibiotics is controversial, although some authors advocate agents effective against Staphylococcus and Streptococcus (e.g., cephalosporins, penicillins). Avoid débridement of tissue in the ED. Give analgesics (IV opioids) as needed.
Cold Water Immersion and Submersion

One of the leading causes of hypothermia remains cold water immersion or submersion. In a retrospective review of AH cases in a 3-year period, submersion hypothermia accounted for the greatest number of cases. Unlike cases of AH caused by cold exposure, risk factors are harder to identify because of the high mortality from drowning. Studies have shown that at cold water temperatures (8°C [46.4°F]), core cooling occurs at slower rates in persons with increased body mass and subcutaneous fat and at faster rates with increased voluntary activity (e.g., treading water). Risk factors for submersion hypothermia include impaired performance and the initial cardiorespiratory response to immersion. Studies have demonstrated that, in healthy volunteers, swimming efficiency and length of stroke decreased whereas the rate of stroke and swim angle increased as the water temperature dropped.

The body’s response to cold water immersion (head out) has previously been described as occurring in three phases. The initial phase involves the “cold-shock response,” which typically occurs within the first 4 to 6 minutes. Signs include peripheral vasoconstriction, gag reflex, hyperventilation, and tachycardia. At this stage, there is a higher incidence of sudden death resulting from hypocapnia, inability to hold one’s breath, and increased cardiac output. After the initial cold-shock response, the body undergoes profound cooling of the peripheral tissues. The peripheral cooling tends to be the greatest in the hands, which leads to incoordination and difficulty grasping. With prolonged immersion in cold water, heat is lost from the body quicker than it is produced, with the individual quickly progressing to hypothermia. In cases of cold water submersion, researchers have found that rapid cooling is protective against neurologic impairment and increases the chance of survival. There are numerous reports in the literature of survival in children after cold water submersion but very few reports in adults. There are also reports of survival after up to 66 minutes of cold water submersion. Children tend to have a better prognosis because of the presence of the mammalian dive reflex and a greater body surface area-to-mass ratio, which allows more rapid cooling. A 2007 case was reported of a 2-year-old boy who suffered from severe hypothermia after falling into ice water. On discovery, cardiac arrest and asystole were present and the first measured temperature was 23.8°C (74.8°F). The patient was rewarmed by ECC with cardiopulmonary bypass and was discharged 9 days later without any sequelae. Five poor prognostic factors for near-drowning in pediatric patients include: (1) maximum submersion time longer than 5 minutes, (2) comatose on arrival at the ED, (3) arterial blood gas pH less than 7.10, (4) age younger than 3 years, and (5) resuscitation not attempted for at least 10 minutes after rescue. Adults tend to have higher mortality rates because of the following: (1) lack of the mammalian dive reflex and (2) slower rates of cooling secondary to lower body surface area-to-mass ratios than in children. Recent reports of hypothermia and drowning in commercial fishing institutions noted a strong protective association with the use of personal floatation devices, particularly immersion suits, in surviving cold water-related events in adults.

Various mechanisms of brain and body cooling during submersion hypothermia have been described, including the mammalian dive reflex, cold-induced changes in release of neurotransmitters, and water ventilation. The mammalian dive reflex prevents or delays aspiration or ventilation until the body has cooled to a point at which protection against hypothermia occurs. Much attention has focused on the theory of water ventilation as a key component of accelerated brain cooling. Animal studies comparing immersed (head out) and submersed dogs found that cooling rates were faster in submersed dogs than in immersed dogs. The submersed dogs cooled by convective heat exchange in the lungs, whereas the immersed dogs cooled by surface conduction only. Laboratory data obtained after the submersion indicates that there was indeed ventilation exchange in the water. The body also undergoes a relative bradycardia as another protective measure. Bradycardia is inversely proportional to the water temperature, with heart rates reaching 18 beats/min in water at 10°C (50°F). Many authors advocate therapies aimed at symptoms resulting from near-drowning rather than severe hypothermia because in fatal cases of submersion, death occurs too rapidly for hypothermia to be a significant contributor. Complications of near-drowning include pneumonia, lung edema, hemorrhagic pancreatitis, and skin edema.

Environmental Hypothermia Conclusion

Mortality rates from AH are decreasing, and this is linked to increased recognition and advanced therapy. Caution should be used when extrapolating published data obtained in adults to children. With the exception of severe hypothermia, the prognosis correlates mostly with the presence or absence of underlying disease states. Studies have shown that the prognosis is excellent in patients in whom no hypoxic event precedes the hypothermia and no serious underlying disease states exist. Previously healthy individuals usually have full recovery with mortality rates lower than 5%, but patients with coexisting medical illnesses reportedly have mortality rates higher than 50%.

As a general guideline, take a conservative approach to rewarming stable hypothermic patients, with avoidance of overtreatment and selective and careful use of invasive monitoring. Evaluate a hypothermic patient’s “physiologic” hypotension, hypoventilation, and bradycardia with regard to that expected for the given core temperature. Because death is related more to underlying illnesses than to hypothermia, some sources do not believe that invasive rewarming modalities are useful for poikilothermic patients with severe underlying disease.

With moderate hypothermia, underlying problems should be sought, passive rewarming and basic support started, and less invasive core rewarming begun. This approach should include mask ventilation with warm humidified air or oxygen in conscious patients and intubation and ventilation in unconscious patients. In selected patients, gastric or peritoneal lavage with warm fluid may be considered. For severely hypothermic, unstable patients, cardiac bypass and thoracic lavage may offer additional benefits, including rapid warming rates and direct heart warming. The benefits should be weighed against the institutional capabilities, time, expense, and the danger for complications that these procedures entail. Although these patients often require tremendous time and resources, the literature is clear on one point: incredible recoveries are possible.

Targeted Temperature Management for Cardiac Arrest

In contrast to environmental hypothermia or AH, therapeutic hypothermia (TH, cooling to 32°C–34°C) or targeted
temperature management (TTM, cooling to 32°C–36°C) refers to the intentional lowering of temperature in patients for clinical benefit. Whereas lowered temperature may be protective for several diseases, particularly following an ischemic injury, the greatest benefit of active cooling has been demonstrated in the clinical context of out-of-hospital cardiac arrest (OHCA). Application of hypothermia as a therapeutic agent dates back to over 5000 years ago. In 1700 the first systematic experimental studies were done on humans to determine the effects of cooling on the body. However, it was not until 1958 that the first clinical trial of hypothermia on comatose cardiac arrest patients was conducted. Since then TTM (a term that now encompasses TH) has proven to be one of the few treatment lines allows for a wider range of target temperatures that the HACA trial. However, the Nielsen study enrolled patients neurological benefit at 180 days comparable to that seen in 72 hours. Both groups demonstrated a similar mortality and control to prevent fever of less than 37.5°C for an additional 36°C for 24 hours following OHCA, with further temperature of cooling on the body. However, it was not until 1958 that mental studies were done on humans to determine the effects OHCA patients in the most recent I study was conducted for pediatric patients through the Thera-utic Hypothermia After Pediatric Cardiac Arrests Out of Hospital (THAPCA-OH) to evaluate for the efficacy of TH (TTM to 33) versus therapeutic normothermia (TTM to 36.8). Primary outcome of good neurological outcome at 12 months resulted in no significant difference between the groups. It must be noted that there are grave differences in the pediatric patients; the leading cause of arrest in pediatric patients was respiratory (72%) and the proportion of patients with shockable rhythms was only 8% versus 80% in the Neilsen study. Therefore, a potentially important clinical benefit cannot be ruled out. Many unanswered questions remain about the depth (higher temperature vs. lower temperature), duration (longer vs. shorter), and timing of TTM in pediatric patients. However, as of 2015, the American Heart Association national recommendations conclude that for comatose children who achieve return of spontaneous circulation (ROSC), temperature should be monitored closely and fever treated aggressively (see Table 65.5 for specific recommendations for OHCA).

The clinical implementation of TTM for OHCA occurs within hours after a patient has had ROSC and is generally maintained for 24 hours. An opportunity to expand the application of cooling during CPR prior to ROSC has been highlighted in the ILCOR systematic review by Donnino and colleagues. Although it remains clinically challenging to implement, there is mounting evidence that such intra-arrest therapeutic hypothermia (IATH) could have a profound benefit that exceeds TTM alone. Animal studies have shown that cooling to 32°C during CPR is highly protective of heart and brain function and that survival protection is lost if active cooling is delayed by even 20 mins. This suggests that an important therapeutic window exists for cooling during CPR. Furthermore, a systematic review of the literature by Scolletta concludes that IATH improves survival and neurological outcome compared to normothermia and conventional hypothermia in experimental models.

In summary, the role of environmental hypothermia and hyperthermia in causing human injury is well known and treatment strategies have advanced. In parallel, interest in intentional cooling and fever control for preventing the human injury of disease has a long history in medicine, with tremendous advances made in its application to OHCA patients. The time-sensitive care of patients that involves achieving optimal temperature for protection from injury remains a unique aspect of emergency medicine, with great potential to save the lives of some of our sickest patients.

A concise overview of all the recent TTM recommendations for adult, pediatric, and neonatal guidelines for OHCA are listed in Table 65.5.

**PROCEDURES PERTAINING TO HYPERThERMIA**

As a result of global climate change, it is projected that worldwide there will be a significant increase in the number and intensity of heat waves with resultant deaths from hyperthermia and heat-related illness. Temperature extremes and variability will remain important determinants of overall health, especially in the vulnerable populations of the elderly, children, and those with chronic illness. The mortality associated with heatstroke accounts for more than 200 deaths per year in the United States. In the United States from 1999 to 2003, a total of 3442 deaths were attributed to extreme heat exposure. During the heat wave of 2003, France reported 15,000 excess deaths. The morbidity associated with heat-related illness is on the rise. Nationally, an estimated 54,983 patients were evaluated in US EDs for exertional heat-related illness from 1997 to 2006. This represents a 133% increase over the 10-year period. Lack of heat acclimatization during extreme environmental conditions is responsible for the increasing percentage of heat-related illness, particularly in younger populations and agricultural workers.

Other important causes of hyperthermia include malignant hyperthermia (MH) and neuroleptic malignant syndrome
(NMS). Both MH and NMS are largely iatrogenic and are mostly triggered by modern pharmacologic therapy.150 There is evidence that MH involves a dose-dependent response, but the minimum dose is unknown.150 The incidence of hyperthermic conditions induced by psychostimulant drugs of abuse, such as morphine and amphetamine derivatives, continues to increase.151

As with hypothermia, it is important to use the proper thermometer capable of detecting a wide range of body temperatures. Heatstroke remains a common clinical problem with significant morbidity and mortality. A variety of cooling techniques have been advocated since World War II. Although some cooling techniques have been compared in controlled human and animal models of heatstroke, our practice decisions are not based solely on the theoretical rate of cooling. Other important factors include the ease of use, rapidity of initiation, and safety.

Before considering the various cooling techniques, it is essential that the underlying disorders of hyperthermia be clearly understood. Heat illness represents a broad spectrum of disease ranging from mild heat exhaustion to severe heatstroke. The latter includes disorders such as MH and NMS. Treatment of this spectrum of disease requires a discriminating approach, including supportive care only for heat exhaustion and rapid cooling for heatstroke. MH requires specific pharmacologic therapy (e.g., dantrolene), in addition to cooling measures. A brief discussion of hyperthermic disorders is necessary before describing cooling techniques.

### Normal Thermoregulation

Body temperature typically follows a diurnal pattern, with an increase from approximately 36°C (96.8°F) in the early morning to 37.5°C (99.5°F) in the late afternoon, and is normally tightly regulated by an effective thermoregulatory system.152 Heat is produced as a by-product of metabolic processes and when ambient temperature exceeds body temperature. Body temperature increases when the rate of heat production exceeds the rate of heat dissipation. The brain’s thermal center is located in the preoptic nucleus of the anterior hypothalamus. In response to rising core temperature, this thermal center activates efferent fibers of the autonomic nervous system to produce vasodilation and increase the rate of sweating. Vasodilation dissipates heat by convection, and sweat dissipates heat by evaporation.

Hyperthermia occurs when the thermoregulatory mechanisms are overwhelmed by excessive metabolic production of heat, excessive environmental heat, or impaired heat dissipation. Different age-related thermoregulation strategies are used when dealing with heat stress. Children have a greater surface area-to-mass ratio and a lower sweating rate and rely more on "dry" heat exchange to dissipate heat. On the contrary, adults use evaporative heat loss as the primary heat dissipation technique. With primary aging, the reflex cutaneous vasconstriction and vasodilation capabilities are impaired, thereby allowing increased susceptibility to complications from heat-related exposure.153 Fever occurs when the hypothalamic
set-point is increased by the action of circulating pyrogenic cytokines, which cause peripheral mechanisms to conserve and generate heat until the body temperature rises to the elevated set-point. Hyperthermia and fever cannot be differentiated clinically on the basis of the magnitude of temperature or on the pattern of its changes.154,155

Types of Hyperthermia

**Mild Heat Illness**

Heat cramps and heat exhaustion are induced by a hot environment.161 The body’s heat dissipation mechanisms are generally able to keep up with heat production and absorption in these disorders. Symptoms are largely due to the mechanisms used by the body to dissipate heat, and body temperatures remain at or near normal. Rapid cooling techniques are not required, and supportive care and hydration in a cool environment are usually adequate therapy.

Heat cramps are intensely painful but generally benign involuntary skeletal muscle spasms. The pain most often occurs in the calf, hamstring, or quadriceps muscles but may also involve the arms and back. The cramps may be severe and prolonged but only rarely lead to rhabdomyolysis. Heat cramps occur after strenuous exercise or heavy labor in a hot environment. Heat cramps were previously thought to be the result of dehydration associated with significant loss of sodium chloride, but some clinical observations have proved that heat cramps can occur at rest or during exercise under any environmental conditions.157 Resting in a cool environment plus vigorous oral fluid replacement with isotonic solutions is usually adequate therapy, but in some cases IV saline is required.157 The benefits of oral rehydration over IV hydration directly relate to oropharyngeal stimulation, which influences the release of antidiuretic hormone (arginine vasopressin), cutaneous vasodilation, thirst sensation, and mean arterial pressure.158 A common mistake is to rely on thirst to indicate dehydration. The pain of severe cramping may be resistant to narcotics in the absence of adequate fluid replacement.

Heat exhaustion, commonly referred to as heat syncope, is a poorly defined syndrome with nonspecific symptoms that occur after heat exposure.199 Many have suggested replacing the current terminology of heat exhaustion with the term exercise-associated collapse.160 Malaise, flulike symptoms, orthostasis, dehydration, nausea, headache, and collapse may all occur. Previously it was believed that heat exhaustion is the result of dehydration-induced heat retention that is not severe enough to cause heatstroke.161 There is modern evidence that postural hypotension developing after exercise is the result of exercise-induced changes in blood pressure regulation. These changes involve recalibration of the arterial baroreflex to lower pressures after exercise, impaired sympathetic vascular regulation, and H1 and H2 receptor–mediated vasodilation.161 When compared with the more severe heat disorder of heatstroke, mental status is normal and body temperature is normal or mildly elevated with heat exhaustion. There does not appear to be any thermoregulatory failure in persons with heat exhaustion. Rehydration, rest, and supportive care in a cool environment are adequate therapy for heat exhaustion.156,157,160 Some authors advocate cooling and placing the patient either in the supine position with the legs elevated or seated with the head between the knees to decrease skin blood flow and increase venous blood flow to the heart.161 Recovery is usually evident within a few minutes to hours. Occasionally, heat exhaustion is accompanied by heat cramps, thus presenting a confusing scenario if the diagnosis is not suspected. Rapid cooling techniques, IV hydration, and advanced therapies are not usually required, but patients should be observed for progression to heatstroke because heat exhaustion and heatstroke are a continuum of one disease process.158,161

Heatstroke

When the body’s normal heat dissipation mechanisms are overwhelmed, core temperature elevation and heatstroke develop rapidly. Heatstroke is a state of thermoregulatory failure.161 Previously, the morbidity and mortality associated with heatstroke were attributed to the magnitude of the hyperthermic response.161 Recent literature has described a more complex interaction between cytokines, coagulation, and the systemic inflammatory response syndrome (SIRS), with endotoxin and cytokines being implicated as key mediators of heat-induced SIRS.162 Two forms of heatstroke are described in the literature. Classic (nonexertional) heatstroke usually occurs during summer heat waves. The poor, urban elderly, infants, homeless, and persons with impaired mobility are at greatest risk.163,164 Dehydration, lack of air-conditioning, obesity, neurologic disorders, hyperthyroidism, cardiovascular disease, impaired mentation, and medications that interfere with heat dissipation (e.g., phenothiazines, diuretics, and anticholinergics) predispose this population to heatstroke.163,164 Exertional heatstroke, a consequence of strenuous physical activity, usually afflicts a younger segment of the population. Highly motivated, poorly acclimatized, or unconditioned athletes and overweight military recruits are common victims, as are individuals who perform heavy physical labor in hot, humid conditions (Fig. 65.7).165–166 With exertional heatstroke, the risk appears to

![Figure 65.7](Image)

be greatest in individuals performing high-intensity exercise for relatively short durations. A retrospective review of long-distance cyclists participating in the California AIDS (acquired immunodeficiency disease) Ride found that as the number of chronic medical illnesses increased, so did the risk for development of an exertional heat-related illness. Human immunodeficiency virus seropositivity alone was not associated with an increased risk for exertional heat-related illness.

The degree of hyperthermia necessary to produce heatstroke in humans is unknown. In tissue culture cells, thermal injury is observed with temperatures in the range of 40°C to 45°C (104°F to 113°F). Studies of hyperthermia in patients undergoing cancer therapy have revealed that tissue sensitivity to heat is increased by relative hypoxia, ischemia, and acidosis.

The key clinical findings in the diagnosis of heatstroke are (1) a history of heat stress or exposure, (2) a rectal temperature higher than 40°C (>104°F), and (3) central nervous system (CNS) dysfunction (altered mental status, disorientation, stupor, seizures, or coma). The cerebellum is very sensitive to heat, and ataxia may be an early clue. Although anhidrosis is described as a classic sign of heatstroke, investigations have demonstrated that cessation of sweating may be a late finding. Failure to consider the diagnosis of heatstroke in a diaphoretic patient with changes in mental status could prove disastrous.

The sequelae of heatstroke are caused by thermal damage to multiple organ systems. Whole-body hyperthermia decreases pulmonary capillary wedge pressure and cerebral vascular conductance and causes an inotropic shift in the Frank-Starling curve. After a hyperthermic event, tissue injury continues. Delirium, seizures, and coma can result from the direct effects of heat on the CNS. Autopsies show profound brain edema after hyperthermic insults. Researchers suggest that in cases of instant death, brain edema from the increased permeability of the blood-brain barrier causes raised intracranial pressure and papilledema, followed by vascular infarction and brain herniation. Cardiovascular collapse results from dehydration, maximal cutaneous vasodilation, and direct heat-induced myocardial depression. Coagulopathies and liver dysfunction (elevated levels of bilirubin and transaminases) occur as a consequence of thermal breakdown, consumption of serum proteins, and direct heat damage to hepatic cells. Children often demonstrate diarrhea. Reduced intestinal blood flow causes barrier dysfunction and endotoxemia. Development of an acute abdomen, bloody diarrhea, dilated loops of bowel on radiographic studies, and unexplained shock should raise suspicion for colon ischemia and pending colonic perforation. Renal failure can result from myoglobinuria (related to rhabdomyolysis) and acute tubular necrosis. Metabolic acidosis is the primary acid-base alteration observed in patients with heatstroke, with the prevalence increasing with the degree of hyperthermia.

Treatment of these sequelae of acute heatstroke does not differ from that of other heat-related disorders, with the sole exception that rapid cooling is necessary to prevent further damage and reverse the heat stress. The more rapidly that rectal temperature is reduced to 38°C (100.4°F), the better the prognosis. The human body tolerates hyperthermia poorly. Unlike patients with hypothermia, in whom slow, gentle rewarming and supportive care often result in a favorable outcome, victims of severe heatstroke must be treated aggressively with measures designed to rapidly lower core temperature. Studies investigating precooling techniques to avoid heatstroke have been relatively unsuccessful in attenuating increases in core body temperature, and such techniques are not recommended. In contrast, whole-body precooling increases overall exercise endurance.

**Malignant Hypertension (MH)**

MH is a pharmacogenetic disease attributable to a medication that triggers a life-threatening, hypermetabolic syndrome. It results from a rare inherited autosomal dominant abnormality in the skeletal muscle membrane and has an incidence of 1 in 50,000 in adults. In response to certain stresses or drugs (Box 65.1), patients with this disorder sustain a potentially lethal hypermetabolic reaction with massive efflux of calcium from the skeletal muscle sarcoplasmic reticulum. This results in contraction of the sarcomeres, skeletal muscle rigidity, increased skeletal muscle metabolism, elevated serum creatine kinase levels, heat production, and finally, systemic hyperthermia. Hyperthermia is a late development and occurs after rigidity has been present for some time and the body's normal heat dissipation mechanisms are overwhelmed. The earliest signs of MH are increased carbon dioxide production, muscle rigidity, and tachycardia. Cardiac output and cutaneous blood flow also increase to maximize the heat loss. Diagnosis of MH is based on the clinical triad of (1) exposure to an agent or stress known to trigger the condition, (2) skeletal muscle rigidity, and (3) hyperthermia.

MH is usually encountered in the operating room while patients are undergoing general anesthesia, particularly with halogenated inhalational agents and depolarizing muscle relaxants. Heat production in anesthetized patients can be profound with as much as a fivefold increase in oxygen consumption. Cases of MH may be encountered anywhere that general anesthetics or neuromuscular blocking agents are used. A massive increase in creatine kinase is a strong indicator of an MH reaction.

As with heatstroke, treatment of MH requires rapid cooling and supportive care for the sequelae described previously. Unlike heatstroke, MH requires specific pharmacologic therapy to stop excessive heat production by skeletal muscle. Dantrolene sodium induces muscle relaxation in patients with MH by blocking release of calcium from the muscle cell sarcoplasmic reticulum. In all cases of MH, the inciting stimulus should be discontinued immediately and dantrolene therapy administered. A dantrolene bolus of 2.5 mg/kg should be started and repeated at 5-min intervals until normalization of the hypermetabolic state is achieved and all MH symptoms disappear.

**BOX 65.1 Triggers of MH**

<table>
<thead>
<tr>
<th>DRUGS</th>
<th>CONDITIONS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Halothane</td>
<td>Heat stress</td>
</tr>
<tr>
<td>Methoxyflurane</td>
<td>Vigorous exercise</td>
</tr>
<tr>
<td>Enflurane</td>
<td>Emotional stress</td>
</tr>
<tr>
<td>Diethyl ether</td>
<td>Ketamine</td>
</tr>
<tr>
<td>Cyclopropane</td>
<td>Trichloroethylene</td>
</tr>
<tr>
<td>Succinylcholine</td>
<td>Chloroform</td>
</tr>
<tr>
<td>Tubocurarine</td>
<td>Gallamine</td>
</tr>
<tr>
<td>Lidocaine</td>
<td>Nitrous oxide</td>
</tr>
</tbody>
</table>

MH, Malignant hyperthermia.
Procainamide has been used successfully when dantrolene is unavailable. It has been suggested that dantrolene administration speeds cooling of heatstroke victims by reducing skeletal muscle heat production. A randomized, controlled trial of the use of dantrolene for heatstroke found no difference between the treatment and placebo groups in terms of cooling time, complications, or length of stay. In 2005, a meta-analysis concluded that there was no role for the use of dantrolene in the management of heatstroke. Currently, dantrolene administration is best reserved for patients with clinical muscle rigidity or suspected MH. Routine use of this drug in heatstroke patients is not recommended. New and promising treatments of MH are being investigated. Researchers have discovered mutations in the gene encoding for the ryanodine receptor calcium release channel (RyR1) in families with MH, which may be the functional basis for MH. Some studies have examined the effects of MH mutations on the sensitivity of the RyR1 to drugs and endogenous channel effectors, including Ca2+ and calmodulin.

**Neuroleptic Malignant Syndrome (NMS)**

First described in the late 1960s, NMS is characterized by hyperthermia, muscle rigidity, altered level of consciousness, and autonomic instability. Mortality from NMS is estimated to be 20% in patients in whom the condition develops. This idiosyncratic disorder follows the therapeutic use of neuroleptic drugs, including phenothiazines, butyrophenones, thioxanthenes, lithium, and tricyclic antidepressants. The reaction is triggered by blockade of dopaminergic receptors and results in skeletal muscle spasticity, which generates excessive heat and impairs hypothalamic thermoregulation and heat dissipation. Muscle rigidity, described as "lead pipe" rigidity in its most severe form, can be manifested as oculogyric crisis, dyskinesia, akinesia, dysphagia, dysarthria, or opisthotonos. NMS occurs in 0.2% of patients who take neuroleptic agents either chronically or acutely. Haloperidol and depot fluphenazine appear to be the most common offending agents. Temperatures can exceed 42°C (107.6°F). The initial agitation often progresses to stupor and coma. Catatonia and mutism may also be present. Autonomic instability is manifested as tachycardia, labile blood pressure, sweating, and incontinence. Ventilations are impaired by the chest wall rigidity.

This syndrome is more likely to occur at the initiation of or after an increase in neuroleptic dosage. Researchers suggest that NMS typically occurs over a period of several days (average in patients taking neuroleptic agents). It may also occur if the use of antiparkinsonian drugs is suddenly discontinued. NMS resembles MH but usually lasts longer (5 to 10 days) after use of the inciting drug is discontinued. The syndrome may be misinterpreted as worsening of an underlying psychiatric disorder, drug intoxication (e.g., cocaine and amphetamines), a severe dystonic reaction, tetanus, or a variety of CNS infections. In addition to agents with increased dopaminergic blocking activity, other risk factors for NMS include dehydration, previous history of dystonia, catatonia, agitation, and iron deficiency. The mortality rate is high, and respiratory failure, renal failure, cardiovascular collapse, or thromboembolic disease usually causes death.

Treatment of severe NMS (i.e., hypotension, hyperthermia, marked rigidity) closely follows that of MH, except that therapy must be maintained for several days until the symptoms resolve. Therapy for NMS involves discontinuation of the triggering agent; rapid cooling; benzodiazepines; a combination of a central dopamine agonist, bromocriptine, or levodopa; dantrolene; and supportive treatment of the ensuing organ failure. Although the effects are not immediate, pharmaceutical therapy is directed at overriding the dopaminergic blockade caused by the offending neuroleptic agent or the dopamine depletion resulting from the cessation of antiparkinsonian medications. There are reports of successful treatment of NMS with subcutaneous apomorphine monotherapy or high-dose lorazepam and diazepam. As with MH, dantrolene (2.5 mg/kg) can be given to treat NMS-induced muscle rigidity. The beneficial response stems not only from effects at the sarcoplasmic reticulum of skeletal muscle but also from central dopamine metabolism of calcium in the CNS. Bromocriptine, a central dopamine agonist, is reported to be efficacious in treating NMS at doses of 2.5 to 10 mg three times a day. Although both these agents have been noted to reduce the duration of hyperthermia, there have been mixed results with the use of bromocriptine and dantrolene.

A more recently described disorder often confused with NMS is serotonin syndrome. This syndrome involves the newer antidepressants fluoxetine, paroxetine, citalopram, fluvoxamine, venlafaxine, and sertraline, which are selective serotonin reuptake inhibitors. These drugs can adversely react with other stronger serotonin receptor agents such as monoamine oxidase inhibitors and nonselective serotonin reuptake inhibitors (clo mipramine and tricyclic antidepressants) to induce a clinical picture similar to that of NMS, only milder. Serotonin syndrome classically occurs when two or more drugs that interfere with serotonin metabolism act synergistically on the 5-HT1A receptor and lead to overstimulation. Drugs that act at any of the other serotonin receptors are not likely to produce the syndrome. The range of symptoms varies from mild gastrointestinal upset, insomnia, and agitation to more severe symptoms that include muscle spasms, seizures, ataxia, rhabdomyolysis, and autonomic instability. Treatment is primarily supportive in milder cases and consists of prompt recognition and withdrawal of the offending agent. Adequate sedation, usually with benzodiazepines, is essential in controlling serotonin syndrome. Most cases resolve spontaneously within 24 hours. For the most severe cases, aggressive intensive care unit management is warranted to prevent renal failure and death.

The drug cyproheptadine (Periactin, TEVA Pharmaceuticals USA, Sellersville, PA) has shown promise in managing the agitation often seen in severe cases. Cyproheptadine is an antihistamine with antiserotonergic properties. Cyproheptadine is available in 4-mg tablets or 2-mg/5 mL syrup. When administered as an antidote for serotonin syndrome, an initial dose of 12 mg is recommended, followed by 2 mg every 2 hours until clinical response is seen. Cyproheptadine is only available in an oral form, but it may be crushed and given through a nasogastric or orogastric tube. There has been limited success with benzodiazepines and β-blockers to treat agitation in these patients. In cases in which serotonin syndrome and NMS cannot be differentiated, benzodiazepines represent the safest therapeutic option. Further study of the newer antipsychotic drugs, such as ziprasidone, a powerful 5-HT1A receptor blocker, may delineate other possible benefits.

**Hyperthermia and Psychostimulant Overdose**

As mentioned previously, the recognized incidence of hyperthermia induced by sympathomimetic psychostimulant drugs of abuse is on the rise. The offending agents most commonly described are

---

**Table 65.1**

<table>
<thead>
<tr>
<th>Drug</th>
<th>Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluphenazine</td>
<td>Commonly described offending agent.</td>
</tr>
<tr>
<td>Procainamide</td>
<td>Used successfully when dantrolene is unavailable.</td>
</tr>
<tr>
<td>Dantrolene</td>
<td>Speeds cooling of heatstroke patients.</td>
</tr>
<tr>
<td>Bromocriptine</td>
<td>Can be used to treat NMS.</td>
</tr>
<tr>
<td>Diclofenac</td>
<td>Known to reduce muscle rigidity.</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>Used to manage agitation.</td>
</tr>
<tr>
<td>Cyproheptadine</td>
<td>Promising in managing agitation.</td>
</tr>
</tbody>
</table>

---

**Chapter 65: Procedures Pertaining to Hyperthermia and Hyperthermia**

1425
cocaine, phencyclidine, amphetamine, and amphetamine derivatives such as 3,4-methylenedioxy-N-methamphetamine (MDMA) (“ecstasy”) and 3,4-methylenedioxy-N-ethylamphetamine (MDEA) (“Eve”). A number of studies have looked specifically at the club drug MDMA and its impairment of heat dissipation. Animal studies in rats suggest that MDMA-induced hyperthermia results not from MDMA-induced release of 5-HT but from increased release of dopamine acting at D1 receptors, thus suggesting a future role for the use of dopamine antagonists in clinical treatment.

Hyperthermia is a common feature of these potentially severe to lethal poisonings with sympathomimetic psychostimulant drugs and may be the primary cause of mortality. Because of the nonlinear pharmacokinetics of MDMA and γ-hydroxybutyrate, it is difficult to estimate a dose-response relationship. Some have applied a pathophysiologic model of exertional heatstroke or NMS to profound cocaine intoxication. In addition to profound hyperthermia (> 42°C [> 107.6°F]), acute rhabdomyolysis, disseminated intravascular coagulation, psychiatric and cognitive dysfunction, renal failure, coma, seizures, and death have been described in these patients. As demonstrated by Roberts and associates, even a patient with a core temperature of 45.5°C (114°F) because of acute cocaine intoxication may survive with aggressive cooling methods. Treatment requires prompt recognition, maintenance of adequate hydration, rapid cooling, and the aggressive use of sedatives or paralyzing agents (or both) to control agitation. Importantly, the longer that psychostimulant-overdosed patients remain hyperthermic, the higher their morbidity and mortality rates. Agitation and seizures must be chemically controlled because they lead to continued generation of heat and muscle injury. Therefore liberal doses of benzodiazepines are recommended. Some have advocated the use of bromocriptine and dantrolene as for MH and NMS, but their efficacy in the setting of drug-associated hyperthermia remains controversial.

**Hemorrhagic Shock and Encephalopathy Syndrome**

The condition of hemorrhagic shock and encephalopathy syndrome in children (mainly infants, but some older children also) resembles heatstroke in adults. The full-blown syndrome includes hyperthermia, coagulopathy, encephalopathy, and renal and hepatic dysfunction. Although there may be an association with concurrent viral illness, the condition generally follows an elevation in temperature, which may be triggered by “bundling” of a child with a low-grade fever. Therapy is largely supportive and includes volume replacement with rapid cooling of the hyperthermic child while sources of bacterial infection are sought and treated.

**Cooling Techniques**

**General Considerations**

Heatstroke mortality is proportional to the magnitude and duration of thermal stress measured in degree-minutes. Delay in cooling may be the single most important factor leading to death or residual disability in those who survive. In addition, advanced age and underlying disease states are significant contributing factors.

Many exertional heatstroke victims are volume-depleted and may exhibit hypotension. Initial stabilization with cooled (room-temperature) IV fluids and correction of electrolyte abnormalities are valuable in hypotensive patients. Traditional sources recommend a rate of 1200 mL over the first 4 hours. Others advise a 2-L bolus over the first hour, with an additional 1 L/hr for the following 3 hours. Seraj and coworkers challenged this more aggressive recommendation. In their study of pilgrims who suffered heatstroke, 65% had normal or above normal central venous pressure (CVP) on arrival. These authors found that an average of 1 L of saline was sufficient to normalize CVP during the cooling period in their patients, who had a mean age of 55 years (range, 31 to 80 years). In older patients, fluid resuscitation should be monitored carefully to avoid pulmonary edema. Regarding antipyretics, there is no indication for either salicylates or acetaminophen in the setting of heatstroke because their efficacy depends on a normally functioning hypothalamus. Overzealous use of acetaminophen could potentiate hepatic damage, and salicylates may promote bleeding tendencies. A study comparing acetaminophen and physical cooling methods found that in patients treated with antipyretics only, mean body temperature increased by 0.2°C (32.4°F) on average.

Given that rapid cooling is accepted as the cornerstone of effective heatstroke therapy, the clinician must choose which cooling technique to use. Studies in animal models are based on the assumption that the fastest cooling technique is the best. In clinical patient care, other factors also influence the choice of technique. Patient access, monitoring, safety, ease of use, availability, and speed are all considerations. A technique that may not be the most rapid but allows easy patient access and is readily available may be preferable to more cumbersome (albeit more rapid once established) cooling techniques in some clinical settings.

The cooling rates achieved in various human and animal studies of heatstroke are summarized in Table 65.6. The advantages and disadvantages of various cooling techniques are outlined in Table 65.7.

In addition to the cooling procedures outlined, it is imperative that the clinician institute the judicious use of sedation, muscle paralysis, or both to control agitation, suppress shivering, reduce energy expenditure, and make the patient receptive to sometimes unpleasant therapies. In general, IV benzodiazepines are the easiest and safest first-line drugs used for sedation.

**Indications for Rapid Cooling**

Rapid cooling should be instituted as soon as the diagnosis of heatstroke (rectal temperature > 40°C [104°F], altered mental status, history of heat stress or exposure) is made. Rapid cooling is also indicated for the treatment of MH and NMS but should be instituted concurrently with discontinuation of the triggering agent or drug and administration of dantrolene. Because studies show that the degree of organ damage correlates with the degree and duration of temperature elevation above 40°C (>104°F), a reasonable clinical goal is to reduce the temperature to below 40°C (<104°F) within 30 minutes to an hour after the start of therapy.

**Contraindications to Rapid Cooling**

Rapid cooling is never contraindicated in patients with heatstroke. Immersion cooling is relatively contraindicated when cardiac monitoring of an unstable patient is required or when limited personnel make constant patient supervision impossible. Iced gastric lavage is contraindicated in patients with depressed airway reflexes unless the airway is protected by ET intubation. Gastric lavage is also contraindicated by conditions that preclude...
placement of an orogastric or nasogastric tube. Cold peritoneal lavage is relatively contraindicated when multiple previous abdominal surgeries make placement of a lavage catheter risky.

**Evaporative Cooling**

Evaporating water is thermodynamically a much more effective cooling medium than melting ice, given an appropriate water-vapor gradient. Evaporating 1 g of water requires 540 kcal. Melting 1 g of ice requires only 80 kcal. In theory, evaporative cooling should be approximately seven times more efficient than ice packing. In practice, evaporative cooling is more efficient. In separate human studies, Weiner and Khogali and Wyndham and colleagues found that evaporative cooling rates were substantially greater than cooling rates with water immersion at 14.4°C (57.9°F). Studies in primate models demonstrated faster cooling rates with evaporative cooling as an adjunct to ice bag placement. Methods using convection and evaporation were more effective than those involving conduction for the treatment of hyperthermia. In clinical practice, ice water immersion or ice packing causes heat loss by conduction and heat consumption by the phase change of melting ice. In healthy volunteers, evaporative cooling techniques (e.g., facial fanning) were associated with decreased thermal sensation and improved thermal comfort.

Despite the continued enthusiasm of some clinicians for ice water immersion, evaporative cooling has been an effective noninvasive cooling technique in human studies. To maximize evaporative cooling rates, several factors must be optimized. Airflow rates must be high and therefore large fans are required. The air must be warm but not humid because evaporation is decreased at lower temperatures. The entire body surface must be exposed to airflow and continuously moistened with water. Ideally, the patient is suspended in a mesh sling to expose the back to airflow and moisture. Finally, the temperature of the water used to moisten the skin must be tepid (15°C [59°F]). Warm forced air is essential for effective evaporation. It maintains good peripheral perfusion and prevents shivering by warming the skin. If the water is ice cold, evaporation will be slowed. Conversely, if it is hot, conductive heat gain may occur. Studies conducted in heat-stressed laying hens demonstrated superior cooling rates with ventral cooling regimes over dorsal cooling.

Weiner and Khogali constructed a sophisticated “body cooling unit” (BCU) to maximize evaporative cooling. Patients in the BCU are suspended in a mesh net. High airflow rates (30 m/min) at a temperature of 45°C (113°F) are maintained both anterior and posterior to the mesh net. Atomized water at 15°C (59°F) is continuously sprayed on all body surfaces.

### TABLE 65.6 Cooling Rates Achieved With Various Cooling Techniques

<table>
<thead>
<tr>
<th>TECHNIQUE</th>
<th>REFERENCE</th>
<th>MODEL</th>
<th>RATE (°C/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Evaporative</td>
<td>Poulton and Walker,195 1987</td>
<td>Human</td>
<td>0.10</td>
</tr>
<tr>
<td></td>
<td>Weiner and Khogali,196 1980</td>
<td>Human</td>
<td>0.31</td>
</tr>
<tr>
<td></td>
<td>Kielblock et al,197 1986</td>
<td>Human</td>
<td>0.09</td>
</tr>
<tr>
<td></td>
<td>Wyndham et al,198 1959</td>
<td>Human</td>
<td>0.07</td>
</tr>
<tr>
<td></td>
<td>Daily and Harrison,199 1948</td>
<td>Rat</td>
<td>0.93</td>
</tr>
<tr>
<td>Immersion (ice water)</td>
<td>Armstrong et al,200 1996</td>
<td>Human</td>
<td>0.20</td>
</tr>
<tr>
<td></td>
<td>Weiner and Khogali,196 1980</td>
<td>Human</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>Wyndham et al,198 1959</td>
<td>Human</td>
<td>0.14</td>
</tr>
<tr>
<td></td>
<td>Magazanik et al,201 1980</td>
<td>Dog</td>
<td>0.27</td>
</tr>
<tr>
<td></td>
<td>Daily and Harrison,199 1948</td>
<td>Rat</td>
<td>1.86</td>
</tr>
<tr>
<td></td>
<td>Costrini,202 1990</td>
<td>Human</td>
<td>0.15</td>
</tr>
<tr>
<td>Selective immersion</td>
<td>Clapp et al,203 2001</td>
<td>Human</td>
<td>0.16</td>
</tr>
<tr>
<td></td>
<td>Torso immersion</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hand/foot immersion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ice packing (whole body)</td>
<td>Kielblock et al,197 1986</td>
<td>Human</td>
<td>0.034</td>
</tr>
<tr>
<td>Strategic ice packs (towels)</td>
<td>Armstrong et al,200 1996</td>
<td>Human</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>Kielblock et al,197 1986</td>
<td>Human</td>
<td>0.028</td>
</tr>
<tr>
<td>Evaporative strategic ice packs</td>
<td>Kielblock et al,197 1986</td>
<td>Human</td>
<td>0.036</td>
</tr>
<tr>
<td>Cold gastric lavage</td>
<td>Syverud et al,204 1985</td>
<td>Dog</td>
<td>0.15</td>
</tr>
<tr>
<td></td>
<td>White et al,205 1987</td>
<td>Dog</td>
<td>0.06</td>
</tr>
<tr>
<td>Cold peritoneal lavage</td>
<td>Horowitz et al,206 1989</td>
<td>Human</td>
<td>0.11</td>
</tr>
<tr>
<td></td>
<td>Bynum et al,207 1978</td>
<td>Dog</td>
<td>0.56</td>
</tr>
<tr>
<td></td>
<td>White,208 1993</td>
<td>Dog</td>
<td>0.14</td>
</tr>
<tr>
<td>Cyclic lung lavage</td>
<td>Harris et al,209 2001</td>
<td>Dog</td>
<td>0.5</td>
</tr>
</tbody>
</table>
Complications of evaporative cooling are rare and more often a result of the underlying disorder than the cooling technique. Wet skin may interfere with electrocardiographic monitoring, but this can usually be avoided by placing electrodes on the patient’s back. Shivering occurs infrequently with this technique when compared with other cooling methods because the water is relatively lukewarm. Because rectal temperature lags behind core (esophageal) temperature, evaporative cooling should be discontinued when rectal temperature reaches 39°C (102.2°F). In cases of mild hyperthermia, tympanic temperature also accurately reflects core temperature and can be useful in this setting. Continued cooling beyond this temperature may lead to subsequent “overshoot hypothermia” as a result of a continued drop in core temperature after active evaporative cooling is discontinued. Shivering indicates that core temperature has decreased to 37°C (89.6°F) or below.

Immersion Cooling

It would seem obvious that the fastest way to cool a heatstroke patient would be immersion in ice water. In a case series of exertional heatstroke patients, iced water immersion cooled patients to lower than 39°C (<102.2°F) within 19.2 minutes. A 2010 study found that cold water immersion for 9 minutes in a 2.0°C circulated water bath until a rectal temperature of 38.6°C was achieved avoided any risk associated with overcooling. Some contemporary sources recommend ice water immersion as the cooling technique of choice for heatstroke. Plattner and associates demonstrated cooling rates with ice water immersion that were six times faster than rates seen with forced air or circulating water.

| TABLE 65.7 Advantages and Disadvantages of Various Cooling Techniques |
|-------------------------------|-----------------|-----------------|
| TECHNIQUE         | ADVANTAGES                      | DISADVANTAGES                        |
| Evaporative      | Simple                         | Constant moistening of skin required |
|                  | Readily available              |                               |
|                  | Noninvasive                    |                               |
|                  | Easy monitoring and patient access |                       |
|                  | Relatively fast                |                               |
| Immersion        | Noninvasive                    | Cumbersome                       |
|                  | Relatively fast                | Patient access and monitoring difficult |
|                  | Low mortality rates            | Shivering                        |
|                  |                                | Poorly tolerated by conscious patients |
| Ice packing      | Noninvasive                    | Shivering                        |
|                  | Readily available              | Poorly tolerated by conscious patients |
| Strategic ice packs | Noninvasive               | Relatively slower cooling          |
|                  | Readily available              | Shivering                        |
|                  | Can be combined with other techniques | Poorly tolerated by conscious patients |
| Cold gastric lavage | Can be combined with other techniques | Relatively slower cooling |
|                  |                               | Invasive                         |
|                  |                               | Requires airway control           |
|                  |                               | Human experience limited          |
| Cold peritoneal lavage | Rapid cooling                  | Invasive                         |
|                  |                               | Human experience limited          |
Costrini reported no fatalities in 252 consecutive young marine recruits with exertional heatstroke who were treated by ice water immersion within 20 minutes of diagnosis. He regarded ice water immersion as superior to other conventional methods described in the literature in reducing mortality rates.

In clinical trials, cold water immersion remains one of the fastest noninvasive cooling techniques available (see Table 65.6). Cold water immersion takes advantage of the high-conductance property of water, which is 25 times that of air. When an adequate evaporative cooling system is not available, immersion may be the cooling technique of choice. Several factors are important in maximizing the rate of immersion cooling. Conductive heat loss depends on cutaneous blood flow to maintain a heat gradient from skin to water. Theoretically, contact with ice water causes skin and subcutaneous vasoconstriction, which blocks heat exchange and turns these structures into insulators. Intense cutaneous vasoconstriction will impede conductive heat loss. Mekjavic and coworkers reported that motion sickness actually potentiates core cooling during immersion by attenuating the vasoconstrictor response to skin and core cooling, thereby augmenting heat loss and the magnitude of the decrease in deep body temperature. Careful monitoring is required because this may predispose patients to hypothermia.

Researchers have suggested that ice water immersion may be superior to cold water immersion because of the establishment of a steeper thermal gradient between the skin and the environment. A study comparing the cooling capacity of ice water immersion (5.2°C [41.4°F], tepid water immersion (14°C [57.2°F]), and passive cooling in experienced distance runners was undertaken. Subjects were randomized to receive high-dose (20-ml) or low-dose (10-ml) ice water immersion. The high-dose ice water immersion resulted in significantly faster decreases in core temperature than the low-dose treatment (6.2°C [37.2°F]/min vs. 3.4°C [38.1°F]/min). The difference in cooling rates with ice water immersion (1°C–5°C [33.8°F–41.0°F]) in heatstroke patients of 0.15°C/min to 0.23°C/min (32.3°F/min to 32.4°F/min). 

Complications

The common complications of immersion cooling are shivering, cutaneous vasoconstriction, discomfort, and loss of monitoring capability. Shivering generates considerable heat through muscle metabolism. Cutaneous vasoconstriction impedes conductive heat loss. If significant shivering does occur, it can be reduced with benzodiazepines. A 2009 study found that high-dose diazepam facilitates core cooling during cold saline infusion in healthy volunteers. Subjects were randomized to receive high-dose (20-mg) or low-dose (10-mg) diazepam or placebo during cold saline infusion. Administration of high-dose diazepam decreased the shivering threshold without compromising respiratory or cardiovascular status. The use of phenothiazines such as chlorpromazine has been advocated for shivering in the past. They are currently discouraged because administration of these agents may impair heat loss through anticholinergic effects on sweat glands, contribute to hypotension via α-adrenergic blockade, lower the seizure threshold, and cause dystonic reactions. In addition, phenothiazines possess central dopamine-blocking effects, which may exacerbate symptoms of NMS. Benzodiazepines are also valuable if the patient is hyperthermic secondary to sympathomimetic agents such as cocaine. Patient monitoring is a problem under water. Electrodes can be used on the nonimmersed upper part of the shoulders, but electrocardiographic artifact often becomes a major problem during vigorous shivering. Immersion cooling is not recommended for patients with unstable cardiac rhythms or those at risk for the development of these rhythms. A significant change in cardiac rhythm might go undetected during the labor-intensive process of immersion cooling.

Patient access for resuscitative procedures is also a major problem when using this technique. Should ventricular fibrillation develop, the patient must be removed from the bath and dried before defibrillation. Invasive and diagnostic procedures (e.g., IV access and radiography) cannot be performed during the cooling period. Care must be taken to avoid displacement of IV lines during placement into and removal from the bath.

As body temperature drops, mental status will improve in many heatstroke victims. When awake, most people find ice water immersion difficult to tolerate. IV sedation may be required. Finally, this technique is labor-intensive. Several caregivers must be present throughout the process. The patient's head must be maintained out of the bath. If massage is used, one or more individuals will need to immerse their own hands in water to continuously massage the patient. Medications should be given intravenously, and constant attention to temperature and electrocardiogram monitors is also necessary. This cooling technique should be used only if adequate personnel are available.

**Whole-Body Ice Packing**

Packing a heatstroke victim in ice may enhance conductive heat loss without the attendant logistic problems caused by water immersion (Figs. 65.8 and 65.9). Constant attendance,
Figure 65.8 It is essential to rapidly lower the core temperature of a severely hyperthermic patient by instituting cooling techniques as soon as possible. Evaporative cooling (see text) is usually quite effective and technically easy. Note the fan for additional cooling. Many such patients require sedation, paralysis, and mechanical ventilation. An alternative aggressive method shown here, in a patient with a rectal temperature of 110°F, is to literally pack the patient in ice. As is often the case, this older patient did not survive.

Figure 65.9 A body bag or plastic sheets may keep water from flooding the floor when packing the patient in ice.

as required for skin moistening with evaporative cooling and as described for immersion cooling, may not be necessary with ice packing. Kielblock and associates demonstrated in a human study of mild, exercise-induced hyperthermia that whole-body ice packing cooled just as fast as evaporative cooling did (see Table 65.6).241

Procedure
For whole-body ice packing, undress the patient completely and then cover the extremities and torso with crushed ice. A fan blown over the patient may increase cooling. As with any cooling technique, monitor the patient’s temperature constantly with an electric thermometer and a long, flexible rectal probe. A large supply of crushed ice will be needed whenever this technique is used. Logistically, ice packing may be problematic. Whole-body ice packing can usually be performed on an ED stretcher without additional equipment. Ideally, the patient is placed in a container that facilitates contact of ice with the skin and prevents water from dripping onto the floor. A body bag makes an ideal device. Iced cooling may also be accomplished by placing the patient in a child’s lightweight plastic pool, available in toy stores. Lacking this equipment, plastic cloths or trash bags may be placed under the patient with the edges curled up to form a slinglike apparatus. As with immersion cooling, electrocardiographic monitoring can potentially be difficult because of shivering artifact and displacement of electrodes. If the patient is alert and cannot tolerate the ice packing, use IV sedation or restraint. Treat excessive shivering with benzodiazepines if needed. Once rectal temperature reaches 39°C (102.2°F), remove the ice and dry the patient off. Studies show cooling rates of 0.34°C/min (32.6°F/min) in heatstroke patients with whole-body ice packing.213,233

Strategic Ice Packs
Noakes suggested that strategic placement of ice packs over areas of the body where large blood vessels run close to the skin may be an effective cooling technique.241 Cooling in these areas occurs despite cutaneous vasoconstriction because of direct conductive heat loss from blood within the vessel and across the vessel wall, subcutaneous tissue, and skin to ice. The most common areas used for strategic ice packing are the anterior aspect of the neck (carotid and jugular vessels), the axilla (axillary artery and vein), and the groin (femoral vessels). There have been numerous reports of successful cooling using ice packs as primary or adjunctive therapy (see Table 65.6).241,242

Procedure
Place large plastic bags filled with crushed ice or an ice and water mixture in both axillae and over both femoral triangles. If the neck is used, place the packs laterally so that they do not compress the trachea or apply excessive weight over the carotid arteries. Do not pack the neck if the patient has carotid bruits or a history of cerebrovascular disease. Some sources advocate rubbing the body surface briskly with plastic bags containing ice after the body has been wet down with water. This is effective, provided that it is combined with evaporation therapy. Studies show ranges of cooling rates of 0.028°C/min to 0.087°C/min (32.0°F/min to 32.2°F/min) in heatstroke patients with strategic ice packs.241

Complications
Complications of strategic ice packing are limited to shivering and patient discomfort, as described previously for whole-body
ice packing. The ice packs are removed when rectal temperature reaches 39°C (102.2°F) to avoid an excessive drop in core temperature.

**External Versus Core Cooling**

All the external cooling techniques described previously are noninvasive and involve heat loss by evaporation or conduction across the skin as the primary cooling mechanism. With each of these techniques, central temperature will continue to drop even after the technique is discontinued and the skin is dried. This is due to a delay in establishment of an equilibrium between the cold skin and the core. The amount of “core afterdrop” can exceed 2°C (> 35.6°F). For this reason, cooling is discontinued when the core temperature reaches 39°C (102.2°F).

Because the sites of significant cell damage with heatstroke are centrally located (e.g., liver, kidney, heart), central cooling techniques are theoretically preferable to external techniques. Core cooling techniques studied in both animal and human models include iced gastric lavage, intravascular cooling, bladder lavage, and peritoneal lavage. Central venous cooling is effective in rapidly decreasing core temperatures. Studies conducted in healthy volunteers have demonstrated that reductions in core temperature vary according to the temperature of the infused fluid. Subjects receiving 30-minute infusions of fluid at 4°C (39.2°F) experienced decreases in core temperature of 2.5°C ± 0.4°C (36.5°F ± 32.7°F). Subjects receiving 30-minute infusions of fluid at 20°C (68°F) experienced decreases of 1.4°C (±0.2°C [34.5°F ± 32.4°F]). Clinical trials investigating this method showed that cooling via the respiratory tract had no significant impact on temperature changes when used exclusively but did demonstrate effectiveness as an adjunctive measure to other external cooling techniques. Cool air (10°C [50°F]) was administered via a hood or mask. Cooling via the respiratory tract has been studied in animals but not investigated clinically. Central cooling techniques are necessarily more invasive than external techniques and have the potential for more significant complications.

**Cold Gastric Lavage**

The stomach lies in close proximity to the liver, great vessels, kidneys, and heart. The gastric mucosa is not subject to the intense vasoconstriction observed on exposure of the skin to ice water. For these reasons, lavage of the stomach might be expected to be an effective central cooling method. Studies of cold gastric lavage in a canine model produced cooling rates five times greater than in controls exposed to ambient air at room temperature (0.15°C/min vs. 0.03°C/min). Human heatstroke victims have been cooled successfully with gastric lavage, but only in combination with external techniques. Cold gastric lavage seems to be best suited for use in patients with severe hyperthermia who are cooled at a slow rate with external techniques alone. The presence of an ET tube and passage of a large-bore gastric tube make rapid lavage without aspiration possible. This technique should be reserved for patients whose airway is protected by ET intubation and who do not have a contraindication to gastric tube placement.

**Procedure**

For cold gastric lavage, instill 10 mL/kg of iced tap water into the stomach as rapidly as possible (usually over a 30- to 60-second period). After a 30- to 60-second dwell time, remove the water by suction or gravity. Cooling will theoretically be faster if a high temperature gradient is maintained in the stomach. A faster lavage rate can be maintained if suction is used to withdraw the instilled fluid. A large container of ice temperature water maintained 1 to 1.5 m above the patient’s body will facilitate the instillation of fluid. Connect this container directly to the lavage tubing and ideally allow passage of water but not ice, which may occlude the tube. Because large volumes of water are needed, it is helpful if additional ice can be added to the container without interrupting the lavage. A large syringe can be used as an alternative to gravity instillation, but this is usually slower.

A simple system that accomplishes this procedure can be devised from equipment readily available in most EDs. Use a standard lavage setup (for use in drug overdoses) and a large-bore gastric tube. Cut the lavage bag open at the top to allow water and ice to be added. Suspend this bag above the patient’s body and connect it to the orogastric tube by Y tubing with clamps. Connect the other arm of the Y tubing to suction. Using the clamps, intermittently instill ice water by gravity and withdraw it by suction.

**Complications**

A major potential complication of cold gastric lavage is pulmonary aspiration. Use of a cuffed ET tube minimizes the incidence of this complication. Because of the large volume of water used and the frequent depression of airway reflexes seen with severe heatstroke, this technique should rarely be used in a patient who is not endotracheally intubated.

If tap water is used, water intoxication, hyponatremia, and other electrolyte disturbances are potential complications, particularly in pediatric or geriatric patients. Water is absorbed from the stomach and, with large-volume lavage, may pass the pylorus into the small intestine. In canine studies, large-volume gastric lavage with tap water did not cause electrolyte abnormalities. The actual incidence of these potential complications in human heatstroke has not been determined. Use of normal saline instead of tap water would eliminate this potential problem.

Theoretically, passage of cold water through the esophagus, located directly behind the heart, has the potential to induce cardiac dysrhythmias. Dysrhythmias have not been observed in canine studies or in case reports of human heatstroke victims cooled with this technique.

**Cold Peritoneal Lavage**

The surface area and blood flow of the peritoneum greatly exceed those of the stomach. Peritoneal lavage is expected to exchange heat much faster than possible with gastric lavage. Peritoneal lavage achieves some of the fastest cooling rates ever reported in large animal or human studies (see Table 65.6). A case report of cooling via cold peritoneal lavage for hyperthermia after the ingestion of ecstasy demonstrated rapid cooling. As with gastric lavage, this central cooling technique offers the advantage of directly cooling the core organs that are most susceptible to thermal damage. Unlike gastric lavage, ET intubation is not required. Peritoneal lavage is used extensively to treat hyperthermia under various conditions and typically decreases core temperatures 5°C/hr to 10°C/hr (41°F/hr to 50°F/hr).

Peritoneal lavage is a more invasive cooling technique. Because heat exchange is more efficient across the peritoneum,
smaller volumes of fluid can be used. Surgical placement of the lavage catheter is necessary. This cooling technique is relatively contraindicated by conditions that preclude placement of a lavage catheter (e.g., multiple abdominal surgical scars).

Peritoneal lavage is the most rapid central cooling technique. It can theoretically be combined with other techniques to speed cooling of heatstroke patients with refractory hyperthermia. As the most invasive cooling technique, it requires time, proper equipment, and surgical expertise to institute. Its use is probably best suited to situations in which heatstroke patients are not responding to external cooling and adequate equipment and personnel are readily available.

Procedure
To institute cooling by peritoneal lavage, immerse 2 to 8 L of sterile saline in an ice water bath to cool while the catheter is being placed. Place a standard peritoneal lavage catheter (as for diagnostic use in trauma patients) via any of the techniques described in Chapter 43. Standard contraindications apply. Use of a larger peritoneal dialysis catheter may speed instillation and withdrawal of fluid. Actual lavage volumes and rates have not been established, however. One approach is to instil and withdraw 500 to 1000 mL every 10 minutes until adequate cooling is achieved. Rectal temperature may be falsely low during lavage because of the presence of cold water around the rectum at the level of the rectal temperature probe. It may be preferable to monitor temperature in the tympanic membrane or esophagus when using this technique. Stop the lavage when core temperature reaches 39°C (102.2°F) to avoid excessive core temperature afterdrop.

Complications
The potential complications of peritoneal lavage cooling are primarily related to placement of the catheter and include bowel or bladder perforation and placement into the rectus sheath rather than the peritoneum.

Other Cooling Techniques
“Rewarming” techniques are used to minimize ongoing heat loss via the respiratory tract in hypothermic patients. Although high-frequency jet ventilation (HFJV) achieves core cooling in critically ill patients, efforts to use the respiratory tract to cool heatstroke victims have been unsuccessful. In a canine model of heatstroke, HFJV was shown to be a relatively ineffective cooling technique. Heat loss by convection (air transfer) is relatively inefficient when compared with the conductive heat loss mechanism used by other cooling techniques. The use of dry, hot air to maximize evaporative heat loss from the lungs might cause respiratory complications.

In human trials, ice water lavage of the bladder (300 mL of iced Ringer’s solution every 10 minutes) provided only minimal cooling at rates of 0.8°C/hr (±0.3°C/hr [33.4°F/hr ± 32.5°F/hr]). Iced water lavage of the rectum would theoretically provide faster cooling rates secondary to the increased surface area and better perfusion, but it has not been investigated in human trials.

Hemodialysis or partial cardiopulmonary bypass could theoretically be used to cool heatstroke patients. Before the availability of dantrolene in 1979, partial cardiopulmonary bypass was one of the treatments of MH. Drawbacks include the need for technical expertise and preparation time for the procedure. A 2005 case report described successful treatment of a heatstroke patient with multiple-organ failure refractory to conventional cooling techniques with cold hemodialysis initially at 30°C (86°F) and later at 35°C (95°F), followed by continuous hemodiafiltration with cold dialysate (35°C [95°F]) at a high flow rate of 18,000 mL/hr. Within 3 hours of starting this particular technique, the patient’s body temperature was below 38°C (100.4°F).

Cyclic lung lavage with cold perfluorochemicals is currently under investigation in animal models. Benefits include rapid cooling rates of 0.5°C/min (32.9°F/min) and minimally invasive nature in already mechanically ventilated subjects.

Intravascular cooling catheters have demonstrated efficiency as cooling devices. They circulate temperature-controlled sterile saline placed in the bladder or inferior vena cava. Although these devices have not been used in heatstroke patients, studies have found the cooling catheters to be very effective for neurologic conditions in both human and animal models.

Another promising cooling technique involves the use of a hypothermic retrograde jugular vein flush (HRJVF) for heatstroke. HRJVF has been studied only in animal models thus far. This technique involves the infusion of 4°C (39.2°F) isotonic sodium chloride solution through the external jugular vein (1.7 mL/100 g of body weight over a 5-minute period). Use of HRJVF was found to increase survival rates during heatstroke by attenuating cerebral oxidative stress, tissue ischemia or injury, systemic inflammation, and activated coagulation.

Pharmacologic agents have demonstrated merit as adjunctive agents in the management of hyperthermia. There are anecdotal reports of enhanced reduction in temperature with IV ketorolac. Cienki and colleagues demonstrated enhanced decreases in temperature with the administration of ketorolac, 30 mg intravenously. All patients received standard hyperthermia treatment (e.g., ice packs, iced lavage, circulating air). Patients were randomized to receive ketorolac versus saline. In the group receiving ketorolac, the average rectal temperature after 90 minutes was two times lower than in those receiving placebo saline (3.7°C vs. 1.6°C [38.7°F versus 34.9°F]).

Conclusion
Rapid cooling is the key step in the emergency management of heatstroke patients. Survival rates approach 90% when elevated temperatures are lowered in a timely fashion. Evaporative cooling appears to be the technique of choice. It combines the advantages of simplicity and noninvasiveness with the most rapid cooling rates achieved with any external technique. It is also logistically easier to institute, maintain, and monitor evaporative cooling measures than with any other cooling technique. If a patient is not cooling rapidly with evaporative cooling, other techniques can be added. Strategic ice packs can be used. If the patient is endotracheally intubated, gastric lavage can be instituted. If facilities and personnel are available, peritoneal lavage cooling can be used as a rapid central cooling technique. If muscle rigidity is present or MH is suspected, dantrolene sodium should be administered. In addition, the clinician should have a heightened index of suspicion for NMS and toxicity from sympathomimetic drugs. Regardless of the cause, a reasonable clinical goal is to reduce rectal temperature to 40°C (104°F) or below within 30 minutes of instituting therapy.

Immersion cooling is best limited to centers with the proper equipment and skilled medical personnel experienced in
managing hyperthermic patients. This method may also be effective in conditions in which electric power for evaporative cooling is unavailable (e.g., in wilderness settings where bodies of cool water are available nearby and the victim is far from more sophisticated medical care). Central venous cooling with iced saline is a promising technique for rapid cooling of patients with severe hyperthermia. Other cooling techniques require further study before a clear recommendation regarding their efficacy can be made.

ACKNOWLEDGMENTS

The authors wish to acknowledge and thank Terry VandenHoek and Pavitra Kotini-Shah for their contributions. We also wish to thank Dwight E. Helmrich, Scott A. Syverud, David Doezema, and David P. Sklar for their contributions and authorship in previous editions.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
60. Hilmo J, Naesheim T, Gilbert M: “Nobody is dead until warm and dead.” Prolonged resuscitation is warranted in arrested hypothermic victims also in remote areas—A retrospective study from northern Norway. Resuscitation 85:1204, 2014.


Bedside ultrasound has become an indispensable tool in the emergency department (ED). It has enabled physicians to make a rapid diagnosis at the bedside and formulate a plan of care. Perhaps most importantly, ultrasound has revolutionized procedures performed in the ED. Use of ultrasound gives the physician the advantage of viewing the anatomy and directly imaging the procedure while it is being performed. Procedures that had previously been performed “blindly” can now be performed with the added assurance of monitoring the procedure while it is in progress. This has resulted in greater safety in both common and uncommon procedures.\(^1\) \(^2\)

Ultrasound is now routinely applied to procedures in the ED ranging from the common, such as incision and drainage of an abscess, to the rare, such as drainage of a pericardial effusion (Videos 66.1-66.15). Use of ultrasound to facilitate each procedure is covered in detail in individual chapters. This chapter covers the basic principles of ultrasound that can be applied to any procedure. Ultrasound applications for specific procedures can be found in the appropriate procedure chapter.

**PHYSICS**

Ultrasound operates on the pulse echo principle. Electrical energy created by the ultrasound machine causes crystals in the tip of the transducer to vibrate, also known as the piezoelectric effect. This vibration emits high-frequency sound waves that travel into the body. The sound waves are reflected back to the transducer at varying intensities and speeds, depending on the nature of the object that they encounter. The ultrasound machine is able to interpret this information and plot an image on the screen.

Objects in the body that are liquid or water-like, such as a full bladder, reflect very few sound waves and allow most of the energy to pass through them. These objects are presented on-screen as black by ultrasound and are described as anechoic (i.e., without echoes) (Fig. 66.1). Conversely, objects that are dense and have very little water content, such as bones, reflect almost all the sound waves back. These objects are presented on-screen as white by ultrasound and are described as hyperechoic (i.e., producing a lot of echoes) (Fig. 66.2). Objects that lie between these two extremes present as varying shades of gray, depending on the water content of the object. For example, the liver contains a large amount of blood, a water-like substance, and is not completely liquid, appearing on-screen as dark gray (Fig. 66.3). An object that contains less water and is not completely solid would appear as a lighter shade of gray.

These properties of objects also account for two important artifacts. Acoustic shadowing is an artifact that is encountered when dealing with hyperechoic objects. These objects reflect almost all the sound waves back to the transducer. As a result, the ultrasound machine “senses” an absence of information deep to the hyperechoic object. The absence of information is represented by a strong, dark vertical line emanating deep to the object (Fig. 66.4). This type of shadowing is referred to as a clean shadow. This shadow can be frustrating to the sonographer when attempting to obtain the best image possible, classically, when imaging over the ribs. Alternatively, it can also be helpful in identifying hyperechoic objects, such as gallstones or foreign bodies.

In contrast to a clean shadow, the presence of air may create a phenomenon known as dirty shadowing (Fig. 66.5). Air causes the ultrasound beam to scatter and creates a hazy, gray appearance on the image. This can be an anticipated finding, such as when viewing bowel gas within the abdomen, or an indication of an abnormality, such as when viewing gas within subcutaneous tissue.

Acoustic enhancement, or an acoustic window, is the artifact created by an anechoic object. As noted earlier, sound waves pass through anechoic objects well and therefore lose less of their energy. This enables more ultrasonic energy to be available when the sound reaches the object on the other side of the fluid. This results in a brighter, clearer image immediately behind the fluid-filled object. As an example, a full bladder enables a clearer image of the pelvic organs (Fig. 66.6).

**INDICATIONS AND CONTRAINDICATIONS**

Although vascular access was one of the first uses of ultrasound for ED procedures, the list of procedures that can be facilitated by ultrasound is growing rapidly and continuously (Box 66.1). Even when a procedure cannot be directly observed with ultrasound, bedside ultrasound can frequently be used to diagnose the abnormality and to plan the approach for the procedure.

Ultrasound is a very safe modality for imaging at the bedside. It can be used in a variety of populations (pediatrics, pregnancy), without concern for excessive radiation exposure. The only absolute contraindication to using bedside ultrasound for procedural guidance is lack of training or experience in its use. Lack of adequate training or experience may result in an incorrect diagnosis and erroneous evaluation of the anatomy. This may result in harm and unnecessary complications in patients.

**EQUIPMENT**

The nature of the image obtained depends on the type of transducer used. In general, transducers fall into two categories: high frequency and low frequency, referring to the type of ultrasound waves generated by the transducer.

High-frequency sound waves do not penetrate very far into the tissue but provide excellent resolution. A linear or vascular transducer and an intracavitary transducer are examples of specific types of high-frequency transducers (Figs. 66.7 and 66.8). Examples of examinations that are best evaluated with a high-frequency transducer are central or peripheral vascular access, evaluation of the pleura for pneumothorax, and evaluation of soft tissue for an abscess or foreign body. These examinations rely on a high degree of resolution (Fig. 66.9).

Low-frequency sound waves penetrate farther into tissues and do not provide as much resolution as high frequency does.
Curvilinear, phased-array, and microconvex transducers are examples of specific types of low-frequency transducers (Figs. 66.10 and 66.11). They are distinguished from each other by the shape of the footprint of the transducer, the part that directly touches the patient, and by the layout of crystals in the tip of the transducer. Examples of the types of examinations that are best evaluated with a low-frequency transducer are evaluation of the pleural or peritoneal cavities for fluid drainage, evaluation of the bladder for aspiration or suprapubic catheter placement, and evaluation of the pericardium for pericardiocentesis. These examinations are not as dependent on resolution and typically require greater penetration (Fig. 66.12).
The growing use of ultrasound for procedures has prompted the development of echogenic needles and needle guidance systems. Both these advances are designed to improve the ease and accuracy of ultrasound-guided procedures, but this depends on the operator's familiarity with their properties and the correct use.

Additionally, many procedures require sterile conditions. Commercially designed sterile transducer covers are available and allow the operator to completely cover both the transducer and the cord. In the absence of these covers, the transducer can be placed inside a sterile glove. To further protect the field, a sterile drape can be wrapped around the cord.
CHAPTER 66  Ultrasound

1437

turned toward the patient's head, a longitudinal image is generated (Fig. 66.16). Objects on the right side of the screen are closer to the patient's feet and objects on the left side of the screen are closer to the patient's head. As with the transverse image, objects near the top of the screen are closer to the skin and objects at the bottom of the screen are deeper in the body.

Once the initial evaluation has been completed, make a decision whether to use ultrasound to directly guide the procedure or to simply “mark the spot” beforehand. Some procedures, such as paracentesis of large fluid collections, may not require direct guidance. In this case, ultrasound can be used to locate the most optimal puncture site, away from important structures. Once that site has been marked, proceed with the procedure in the usual fashion. For other procedures, such as central line placement, direct visualization throughout the entire procedure may be more important.

When direct visualization is desired, the orientation of the needle to the transducer must be considered. When the needle is introduced at the midpoint of the transducer in the transverse

---

Figure 66.12 Example of an image as seen with a low-frequency transducer. In this case the liver and kidney can be seen in the right upper quadrant.

Figure 66.13 Indicator on the transducer (arrow).

Figure 66.14 Indicator on-screen (arrow), which corresponds to the indicator on the side of the transducer.

Figure 66.15 Relationships in the transverse orientation. As the indicator points toward the patient's right, objects on the left side of the screen are toward the patient's right (toward the indicator). Objects on the right side of the screen are toward the patient's left (away from the indicator). As in all images, objects closer to the top of the image are closer to the surface, whereas objects at the bottom of the image are deeper in the body.

---

GENERAL APPROACH

Although each procedure has unique techniques, there are several general principles that apply when using ultrasound. Selection of the correct transducer is of the utmost importance for adequate visualization of the anatomy and pathology. This process is described in the preceding section, and in general the degree of resolution and depth of penetration should guide selection of the transducer.

Once the correct transducer has been selected, evaluate the area of interest. This step is key when evaluating the nature of the pathology. For example, understanding the size and position of a subcutaneous abscess and the surrounding anatomy is essential. Determining the correct orientation of objects on the screen can be challenging, and it is crucial for success with ultrasound-guided procedures. All transducers have a dot or mark on one side (Fig. 66.13). This mark corresponds to an indicator on the left-hand side of the screen (Fig. 66.14). When the indicator on the transducer is pointing toward the patient's right side, a transverse image will be generated (Fig. 66.15). Objects on the right side of the screen are closer to the patient's right side, near the on-screen indicator, and objects near the top of the screen are closer to the skin and objects near the bottom of the screen are deeper in the body. When the indicator is turned toward the patient's head, a longitudinal image is generated (Fig. 66.16). Objects on the right side of the screen are closer to the patient's feet and objects on the left side of the screen are closer to the patient's head. As with the transverse image, objects near the top of the screen are closer to the skin and objects at the bottom of the screen are deeper in the body.

---

Figure 66.12 Example of an image as seen with a low-frequency transducer. In this case the liver and kidney can be seen in the right upper quadrant.

Figure 66.13 Indicator on the transducer (arrow).

Figure 66.14 Indicator on-screen (arrow), which corresponds to the indicator on the side of the transducer.

Figure 66.15 Relationships in the transverse orientation. As the indicator points toward the patient's right, objects on the left side of the screen are toward the patient's right (toward the indicator). Objects on the right side of the screen are toward the patient's left (away from the indicator). As in all images, objects closer to the top of the image are closer to the surface, whereas objects at the bottom of the image are deeper in the body.
Figure 66.16 Relationships in the longitudinal, or sagittal, orientation. As the indicator on the transducer points toward the patient’s head, objects on the left side of the screen are toward the patient’s head. Objects on the right side of the screen are toward the patient’s feet. As in all images, objects closer to the top of the image are closer to the surface, whereas objects at the bottom of the image are deeper in the body.

Figure 66.17 Inserting a needle in the transverse approach. In this technique the needle is inserted at the midpoint of the transducer. This image is for demonstration purposes only and does not demonstrate sterile technique.

Figure 66.18 As the needle is advanced the transducer can be seen to transect only a small portion of the needle. The tip of the needle is now away from the transducer and will not appear on-screen.

Figure 66.19 Real-time image of the needle as it approaches the vein. The needle (arrow) is seen as a hyperechoic object with strong shadowing extending behind it. The portion of the needle seen will correspond to the portion of the needle that is underlying the transducer, as shown in Fig. 66.18.

Figure 66.20 Inserting a needle in the longitudinal approach. In this technique the needle is inserted at the end of the transducer. This image is for demonstration purposes only and does not demonstrate sterile technique.

Figure 66.21 As the needle is advanced the transducer can be seen to transect only a small portion of the needle. The tip of the needle is now away from the transducer and will not appear on-screen.

approach, the tip of the needle may be difficult to visualize (Fig. 66.17). In this orientation the operator may have difficulty following the tip of the needle because only a small portion of the needle intersects the ultrasound beam at any given time (Figs. 66.18 and 66.19). Conversely, when the needle is introduced from either end of the transducer, the needle can be visualized in its entirety (Figs. 66.20 and 66.21). Slight movement of the transducer may result in losing the view of the needle. Consider each potential approach in the context of the particular procedure.

The oblique approach is a combination of the transverse and longitudinal approaches. Primarily, this has been described for central venous access and can also be applied to other procedures. In this method, place the transducer on an oblique axis to the target object (Fig. 66.22). This allows an oblique view of the target object while enabling the operator to introduce the needle along the long axis of the transducer (Fig. 66.23).
In general, the incidence of complications should be reduced with the use of ultrasound, but complications may occur even when performed by an experienced sonographer. Complications may be related to one of several factors. To maximize success, obtain an optimal image of the structure in question. Evaluate the anatomic relationships to ensure that the target object (e.g., the femoral vein versus the femoral artery) is correctly identified and pursued. Once the procedure begins, pay constant attention to the orientation and location of the needle. Errors, including inadvertent arterial puncture, can occur when the position of the tip of the needle is not closely followed.4,5 Techniques to minimize these errors are addressed in the individual chapters.

FAST, E-FAST, AND RUSH EXAMS

No discussion about bedside ultrasound would be complete without a reference to the FAST, E-FAST, and RUSH exams.

The FAST (Focused Assessment with Sonography for Trauma) exam was one of the first applications of ultrasound at the bedside, performed by non-radiology physicians. The FAST exam was first developed in the 1970s outside of the United States and spread to the United States in the 1980s, with rapid growth thereafter.6,7 The FAST exam was first developed as an alternative to diagnostic peritoneal lavage (DPL) in patients with blunt trauma. In contrast to DPL, FAST is noninvasive, repeatable, and can be performed on any patient, from infants to pregnant women. Due to these advantages, FAST has not only largely replaced DPL for evaluation of hemorrhage following blunt trauma, but has been adapted for use in other scenarios, such as penetrating trauma and nontrauma situations (such as in the case of a ruptured ectopic pregnancy). The E-FAST (Extended Focused Assessment with Sonography for Trauma) exam builds upon the traditional thoracoabdominal views of the FAST to include evaluation of the pleura and lung.

The effectiveness of the FAST and E-FAST exams as diagnostic tools varies depending on the scenario in which they are used. The expertise and experience of the examiner plays a significant role and although the basic exam appears to be easily mastered, examiners with more experience are more capable of diagnosing small free-fluid collections.8 FAST appears to be most sensitive when evaluating hypotensive blunt trauma patients.9,10 Multiple studies have determined that the sensitivity for detecting free fluid in these patients approaches 100%.11–13 In patients who are normotensive following blunt trauma, the sensitivity and specificity for FAST is more questionable, with most studies finding between 42% to 86% and 98% to 99%, respectively.14–16 The FAST exam in penetrating trauma can also be helpful, although the sensitivity is not as high as in blunt trauma patients.17 In an unstable penetrating trauma patient, a quick FAST exam to evaluate the pericardium can yield important information to help aid the operating room course.18 Ultrasound for pneumothorax has shown to be extremely sensitive, particularly in comparison to traditional supine chest radiography.19 Supine x-rays have sensitivities ranging from 47% to 75%, even with moderately sized pneumothoraces.20,21 Ultrasound conveys the advantage of quickly identifying even a small pneumothorax. Ultrasound is also reliable for identifying pleural fluid.22 Identifying a
evaluation of the abdomen. A curvilinear or phased-array transducer can be used, depending on availability and examiner preference. For evaluation of the pleura, a high-frequency transducer (6 to 13 mHz) is optimal for maximum resolution, although a low-frequency transducer will suffice if no high-frequency transducer is available. The pericardium can be assessed for effusion from either the subxiphoid region of the chest or from the left anterior chest wall (Fig. 66.26). In the supine patient, a pneumothorax will rise and be best seen on the anterior-medial aspect of the chest wall. A more detailed description of components of this exam can be found within this text:

- Evaluating the dependent portions of the abdomen in trauma is similar to evaluating dependent portions of the abdomen for ascites and paracentesis; see Chapter 43.
- Evaluating the pericardium for effusion following trauma is similar to evaluating effusions in anticipation of pericardiocentesis; see Chapter 16.
- Evaluating the pleura for evidence of a pneumothorax is covered in Chapter 10.

The RUSH (Rapid Ultrasound in Shock) exam was developed as a means to quickly assess a hypotensive patient at the bedside, utilizing ultrasound. The goals of the RUSH exam are to attempt to identify treatable causes of shock, such as pericardial tamponade, and also to attempt to classify the type of shock present. Distinguishing between cardiogenic, obstructive, distributive, and hypovolemic shock is important, as the treatment strategies can vary widely. For example, determining that cardiogenic shock is present early in the evaluation of a hypotensive patient can assist the physician in narrowing the

hemothorax can be difficult in the supine patient, as fluid can layer on chest x-ray, making rapid diagnosis challenging. Ultrasound at the costophrenic angles can quickly identify as little as 10 cc of fluid. The basis of the FAST exam relies on the principle that intraperitoneal free fluid will settle into dependent areas within the peritoneum when the patient is supine. These dependent areas form the foundation of a basic FAST exam. The right and left upper quadrants, as well as the pelvis, are evaluated for free fluid. The pericardium is easily viewed from either the subxiphoid region or from the anterior chest wall, identifying pericardial effusions. This view is helpful in determining if cardiac motion is present, particularly when deciding whether to continue resuscitation. Quick views of the pleura from the anterior chest and the costophrenic angles can be added to evaluate for pneumothorax and hemothorax.

A full description of the FAST and E-FAST exams is outside of the scope of this text, but the dependent areas of the abdomen should be evaluated for the presence of free fluid, which will typically appear anechoic or black (Figs. 66.24 and 66.25). A low-frequency transducer (2 to 5 mHz) is usually optimal for evaluation of the abdomen. A curvilinear or phased-array transducer can be used, depending on availability and examiner preference. For evaluation of the pleura, a high-frequency transducer (6 to 13 mHz) is optimal for maximum resolution, although a low-frequency transducer will suffice if no high-frequency transducer is available. The pericardium can be assessed for effusion from either the subxiphoid region of the chest or from the left anterior chest wall (Fig. 66.26). In the supine patient, a pneumothorax will rise and be best seen on the anterior-medial aspect of the chest wall. A more detailed description of components of this exam can be found within this text:

- Evaluating the dependent portions of the abdomen in trauma is similar to evaluating dependent portions of the abdomen for ascites and paracentesis; see Chapter 43.
- Evaluating the pericardium for effusion following trauma is similar to evaluating effusions in anticipation of pericardiocentesis; see Chapter 16.
- Evaluating the pleura for evidence of a pneumothorax is covered in Chapter 10.

The RUSH (Rapid Ultrasound in Shock) exam was developed as a means to quickly assess a hypotensive patient at the bedside, utilizing ultrasound. The goals of the RUSH exam are to attempt to identify treatable causes of shock, such as pericardial tamponade, and also to attempt to classify the type of shock present. Distinguishing between cardiogenic, obstructive, distributive, and hypovolemic shock is important, as the treatment strategies can vary widely. For example, determining that cardiogenic shock is present early in the evaluation of a hypotensive patient can assist the physician in narrowing the

<table>
<thead>
<tr>
<th>Pump</th>
<th>Tank</th>
<th>Pipes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart</td>
<td>Inferior vena cava</td>
<td>Aorta</td>
</tr>
<tr>
<td>FAST exam</td>
<td>Pleural/lungs</td>
<td>Lower extremity veins</td>
</tr>
</tbody>
</table>

BOX 66.2 Components of the RUSH Exam

FAST, Focused Assessment with Sonography for Trauma; RUSH, Rapid Ultrasound in Shock.
### BOX 66.3 Findings Sought in Each Component of the RUSH Exam

<table>
<thead>
<tr>
<th>Pump</th>
<th>Tank</th>
<th>Pipes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Estimate contractility of the heart to evaluate for cardiogenic shock and to determine ability to tolerate volume resuscitation. Identify the presence of pericardial effusion and tamponade.</td>
<td>Determine the size and collapsibility of the inferior vena cava as a marker of volume status. Assess for the presence of free fluid in the peritoneum to indicate hypovolemic shock.</td>
<td>Evaluate the thoracic and abdominal aorta for the presence of either an aneurysm (AAA) or dissection. Study the deep veins of the lower extremities to identify thrombus (DVT) when PE is a consideration.</td>
</tr>
<tr>
<td>Evaluate for signs of right heart strain to indicate possible PE.</td>
<td>Identify the presence of pneumothorax/tension pneumothorax. Evaluate the lung parenchyma for evidence of pulmonary edema associated with cardiogenic shock. Assess the costophrenic angles for evidence of pleural effusion/hemothorax.</td>
<td></td>
</tr>
</tbody>
</table>

AAA, Abdominal aortic aneurysm; DVT, deep vein thrombosis; PE, pulmonary embolus; RUSH, Rapid Ultrasound in Shock.

### BOX 66.4 Sonographic Views Used to Evaluate the Components of the RUSH Exam

<table>
<thead>
<tr>
<th>Pump</th>
<th>Tank</th>
<th>Pipes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parasternal long axis</td>
<td>IVC near entry into the right atrium</td>
<td>Thoracic aorta: parasternal long axis</td>
</tr>
<tr>
<td>Parasternal short axis</td>
<td>FAST: RUQ</td>
<td>Thoracic aorta: apical</td>
</tr>
<tr>
<td>Apical</td>
<td>FAST: LUQ</td>
<td>Abdominal aorta: proximal through bifurcation at umbilicus</td>
</tr>
<tr>
<td>Subxiphoid</td>
<td>FAST: pelvis</td>
<td>DVT: common femoral vein, superficial femoral vein, and popliteal veins</td>
</tr>
<tr>
<td></td>
<td>Anterior pleura for sliding</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Anterior and lateral chest wall for signs of pulmonary edema</td>
<td></td>
</tr>
</tbody>
</table>

DVT, Deep vein thrombosis; FAST, Focused Assessment with Sonography for Trauma; IVC, inferior vena cava; LUQ, left upper quadrant; RUQ, right upper quadrant; RUSH, Rapid Ultrasound in Shock.

differential diagnosis and in avoiding unnecessary (and potentially harmful) initial treatments such as large fluid boluses. The RUSH exam is divided into components: the “pump”, the “tank”, and the “pipes” and should be evaluated in a step-wise fashion (Box 66.2). Each component of the RUSH exam should be evaluated for specific findings to guide management (Box 66.3). A detailed overview of the technique is outside of the scope of this text. Box 66.4 lists the specific views that can be used to evaluate each component.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


Bedside Laboratory and Microbiologic Procedures
Anthony J. Dean and David C. Lee

ASSESSMENT OF URINE

Obtaining a Urine Specimen
Several methods are available for obtaining a urine specimen. They can be found in Table 67.1 and are listed in order of increasingly precise collection techniques. They come at the cost of increasing difficulty, patient discomfort, or both.

General Considerations Regarding Urine Collection
The advantages and disadvantages of each of the techniques listed in Table 67.1 can be determined only by the purpose of the urine test and the clinical context. The clinical context influences interpretation of the results. For the great majority of clinical scenarios, the basic dichotomy is between specimens obtained for infectious versus noninfectious reasons. With the exception of testing for red blood cells (RBCs) and white blood cells (WBCs), most of the noninfectious tests (e.g., ketones, glucose, bilirubin, protein) are not affected by the collection method. Urine specimens collected to diagnose infection can be contaminated in a number of ways, and the clinical scenario intricately influences the choice and interpretation of tests.

Urinary tract infections (UTIs) are either symptomatic or asymptomatic, and the symptoms determine which collection method is required (Fig. 67.1). Symptomatic female patients without potential complications who present for the first time require no further urine testing, and should be treated empirically based on local patterns of susceptibility for urinary pathogens. However, in symptomatic patients who have failed a course of therapy or have potential complicating factors, testing is recommended and extremely low levels of bacteriuria (10^2 colony-forming units [CFUs]/mL) and pyuria are of clinical significance.1,2 This may be obscured in some clinicians' minds by an alternative, more widely promulgated fact: In asymptomatic patients the threshold for “significant bacteriuria” is 1000-fold higher at greater than 10^5 CFUs/mL.3,4 Symptomatic patients constitute a clinically distinct group who require a urine test and the clinical context. The clinical context influences interpretation of the results. For the great majority of clinical scenarios: pregnant women and all patients scheduled for urologic surgery.14,15 If only a urine culture is to be performed, some would argue that any spontaneously voided specimen would suffice because the diagnosis of asymptomatic bacteriuria depends on 10^2 or more CFUs of a single pathogen per milliliter of urine. With such criteria, contaminants are usually easily identified.9 Because performing cultures on all such patients is prohibitively costly, dipstick testing, urinalysis (UA), or both are commonly used for screening.16,17 With these tests, contamination by bacteria, leukocytes, or erythrocytes results in diagnostic confusion. The advantages of a less-contaminated specimen are worth the minimal, extra effort of asking the patient to provide an MSCC specimen. The approach in Fig. 67.1 covers the vast majority of situations. A few circumstances and techniques deserve special mention.

Bladder Percussion and the Midstream Specimen in Infants
The emergency clinician is familiar with how frequently a urine stream is generated in infants confronted by the alarming emergency department (ED) environment and a cold stethoscope. Rather than wasting a potentially perfect MSCC specimen on a laboratory coat with an ensuing delay in obtaining urine, the clinician can exploit the situation by approaching an infant with an open sterile urine container in hand in case
the urine stream is spontaneously forthcoming. The process is facilitated by the application of cold povidone-iodine to the genitalia. Such an approach has been shown to generate a urine sample in a median time of 10 minutes. This is less than the typical time needed for straight catheterization or suprapubic aspiration (SPA), and it can be performed concomitantly with the history and physical examination and thereby circumvent an invasive procedure. If the urine specimen is not immediately forthcoming, a parent can be equipped with a sterile container and be instructed on collection of an ensuing specimen to free up ED staff for other tasks. Two techniques to actively induce voiding in infants have been described. The first, which is useful in newborns, exploits the Perez reflex. After cleansing the genitalia, hold the infant in one hand while stroking the paraspinal muscles in a cephalad to caudal direction. This causes extension of the back and flexion of the hips and induces micturition in less than 5 minutes in most cases. The second technique is known as “bladder tapping.” After urethral cleansing, if there is still no urine, use two fingers to tap on the suprapubic area at a rate of approximately once per second for a full minute, followed by a minute’s rest. Repeat the cycle until urine is produced. The mean time before the production of a urine sample is approximately 5 minutes. This technique, though not practicable for the staff in a busy ED, can provide an infant’s parents with a task that invests them in the clinical process. This clinical pearl may expeditiously furnish a specimen with significantly less investment of staff time than required for more invasive techniques. A combined method using finger-tap for 30 seconds alternating with lumbar stimulation with a light circular motion for 30 seconds after oral hydration achieved success in 86% of infants younger than 30 days old in less than 2 minutes.

**Bag Collection in Non-Toilet-Trained Children**

The incidence of unsuspected UTI in a febrile neonate or infant is approximately 5%. Similar rates are found in asymptomatic children less than 5 years of age who present to the ED with nonspecific symptoms of acute illness. A true UTI in an infant or child requires subsequent evaluation for urinary tract pathology, and the disease may produce significant morbidity (e.g., hypertension, renal disease). One must be certain of the presence or absence of infection in this subgroup. The costs of overdiagnosis with a false-positive UA are significant and include bacterial resistance that leads to the use of increasingly expensive antibiotic medications. Numerous studies have demonstrated the dissility of urine specimens obtained for culture from a collection bag stuck to an infant’s perineum. Bag specimens may be more sensitive than catheter specimens when used for UA or microscopy to identify infection in children at low or moderate risk for UTI. In this group it is acceptable to perform screening UA, microscopy, or both on a bag specimen. If negative, UTI is ruled out.

### Table 67.1: Urine Collection Methods Listed in Order of Increasing Precision

<table>
<thead>
<tr>
<th>METHOD</th>
<th>DESCRIPTION AND COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random voided</td>
<td>Any specimen provided by the patient.</td>
</tr>
<tr>
<td>Midstream voided</td>
<td>No skin preparation, container placed in the urinary stream 2–3 sec after initiation of micturition.</td>
</tr>
<tr>
<td>Clean catch</td>
<td>Same as above, plus antiseptic cleansing of the urethral area. This is performed by retraction of the prepuce in males and the labia in females and then cleansing the meatus in an anterior-to-posterior direction. Use three swabs soaked in povidone-iodine (or some other antiseptic solution). For female patients who are physically capable, the ideal position is sitting astride a toilet, facing backward. This helps separate the labia and position the cup for collection of the specimen.</td>
</tr>
<tr>
<td>Midstream clean catch</td>
<td>Cleansing as for clean catch, with midstream collection as for midstream voided.</td>
</tr>
<tr>
<td>Catheterized</td>
<td>Obtained from a newly placed catheter after cleansing of the meatus.</td>
</tr>
<tr>
<td>Suprapubic aspiration</td>
<td>See Chapter 55.</td>
</tr>
</tbody>
</table>

**Figure 67.1** Algorithm for deciding method of obtaining urine specimen for evaluation of possible urinary tract infection. For initial presentation with symptoms in otherwise healthy females: no urine testing indicated; see text. MSCC, Midstream clean catch; SPA, suprapubic aspiration; UTI, urinary tract infection.
If positive (leukocyte esterase or nitrite present, more than 5 WBCs/high-power field [HPF] on spun urine, or bacteria on an unspun Gram-stained specimen), it is followed by catheterization and culture, with treatment usually pending the results of culture.29,30,31 If a urine specimen is needed solely for chemical analysis (e.g., glucose, ketones, specific gravity), a bag specimen will suffice.

**Urine Specimens From Patients With Chronic Urinary Drainage Systems**

Urine obtained from any part of a chronic urinary drainage system is highly inaccurate for bacteriologic purposes. If UTI is suspected, insert a new catheter and obtain a fresh bladder urine specimen.32 A small study advocating replacement of a chronically applied Texas sheath catheter with a fresh one was performed on patients who did not have symptoms of UTI.33 Such a method might be sufficiently accurate for screening asymptomatic patients, but chronic asymptomatic bacteriuria is very common in such patients, and treatment is not recommended.34 In most cases, a Foley catheter should be used to obtain urine from patients with sheath catheters who have signs or symptoms of acute UTI and are unable to provide an MSCC specimen.

**Catheterization and SPA**

The low levels of bacteriuria found in 2% to 8% of patients after straight catheterization are generally below the threshold that defines the presence of UTI.35 Catheterization can cause minor local injury, as reflected by low-level hematuria in 15% of patients.33,36 SPA continues to be advocated by some for neonates in cases in which accurate diagnosis is essential and the risk for infection must be minimized. Both SPA (see Chapter 55) and catheterization have an approximately 25% failure rate as a result of an empty bladder.37 This problem and associated complications can be avoided by performing bedside ultrasound before the procedure.37-39 SPA may sparsely lower leukocyte or bacterial colony counts because of the necessity of filling the bladder before performing the procedure.40 A study in infants demonstrated that the discomfort associated with SPA is greater than that with catheterization.41 Surprisingly, however, older men who underwent both catheterization and SPA strongly preferred SPA.3

**Urine Dipstick**

Urine dipstick tests are available to test 10 separate parameters. The unassuming appearance and commonplace use of the urine dipstick might lead one to mistakenly underestimate its technical sophistication. Each colored square on a urine dipstick involves a biochemically complex assay, and therefore it is essential to meticulously follow the manufacturer’s instructions for storage and use.42 Even with optimal storage and testing conditions, the false-negative and false-positive rates of these tests are problematic. In addition, most of the tests are susceptible to interference from a variety of substances (Table 67.2).

**Method**

Test urine specimens as soon as possible after they are collected. If the urine has been standing, stir or shake the specimen well because cells sink rapidly in a container. Immerse the test strip completely for 1 second or less. Draw the edge of the strip along the rim of the specimen container and lightly tap it to remove excess urine, thus avoiding mixing the reagents between different test patches. Next, hold the strip horizontally or place it on a clean gauze pad until the recommended time has elapsed. Most strips are designed so that all the test results can be read together after 1 to 2 minutes (Fig. 67.2).

**Interpretation**

**Glucose**

The urine glucose test is normally negative. Urine glucose testing has limited usefulness in quantitative testing because the serum glucose level at which spillage occurs varies (although in most patients it starts at between 180 and 200 mg/dL).43 Changes in urine glucose lag behind changes in blood glucose by approximately half the interval between voids.43 Glycosuria in the absence of hyperglycemia suggests renal tubular dysfunction. Glycosuria may occur in hypothermic patients in the absence of hyperglycemia and indeed may actually occur with hypoglycemia in such patients.45

**Ketones**

Ketones are found in the urine of patients with starvation, inadequate carbohydrate intake, diabetic and alcoholic ketoacidosis, isopropyl alcohol poisoning, or glycogen storage disease. Tests for urine ketones are 5 to 10 times more sensitive to acetoacetate than to acetone, similar to the serum tests. Dipsticks do not detect 5-hydroxybutyrate, which accounts for 80% to 95% of the three ketone bodies and is the predominant form in the setting of ketoacidosis. Urine ketone testing is significantly more sensitive than serum ketone testing.46 There is generally no need to obtain serum acetones to diagnose or manage diabetic ketoacidosis when urine ketone monitoring is coupled with blood gas and anion gap analysis.

**Leukocyte Esterase**

This portion of the dipstick test is designed to detect enzymes from the azurophilic granules in neutrophils. Normally the test is negative. Studies report a wide and clinically important range of thresholds for the sensitivity of dipstick testing, from 10 to 100 WBCs/µL urine.46,47 Studies suggest that the test is between 50% and 96% sensitive in detecting infection.47,48 Its specificity for the presence of WBCs is between 91% and 99%. The most common cause of a false-positive leukocyte test is vaginal contamination.

**Nitrites**

Normally, urine does not contain nitrites. Nitrites are specific (≈ 95%), but not sensitive (≈ 45%) indicators of UTI.14,49,50 Urinary nitrates are converted to nitrites most strongly by enteric coliform bacteria, thus explaining the nitrite test’s 90% sensitivity in detecting UTI caused by *Escherichia coli*. *Enterococcus*, a moderately frequent urinary pathogen, *Pseudomonas* species, and *Acinetobacter* lack the reductase enzyme and are not detected by nitrite testing. False-negative results also occur because of the lack of dietary nitrate, frequent voiding, and diuresis. Early morning–voided specimens are ideal because they allow time for conversion of nitrate to nitrite, but they are rarely available in the ED. If possible, a specimen obtained more than 4 hours after the last voiding is preferred.

**Protein**

Proteins with a molecular weight below 50,000 to 60,000 daltons can pass through the glomerulus to be reabsorbed in the proximal tubule. Normal passage of protein in urine is less
than 150 mg/24 hours, or approximately 10 mg/dL of urine. Approximately 10% to 33% of urinary protein is albumin, 33% is Tamm-Horsfall glycoprotein (secreted by renal tubular cells), and the balance is made up of a variety of immuno-globulins and other proteins. Proteinuria is a finding noted in approximately 5% of routine urine screens in men.\(^5\) This may represent a normal variant as 3% to 5% of healthy adults have postural proteinuria (proteinuria when standing but not when recumbent).\(^4\) Proteinuria is rarely clinically significant unless 3+ or greater is seen on the dipstick, although lower levels are still indicative of some degree of renal dysfunction (see later). The dipstick detects negatively charged proteins more strongly than positively charged ones; it is therefore most sensitive to albumin. A study of ED patients with severe acute hypertension identified renal dysfunction, defined as elevated serum creatinine, with 100% sensitivity when using urine dipstick detection of 1+ proteinuria or hematuria.\(^52\) Along similar lines, in patients being considered for radiographic contrast-enhanced studies who do not have a serum creatinine measurement, a negative dipstick test for protein or blood combined with the absence of postural proteinuria is associated with an almost 100% negative predictive value for renal dysfunction.\(^53\)

### TABLE 67.2 Overview of Urine Dipstick Tests

<table>
<thead>
<tr>
<th>SOURCE OF ERROR AND ARTIFACT</th>
<th>COMMENTS</th>
</tr>
</thead>
</table>
| **Glucose**                 | False positive with peroxide, hypochlorite, ketonuria, levodopa, and the dipstick exposed to air | Hypothermia may cause glycosuria despite hypoglycemia  
Glycosuria without hyperglycemia suggests renal tubular dysfunction |
| False negative with ascorbate, ketones, uric acid, and high specific gravity | |
| **Ketones**                 | False positive with ascorbate, low pH urine, high specific gravity, levodopa, valproate, phenazopyridine, N-acetylcysteine, high-protein diet, phenylketonuria, phthalein compounds | Very susceptible to deterioration with humidity and delay in analysis, which can cause false-negative results |
| False negative with high specific gravity, frequent urination, ascorbate, high urine pH, and urine standing in the specimen cup >2 hr | |
| **Nitrites**                | False positive with phenazopyridine  
False-negative with high specific gravity, frequent urination, ascorbate, high urine pH, and urine standing in the specimen cup >2 hr | 75% false-negative rate when exposed to air for 15 days  
Does not detect reductase-negative bacteria |
| **Protein**                 | False positive with pH >7 and chlorhexidine  
False negative with low pH, very dilute urine | Only reliable for albumin (glomerular proteinuria)  
Does not detect Bence-Jones protein  
Positive with pyuria, rarely with hematuria |
| **Blood**                   | False positive with povidone-iodine, certain (peroxidase-producing) bacteria, hypochlorite  
False negative with high specific gravity and high concentrations of urinary nitrates, ascorbate, or captopril | Positive test with speckles or dots implies nonhemolyzed blood  
Positive test with a diffuse pattern implies hemolyzed red blood cells or high levels of myoglobin |
| **Bilirubin**               | False positive with iodine, stool contamination, chlorpromazine, mafenamic acid  
False negative after prolonged standing | Hard to read with agents causing marked urine discoloration |
| **Urobilinogen**            | False positive with phenazopyridine, sulfisoxazole, sulfonamides, porphyrin, methylldopa, procaine, aminosalicylic acid, 5-hydroxyindolacetic acid  
False negative with sulfisoxazole and phenazopyridine | Use a fresh specimen: rapidly broken down by light and in acidic urine |
| **Leukocyte Esterase**      | False positive with vaginal contamination, oxidizing agents, eosinophils in urine, \textit{Trichomonas}  
False negative with high glucose, ketones, protein (especially albumin), pH, and specific gravity, as well as with cephalaxin, tetracycline, oxalates, ascorbic acid, neutropenia | Sterile pyuria seen with tuberculosis, nephrolithiasis, interstitial nephritis |
| **pH**                      | Urea-splitting bacteria elevate pH  
Runoff from the protein strip can falsely lower pH | Use a fresh specimen: standing raises pH by loss of CO$_2$ |
| **Specific Gravity**        | Overestimates specific gravity with low pH, ketoadidosis, and protein  
Underestimates specific gravity with glucose, urea, or pH >7 | Not reliable at specific gravity >1.025  
Elevated with use of dextran, intravenous contrast material, proteinuria |
with a molecular weight of 32,000 daltons, which easily passes through the glomerulus. Myoglobin has a molecular weight of 17,000 daltons, which also allows easy glomerular passage, but the dipstick has been shown to have a sensitivity of only 14% with heat-induced rhabdomyolysis, as reflected by serum creatine phosphokinase levels of up to 1000 U/L.

Guidelines for distinguishing hematuria, hemoglobinuria, and myoglobinuria are outlined in Table 67.3. In asymptomatic men older than 50 years, significant disease can be signaled by intermittent hematuria, thus mandating follow-up of patients with this incidental finding. Dipsticks are vitiated by humidity and air, which can cause false-negative results after improper storage. Because RBCs may lyse rapidly, delays in performing UA may misleadingly suggest myoglobinuria or hemoglobinuria. Microscopy or dipstick testing of a freshly obtained specimen can clarify this issue. Conversely, high specific gravity or low pH can inhibit lysis of erythrocytes, which is necessary for the dipstick chemical reaction to occur, thus causing false-negative results. A study reproducing clinical conditions has demonstrated that povidone-iodine does not cause false-positive results on the dipstick. Iatrogenically caused trace positive results may occur after catheterization in 15% of cases.

Urine Bilirubin
Urine bilirubin represents the filtered, soluble, conjugated form of bilirubin. Unconjugated bilirubin is bound to protein and does not pass through the glomerulus. Bilirubinuria is therefore due to intrahepatic or extrahepatic cholestasis. Bilirubinuria will be detected significantly earlier than clinical jaundice.
Urinary bilirubin excretion is enhanced by alkalosis. A fresh sample of urine should be tested because bilirubin glucuronide is hydrolyzed when exposed to light. Ascorbic acid and high levels of urinary nitrites decrease the sensitivity of the test to bilirubin.

Urobilinogen
In a healthy person, conjugated bilirubin is excreted in bile. In the colon it is broken down into a number of compounds, including urobilinogen. Most of these compounds are excreted in stool, which is the source of its characteristic color. A small amount of urobilinogen is absorbed from the colon, and if it is not taken up on the first pass through the liver, it enters the circulation. Ultimately, some of this urobilinogen may enter the urine, so it is normal to have zero to moderate levels of urinary urobilinogen on dipstick testing. Most diseases that cause hepatocyte dysfunction (hepatitis, cirrhosis, passive liver congestion, etc.) increase urinary urobilinogen excretion by impairing hepatic uptake of urobilinogen. As a qualitative test with a wide range of normal values it is rarely helpful, but in evaluating a patient with jaundice it can have diagnostic significance (Table 67.4).

pH
The average daily excretion of 50 to 100 mmol of H⁺ in urine gives rise to a typical urine pH of approximately 6 with a range from 4.5 to 8. Dietary protein lowers urinary pH, whereas fruit (especially citrus) and vegetables tend to raise it. The significance
of pH testing is in the assessment of normal renal function. In most states of alkalosis and acidosis, healthy kidneys maintain homeostasis by conserving or excreting H⁺. Failure to do so suggests renal disease, especially renal tubular acidosis. An exception is the paradoxical aciduria of hypokalemie alkalosis secondary to volume contraction, hypercorticism, or diuretics, where the highest priority of the renal tubule is to conserve sodium. pH is elevated by the action of urea-splitting bacteria, especially Proteus species. This can occur with stasis of urine, either in the bladder or in specimen cups. A persistently alkaline urine is seen in patients with struvite (triple phosphate) urolithiasis.

### Specific Gravity

The dipstick test for urine specific gravity assays for the primary urinary cations, sodium, and glucose, is therefore not measured. Artifically low specific gravity readings are obtained with alkaline urine, whereas acidic urine and albumin falsely elevate the specific gravity reading. Consequently, some investigators believe that these strips on the dipstick test are of marginal clinical utility. Other clinical indicators of a patient’s hydration status are probably more reliable. If necessary, a refractive specific gravitometer or a hygrometer should be used.

### Microscopic UA

Microscopic UA is performed to identify cells, bacteria, and other microbes, as well as formed elements such as casts and crystals (Fig. 67.3). The following discussion focuses on the findings of significance in diagnosing UTI: WBCs, bacteria, and RBCs casts. The presence of WBCs with bacteria distinguishes infection from colonization (bacteriuria without pyuria). Some authorities state that significant infection without pyuria occurs in less than 5% of cases, thus making pyuria alone a sensitive marker of infection. Other studies do not support reliance on pyuria as an indicator of infection. The presence of WBC casts distinguishes pyelonephritis (“upper UTI”) from cystitis (“lower UTI”).

Microscopic UA is performed by one of five methods. Traditionally, the most common technique has been examination of unstained centrifuged urine. It has the advantage of concentrating formed elements that might otherwise be missed. Its disadvantage is that the presence and quantity of elements in a specimen will depend on many uncontrolled factors: the volume of the specimen, the duration and speed of centrifugation, the fragility of the formed elements, the volume of the “drop” in which the pellet is resuspended, and the size of the microscope’s HPF.

1. Examination of unspun urine in a hemocytometer counting chamber. A hemocytometer is a precisely milled slide etched with measured squares, which allows exact enumeration of the cells in each square. Because the distance between the etched surface and the coverslip is known exactly, it is possible to determine the number of cells per unit volume of specimen. Enough fresh unspun urine is placed on the slide to fully cover the counting area, and the cells are counted. There should usually be less than 1 WBC/µL, although more than 10 WBCs/µL and more than 5 RBCs/µL are unequivocally abnormal. The threshold for diagnosing UTI is usually set at 8 or more WBCs/µL. Bacteria do not sink to the surface of the hemocytometer, so counting them through the many focal planes of the chamber is not possible, although methods to estimate the bacterial count per unit volume have been described. The hemocytometer is accurate, fast (time need not be spent on staining or centrifugation), and relatively easy to master. Its major drawbacks are cost (approximately $150 for the slide and coverslip) and fragility (easily destroyed if dropped). These characteristics are problematic given the conditions and resources in many EDs, although their use has been advocated by some. Formed elements other than WBCs and RBCs (e.g., casts) occur in such low concentrations that they are encountered in the hemocytometer only by chance, and therefore microscopy of spun urine is needed for their identification.

2. Examination of unspun, unstained urine placed on a regular microscope slide. This qualitative method is sometimes used for the diagnosis of UTI. Using 1 organism/HPF as a positive result, the sensitivity and specificity of detecting 10⁵ CFUs/mL are between 60% and 90%. This method identifies only 1 WBC/HPF with the highly pyuric state of 250 WBCs/µL, which has led some to advocate the use of more than 1 WBC per low-power field as a criterion for infection.

3. Examination of unstained, centrifuged urine. With this method, 10 mL of urine is centrifuged at approximately 450g (1000 to 4000 rpm) for 3 to 5 minutes. Roughly 9 mL of supernatant is poured off and the pellet is resuspended in the remaining fluid. This suspension is placed on a slide with a coverslip and examined. The larger formed elements, especially casts, tend to migrate to the edge of the coverslip, and they can be seen with low magnification. One or two casts, depending on the clinical context, may be normal; more are not. The morphology of the more commonly encountered formed elements of urine sediment is shown in Fig. 67.3. The significance of each is beyond the scope of this text, but this information can be found in a standard textbook of clinical laboratory procedures and diagnostic testing. When examining centrifuged urine, more than 5 WBCs/HPF in the middle of the coverslip has traditionally been taken as being indicative of abnormal pyuria. Most authors have estimated that 10 WBCs/µL is equivalent to

<table>
<thead>
<tr>
<th>TABLE 67.4 Relationship Between Urinary Bilirubin, Urobilinogen, and Stool Color in Jaundiced Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HEALTHY NORMAL</strong></td>
</tr>
<tr>
<td>---------------------</td>
</tr>
<tr>
<td>Urinary Bilirubin</td>
</tr>
<tr>
<td>Urinary Urobilinogen</td>
</tr>
<tr>
<td>Stool Color</td>
</tr>
</tbody>
</table>

...
of expense, delay, and resources. A brief discussion of their relative strengths and weaknesses ensues.

**Urine Dipstick**

Dipstick testing of urine is faster than microscopic UA, is less labor-intensive, cheaper, and circumvents multiple sources of potential and proven error. Is it sufficiently accurate to replace UA; however, if either leukocyte esterase or nitrites are used to indicate infection, the dipstick is only 50% to 90% sensitive for culture-proven infection. Is it sufficiently accurate to replace UA; however, if either leukocyte esterase or nitrites are used to indicate infection, the dipstick is only 50% to 90% sensitive for culture-proven infection.16,48,49,70,72 This is not adequate to rule out infection in symptomatic patients, in whom the prevalence of disease is high, but may be acceptable in asymptomatic patients, in whom the test has a serviceably high negative predictive value of 95% to 99%. A large retrospective series demonstrated laboratory-performed UA (with microscopy) in children to be 8% more sensitive than dipstick testing in the ED, with similar specificity of approximately 80% for both tests.24 In symptomatic men, sensitivity is enhanced by taking a positive result in any one (or more) of either leukocyte esterase, nitrites, protein, or blood as an indication of UTI.16,73–75 This process can be augmented by allowing extra time before reading the strip. For women with classic symptoms of UTI, empirical treatment is recommended because no test can rule out infection.71 A negative dipstick test in a patient with a high pretest probability of UTI should prompt a search for an alternative source of the patient’s symptoms. Maneuvers to enhance dipstick sensitivity diminish its specificity, which may be as low as 26%.10 This has led some to advocate microscopic UA on all urine specimens that are found to be abnormal by dipstick,74 but this probably adds little to a carefully performed dipstick test.

**Summary of Tests Used in the Diagnosis of UTI**

The three tests commonly used to evaluate a patient for the presence or absence of UTI are urine dipstick, microscopic UA, and urine culture. Each represents an increasing degree of expense, delay, and resources. A brief discussion of their relative strengths and weaknesses ensues.

4. Examination of Gram-stained, uncentrifuged urine. Also a semiquantitative measurement, it is estimated that 1 bacterium/HPF is equivalent to 10^4 CFUs/mL in bacterial culture. As the gold standard for the diagnosis of UTI is more than 8 WBCs/µL of unspun urine, many infections will escape detection with this method. Various numbers of bacteria per HPF have been used as criteria for the diagnosis of UTI. A threshold of 10 to 20 organisms/HPF has been recommended to rule out bacteriuria at the 10^4-CFU/mL level. As noted previously, this threshold would not exclude infection in symptomatic patients.

5. Examination of Gram-stained, centrifuged urine. This method is probably the optimal technique, short of culture, for the assessment of bacteriuria. It is more than 95% sensitive and more than 60% specific at 10^4 CFUs/mL, a concentration of bacteriuria that is an order of magnitude lower than the previously described methods. Detection of 1 organism per oil-immersion field constitutes a positive result. Specificity is increased to 95% if 5 organisms/HPF are seen.

Microscopic UA
If the urine dipstick has such poor specificity when it is used as a test with adequate sensitivity, should it be discarded altogether and microscopic UA be relied on instead? Apart from the hemocytometer, the most reliable method for identifying significant bacteriuria is oil-immersion microscopy of Gram-stained, centrifuged urine. Nonetheless, the practice in most hospital laboratories is to examine a resuspended pellet of unstained, centrifuged urine. The problems with this method have been discussed. In various studies a range between 1 and 10 organisms/HPF or 5 WBCs/HPF has been considered a “positive” test (as the threshold number rises, so does specificity, at the price of sensitivity). In aggregate, the accuracy of microscopy in the diagnosis of UTI is similar to that of the dipstick alone, with 22% false-positive and 23% false-negative rates when compared with culture. Microscopic UA, like the dipstick, cannot rule out infection in symptomatic patients. The specificity of pyuria is improved when viewed as a marker of all genitourinary infections, including urethritis, prostatitis, epididymitis, vaginitis, and cervicitis. These diagnoses should always be entertained in patients with urinary symptoms, especially those with sterile pyuria.

Urine Culture
Cultures are indicated for any potentially complicated UTI, including infections in children; men; women with recurrences or relapses; immunocompromised individuals; patients with urinary tract pathology, including stones and possible pyelonephritis; and pregnant patients. Cultures are usually recommended at approximately the 16th week of gestation. Every effort should be made to obtain a high-quality specimen because of the difficulty in distinguishing contamination from significant low-count bacteriuria in asymptomatic patients and in view of the expense of cultures and treatment if the culture is positive.

The Bottom Line
It is well established that female patients with symptoms of uncomplicated lower UTI can be treated without culture because the prevalence of disease is 50% or greater in women with classic urinary symptoms. Treatment is generally benign, and 95% of urine cultures that are positive will yield a limited number of organisms with predictable antibiotic susceptibilities. Some authors recommend dipstick or microscopic UA (or both) because a negative test might prompt more careful consideration of alternative diagnoses in this group, with a high incidence of sexually transmitted disease. If none is found, such patients can be treated empirically.

Fluoroquinolone antibiotics are no longer recommended as empiric treatment for uncomplicated UTI. Traditional choices have included trimethoprim-sulfamethoxazole or macrodantin, but both of these agents have high resistance rates in many locations, so choices should be based on local microbial susceptibility patterns, if available.

Urine testing is more likely to influence clinical management in patients with an intermediate pretest probability of infection. Urine microscopy may also help in distinguishing pyelonephritis (WBC casts), vaginitis (absence of pyuria or hematuria in a meticulous MSCC or catheterized specimen), and urethritis (pyuria, rarely hematuria) from cystitis (pyuria and often hematuria). In the ED, the ease of performing a dipstick test argues for its use without UA because it is equally sensitive unless formed elements such as casts, crystals, Trichomonas, or other parasites are suspected. In patients with pyelonephritis, urine cultures are warranted because they alter therapy in approximately 5% of cases. In most asymptomatic patients, a negative dipstick or UA result has sufficient negative predictive value to rule out disease unless the patient is pregnant or undergoing urologic surgery. Special clinical considerations in some patients (e.g., the immunocompromised, diabetic females at high risk for UTI) may mandate adjustments to this approach. Recommendations vary regarding infants. Cultures are recommended for all febrile infants younger than 2 months and for children who appear sick or have a high pretest probability of infection. In a 2009 study, combined microscopic and dipstick UA was only 64% sensitive (and 91% specific) for culture-proven UTI in febrile infants younger than 24 months, thus suggesting the use of culture unless an alternative source of infection is clear.

Testing for Pregnancy
Some investigators have found female patients reliable in determining their own pregnancy status. Others have not. With respect to ordering radiographic studies on women of childbearing age, clinicians should determine from their own practice experience the reliability of patient history in this regard. If in doubt and with the potentially lethal outcome of an ectopic pregnancy, it might be prudent to use an objective test. Pregnancy tests are based on the detection of β-human chorionic gonadotropin (β-hCG) in serum or urine. β-hCG is secreted by trophoblastic cells of the placenta starting from the time of implantation of the blastocyst. Qualitative serum and urine tests can detect β-hCG levels of between 15 and 25 mIU/mL (Fig. 67.4). The concentration of β-hCG is usually lower in urine than in serum, which accounts for the slight advantage of serum tests in detecting early pregnancy. Optimal results on urine pregnancy tests are obtained with first-voided, concentrated morning specimens. It has recently been established that whole blood is a reliable specimen for the bedside dipstick tests used in most EDs. The same volume of blood is placed in the testing well(s) as when performed with urine. This alternative may be lifesaving in timely recognition of an ectopic pregnancy in a hypotensive female with pelvic complaints and unknown pregnancy status when a urine specimen is unavailable or results in significant delay.

If fertilization has occurred, β-hCG levels of 5 to 8 mIU/mL (the threshold of the quantitative serum test) correspond to the 9th to 11th day after ovulation (23 to 25 days after the first day of the last normal menstrual period). In a viable intrauterine pregnancy (IUP), the β-hCG level doubles approximately every 2.5 days during the first 4 weeks of gestation and reaches a serum level of greater than 25 mIU/mL, which is detectable by virtually all pregnancy tests on the first day of the missed menstrual period. The doubling rate declines to every third day thereafter. β-hCG reaches a peak of between 100,000 and 200,000 mIU/mL between the 10th and 14th gestational weeks and declines to 10,000 to 20,000 mIU/mL for the rest of the pregnancy (Table 67.5). There is a wide range of β-hCG levels in different women at the same stage of gestation, thus making definite clinical determination on the basis of a single quantitative test impossible. False-positive test results have been described in association with molar pregnancy, choriocarcinoma, teratoma, occasional
malignancies outside the genitourinary tract, and very high levels of proteinuria. There is one report of a positive urine β-hCG finding (but normal serum β-hCG) associated with a tuboovarian abscess.92

Although a previous quantitative β-hCG level is rarely available to the emergency clinician, doubling rates are an important part of the assessment of a healthy first-trimester pregnancy. Fetal nonviability, ectopic pregnancy, and intrauterine demise are signaled by abnormalities in the predicted rise in quantitative β-hCG.93,94 A serum quantitative β-hCG level that does not increase by 66% every 48 hours has a 75% chance of being a nonviable pregnancy.94,95 β-hCG levels in a healthy IUP and associated sonographic findings are listed in Table 67.5.

The rate of decline in quantitative β-hCG after gestation depends on the reason for the conclusion of the pregnancy. After a term delivery, β-hCG falls to zero in 2 weeks; after surgery for an ectopic pregnancy, the range is 1 to 31 days, with a median of 8.5 days; after a first-trimester spontaneous abortion, the range is 9 to 35 days (median, 19 days); and after a first-trimester elective abortion, the range is 16 to 60 days (median, 30 days).43

The association between β-hCG levels and gestational dates has led to the concept of the “discriminatory zone.” In a normal pregnancy, a quantitative β-hCG level of 1000 to 2000 mIU/mL should be reflected by the presence of a double decidual sac (at the least) on transvaginal ultrasound (and on transabdominal ultrasound with levels higher than 6500 mIU/mL).96 The clinician should beware of several potential pitfalls in applying this concept. First, it applies only to the double decidual sign, itself subject to interobserver variation among sonographers. The discriminatory zone cannot be extrapolated to the expected levels at which any of the more definitive sonographic signs of IUP such as yolk sac, fetal pole, or cardiac motion should be seen. Second, although it is true that if the double decidual sign is absent at these thresholds, the pregnancy is almost certainly abnormal with a significant possibility of

### Table 67.5 Relationship Between Gestational Age, Qβ-hCG Levels, and Ultrasound Findings

<table>
<thead>
<tr>
<th>TIME ELAPSED FROM THE FIRST DAY OF THE LAST NORMAL MENSTRUAL PERIOD</th>
<th>Qβ-hCG LEVEL (mIU/mL) USING THE IRP</th>
<th>ULTRASOUND FINDINGS</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 28 days</td>
<td>5–50</td>
<td></td>
</tr>
<tr>
<td>4–5 wk</td>
<td>50–500</td>
<td>From approx. 4.5 wk and Qβ-hCG level of 1000–1500 EVU can show a nonspecific intrauterine sac</td>
</tr>
<tr>
<td>5–6 wk</td>
<td>100–10,000</td>
<td>Definitely abnormal if no DDS seen on TVU with a Qβ-hCG level &gt;2000 or by TAU with a Qβ-hCG level &gt;6500 Abnormal if no YS with an MSD &gt;10 mm</td>
</tr>
<tr>
<td>6–7 wk</td>
<td>1000–30,000</td>
<td>FP, cardiac activity 5.5–7 wk, Qβ-hCG level &gt;10,000 Abnormal if no FP with an MSD &gt;18 mm</td>
</tr>
<tr>
<td>7–8 wk</td>
<td>3500–115,000</td>
<td></td>
</tr>
<tr>
<td>8–14 wk</td>
<td>12,000–270,000</td>
<td></td>
</tr>
<tr>
<td>&gt;10 wk</td>
<td>270,000–15,000</td>
<td></td>
</tr>
</tbody>
</table>

DDS, Double decidual sac; EVU, endovaginal ultrasound; FP, fetal pole; IRP, international reference preparation; MSD, mean sac diameter; Qβ-hCG, quantitative β-human chorionic gonadotropin; TAU, transabdominal ultrasound; TVU, transvaginal ultrasound; YS, yolk sac.
being ectopic, the converse—that it is pointless to perform ultrasonography if the quantitative β-hCG level is lower than 1000 mIU/mL—is not true. This is because the discriminatory zone does not preclude the possibility of identifying an ectopic pregnancy (or an IUP) before the quantitative β-hCG level reaches 1500 mIU/mL. Ectopic gestational sacs are pathological and therefore do not display β-hCG levels according to nomograms established for normal IUPS, and advanced ectopic pregnancies may have low β-hCG levels. One percent of ectopic pregnancies have a quantitative β-hCG level of less than 10 mIU/mL, and approximately a third of ectopic pregnancies are diagnosed in patients with a quantitative β-hCG level lower than 1000 mIU/mL. Even if no definitive diagnosis can be made, the clinician should be aware that pregnant ED patients with pelvic complaints and a β-hCG level higher than 1000 mIU/mL is predictive of a positive blood culture.105–109 The subjective impression of “having fever” in adults is not a reliable indicator of the presence of fever, although the subjective impression of “no fever” is much more likely to be accurate.110 Prediction models to optimize the use of this costly test have been explored but are cumbersome, add little to educated clinical judgment, and lack widespread validation or acceptance.111–113 Many studies have shown that blood cultures in patients with uncomplicated pneumonia or pyelonephritis are of very limited clinical value,114–117 although a 2010 prospective investigation of patients with upper UTI found that malignancy, an indwelling urinary catheter, and ongoing antimicrobial treatment were clinical conditions that made it significantly more likely that blood cultures would be discordant with urine cultures and that they would reveal clinically important additional information.126 Blood cultures are obtained during 2.8% of all ED visits in the United States, and in a single 3-year period this rate increased by 33%.127 As disease prevalence is unlikely to have changed so rapidly, this increase is likely due to changes in practice. These changes have been coincident with regulatory agency-mandated blood cultures in patients being evaluated for pneumonia and have led to calls for more stringent criteria for obtaining blood cultures.

In children, the traditional teaching that blood cultures are indicated for all patients younger than 2 years with fever higher than 38.6°C (>101.5°F) and without an obvious source is being modified by the widespread use of pneumococcal conjugate (PC) and *Haemophilus influenzae* type b (HIB) vaccines.128–129 In the post-PC and post-HIB vaccine era, the incidence of occult bacteremia in otherwise well-appearing febrile children is probably lower than 1%, thus making the false-positive rate at least four times as high.130–134 It seems reasonable to withhold blood cultures and empirical antibiotic coverage in otherwise well-appearing febrile children with reliable parents. The risk for bacteremia in a child is positively correlated with the degree of fever, WBC count, and rapidity of onset of the illness. It is inversely proportional to the patient’s age.131–133 In infants younger than 2 months with temperatures higher than 38°C (>100.5°F), some authorities would recommend blood cultures regardless of the presence or absence of a source, although this approach is subject to modification by experienced clinicians based on the patient’s age and clinical setting. A child with a normal temperature in the ED and a history from the parents of a tactile fever needs to be approached in the same way as a patient with fever documented on physical examination. Parents’ tactile impression of fever is reliable, and bacteremic children, like adults, have intermittent fever, with up to 50% afibrile rates in children with demonstrated bacteremia.135,136

**BLOOD CULTURES IN THE ED**

**Indications**

Blood cultures are indicated when the clinical findings suggest an otherwise unidentifiable bacteremic state (Box 67.2). Twenty-five percent of patients with documented bacteremia have periods without fever.104 In the elderly the proportion is even higher, with 50% of bacteremic patients older than 65 having a temperature between 97.1°F (36.2°C) and 100.9°F (38.3°C) and at least 13% with no documented temperature higher than 99.1°F (37.3°C) at any time.105–108 In the elderly, increasing age, vomiting, altered mental status, urinary incontinence, presence of a Foley catheter, or greater than 6% band forms is predictive of a positive blood culture.106,109 The subjective impression of “having fever” in adults is not a reliable indicator of the presence of fever, although the subjective impression of “no fever” is much more likely to be accurate.110 Prediction models to optimize the use of this costly test have been explored but are cumbersome, add little to educated clinical judgment, and lack widespread validation or acceptance.111–113 Many studies have shown that blood cultures in patients with uncomplicated pneumonia or pyelonephritis are of very limited clinical value,114–117 although a 2010 prospective investigation of patients with upper UTI found that malignancy, an indwelling urinary catheter, and ongoing antimicrobial treatment were clinical conditions that made it significantly more likely that blood cultures would be discordant with urine cultures and that they would reveal clinically important additional information.126 Blood cultures are obtained during 2.8% of all ED visits in the United States, and in a single 3-year period this rate increased by 33%.127 As disease prevalence is unlikely to have changed so rapidly, this increase is likely due to changes in practice. These changes have been coincident with regulatory agency-mandated blood cultures in patients being evaluated for pneumonia and have led to calls for more stringent criteria for obtaining blood cultures.

In children, the traditional teaching that blood cultures are indicated for all patients younger than 2 years with fever higher than 38.6°C (>101.5°F) and without an obvious source is being modified by the widespread use of pneumococcal conjugate (PC) and *Haemophilus influenzae* type b (HIB) vaccines.128–129 In the post-PC and post-HIB vaccine era, the incidence of occult bacteremia in otherwise well-appearing febrile children is probably lower than 1%, thus making the false-positive rate at least four times as high.130–134 It seems reasonable to withhold blood cultures and empirical antibiotic coverage in otherwise well-appearing febrile children with reliable parents. The risk for bacteremia in a child is positively correlated with the degree of fever, WBC count, and rapidity of onset of the illness. It is inversely proportional to the patient’s age.131–133 In infants younger than 2 months with temperatures higher than 38°C (>100.5°F), some authorities would recommend blood cultures regardless of the presence or absence of a source, although this approach is subject to modification by experienced clinicians based on the patient’s age and clinical setting. A child with a normal temperature in the ED and a history from the parents of a tactile fever needs to be approached in the same way as a patient with fever documented on physical examination. Parents’ tactile impression of fever is reliable, and bacteremic children, like adults, have intermittent fever, with up to 50% afibrile rates in children with demonstrated bacteremia.135,136

**THE CONTROVERSY REGARDING OUTPATIENT BLOOD CULTURES**

There is a long-standing debate regarding the utility of outpatient blood cultures (i.e., blood cultures on patients who are discharged from the ED pending results). Arguments for and against outpatient blood cultures are summarized in Box 67.3. Opponents cite medicolegal issues, problems with follow-up, high contamination rates, low rates of positive cultures, and
ARGUMENTS AGAINST OUTPATIENT BLOOD CULTURES
Low true-positive rates
True positives are rarely clinically significant.
False positives are expensive and time-consuming for both the patient and the health care system.
Unreliability of emergency department follow-up makes positive results a medicolegal liability.

ARGUMENTS FOR OUTPATIENT BLOOD CULTURES
Permits outpatient evaluation and management of patients with a low probability of disease (especially infectious endocarditis).
Patients: financial, psychological, and nosocomial cost savings from spared admissions
Society: financial and nosocomial cost savings from spared admissions
Allows initiation of antibiotics for possible alternative infectious diagnoses without irrevocable loss of the opportunity for blood cultures

ARGUMENTS AGAINST OUTPATIENT BLOOD CULTURES
Even lower rates of frequency in patients in whom therapy is changed because of culture results. Proponents also cite medicolegal concerns, positive rates similar to those seen with inpatient blood cultures, cost savings, and the benefit of diagnosing significant, yet subtle bacteremic states (such as endocarditis) Patients discharged with occult bacteremia generally have a benign clinical course. It should be noted that “occult” may be something of a misnomer in this setting, because some aspect of the patient’s clinical state prompted the decision to obtain blood cultures. Proponents of outpatient blood cultures also point out that the high false-positive rates seen in many ED series should be an indictment of poor technique, not of the test itself.

On the basis of current data, it seems fiscally extravagant to admit all patients in whom a bacteremic state is possible and injudicious to deny blood cultures solely on the basis of a patient not appearing “toxic enough” to warrant admission. Societal and economic pressures to avoid hospital admission buttress these clinical considerations. Outpatient blood cultures, with due attention to collection technique, patient selection, and arrangements for follow-up, have a place in emergency practice. This approach is reflected by the fact that fully half of ED blood cultures are obtained from patients who are not admitted to the hospital.

Technique for Obtaining Blood for Culture
False-positive blood cultures obtained in the ED can result in up to $88000 in additional costs, with an average increase in hospital length of stay of 1 day per patient. Studies have demonstrated sources of contamination at every stage of the process of obtaining and inoculating blood cultures. In addition to obvious sources of contamination from the patient’s and phlebotomist’s skin, antiseptic agents and gloves have been implicated. Some authorities have argued that the primary source of contamination is in the laboratory processing of specimens. The consensus is that the most common source of contamination is the process of phlebotomy and inoculation of blood culture bottles. Obviously, this is the single step over which emergency clinicians have control, either directly or via protocols of technique for blood culture phlebotomy. Contamination rates are typically between 1.5% and 3%, although many ED series show much higher rates. Conversely, settings in which there are rigorous protocols for obtaining blood cultures have contamination rates as low as 1%. A high degree of sensitivity is required for blood cultures. Many significant bacteremic illnesses have been documented with as little as 1 CFU/10 mL of blood. In a cadaver study, human skin has been shown to have a bacterial concentration of between 10³ and 10⁶ CFUs/mL on the forearm and groin, respectively. Designed to detect vanishingly low concentrations of bacteria, the test is clearly susceptible to false-positive results (impaired specificity) when blood must necessarily be obtained by passing a needle through the skin. Eighty percent of the skin flora is transient, superficial, and removable; 20% inhabit the sebaceous ducts and hair follicles and cannot be removed without destroying the skin. The former group consists of predominantly gram-positive and gram-negative aerobes and is the target of skin disinfectants.
The primary agents for skin disinfection are iodine compounds, alcohols, chlorhexidine, and hexachlorophene. Iodine solution remains a gold standard and kills bacteria, fungi, protozoa, and viruses but has been replaced in many institutions because of concern about skin burns and allergic reactions. Reports of the former were probably due to the use of 7% solution. The risk for a burn or an allergic reaction is thought to be negligible with the currently available 2% preparation. The most effective cleansing agent is tincture of iodine, which is a mixture of 2% iodine solution and 70% alcohol. Povidone-iodine 10% solution (Betadine, Purdue Pharma L.P., Stamford, CT) has a much lower free-iodine concentration than iodine solution and is therefore less potent. Iodine is superior to hexachlorophene and chlorhexidine in killing gram-negative bacteria. Iodine, like other antiseptic agents, is inhibited by the presence of organic matter and thus requires thorough skin cleansing before the application of any skin disinfectant.
Ethyl or isopropyl alcohol should be used in a 60% to 80% solution. Alcohol prep pads, which generally contain 70% isopropanol, have solved traditional concerns regarding evaporation of alcohol from cotton balls stored in jars. Alcohol is a less powerful germicide than iodine in vitro and kills only 90% of surface bacteria after a full 2 minutes with reapplication to prevent drying. Alcohol foam applicators can avoid premature drying. Alcohol is inactive against fungi, spores, and viruses, but in vivo studies of blood culture contamination rates have shown it to compare favorably with iodine. Because iodine solution is often not available and iodophor solutions are less potent, alcohol still has an important place in skin antisepsis. In addition, alcohol is an excellent solvent, so alcohol pads may assist in skin preparation by removing dirt- and microbe-laden skin oils before the application of iodine compounds.
Chlorhexidine (Hibiclens, Mölnlycke Health Care, Norcross, GA) and hexachlorophene (pHisohex, Aspen Pharmacare Australia Pty Ltd) are antiseptics that are more effective against gram-positive than gram-negative bacteria. Both agents are absorbed intradermally, thereby offering prolonged antimicrobial activity, which is the basis of their popularity as surgical.
Studies show chlorhexidine to be more potent than hexachlorophene. The most important concept in skin disinfection is that bacteria do not die at the instant of contact with disinfectant agents. Iodine (2%), which is twice as potent as 10% povidone-iodine, requires at least 90 seconds in contact with skin for inactivation. Most studies show that newly placed intravenous (IV) catheters are an acceptable source of blood specimens for culture, provided that the usual measures are taken in skin preparation. Chronically placed lines either trend toward or show statistically significant increases in contamination rates. An exception can be made for carefully tended central venous access ports in cancer patients. These cultures may have increased sensitivity in identifying bacteremia, possibly because the catheters themselves are often a source of bacteremia in these patients.

### Box 67.4

**Skin Preparation and Technique for Drawing Blood for Culture**

Cleanse the skin with alcohol swabs three times or until the swabs appear entirely free of surface dirt. Allow to dry. Apply 10% povidone-iodine or (preferably) 2% iodine solution or (ideally) 2% tincture of iodine in 70% alcohol three times in centrifugal circles from the anticipated site of venipuncture.

After the third swab, allow to dry for at least 60 seconds. During this period:

- Remove the covers and sterilize the rubber stoppers of blood culture bottles with iodine, alcohol, or both.
- Lay out sterile gloves.
- Use a paper glove wrapper as a sterile field for the needle and syringe (not necessary if using a Vacutainer system). Wipe off dry iodine at the venipuncture site with alcohol. Substitute chlorhexidine if placing an indwelling catheter.
- Obtain at least 20 mL of blood to place in two bottles.
- If short, use at least 10 mL for an aerobic bottle.
- If more than 20 mL, use additional aerobic bottles. Place no more than 10 mL of blood per bottle.
- Inoculate bottles without changing needles between bottles. However many bottles are filled, this is one set of blood cultures. Another set cannot be obtained from this site, regardless of how many bottles are filled.

During this period:

- Use a paper glove wrapper as a sterile field for the needle and syringe (not necessary if using a Vacutainer system). Wipe off dry iodine at the venipuncture site with alcohol. Substitute chlorhexidine if placing an indwelling catheter.
- Obtain at least 20 mL of blood to place in two bottles.
- If short, use at least 10 mL for an aerobic bottle.
- If more than 20 mL, use additional aerobic bottles. Place no more than 10 mL of blood per bottle.
- Inoculate bottles without changing needles between bottles. However many bottles are filled, this is one set of blood cultures. Another set cannot be obtained from this site, regardless of how many bottles are filled.

**Special Considerations in Obtaining Blood for Culture**

### Changing the Needle After Phlebotomy

In considering this issue it is important to emphasize the distinction between needle changing and needle recapping. The latter is a well-established risk to health care workers. It contravenes standard recommendations for universal precautions and should not be performed. Needle replacement using the standard needle removal device on sharps containers is an unquantified risk, but clearly much less dangerous than recapping.

Based on little scientific data it was long considered essential to change the phlebotomy needle before inoculating blood culture bottles. With increasing awareness of the risks associated with needlestick injuries, this practice has come under scrutiny. Studies generally show trends toward lower contamination rates with needle change, but without reaching statistical significance. Thus, not changing needles before inoculation of blood culture bottles is acceptable practice for routinely obtaining blood for culture. In situations in which the results of blood cultures are of paramount importance (e.g., suspected infectious endocarditis, where empirical antibiotics are to be started immediately), needles can be changed before inoculation of culture bottles (without recapping).

### Special Access Sites

Most studies show that newly placed intravenous (IV) catheters are an acceptable source of blood specimens for culture, provided that the usual measures are taken in skin preparation. Chronically placed lines either trend toward or show statistically significant increases in contamination rates. An exception can be made for carefully tended central venous access ports in cancer patients. These cultures may have increased sensitivity in identifying bacteremia, possibly because the catheters themselves are often a source of bacteremia in these patients.

### Heel Stick in Neonates

This technique resulted in recovery rates of bacteria equivalent to those with phlebotomy in two studies. As approximately 25% of bacteremic infants have fewer than 5 CFUs/mL of blood, this proportion (25%) will be missed if less than 0.2 mL is obtained for culture. For this reason, heel stick should be considered a source of last resort for blood culture.

### Intraosseous Specimens

This technique may also be used when phlebotomy is impossible.

### Timing of Blood Cultures

In most circumstances the timing of blood cultures is moot in the ED. Patients are sick enough to warrant the initiation of empirical antibiotics or are well enough for discharge, so two or more sets of blood need to be drawn immediately. The timing of blood cultures might become a consideration in a patient requiring admission but in whom the diagnosis of bacteremia is in doubt such that empirical antibiotic therapy is withheld. Contrary to medical lore, true-positive blood cultures are more likely if blood is drawn in the 12 hours before a fever spike. Furthermore, except for infectious endocarditis, most clinically significant bacteremia is thought to be intermittent, so multiple sets of specimens obtained at one time for culture would heighten the risk for missing the period of bacteremia. For patients admitted to the hospital with the tentative diagnosis of sepsis, it is theoretically advantageous to draw the three sets of blood for culture over the first 12 to 24 hours of admission. If immediate administration of antibiotics is indicated, the two or three sets of blood for culture should be obtained before initiation of antibiotic therapy.

### Blood Culture Volumes

#### Volumes in Adults

A large number of studies almost uniformly demonstrate that the sensitivity of blood cultures is directly related to the volume...
of blood cultured. In a representative study, Ilstrup and Washington showed that 20 mL and 30 mL of blood yielded, respectively, 38% and 62% more true-positive results than 10 mL did. Mermel and Maki showed that each additional milliliter of blood yields an average of 3% more true-positive results. This finding is also consistent with the fact that 40% of adults with bacteremia have less than 1 CFU/mL of blood and that 20% have less than 1 CFU/10 mL. Alternatively expressed, if 10 mL of blood is obtained for culture, 20% of patients with continuous bacteremia will be missed. As most bacteremia is intermittent and endogenous factors in blood will cause some inhibition of bacterial growth even with modern lysis and filtration centrifugation techniques, the false-negative rate in clinical practice will always be significantly higher. On purely mathematical grounds, 10 mL per set of blood cultures is a bare minimum for culture. In adults, most authorities recommend at least 30 mL of blood per culture site or set. To ensure dilution of the blood’s antibacterial properties (e.g., immunoglobulins, complement, WBCs), culture bottles should contain a concentration of blood of less than 1 part blood to 10 parts medium. If 30 mL of blood is obtained from one site, it should be divided equally into three of the usual 100-mL broth bottles.

**Volumes in Children**

A blood volume of 30 mL from a 70-kg adult is equivalent to 0.5 mL of blood from a 3.5-kg neonate. Fortunately (for the utility of blood cultures), levels of bacteremia are typically 10-fold higher in neonates than in adults. The sicker the child, the greater the likelihood of a high level of bacteremia. As the immune system matures during infancy, levels of bacteremia might be expected to fall toward those seen in adults, so small culture volumes are at increased risk for false-negative results. As a rule of thumb, a similar volume of blood with respect to body mass should be drawn from children as is drawn from adults: approximately 1 mL/2.5 kg, or 4 mL blood per 10-kg body mass.

**How Many Sets of Blood Cultures Are Needed?**

A set of blood cultures is the sample obtained from a single site. A 1-mL specimen from a neonate placed in an aerobic bottle and a 30-mL specimen from an adult divided between fungal, aerobic, and anaerobic bottles are both a single set of blood cultures (Fig. 67.5). Two or more sets of blood cultures make up a series. The information derived from the blood culture sets is pooled in such a way to make both the sensitivity and specificity of the series greater than that of the component sets. Sensitivity is enhanced because, as discussed earlier, an individual set is typically not more than 80% sensitive. Specificity is improved for microbes that can act as both pathogens and contaminants by determining whether they appear in all sets (pathogen) or in only some (more likely a contaminant).

Although this conceptual process is applied to all blood culture series, the focus of inquiry depends on the infectious process being ruled in or out. For example, in an elderly patient with sepsis and a chronic indwelling Foley catheter, it is extremely unlikely that the causative organism is a typical skin contaminant. The usual causes of false-positive blood cultures will therefore be easily recognized, thus lowering the false-positive rate for the series and yielding a test with intrinsically higher specificity. At the same time, with the typical pathogens in this clinical context being nonfastidious organisms, sensitivity is typically at approximately 99% with two sets consisting of 20 mL of blood per set. Conversely, in a patient with a prosthetic heart valve, fever, and signs of septic emboli, many probable pathogens are also skin contaminants, thereby lowering the specificity of each individual blood culture set. Thus, at least two sets of cultures must be positive with such organisms before the overall test (i.e., the series) is considered positive. At the same time, this clinical picture makes the pretest probability of disease very high (diminishing the negative predictive value of a negative set), so an extremely sensitive overall test (i.e., series) will be needed to adequately rule out disease. In this setting, most authorities would recommend four sets of blood culture bottles, with good volumes in each. Except in infants, single sets of blood for culture are of insufficient sensitivity or specificity to be of any utility and should not be drawn. The recommended numbers of sets of blood cultures as they relate to the pretest probability of disease and causative organism are summarized in Table 67.6.

**Aerobic Versus Anaerobic Versus Other Bottles**

Anaerobic infections tend to occur in poorly perfused tissues or locations and frequently evolve into abscesses. Both characteristics mean that these infections tend to be isolated or separated from the bloodstream, thereby decreasing the likelihood of bacteremia and detection by blood culture. For these reasons it is not surprising that anaerobic isolates account for 0.5% to 12% of positive blood cultures. Blood cultures in general have a typical true-positive rate of approximately 5% with 5% or less of positive blood cultures being anaerobic, more than 400 patients need to have a complete series of blood drawn for culture to detect one case of anaerobic bacteremia. If more than 50% of anaerobic infections are clinically evident before culture, at least 800 blood culture series are needed to generate a single anaerobic result that would alter clinical management. Anaerobic cultures may also have the unintended consequence of compromising the sensitivity of aerobic cultures in situations in which less than the ideal 20 mL of blood is
Identifying Contaminants

The emergency clinician may receive calls from the laboratory about the results of positive blood cultures obtained on previous shifts. These may be true positives as a result of true contamination or may be caused by the intermittent bacteremia that occurs in normal, healthy people. This situation has been complicated by the increasingly common identification of Staphylococcus epidermidis, Streptococcus viridans, and fungi as real pathogens in blood culture series.\(^214\text{–}217\) The expense associated with false-positive blood cultures has been noted earlier. These costs emphasize the importance of good technique in obtaining blood for culture.\(^218\text{,}\,219\)

Distinguishing contaminants from clinically significant bacteremia is based on both microbiologic information and the patient’s clinical condition.\(^212\text{–}213\) Features of false-positive blood culture results are listed in Box 67.6.\(^189\text{,}\,198\text{,}\,220\) It would be prudent to contact discharged patients with positive blood cultures, even when contamination is suspected on a microbiologic basis, to ensure that their condition is improving.

<table>
<thead>
<tr>
<th>TABLE 67.6 Numbers of Blood Culture Sets to Be Obtained in Various Clinical Situations in Adults</th>
</tr>
</thead>
<tbody>
<tr>
<td>NUMBER OF SETS (MINIMUM)</td>
</tr>
<tr>
<td>Two sets</td>
</tr>
<tr>
<td>Three sets</td>
</tr>
<tr>
<td>Four sets</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TABLE 67.7 Blood Culture Types to Be Used in Various Clinical Settings</th>
</tr>
</thead>
<tbody>
<tr>
<td>CLINICAL SITUATION</td>
</tr>
<tr>
<td>Children &lt;12 yr</td>
</tr>
<tr>
<td>Adults and children &gt;12 yr</td>
</tr>
</tbody>
</table>

Coagulase-negative staphylococci (\textit{Staphylococcus epidermidis}) and \textit{Staphylococcus viridans} in a single bottle in patients not suspected of having infectious endocarditis and without chronic indwelling intravenous access catheters are usually contaminants. Corynebacteria (previously known as diphtheroids), \textit{Propionibacterium acnes}, and \textit{Bacillus} species are usually contaminants but can be pathogenic in the immunocompromised. Multiple organisms in a series suggest contamination. Species that grow after prolonged culture have a higher likelihood of being contaminants. Conversely, early-growing bacteria have a much higher likelihood of being pathogens. The patient’s symptoms have resolved or are inconsistent with sepsis (beware with infectious endocarditis, which can have an indolent course). A primary source (e.g., sputum or urine) has a different pathogen isolated.

\textbf{Identifying Contaminants}

The emergency clinician may receive calls from the laboratory about the results of positive blood cultures obtained on previous shifts. These may be true positives as a result of true contamination or may be caused by the intermittent bacteremia that occurs in normal, healthy people. This situation has been complicated by the increasingly common identification of \textit{Staphylococcus epidermidis}, \textit{Streptococcus viridans}, and fungi as real pathogens in blood culture series.\(^214\text{–}217\) The expense associated with false-positive blood cultures has been noted earlier. These costs emphasize the importance of good technique in obtaining blood for culture.\(^218\text{,}\,219\) Distinguishing contaminants from clinically significant bacteremia is based on both microbiologic information and the patient’s clinical condition. Features of false-positive blood culture results are listed in Box 67.6.\(^189\text{,}\,198\text{,}\,220\) It would be prudent to contact discharged patients with positive blood cultures, even when contamination is suspected on a microbiologic basis, to ensure that their condition is improving.
**Fungal Cultures**

Generally, fungi are difficult to isolate in blood cultures, and it may take 4 to 6 weeks to obtain a positive yield. If fungemia is suspected, it is best to discuss culture media and technique with the laboratory before blood is taken for culture. Cultures of bone marrow are occasionally positive for mycoses when blood cultures are negative.

**PRINCIPLES AND PITFALLS IN PHLEBOTOMY FOR BLOOD TESTING**

A number of blood tests are ordered as a part of emergency practice, but by far the most common are the complete blood count (CBC) and serum chemistries. Many medications, substances, and diseases have been identified as causes of hematologic abnormalities besides the familiar categories of infectious, inflammatory, stress-related, neoplastic, and hematopoietic processes. These include antibiotics (especially sulfonamides), antineoplastic and therapeutic drugs, immuno-suppressives, and toxins (mercury and black widow spider envenomation causing leukocytosis and arsenicals causing leukopenia). There are also substances and disease processes that cause purely artifactual errors by interfering with the equipment or procedures used to perform the tests. Examples include in vivo and in vitro hemolysis, cellular clumping, and markedly elevated platelet, leukocyte, or triglyceride levels, all of which can perturb proper functioning of the machinery used to perform blood assays. The less common causes of laboratory abnormalities (both pathophysiologic and artifactual) are legion. It is necessary for most clinicians, when encountering a confirmed laboratory abnormality, to resort to standard reference texts or online sources to review potential causes or sources of error. Some of the analytical causes of false laboratory values on the CBC are listed in Table 67.8.

Before checking for obscure or uncommon causes of laboratory abnormalities, one should bear in mind that the most common source of error in laboratory blood tests is in the preanalytical phase. Improper collection techniques are the most common cause of errors in all categories.221–224 This is the part of the laboratory process that actively involves the practicing clinician. Preanalytical errors have been broken down into problems with specimen loss and handling, clotting, hemolysis, inadequate volume, and patient identity.225 In one series, more than 90% of these errors related to specimen collection.223 These key components and their most common pitfalls are as follows:

1. **Preparation of the site.** The importance of aseptic technique in the preparation of a site to draw blood for culture has been discussed. For hematologic and serum analysis, enough time should be allowed for drying of the alcohol because trace amounts can cause hemolysis. Povidone-iodine can cause errors in several chemistry assays. When used, it should be cleaned off with alcohol, as described in the section on blood cultures.

2. **Venous occlusion.** Both intracellular and chemical changes start occurring in blood as soon as a tourniquet is applied, not, as is often thought, only after it has passed through a needle into a specimen container. Serum potassium levels may increase by 6% in a vessel that has been occluded for only 3 minutes.224 Venous access sites, if visually identifiable, should be prepared before placement of a tourniquet. After tourniquet application, phlebotomy and sample acquisition should proceed as rapidly as possible, ideally within 30 seconds.225 Pumping the fist should be avoided if possible because it increases serum concentrations of potassium, lactate, and phosphate.224

3. **Routine phlebotomy.** Smaller-gauge needles and higher suction pressure are associated with hemolysis. Overexuberant application of suction on a syringe is doubly counterproductive because in addition to causing hemolysis, the needle is likely to be occluded by the wall of the vein. This increases the likelihood of unsuccessful phlebotomy and iatrogenic injury to the patient. When applying negative or positive pressure to a syringe, a given amount of force on the plunger causes higher pressure within the chamber of a smaller-diameter syringe (pressure is proportional to 1/radius^3). Blood drawn into either a standard Vacutainer or a syringe via a butterfly needle and tubing causes similar hemolysis rates (Fig. 67.6).226 If a phlebotomy site is tenuous, specimens are obtained in order of clinical priority. In most cases, tubes should be filled in the following order to avoid cross-contamination of chemicals: blood cultures, red, blue, speckled red, green, lavender, gray.225 Consistent with experience, the perception of difficult access or phlebotomy is associated with higher hemolysis rates.227,228

**Table 67.8** Artifactual Causes of False Values of the CBC

<table>
<thead>
<tr>
<th>ARTIFACTUAL INCREASE</th>
<th>ARTIFACTUAL DECREASE</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>White Blood Cell Count</strong></td>
<td>Nucleated red blood cells</td>
</tr>
<tr>
<td><strong>Hemoglobin and Hematocrit</strong></td>
<td>Severe leukocytosis (&gt; 30,000), hyperlipemic serum, giant platelets, cryoproteins</td>
</tr>
</tbody>
</table>

*Improper collection techniques are the most common cause of errors in all categories. CBC, Complete blood count; RBC, red blood cell.
4. Phlebotomy through a freshly placed IV catheter. Routine
phlebotomy typically causes a hemolysis rate of less than
2%.221 Reported hemolysis rates in ED patients are often
7% to 15%,228–230 This may be due to any or all of the
reasons considered here, but one contributing factor is the
number of blood specimens drawn in the ED through a
catheter being placed for therapeutic use (Fig. 67.7). Several
studies have shown that this arrangement significantly
increases rates of hemolysis.228,230,231 The popularity of this
technique in emergency care is most probably due to limita-
tions of personnel and temporal resources and an attempt
to enhance patient comfort by avoiding an extra needlestick.
As tourniquet times are probably longer with catheter
placement than with simple phlebotomy, it is likely that
this contributes to hemolysis, although it has never been
experimentally verified. Perceived ease of blood aspiration,
larger-bore catheters, and small aliquots drawn through the
catheter are associated with lower rates of hemolysis.230,231
Laboratory results of specimens obtained in this manner
are accurate within clinically acceptable margins of error.
5. Phlebotomy through an established IV catheter. An estab-
lished IV line appears to be a reliable source of blood for
analysis, although success rates are lower and hemolysis
rates are higher than for phlebotomy from a fresh site.212,233
Box 67.7 lists common blood tests that are accurate when
drawn through an established IV line.233,234 As with the use
of a freshly placed catheter, results are accurate within clinical
tolerances.232,235 If fluids are being administered through
the catheter, it should be turned off for at least 3 minutes
before the application of a tourniquet for phlebotomy. A
typical 18-gauge 30-mm-long plastic IV catheter without
a heparin well has a volume of 0.07 mL. The heparin well
adds a volume of 0.05 mL to make a combined volume of
0.12 mL. Thus, when using a syringe directly attached to
the heparin well (IV tubing detached), just 1 mL of aspirated
fluid should have replaced the entire volume of the heparin
well at least six times. This is demonstrated in a 2013 study
that showed consistent and accurate results with aspiration
and discard of 1 mL of blood through a catheter that
was also attached to a 6-inch extension tube.234 For a catheter
that is being used for active infusion, a volume of 1.6 mL
should be discarded.235 This suggests that the standard
practice of discarding 3 mL to 5 mL is wasteful and a
potential cause of iatrogenic anemia. These studies do not
support the use of a “push-pull” technique (aspirating a
small volume into the syringe and reinjecting prior to
drawing blood to be discarded, out of concern for sequestered
pockets of fluid).237 Use of the standard distal port of the
IV tubing (usually 25 cm from the end) is not recommended.
The IV tubing has a volume (combined with the heparin
well and catheter) of approximately 1.6 mL, thus making
it necessary to discard large volumes of blood, in addition
to the risk of contamination from IV fluids entrained in
the specimen from the main IV line during aspiration
through the port.
6. Disposition of the specimen after phlebotomy. If blood has
been drawn into a syringe, it should be promptly decanted
into the appropriate containers for the laboratory. If it has
already started to clot, it should not be forced because this
can cause hemolysis of the specimen. As cells and platelets
are fragile, specimens requiring agitation (all except red
and speckled red-topped tubes) should be rocked gently,
not shaken. If specimens are sent to the laboratory in
pneumatic tubes, they should be surrounded by shock-
absorbing material. Artifactual increases in measured HCO3
occur fairly rapidly at room temperature.212 Progressive
hemolysis of blood occurs with prolonged standing, and it
is likely that specimens are of little utility after more than
2 to 3 hours of typical ED storage conditions. One study
showed that unrefrigerated, nonagitated samples were reliable
for up to 8 hours.238

**Figure 67.7** Phlebotomy via a freshly placed intravenous catheter
and Vacutainer system.

**Box 67.7** Accurate Values Using Venous Blood
Obtained From an Indwelling Intravenous Catheter or Saline Lock

<table>
<thead>
<tr>
<th>CBC</th>
<th>Serum Chemistry</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin</td>
<td>Sodium</td>
<td>Creatine phosphokinase</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>Potassium</td>
<td>Troponin I</td>
</tr>
<tr>
<td></td>
<td>Bicarbonate</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Chloride</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Glucose</td>
<td></td>
</tr>
</tbody>
</table>

CBC, Complete blood count.
These laboratory values will be accurate when the procedures outlined in the text are followed.

**BEDSIDE TESTS FOR GASTROINTESTINAL HEMORRHAGE**

**Detection of Blood in Stool**

Bedside fecal blood tests make use of the peroxidase-like activity of Hb. The test card is impregnated with a dye that exhibits
a blue color reaction when oxidized. The developer contains
5% hydrogen peroxide and 75% alcohol. The original test
used guaiac, but current tests use more sensitive and more
reliable quinolone compounds. The addition of hydrogen
peroxide developer solution will oxidize the dye to a blue color
in the presence of a peroxidase (e.g., Hb).

Testing for occult blood in stool is associated with false-
positive and false-negative results, but in its primary role in
emergency medical practice, the test is usually reliable in
Normal GI blood loss is limited to less than 2.5 mL/day, which translates to less than 2 mg of Hb per gram of stool (0.2% by weight). The sensitivity of the Hemoccult test varies both with the concentration of Hb present in stool and with the extent of Hb exposure to the proteolytic effects of the digestive tract. The Hemoccult test is 37% sensitive to stool containing 2.5 mg Hb/g of stool but 95% sensitive when the concentration is 20 mg Hb/g of stool, thus indicating that low to moderate levels of blood may be missed. The test is much more likely to detect lower GI hemorrhage than an identical rate of upper GI bleeding because of the 100-fold diminution in the peroxidase activity of blood during transition through the GI tract. Impaired detection of Hb may also occur as a result of dilution because of diarrheal illness. Regulatory attempts in the United States to improve quality control in bedside occult blood testing seem to have had the unintended consequence of discouraging digital rectal examination in patients for whom they are indicated. In the event of a trace positive Hemoccult test that is not the source of an emergency illness, notification to the patient and advice regarding further outpatient evaluation should not be overlooked.

**Method**

Smear the stool specimen onto the reagent area on the card and add a drop of developer. Because the reaction must occur
Testing for Gastric Blood

Heme tests designed for use on stool specimens can be unreliable when applied to gastric juices, with increasing inaccuracies being reported as the pH decreases. Although a positive test of gastric contents with a fecal Hemoccult card is likely to be accurate, a negative result with the fecal Hemoccult card does not rule out the presence of blood. The Gastroccult card uses a modified guaiac developer containing buffers to neutralize gastric acid, thereby facilitating accurate detection of Hb. The test works on the same basis as the fecal guaiac test in that it uses the properties of Hb as a peroxidase. In product testing, the Gastroccult card was 100% sensitive in detecting specimens with greater than 500 ppm of blood by volume, equivalent to 0.05% or 0.25 mL of blood in 500 mL of gastric contents.

Method

Apply a drop of gastric aspirate onto the test area and two drops of developer onto the sample. Look for the formation of a blue color within 1 minute. Do not use fecal blood test developer. In a specimen that is already a bilious green, the test is considered positive only if new blue color is formed. The Gastroccult card also contains a pH testing strip located close to the occult blood testing area, which might be useful in testing emesis after an acidic or alkaline ingestion. Inaccurate results might be anticipated in the presence of the same substances that can confound the Hemoccult test: meats, peroxidase-rich foods, and reducing substances such as ascorbic acid. The accuracy of Gastroccult should not be affected by the presence of cimetidine or sucralfate.

BLOOD GLUCOSE METERS

Bedside testing of capillary blood for glucose levels is a common procedure performed in the ED and is generally an excellent substitute for venous blood testing. A variety of meters are available and they are reasonably accurate (±10%) when a small drop of blood is electronically analyzed (Fig. 67.9). In patients with poor tissue perfusion, the accuracy of determining hypoglycemia is less precise and may vary up to approximately 5% from venous blood. In the setting of hypoperfusion, bedside measurement of whole blood is preferable. Bedside testing is also less accurate in patients with extreme hypoglycemia or hyperglycemia, but readings are sufficiently accurate to alert the clinician to very high or very low glucose levels. Fingertip capillary blood is the preferred specimen for bedside glucose meter testing. Blood from alternative sites, such as the skin of the forearm, may give slightly lower results than those taken at the fingertips as they may sample venous blood rather than capillary blood. When blood glucose concentrations are rising rapidly or falling rapidly (such as a hypoglycemic response secondary to rapidly acting insulin), blood glucose results from alternative sites may yield significantly delayed results (up to 30 minutes) when compared with finger stick readings, which are generally accurate at all time points.

Figure 67.9 Bedside glucose meters analyze capillary blood samples and are generally an excellent substitute for venous blood testing. Errors may occur in patients with poor tissue perfusion or in cases of extreme hypoglycemia or hyperglycemia. Additional errors may be due to the operator, including improper calibration, dirty meters, and improper storage of test strips.

Older glucose meters reported whole blood glucose values, which made it difficult to compare finger stick results with those from venous blood testing by the laboratory, which measures plasma glucose. Plasma glucose levels are 10% to 15% higher than whole blood glucose levels. The majority of glucose meters now available provide plasma-equivalent values rather than whole blood glucose values, so glucose meters and the results from venous blood analyzed in hospital laboratories should be comparable.

Most errors in bedside glucose testing are, however, due to operator error, including improper calibration, dirty meters, and improperly stored test strips. Meters should be calibrated frequently to ensure quality readings. Glucose strips have batch-to-batch variation. A common error in testing is due to leaving the lid off glucose strips for prolonged periods because inaccuracies on test strips can result from exposure to heat, moisture, and humidity.

DIAGNOSTIC AND THERAPEUTIC TOXICOLOGIC BEDSIDE PROCEDURES

Management of patients with altered mental status can be challenging, especially if the clinician suspects a drug overdose or poisoning. These patients often have no available history or provide an inaccurate history. Clinicians must rely heavily on the findings on physical examination and other sources of information to diagnose or confirm their clinical suspicion of poisoning or overdose. The hospital-based toxicology laboratory can be valuable in select cases. Screening tests for commonly ingested mind-altering substances are available but limited in scope. Determining concentrations of specific drugs (e.g., acetaminophen, lithium, digoxin, phenytoin) tend to be more
useful in guiding management. However, hospital-based laboratories are not equipped to perform timely analytical procedures for the thousands of possible drugs or toxins. In fact, the screening panels for drugs of abuse that most hospitals use have been shown to rarely influence medical management of adult patients in the ED. Use of these drug screens in selected pediatric patients may have more of an impact on medical management.

Diagnostic bedside testing (point-of-care testing) for specific poisons or toxins may have the advantage of being cost-effective and timely. When applied appropriately, certain bedside tests provide immediate information to the clinician and can significantly influence medical management in a timely manner. Unfortunately, with establishment of the Clinical Laboratory Improvement Amendments of 1988 (CLIA), bedside laboratory testing has undergone stricter oversight. This federal regulation has jurisdiction on any laboratory tests performed on humans, or specimens obtained from humans, and has added a layer of complexity to bedside testing. Tests that are not performed on humans, such as the Meixner test or mothball testing, are not covered by the CLIA act. This section discusses bedside diagnostic and therapeutic toxicologic procedures.

Noninvasive Diagnostic Procedures

**Amatoxin: Meixner Test**

Ingestion of several types of mushrooms (e.g., *Amanita phalloides*) can be fatal. The most poisonous of these mushrooms contain amatoxins. Patients who have ingested amatoxins often complain of GI symptoms consisting of nausea, vomiting, diarrhea, and abdominal cramping beginning 6 to 8 hours after ingestion. They often bring in specimens of the mushrooms chopped, crushed, cooked, or mixed with stool or gastric contents. Standard hospital laboratories cannot confirm or exclude the diagnosis of amatoxin poisoning; therefore treatment decisions to be effective with other bodily secretions, such as blood or urine.

**Mothball Identification**

At present, commercial mothballs are composed of either nontoxic paradichlorobenzene or possibly toxic naphthalene. Naphthalene can cause a hemolytic reaction in neonates and patients with glucose-6-phosphate dehydrogenase (G6PD) deficiency.255 In the past, mothballs have also been produced from camphor, which can cause central nervous system depression and seizures. Fortunately, camphor mothballs are no longer commercially available in the United States, although they may still exist in older households and may be obtained in other countries. Rapid differentiation between these groups of mothballs can expedite patient management and disposition. Several bedside tests have been reported to facilitate this.

1. **Paradichlorobenzene** is heavier than naphthalene. In turn, naphthalene is heavier than camphor. In lukewarm tap water, camphor will float and naphthalene and paradichlorobenzene will sink. In a solution of 3 tbsp of table salt thoroughly dissolved in 4 oz of lukewarm water, camphor and naphthalene will float and paradichlorobenzene will sink.

2. **Paradichlorobenzene** has a lower melting point than naphthalene does. Paradichlorobenzene mothballs will melt in a water bath at 33°C, whereas naphthalene requires a water bath hotter than 80°C.

![Figure 67.10](http://blog.mycology.cornell.edu/) The Meixner test is a crude assay for amatoxins. Place portions of the unknown dried mushroom on low-grade newsprint and add 10-N hydrochloric acid. The dried-up mushroom (left) is *Galerina marginata*, which yields a blue reaction (positive, probably deadly poisonous); the little brown mushroom (right) does not (negative, toxicity uncertain). This test is the only one readily available but has varying accuracy and depends on the paper being used (regular newsprint is shown here). (Courtesy Kathie T. Hodge and Kent Loeffler [photographer], Cornell University. Available at http://blog.mycology.cornell.edu.)
3. Paradichlorobenzene is described as wet and oily, whereas naphthalene is described as having a dry appearance. Paradichlorobenzene is familiar to many people as a cake of disinfectant used in urinals and diaper pails.

**Body Secretion Analysis**
Careful analysis of bodily secretions, the odor emanating from poisoned patients, and the color of their urine can help identify certain toxins. Some characteristic smells and urine colors are listed in Table 67.9 and Box 67.8.

**Bedside Toxicologic Tests on Urine**

**Ethylene Glycol**
Evaluation of the urine of patients who may have ingested ethylene glycol can be helpful. Urine should be tested for fluorescence (an additive in many commercial antifreeze products) under an ultraviolet light and for the presence of calcium oxalate crystals (a metabolic by-product of ethylene glycol metabolism).

The presence of calcium oxalate crystals (either envelope-shaped calcium dihydrate or needle-shaped calcium monohydrate) in urine on microscopic inspection is indicative of high oxalate levels in serum (Fig. 67.11; see also Fig. 67.3F). Calcium monohydrate crystals can easily be confused with sodium urate crystals; therefore the presence of the dihydrate crystal tends to be more specific for ethylene glycol ingestion. Lack of these crystals does not rule out significant ethylene glycol ingestion because excretion of these crystals may occur late in the ingestion (> 6 hours) and occasionally does not occur at all.²⁵⁶,²⁵⁷

Visual inspection of urine under a Wood lamp or ultraviolet light to ascertain fluorescence may also be helpful in the diagnosis of ethylene glycol exposure. Antifreeze is the most common source of ingested ethylene glycol. Fluorescein, the actual fluorescing material, is often placed in commercially available antifreeze to enable mechanics to detect radiator leaks with a Wood lamp or other ultraviolet light source. Fluorescein is a nontoxic, inert vegetable dye that is eliminated unchanged in urine. Therefore high levels of fluorescein in urine suggest significant ethylene glycol ingestion. However, lack of fluorescein does not rule out a significant exposure because not all antifreezes contain fluorescein or high concentrations of fluorescein in relation to ethylene glycol. False-positive findings can occur if certain plastic urine containers are used.²⁵⁸

To perform this procedure, place the test urine sample and control samples into separate glass test tubes. Inspect for fluorescence under a Wood lamp in a dark room. Always perform this test with controls that include urine that does not contain fluorescein and urine that does contain fluorescein. The use of controls may increase the sensitivity and specificity from 49% and 75%, respectively, to a sensitivity and specificity of 100%.²⁵⁹ Fluorescein is readily available because fluorescein-containing strips are commonly used in ophthalmologic procedures (see Chapter 62).

**Salicylates**
Several bedside tests have been developed to qualitatively detect salicylates in urine, including 10% ferric chloride solution, Trinder’s solution, and Phenistix reagent strips. All are rapid, inexpensive, sensitive tests that give a qualitative rather than a quantitative result.

Ferric chloride and Trinder’s solution both have sensitivities of 100% with serum salicylate levels of 5 mg/dL. False positives can occur with both tests. Acetoacetic acid, acetone, and phenylpyruvic acid will cause false-positive results. Thus, this test may be falsely positive in patients with diabetic, alcoholic, or starvation ketoacidosis. Phenol-containing drugs such as diflunisal, sulfasalazine, and salicylamide may also produce false positives. Any positive result requires a confirmatory quantitative serum salicylate assay.²⁶⁰

The ferric chloride test is a commonly used rapid, qualitative, urinary screening procedure. To perform this test, add several drops of 10% ferric chloride to 1 or 2 mL of urine that has been collected in a test tube. The immediate appearance of a bluish-purple color signifies that salicylates are present in the urine (Fig. 67.12). This test is very sensitive, and just one aspirin tablet taken within 12 to 24 hours will give a positive

---

**Figure 67.11** A, Monohydrate (needle- or prism-shaped) and dihydrate (envelope- or tent-shaped) calcium oxalate crystals from a patient poisoned with ethylene glycol. The dihydrate form is more specific for ethylene glycol toxicity because the monohydrate form can easily be confused with urate crystals. B, Calcium dihydrate crystal (arrow) and a pseudocast of a crystal.
Phenistix reagent strips were originally developed to detect phenylketonuria. However, Phenistix strips also turn brown in the presence of salicylates. False-positive findings for salicylates can occur if phenothiazines are present.

**Bedside Toxicologic Tests on Oral Secretions and Breath: Ethyl Alcohol**

Several bedside devices have been developed to measure alcohol concentrations in body fluids. Measurements of the alcohol result. It will require 60 to 120 minutes from the time of ingestion for this reaction to become positive in patients with normal renal function, so early test results may be misleading.

The Trinder test uses a mixture of mercuric chloride and ferric nitrate in deionized water. To perform this test, mix 1 mL of urine with 1 mL of Trinder's solution. A violet or purple color signifies the presence of salicylates. Acetoacetic acid and high levels of phenothiazines may give false-positive results.

<table>
<thead>
<tr>
<th>CHARACTERISTIC ODOR</th>
<th>RESPONSIBLE DRUG OR TOXIN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetone (sweet, fruity; pearlike)</td>
<td>Lacquer, ethanol, isopropyl alcohol, chloroform, diabetic ketoacidosis, alcoholic ketoacidosis, trichloroethane, paraldehyde, chloral hydrate, methylbromide, <em>Pseudomonas</em> infection</td>
</tr>
<tr>
<td>Alcohols</td>
<td>Ethanol (congeners), isopropyl alcohol</td>
</tr>
<tr>
<td>Ammonia-like</td>
<td>Uremia</td>
</tr>
<tr>
<td>Automobile exhaust</td>
<td>Carbon monoxide (odorless, but associated with exhaust)</td>
</tr>
<tr>
<td>Beer (stale)</td>
<td>Scrofula</td>
</tr>
<tr>
<td>Bitter almond</td>
<td>Cyanide</td>
</tr>
<tr>
<td>Carrots</td>
<td>Cicutoxin (or water hemlock)</td>
</tr>
<tr>
<td>Coal gas (stove gas)</td>
<td>Carbon monoxide (odorless, but associated with coal gas)</td>
</tr>
<tr>
<td>Disinfectants</td>
<td>Phenol, creosote</td>
</tr>
<tr>
<td>Eggs (rotten)</td>
<td>Hydrogen sulfide, carbon disulfide, mercaptans, disulfiram, N-acetylcysteine</td>
</tr>
<tr>
<td>Feculent</td>
<td>Intestinal obstruction</td>
</tr>
<tr>
<td>Fish or raw liver (musty)</td>
<td>Hepatic failure, zinc phosphide, hypermethioninemia, trimethylaminuria</td>
</tr>
<tr>
<td>Fruitlike</td>
<td>Nitrites (e.g., amyl, butyl), ethanol (congeners), isopropyl alcohol</td>
</tr>
<tr>
<td>Garlic</td>
<td>Phosphorus, tellurium, arsenic, parathion, malathion, selenium, dimethyl sulfoxide, thallium</td>
</tr>
<tr>
<td>Halitosis</td>
<td>Acute illness, poor oral hygiene</td>
</tr>
<tr>
<td>Hay</td>
<td>Phosgene</td>
</tr>
<tr>
<td>Mothballs</td>
<td>Naphthalene, <em>p</em>-dichlorobenzene, camphor</td>
</tr>
<tr>
<td>Peanuts</td>
<td><em>N</em>-3-pyridyl-methyl-<em>N</em>-p-nitrophenyl urea (<em>Vacor</em>)</td>
</tr>
<tr>
<td>Pepper-like</td>
<td>O-chlorobenzylidene malononitrile</td>
</tr>
<tr>
<td>Putrid</td>
<td>Anaerobic infections, esophageal diverticulum, lung abscess, scurvy</td>
</tr>
<tr>
<td>Rope (burned)</td>
<td>Marijuana, opium</td>
</tr>
<tr>
<td>Shoe polish</td>
<td>Nitrobenzene</td>
</tr>
<tr>
<td>Sweating feet</td>
<td>Isovaleric acid acidemia</td>
</tr>
<tr>
<td>Tobacco</td>
<td>Nicotine</td>
</tr>
<tr>
<td>Vinegar</td>
<td>Acetic acid</td>
</tr>
<tr>
<td>Vinyl-like</td>
<td>Ethchlorvynol (Placidyl)</td>
</tr>
<tr>
<td>Violets</td>
<td>Turpentine (metabolites excreted in urine)</td>
</tr>
<tr>
<td>Wintergreen</td>
<td>Methyl salicylate</td>
</tr>
</tbody>
</table>

Alcohol concentrations in saliva have been shown to correlate well with serum concentrations. Bedside measurement of salivary alcohol concentrations can also be obtained with a dipstick-like device. These devices use an enzymatic reaction involving alcohol dehydrogenase to measure alcohol concentrations.

Patients who are dehydrated (a common occurrence in alcohol-intoxicated patients) are often unable to provide adequate saliva samples, and inaccurate readings have occurred in patients with high blood alcohol concentrations.

**BOX 67.8 Drugs That Color Urine**

**YELLOW**  
Quinacrine (Atabrine) in acidic urine  
Riboflavin (large doses)

**YELLOW-GREEN**  
Methylene blue; see Blue

**YELLOW-ORANGE**  
Fluorescein sodium

**YELLOW-PINK**  
Cascara in alkaline urine; see Yellow-Brown, Brown, Black  
Senna in alkaline urine; see Yellow-Brown, Brown

**YELLOW-BROWN**  
Cascara in acidic urine; see Yellow-Pink, Brown, Black  
Nitrofurantoin (Furadantin [Custom Pharmaceuticals Ltd., East Sussex, United Kingdom] and others); see Brown

**ORANGE**  
Phenazopyridine (Pyridium, Marlex Pharmaceuticals, Inc., New Castle, DE); see Red

**ORANGE-RED**  
Rifampicin (Rifadin, Rifamycin, Rimactane)

**PINK**  
Phenothiazines; see Red, Red-Brown  
Phenytoin (Dilantin, Pfizer Inc., New York, NY); see Red, Red-Brown

**RED**  
Antarquinone in alkaline urine  
Deferoxamine (Desferal, Novartis Pharmaceuticals UK Limited, Surrey, United Kingdom)  
Methyldopa (Aldomet); see Brown, Black  
Phenazopyridine (Pyridium, Marlex); see Orange  
Phenothiazines; see Pink, Pink-Brown  
Phenytoin (Dilantin, Pfizer); see Pink, Red-Brown

**RED-PURPLE**  
Phenacetin; see Brown

**RED-BROWN**  
Phenothiazines; see Pink, Red  
Phenytoin (Dilantin, Pfizer); see Pink, Red

**BROWN**  
Cascara in alkaline urine; see Yellow-Brown, Yellow-Pink, Black  
Levodopa (Dopar)  
Methocarbamol (Robaxin, Pfizer); see Green, Black  
Metronidazole (Flagyl, Famar Health Care Services Madrid S.A.U., Madrid, Spain)  
Methyldopa (Aldomet); see Red, Black  
Nitrofurantoin (Furadantin [Custom Pharmaceuticals] and others); see Yellow-Brown  
Phenacetin; see Red-Purple  
Quinina; see Black  
Senna in alkaline urine on standing; see Yellow-Brown, Yellow-Pink

**BLUE**  
Methylene blue; see Green  
Triamterene (Dyrenium, Concordia Pharmaceuticals Inc., St. Michael, Barbados), fluorescent

**BLUE-GREEN**  
Amitriptyline (Elavil, Endep)

**GREEN**  
Indomethacin (Indocin, MERCK & CO., Inc., Whitehouse Station, NJ) from liver damage  
Methocarbamol (Robaxin, West-Ward Pharmaceuticals, Eatontown, NJ); see Brown-Black

**BLACK**  
Cascara in alkaline urine on standing; see Yellow-Brown, Yellow-Pink, Brown  
Iron sorbitex (Jectofer); see Brown  
Methocarbamol (Robaxin, West-Ward); see Brown, Green  
Methyldopa (Aldomet); see Red, Brown  
Quinina; see Brown

*The drug imparts more than one color to urine and is listed under each color that it adds.*


Concentration in expired air or saliva have been shown to correlate well with blood alcohol concentrations in the appropriate settings.

Breath alcohol analyzers have been developed since the 1950s and are currently used in law enforcement. These devices typically use infrared spectral analysis to determine the concentration of alcohol in expired air. Almost all the alcohol found in expired air at the level of the mouth is secondary to alcohol diffused from the bronchial system rather than the alveolar system. Minor alterations in breathing patterns can cause large variations in readings. Thus, uncooperative patients who do not exhale properly will cause an inaccurate reading. Other causes of inaccurate readings include the recent ingestion of alcohol-containing products, belching or vomiting, use of inhalers, poor technique, or restrictive pulmonary pathology.

Alcohol concentrations in saliva have been shown to correlate with serum concentrations. Bedside measurement of salivary alcohol concentrations can also be obtained with a dipstick-like device. These devices use an enzymatic reaction involving alcohol dehydrogenase to measure alcohol concentrations. Patients who are dehydrated (a common occurrence in alcohol-intoxicated patients) are often unable to provide adequate saliva samples, and inaccurate readings have occurred in patients with high blood alcohol concentrations.

**Bedside Toxicologic Tests on Blood: Methemoglobinemia**

Patients with methemoglobinemia will often have a normal partial pressure of oxygen on routine arterial blood gas analysis, a normal calculated Hb saturation, a nondiagnostic pulse...
is quite accurate. At this level, almost all patients are obviously methemoglobinemia by visual inspection of the color of blood in patients. With levels of 35% or greater, identification of greater than 15% are reported to cause a cyanotic appearance a false-negative reading 50% of the time. Methemoglobin levels of between 12% and 14% may cause Methemoglobin levels of less than 10% may alter the color color and may therefore have a degree of interobserver variance. relies on the ability of the examiner to distinguish changes in globinemia appears darker or chocolate brown. This method molecule.

To evaluate for methemoglobinemia, place a drop of sample blood on a white background (a white coffee filter is appropriate) in a well-lit environment. Next to this, place a drop of normal blood as a comparison control sample. Blood with methemoglobinemia appears darker or chocolate brown. This method relies on the ability of the examiner to distinguish changes in color and may therefore have a degree of interobserver variance. Methemoglobin levels of less than 10% may alter the color of blood only slightly and thereby cause a false-negative finding. Methemoglobin levels of between 12% and 14% may cause a false-negative reading 50% of the time. Methemoglobin levels greater than 15% are reported to cause a cyanotic appearance in patients. With levels of 35% or greater, identification of methemoglobinemia by visual inspection of the color of blood is quite accurate. At this level, almost all patients are obviously cyanotic and symptomatic.

Invasive Diagnostic Procedures

Several invasive diagnostic bedside procedures can be useful in the assessment of possible drug overdoses. The basic premise of these procedures is that patients who have been exposed to a certain drug or poison will respond in a particular fashion if given a diagnostic challenge dose of another particular drug or true antidote.

Naloxone

Naloxone hydrochloride (Narcan, Adapt Pharma, Inc., Radnor, PA) is a µ-opioid receptor antagonist. A diagnostic challenge with IV naloxone has been recommended for all patients with central nervous system depression. Certain clinical findings such as miosis, decreased respiratory rate, and evidence of illicit drug use can predict many patients who will respond to a challenge dose of naloxone. If a patient's mental status improves significantly after a dose of naloxone, the patient should be considered to have been exposed to an opioid-like substance. This is true even if a laboratory drug screen is negative for opioids. Furthermore, the routinely used immunoassay drug screen may not detect many of the synthetic opiates, such as fentanyl and fentanyl derivatives, tramadol, meperidine, methadone, and buprenorphine. Although cases have been reported of patients with nonopioid overdoses (such as alcohol or phencyclidine) responding to naloxone, these single observations have not been confirmed in controlled animal or human studies.

The traditional challenge dose of naloxone in an adult or child is 2 mg every 2 minutes intravenously until a response is achieved or 10 mg is given. Some clinicians prefer to use much smaller doses (0.1 to 0.2 mg) and titrate to effect. This may partially reverse the opioid overdose–related symptoms and confirm the diagnosis without precipitating a withdrawal syndrome in patients with opioid dependency. Most patients with an opioid overdose will exhibit some response to 1 to 4 mg of naloxone, but some massive overdoses might require larger doses. A patient who does not respond at all to 10 mg of naloxone probably does not have a pure opioid overdose. High doses of naloxone may be needed to reverse many synthetic opiates, such as buprenorphine and methadone. Lower doses can be given (0.4 to 0.8 mg in adults or 0.01 mg/kg in children) to reverse known opioid-induced respiratory depression without reversing the analgesia. Because naloxone has a half-life of between 30 and 60 minutes, a continuous drip of naloxone can be used to avoid resedation. A reasonable choice is to use two-thirds of the initial bolus dose that achieved the desired reversal effect as the hourly IV dose. For example, a patient who satisfactorily responded to 1.5 mg of naloxone might receive a naloxone solution of 10 mg of naloxone in 500 mL of normal saline at a rate of 1 mg (50 mL) per hour intravenously. Nalmefene, a long-acting opioid receptor antagonist that has a terminal half-life of roughly 11 hours, can also be given to patients with suspected overdose. Theoretically, a single dose of nalmefene will be effective longer than the effects of heroin or most abused opiate substances. The initial recommended dose is 1.0 to 1.5 mg intravenously.

Naloxone and nalmefene have minimal significant side effects, but they can precipitate withdrawal in patients addicted to opioids. Unlike alcohol withdrawal, naloxone-induced opioid withdrawal in adults is short-lived and not usually life-threatening. Withdrawal can be avoided if lower initial doses of naloxone or nalmefene are given and then slowly titrated upward to the desired effect.

Resedation may occur if the ingested drug (e.g., methadone, oxycodone [OxyContin, Purdue Pharma L.P.], morphine sulfate [MS Contin, Purdue Pharma L.P.]) has a clinical effect longer than that of naloxone. Reported drug half-lives may have significant variability in the clinical setting. If no narcotic effect is evident in 60 to 120 minutes after standard doses of naloxone (common with heroin, for example), no clinically significant resedation is expected. Larger naloxone doses may prolong the expected antidote effect of naloxone, and longer observation is required. All timing and dose recommendations are guidelines, and all clinical decisions with regard to resedation should be individualized.
**Flumazenil**

Flumazenil (Romazicon, Roche Pharmaceuticals, Nutley, NJ) is a competitive benzodiazepine receptor antagonist that has the ability to reverse the central nervous system depression caused by all currently commercially available benzodiazepines. However, flumazenil is no longer recommended as empiric treatment (part of the coma cocktail) of all sedated patients. The most supported use of flumazenil is to reverse excessive physician-initiated conscious sedation with benzodiazepines and for children with suspected benzodiazepine overdose. Its routine use in the setting of possible benzodiazepine overdose is controversial but is supported as a diagnostic and therapeutic agent in selected cases. Unlike naloxone, flumazenil can have significant side effects, but only in certain subsets of patients. Complications include precipitation of seizures or a withdrawal syndrome in benzodiazepine-dependent patients. To minimize the chance of seizures, flumazenil should be avoided in known benzodiazepine-dependent patients and those who may have ingested epileptogenic drugs (e.g., cyclic antidepressants, cocaine, theophylline, or isoniazid).

In suspected benzodiazepine overdoses in which patients are obtunded and have no history of seizures or suspicion of involvement of epileptogenic agents, flumazenil can be administered intravenously at a dose of 0.2 to 0.5 mg/min. Most benzodiazepine-overdosed patients show improvement in mental status with 1 mg of flumazenil and almost all respond to 3 to 5 mg. Small, escalating doses given slowly (maximally, 0.5 mg/min) have been recommended. Larger doses can be given at one time as a bolus, although this increases such side effects as anxiety, agitation, and emotional lability; it also increases the chances of precipitating withdrawal in benzodiazepine-dependent patients. Fortunately, seizures that occur after flumazenil use are usually transient. In rare cases, higher doses of benzodiazepines, barbiturates, and phenytoin might be required.

If a patient responds to flumazenil with an improvement in depressed mental status, this suggests only that the patient is under the influence of a benzodiazepine. Flumazenil can partially reverse the effects of many of the newer nonbenzodiazepine sleeping agents that affect the γ-aminobutyric acid pathway, such as zolpidem, zopiclone, and eszopiclone. Flumazenil can improve mental status in patients with hepatic encephalopathy. It does not have any significant effect on alcohol, barbiturates, and other nonbenzodiazepine sedative-hypnotics.

Resedation is possible if the ingested drug has a clinical duration longer than that of flumazenil. If no resedation has occurred 60 to 90 minutes after standard doses of flumazenil, clinically significant resedation is not expected. If higher flumazenil doses have been used, additional observation may be warranted. All timing and dose recommendations are guidelines, and all clinical decisions with regard to resedation should be individualized.

**Physostigmine**

Physostigmine is an acetylcholinesterase inhibitor that can penetrate into the central nervous system and thus can reverse both the central and peripheral effects of anticholinergic agents. In the majority of patients with anticholinergic toxicity, no laboratory tests are available to rapidly confirm the diagnosis, and testing for specific drugs is limited or unavailable. A clinical picture that may consist of mydriasis, dry and flushed skin, dry mucous membranes, urinary incontinence, absent bowel sounds, tachycardia, hyperthermia, hallucinations, agitation, and seizures suggests an anticholinergic syndrome. In some cases (low-dose antihistamines and others), only a central nervous system syndrome characterized by hallucinations, agitation, and confusion exists. A dramatic response to physostigmine frequently confirms a diagnosis of anticholinergic toxicity. In these patients, physostigmine often decreases the degree of agitation and confusion. The use of physostigmine as a diagnostic challenge can be helpful in selected situations, but similar to flumazenil, routine use of physostigmine as a diagnostic bedside challenge in all obtunded patients, especially in those without anticholinergic findings, should be discouraged. In some cases the agitation produced by potent anticholinergics, such as scopolamine, can be resistant to benzodiazepines. Judicious use of physostigmine is warranted to avoid excessive sedation, chemical paralysis, or impaired ventilation.

As a diagnostic challenge or therapeutic intervention, physostigmine can be administered intravenously under constant cardiac monitoring at a dose of 1 to 2 mg in adults and 0.02 mg/kg in children over a 3- to 5-minute period. It will take 3 to 6 minutes for the central nervous system effect to become apparent. Some clinicians empirically pretreat with a benzodiazepine to prevent possible seizures. Because the half-life of physostigmine is 30 to 60 minutes, a repeated dose of 0.5 to 2 mg can also be given as clinically indicated.

Similar to flumazenil, physostigmine has been reported to interact detrimentally with cyclic antidepressants. In this setting, physostigmine has been associated with life-threatening dysrhythmias. Physostigmine can also cause an excess of acetylcholine and a resultant cholinergic crisis. This syndrome includes salivation, lacrimation, urination, defecation, bradycardia, bronchorrhea, and seizures. For this reason, 1 mg of atropine intravenously should be readily available to reverse potential cholinergic excess when using physostigmine.

**Deferoxamine**

Deferoxamine is an organic compound derived from the bacterium *Streptomyces pilosus* and can chelate iron. It can be used as a therapy or as a diagnostic challenge in patients with iron overdoses. Patients who have unstable vital signs or significant GI or central nervous system symptoms usually require therapeutic doses of deferoxamine. Asymptomatic patients with a history of iron overdose typically require supportive care only. Patients with persistent but mild symptoms, such as vomiting and diarrhea, may be given a diagnostic challenge dose of deferoxamine. A diagnostic challenge is preferential over ancillary laboratory testing because tests such as iron levels and total iron-binding capacity in the setting of iron overdose can be inaccurate, misleading, and time-consuming.

A diagnostic challenge dose of deferoxamine is administered intramuscularly or intravenously over a 45-minute period at doses of 40 to 90 mg/kg up to a maximum of 1 g in children and 2 g in adults. Deferoxamine can also be administered intravenously as a constant infusion of 15 mg/kg per hour. A positive result occurs when chelated iron in the form of ferric oxide appears in the urine. This usually causes the urine to turn a reddish orange or “vin rose” color in 2 to 3 hours after the initiation of treatment. The change in color is qualitative only and has no prognostic significance. The color change caused by ferrioxamine is dependent on pH and concentration, and false-negative test results can occur.

Chronically administered deferoxamine has been reported to have multiple adverse effects, such as acute respiratory distress...
syndrome, visual defects, and enhancement of *Yersinia enterocolitica* infection. In the setting of a single challenge dose, flushing, erythema, tachycardia, urticaria, and hypotension caused by rapid administration of deferoxamine are the most serious side effects.

**Invasive Therapeutic Procedures**

The indications and rationale for the use of certain therapeutic procedures in toxicology are often misunderstood.

**Alkalization of Urine and Blood**

Alkalization of urine consists of manipulating the pH of urine to enhance the excretion of certain drugs (Box 67.9). Weak acids remain in ionic form in a basic milieu. The ionic form often prevents reabsorption of that drug in the proximal tubule, and urinary alkalization can therefore promote elimination in urine. For certain drugs this can play a significant role in their elimination. For example, salicylate elimination increases proportionately to the urinary flow rate, but it increases exponentially with increases in urinary pH. The increased serum clearance attributed to alkalization does not correlate well with outcome or length of hospitalization.

Recommendations differ on the actual method or formula to achieve urinary alkalization. No body of literature supports one method of urinary alkalization over another. In general, this procedure should be titrated to the patient’s fluid and

---

**TABLE 67.10 Use of Ethanol for Methanol or Ethylene Glycol Poisoning**

<table>
<thead>
<tr>
<th>Intravenous Ethanol: Loading Dose (Using a 10% Ethanol Solution) a</th>
<th>Volume of Loading Dose (Given Over 1-2 hr as Tolerated)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loading dose of 1000 mg/kg of 10% ethanol (infused over 1–2 hr as tolerated); assumes a zero-ethanol level to start. Aim is to produce a serum ethanol level of 100–150 mg/dL.</td>
<td>10 kg</td>
</tr>
<tr>
<td>100 mL</td>
<td>150 mL</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Oral Ethanol: Loading Dose (a 20% Volume/Volume Concentration Yields Approximately 200 mg/mL)</th>
<th>Volume of Loading Dose</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loading dose of 1000 mg/kg of 20% ethanol b diluted in juice; may be administered orally or via nasogastric tube; assumes a zero ethanol level to start. Aim is to produce a serum ethanol level of 100–150 mg/dL.</td>
<td>10 kg</td>
</tr>
<tr>
<td>50 mL</td>
<td>75 mL</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Intravenous Ethanol: Maintenance Dose (Using a 10% Ethanol Solution) c</th>
<th>Infusion Rate (mL/hr for Various Weights)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal Maintenance Range (mg/kg/hr)</td>
<td>10 kg</td>
</tr>
<tr>
<td>80</td>
<td>8</td>
</tr>
<tr>
<td>110</td>
<td>11</td>
</tr>
<tr>
<td>130</td>
<td>13</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Approximate Maintenance Dose for Chronic Alcoholics</th>
<th>15 kg</th>
<th>30 kg</th>
<th>50 kg</th>
<th>70 kg</th>
<th>100 kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>150 e</td>
<td>15</td>
<td>22</td>
<td>45</td>
<td>75</td>
<td>105</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Range Required during Hemodialysis</th>
<th>25 kg</th>
<th>30 kg</th>
<th>35 kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>250 e</td>
<td>25</td>
<td>38</td>
<td>75</td>
</tr>
<tr>
<td>300 e</td>
<td>30</td>
<td>45</td>
<td>90</td>
</tr>
<tr>
<td>350 e</td>
<td>35</td>
<td>53</td>
<td>105</td>
</tr>
</tbody>
</table>

---

*Continued*
The formula is to begin with a bolus of two ampules of IV sodium magnesium to the diuresis fluid. Some will empirically add potassium and the infusion rate of ethanol must be increased during dialysis. Prolonged ethanol administration may lead to hypoglycemia. Note that 10% ethanol for infusion may be difficult to achieve. Urinary alkalinization may be difficult, if not impossible, to achieve. Other theoretical causes are relative hypocalcemia, fluid overload, and alkalemia. All of these are potential adverse effects of aggressive urinary alkalinization. The actual amount of fluids and bicarbonate administered requires titration to the patient's clinical condition, and careful monitoring of electrolyte, pH, and fluid status is encouraged.

Urinary alkalinization can sometimes be difficult to achieve or maintain. Hypovolemia is probably the leading cause of an inability to achieve alkaline urine. Other theoretical causes are hypokalemia, hypomagnesemia, and hypochloremia. Several authors have suggested that in patients with severe salicylate poisoning, urinary alkalinization may be difficult, if not impossible, to achieve. Some will empirically add potassium and magnesium to the diuresis fluid.

### TABLE 67.10 Use of Ethanol for Methanol or Ethylene Glycol Poisoning—cont’d

<table>
<thead>
<tr>
<th>Oral Ethanol: Maintenance Dose (a 20% Volume/Volume Concentration Yields Approximately 200 mg/mL; Infusion to be Given Each Hour Immediately Following a Loading Dose; Aim is to Maintain a Serum Ethanol Level of 100–150 mg/dL; Each Dose may be Diluted in Juice and Given Orally or via Nasogastric Tube)</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Infusion Rate (mL/hr for Various Weights)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal Maintenance Range (mg/kg/hr)</td>
</tr>
<tr>
<td>80</td>
</tr>
<tr>
<td>110</td>
</tr>
<tr>
<td>130</td>
</tr>
<tr>
<td>Approximate Range for Chronic Alcoholics or for Patients Receiving Continuous Oral Activated Charcoal</td>
</tr>
<tr>
<td>150</td>
</tr>
<tr>
<td>Range Required During Hemodialysis</td>
</tr>
<tr>
<td>250</td>
</tr>
<tr>
<td>300</td>
</tr>
<tr>
<td>350</td>
</tr>
</tbody>
</table>

- If a 5% ethanol solution is used, double the volume of the loading dose.
- Equivalent to a 40-proof solution.
- If a 5% ethanol solution is used, double the volume rate; monitor closely for potential volume overload.
- Serum ethanol levels should be monitored closely.
- At higher infusion rates it may be necessary to administer by volume rather than by milliliters per hour.
- For a 30% concentration, divide the amount by 1.5.
- Rounded off to the nearest milliliter.

Note: Concentrations higher than 10% are not recommended for intravenous administration. Concentrations higher than 30% are not recommended for oral administration. The dose schedule is based on the premise that the patient initially has a zero-ethanol level. The aim of therapy is to maintain a serum ethanol level of 100 to 150 mg/dL, but constant monitoring of the ethanol level is required because of wide variations in endogenous metabolic capacity. Ethanol is removed by dialysis, and the infusion rate of ethanol must be increased during dialysis. Prolonged ethanol administration may lead to hypoglycemia.

Ethanol Infusion

Fomepizole (4-methylpyrazole) has been approved by the US Food and Drug Administration for the treatment of ethylene glycol poisoning. It has also been used successfully in treating methanol poisoning. When compared with the traditional treatment of toxic alcohol poisoning, namely, ethanol, fomepizole has the advantages of ease of use, fewer side effects (specifically hypoglycemia), and ability to maintain therapeutic levels. Fomepizole is considered the antidote of choice; however, because of the cost and the logistics of stocking this antidote, many hospitals do not have this drug readily available.

Ethanol can be used as a therapeutic intervention in patients with methanol or ethylene glycol poisoning because of ethanol’s much greater affinity for alcohol dehydrogenases. These enzymes metabolize methanol and ethylene glycol to toxic by-products. However, with serum ethanol levels of 100 mg/dL, minimal amounts of ethylene glycol or methanol are metabolized by alcohol dehydrogenases. Ethanol infusions are not useful in the treatment of isopropyl alcohol poisoning.

Ethanol can be administered orally or intravenously (Table 67.10). IV ethanol has the advantages of achieving therapeutic levels rapidly, ensuring complete absorption, limiting the chance of aspiration, and avoiding gastritis. A 5% concentration of ethanol, which can be given in a peripheral vein, requires the use of large fluid volumes. In a 70-kg patient, a loading dose...
requires 1.4 L of a 5% solution, with a maintenance dose of 700 mL/hr. If IV ethanol is given, maintain careful attention to cardiopulmonary status. In contrast, oral loading can be achieved with much lower volumes. However, oral loading can be difficult in an uncooperative or unconscious patient or if vomiting or GI hemorrhage is present. A therapeutic level is reached more slowly with oral loading.

Ethanol metabolism can vary widely, and ethanol is dialyzable. Therefore it may be difficult to maintain appropriate ethanol levels during dialysis therapy for ethylene glycol or methanol poisoning. Frequent measurements of ethanol should be obtained and the infusion adjusted accordingly. When patients are given ethanol infusions, central nervous system depression, hypothermia, hypotension, hypoglycemia, and phlebitis are common adverse effects, especially in children. Serial levels of ethanol and glucose should be obtained.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


1469.e2 SECTION XII SPECIAL PROCEDURES


Contamination of health care workers (HCWs) with body fluids is a frequent occurrence in the emergency department (ED). A survey of ED and trauma staff found that almost half reported a needlestick injury within the preceding year. Body fluids often contain various transmissible infectious diseases, and the prevalence of human immunodeficiency virus (HIV) infection, hepatitis, and other communicable diseases has been shown to be high in ED patient populations.

Routine opt-out HIV screening of ED patients identifies approximately 1% of patients with previously undiagnosed HIV infections. Of such patients, many (20%) have no identifiable HIV risk factors, making it more difficult to identify patients who pose a risk to HCWs. This makes widespread identification of pediatric trauma resuscitations, compliance with barrier precautions was 81.3%. Similarly, ED staff have low observed compliance rates with standard precautions during cardiopulmonary resuscitation, with the lowest rates for wearing gowns (20%) and highest for wearing masks (90%).

In 1985, the combination of high-risk illness with low-compliance barrier use prompted the Centers for Disease Control and Prevention (CDC) to recommend guidelines for the protection of HCWs. In 1991, these recommendations were enacted into law by mandate of the Occupational Safety and Health Administration (OSHA). The primary focus of the CDC guidelines is to reduce mucocutaneous exposure to body fluids by encouraging hand washing and barrier protection. These measures do little to protect from percutaneous exposure, which is the most efficient method of transmission of hepatitis and HIV.

The current strategy for risk reduction in the ED includes immunization against hepatitis B virus (HBV), use of standard precautions (including reengineered safety products), and prompt initiation of postexposure prophylaxis (PEP) when appropriate.

GUIDELINES FOR STANDARD PRECAUTIONS

Appropriate precautions for all patient contact must be viewed as a consistent practice or “way of life” in the ED. The following guidelines, based on CDC recommendations, should be used when there is any possibility of contact with body fluids.

**Barrier Precautions**

1. Use gloves for any patient contact with a risk for exposure to body fluids (Fig. 68.1). Both cutaneous and percutaneous exposure can be reduced by the use of gloves. Gloves have been shown to reduce disease transmission in needlestick injuries, with greater reduction seen with double-gloving in animal models and in a case crossover study of HCWs.

2. Wear a mask and protective eyewear when exposure to body fluid aerosols is possible (e.g., wound irrigation, traumatic chest wound) (Fig. 68.2).

3. Wear a gown and shoe covers when there is a risk for large volumes of splashed body fluids (e.g., chest tube, thoracotomy) (Fig. 68.3).

**Respiratory Precautions**

ED patients commonly present with respiratory syndromes that require respiratory precautions in addition to standard precautions (Table 68.1). For example, patients with suspected
CHAPTER 68  Standard Precautions and Infectious Exposure Management

Figure 68.1 Gloves are mandatory for any patient contact with a risk for exposure to body fluids. Double-gloving may confer additional protection and should be considered in high-risk situations.

Figure 68.2 A mask and protective eyewear are mandatory when exposure to body fluid aerosols is possible. A gown is also suggested. This patient vomited profusely during intubation, which may have led to an exposure if proper protective gear had not been worn.

Figure 68.3 Full protective gear (eye protection, face mask, impervious gown, gloves, and shoe covers) should be worn when there is a risk for large volumes of splashed bodily fluids (e.g., emergency thoracotomy).

Figure 68.4 A, Recapping a needle by holding the cap in the hand is the most common way to sustain a needle puncture and should never be done! B, It is best to discard the needle/syringe without recapping, but if deemed absolutely necessary, use a single-handed technique to partially recap without holding the needle cap. C, Make sure that at least 80% of the needle is covered before completing the recapping with the second hand (by holding the base of the needle cap).

or confirmed pulmonary tuberculosis (TB), rubeola virus (measles), severe acute respiratory syndrome virus (SARS), or avian influenza should be isolated with airborne precautions, plus contact precautions with eye/face protections during any aerosol-generating procedures or if any contact with respiratory secretions is anticipated. During contact with patients who require airborne precautions, wear a National Institute for Occupational Safety and Health (NIOSH)–approved N-95 particulate respirator (Fig. 68.6). These masks are designed to efficiently filter 1- to 5-µm particles and are less costly and more comfortable than high-efficiency particulate air
SECTION XII  SPECIAL PROCEDURES

Figure 68.5 Using safety systems such as auto-recapping needles and retracting scalpels dramatically reduces percutaneous injuries to health care workers, and US federal law now mandates their use.

Figure 68.6 Respiratory precautions. Surgical masks are appropriate to protect against pathogens that are larger than 5 µm, such as influenza and Neisseria meningitidis. N-95 respirators are required for pathogens smaller than 5 µm, such as tuberculosis.

### TABLE 68.1 Respiratory Syndromes That Require Respiratory Precautions in Addition to Standard Precautions

<table>
<thead>
<tr>
<th>CLINICAL SYNDROME OR CONDITION</th>
<th>POTENTIAL PATHOGENS</th>
<th>EMPIRIC PRECAUTIONS (ALWAYS INCLUDES STANDARD PRECAUTIONS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cough/fever/pulmonary infiltrate in any lung location in an HIV-infected patient or a patient at high risk for HIV infection.</td>
<td><em>M. tuberculosis</em>, respiratory viruses, <em>S. pneumoniae</em>, <em>S. aureus</em> (MSSA or MRSA).</td>
<td>Airborne precautions plus contact precautions. Use eye/face protection if aerosol-generating procedure performed or contact with respiratory secretions anticipated. If tuberculosis is unlikely and there are no AIIRs and/or respirators available, use droplet precautions instead of airborne precautions. Tuberculosis more likely in HIV-infected individual than in HIV-negative individual.</td>
</tr>
<tr>
<td>Cough/fever/pulmonary infiltrate in any lung location in a patient with a history of recent travel (10–21 days) to countries with active outbreaks of SARS, avian influenza.</td>
<td><em>M. tuberculosis</em>, <em>SARS-CoV</em>, avian influenza</td>
<td>Airborne plus contact precautions plus eye protection. If SARS and tuberculosis unlikely, use droplet precautions instead of airborne precautions.</td>
</tr>
<tr>
<td>Respiratory infections, particularly bronchiolitis and pneumonia, in infants and young children.</td>
<td>Respiratory syncytial virus, parainfluenza virus, adenovirus, influenza virus, human metapneumovirus.</td>
<td>Contact plus droplet precautions; droplet precautions may be discontinued when adenovirus and influenza have been ruled out.</td>
</tr>
</tbody>
</table>

---

*Infection control professionals should modify or adapt this table according to local conditions. To ensure that appropriate empiric precautions are always implemented, hospitals must have systems in place to evaluate patients routinely according to these criteria as part of their preadmission and admission care. Patients with the syndromes or conditions listed below may present with atypical signs or symptoms (e.g., neonates and adults with pertussis may not have paroxysmal or severe cough). The clinician’s index of suspicion should be guided by the prevalence of specific conditions in the community, as well as clinical judgment.

The organisms listed under the column “Potential Pathogens” are not intended to represent the complete, or even most likely, diagnosis, but rather possible etiologic agents that require additional precautions beyond the standard precautions until they can be ruled out.

AIIR, Airborne infection isolation room; HIV, human immunodeficiency virus; CoV, coronavirus; MRSA, methicillin-resistant *S. aureus*; MSSA, methicillin-sensitive *S. aureus*; SARS, severe acute respiratory syndrome.

(HEPA)-filtered masks. In addition, place such patients in a respiratory isolation room with negative pressure, high circulation (optimally at least 12 air changes per hour), and external exhaust. When performing aerosol-generating procedures (such as intubation or sputum induction) that result in increased release of infectious droplets, add extra face/eye protection. However, even during outbreaks, adherence to isolation precautions is variable. A survey of medical students and residents at a Washington, DC, hospital (November–December 2009) showed that only 13% of medical students and 21% of residents would wear an N-95 respirator when caring for a patient with influenza symptoms.

For patients with respiratory infections such as suspected or confirmed respiratory syncytial virus (RSV), parainfluenza virus, adenovirus, or human metapneumovirus, the CDC recommends droplet precautions (see Table 68.1). Droplet precautions consist of placing the patient in a private room or a special separation of >3 feet. HCWs should don a mask on entering the patient’s room. If transport or movement is necessary, such as to radiology, instruct the patient to wear a mask and follow Respiratory Hygiene/Cough Etiquette.

**Personal Protective Equipment**

Isolation gowns are worn in combination with gloves only if contact with blood or body fluid is anticipated. The need for and the type of isolation gown selected is based on the nature of the patient interaction, including the anticipated degree of contact with infectious material and the potential for blood and body fluid penetration of the barrier. The CDC updated its personal protective equipment (PPE) guidelines when contacting haemorrhagic fever viruses in response to new data gained during an outbreak of the Ebola virus in several West African nations. CDC guidance emphasizes the importance of training, practice, competence, and observation of HCWs during an outbreak of the Ebola virus in several West African nations. CDC guidance emphasizes the importance of training, practice, competence, and observation of HCWs when evaluating persons under investigation, who are clinically stable and do not have bleeding, vomiting, or diarrhea. There is a lower risk of splashes and contamination in these situations. An established protocol, combined with proper training of the HCW, helps to facilitate compliance with PPE guidance.

1. Remove personal clothing and items. The HCW should wear surgical scrubs. No personal items (e.g., jewelry [including rings], watches, cell phones, pagers, pens) should be worn under PPE or brought into the patient room. Long hair should be tied back. Eyeglasses should be secured with a tie.
2. Inspect PPE prior to donning. Visually inspect the PPE ensemble to ensure that it is in serviceable condition (e.g., not torn or ripped), that all required PPE and supplies are available, and that the sizes selected are correct for the HCW.
3. Perform hand hygiene. Perform hand hygiene with alcohol-based hand rub (ABHR). When using ABHR, allow hands to dry before moving to the next step.
5. Put on gown or coverall. Ensure gown or coverall is large enough to allow unrestricted movement. Ensure cuffs of inner gloves are tucked under the sleeve of the gown or coverall.
6. Put on face mask.
7. Put on outer gloves. Put on second pair of gloves (with extended cuffs). Ensure the cuffs are pulled over the sleeves of the gown or coverall.
8. Put on face shield. Put on full face shield over the surgical face mask to protect the eyes, as well as the front and sides of the face.
9. Verify. After completing the donning process, the integrity of the ensemble should be verified by the HCW (e.g., there should be no cuts or tears in the PPE). The HCW should be comfortable and able to extend the arms, bend at the waist, and go through a range of motions to ensure there is sufficient range of movement while all areas of the body remain covered. A mirror in the room can be useful for the HCW while donning PPE.

**Doffing PPE**

1. PPE is doffed in the designated PPE removal area in the health care facility. As with all PPE doffing, meticulous care should be taken to avoid self-contamination. Place all PPE waste in a leak-proof infectious waste container.
2. Inspect. Inspect the PPE for visible contamination, cuts, or tears before starting to remove. If any PPE is visibly contaminated, disinfect by using an Environmental Protection Agency (EPA)-registered disinfectant wipe. If the facility conditions permit and appropriate regulations are followed, an EPA-registered disinfectant spray can be used, particularly on contaminated areas.
3. Disinfect and remove outer gloves. Disinfect outer-gloved hands with either an EPA-registered disinfectant wipe or ABHR. Remove and discard outer gloves, taking care not to contaminate inner gloves when removing the outer gloves. Dispose of outer gloves into the designated leak-proof infectious waste container.
4. Inspect and disinfect inner gloves. Inspect the inner gloves’ outer surfaces for visible contamination, cuts, or tears. If an inner glove is visibly soiled, disinfect the glove with either an EPA-registered disinfectant wipe or ABHR, remove the inner gloves, perform hand hygiene with ABHR on bare hands, and don a new pair of gloves. If a cut or tear is seen on an inner glove, immediately review occupational exposure risk per hospital protocol. If there is no visible contamination and no cuts or tears on the inner gloves, then disinfect the inner-gloved hands with either an EPA-registered disinfectant wipe or ABHR.
5. Remove face shield. Remove the full face shield by tilting the head slightly forward, grabbing the rear strap and pulling it over the head, gently allowing the face shield to fall forward. Avoid touching the front surface of the face shield. Discard the face shield into the designated leak-proof infectious waste container.
Change into scrubs, and remove all personal items such as jewelry and watches.

Visually inspect the PPE prior to donning to confirm correct sizing and ensure its structural integrity.

Perform hand hygiene and allow hands to dry. Alcohol-based hand rub is preferred if available.

Put on the gown or coveralls. Ensure the gown is large enough to allow unrestricted movement.

Apply the face mask or respirator. This should fit snug to the face and below the chin.

Apply gloves. Inner standard gloves and outer long-cuff nitrile gloves should be worn.

Put on full face shield over the surgical face mask to protect the eyes and front and sides of the face.

Have an assistant check the PPE to assure that all body areas are properly protected.

**Figure 68.7** Donning personal protective equipment. A trained safety observer should be present to help apply and inspect the equipment. *PPE, Personal protective equipment.*
Inspect the PPE for visible contamination, cuts, or tears before starting to remove.

Disinfect outer-gloved hands with either a disinfectant wipe or ABHR, and then remove.

Inspect and disinfect the inner gloves. Replace them with a new pair if visibly soiled.

Remove the outer gown, avoiding contact with the outer surface. Disinfect and replace gloves.

Remove the face mask and discard. Disinfect hands and replace gloves after face mask removal.

Remove the surgical or respiratory mask. Hold it by the straps and avoid touching the front of the mask.

Disinfect and remove the inner gloves.

Perform hand hygiene with ABHR, and have your safety observer inspect your scrubs for contamination.

**Figure 68.8** Doffing personal protective equipment. *ABHR*, Alcohol-based hand rub; *PPE*, personal protective equipment.
6. Disinfect inner gloves. Disinfect inner gloves with either an EPA-registered disinfectant wipe or ABHR.
7. Remove gown or coverall. Remove and discard.
   a. Depending on the gown design and location of fasteners, the HCW can either untie or gently break fasteners. Avoid contact of scrubs or disposable garments with the outer surface of the gown during removal. Pull the gown away from the body, rolling it inside out and touching only the inside of the gown.
   b. To remove the coverall, tilt the head back to reach zipper or fasteners. Unzip or unfasten the coverall completely before rolling down while turning inside out. Avoid contact of scrubs with outer surface of coverall during removal, touching only the inside of the coverall. Dispose of gown or coverall into the designated leak-proof infectious waste container.
8. Disinfect and change inner gloves. Disinfect inner gloves with either an EPA-registered disinfectant wipe or ABHR.
   a. Remove and discard gloves, taking care not to contaminate bare hands during the removal process.
   b. Perform hand hygiene with ABHR.
   c. Don a new pair of inner gloves.
9. Remove surgical face mask. Remove the surgical face mask by tilting the head slightly forward, grasping first the bottom tie or elastic strap, then the top tie or elastic strap, and remove the front of the surgical face mask without touching it. Discard the surgical face mask into the designated leak-proof infectious waste container.
10. Disinfect and remove inner gloves. Disinfect inner-gloved hands with either an EPA-registered disinfectant wipe or ABHR. Remove and discard gloves, taking care not to contaminate bare hands during the removal process. Dispose of inner gloves into the designated leak-proof infectious waste container.
11. Perform hand hygiene. Perform hand hygiene with ABHR.
12. Inspect. The HCW should inspect for any contamination of the surgical scrubs or disposable garments. If there is contamination, shower immediately, and then immediately inform the infection preventionist, occupational safety and health coordinator, or their designee.

Hand Hygiene
Immediately wash any skin surface coming in contact with body fluids with soap and water. If performed properly, both soap and water and alcohol-based products are generally efficacious in removing bacteria and preventing the spread of communicable diseases.25,26 Alcohol-based formulations have shown both bactericidal and virucidal activity.27,28 However, soap and water perform better in removing Clostridium difficile spores.29 Hand hygiene adherence is frequently low in emergency settings.10-32

OCCUPATIONAL DISEASE EXPOSURE
Occupationally acquired infections cause considerable morbidity and mortality among HCWs, despite OSHA requirements for precautions. Given the often occult manifestation of disease in the ED patient population, emergency HCWs are at high risk for exposure to infectious diseases. There is a high prevalence of HIV, HBV, hepatitis C virus (HCV), and pulmonary TB in ED patients, as well as associated high morbidity and mortality. This chapter focuses on these particular diseases and provider risks.

HBV
Transmission
HBV is a well-recognized occupational risk for health care providers, and multiple studies have documented the high prevalence of hepatitis in ED patients.13,14 Despite the attention focused on transmission of HIV, the infectivity of HBV is significantly higher. HBV is a more virulent organism and requires a relatively small inoculum for transmission.31 Percutaneous injuries are the most efficient mode of HBV transmission, and many infected HCWs do not recall a specific injury.32 Many body fluids other than blood contain hepatitis B surface antigen (HBsAg), and levels of infectious HBV particles in blood-free body fluids are 100 to 1000 times lower than in blood itself. Implementation of the CDC's standard precautions, along with the OSHA regulations for barrier protection and preexposure vaccination, has led to a decrease in the incidence of HBV transmission.37

To understand the risk of HBV transmission resulting from occupational exposure, an understanding of a few key serologic markers for HBV is essential. HBsAg is a marker of active infection in the source patient. From a practical standpoint, HBV can be transmitted when HBsAg is present, and it is not generally transmissible when this marker is absent. Hepatitis B surface antibody is a protective antibody against HBV. In vaccinating HCWs, the goal is to stimulate the immune system prior to accidental exposure, to produce a sufficient quantity of this antibody to protect the individual if exposed to the virus. Hepatitis B e antigen (HBeAg) can be found in the bloodstream of HBV-infected individuals during times of peak virus replication. The risk for clinical hepatitis is approximately 2% (range, 1% to 6%) if HBeAg is absent, as opposed to a risk of 22% to 31% if HBeAg is present.38 Yet testing occupational exposure–source patients for HBeAg is not recommended.19

Postexposure Management
PEP following exposure to an HBsAg-positive source may require hepatitis B vaccine, hepatitis B immunoglobulin (HBIG), both, or neither (Table 68.2). This depends on the vaccination and antibody response status of the exposed HCW. HBIG is derived from pooled human plasma and provides passive immunization for nonimmune exposed individuals. This preparation is very safe and not known to transmit disease.17,18 When HBIG is used for PEP, give it ideally within 24 hours after exposure, and note that it is of questionable value beyond 7 days.18 PEP for HBV is remarkably effective, and infection is unlikely to develop in individuals who receive PEP.18 Hepatitis B vaccine may also be given with PEP. Individuals who have not previously been vaccinated or who have not demonstrated an adequate response should receive the hepatitis B vaccine. Adverse reactions to the hepatitis B vaccine are generally quite mild, and it is safe to give during pregnancy. For primary immunization, give an initial intramuscular injection, followed by subsequent intramuscular vaccinations at 1 and 6 months. There is no need to provide vaccination or to check titers in individuals who have previously had an adequate titer.17 PEP with these agents is not contraindicated during pregnancy or lactation. HCWs who have previously been infected with HBV are immune to reinfection, so PEP is not indicated.
HCV

Transmission

The estimate of the number of US residents with HCV is at least 4.6 million, and many individuals are unaware of their infection. ED patients have a high prevalence of HCV. HCV is often acquired from injection drug use. It was once commonly transmitted by blood transfusion, but fortunately, with modern screening, this is now rare. Although HCV can be transmitted sexually, this is a minor route. Mucous membrane transmission of HCV is possible but much less common. Percutaneous transmission is the most efficient route. The incidence of seroconversion after an HCV-positive needlestick is approximately 1.8% (estimates range from 0% to 7%). It is useful to remember that the risk positive needlestick is approximately 1.8% (estimates range from 0% to 7%). The risk of HCV transmission after a needlestick injury is much less common. Percutaneous transmission is the most efficient route. The incidence of seroconversion after an HCV-positive needlestick is approximately 1.8% (estimates range from 0% to 7%).

HIV

Transmission

The number of occupational exposures to HIV reported to the CDC have steadily decreased. As of December 2013, 58 confirmed occupational transmissions of HIV have been reported. In 2010 one case of occupational HIV transmission was reported to the CDC. The risk for contracting HIV from working in the ED depends on the prevalence of HIV in the local patient population. The overall risk for HIV seroconversion is approximately 0.23% after a needlestick injury and 0.03% for mucocutaneous transmission. The risk of HIV infection associated with intact skin is too low to be detected. Cardo and colleagues demonstrated that the risk for HIV seroconversion after needlestick injuries is not uniform. Seroconversion was found to be more likely for deep injuries.
(odds ratio [OR] = 15), if blood was visible on the device (OR = 6.2), if the needle had been used in a source patient’s artery or vein (OR = 4.3), or if the source patient suffered from terminal acquired immunodeficiency syndrome (OR = 5.6). It is essential to gather information regarding the nature of the injury to risk-stratify the exposure.

When seroconversion occurs, HIV antibodies can be detected as early as 3 days after exposure and are almost always present by 6 months. Acute retroviral syndrome is a clinical manifestation of HIV seroconversion that occurs in approximately 80% of newly infected individuals at a median of 25 days after exposure. The signs and symptoms of acute retroviral syndrome are similar to those of mononucleosis and consist of fever, lymphadenopathy, and rash.

**Postexposure Management**

Evidence Supporting PEP

In 1998 the US Public Health Service recommended using PEP for selected HIV exposures. These recommendations were based on a single case-control study. Currently, a large placebo-controlled trial would be considered unethical. The CDC-sponsored case-control study, undertaken in the United States, France, the United Kingdom, and Italy, compared 33 HCWs who seroconverted after exposure to HIV with 665 control HCWs who did not seroconvert after exposure to HIV. Approximately 90% of the patients in this study were exposed via hollow-bore needles. When postexposure zidovudine (azidothymidine) was used, the risk for HIV infection was reduced by 81% (95% confidence interval, 48% to 94%).

Selecting Patients for PEP

In 2013 the US Public Health Service published updated recommendations regarding the use of HIV PEP. In general, the decision to use PEP depends on the type of exposure and the source’s HIV status. The first step in determining whether PEP is indicated is to assess the severity of exposure. Percutaneous exposure can be categorized as “less severe” or “more severe.” A less severe exposure involves a solid needle, a superficial injury, and no blood visible on the device. All other percutaneous injuries are categorized as more severe. Exposure to mucous membranes and nonintact skin is categorized as either “small volume” (a few drops of blood) or “large volume” (a major blood splash). There are no reported cases of HIV seroconversion after exposure of intact skin to blood.

After assessing the severity of exposure, determine the potential infectivity of the source. The US Public Health Service emphasizes the importance of determining the HIV status of the exposure source. Consider PEP only for exposure to blood and body fluids from a source known or likely to be HIV positive. Exposure from an HIV-negative source does not require PEP. Do not test sharp instruments for HIV because this is not reliable or recommended. Note that drug development is ongoing and recommendations change rapidly. The clinician is urged to periodically check for updates. The most reliable source is the CDC website or the *Morbidity and Mortality Weekly Report* journals, most easily found via an Internet search under various topics.

A number of special circumstances may arise when determining the need for HIV PEP. When a source is known but an individual’s HIV status is pending, decide about the use of PEP on a case-by-case basis. When the source is high risk, initiate PEP and then stop or modify it later once the HIV status of the source is determined. When a source can be identified but the HIV status of the source is unknown and will not become available, PEP is not generally recommended. However, if it is unknown whether the source constitutes a significant risk and consultation is not available within a few hours; it is recommended to begin PEP and obtain consultation later.

**Choice of PEP Medications**

The US Public Health Service recommends that PEP medication regimens begin as soon as possible after the exposure and should be continued for 4 weeks. When HIV PEP is administered, a minimum of three drugs is recommended. The current preferred three-drug HIV PEP regimen includes raltegravir, tenofovir, and emtricitabine (Table 68.3).

A number of second-line and alternative agents may be chosen for HIV PEP. Expert consultation is recommended, especially if antiretroviral (ARV) resistance is suspected. Pregnancy and breastfeeding are not contraindications to taking ARV agents and PEP is recommended. Consult an infectious diseases or occupational health specialist when treating pregnant and lactating patients, as alternative regimens may be considered. Enter all pregnant women starting ARVs in the Antiretroviral Pregnancy Registry, a database designed to collect information on the outcomes of ARV-exposed pregnancies regardless of HIV status: http://www.apregistry.com.

**TABLE 68.3 HIV PEP Regimens**

<table>
<thead>
<tr>
<th>Preferred HIV PEP regimen</th>
<th>Raltegravir (Isentress; RAL) 400 mg po bid Plus Tenofovir DF (Viread; TDF) 300 mg po qd plus Emtricitabine (Emtriva; FTC) 200 mg po qd</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alternative agents for use with expert consultation</td>
<td>Abacavir (Ziagen; ABC) Efavirenz (Sustiva; EFV) Efavirenz (Fuzone; T20) Fosamprenavir (Lexiva; FOSAPV) Maraviroc (Selzentry; MVC) Saquinavir (Invirase; SQV) Stavudine (Zerit; d4T)</td>
</tr>
<tr>
<td>Agents generally not recommended for use as PEP</td>
<td>Didanosine (Videx EC; ddI) Nelfinavir (Viracept; NFV) Tipranavir (Aptivus; TPV)</td>
</tr>
<tr>
<td>Agents contraindicated as PEP</td>
<td>Nevirapine (Viramune; NVP)</td>
</tr>
</tbody>
</table>

HIV, human immunodeficiency virus; PEP, postexposure prophylaxis.

Note: for consultation or assistance with HIV PEP contact the National Clinicians’ Post-Exposure Prophylaxis Hotline at telephone number 888-448-4911 or visit its website at http://ncce.ucsf.edu/clinician-consultation/pep-post-exposure-prophylaxis/.

agents can be used in pediatric patients for occupational and nonoccupational exposure, however consultation is recommended. For consultation or assistance with HIV PEP, contact the National Clinicians’ Post-Exposure Prophylaxis Hotline, telephone number 888-448-4911 or visit its website at http://nccc.ucsf.edu/clinician-consultation/pep-post-exposure-prophylaxis/.

Timing, Duration, and Side Effects of PEP
HIV exposure should be considered a true emergency. Administer PEP as soon as possible after exposure, ideally within 1 hour. Animal studies indicate that the efficacy of PEP diminishes with a delay in initiation. HIV PEP regimens consist of a 4-week course of therapy. In the ED, patients can be prescribed the first 3 days of medications, and outpatient follow-up should be arranged within 72 hours (Fig. 68.9). Side effects from the medications used for HIV PEP are not insignificant. However, the current recommended three-drug regimen is better tolerated than earlier zidovudine-containing regimens. Depending on the choice of PEP medications, patients should also be prescribed antiemetics and antidiarrheal agents when PEP is initiated.

TB
Transmission
During the mid-1980s the United States experienced a resurgence in TB, especially among HIV-positive patients. Although the incidence of TB cases in the United States has continued to decline, more than 9000 cases were reported in 2014. TB continues to pose a serious risk to both public health and HCWs. Baussano and colleagues estimated that the annual risk for TB infection in HCWs, attributable to occupational exposure, is between 3.8% and 8.4% depending on local prevalence. TB is transmitted by infectious droplets 1 to 5 µm in size. Primary infection occurs when one to three organisms are inhaled into the alveoli, where they begin to replicate. Host defenses usually stop infection within 2 to 10 weeks, and the patient enters the latent period. During this time, patients are asymptomatic and not contagious. Reactivation occurs when cell-mediated immunity wanes, and patients are again contagious. This can be due to advancing age, HIV infection, steroid use, malignancy, malnutrition, or other causes of suppression of the immune system. The lifetime risk for reactivation is 5% to 10%, with approximately half this risk occurring in the first few years after primary infection. Patients with increased infectivity include those with pulmonary or laryngeal TB, an active cough, positive sputum smears for acid-fast bacilli, cavitary lesions on chest radiographs, and inadequate therapy. Overall, children are less contagious than adults, but can still transmit the disease. Extrapulmonary TB is contagious only in cases of an open skin lesion or involvement of the oral cavity.

Depending on the patient population and geographic location, ED personnel can be at high risk for occupational TB infection. In a 1993 study at a county hospital in Los Angeles it was reported that 31% of ED workers became positive for purified protein derivative (PPD) during their employment, including 20% of attendings, 32% of nurses, and 33% of residents. The risk for PPD conversion was found to be 6% after 1 year of ED employment, 14% after 2 years, and 27% after 4 years. EDs typically care for higher-risk patients: those who are homeless, foreign born, recently incarcerated, or chronically debilitated. Overcrowding can lead to extended waiting periods and delays in admission. The clinical manifestation of TB in ED patients is often atypical, which can lead to a delay in diagnosis. Delayed ED diagnosis of pulmonary TB is associated with factors such as recent antibiotics, older age, renal insufficiency, preexisting chronic lung disease, HIV, homelessness, or substance abuse. A 14-year review of the US National Tuberculosis Surveillance System showed that rates of advanced pulmonary TB are on the rise, especially in populations without traditional risk factors. These data suggest that substantial delays in diagnosis and misdiagnosis of pulmonary TB may be increasing.

Preventing exposure to TB requires a multifaceted approach. Proper ED ventilation plays a key role; inadequate ventilation has been a contributing factor in many nosocomial outbreaks. Ideally, install single-pass airflow from waiting rooms to the outside. Within the ED, make sure that air flows from clean areas to less clean areas, not vice versa. If patients with TB are seen frequently, provide at least one true respiratory isolation room in the ED. Make sure that such rooms have at least 12 air changes per hour and that they are “negative pressure” rooms in which air flows into the room from other ED areas. Other engineering approaches for control of TB infection include using HEPA filters and upper-room ultraviolet light irradiation.

Familiarize all ED personnel with the appropriate use of respiratory protection against TB. Provide surgical masks (e.g., string-tie masks) for source control. Place these masks on potentially contagious patients to decrease the passage of infectious droplets into the air. Because air can leak around such masks, they are not optimal for protection of HCWs. In late 1995, NIOSH certified a new class of masks known as N-95 particulate respirators. These masks filter particles 1 µm in size with at least 95% efficiency and are generally the
findings of an apical infiltrate, cavitary lesions, extensive hilar or mediastinal lymphadenopathy, pleural disease, or a miliary pattern.  

**Postexposure Management**

If HCWs are exposed to patients with active pulmonary TB, refer them to either employee health care or their primary care clinician for follow-up testing and treatment. To establish a HCW’s baseline PPD status, perform skin testing within days after exposure. If the baseline test is negative, perform a follow-up skin test 8 to 10 weeks later to determine whether PPD conversion has occurred. A positive baseline test (≥10-mm induration) indicates previous exposure or infection. A positive test consists of 5-mm induration after a negative baseline test or an increase to at least 10-mm induration after a baseline test of 1- to 9-mm induration.

HCWs in whom PPD converts to positive after an exposure should undergo chest radiography to screen for active pulmonary TB. If active disease is present, initiate treatment with at least four antituberculous medications. In PPD converters who do not have active disease, consider chemoprophylaxis. When deciding whether to initiate chemoprophylaxis, balance the potential benefit of TB prevention with the risk for medication-associated hepatitis. In general, give chemoprophylaxis in the case of a recent (≤2 years) PPD conversion, a known TB contact, a patient who is medically predisposed to TB, HIV-infected patients, intravenous drug users, or those younger than 35 years. For occupationally exposed HCWs who are PPD converters, the preferred regimen for HIV-negative persons is daily isoniazid (INH) for 9 months, although acceptable alternative regimens can be considered. Some HCWs may become exposed to strains of TB that are resistant to INH because of the continued emergence of multidrug-resistant and extensively drug-resistant strains of TB. For exposure to multidrug-resistant TB, consult an expert when selecting an individualized chemoprophylaxis regimen. Baseline and serial liver function testing is not necessary for administration of chemoprophylaxis in most cases, but monitor closely for clinical symptoms suggestive of hepatotoxicity.

**REFERENCES ARE AVAILABLE AT** www.expertconsult.com


Emergency clinicians often face the challenge of caring for agitated, uncooperative, combative, and violent patients who are unable to participate in their care, make rational health care decisions, and/or are a danger to themselves and others.\(^1\)\(^,\)\(^2\) Psychiatric illness, acute chemical intoxication or withdrawal, delirium, medical illness, uncontrolled rage, hypoxia, and rarely, central nervous system infection are causes of agitated or violent behavior in the emergency department (ED).\(^1\)

**BACKGROUND**

Patients more likely to exhibit severe uncontrolled agitation are those who abuse alcohol; sympathomimetic agents such as ecstasy (MDMA), mephedrone, or methylone (bath salts); and hallucinogenic drugs such as phencyclidine (PCP). Patients can also experience toxicity after accidental and intentional ingestion of medications. Drug-associated delirium may be related to anticholinergic toxicity, serotonin syndrome, or intoxication with sympathomimetics. Alcohol withdrawal may progress to delirium tremens, a condition typified by excessive agitation and confusion.

Patients with agitated delirium are often the most difficult to manage and at high risk for significant morbidity and mortality. This is especially true for patients with excited delirium syndrome (ExDS), a variant of and the extreme form of agitated delirium (see later section on ExDS).\(^3\) Causes of agitated delirium include manic-depressive disorder, psychosis, chronic schizophrenia, intoxication with sympathomimetics or anticholinergics, cocaine intoxication, alcohol or benzodiazepine withdrawal, hypoglycemia, the postictal state, or head trauma. Some causes are idiopathic, but many are due to a combination of underlying psychiatric illness and stimulant drugs or alcohol (or both). Because of the variety of causes, management of agitated and combative patients requires a systematic approach (Fig. 69.1).

Schizophrenia, schizoaffective disorder, and the manic phase of bipolar disorder can all lead to an impaired perception of reality. Paranoid delusions, hallucinations, and hostile moods that can easily lead to severe agitation often develop in patients with these disorders.

Any medical condition that leads to brain dysfunction may also result in agitated, combative, or violent behavior (Box 69.1). Well-documented examples include hypoglycemia, hypoxia, medication intoxication, encephalitis, meningitis, intracranial hemorrhage, thyrotoxicosis, traumatic brain injury, febrile illnesses in the elderly, and dementia.\(^4\)

It is essential to achieve control of an agitated patient as quickly as possible. Goals of management include preventing patients from harming themselves and others, identifying the underlying cause, and initiating medical treatment. Early intervention allows caregivers to implement appropriate monitoring, perform a physical examination, initiate diagnostic testing, and institute timely treatment. Successfully achieving these goals often necessitates physical and chemical restraint.

During the late 1980s and early 1990s, reports of restraint-associated deaths in psychiatric and extended care facilities\(^5\) led lawmakers to pass legislation establishing regulations for the use of restraint.\(^6\) Since then, the Joint Commission for Accreditation of Healthcare Organizations (JCAHO) and the Center for Medicare and Medicaid Services (CMS) have created standards governing the use of restraint in a variety of health care settings, including EDs and tertiary care facilities (Box 69.2).\(^7\)\(^,\)\(^8\) The aforementioned standards allow the use of physical restraint for a limited period. A licensed practitioner must evaluate the patient and determine that less restrictive interventions have been ineffective and physical restraint is required to ensure the patient’s well-being. The standards also include requirements for written policies and procedures governing the use of restraint that must address indications, staff training and education, patient assessment and reevaluation, appropriate documentation, and patient-focused issues such as maintaining dignity and respect. Hospitals and extended care facilities are required by law to report any death or adverse event related to the use of restraint.\(^8\)

Some authors have advocated “restraint-free” environments in nursing homes, extended care facilities, and acute care hospitals.\(^9\) Although this concept is laudable, mandating a restraint-free ED is impractical and potentially dangerous. Clinicians should recognize the risks associated with restraining a patient; however, they should not be deterred from using either physical or chemical restraint when patients demonstrate dangerous behavior toward themselves or others. Indeed, the American College of Emergency Physicians (ACEP) recently revised and approved their policy statement in 2014 by supporting the careful and appropriate use of physical restraints or seclusion.\(^10\)

**EXCITED DELIRIUM SYNDROME (EXDS)**

In September 2009, a task force of the ACEP published a multi-authored White Paper Report on ExDS. This was a landmark report because it legitimized a heretofore poorly defined serious medical scenario that was familiar to law enforcement, emergency medical services, and emergency physicians. In addition, it supported the aggressive and potentially lifesaving use of chemical sedation.

It was the consensus of the task force that ExDS is a unique syndrome that may be identified by the presence of a distinctive group of clinical and behavioral characteristics that can be recognized in the premortem state. ExDS, though potentially fatal, may be amenable to early therapeutic intervention in some cases.

The task force defined the existence of excited delirium as a true disease entity; described the signs, symptoms, and risk for death; and reviewed current and emerging methods of control and treatment. Before this report, agitated delirium and various poorly defined related terms were used to describe a serious form of agitated delirium that could culminate in death. Such deaths were often high-profile news events that
inexplicably occurred in seemingly healthy young males and often involved interactions with seriously deranged and violent individuals and law enforcement officers.

The majority of affected individuals are young males. Features of ExDS include uncontrolled aggression and agitation, tremendous pain tolerance, tachypnea, tachycardia, sweating, tactile hyperthermia, pacing, grunting, noncompliance with police orders, unusual and uniriting strength, being inappropriately clothed, extreme paranoia, and having an attraction to mirror or glass. Individuals are unable to engage in rational discussion or understand or deescalate their abnormal aggressive, violent, and threatening behavior. A common fatal scenario is characterized by a period of extreme delirium, increasing agitation, and inability to reason or comply with efforts to control their agitated state, followed by a struggle with law enforcement that involves vigorous physical restraint (choke holds, hog-tying, prone positioning, knee to the throat), and noxious chemicals. The actual role of physical restraint is unproven. The use of the TASER device (TASER International, Inc., Scottsdale, AZ) to control violent and markedly agitated individuals has been temporally related to death, but the role of this device in causing death is also unproven. Intervention may be followed by a sudden cessation of struggling that may herald death by cardiopulmonary arrest.
The exact cause of death in ExDS cannot be identified at autopsy. Death may be related to underlying but unknown pathology (such as cardiomyopathy, conduction abnormalities, and metabolic disturbances). Lack of complete prior medical information, especially underlying cardiac abnormalities, hampers ascertainment of the actual cause of death when only the autopsy results are interpreted. Unfortunately, law enforcement may be blamed for using excessive force when no autopsy proven cause of death is forthcoming.

Stimulant drug use, including cocaine, methamphetamine, and PCP, has a well-established association with ExDS and is usually implicated as being closely linked to if not causative of death from ExDS. Persons with psychiatric illnesses represent the second largest, but distinctly smaller cohort of ExDS cases. The combination of drugs, alcohol, and psychiatric illness is well recognized. Some deaths are temporally related to the application of a conducted electrical weapon (CEW; e.g., a TASER), but there are well-documented cases of ExDS-associated deaths with minimal restraint such as handcuffs without TASER use.

No definitive diagnostic tests are available for ExDS. Currently, it must be identified by clinical features, which renders it very difficult to ascertain the true incidence. Though not universally fatal, it is clear that a proportion of individuals with ExDS progress to cardiac arrest and death regardless of medical intervention.

Although the specific precipitants of fatal ExDS remain unclear, epidemiologic and clinical reports provide some understanding of the underlying pathophysiology. The clinical picture is one of an agitated and delirious state with severe autonomic dysregulation. When available, cardiac rhythm analysis demonstrates bradyarrhythmic asystole; ventricular dysrhythmias are rare and have occurred in only a single patient in one study. Severe acidosis and hyperthermia appear to play a prominent role in lethal ExDS-associated cardiovascular collapse. One potential cause is thought to be excessive central dopamine stimulation.

Based on the available evidence, it was the consensus of the task force that ExDS is a real syndrome of uncertain etiology. It was also the consensus of the panel that rapid and appropriate, albeit limited, physical restraint measures and immediate administration of benzodiazepines intravenously, ketamine intravenously, or midazolam intranasally may be lifesaving.

Importantly, insufficient data were available to the task force to determine whether fatal ExDS is preventable or whether there is a point of no return after which the patient will die regardless of advanced life support interventions.

**MEDICOLEGAL CONCERNS**

Emergency clinicians need to be cognizant of the potential legal ramifications stemming from physically restraining patients. The risk for litigation may be mitigated by strict adherence to written institutional and departmental policies regarding the use of restraint and medications. Unfortunately, no one-size-fits-all protocol is realistic, and each case must be individualized by a clinician at the bedside, often with little data or background on which to base a specific intervention. Emphasis should be placed on timely and comprehensive assessment before restraint, regular patient reevaluation, limitations on the time spent in restraint, and detailed documentation in the ED record. Furthermore, clinicians need to recognize that competent patients do have the legal right to refuse medical treatment even if the result of their refusal is death or serious bodily harm. Competence, however, may be difficult to evaluate or ascertain in the time frame required to make important clinical decisions. There are many gray areas that cannot simply be solved by attempting to obtain a formal psychiatric consultation. In fact, there are no data proving that a psychiatrist is any better than an emergency physician in determining competence in an ED patient within the time frame of the ED evaluation, when important decisions must be made. Individuals with true ExDS are clearly mentally incompetent, but coercive measures, including physical restraint or the threat of physical restraint, should not be used simply because a competent patient refuses treatment or as retaliation for perceived disruptive behavior.

**PATIENT ASSESSMENT**

Emergency clinicians must avoid ascribing agitated or abusive behavior to drug or alcohol intoxication or underlying psychiatric disease before considering severe, life-threatening diagnoses (see Box 69.1). Every attempt should be made to obtain a detailed history of issues related to the patient’s condition, as well as the patient’s past medical and psychiatric history, including medications, drug and alcohol use, and previous similar events. In reality, such information is rarely available or accurate, thus leaving only theclinician’s clinical judgment to guide therapy.

Five groups of patients have been identified as being at increased risk for an underlying medical problem: the elderly, those with a history of substance abuse, patients without a previous psychiatric disorder, those with preexisting or new medical complaints, and individuals from lower socioeconomic groups. Of these, patients with new psychiatric symptoms are especially worrisome and require careful evaluation for underlying medical illness.

Physical examination should be aimed at identifying organic causes of the patient’s behavior such as trauma, infection, and metabolic derangements. Rapid bedside serum glucose
restraint is necessary. Indeed, several de-escalation techniques in defusing a difficult situation before physical or chemical recognition of this somewhat predictable pattern may help anxiety, defensiveness, and finally, physical aggression.\textsuperscript{16}

Coburn and Mycyk described three phases of escalating violence:

1. Recognition
2. Increased defensive behavior
3. Aggression

DE-ESCALATION TECHNIQUES

Coburn and Mycyk described three phases of escalating violence: anxiety, defensiveness, and finally, physical aggression.\textsuperscript{16} Recognition of this somewhat predictable pattern may help in defusing a difficult situation before physical or chemical restraint is necessary. Indeed, several de-escalation techniques have been shown to assist in quelling agitated and violent patients.\textsuperscript{7}

One simple, yet effective technique is verbally engaging the patient by asking, “how can we help you?” This display of compassion on the part of the treating clinician and staff may calm the patient. Similarly, offering food and drink will often soothe an agitated patient. Along with these displays of caring and empathy, it is important to impress on the patient that violent behavior will not be tolerated and will be dealt with quickly and firmly.\textsuperscript{7} If the patient continues to demonstrate agitated or violent behavior, enlist the aid of a family member or a specially trained individual, such as a social worker, psychiatric counselor, or member of the clergy.

If these conservative measures fail, summon hospital security to ensure the safety of the patient and staff. In these situations, security staff should not rush to the patient’s bedside but instead should gather outside the door or close by, within eye contact of the patient’s room. A strong show of force may calm a potentially violent patient without the need for restraint.

TYPES OF PATIENT RESTRANST

Seclusion

The utility of seclusion in psychiatric evaluation units and inpatient hospital wards is well documented in both adult and pediatric populations.\textsuperscript{17-20} In the mid-1980s, seclusion was commonly used to calm combative and violent ED patients, but its popularity has declined since then.\textsuperscript{1,21,22} Reasons for this decline are unclear but probably include lack of adequate space and concerns regarding provision of medical care and compliance with regulatory agencies.\textsuperscript{22} Nevertheless, at institutions with adequate physical space and well-designed policies and procedures, experience has shown that seclusion is an effective ED practice for selected patients.\textsuperscript{22,23}

Seclusion involves placing agitated and violent patients in an isolated space and confining them to dedicated hospital stretchers or beds that are secured in place to prevent injury. Seclusion is often used in conjunction with chemical sedation, physical restraint, or both.\textsuperscript{22} The seclusion room should be located near ED staff to allow continuous patient observation. Regularly reassess patients who are placed in seclusion, similar to physically restrained patients (see Box 69.3).

Physical Restraint

Research regarding the application of physical restraint in the ED is limited. In one prospective study of restraint-associated complications in 298 ED patients, minor complications occurred in only 7% of patients, and there were no serious complications or deaths.\textsuperscript{24} The author of this study concluded that the dangers of ED patient restraint promulgated by professional organizations and health care regulators may be overstated.\textsuperscript{24} Moreover, this study supports the safe use of restraint by emergency clinicians, who by virtue of their training and experience are experts in recognizing signs of deterioration and skilled in airway management and resuscitation. It should also be noted that TJC sentinel event tracking has demonstrated a decreased rate of complications when patients are restrained on their sides.\textsuperscript{24,25}

Although the actual prevalence of patient restraint in EDs is unknown, it has been estimated that 25% of teaching hospitals physically restrain at least one person per day.\textsuperscript{1}

Restraint Devices

Limb Holders (Restrains)

Restraining a patient's extremities is the primary method of physical restraint used in the ED. Limb restraints are constructed from a variety of materials, including leather, synthetic leather, cotton, and single-use foam material. These materials provide restraint that differs in strength, ease of removal, and cleaning.

Hard leather and synthetic leather limb holders are virtually impossible to break or tear but are difficult to sterilize if they become soiled with blood or body fluids (Fig. 69.2). They are more rigid than soft restraints, which also makes them somewhat more difficult and time-consuming to apply. More importantly, most leather limb holders require a key to unlock and, as a result, might take longer to remove after an adverse event such as vomiting or respiratory arrest. Leather limb holders are generally used to restrain combative and violent patients in whom the need for indestructible secure restraints outweighs the more time-consuming application and removal process.
Soft limb restraints are usually made from cotton or foam material, or both (Fig. 69.3). They are single-use devices, which obviates the need for cleaning and sterilization. Soft limb restraints are less rigid than leather limb restraints, which makes them easier to apply. In addition, because they are fastened without the use of a key, soft limb restraints are more easily removed. Soft restraints are typically used for agitated but less combative patients because they are not as secure as leather limb holders.

Belts/Fifth-Point Restraint
A fifth-point restraint is a belt apparatus used to supplement the use of four limb restraints by holding down the thighs, chest, or pelvis (Fig. 69.4). A fifth-point restraint is used on patients who continue to be at risk for harming themselves despite adequate limb restraint and whose continued combative behavior interferes with diagnostic or therapeutic interventions. When patients are restrained across the thighs, chest, or pelvis, they may not be able to sit up or turn onto their sides, thus placing them at higher risk for aspiration should vomiting occur. Keep the side rails of the stretcher in the upright position at all times and place the belt snugly enough to prevent the patient from slipping under the device and thereby increasing the risk for accidental suffocation. Fifth-point restraints are usually made of synthetic material and are available with both quick-release and key-release locks.

Jackets and Vests
Jackets and vests are generally used on inpatient wards and extended care facilities for the prevention of falls; they have little utility in the ED (Fig. 69.5). Moreover, these products have been implicated in a number of restraint-associated deaths secondary to choking and suffocation. The use of restraint jackets and vests in the ED is not recommended.

Hobble Leg Restraints
Hobble leg restraints limit movement by securing the patient’s ankles with connecting locking cuffs (Fig. 69.6). Hobble leg restraints are commonly used by law enforcement agencies because they impede running and kicking, which makes them...
an effective method of transporting potentially violent patients and those who pose a risk for flight. Hobble leg restraints are seldom used in the ED, but their use by law enforcement agencies and prison authorities means that most emergency clinicians will encounter patients placed in these devices. The combination of prone positioning, hobble leg restraints, and binding a patient’s hands behind the back, commonly referred to as hog-tying, was a common method of restraining prisoners and violent psychiatric patients (Fig. 69.7). However, this practice is no longer widely used because of the risk for suffocation (see later section on Positional Asphyxia).

**Indications**

Use limb restraints to prevent agitated, combative, or violent patients from harming themselves or others. Frequently, the use of restraints can be delayed while verbal de-escalation techniques are attempted (see earlier section on De-escalation Techniques). However, if a patient is deemed an immediate threat to himself/herself or others, restrain him/her without delay. Patients with altered mental status may require limb restraints so that diagnostic testing can be completed or appropriate treatment rendered, or both. Use limb restraints also to prevent patients from interfering with devices, such as endotracheal tubes, cardiac monitors, and indwelling intravenous (IV) lines and catheters.

**Contraindications**

When appropriate, delay physical restraint in favor of trying verbal de-escalation techniques. Do not place limb restraints on extremities with fractures, open wounds, or acute skin and soft tissue infections. Use caution to avoid ischemia in patients who exhibit tenuous perfusion of extremities, such as those with peripheral vascular disease or previous arterial injury or surgery.

Avoid fifth-point restraint of the abdomen and pelvic region in patients with pelvic fractures, suprapubic tubes, ostomies, and percutaneous feeding tubes. In addition, it is important to note that patients with underlying pulmonary or cardiac disease may not tolerate restraint of their thorax.

**Procedure**

Use a minimum of five people, all trained in restraint techniques and patient safety, to restrain the patient if possible (Fig. 69.8). This show of force may help discourage the patient from resisting and is an important part of the restraint process. Always restrain patients in the supine rather than the prone position because the prone position increases the risk for suffocation. Assign one person to hold each limb firmly against the stretcher by applying direct pressure proximal to the elbows and knees. The fifth member of the team places restraints around the wrists and ankles (Fig. 69.9). Control above the elbows and knees reduces the risk for injury to these joints and concentrates force closer to the patient’s center of gravity for better control.

Apply the limb holders snugly enough to control movement and prevent escape, but not so tight that they cause pain or impair circulation. If necessary, place a fifth-point restraint across the patient’s thighs, pelvis, or chest to further limit motion. Place a surgical mask temporarily over the patient’s mouth to prevent the patient from spitting at members of the restraint team or ED staff. If leather restraints are used, keep
Complications

Increased Agitation

For some patients, placement in physical restraints is so emotionally disturbing that it actually increases agitation and combative behavior. Patients who continue to struggle despite restraints are at risk for a number of potentially serious adverse events, including skin damage, ischemia, metabolic acidosis, rhabdomyolysis, hyperthermia, and even death (see later section on Other Complications). In these patients, the addition of chemical sedation is highly recommended (see later section on Chemical Restraint). In contrast, the use of extremity restraints alone is often effective in an alcohol-intoxicated patient because the natural progression of alcohol intoxication is sedation and sleep. Intoxicated patients may benefit from a brief period of observation before a decision is made to administer chemical sedation. Most of these patients will fall asleep, thereby obviating the need for sedation and the accompanying risks.

Local Skin

Restraints may cause skin irritation or breakdown. Risk factors include restraints that are too tight and those that have been left on for prolonged periods. Leather restraints are more likely than soft restraints to cause skin damage, particularly in patients who continue to struggle despite being placed in restraints. To help prevent skin complications, use soft restraints whenever possible, avoid overly tight restraints, limit restraint time, and reevaluate the patient frequently. In addition, use chemical restraint judiciously to help avoid skin damage in patients who continue to struggle. It is important to not ignore a patient’s complaints of restraint pain without first evaluating the possibility of an adverse event.

Vascular Compromise

Restraints have the rare potential to impede blood flow to the hands and feet. In extreme cases, this could result in ischemia of the distal end of the extremity. Ischemia is more likely to occur in patients whose restraints are placed too tightly and in those in whom swelling develops as a result of an occult injury. Struggling against the restraints may further increase this risk. A thorough search for occult injuries, attention to proper fit, and limited restraint time will help avoid ischemia. Frequent assessment of pulses, capillary refill, skin color and temperature, and motor and sensory function is also extremely important. As mentioned previously, it is also important to not ignore a restrained patient’s complaints regarding extremity pain without first evaluating for the possibility of ischemia.

Respiratory Compromise

Restraints may impair respiratory mechanics in some patients. This is more likely to occur in patients restrained in the prone or hog-tied positions and in those with underlying pulmonary disease. Patients with chronic obstructive pulmonary disease (COPD) may not tolerate a fifth-point restraint across the chest. Avoid respiratory complications by not restraining patients in the prone or hog-tied position. In addition, use adequate chemical sedation to help negate the need for a supplemental restraint belt in patients with underlying COPD.

Positional Asphyxia

Positional asphyxia is a poorly elucidated respiratory complication often attributed to the use of restraints that results in asphyxia and eventually death. The specifics of respiratory embarrassment...
secondary to physical restraint are vague and unproven and have not been reproduced in volunteers, who do not experience severe pulmonary compromise from restraint. Thus, though often implicated, the exact contribution of restraint to sudden death is unclear. Obesity, underlying cardiac and pulmonary disease, prone positioning, and concurrent stimulant use are thought to be contributing factors.26–28,33–36 The hog-tied position (see Fig. 69.7), in which a patient’s hands and feet are bound behind the back, places a patient at theoretical risk for positional asphyxia and should therefore not be used in the ED.26–28,33–36 Placing restrained patients in the supine position and frequently reevaluating them will help prevent positional asphyxia. Continuous monitoring of respiratory status, including oxygenation, ventilation, and tidal volume, is indicated in all physically restrained and chemically sedated patients, especially those with obesity, COPD, or intoxication from stimulant drugs.

**Cocaine-Associated Agitated Delirium**

*Cocaine-associated agitated delirium* is a syndrome consisting of hyperthermia with delirium and severe agitation that can progress to multisystem failure, coagulopathy, respiratory arrest, and death.28,30,32,14,17 Much of the pathology may be due to cocaine alone, but restrained patients appear to be at particularly high risk for this syndrome. The syndrome was first described in 1985, but its incidence increased significantly in the 1990s because of the popularity of crack cocaine.14,15

The pathophysiology of cocaine-associated agitated delirium is a complex process involving downregulation of dopamine receptors with subsequent dopamine excess during times of cocaine binges.14,18–20 When patients with cocaine-associated agitated delirium are restrained, especially in the prone position, interference with normal respiratory mechanics increases the likelihood of hypoventilation, hypercapnia, and hypoxemia and ultimately leads to asphyxia and death. It has also been suggested that the stress caused by the restraining process increases the risk for fatal cardiac arrhythmias secondary to catecholamine surge in an already cocaine-sensitized myocardium.14 Chronic stimulant use leads to adrenergic-induced cardiomyopathy, which is often clinically silent until the individual is severely stressed. The potential for malignant arrhythmias is unknown, but they have been implicated in some cases of sudden death in restrained patients.

**Metabolic Acidosis**

In patients who have been restrained, continued agitation and struggling can lead to severe metabolic acidosis.15 Their pH is often lower than 7.0. The etiology of this acidosis is unclear but probably involves the production of lactic acid from physical exertion compounded by sympathetic-induced vasoconstriction. Such vasoconstriction may result from agitation or cocaine (and other stimulant) use and is believed to enhance exercise-induced lactic acidosis by impeding clearance of lactate by the liver.41,42 In some patients the buildup of lactate is further increased by the presence of psychosis and delirium, which may alter pain sensation and allow exertion far beyond normal physiologic limits.17 In addition, some restraint positions (e.g., prone, hog-tied) may not allow adequate respiratory compensation, thereby resulting in further enhancement of the acidosis. A common scenario is an out-of-hospital cardiac arrest in which a severely agitated individual suddenly stops struggling and experiences a bradycardic, asystolic death that is not immediately recognized. Such patients may have been subdued by force, by TASER, by mace or by pepper spray, which has led to unproven speculation that these interventions may have caused the change in patient status.

Regardless of the etiology, profound metabolic acidosis has been associated with cardiovascular collapse and sudden and unexpected death in restrained patients.15 Individuals suffer a bradycardic, pulseless electrical activity, asystolic cardiac arrest, and resuscitation is unlikely once cardiac arrest ensues. Patients who remain combative despite restraints, especially those who have used cocaine or other sympathomimetic agents, are at particularly high risk for death.15 Clues to the presence of metabolic acidosis include severe agitation, abnormal vital signs (e.g., persistent tachycardia, tachypnea, and hyperpyrexia), and decreased or concentrated urine output despite adequate IV fluid administration. Laboratory testing, including arterial or venous blood gas analysis, serum electrolytes, and serum creatine phosphokinase, is recommended in patients with signs and symptoms suggestive of metabolic acidosis and in those with potentially lethal co-ingestion (e.g., salicylates and toxic alcohols).

Restrained patients with severe metabolic acidosis should receive aggressive saline hydration and sedation with benzodiazepines to counteract the sympathetic hyperactivity.15 The utility of sodium bicarbonate is unknown and it is probably best reserved for patients with a pH below 7.0.

**Chemical Restraint**

Agitation leading to delirium and culminating in ExDS is a continuum with no rigidly defined parameters. This chapter discusses the treatment of mild agitation, as well as life-threatening ExDS, but it is important to realize that the recommendations are subject to alteration or modification depending on the degree of impairment and the necessity of the clinician to intervene differently in individuals with varying degrees of impairment. The exact role of chemical restraint will always be a clinical judgment call made at the bedside and based on the scenario at hand. It is, however, axiomatic that chemical restraint is mandated when physical restraint and other modalities have failed. Medically compromised patients who appear mentally incompetent, are unable to comprehend or participate in the evaluation or treatment of a serious medical condition, or are otherwise unable to cooperate with required diagnostic or therapeutic medical interventions should be chemically restrained.

Chemical restraint, more aptly termed chemical sedation, describes the act of quelling an agitated patient by the administration of approved sedative-hypnotic, antipsychotic, or dissociative medications. Early, liberal use of appropriate anxiolytic and sedating medications permits thorough evaluation of a patient’s medical condition. In some cases, administration of anxiolytic drugs (e.g., benzodiazepines) may be the optimal treatment of a person with an undifferentiated delirious state such as delirium tremens.43 This is supported by Khan and associates, who identified an association between the use of physical restraint and death in patients with delirium tremens.44 Although a clear causal relationship between restraint and death is lacking, early use of chemical sedation may be safer than physical restraint for treatment of an undifferentiated delirium.

The CMS states that chemical restraint is “a medication used to control behavior or to restrict the patient’s freedom of movement and is not a standard treatment for the patient’s
medical or psychiatric condition.” Curiously, TJC defines chemical restraint as “the inappropriate use of a sedating psychotropic drug to manage or control behavior.” These definitions lack perspective on the use of chemical sedation in the ED, where these medications are typically used after all other measures fail and the health and safety of the patient or staff are threatened. Such statements do not reflect standard care in the ED and should not be interpreted as prohibition of the appropriate short-term use of chemical sedation.

The pathogenesis of agitation is poorly understood; however, the advent of anxiolytic and antipsychotic medications has revolutionized the treatment of acute agitation. Not only has sedation become safer, but many of these new agents also have the added benefit of treating underlying psychotic states. Benzodiazepines, with rare exception, have replaced barbiturates for the treatment of acute agitation. Recently, intramuscular (IM) preparations of “atypical antipsychotic” medications such as olanzapine, ziprasidone, and aripiprazole have provided additional treatment options for rapid control of acute psychosis.

Oral administration is a reasonable first-line option in the cooperative patient and is the preferred route as consent is implied when he or she willingly ingests it. Previous reports have shown that patients prefer oral formulations during the treatment of psychotic episodes because they are participating in their own care plan. Oral lorazepam and oral risperidone have demonstrated increased efficacy when compared with the IM administration of haloperidol. However, in the uncooperative agitated patient, where rapid tranquilization is usually the goal, medications are typically administered IV or IM. Obtaining early IV access allows titration of rapidly acting medications but may not be possible in some patients. When IV access is not possible, IM administration is recommended and, in some cases, has been shown to be as efficacious and safe as IV administration. Other routes (oral, transmucosal) may be used when parenteral administration is not practical or feasible.

An ideal drug for chemical sedation in the ED should have multiple routes of administration (e.g., IV, IM, transmucosal), a rapid onset of action, negligible hemodynamic effects, and a good safety record with minimal adverse effects. Although no medication fits this profile perfectly, with proper patient assessment and careful drug selection, most ED patients can be rapidly and safely sedated. The remainder of this chapter discusses the safety, efficacy, side effect profile, and recommended dosages of the medications most commonly used for chemical sedation. Recommendations for drug selection are also discussed (Table 69.1).

**Other Methods of Drug Delivery**

Under most circumstances, sedative medications are best administered IV. A peripheral vein is adequate but may be difficult to access or maintain in a struggling patient. Although an indwelling catheter is preferable, to titrate escalating doses, the first dose may be directly administered into a peripheral vein via a syringe/needle, so-called “mainlining” (Fig. 69.10). A large extremity vein in the arm or leg is usually available if the extremity can be adequately immobilized. The external jugular vein presents another route. The head can often be more easily stabilized than a muscular extremity, and the external jugular vein is usually quite prominent in a struggling patient. Note that some antipsychotics, such as olanzapine and ziprasidone, have indications only for use IM. Haloperidol is universally administered IV, although it has no formal indication for this route. Intranasal midazolam is another alternative route of administration that may have prehospital utility.

**Indications**

Chemical sedation is used to prevent patients from injuring themselves or others, attenuate psychosis, decrease the time
<table>
<thead>
<tr>
<th>AGENT</th>
<th>DOSAGE</th>
<th>ONSET/DURATION OF ACTION</th>
<th>INDICATIONS</th>
<th>CONTRAINDICATIONS</th>
<th>ADVERSE EVENTS</th>
<th>COMMENTS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haloperidol</td>
<td>Adults: 0.5–10 mg IM/IV/PO q 1–4 hr&lt;br&gt;Children/Adolescents: 0.05–0.15 mg/kg IM/IV q 1 hr (max 5 mg)</td>
<td>Onset: IM/IV: 30–45 min Duration: IM/IV: 4–24 hr</td>
<td>All forms of agitation</td>
<td>Patients with a history of QT prolongation, thyrotoxicosis, Parkinson's disease, and severe hepatic disease</td>
<td>EPSs, QTc prolongation, NMS, hypotension, cholinergic blockade</td>
<td>Can be given alone or in combination with a benzodiazepine; may lower the seizure threshold</td>
</tr>
<tr>
<td>Droperidol</td>
<td>Adults: 5 mg IM/IV</td>
<td>Onset: IM/IV: 3–10 min Duration: IM/IV: 2–12 hr</td>
<td>All forms of agitation</td>
<td>Patients with a history of QT prolongation, thyrotoxicosis, Parkinson's disease, severe hepatic disease, and intoxication with LSD</td>
<td>EPSs, QTc prolongation, NMS, hypotension, cholinergic blockade</td>
<td>Black box warning regarding QT prolongation and serious arrhythmias; rapid-onset IM; use with caution in children</td>
</tr>
<tr>
<td>Lorazepam</td>
<td>Adults: 1–3 mg IM/IV/PO q 30–60 min&lt;br&gt;Children: 0.05–0.1 mg/kg (max 4 mg/dose) Geriatric: &lt; 2 mg/dose</td>
<td>Onset: IM: 15–20 min IV: 2–3 min Duration: IM/IV: 8–10 hr</td>
<td>All forms of agitation</td>
<td>Patients with respiratory depression and pregnant women</td>
<td>Respiratory depression, ataxia, hypotension</td>
<td>First-line therapy for children; use with caution in patients with alcohol intoxication because of the risk for respiratory depression</td>
</tr>
<tr>
<td>Diazepam</td>
<td>Adults: 2–10 mg IV/IM/PO&lt;br&gt;Children: 0.04–0.3 mg/kg per dose IM/IV (max 0.6 mg/kg in 8 hr) Geriatric 2 mg PO</td>
<td>Onset: IV: 1–5 min IM: unknown Duration: IM/IV 15–60 min</td>
<td>All forms of agitation</td>
<td>Patients with respiratory depression and pregnant women</td>
<td>Respiratory depression, ataxia, hypotension</td>
<td>Use with caution in patients with alcohol intoxication because of the risk for respiratory depression</td>
</tr>
<tr>
<td>Midazolam</td>
<td>Adults: 1–5 mg IM/IV&lt;br&gt;Children: 0.1–0.2 mg/kg IV/IM (max 0.5 mg/kg or 10 mg) Geriatric: 1–3 mg IM/IV</td>
<td>Onset: IV: 3 min IM: 5 min Duration: IM/IV: 30–120 min</td>
<td>All forms of agitation</td>
<td>Patients with respiratory depression and pregnant women</td>
<td>Respiratory depression, ataxia, hypotension</td>
<td>First-line therapy for children; rapid onset IM; use with caution in patients with alcohol intoxication because of the risk for respiratory depression</td>
</tr>
<tr>
<td>Agent</td>
<td>Dosage</td>
<td>Onset: IM/IV</td>
<td>Action</td>
<td>Indications</td>
<td>Contraindications</td>
<td></td>
</tr>
<tr>
<td>--------</td>
<td>-----------------------------------------------------</td>
<td>--------------</td>
<td>-----------------</td>
<td>------------------------------------</td>
<td>-----------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Haloperidol</td>
<td>Adults: 10 mg IM q 2 hr, 20 mg IM q 4 hr (max 40 mg/day) Children: not indicated</td>
<td>Onset: 15–30 min Duration: 4 hr</td>
<td>Psychoses, intoxications</td>
<td>Patients with dementia</td>
<td>QT prolongation, somnolence, EPSs Lower incidence of EPS than with haloperidol and droperidol; black box warning regarding use in elderly patients with dementia</td>
<td></td>
</tr>
<tr>
<td>Droperidol</td>
<td>Children: 0.05–0.15 mg/kg IM/IV q 30–60 min</td>
<td>Onset: IM/IV: 30–45 min Duration: IM/IV: 4–24 hr</td>
<td>Psychiatric illness</td>
<td>Patients with dementia</td>
<td>Somnolence, EPSs Lower incidence of EPS than with haloperidol and droperidol; black box warning regarding use in elderly patients with dementia</td>
<td></td>
</tr>
<tr>
<td>Lorazepam</td>
<td>Adults: 2–10 mg IV/IM/PO (max 10 mg) Children: 0.04–0.3 mg/kg IV/IM (max 0.5 mg/kg)</td>
<td>Onset: IM: unknown Duration: IM: 5 min</td>
<td>Psychiatric illness</td>
<td>Patients with dementia</td>
<td>Somnolence, EPSs Black box warning to avoid use for dementia-related psychosis; may cause or increase suicidal ideation</td>
<td></td>
</tr>
<tr>
<td>Ketamine</td>
<td>Adults/children: 1–2 mg/kg IV, 4–5 mg/kg IM</td>
<td>Onset: IV: &lt; 1 min IM: 3–5 min Duration IV:15 min (recovery 1–2 hr) IM: 30 min (recovery 3–4 hr)</td>
<td>Acute agitation</td>
<td>Acute upper respiratory infection, children &lt; 3 mo of age</td>
<td>Emergence phenomenon, potential increased intracranial and intraocular pressure, hypertension, tachycardia, salivation, vomiting, laryngospasm Doses provided are for achieving a dissociative state; use ketamine when rapid sedation is required for lifesaving interventions</td>
<td></td>
</tr>
</tbody>
</table>

*Many of the caveats concerning the safety and efficacy of long-term use cannot be equated to short-term ED use. These medications are currently often used in doses far exceeding those recommended by the manufacturer or under other clinical circumstances; safety and efficacy for short-term sedation have been demonstrated in clinical ED practice. ED, Emergency department; EPSs, extrapyramidal symptoms; IM, intramuscularly; IV, intravenously; LSD, lysergic acid diethylamide; NMS, neuroleptic malignant syndrome; PO, per os (oral).
spent in physical restraint, and calm patients enough to permit a medical history, physical examination, diagnostic testing, and procedures.

Contraindications and Adverse Effects
Absolute and relative contraindications, as well as adverse effects, vary by medication and are discussed separately for each drug.

Neuroleptic Agents
Neuroleptic medications or “typical antipsychotics” have been used safely and effectively for years to manage patients with undifferentiated agitated delirium in the ED. The antipsychotic effects of neuroleptic agents do not usually take place for 7 to 10 days, but the onset of sedation is rapid, thus making them useful to calm an acutely agitated patient. Potent neuroleptic agents such as haloperidol and droperidol are preferred because they lack tolerance after repeated uses, have a low addiction potential, and possess a high therapeutic index. Low-potency neuroleptics such as chlorpromazine are less desirable because of a higher incidence of hypotension, seizures, and anticholinergic effects.

Contraindications
Haloperidol and droperidol are contraindicated in patients with thyrotoxicosis (neurotoxicity may develop), Parkinson's disease, or severe hepatic disease. These drugs can lower the seizure threshold, so they should be used with caution or be avoided altogether in patients with a history of seizures or those who may be at known risk for the development of seizures (i.e., meningitis, sympathomimetic intoxication). Nevertheless, droperidol has been used safely to manage patients with known seizure disorders. In addition, like all neuroleptic agents, haloperidol and droperidol can cause QT prolongation. It is not standard, however, to obtain an electrocardiogram prior to their use. However, they should be used with caution in patients at risk for QT prolongation and torsades de pointes (Box 69.4). Droperidol has also been associated with serotonin syndrome in patients taking lysergic acid diethylamide (LSD) and should be avoided in these patients.

Adverse Effects
Adverse effects common to all neuroleptic agents include extrapyramidal symptoms (EPSs), QTc prolongation, neuroleptic malignant syndrome (NMS), hypotension, and cholinergic receptor antagonism.

EPSs include akathisia (restlessness), dystonia (muscular spasms of the neck, eyes [oculogyric crisis], tongue, or jaw), drug-induced parkinsonism (muscle stiffness, shuffling gait, drooling, tremor), and tardive dyskinesia. These effects are due to the drugs’ antidopaminergic action and, though distressing to the patient, are seldom if ever life-threatening. Anticholinergic agents such as diphenhydramine (25 to 50 mg orally [PO], IM, or IV) and benztropine (1 to 2 mg PO, IM, or IV) are very effective in preventing or minimizing EPSs.

A potentially deadly, but exceedingly rare effect of neuroleptic drug use is prolongation of the QTc interval, which can lead to torsades de pointes, a polymorphic ventricular arrhythmia that can progress to ventricular fibrillation and sudden death. Droperidol has been the most publicized agent associated with QTc prolongation. It is currently the only neuroleptic agent that has received a “black box warning” for its propensity to cause QTc prolongation and sudden death. In addition, a number of case reports and small case series have documented QTc prolongation and torsades de pointes after the administration of haloperidol. Risk factors for QTc prolongation and torsades de pointes are listed in Box 69.4.

Patients in whom symptoms of NMS develop exhibit autonomic instability, including rigidity of the extremities, hyperthermia, and delirium. Treatment includes cooling measures, sedation with benzodiazepines, and in severe cases, bromocriptine, dantrolene, neuromuscular paralysis, and endotracheal intubation.

Hypotension is usually orthostatic in nature and tends to be more pronounced when the drugs are administered IV. In general, haloperidol and droperidol have a lower incidence of hypotension than do lower-potency neuroleptic agents. Likewise, the anticholinergic effects (e.g., confusion, dry mouth, blurred vision, urinary retention) of haloperidol and droperidol are much less severe than those of the lower-potency agents.

Haloperidol
Haloperidol is a very commonly used neuroleptic and a butyrophenone. It is categorized as a high-potency neuroleptic because of its strong antidopaminergic activity. The antidopaminergic activity is responsible for both its intended effects against delusions, hallucinations, and psychomotor agitation and its unintended parkinsonian symptoms. Administration of haloperidol both alone and in combination with benzodiazepines has been evaluated in a large number of clinical trials. These studies have demonstrated the use of haloperidol IM to be both safe and effective for the management of acute agitation from virtually any cause.

Dosage and Administration. Haloperidol can be administered PO, IV, and IM and has a low incidence of oversedation regardless of which route is chosen. Despite a lack of US Food and Drug Administration (FDA) approval for use IV, haloperidol is typically administered IV. The recommended

---

**Box 69.4 Risk Factors for QTc Prolongation and Torsades de Pointes**

**NONPHARMACOLOGIC**
- Congenital long QT syndromes
- Cardiac disorders (ventricular hypertrophy, heart failure, bradycardia)
- Electrolyte imbalance (especially hypokalemia)
- Overdose of an antipsychotic drug
- Female sex
- Use of restraints and psychological stress
- Substance abuse
- Miscellaneous factors (obesity, hypothyroidism)
- Elderly patients
- Renal and hepatic impairment

**PHARMACOLOGIC**

**Pharmacokinetic Factors**
- Inhibition of specific cytochrome P-450 enzymes
- Competition for specific cytochrome P-450 enzymes

**Pharmacodynamic Factors**
- Independent QTc prolongation
starting dosage to achieve sedation is 5 mg IM or IV, titrated to effect. There is no absolute maximum dose, and in cases of severe ExDS, doses of 10 to 30 mg are common. Occasionally, extrapyramidal reactions occur but are easily treated. The dose should be halved when administered to elderly patients. In children 6 to 12 years of age, the dosage is 1 to 3 mg IM every 4 to 8 hours with a maximum of 0.15 mg/kg per day. Children older than 12 years can receive the adult dosage. Haloperidol administered IM has a peak clinical effect within 30 to 45 minutes and may last up to 24 hours when given for acute agitation. Haloperidol is metabolized by the liver and excreted by the kidneys.54

Droperidol
Droperidol, an analogue of haloperidol, is a high-potency butyrophenone with rapid sedating effects. It also possesses significant antidopaminergic activity. Droperidol acquired FDA approval in 1970 first as an antiemetic and antipsychotic agent. Soon thereafter, psychiatric EDs found it useful for chemical sedation.65 Many physicians prefer droperidol to haloperidol because of its more rapid onset and shorter duration of action.

Use of droperidol as a chemical sedative continued until 2001, at which time the FDA issued a black box warning regarding the potential for fatal dysrhythmias.66 Many hospitals and pharmacies subsequently sought to restrict or prohibit its use. However, two large retrospective studies encompassing more than 15,000 patients failed to demonstrate increased morbidity or mortality associated with the use of droperidol for the management of acute agitation.54,65 This controversy prompted an independent review of the data submitted to the FDA that led to the black box warning. The authors of this review found a number of anomalies and duplicate reports. They concluded that droperidol is a safe drug when used at the recommended dosage (i.e., 5 to 10 mg).67 Despite an apparent lack of evidence behind the black box warning, use of droperidol has declined dramatically.68

Studies comparing a variety of agents for rapid sedation have found that droperidol provides more rapid and effective control than do lorazepam,69 haloperidol,49 midazolam,70,71 and ziprasidone.72 In addition, droperidol has been proved to be safe in patients with head injuries, alcohol and cocaine intoxication, and seizure disorders.54

Dosage and Administration. The initial dosage used for sedation is 5 mg IV or IM. Its onset of action is 3 to 10 minutes with a peak clinical effect achieved in 30 minutes. The elimination half-life is 2 to 4 hours, but the sedative effects of droperidol may last up to 12 hours.72

Benzodiazepines
All benzodiazepines enhance the neurotransmission of γ-aminobutyric acid and thereby result in anxiolysis, sedation, hypnosis, and muscle relaxation. This combination makes them an excellent choice for tranquilization of agitated patients. Because of a desirable safety profile, benzodiazepines are an excellent choice for sedating medically undifferentiated patients. The differences in clinical effects (e.g., onset, duration, adverse effects) are primarily related to dosage, route of administration, and pharmacokinetics. Benzodiazepines can be used as single agents or in combination with an antipsychotic drug. Lorazepam and midazolam are the benzodiazepines most commonly used for chemical sedation. This is probably due to their rapid and predictable absorption when given IV or IM, as well as a long history of safety and efficacy. Unlike neuroleptics, benzodiazepines do not treat underlying psychiatric disorders. Some clinicians prefer to administer diazepam IV for the treatment of delirium related to sedative-hypnotic withdrawal.43

Contraindications
There are few contraindications to the use of benzodiazepines as a chemical sedating agent. Because of the possibility of respiratory depression, use benzodiazepines with caution in patients in respiratory distress or identified pulmonary pathology like obstructive sleep apnea.

Adverse Effects
As discussed previously, benzodiazepines may cause respiratory depression. Additionally, hypotension, deep sedation, and paradoxical agitation have been reported. However, when administered in the doses recommended for agitation, adverse events are rare, thus making benzodiazepines the drugs of choice in most circumstances.

Respiratory compromise is dose dependent and typically occurs only in the presence of other respiratory depressants.73,74 Because of an increased risk for respiratory depression, administer benzodiazepines cautiously to elderly patients and those with underlying pulmonary diseases such as apnea and COPD. In healthy patients, particularly those suffering from agitated delirium, respiratory depression is very unlikely to occur, even when large doses of benzodiazepines are used. If available, end-tidal carbon dioxide monitoring (e.g., capnography) may assist the practitioner in detecting the onset of respiratory depression before it becomes clinically significant.75,76

The most appropriate benzodiazepine is the one that the clinician is most comfortable administering and that has a duration of action most appropriate for the clinical situation. Titrate all benzodiazepines administered IV until the desired level of sedation is reached. In the setting of severe delirium, there is no maximum dose described for any benzodiazepine, and doses well in excess of the manufacturer's recommendation, and even those considered toxic under other circumstances, are commonly used. One difference that does bear mentioning is the use of propylene glycol, a solvent needed to keep non–water-soluble benzodiazepines (e.g., lorazepam, diazepam) in solution. In large doses, propylene glycol can cause an increased osmolar gap and metabolic acidosis and may precipitate or contribute to hypotension in some patients.

Lorazepam
Lorazepam is the benzodiazepine most frequently studied for the management of acute agitation. It enjoys popularity as part of the well-known “five and two” treatment regimen, which consists of haloperidol, 5 mg IM, and lorazepam, 2 mg IM.77 In a number of clinical trials, lorazepam was shown to be an effective drug for rapid chemical sedation.59,62,63,69 In these studies, lorazepam had no extrapyramidal effects and was better tolerated than the neuroleptics. However, when compared with haloperidol, droperidol, midazolam, and a combination of haloperidol and lorazepam, the onset of sedation after administering lorazepam IM was more delayed.59,62,69

Dosage and Administration. When used alone, lorazepam is usually given in 2- to 4-mg doses and can be administered PO, sublingually, IM, IV, or rectally. No maximum dose has been established. Following an injection of lorazepam IM, adequate sedation is usually achieved in 30 to 45 minutes.59
When given IV, sedation occurs in 15 to 20 minutes. 62 The elimination half-life is 12 to 15 hours, which produces a duration of effect of 8 to 10 hours. This makes lorazepam a better choice when long-term sedation is the goal. Lorazepam is rapidly conjugated to an inactive glucuronide. This does not require involvement of the cytochrome P-450 system, so lorazepam has few drug-drug interactions. Preparations of lorazepam intended for use IV or IM must be refrigerated, thus potentially limiting its use in underdeveloped countries and in the prehospital setting.4

Midazolam

Midazolam has been shown to be effective for rapid sedation. 63,64,70,71 It has a more rapid onset and shorter duration of action than other benzodiazepines do, which makes it a good choice when rapid short-term sedation is desired. Midazolam also compares favorably with haloperidol, droperidol, and ziprasidone for the treatment of acute agitation in the ED. 61,62,71 With the exception of droperidol, which has a similar time until onset (5 to 10 minutes), midazolam has a more rapid onset of sedation than the other drugs in these studies. The studies also noted that midazolam, as expected, has a shorter duration of sedation.

**Dosage and Administration.** The initial dosage of midazolam for an agitated adult is 5 mg IV or IM, and it may be repeated at 5- to 10-minute intervals. No maximum dose has been established. Nasal administration is also an option. The onset of sedation occurs approximately 3 minutes after IV administration, 5 minutes after an IM injection, and 15 minutes after nasal administration. Midazolam is hydroxylated by the cytochrome P-450 system to its primary metabolite, α-hydroxymidazolam, which undergoes glucuronide conjugation before being excreted in urine. Its duration of action is between 30 and 120 minutes and does not vary significantly by route of administration.62,78

Diazepam

Diazepam has a long history of safe use in the management of agitated delirium, especially delirium tremens and alcohol withdrawal.4,5,74 For patients experiencing delirium tremens the usual dose is 5 to 10 mg IV every 5 to 10 minutes until the desired level of sedation is achieved (lower doses are appropriate for less severe agitation or agitation from other causes). No maximum dose has been established, and doses of up to 2000 mg have been used safely over a 24-hour period in patients experiencing delirium tremens.4,81 Because of erratic absorption, IM administration of diazepam is not recommended. Following the administration of diazepam, peak sedation is reached in approximately 5 to 6 minutes, which allows additional titrated doses if the desired clinical effect is not achieved initially.

**Atypical Antipsychotic Agents**

Atypical antipsychotic agents have high affinity for 5-HT (serotonin) receptors and less affinity for D1 and D2 receptors. As a result, they have a lower incidence of EPSs than haloperidol and droperidol.84 In the past, medications in this class were available only in an oral formulation, thus limiting their use in the management of acute agitation. Recently, formulations of ziprasidone, olanzapine, and aripiprazole have been developed and are intended for IM administration. When administered IM, all three drugs are effective for the treatment of acute agitation.81 However, the delay in onset from IM medications may be a disadvantage of these medications in circumstances that require rapid sedation. Often sedation is markedly prolonged from the relatively high doses required. However, the combination of IM administration and the low incidence of EPSs makes these newer agents an attractive option for rapid sedation of ED patients with undifferentiated agitation. This is particularly true for patients with a history of mental illness, for whom this class of medications is now considered appropriate first-line management.82

**Contraindications**

In 2005, a meta-analysis of placebo-controlled trials demonstrated an increased risk for death associated with the atypical antipsychotic agents used to treat elderly patients with dementia-related psychosis.83 This report led the FDA to issue a black box warning regarding the use of atypical antipsychotics for “behavioral disorders” in elderly patients with dementia. Such warnings have not resulted in the prohibition of such agents for short-term use in the ED. The FDA has also advised caution with ziprasidone because of its tendency to prolong the QTc interval, especially when used in patients taking other drugs or with medical conditions that increase the risk for prolongation of the QTc interval (see Box 69.4).81 Ziprasidone prolongs the QTc interval more frequently than haloperidol, droperidol, and olanzapine do, but the degree of prolongation is considered minor and is rarely greater than 500 msec.81 To date, there have been no clinical reports of adverse events as a result of QTc prolongation with the short-term ED use of atypical antipsychotic agents. Experience with these agents in undifferentiated acutely agitated patients in the ED is limited, but their use is increasing, and although initial reports are supportive, they are not yet definitive.

**Adverse Effects**

Atypical antipsychotic agents may cause somnolence, EPSs (though less often than with haloperidol and droperidol), QTc prolongation, and rarely, anticholinergic symptoms and NMS.

**Ziprasidone**

Ziprasidone is a benzylisothiazolypiperazine antipsychotic agent whose effects are the result of dopamine and serotonin 5-HT2A receptor antagonism.5,81,86 This was the first atypical antipsychotic agent available in a fast-acting preparation administered IM. In a double-blind, randomized study, Martel and associates noted that ziprasidone was as effective as midazolam and droperidol in controlling acute agitation. Patients receiving ziprasidone and droperidol took longer to be sedated (30 minutes as compared with 15 minutes for midazolam) but were more deeply sedated at 60 and 120 minutes.79 In a prospective, open-label study, ED patients receiving ziprasidone exhibited progressive improvement in anxiety, hostility, and cooperativeness starting at 15 minutes and continuing through the 90-minute study period.86 In an observational study of agitated psychiatric ED patients with nonspecific psychosis, alcohol intoxication, or substance-induced psychosis, ziprasidone, 20 mg IM, was effective in sedating patients as quickly as 15 minutes.86 Ziprasidone has not been approved by the FDA for the treatment of dementia-related psychosis.87

**Dosage and Administration.** The recommended dosage of ziprasidone is 10 mg IM, which can be repeated at 2 hours, or 20 mg IM, which may be repeated at 4 hours. The drug...
reaches peak plasma concentrations in 30 to 45 minutes and has an elimination half-life of 2 to 4 hours. Following an IM injection, sedation usually begins within 15 to 30 minutes and peaks at approximately 2 hours. The clinical effects usually last at least 4 hours.87

Olanzapine
Olanzapine is a second-generation thienobenzodiazepine antipsychotic agent that is thought to exert its effects through antagonism of both dopamine and serotonin type 2 receptors. Olanzapine in an IM formulation has been approved by the FDA for the treatment of agitation in acutely psychotic patients. IM olanzapine is comparable to haloperidol or lorazepam monotherapy for acute agitation associated with schizophrenia and dementia.50,58,87 and superior to lorazepam monotherapy in the management of agitation associated with bipolar disorder.90 To date, there have been no clinical trials using IM olanzapine for undifferentiated agitation in the ED, but its use is increasing.

**Dosage and Administration.** The recommended dosage is 5 to 10 mg IM. Additional doses may be considered 2 to 4 hours after the preceding dose with a maximum recommended dose of 30 mg/day IM. Olanzapine reaches peak plasma concentrations in 15 to 45 minutes and has an elimination half-life of 21 to 54 hours. The onset of sedation usually begins 15 to 30 minutes after IM administration and typically lasts at least 2 hours. In some patients, the clinical effects have lasted as long as 24 hours. The drug is metabolized via direct glucuronidation and cytochrome P-450-mediated oxidation.91

Aripiprazole
Aripiprazole is an atypical antipsychotic that acts at both serotonin and dopamine receptors. Rates of EPSs are lower than with other antipsychotics, which may be a result of the drug’s partial agonistic activity at dopamine receptors. IM administration has been approved for acute agitation related to schizophrenia and bipolar mania. To date, there have been no clinical trials using aripiprazole IM for undifferentiated agitation in the ED.

**Dosage and Administration.** The recommended dosage is 5.25 to 9.75 mg IM. Additional doses may be considered 2 to 4 hours after the preceding dose with a maximum recommended dose of 30 mg/day IM. Peak serum levels occur within 1 to 3 hours, with an elimination half-life of approximately 75 hours.

**Dissociative Agents**
Ketamine
Ketamine is a dissociative agent that has been used safely throughout the world for major surgery and with minimal monitoring.92 Many clinicians are familiar with ketamine as a safe and rapidly effective dissociative agent for tracheal intubation and for both adults and children undergoing painful procedures in the ED. When given IM or slow IV (e.g., greater than 60 seconds), ketamine has no significant adverse effects on respiration. It inhibits the reuptake of catecholamines promoting bronchodilation and increases in both heart rate and blood pressure. Commonly raised as a caution, there is no proven issue with ketamine causing harmful increased intracranial pressure.91,94 Possible side effects of ketamine include salivation, vomiting, laryngospasm, and emergence phenomena consisting of nightmares, short-lived bizarre thoughts, and hallucinations. Fortunately, serious emergence phenomena have not been described as significant issues following ED use.

Literature supporting ketamine for the control of agitation and delirious patients has grown over the past decade. The drug’s pharmacologic profile lends itself for use in acute, potentially violent situations. For example, ketamine has been used successfully in aeromedical transport as a sedative agent for patients with agitation.93 It has also been effective in the prehospital management and transport of patients with ExDS and in combative trauma patients (including those with head injury),96,97 acutely agitated cocaine-intoxicated patients,98 and agitated suicidal patients.99

**Dosage and Administration.** The dose of ketamine to produce profound dissociation is 1.5 to 2.0 mg/kg IV or 4 to 5 mg/kg IM. IM administration is ideal in the ED when access for IV administration is not readily available. The anterior aspect of the thigh is a preferred site of injection for rapid tranquilization. Time of onset may vary slightly, but both routes have a rapid onset of action (30 seconds IV, 3 to 5 minutes IM). At the current time, ketamine is undergoing resurgence in popularity among emergency physicians, and the role of ketamine for the chemical control of acutely agitated and delirious patients in the ED is evolving.

**Choosing the Best Agent**
Undifferentiated Agitation
The safest agent for undifferentiated agitation in the ED has traditionally been a benzodiazepine administered IV, often combined with judicious doses of haloperidol. Chan and associates demonstrated the combination of an antipsychotic and midazolam had a shorter time to sedation than midazolam alone.100 Ketamine and newer atypical antipsychotics are gaining favor, but their track record is short. Administering escalating doses of benzodiazepines is a prudent choice in such circumstances when the clinician is comfortable prescribing a drug from this class. There is little concern for respiratory or cardiovascular depression with the prudent use of carefully titrated IV benzodiazepines in the monitored setting. Fig. 69.1 suggests a possible management approach to an acutely agitated patient in the emergency setting. It must be emphasized that benzodiazepine doses considered exceeding high or even toxic by other standards are routinely and safely administered in the ED when aggressive sedation and chemical control are mandated as clinical priorities. In severe cases, benzodiazepines alone may not be totally effective.

**Agitation Caused by Alcohol and Drugs of Abuse**
For patients who are suspected of intoxication from alcohol or other sedative agents, haloperidol, droperidol, or ziprasidone will provide rapid, safe, and effective tranquilization.54,65,71 Benzodiazepines should be administered with caution to alcohol-intoxicated patients and those taking sedative agents because of the possibility of respiratory depression.72,84 In contrast, benzodiazepines are the drugs of choice in patients who are agitated as a result of alcohol or benzodiazepine withdrawal. Large doses may be required, and the safety profile is wide. Patients who are agitated because of sympathomimetic agents such as cocaine, methamphetamine or hallucinogens, such as PCP or LSD, may be treated safely with large doses of benzodiazepines, butyrophenones, or a combination of the two.54,99,99 Droperidol has been associated with serotonin syndrome in patients taking LSD and should be avoided in these patients.51,55
Agitation Caused by Medical Illness
In patients whose agitation is due to medical illness, treatment should be aimed at correcting the underlying pathology. If rapid sedation is required, typical antipsychotics or benzodiazepines should be used as first-line therapy. If the patient is frail or elderly or is known to have renal impairment, consider using smaller doses of a single agent.

Agitation Caused by an Underlying Psychiatric Disorder
Patients with an established psychiatric history and agitation attributed to schizophrenia, schizoaffective disorder, or the manic phase of bipolar disorder may be treated with typical antipsychotic agents, atypical antipsychotic agents, or benzodiazepines. However, a growing body of evidence seems to support the use of atypical antipsychotic agents in this circumstance.104,60,70,86,88,89,98,101,102

Agitation in Children
Because of years of experience and a proven safety record, benzodiazepines are considered first-line therapy for the management of agitation in children. The IM dose of lorazepam is 0.05 to 0.1 mg/kg and the IM dose of midazolam is 0.1 to 0.2 mg/kg. Neuroleptic agents have also been used to manage agitation in children. The IM dose of haloperidol is 0.025 to 0.075 mg/kg, with a maximum dose of 2.5 mg. Children older than 12 years can receive the adult dose, usually 2.5 to 5.0 mg IM (see Table 69.1). Although droperidol is a highly effective drug for rapid sedation of adults, there is a paucity of literature supporting its use in children. The pediatric dose of droperidol is 0.03 to 0.07 mg/kg with a maximum initial dose of 2.5 mg.11 Combinations of haloperidol and lorazepam have also been used successfully in children.111

Agitation in Pregnancy
Psychotropic and anxiolytic medications should be used during pregnancy only when the potential risk to the fetus from exposure is outweighed by the risk of not treating the disorder in the mother.104 Based on years of accumulated clinical experience but very little scientific data, conventional antipsychotic agents such as haloperidol and droperidol (pregnancy class C) are recommended to control agitation in pregnant women.105 In pregnancy, benzodiazepines (pregnancy class D) may be associated with teratogenicity (especially craniofacial abnormalities) when used for an extended period.106 Although clear causation has not been proved, benzodiazepines are generally avoided during the first trimester to minimize the risk for fetal malformation.107 Judicious benzodiazepine administration should be considered a viable option in the acute ED setting in the second and early third trimesters.

Agitation in Older Patients
Patients 65 years or older are particularly susceptible to adverse drug reactions because of coexisting medical illness, use of multiple prescription medications (which increase the risk for drug-drug interactions), and age-associated changes in pharmacokinetics and pharmacodynamics. Research suggests that conventional antipsychotic medications such as haloperidol and droperidol are safe and effective for both psychotic symptoms and nonpsychotic agitated behavior.108 Low doses (e.g., half the usual dose) of benzodiazepines can also be used but require close observation for respiratory depression. Continued use (> 8 to 10 weeks) of atypical antipsychotic agents has been associated with increased rates of death in cases of dementia-related psychosis.83

Conducted Electrical Weapons (CEWs)
CEWs, including TASERs (TASER International, Inc., Scottsdale, AZ) and stun guns, are used by law enforcement agencies throughout the world. These nonlethal weapons use a temporary high-voltage low-current electrical discharge to overcome a body’s voluntary muscle-triggering mechanisms, which results in widespread involuntary muscle contractions that incapacitate the victim. The current is delivered by direct contact with a handheld device (i.e., electric shock prods) or via a small dart-like electrode fired from a gun using small gas charges (e.g., TASER) similar to some air rifle propellants. There are also electrical weapons that cause intense pain without incapacitating the target, so-called drive stun devices. Law enforcement and correctional personnel typically use drive stun devices as a pain compliance technique. A TASER may deliver approximately 1200 V with direct skin contact.109

The exact street scenario of ExDS, physical restraint, and a TASER-linked death will probably never be studied with unchallenged scientific accuracy, and hence one is left with animal and human volunteer data to intuit the cause and effect of untoward events. Despite claims of lethality in the press, emotional testimony and unscientific reports from human rights organizations, and poorly documented claims in case reports, there have been no documented deaths directly and indisputably attributable to a TASER discharge in humans.110 Volunteer studies and animal models failed to provoke cardiac arrest, significant cardiopulmonary, or metabolic derangements after standard electrical discharges, even when animal models included stimulant toxicity and human volunteers were exercised to exhaustion.109 Nevertheless, CEWs have been temporarily linked to a number of fatalities.109–111 The majority of these cases involved sympathomimetic drug intoxication (e.g., cocaine, methamphetamine), prolonged physical exertion, or both.111 The role of severely altered metabolic and autonomic parameters, drug intoxication, or underlying comorbid conditions, including cardiomyopathy, obscures the exact contribution of CEW use in unexpected deaths. A complete discussion of this topic is beyond the scope of this chapter, but it has been well reviewed elsewhere.112,113 Darts from a CEW often penetrate the skin and must be removed, so the remainder of this section focuses on the procedure for removing embedded TASER electrodes.

Electronic Control Devices
TASER is an acronym for “Thomas A. Swift’s Electric Rifle” and has been in existence since the early 1990s (Fig. 69.11). More than 11,000 law enforcement, correctional, and military agencies in 44 countries deploy TASER devices, and many municipalities in the United States allow civilians to purchase and carry these weapons for personal protection.114 A TASER uses compressed nitrogen to propel two electrode-tipped barbs at 180 ft/sec at the target. The electrode-tipped barbs are attached to the electric device via two thin 21-foot wires and are similar in size to a No. 8 fishhook measuring 4 mm in length (Fig. 69.12). The barbs may attach to clothing and fail to penetrate the skin, or they may become embedded in skin and must be removed.
the barb was difficult to remove, or if there is concern for a
retained foreign body. Tetanus status should be reviewed and
updated if necessary. Significant infection after barb removal
is rare, and prophylactic antibiotics are unnecessary. Following
removal of a TASER dart, the disposition of the patient is
based on the individual social scenario, and long-term ED
observation is not required.

Complications
The risk for penetrating injury to the heart, lungs, or bowel
from a TASER device is minimal, because of the small size of
the barb. There are, however, increasing case reports of scrotal
trauma, frontal sinus penetration, intraocular, and intracranial
injuries resulting from TASER barbs.\(^{116-123}\) Management will
depend on the specific structure affected and can carry sig-
nificant morbidity if not recognized. A barb embedded in a
vascular structure can probably be removed with manual traction
followed by direct pressure on the wound because the size of
the barb is similar to the size of devices used to obtain central
venous access.\(^{115}\) Consultation with an appropriate consultant
can help manage these rare but potentially serious penetrating
injuries.

Severe involuntary muscle contractions from the electrical
discharge has been implicated as a cause of acute thoracic
compression fractures.\(^{124,125}\) Based on a literature review in
2011 by Vilke and colleagues, there is currently no indication
for routine diagnostic or laboratory testing in asymptomatic
patients following short-duration exposure (<15 seconds) to
a CEW.\(^{126}\)

TASER Use in the ED
Although data are difficult to confirm, it has been estimated
that approximately 150 hospitals allow hospital security guards
to carry TASERS. Those equipped with TASERS should have
specific training, and policies on the use of the TASER in the
hospital setting should be in place and training periodically
updated. CMS regulations do not disallow TASERS in the
hospital, but CMS and most states do not sanction the use of
such devices to restrain a patient or to make them comply
with medical treatment. Whenever possible, the least restrictive
methods should be used to de-escalate aggressive behavior, or
to calm agitated or disruptive individuals, such as a quiet and
low-stimulation environment, reasonable bargaining, redirection
of the patient, involvement of family, reality orientation, talk
down, or a show of force. It is a gray area, indeed, as to when,
or to what extent, any intervention is considered necessary to
restrain a patient, or to protect a patient or medical personnel
from harm. Every situation is somewhat unique so dogmatic
approaches cannot be used. When a mentally incompetent or
potentially suicidal patient wants to leave the ED against medical
advice, and does not have insight into the adverse effects of
such an egress, involuntary commitment is initiated. Effective
measures are usually initiated by the emergency physician
because psychiatric evaluation on such short notice is impractical
or unavailable and important decisions must be made imme-
diately with limited data. Such patients may assault those
attempting to keep them from leaving the ED in their
incompetent mental state, when medical consequences, suicide,
or harm to others could be the ultimate outcome.

Per CMS guidelines, the use of weapons by security staff
is considered a law enforcement action, not a health care
intervention. CMS does not support the use of weapons by
any hospital staff as a means of subduing a patient to place
that patient in restraint or seclusion. CMS interpretive guidelines to section 482.13(e) of the State Operations Manual say: “If a weapon is used by security or law enforcement personnel on a person in a hospital (patient, staff, or visitor) to protect people or hospital property from harm, we would expect the situation to be handled as a criminal activity and the perpetrator placed in the custody of local law enforcement.”

In one small nonvalidated study, Ho and colleagues concluded that the introduction of the TASER into the health care setting (a large urban tertiary care teaching hospital) demonstrated the ability to avert and control situations that could result in injury to medical personnel and patients, by simple TASER presentation or rarely actual use. In that study there was a reduction in personnel injury rates and the contention that one suicide was averted. Further study is required before definitive statements can be made. In summary, the use of the TASER in the ED has not yet been clarified, and there are advocates as well as critics. Any weapon, including the TASER, should not be used to force a patient to comply or to induce restraint in the absence of reasonable suspicion of impending harm or actual assault of medical personnel.

ACKNOWLEDGMENT

We would like to thank the previous author, Adam K. Rowden, for his work on earlier editions of this chapter.

REFERENCES ARE AVAILABLE AT www.expertconsult.com
REFERENCES


64. American Society of Health System Pharmacists: Butyrophenones, Bethesda, MD, 2006, ASHSP.
Noncardiac Implantable Devices

Paul Jhun and Christopher R. Peabody

In addition to cardiac pacemakers and defibrillators, a number of noncardiac devices have been developed for electronic neuromodulation and drug delivery. Although these devices are placed by a variety of subspecialists for the treatment of chronic illnesses, if the devices malfunction, patients may arrive at the emergency department in an acute state, thereby necessitating intervention by the emergency provider.

INSULIN INFUSION DEVICES

Background

External insulin infusion devices have become increasingly popular since their introduction in 1974. By 2007, there were more than 375,000 external insulin infusion pumps in use in the United States.

Anatomy

An external insulin infusion pump device consists of a portable, programmable infusion pump connected to a subcutaneously implanted catheter that is maintained in place with adhesive tape (Fig. 70.1). The implanted catheter site varies, but it is commonly placed in the subcutaneous tissue of the abdomen in adults and the buttocks in young children. The thighs, hips, and upper part of the arms are other sites. It is recommended that the implanted catheter be replaced every 2 to 3 days.

Device Complications

In 2010, the Food and Drug Administration (FDA) published a panel report that highlighted problems associated with insulin infusion devices. From the top five device manufacturers, the FDA noted 16,640 adverse events, including 310 deaths, 12,093 injuries, and 4294 malfunctions. Box 70.1 lists the most frequently reported problems with devices in descending frequency. Box 70.2 lists the most frequently reported patient-oriented adverse reactions, which include hyperglycemia and hospitalization. Similar findings were reported in a 2014 prospective study, in which the most clinically significant adverse events related to insulin infusion devices were hyperglycemia and ketosis.

Of the 310 deaths reportedly related to insulin infusion pumps in the FDA report, the vast majority of problems with the device were not known to the patient or providers at the time, and the root cause of failure of the device was not identified by the manufacturer. In 29 deaths, problems with the device that were identified included overinfusion, bent cannula, disconnection, pump alarming, failure to deliver, suspected electromagnetic interference, and display failure.

Procedure

If the device appears to be functioning normally, the American Association of Clinic Endocrinologists recommends that, should the patient be unable to manage his/ her own pump while in the emergency department (ED), “the specialist(s) responsible for the patient's ambulatory pump management should be contacted promptly to make decisions about infusion adjustments.” Moreover, device malfunction should be considered in patients with insulin infusion devices who present to the ED with symptoms that may be related to blood glucose. In addition to medical management, troubleshoot the device and remove it in cases of uncertainty or emergency.

To remove the catheter, simply peel off the adhesive and embedded catheter together to discontinue the flow of injected medication into the patient. There are a variety of proprietary pump manufacturers, each with their own device programming. Call the appropriate manufacturer to troubleshoot the device, visit the manufacturer's website for an online troubleshooting manual, or simply instruct the patient to discontinue use of the device and return to a standard calculated insulin schedule by manual injection until the device-related complication is investigated and resolved.

INTRATHECAL DRUG DELIVERY SYSTEMS

Background

Intrathecal (spinal canal/subarachnoid space) drug delivery systems (IDDSs) have been in clinical use since the 1980s. The FDA approved the use of intrathecal baclofen in 1992 for severe spasticity secondary to spinal cord injury, multiple sclerosis, cerebral palsy, or stroke. Subsequently, the FDA approved the use of intrathecal morphine in 1995 and intrathecal ziconotide in 2004 for chronic pain refractory to traditional medical therapies. Ziconotide is a non-opioid calcium channel blocker, but its use may be complicated by confusion, somnolence, and other neurologic side effects. Although limited data demonstrate efficacy of intrathecal morphine and ziconotide in relieving refractory pain and efficacy of intrathecal baclofen in reducing spinal cord injury-induced spasticity, more quality clinical evidence is needed. IDDS devices are also used for the delivery of chemotherapeutic medications for specific oncologic conditions. Moreover, there are several off-label uses of various intrathecal medications. Pumps are often used to deliver analgesics to patients with intolerable side effects from oral or parenteral analgesics, such as nausea, pruritis, and cognitive dysfunction. Other medications used include butyricaine, hydromorphone, fentanyl, sufentanil, clonidine, midazolam, and meperidine. Some patients will receive a combination of medications. There is minimal data comparing various medications and current regimens have been empirically derived.

The advantage of IDDSs for chronic pain is the ability to get medication directly to receptors in the spinal cord by placing the medication into the subarachnoid space. For intrathecal morphine infusions, approximately 1% of the total daily morphine dose is a standard starting point. Such small intrathecal doses reduce systemic concentrations and minimize side effects. Patients in chronic pain may also use oral analgesics on a rescue basis.
Anatomy

There are two types of IDDS: continuous infusion devices and programmable devices. In the United States, the Codman 3000 (Codman & Shurtleff, Inc., Raynham, MA) is an FDA-approved continuous infusion device, whereas the Medtronic (Minneapolis, MN) SynchroMed II and the Flowonix (Mount Olive, NJ) Promeda are FDA-approved programmable devices. IDDS devices consist of an infusion pump connected to an intrathecal catheter (Fig. 70.2A). Codman's infusion pump is available with a refillable reservoir of either 15, 30, or 50 mL. Medtronic's refillable reservoir is available in either 20 or 40 mL, whereas Flowonix's refillable reservoir has a 20-mL capacity. The Codman 3000 does not utilize a battery and does not need to be replaced. SynchroMed II's battery life ranges between 4 to 7 years, whereas Flowonix's battery life is reportedly more than 10 years. Normal refill intervals vary based on usage, typically on the order of every few months.

To place an intrathecal catheter, a small incision is made in the back and the catheter tip is placed into the cerebrospinal fluid. The catheter is advanced to the desired level of medication.

---

**Figure 70.1** External insulin infusion pump. Components include a subcutaneously implanted catheter (A) and a programmable infusion pump (B) (OneTouch Ping, Animas Corporation, West Chester, PA). To discontinue the flow of insulin into the patient, simply peel off the adhesive and embedded catheter together. Also shown is a wireless continuous glucose monitoring receiver (C), along with its sensor and transmitter (D) (extCom SEVEN PLUS, DexCom, Inc., San Diego, CA).

**Figure 70.2** A, The Medtronic (Minneapolis, MN) SynchroMed II Programmable Infusion Pump is used primarily for intrathecal (cerebrospinal fluid) administration of opioids, baclofen, calcium channel blockers, or other medications used alone or in combinations. B, The pump is usually implanted in a subcutaneous pocket in a lower abdominal quadrant, with the catheter tunneled subcutaneously to an appropriate lumbar interspace. Severe and difficult to treat withdrawal may occur if opioids or baclofen infusions are interrupted by a drained drug reservoir or pump malfunction.
delivery, based on symptoms. The catheter is then tunneled around the abdomen, placed in the lower abdominal wall, and attached to the subcutaneous pump. The pump has a port in the center that can be accessed by a needle placed through the skin. Medications are placed into the pump and refilled as needed (see Fig. 70.2B).

Continuous infusion devices, such as those made by Codman, are regularly refilled via manual bolus injections. Programmable devices are interrogated and programmed via an external manufacturer-specific device.

Device Complications

IDDS device complications requiring surgical intervention were found to occur with a frequency of 10.5% per year, with 35% pump related and 65% catheter related.13,16

The most frequent and serious adverse events related to the device and implant procedures are catheter dislodgement from the intrathecal space, catheter fracture, implant site infection, and meningitis. Internal device programming errors that cause underdosing or overdosing are less common. If the drug supply is depleted, systemic symptoms related to acute drug withdrawal will occur. This is characteristic for the specific drug being delivered and should be anticipated.

A unique complication related to IDDSs that is the subject of a warning by the FDA is the formation of a granuloma at the tip of the intrathecal catheter. This can obstruct infusion of the medication and lead to withdrawal symptoms.17,18 Interestingly, catheter-tip granuloma formation appears to be primarily associated with intrathecal opioids, and most granulomas regress spontaneously when the offending agent is discontinued.13

Common adverse effects related to intrathecal opioids include centrally mediated respiratory depression, nausea, vomiting, sedation, pruritis, constipation, urinary retention, cognitive impairment, and headache.10–21 One study using Medtronic’s device registration system found a mortality rate of 3.9% at 1 year in IDDS patients being treated with intrathecal opioid. The deaths were associated with respiratory arrest, although not causally proven.22

Intrathecal ziconotide may result in dizziness, nausea, vomiting, urinary retention, gait imbalance, nystagmus, and confusion. Although respiratory depression is uncommon, ziconotide may be associated with rare cases of rhabdomyolysis and psychosis.8,9,19

Common adverse effects related to intrathecal baclofen include hypotonia, somnolence, headache, dizziness, urinary retention, nausea, and paresthesias. It is more common to have an underdose of medication due to pump malfunction or a break or dislodgement of the catheter than to have an overdose. In the ED, it is usually impossible to accurately determine the exact cause of reduced medication delivery. In such cases, withdrawal syndromes will occur. In patients experiencing opioid, baclofen, or other medication withdrawal, treatment may be quite difficult because large doses of systemic medications are required to equal the effects of intrathecal medications. Hospital admission and careful monitoring is often the most prudent tactic. Acute intrathecal baclofen withdrawal, which may resemble alcohol or benzodiazepine withdrawal can be serious and difficult to manage.14,19

Baclofen withdrawal can lead to hemodynamic instability or seizures. As withdrawal is due to loss of medication at the spinal level, it may not be possible to safely deliver enough medication systemically to quell baclofen withdrawal symptoms. A propofol infusion has been suggested to treat baclofen withdrawal, but the patient may need to be intubated to facilitate a sufficient amount of propofol. Specialty consultation is required to address most complications of intrathecal pumps.

Procedure for Treating Medication Overdose

When there is concern for medication overdose secondary to the rare malfunction of the device, in addition to emergency medical management, the emergency physician can empty the pump reservoir by using the steps outlined in Box 70.3.

One may also access the manufacturer-specific online manuals to review the steps necessary to remove the contents of the pump reservoir and thus prevent further infusion of intrathecal medication (Fig. 70.3).13 Published guidelines suggest, in the absence of contraindications, withdrawal of an additional 30 to 40 mL of cerebrospinal fluid (CSF) by lumbar puncture to reduce the drug’s concentration in CSF.21

**VAGAL NERVE STIMULATORS**

**Background**

Since the 1930s, several studies have shown the effects of vagal nerve stimulation on cerebral activity. In 1985, Zabara demonstrated the anticonvulsant effect of vagal nerve stimulation through animal studies.24 In 1997, the FDA approved use of the vagal nerve stimulator (VNS) as an adjunctive treatment of medically refractory partial-onset epilepsy.25 Subsequently, in 2005 the FDA approved use of the VNS as an adjunctive therapy for treatment-resistant depression.26,27
1502  SECTION XII  SPECIAL PROCEDURES

**Device Complications**

The device is typically turned on 10 to 14 days after implantation to allow adequate wound healing. Be aware that patients seen postoperatively during the first 2 weeks after implantation may not have had their device activated yet.

In addition to surgery-related risks and complications, if the patient complains of severe neck pain, worsening hoarseness, choking, or difficulty breathing, consider device-specific complications. Voice alteration or hoarseness is the most common device-specific adverse effect, with more than 50% of patients being affected.\(^29\)\(^-\)\(^31\) Other common side effects include increased coughing, shortness of breath, and pharyngitis.

Less common device-specific adverse effects reported during clinical studies include ataxia, dyspepsia, dysphagia, hypoesthesia, infection, insomnia, laryngismus, nausea, pain, paresthesia, and vomiting.\(^28\)

**Procedure**

In cases of emergency, diagnostic uncertainty, or significant adverse effects, turn off the pulse generator temporarily by holding a magnet over the generator (Box 70.4). Holding a magnet over the pulse generator for at least 65 seconds (5 seconds in the Model 106) will turn off any ongoing stimulation while the magnet is held in place. Once the magnet is removed, the normal operation will resume. Alternatively, manual on-demand stimulation may be triggered by passing the external VNS magnet over the generator for at least 1 second and then immediately removing it. This on-demand stimulation, if initiated at the onset of an aura or seizure, may abort the attack or, if initiated during a seizure, may halt its progression.\(^28\)

Unlike the typical donut-shaped magnet of a cardiac pacemaker, the VNS magnet is bar shaped, as shown in Fig. 70.5. Be aware that patients with VNSs are typically provided with two magnets: one fitted with a wristband and the other fitted with a belt clip.

**OTHER IMPLANTABLE DEVICES**

There are several additional noncardiac implantable devices that the emergency provider may encounter, which are outlined.
magnetic resonance imaging (MRI) is contraindicated, and indeed, manufacturers of gastric, urinary, diaphragmatic, and spinal cord stimulators list these devices as absolute contraindications to MRI. However, there are a number of clinical series and case reports indicating the compatibility of certain devices with MRI. For VNSs, the manufacturer recommends avoiding the use of any transmit coil for 3T or 1.5T imaging, as it may cause heat injury and damage the device. For deep brain stimulation, some studies have suggested that MRI is safe in certain circumstances, but reports of patient injury can be found in the literature. For IDDS devices, pumps have been shown to stop operating while in the magnetic field but resume functioning once removed. Both SynchroMed II and Prometra pumps are MRI compatible. However, the Prometra device is only MRI compatible after complete removal of medication from the reservoir, whereas there is no need to remove medication from the SynchroMed II device. Insulin pumps may easily be removed in the event that MRI is required.

In summary, the issue of MRI compatibility with implantable devices is very complex. In the event that MRI appears to be necessary, a case-by-case risk-benefit analysis involving the relevant subspecialist and radiologist is recommended.

**REFERENCES ARE AVAILABLE AT www.expertconsult.com**
REFERENCES

CHAPTER 71

Radiation in Pregnancy and Clinical Issues of Radiocontrast Agents

Phillip R. Peterson

The annual, population-wide exposure to ionizing radiation from medical sources increased a staggering 750% from 1980 to 2009, and medical imaging now comprises nearly one-half of Americans’ radiation exposure. The World Health Organization classifies x-rays officially as carcinogens, and indeed a report by the National Academy of Science indicates that a 100 mGy dose is associated with a lifetime attributable risk for development of a solid cancer or leukemia of 0.1%. Efforts were undertaken to reduce unnecessary exposure, and in 2011 the Joint Commission on Accreditation of Healthcare Organizations published a report that suggests risks can be reduced by the following: (1) raising awareness among staff and patients of the increased risk associated with cumulative doses; (2) providing the right test and the right dose (as low as reasonably achievable [ALARA]) through efficient processes and safe technology; and (3) promoting a culture of safety to make sure that doses are as low as possible while achieving the purpose of the study.

Such risks associated with exposure to ionizing radiation during medical care are sources of anxiety and concern in the pregnant patient, based on questions submitted to Health Physics Society’s “Ask the Expert” website. Furthermore, clinicians add a great deal of confusion and fear by providing exposed women with potential erroneous or inadequate information. Clinicians, nurses, and even radiologists may be ignorant of the effects of ionizing and nonionizing radiation. This misinformation contributes not only to undue and unnecessary anxiety and confusion but may lead to inappropriate abortions and litigation. For example, 23% of pregnancies in Greece were terminated after the Chernobyl incident because of unsubstantiated fears. A better understanding of the risks and effects enables providers to make informed decisions and counsel patients that appropriate radiology examinations provide more benefit than harm.

According to the American College of Obstetricians and Gynecologists, “With few exceptions, radiation exposure during medical care is sources of anxiety and concern in the pregnant patient, based on questions submitted to Health Physics Society’s “Ask the Expert” website. Furthermore, clinicians add a great deal of confusion and fear by providing exposed women with potential erroneous or inadequate information. Clinicians, nurses, and even radiologists may be ignorant of the effects of ionizing and nonionizing radiation. This misinformation contributes not only to undue and unnecessary anxiety and confusion but may lead to inappropriate abortions and litigation. For example, 23% of pregnancies in Greece were terminated after the Chernobyl incident because of unsubstantiated fears. A better understanding of the risks and effects enables providers to make informed decisions and counsel patients that appropriate radiology examinations provide more benefit than harm.

When evaluating the pregnant patient who has been or may be exposed to ionizing radiation and attempting to estimate risk, consideration must be given to a number of factors. These include not only factors related to gestational age, dose and type of radiation, and the studies performed or to be performed, but also to the mother herself, as studies raise concern about exposure of maternal breast tissue to ionizing radiation and the future risk of breast cancer. This chapter provides a review of ionizing and nonionizing radiation, its uses in diagnostic radiological examinations, the risks of its use, and strategies for minimizing risk to assist the clinician in counseling the patient to arrive at the most appropriate shared decision.

TYPES OF RADIATION

Simply stated, radiation is the emission or transmission of energy, which can take the form of electromagnetic radiation, such as visible light and x- and gamma-rays (the difference lying in the origin of the photon: electron orbital and nucleus, respectively); particle radiation, such as alpha particles (helium nuclei), beta particles (electrons, positrons), and neutrons; and acoustic radiation. As radiation propagates through space, it can impart or give-up some or all of its energy to surrounding material or tissues, and detection of this energy or its absence forms the basis of diagnostic medical imaging. Radiation is further classified by its effects on the absorbing material. Non-ionizing radiation lacks sufficient energy to liberate electrons, and its deposited energy subsequently manifests as heat, whereas ionizing radiation is of sufficient energy to liberate electrons, thereby breaking chemical bonds, forming highly reactive free radicals, and damaging DNA and cellular machinery.

RADIOBIOLOGY AND RADIOPHYSICS

The detrimental effects of radiation on health are grouped broadly into tissue reactions (also called deterministic effects, nonstochastic effects) and stochastic effects (Table 71.1). Tissue reactions are the result of cell killing and manifest as lethality, central nervous system (CNS) abnormalities, cataracts, growth retardation, malformations, and behavioral disorders. Tissue effects are dependent on the energy imparted to the tissue and demonstrate a threshold below which they are not observed but above which they are reliably observed, with their severity rising with dose. Stochastic effects include cancer and potential hereditary effects. Unlike tissue reactions, they have no identifiable threshold. Their probability of occurrence rises with dose, and their severity is independent of the dose.

The quantity of energy that is imparted by radiation to the material or tissue is a physical quantity, can objectively be measured, and is called the absorbed dose (D). It is defined as the mean energy imparted to matter per unit mass, and the mean absorbed dose (D) in an organ or tissue is the average absorbed dose over that organ or tissue. The Systeme International (SI) unit for both absorbed dose and mean absorbed dose is the joule per kilogram (J/kg) with the special name grey (Gy). In the centimeter-gram-second (CGS) variant of the metric system, the unit is given the special name rad: 1 rad = 100 erg/g = 0.01 J/kg = 0.01 Gy = 10 mGy. This unit is deprecated but remains in use in some countries (notably the United States).
TABLE 71.1 Stochastic and Tissue Reaction (Threshold) Comparison

<table>
<thead>
<tr>
<th>PHENOMENON</th>
<th>PATHOLOGY</th>
<th>DISEASES</th>
<th>RISK</th>
<th>DEFINITION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stochastic</td>
<td>Damage to a single cell may result in disease</td>
<td>Cancer, germ cell mutation</td>
<td>Some risk exists at all dosages; at low doses the risk may be less than the spontaneous risk</td>
<td>The incidence of the disease increases with dose, but the severity and nature of the disease remain the same</td>
</tr>
<tr>
<td>Tissue Reactions</td>
<td>Multicellular injury</td>
<td>Intrauterine death, organ malformations, mental impairment, growth retardation</td>
<td>No increased risk below the threshold dose</td>
<td>Both the severity and the incidence of the disease increase with the dose</td>
</tr>
</tbody>
</table>

Modified from Brent RL: Utilization of developmental basic science principles: the evaluation of reproductive risks from pre- and postconception environmental radiation exposures, Teratology 54:182, 1999.

The many forms of radiation differ in their abilities to impart energy to the absorbing material and cause ionizing events. Neutrons, alpha particles, and other heavy charged particles (e.g., carbon ions) impart their energy more readily, thereby producing more densely spaced ionizing events, whereas the converse is true for electromagnetic radiation and light charged particles (e.g., beta particles). Another quantity, the equivalent dose (HÉ), is therefore necessary and derived from the mean absorbed dose by scaling with a radiation weighting factor (wR), \( HÉ = wRDÉ \) and the effective dose for an organism is the tissue-weighted sum of the equivalent doses for all tissues, radiation sources, and energies. The radiation weighting factor, previously called the quality factor, is dependent only on the type of radiation (and for neutrons, its energy) and is dimensionless. Per the International Committee on Radiological Protection, \( wP = 1 \) for all photons and electrons; \( wP = 2 \) for protons and charged pions; \( wP = 20 \) for alpha particles, heavy ions, and fission fragments; and \( wP \) is a continuous function of energy for neutrons. Equivalent dose and effective dose have essentially the same magnitude as the absorbed dose. As an example, a mean absorbed dose of 100 mGy results in an effective dose of 100 mSv if the source is electromagnetic radiation or 2 Sv if the source is alpha radiation.

Radioactive decay or radioactivity is the process by which the nucleus of an unstable atom emits energy in the form of radiation. At the quantum level, the process is stochastic, but at the macroscopic level, the decay of a sufficiently large collection of unstable atoms can be calculated and is represented by the half-life, or the time it takes for the collection of unstable atoms to reduce by half. The SI unit of radioactivity is the becquerel (Bq) and is defined as one nuclear decay event per second. Its units are therefore inverse seconds (1 Bq = 1 s⁻¹), as a nuclear decay event is dimensionless. A frequently encountered, non-SI unit of radioactivity is the curie (Ci). Its definition is based on the amount of radioactivity in one gram of radium and also has units of events per second; 1 Ci = 3.7 × 10¹⁰ Bq = 37 GBq. Doses of radioactive substances used in medicine are typically on the MBq and mCi scale (e.g., 1 mCi = 37 MBq). These units encode neither the type nor energy of the radiation emitted by the nuclear decay process.

**EFFECTS OF RADIATION**

Exposure to radiation is among many recognized agents and environmental exposures which can produce deleterious effects, resulting in mental retardation, neurobehavioral effects, convulsive disorders, congenital malformations, fetal growth retardation, embryonic death, and cancer.⁵⁶ The significant background prevalences of these events are 0.5% to 1% for mental retardation, 3% for major congenital malformation, 3% for growth retardation,¹ ¹⁵% for miscarriage (with a reported variation as wide as 30% to 50%),⁵ ¹¹% for genetic disease,⁵ and 7% for prematurity,⁵ and in the United States there is a 21% background lifetime risk of fatal cancer in the offspring.⁵

As previously discussed, the effects of radiation are broadly divided into two categories (see Table 71.1): stochastic effects (which include cancer and hereditary disease) have no dose threshold, and the frequencies of which rise with increasing dose; and tissue reactions (which include the remaining effects) have identifiable thresholds above which they are reliably observed, the severities of which do not depend on dose, and whose manifestations depend on the stage of fetal development in which the exposure occurs.⁶,¹² Dependence on the duration of time over which the exposure occurs is also noted,¹ as a protracted exposure is less likely to produce an effect than a single exposure of high intensity (e.g., the dose from continuous exposure during flight).³,¹² Fractionation of the dose (e.g., multiple radiographs over a period of hours or days), like protraction, also decreases the severity of and increases the threshold for tissue reactions.⁵ The dependence of the effects of radiation on the stage of development is demonstrated in Fig. 71.1.¹¹

**Stages of Fetal Development**

Development of the fetus is expressed as postconception age and divided into three major stages: (1) the preimplantation and implantation phase (conception to implantation), (2) organogenesis (3RD to 8th week), and (3) fetal development (9th week until birth), which includes the majority of CNS development (8th to 25th week). The risks of radiation are most significant during organogenesis and the early fetal period, somewhat less in the second trimester, and least in the third trimester.¹⁷ This relationship is demonstrated in Fig. 71.1, and the main effects and their thresholds are summarized in Table 71.2.
**Preimplantation and Implantation Phase**

The preimplantation and implantation phase has been designated an “all or none” period, as embryonic cells are few in number and totipotential, and can replace damaged neighboring cells. It is in this stage of development that the embryo is least likely to be malformed, and the predominant effect of radiation exposure is failure to implant, resulting in abortion. The threshold for mortality in this stage of development is quite high, estimated at 100 to 200 mGy and rises with fetal age such that at term the threshold for mortality is 1 Gy to 1.5 Gy (see Table 71.2). Embryos that survive exposure in this period of development demonstrate no increased risk for malformations or growth retardation.

**Organogenesis**

Exposure to high doses of radiation during the period of organogenesis is likely to affect the organ system or systems under development at the time of the exposure and results from reduced ability to replace those cells damaged or killed by the exposure. Although any organ system can be affected, the predominantly observed effects are microcephaly and growth retardation, and have a dose threshold of 50 to 200 mGy, which is significantly higher than doses obtained from diagnostic radiology examinations. Temporary growth retardation can be observed with doses of 100 to 250 mGy, but these infants recover fully and attain normal adult stature and weight. The dose threshold for permanent growth retardation depends on postconception age and is 200 to 500 mGy in the 3rd to 5th weeks, 250 to 500 mGy in the 6th to 13th weeks, and greater than 500 mGy from the 14th week onward (see Table 71.2). Exposure to less than 50 mGy during the period of organogenesis results in no change in the risk for organ malformations or growth retardation.

**Fetal Period**

At the end of the period of organogenesis, the fetus has diminished vulnerability to multiple organ system teratogenesis, but beginning in the 8th week and continuing through the 25th week is development of the CNS, during which it can be seriously affected by high doses of ionizing radiation, resulting in mental impairment, mental retardation, neurobehavioral effects, and convulsive disorder. CNS sensitivity to ionizing radiation is not uniform throughout its development, with highest sensitivity from the 8th through 15th weeks, during which there is a rapid increase in the number of neurons that migrate and differentiate, losing their ability to divide. This process is largely concluded by the 15th week, at which point and lasting until the 25th week, further differentiation

---

**TABLE 71.2** Risks and Threshold Doses of the Main Effects of Prenatal Irradiation

<table>
<thead>
<tr>
<th>HUMAN GESTATIONAL AGE [POSTCONCEPTION AGE]</th>
<th>MINIMUM LETHAL DOSE TO EMBRYO OR FETUS (Gy)</th>
<th>APPROXIMATE LD50</th>
<th>MINIMUM DOSE FOR PERMANENT GROWTH RETARDATION IN THE ADULT (Gy)</th>
<th>MINIMUM DOSE FOR GROSS ANATOMIC MALFORMATIONS (Gy)</th>
<th>INCREASED INCIDENCE OF MENTAL RETARDATION (Gy)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3rd to 4th [1st to 2nd pc]</td>
<td>0.15 to 0.2</td>
<td>&lt; 1</td>
<td>No increased incidence of GR in survivors</td>
<td>No increased incidence of congenital malformation in survivors</td>
<td></td>
</tr>
<tr>
<td>5th to 7th [3rd to 5th pc]</td>
<td>0.25 to 0.5</td>
<td>1.4–2</td>
<td>0.2–0.5</td>
<td>&gt; 0.2 but most malformations require &gt; 0.5</td>
<td></td>
</tr>
<tr>
<td>8th to 15th [6th to 13th pc]</td>
<td>&gt; 1</td>
<td>&gt; 2</td>
<td>0.25–0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>16th to term [14th pc to term]</td>
<td>&gt; 1.5</td>
<td>Same as for the mother</td>
<td>≥ 0.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10th to 27th [8th to 25th pc]</td>
<td></td>
<td></td>
<td></td>
<td>SMR at doses &gt; 0.5; (lower 95% CI value of ~ 0.3)</td>
<td></td>
</tr>
</tbody>
</table>

CI, Confidence interval; GR, growth retardation; pc, postconception; SMR, severe mental retardation.

and architectural definition occur, and the CNS becomes slightly less susceptible to damage.

Mental Impairment and Retardation
It is important to note that all clinical observations of significant IQ reduction and severe mental retardation (SMR) are associated with high doses of radiation in excess of 500 mGy, a level not easily approached with diagnostic radiology examinations. The high risk for SMR exists in the 8th through 15th weeks and with a high dose threshold of 1 Gy, and the severity of the retardation does appear to have some dependence on the dose. For survivors of the atomic bombings of Japan who were between 8 and 15 weeks after conception at the time of their exposure, a drop of 0.3 IQ points per 10 mGy of exposure, with a threshold of 500 mGy, was observed. There is no documented risk for mental retardation in humans at gestational ages less than 8 weeks or greater than 25 weeks with exposure less than 500 mGy. Furthermore, the risk of exposure to 100 mGy or less is far smaller than the 3% background risk of a child being born with an IQ less than 70, and exposure to less than 100 mGy cannot be clinically identified with reduction in IQ.

Convulsive Disorders
Convulsive disorders are a broad category encompassing various diagnoses, etiologies, and manifestations. The risk for convulsive disorder following in utero radiation exposure was evaluated in several studies on Japanese survivors of the atomic bombing. Among these survivors, no convulsive disorders were identified in those receiving a dose less than 100 mGy. Those at highest risk for seizure disorder were those exposed in the 8th through 15th weeks postconception and who were also mentally retarded. The association of convulsive disorders with radiation exposure between the 8th and 15th weeks postconception without mental retardation was borderline significant. There is no increase in convulsive disorders from exposure before the 8th week or after the 25th week.

Growth Retardation
Growth retardation can occur with exposure to ionizing radiation in the fetal period, though exposure is more likely to result in microcephaly. Studies on the atomic bombing survivors who were exposed in utero show divergent thresholds for microcephaly. For survivors of Nagasaki, the threshold is estimated at 1.5 Gy, whereas for the survivors of Hiroshima, a threshold of 100 to 190 mGy is reported. This latter finding is consistent with a threshold of 100 to 200 mGy for microcephaly that was observed in mammalian animal models.

Irradiation of the human fetus at doses less than 100 mGy has not been observed to cause congenital malformations or growth retardation (see Table 71.2).

Stochastic Risks
The remaining effects (cancer and hereditary effects) are stochastic, notably in that risk exists at all levels of exposure (see Table 71.1). Childhood malignancies account for less than 1% of total cancer diagnoses in the United States, and the cumulative risk to 15 years of age is 1 to 2.5 per 1000 in most western countries. Additionally, lymphoma, leukemia, and brain tumors account for 70% of childhood cancers, which stands in contrast with adult malignancies, the majority of which are epithelial in origin.

Cancer
The Oxford Survey was published 60 years ago and reported a twofold increase in risk in total childhood cancer mortality in the offspring of mothers undergoing abdominal diagnostic radiographs. These results were met initially with skepticism and criticized on the basis of recall bias, but the effect persisted in later studies in which a 1.3- to 1.5-fold risk was reported. Aside from the Oxford Survey, which reported a relative risk of 1.5 for all solid tumors and 1.4 for CNS tumors, all other studies on CNS tumors did not find a statistically significant increase. These findings are consistent with an analysis of later epidemiologic studies that found a relative risk of 1.4 for childhood cancer after exposure to 10 mGy. Efforts were undertaken to estimate the risk per gray from exposure and to determine the excess absolute risk coefficient of ~ 6% per Gy (2.5% per Gy for leukemia). These results must be interpreted in the context of the background incidence; for a fetus exposed to 1 mGy, the risk for carcinogenesis is 0.006%, or 3 additional diagnoses for every 50,000 exposures, as compared with the background incidence of 0.2% to 0.3%, or 100 to 150 per 50,000. Exposure to 50 mGy, the generally accepted upper limit for exposure in utero, may double the risk for childhood malignancy in the offspring, but this absolute risk remains low (0.5% to 0.6%).

Hereditary Effects
Early research with flies and animals demonstrated ionizing radiation can induce germ-cell mutations, but these findings have failed to be demonstrated in humans and were so small in the irradiated Japanese survivors as to be statistically insignificant. Others report an excess absolute risk for hereditary disease of 1% per Gy. In the context of a background incidence of 3%, an exposure of 50 mGy might then be expected to raise this risk to 3.05%.

Radiography and Computed Tomography
Table 71.3 lists the absorbed dose of the fetus or embryo and the gonads from ionizing radiation of various studies. The vast majority of diagnostic imaging studies do not result in significant fetal exposure. Computed tomography studies (CT scans) of the maternal abdomen and pelvis in which the fetus or embryo lies directly in the field of study can deliver doses approaching the safe threshold of 50 mGy, and CT of the pelvis may have the potential to exceed this threshold. Alternative diagnostic imaging algorithms have therefore been developed to assess the pregnant patient in clinical scenarios where CT might otherwise have been indicated.

Whereas fetal position in relationship to the field of study is a major factor determining fetal exposure from CT scans, variables such as slice thickness, number of cuts, and helical movement pattern (i.e., spiral CT) also affect the amount of ionizing radiation used. In helical or spiral CTs, the study is completed in less time, resulting in less radiation exposure, and the use of 16-slice over 4-slice equipment can also lead to a reduction in the dose by 20% to 30%. The principal concern in obtaining a CT scan in the pregnant patient is exposing the fetus or embryo to ionizing radiation. Situations may also arise in which it is necessary to use iodinated contrast material in the study, and the potential harmful effects of the contrast must also be considered. The most important potential effect of iodinated contrast material is depression of fetal thyroid function. By 12 weeks, the fetus is producing...
thyroid function has not been observed with high doses of nonionic contrast. However, neonates are routinely screened for hypothyroidism in the United States. Lactating and nursing mothers may also inquire about interrupting nursing after receiving intravenous (IV) contrast. Very little of the contrast reaches the milk, and only a tiny fraction of that is absorbed in the gut. Interruption of nursing is therefore unnecessary.

Table 71.3 Estimated Fetal and Gonadal Exposure From Various Diagnostic Imaging Methods and Radiopharmaceuticals

<table>
<thead>
<tr>
<th>TYPE OF PROCEDURE</th>
<th>DESCRIPTION</th>
<th>TYPICAL DOSE TO EMBRYO OR FETUS (mGy)</th>
<th>DOSE TO GONADS (OVARIES, TESTES) (mGy)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Radiography and Fluoroscopy</td>
<td>Skull</td>
<td>&lt; 0.01</td>
<td>&lt; 0.01, &lt; 0.01</td>
</tr>
<tr>
<td></td>
<td>Chest</td>
<td>&lt; 0.01</td>
<td>&lt; 0.01, &lt; 0.01</td>
</tr>
<tr>
<td></td>
<td>Thoracic spine</td>
<td>&lt; 0.01</td>
<td>&lt; 0.01, &lt; 0.01</td>
</tr>
<tr>
<td></td>
<td>Mammography</td>
<td>&lt; 0.1</td>
<td>&lt; 0.01, NA</td>
</tr>
<tr>
<td></td>
<td>Pelvis</td>
<td>0.1–1.1</td>
<td>2, 4</td>
</tr>
<tr>
<td></td>
<td>Lumbar spine</td>
<td>1–2</td>
<td>4, 0.6</td>
</tr>
<tr>
<td></td>
<td>Abdomen</td>
<td>1–3</td>
<td>2.5, 0.7</td>
</tr>
<tr>
<td></td>
<td>Upper GI series</td>
<td>7</td>
<td>1, 0.1</td>
</tr>
<tr>
<td></td>
<td>Barium enema</td>
<td>7–8</td>
<td>10, 4</td>
</tr>
<tr>
<td>CT</td>
<td>Brain</td>
<td>&lt; 0.1</td>
<td>&lt; 0.1</td>
</tr>
<tr>
<td></td>
<td>Chest, angiography of coronary arteries</td>
<td>0.1–1</td>
<td>&lt; 0.1, &lt; 0.1</td>
</tr>
<tr>
<td></td>
<td>Abdomen</td>
<td>4–16</td>
<td>1, 1</td>
</tr>
<tr>
<td></td>
<td>Pelvis</td>
<td>(avg: 8; max: 49)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lumbar spine</td>
<td>Avg: 25; max: 79</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Avg: 2.4; max: 8.6</td>
<td></td>
</tr>
<tr>
<td>Nuclear Medicine</td>
<td>Lung-ventilation study; $^{133}$Xe (740 MBq)</td>
<td>0.02–0.3</td>
<td>0.5, 0.5</td>
</tr>
<tr>
<td></td>
<td>Lung-perfusion study; $^{99m}$Tc-MAA (148 MBq)</td>
<td>0.4–0.7</td>
<td>0.27, 0.16</td>
</tr>
<tr>
<td></td>
<td>White cell scan; $^{111}$In (18.5 MBq)</td>
<td>1.7–2.4</td>
<td>2.2, 0.8</td>
</tr>
<tr>
<td></td>
<td>Renal scan; $^{99m}$Tc-MAG3 (370 MBq)</td>
<td>1.9–6.7</td>
<td>2, 1.4</td>
</tr>
<tr>
<td></td>
<td>Bone scan; $^{99m}$Tc-MDP (740 MBq)</td>
<td>1.5–4.4</td>
<td>2.7, 1.8</td>
</tr>
<tr>
<td></td>
<td>Cerebral blood flow; $^{99m}$Tc-HMPAO (740 MBq)</td>
<td>3–6.7</td>
<td>4.9, 1.9</td>
</tr>
<tr>
<td></td>
<td>Positron-emission tomography; $^{18}$F-FDG (740 MBq)</td>
<td>13–16</td>
<td>10, 8.1</td>
</tr>
<tr>
<td></td>
<td>Myocardial perfusion; $^{99m}$Tc-sestamibi (1110 MBq)</td>
<td>5–6</td>
<td>9.0, 4.1</td>
</tr>
<tr>
<td></td>
<td>Cancer therapy – 5% uptake; $^{131}$I (3,700 MBq)</td>
<td>66–1000</td>
<td>163, 107</td>
</tr>
<tr>
<td></td>
<td>Benign therapy – 50% uptake; $^{131}$I (555 MBq)</td>
<td>40–150</td>
<td>23, 14</td>
</tr>
</tbody>
</table>


CT, Computed tomography; FDG, fluodeoxyglucose; GI, gastrointestinal; HMPAO, hexamethylpropyleneamine oxime; MAA, macroaggregated human albumin; MAG3, mercaptoacetyltriglycine; MDP, methylene diphosphonate; NA, not applicable.
NUCLEAR MEDICINE STUDIES

Unlike radiographs and CT, which rely on an external source of radiation that is passed through the body before detection, nuclear medicine detects radiation from unstable isotopes that are introduced into the body within which they emit their radiation. Furthermore, whereas the other ionizing modalities are primarily concerned with imaging specific anatomy, nuclear medicine studies are concerned with the assessment of the physiologic (or pathophysiologic) function of a tissue or organ system. Like studies involving external sources of ionizing radiation, nuclear medicine studies can extract information in two dimensions, such as in scintigraphy, or in three dimensions, such as single photon emission CT (SPECT) and positron emission tomography (PET).

The radioactive isotopes are bound to substrates or tracers that are processed differently in pathologic states. The detection of the altered distribution in time and space of radiation from these radiotagged substrates (radiopharmaceuticals) forms the basis for the nuclear medicine examination. The metastable nuclear isomer of technetium-99 (99mTc) is frequently encountered in diagnostic nuclear medicine studies and emits radiation in the form of gamma-rays.10 Many different isotopes, however, are employed in nuclear medicine studies (see Table 71.3) and the absorbed dose to the fetus or embryo is dependent on the activity and amount of the isotope and the specific metabolism of the substrate to which it is bound.

Some radiopharmaceuticals, such as 131I and 123I, cross the placenta and are generally not used in pregnant patients.12 For the remainder, the dose to the fetus is derived from the activity in maternal tissues, the proximity of these tissues to the fetus, and the excretion of the radiopharmaceutical. For example, the ventilation portion of a ventilation-perfusion (V/Q) study for pulmonary embolism (PE) can be done with either 133Xe or 99mTc-DTPA. Use of 131I Xe results in little fetal exposure, as its activity is confined to the maternal lungs, remote from the fetus or embryo, but 99mTc-DTPA is excreted by the kidneys and can collect in the urinary bladder, where it and other renally excreted radiopharmaceuticals can expose the fetus or embryo.12 Strategies for reducing fetal exposure therefore include maternal hydration and frequent voiding, considering the use of smaller administered doses and longer imaging times, selecting alternative radiopharmaceuticals (e.g., 133Xe vs. 99mTc-DTPA), and adjusting the examination protocol (e.g., obtaining the perfusion portion of a V/Q study first which, if normal, precludes obtaining the ventilation portion).

Nonpregnant women may inquire about delaying pregnancy after undergoing a nuclear medicine examination. It is advised that pregnancy be delayed until the dose from the remaining radiopharmaceutical is less than 1 mGy, which is generally not a consideration except for 59Fe (metabolism studies) and 75Se (adenal studies), for which delays of 6 and 12 months, respectively, are advised.12 Lactating women must also be counseled to cease breastfeeding for at least 12 hours for 99mTc, except for labeled red blood cells, phosphates, and DTPA, for which a period of at least 4 hours is advised. For other less frequently encountered isotopes in the emergency department, it is advised that breastfeeding be ceased for a period of 3 weeks for most 111In and 123I compounds, except hippurate (12 hours) and for 22Na, 67Ga, and 201Tl.

The clinician may also encounter a patient undergoing a nuclear medicine examination who is also the household contact of a pregnant woman. These individuals pose no risk to the pregnant woman, the fetus, or any other household contact, as the total decay dose measure at 0.5 m and 1 m is 0.02 to 0.25 mGy and 0.05 to 0.1 mGy, respectively.12

DIAGNOSIS OF PE

In the developed world, PE is the leading cause of maternal death, accounting for some 20% of maternal deaths in the United States,13 and for every 2000 pregnancies there is one PE.14 Mortality for untreated, acute PE is 30% compared with 2% to 10% in those diagnosed and treated in a timely fashion.12,15

Both CT of the pulmonary arteries (CTPA) and V/Q nuclear medicine studies can be used to assess for PE but employ ionizing radiation in different manners, leading to different risks and benefits, though exposure to the fetus in both studies is minimal.26 Fetal or embryological exposure from a CT chest scan varies with gestational age and is 0.013 to 0.026 mGy in early pregnancy, rising to 0.01 to 0.1 mGy in late pregnancy.14 Fetal exposure is also dependent on the equipment and is less for multidetectors than for single detectors.9,12 A combined 185 MBq 99mTc perfusion and 370 MBq 133Xe ventilation study equates to 2.25 mGy fetal exposure,13 which is greater than that for a CTPA26 but remains a small exposure. The dose in a V/Q study can be lowered, resulting in a suitable study, but one in which the fetal exposure from the 99mTc perfusion portion is 0.1 to 0.6 mGy in early pregnancy and 0.6 to 0.8 mGy in later pregnancy and from the 99mTc ventilation portion is 0.1 to 0.3 mGy for a total exposure less than 1.1 mGy12 (Box 71.1). And whereas pulmonary angiography was long considered the gold standard in the diagnosis of PE, it is now thought to be no more accurate than CTPA and results in no less fetal exposure: 2.2 to 3.7 mGy when done via the femoral route or 0.5 mGy when performed by the brachial route.15

The two modalities also differ, not only in the radiation exposure to the fetus or embryo, but also in the exposure to the mother, in that CTPA exposes a woman’s breasts to a higher level of radiation and elevates her risk for cancer.12 It is estimated that every 10 mGy of exposure to the breasts of a woman aged 35 years or more increases her risk by 14% over the background.67 By this exposure and risk to the mother alone, some authors justify the use of V/Q studies over CTPA as the primary examination in a pregnant patient.18

The American Thoracic Society therefore recommends, and the American College of Obstetricians and Gynecologists has endorsed, an algorithm in which pregnant women suspected of PE are assessed first for clinical evidence of deep venous thrombosis (DVT). If DVT is suspected, an ultrasound (US) of the bilateral lower extremities is obtained, followed by treatment if positive. If the US is negative, or the patient has no evidence of DVT, a chest x-ray (CXR) is performed. Patients with normal CXRs undergo V/Q studies in which the dose is reduced and the perfusion portion is performed first, followed by the ventilation portion if necessary. Patients with nondiagnostic V/Q studies or abnormal CXRs undergo CTPA. This algorithm is depicted in Fig. 71.2. The clinician may also consider obtaining a bedside echocardiogram initially in the unstable or critically ill patient, and it is worthwhile to note that strategies exist to reduce the exposure from CTPA19 (Box 71.2).

The utility of a D-dimer level in excluding PE in pregnant patients remains under investigation.16,17 D-dimer levels are
**BOX 71.1 The Technique of V/Q Scanning**

This test uses both intravenous (perfusion) and aerosolized (ventilation) agents.

**PERFUSION**

1. Before injection, prepare the intravenous technetium. Mix sodium pertechnetate $^{99m}$Tc with macroaggregated human albumin (MAA) to form $^{99m}$Tc-MAA, the substance that is injected intravenously to investigate blood flow in the lungs. If the preparation is not used within 8 hours, discard it.
2. The usual dose is 37–111 MBq (1–3 mCi). Doses as low as 37 MBq are used in pregnancy.
3. Within 5 minutes of injection, more than 90% of the Tc-albumin aggregate is trapped in the arterioles and capillaries of the lung. The particle size determines where the $^{99m}$Tc will be localized in the body.
4. Accumulation in the lung is temporary, and the fragile albumin aggregate quickly breaks down, thereby allowing Tc to enter the general circulation.
5. Once in the body, the half-life of $^{99m}$Tc is 6 hours.
6. The majority of $^{99m}$Tc is excreted in urine. If it remains in the urinary bladder, it is in close proximity to the fetus.
7. Technetium in the bladder exposes the fetus to small amounts of radiation.
8. Frequent voiding or bladder catheterization after the study will lessen exposure of the fetus to radiation.
9. Technetium is relatively contraindicated in patients with severe pulmonary hypertension (because $^{99m}$Tc-MAA temporarily blocks blood flow in the lungs).
10. Allergic reactions to technetium and human serum albumin are extremely rare.
11. The radiation exposure to the total body from 92.5 MBq (2.5 mCi) is extremely low: less than 1 mGy.

**VENTILATION**

1. Some radiologists forgo the ventilation portion of the V/Q scan in pregnancy to limit the total radiation exposure.
2. Most hospitals use $^{133}$Xe for the ventilation portion of the V/Q scan.
3. Fetal radiation exposure from xenon is extremely low, and the 50 mGy threshold is not reached until more than 125 scans have been performed.
4. The estimated dose to the fetus of the standard 370 MBq (10 mCi) of xenon used in a V/Q scan is 0.4 mGy.
5. If Tc-based aerosol is the marker used for the ventilation portion, fetal exposure is higher than with xenon aerosol.


known to rise throughout pregnancy such that 79% of non-pregnant women attempting to conceive have a level lower than 0.5 mg/L, and 50% of women in the first trimester, 22% in the second trimester, and 0% in the third trimester have a level lower than 0.5 mg/L. Use of the D-dimer in excluding PE may be useful in low- and intermediate-risk individuals in the first trimester and becomes useful again 4 weeks after delivery, but further investigation is necessary to establish appropriate thresholds in the three trimesters and test them in the clinical setting.
CHAPTER 71  Radiation in Pregnancy and Clinical Issues of Radiocontrast Agents

A B

Figure 71.3  Magnetic resonance imaging (MRI) for appendicitis in pregnancy. Right lower quadrant pain presents a diagnostic dilemma in pregnant patients, and abdominal computed tomography is generally eschewed in favor of modalities that do not use ionizing radiation. Ultrasound may be utilized but the appendix is often difficult to visualize, especially in the third trimester. A, MRI (fast spin echo, fat saturated) of a patient who is 33 weeks pregnant. A dilated appendix (12 mm) and periappendiceal edema are noted adjacent to the right psoas muscle (arrow), indicative of acute appendicitis. B, A coronal section from the same study reveals the fetus in a frank breech presentation. MRI is thought to present no risk to the fetus at any stage of pregnancy.

DIAGNOSIS OF APPENDICITIS

Acute appendicitis is the most common nonobstetric emergency requiring surgery during pregnancy. Appendicitis is associated with premature labor, fetal morbidity and mortality, and an increased rate of perforation. Concern for appendicitis in a pregnant patient warrants early surgical consultation and discussion of the need and type of imaging. Patel and coauthors36 published an algorithm that uses US and magnetic resonance imaging (MRI) techniques before exposing patients to ionizing radiation. They recommended the use of graded-compression US followed by abdominal/pelvic US to search for other pathology if needed. If the US studies are negative, the authors recommend MRI of the abdomen and pelvis (Fig. 71.3). Notably, some authors have found that using a nonionic oral contrast agent (a mixture of ferumoxsil [category B] and barium sulfate) improved sensitivity and specificity in detecting appendicitis in pregnancy.42 The authors noted that US imaging of the appendix was more easily done during the first and early second trimester and that the left lateral decubitus position assisted in visualization of the appendix in third-trimester patients. Finally, if US and MRI are still inconclusive, CT of the abdomen and pelvis may be considered. Definitive surgical exploration should be discussed with a general surgeon before proceeding to ionizing radiation. If CT is required, the authors reported an estimate of approximately 1 cancer per 500 fetuses exposed to 30 mGy. These guidelines are consistent with the American College of Radiology’s appropriateness criteria that suggest US should be performed first, followed by MRI if inconclusive, and reserves CT as a last resort.

DIAGNOSIS OF PREGNANCY AND CONSENT

If exposure to less than 50 mGy does not measurably affect the exposed embryo, why should the clinician determine the pregnancy status of the patient? Brent6 reported sound reasoning for diagnosing pregnancy before a radiographic study. The principle of informed consent must remain paramount. It is beneficial and more sound ethically to have the patient informed...
of her pregnancy status before imaging. An informative discussion about the risk-benefit aspects of the test before the study conveys concern for the patient and fetus. Discussing the risk-benefit aspects of imaging after the study may be misconstrued as “backpedaling” and has the potential to upset the patient. Many lawsuits are stimulated by the factor of surprise. Frank discussion before imaging may prevent misguided litigation. More importantly, having patients both understand the problem (imaging in pregnancy) and take part in the management discussion can help them become more empowered and potentially reduce the anxiety associated with their condition and with their pregnancy.

Determination of pregnancy by the history and physical examination alone can be problematic. The menstrual history by itself may not be totally reliable in determining pregnancy. Amenorrhea and physical changes in the size and shape of the uterus may be consistent with pregnancy. A history of recent mensturaluation, use of an intrauterine device, tubal ligation, absence of coitus, or proper use of birth control pills can result in a suggestion of nonpregnant status more than 90% of the time, but these parameters are not 100% accurate. If the diagnosis of pregnancy is in the differential or imaging is ordered, or both, definitive determination of the patient’s pregnancy status should be strongly considered if the clinical scenario is reasonable. A menstrual history and other information should be obtained whenever possible, and a confirmatory urine pregnancy test should be considered. Urine pregnancy tests to detect early pregnancy are quite sensitive and reliable, and it is not necessary to routinely order a quantitative serum test. Theoretically, there will be a few days’ window between fertilization and implantation, a period when no method will confirm the presence or absence of early pregnancy.

A pregnant patient has the right to know the magnitude and type of risks that might result from in utero exposure to radiation. The Annals of the International Commission on Radiological Protection Publication 8412 summarized the need for informed consent as follows:

The need and degree of disclosure is usually measured by what a reasonable person believes is material to the mother’s decision to be exposed to radiation. The level and degree of disclosure should be related to the level of risk. For low-dose procedures, such as chest x-rays, <100 mrad, the only information that may be needed is a verbal assurance that the risk is low. If the diagnosis of pregnancy is in the differential or imaging is ordered, or both, definitive determination of the patient’s pregnancy status should be strongly considered if the clinical scenario is reasonable. A menstrual history and other information should be obtained whenever possible, and a confirmatory urine pregnancy test should be considered. Urine pregnancy tests to detect early pregnancy are quite sensitive and reliable, and it is not necessary to routinely order a quantitative serum test. Theoretically, there will be a few days’ window between fertilization and implantation, a period when no method will confirm the presence or absence of early pregnancy.

A pregnant patient has the right to know the magnitude and type of risks that might result from in utero exposure to radiation. The Annals of the International Commission on Radiological Protection Publication 8412 summarized the need for informed consent as follows:

The need and degree of disclosure is usually measured by what a reasonable person believes is material to the mother’s decision to be exposed to radiation. The level and degree of disclosure should be related to the level of risk. For low-dose procedures, such as chest x-rays, <100 mrad, the only information that may be needed is a verbal assurance that the risk is low. If the diagnosis of pregnancy is in the differential or imaging is ordered, or both, definitive determination of the patient’s pregnancy status should be strongly considered if the clinical scenario is reasonable. A menstrual history and other information should be obtained whenever possible, and a confirmatory urine pregnancy test should be considered. Urine pregnancy tests to detect early pregnancy are quite sensitive and reliable, and it is not necessary to routinely order a quantitative serum test. Theoretically, there will be a few days’ window between fertilization and implantation, a period when no method will confirm the presence or absence of early pregnancy.

**PATIENT COUNSELING**

When a pregnant patient requires an imaging study, be prepared to discuss the risk associated with the test. Counseling can be done after attempting to estimate the dose received by the patient. It is important to use terminology that is easily understood by the patient. Fig. 71.4 depicts three different strategies to inform the patient about the level of exposure from her study and established limits.
Table 71.4 compares the level of exposure with established background risks.

The main bar graph (A) in Fig. 71.4 compares the fetal exposure level for various radiographic studies with the maximum accepted fetal dose during pregnancy (50 mGy). A patient's particular study may be plotted on this graph to show the clear margin of safety that exists for all single diagnostic tests.

The middle graph (B) equates the exposure from low-level diagnostic studies to the number of hours needed to accumulate a similar exposure dose from background terrestrial radiation. One of the most commonly ordered studies in pregnancy is a chest radiograph. The potential risk to the fetus can be put into perspective for the patient by comparing the absorbed dose for the chest radiograph with the natural background radiation exposure. The environmental background radiation over a 9-month period results in a cumulative dose of 1 mGy.43 The fetal dose exposure for a chest radiograph (two views) is estimated to be less than 0.1 mGy. Therefore the exposure dose to the fetus from a chest radiograph is equivalent to the same amount of naturally occurring background radiation to which the patient was exposed in the previous 2.7 days.

The lower graph (C) depicts the upper limit of the Nuclear Regulatory Commission (NRC) for cumulative gestational dose versus various diagnostic studies. The NRC has established occupational radiation dose limits for pregnancy. Its recommendation is that the dose to the fetus not be allowed to exceed 5 mSv during gestation. Brent noted that this factor-of-10 lowering of the widely accepted threshold is "extremely conservative." One can explain to the patient that the level of exposure from her radiograph is below the conservative cumulative acceptable dose for a pregnant employee at a nuclear facility in the United States.

Another useful approach is to indicate to the patient the probability of not having a child with either a malformation or cancer and how that probability is affected by radiation. Table 71.4 depicts the probability of bearing healthy children as a function of radiation dose. This discussion should be coupled with the fact that a nonexposed fetus has a baseline incidence of spontaneous abortion, multiple developmental abnormalities, and subsequent childhood cancer.

Numerous organizations have declared fetal exposure to less than 50 mGy as being safe. Box 71.3 presents various conclusions from key organizations on the use of radiation and pregnancy. The International Commission on Radiological Protection concluded that fetal doses below 100 mGy should not be considered a reason for terminating a pregnancy.12 If a patient still has considerable concern or has possibly received greater than 50 mGy, referral to a radiation physicist or genetic specialist for further counseling is reasonable.

### NONIONIZING RADIATION

Despite the United States Food and Drug Administration’s (FDA) caution that “the safety of magnetic resonance examinations has not been completely established for embryos and fetuses,” MRI and US are often the preferred imaging modalities in the pregnant patient, provided the diagnostic information can be obtained satisfactorily, as they do not involve the use of ionizing radiation.
Magnetic Resonance Imaging

MRI functions by using a strong magnetic field to align the spins of the proton in a hydrogen atom. A nonionizing radio-frequency (RF) field then excites some of these protons to a higher energy level by realigning their spins. Several gradient magnets are then employed to alter the local magnetic field at the anatomic location of interest as the RF field is turned off, thereby allowing the excited protons to relax to their base spin-orientation state, releasing the energy absorbed from the, again, nonionizing RF pulse. This energy is subsequently detected and transformed by a computer into anatomic images. The fetus is therefore exposed to both magnetic and electromagnetic fields from three sources: (1) the RF field with frequencies in the 10s to 100s of MHz, (2) time-dependent magnetic field gradients with changes on the order of 10 T/s, and (3) the static magnetic field (i.e., “field strength”) on the order of 1.5 to 3 T.

As MRI employs the use of a nonionizing RF field, the primary concern for effects of the RF field arises from tissue heating, as the dissipated energy manifests as heat. As with any thermal injury, the greater the amount of energy dissipated as heat over the shorter interval of time, the greater the rise in temperature and the greater the potential for injury. The amount of energy deposited by the RF field is called the average specific absorption rate (SAR), which depends on the specifics of the MRI protocol in use and is measured in energy per unit time (i.e., power) per unit mass, J/s/kg per kg = W/kg. SAR is calculated using the patient’s entire mass rather than that of a specific tissue. The FDA limits SAR to no more than 4 W/kg for the whole body averaged over 15 or more minutes and 3.2 W/kg for the head averaged over 10 or more minutes. These limits are the same for fetuses or embryos and for mothers. The International Electrotechnical Commission (IEC) has established three levels of operation which are programmed into the MRI: (1) normal, which permits whole-body SARs up to 2 W/kg, (2) first-level controlled, which permits whole-body SARs up to 4 W/kg and requires approval of the radiologist, and (3) second-level controlled in which whole-body SARs can exceed 4 W/kg but which requires signed patient consent and can be used only under an institutional review board–approved protocol. Whole-body SARs of 2 W/kg and 4 W/kg are not expected to raise core body temperature by more than 0.5°C and 1°C, respectively.

Implied in these estimates are the assumptions of an ideal study environment for the dissipation of heat and normal thermoregulatory mechanisms in the patient. The embryo or fetus, however, exists in a well-insulated environment with more limited heat-removal systems and maintains a temperature of approximately 0.5°C to 1°C above the maternal temperature.

Furthermore, the amount of energy deposited by the RF field is heterogeneous, leading to local SARs that differ from the whole-body SAR. The FDA therefore limits local SAR to 8 W/kg for any 5-minute period in 1 g of tissue, and the IEC limits local SAR to 10 W/kg for any contiguous 10 g of tissue. A 2006 analysis concludes that fetal SAR rises through pregnancy, peaking in the 5th month, and operation in normal mode does not violate the IEC local SAR limits, but it finds that operation in first-level controlled may exceed the IEC limit on local SAR beginning in the 3rd month. It is therefore recommended that scans above normal mode are avoided, the maternal abdomen is kept out of the RF field when possible, care is exercised to identify fetuses with poor placental function (i.e., growth retardation), and to avoid scanning pregnant women with impaired thermoregulatory function (e.g., febrile pregnant women). The remaining two sources of exposure in MRI include the time-dependent magnetic fields and the static magnetic field. The time-dependent magnetic field produces an electric field that can cause painful stimulation of nerves and muscles. The FDA imposes a limit for the rate of change on these time-dependent magnetic fields at a threshold “sufficient to produce severe discomfort or painful nerve stimulation.” The IEC has
imposed more technical limitations, and the FDA limits for the fetus are the same as for the mother. Finally, at high–static magnetic field strengths, various nonharmful side effects (e.g., vertigo) and the induction of electric currents have been observed. The FDA therefore limits the field strength of the static magnetic field to 4 T for neonates and infants aged 1 month or less, and to 8 T for patients aged more than 1 month.

Gadolinium–based contrast agents are sometimes used intravenously during MRI studies. The FDA gives these contrast agents a pregnancy risk factor classification of C (i.e., “risk cannot be ruled out”), but a review by the European Society of Urogenital Radiology concludes gadolinium–based agents are likely safe for pregnant and lactating women.

ULTRASONOGRAPHY
Ultrasonography continues to be the screening modality of choice for evaluation of the maternal pelvis and the fetus because of its safety profile, relatively low cost, and real-time capability. Obstetric and gynecologic US accounts for more than half of the US imaging volume in the United States. In the 40 years since its introduction into clinical practice, US has not been shown to convey any significant health risk to the fetus or mother, although most safety data were collected before 1992, when the permissible power output of scanners for fetal use was increased from 94 mW/cm² to 720 mW/cm². Generally, the increasing power output raises concern for thermal and mechanical effects on developing tissue. A large fraction of US energy is absorbed as it passes through tissue, approximately 20% for each centimeter traveled. Damage can occur if the amount of deposited energy exceeds the tissues’ ability to dissipate the heat, but significant temperature increases are not seen until ultrasonic intensities exceed 10 W/cm². These issues are small with standard B-mode imaging, which operates with power less than 50 mW/cm², and more concerning with the use of higher-power Doppler devices, which have the capacity to raise temperatures by several degrees and use powers approaching the established limit. US societies have developed unitless output display standards, namely, a thermal index and mechanical index, to allow the operator to determine whether the study exceeds a generally accepted safe range. The radiologic principle known as ALARA is generally supported and promotes a balance between obtaining the necessary medical information while using minimal settings and examination time. Human data accumulated over a 25-year period have revealed no consistent adverse effects from prenatal diagnostic US examination. US in pregnancy is considered a safe procedure.

The American College of Obstetricians and Gynecologists has reviewed the effects of radiography, US, and MRI during pregnancy and suggested guidelines for radiographic examination during pregnancy (Box 71.4).

SUMMARY
In summary, the threshold dose for the nonstochastic effects of radiation throughout the gestational period is greater than 50 mGy. Prenatal doses of less than 50 mGy present no measurable increased risk for prenatal death, malformation, growth retardation, or impairment of mental development over the background incidence of these entities. The risk for stochastic effects, carcinogenesis, or mutagenesis, is related to the fetal absorbed dose and is very low in comparison to the natural background incidence of childhood cancer and genetic disease for most diagnostic procedures.

The vast majority of radiographic imaging obtained in the emergency department exposes the fetus to 100 times less than the threshold for adverse effects. The 50 mGy threshold for onset of concern for adverse fetal effects is quite conservative, and any statistically significant change in fetal outcome probably requires at least several times this dose. One of the methods put forth in this chapter can be used to counsel pregnant patients in need of diagnostic imaging. For women inadvertently exposed to radiation before pregnancy is recognized, open and frank discussion can help educate patients and alleviate their fear.

BOX 71.4 Guidelines for ED Diagnostic Imaging during Pregnancy

1. Fetal risk of anomalies, growth restriction, or abortion have not been reported with radiation exposure of less than 50 mGy, a level above the range of exposure for diagnostic procedures.
2. If these techniques (radiographs, CT, NM) are necessary in addition to ultrasonography or magnetic resonance imaging or are more readily available for the diagnosis in question, they should not be withheld from a pregnant patient.
3. US and MRI are not associated with risk and are the imaging techniques of choice for the pregnant patient, but they should be used prudently and only when use is expected to answer a relevant clinical question or otherwise provide medical benefit to the patient.
4. Consultation with an expert in dosimetry calculations may be helpful in calculating the estimated fetal dose when multiple diagnostic radiographs are performed on a pregnant patient.
5. Radioactive isotopes of iodine are contraindicated for therapeutic use during pregnancy. Technetium-99m is the isotope of choice.
6. Radiopaque and paramagnetic contrast agents are unlikely to cause harm and may be of diagnostic benefit, but these agents should be used during pregnancy only if the benefit justifies the potential risk to the fetus.

CT, Computed tomography; ED, emergency department; MRI, magnetic resonance imaging; NM, nuclear medicine; US, ultrasonography.
Modified from American College of Obstetricians and Gynecologists (ACOG), Committee on Obstetric Practice: Guidelines for diagnostic imaging during pregnancy and lactation. [ACOG Committee Opinion No. 656], Washington, DC, 2016, ACOG.

CLINICAL USE OF RADIOCONTRAST MATERIAL
Emergency clinicians must frequently initiate studies with the use of radiocontrast material. A full discussion of these procedures is not within the scope of this chapter, but basic issues of contrast material–induced nephropathy, the possible prevention thereof, and recent concern over the use of gadolinium for MRI studies have been included for completeness and ready reference (Table 71.5 and Box 71.5).
TABLE 71.5  Prevention of Contrast-Induced Nephropathy

Renal Failure: Radiocontrast Agents

Radiocontrast-induced acute renal failure is more likely to occur in the presence of advanced age, renal insufficiency, diabetes mellitus, severe congestive heart failure, multiple myeloma, volume depletions, low-cardiac output states, and high-dose contrast-enhanced studies (> 125 mL). The incidence of nephrotoxicity depends on the underlying risk factors and the sensitivity of the measure used to determine nephrotoxicity. When a rather sensitive index of renal dysfunction (an increase in the level of serum creatinine to > 0.3 mg/dL and > 20% on day 1, 2, or 3 and day 5, 6, or 7) is used, the incidence of nephrotoxicity is approximately 2% in nondiabetic, nonazotemic patients and 16% in diabetic, nonazotemic patients. Diabetic patients with azotemia had approximately a 38% incidence of nephrotoxicity. In a study of 59 diabetic patients with advanced azotemia (mean serum creatinine level, 5.9 mg/dL) undergoing coronary angiography, contrast-related nephrotoxicity developed in 30 (51%), as defined by a serum creatinine level that was 25% above baseline 48 hr after angiography. Nine patients (15%) required hemodialysis.

Risk Factors for Radiocontrast Nephrotoxicity

- Advanced age
- Renal insufficiency
- Decreased absolute and effective circulatory volume
- Diabetes mellitus
- Multiple myeloma
- Coadministration of other nephrotoxic agents

(From Brenner BM, editor: *Brenner and Rector’s the kidneys*, ed 7, Philadelphia, 2004, Saunders, Table 34.3; from MD Consult.)

Note: Renal failure may be oliguric or nonoliguric, with nonoliguric renal failure being more common in patients with previously nearly normal renal function. Most episodes of contrast-induced nephrotoxicity are mild and characterized by a reversible 1- to 3-mg/dL rise in serum creatinine; dialysis therapy is rarely needed and usually only in patients whose baseline serum creatinine level is high, for example, > 3 mg/dL.

Prevention of Radiocontrast Nephropathy

The development of acute renal failure significantly complicates the use of intravascular contrast medium (CM) and is linked with high morbidity and mortality. The increasing use of CM, an aging population, and an increase in chronic kidney disease (CKD) will result in an increased incidence of contrast-induced nephropathy (CIN), unless preventive measures are used. At the current time there is no universally accepted, agreed, or proven intervention to totally prevent CIN. However, the Canadian Association of Radiologists has developed guidelines as a practical approach to risk stratification and potential prevention of CIN. The major risk factor predicting CIN is preexisting CKD, which can be predicted from the glomerular filtration rate (GFR). In terms of being an absolute measure, serum creatinine is an unreliable measure of renal function. Patients with a GFR > 60 mL/min have a very low risk for CIN, and preventive measures are generally unnecessary. With a GFR < 60 mL/min, preventive measures should be instituted. The risk for CIN is greatest in patients with a GFR < 30 mL/min. As preventive measures, alternative imaging that does not require CM should be considered. Fluid volume loading is the single most important protective measure. Nephrotoxic medications should be discontinued 48 hr before the study. CM volume and the frequency of administration should be minimized, but satisfactory image quality should still be maintained. High-osmolar CM should be avoided in patients with renal impairment. There is some evidence to suggest that isosmolar CM reduces the risk for CIN in patients with renal impairment, but further study is necessary to determine whether isosmolar CM is superior to low-osmolar CM. N-acetylcysteine (NAC) has been advocated to reduce the incidence of CIN; however, not all studies have shown a benefit, and it is difficult to formulate evidence-based recommendations at this time. Use of NAC may be considered in high-risk patients but is not thought to be mandatory.

Additional Caveats

1. Metformin use is not a contraindication to the use of CM, but metformin should be withheld for 48 hr after CM and after evaluation of renal function.

(From Benko A: Canadian Association of radiologists: consensus guidelines for the prevention of contrast-induced nephropathy, *Can Assoc Radiol J* 58:79, 2007.)

Potential Prevention of Contrast-Induced Nephropathy

**NAC**

Suggested regimen for oral NAC: 600 mg (3 mL of a 20% solution in liquid) twice a day for 24 hr before and 24 hr after procedure.

Suggested IV (intravenous) NAC: 150-mg/kg IV bolus over 30 min, followed by a 50-mg/kg infusion over 4 hr.

Example in a 80-kg patient: 12,000 mg NAC in 500 mL normal saline over 30 min, followed by 4000 mg NAC in 500 mL normal saline over 4 hr.
### TABLE 71.5 Prevention of Contrast-Induced Nephropathy—cont’d

#### Hydration

Suggested regimen for fluid therapy in elective cases: normal saline, at least 1 mL/kg per hr 12 hr before and 12 hr after the procedure.

Alternative if an emergency procedure is required: 5-mL/kg bolus of normal saline 1 hr before and 1 mL/kg per hr for 12 hr after the procedure.

Alternative fluid regimen with bicarbonate: add 154 mL of 1000 mEq/L sodium bicarbonate to 850 mL of 5% dextrose in water (D₅W) (or add 3 ampules of standard bicarbonate to 1 L D₅W). Initial bolus of 3 mL/kg for 1 hr before injection of contrast material, followed by 1 mL/kg per hr for 6 hr after the procedure.

*Suggested but unproven, minimal downside.*


### Dispensing and Administration Guidelines for CIN

#### NAC + Hydration

**ORAL NAC DOSING**

Give NAC, 600 mg liquid (3 mL of a 20% solution) in 9 mL ginger ale or cola

**ORAL NAC DOSING**

0.9% sodium chloride IV fluid at 1 mL/kg per hr 12 hr before and after catheterization (normal saline preferred, but 0.45% has also been used with success)

#### IV Sodium Bicarbonate (154 mEq/L) Mixed in 1 L of D₅W

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>3-mL/kg IV BOLUS OVER 1 hr</th>
<th>1 mL/kg per hr × 6 hr IV INFUSION</th>
<th>TOTAL mL INFUSED</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>180 mL</td>
<td>360 mL (60 mL/hr)</td>
<td>540 mL</td>
</tr>
<tr>
<td>70</td>
<td>210 mL</td>
<td>420 mL (70 mL/hr)</td>
<td>630 mL</td>
</tr>
<tr>
<td>80</td>
<td>240 mL</td>
<td>480 mL (80 mL/hr)</td>
<td>720 mL</td>
</tr>
<tr>
<td>90</td>
<td>270 mL</td>
<td>540 mL (90 mL/hr)</td>
<td>810 mL</td>
</tr>
<tr>
<td>100</td>
<td>300 mL</td>
<td>600 mL (100 mL/hr)</td>
<td>900 mL</td>
</tr>
<tr>
<td>≥ 110</td>
<td>330 mL</td>
<td>660 mL (110 mL/hr)</td>
<td>990 mL</td>
</tr>
</tbody>
</table>


#### IV NAC

<table>
<thead>
<tr>
<th>Weight (kg)</th>
<th>150-mg/kg IV BOLUS OVER 30 min</th>
<th>50-mg/kg IV INFUSION OVER 4 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>9000 mg/500 mL NS</td>
<td>3000 mg/500 mL NS</td>
</tr>
<tr>
<td>70</td>
<td>10,500 mg/500 mL NS</td>
<td>3500 mg/500 mL NS</td>
</tr>
<tr>
<td>80</td>
<td>12,000 mg/500 mL NS</td>
<td>4000 mg/500 mL NS</td>
</tr>
<tr>
<td>90</td>
<td>13,500 mg/500 mL NS</td>
<td>4500 mg/500 mL NS</td>
</tr>
<tr>
<td>≥ 100</td>
<td>15,000 mg/500 mL NS</td>
<td>5000 mg/500 mL NS</td>
</tr>
</tbody>
</table>

NS, Normal saline.

CONTRAINDICATIONS TO MRI

Overview
There are few contraindications to MRI. Overall, no biologic adverse effects are associated with conventional MRI. Most contraindications to MRI are relative and essentially precautions related to the effect of MRI on devices and material within the body that may be affected by the magnetic field of MRI.

Implanted Devices and Foreign Bodies
Electronic devices and magnetizable material represent potential hazards to the patient. Titanium objects are safe for MRI.

- **Intracoronary stents**: It is considered safe to perform MRI at any time after placement of coronary artery stents of any type.
- **Sternal wires after sternotomy**: Sternal wire sutures are considered safe for MRI.
- **Mechanical cardiac valves**: It is safe to scan most prosthetic cardiac valves because at most, they experience only a mild torque. An exception involves the pre-6000 series Starr-Edwards caged ball valves; devices rarely used now.
- **Pacemakers, implantable defibrillators, and implanted electronic devices**: The risks of scanning patients with cardiac pacemakers are related to possible movement of the device, magnetically induced changes in programming, electromagnetic interference, and induced currents in lead wires leading to heating and/or cardiac stimulation. It is currently considered inadvisable for patients with pacemakers or other intracardiac wires to undergo MRI. Nerve stimulators, insulin pumps, cochlear implants, and other implanted electronic devices may also be affected by MRI and are considered unsafe.
  - **Implanted vagal nerve stimulator**: Brain MRI performed at less than 2 T, with a send and receive head coil and the stimulator turned off, appears to be safe under guidelines published by the manufacturer. Other MRI studies are not known to be safe.
  - **Aneurysm clips and magnetizable material**: Any ferromagnetic object within the body represents a potential hazard when exposed to the large magnetic field of an MRI system. The hazard primarily reflects the possibility of deflecting the foreign body sufficiently to injure vital structures. For example, certain older-model vascular clips

### INFORMATION FOR HEALTH CARE PROFESSIONALS:
**GADOLINIUM-BASED CONTRAST AGENTS FOR MRI (MARKETED AS MAGNEVIST, MULTIHANCE, OMNISCAN, OPTIMARK, PROHANCE)**

FDA ALERT (6/2006, updated 12/2006 and 5/23/2007): This updated alert highlights the FDA's request for the addition of a boxed warning and new warnings about the risk for nephrogenic systemic fibrosis (NSF) and full prescribing information for all gadolinium-based contrast agents (GBCAs) (Magnevist, MultiHance, Omniscan, OptiMARK, ProHance). This new labeling highlights and describes the risk for NSF following exposure to a GBCA in patients with acute or chronic severe renal insufficiency (glomerular filtration rate ≤ 30 mL/min per 1.73 m²) and patients with acute renal insufficiency of any severity as a result of hepatorenal syndrome or in the perioperative liver transplantation period. In these patients, avoid the use of a GBCA unless the diagnostic information is essential and not available with non–contrast-enhanced MRI. NSF may result in fatal or debilitating systemic fibrosis. The requested changes in GBCA product labeling are summarized below.

- Evaluate renal function in all patients before administering a GBCA.
- Whenever possible, avoid GBCAs for MRI and MRA in patients with moderate to end-stage renal failure. Contrast agents that contain gadolinium include Magnevist, MultiHance, Omniscan, OptiMARK, and ProHance.
- If GBCAs must be used, consider prompt dialysis after the procedure to eliminate circulating gadolinium. However, it is unknown whether dialysis can prevent or treat NSF/NFD.
- Encourage patients to contact their health care provider if they have signs of NSF/NFD.
- Report cases of NSF/NFD to the FDA’s MedWatch program at www.fda.gov/medwatch/index.html or by calling 1-800-332-1088 (1-800-FDA-1088).

### CONTRAINDICATIONS TO MRI

(Because this is an area of continuing change and there are rapid advancements in the technology to produce MRI-safe materials, consultation with the radiology department is suggested if any questions arise concerning the safety of MRI.)

---

**TABLE 71.5** Prevention of Contrast-Induced Nephropathy—cont’d

| UMMC Pharmacy Cost Comparisons with Treatment Regimens for CIN |
|---|---|
| **Regimens** | **Regimens** |
| 600 mg NAC × 8 doses (oral) | $5.82 |
| 2 L NS | $1.60 |
| Sodium bicarbonate, 154 mEq/L D$_2$W | $1.32 (drug only) |
| 70-kg patient: 10,500 mg NAC/500 mL NS by IV bolus | $183.00 (drug only) |
| 70-kg patient: 3500 mg NAC/500 mL NS by IV infusion | $61.00 (drug only) |

*UMMC, University of Maryland Medical Center.*
used for cerebral aneurysms are ferromagnetic and could be moved by the magnetic field, with obviously dire consequences.

- **Intraorbital or intraocular metallic fragments**, such as might be acquired from machining, are a potential risk and generally a contraindication to MRI.

- **Cutaneous metal objects**: Although most metallic biomaterial is now nonferrous and nonmagnetizable, any metallic device within or connected to the patient needs to be evaluated for safety. Dental alloys, wires, splints, dental braces, and prostheses do not appear to pose a risk to the patient, although such material may result in artifactual changes. Cutaneous burns can result from contact of the skin with metal objects, including neurosurgical halo pins, pulse oximetry probes, and drug-eluting medical patches that contain metal foil (e.g., nicotine patch), although the mechanism of this injury is unclear.

- **Orthopedic/neurosurgical hardware**: It is safe to perform MRI in patients with titanium implants, screws, rods, and artificial joints.

- **Bullets/shrapnel**: These foreign objects within the body are relative contraindications to MRI. Many bullets are safe, but those with metal (specialized bullets, such as metal jackets) may pose a risk.

- **Tattoos**: The majority of professionally obtained tattoos are safe for MRI; however, tattoos containing lead, such as those obtained in prison, can burn the skin if exposed to MRI.

- **Oxygen cylinders**: Standard metal oxygen cylinders should not be used in the MRI suite. Safe oxygen cylinders are available.

- **Credit cards**: Credit card and other information-containing strips may be destroyed in the MRI scanner.

**REFERENCES ARE AVAILABLE AT www.expertconsult.com**

FDA, United States Food and Drug Administration; MRA, magnetic resonance angiography; MRI, magnetic resonance imaging; NFD, nephrogenic fibrosing deformity.
REFERENCES

30. Mallinckrodt: Product insert for TechniScan MAA Kit for the preparation of technetium Tc 99m albumin aggregated, 093 (Form No. A09310), St. Louis, 2000, Mallinckrodt.
INTRODUCTION

Performing an invasive procedure on an anticoagulated emergency department (ED) patient can be challenging for the emergency provider (EP). Regardless of the procedure, some anticoagulated patients are at potential significant risk of hemorrhage from the procedure. However, emergency reversal of anticoagulation in order to perform the procedure may also place the patient at risk for serious thrombotic complications. When deciding to perform a procedure on an anticoagulated ED patient, the EP must weigh the risks of bleeding with the risk of disrupting anticoagulation, emergency reversal of anticoagulation, and delaying a potentially critical intervention. If the procedure is not needed for life-saving therapy, postponing the procedure or providing empiric treatment may be a reasonable choice. In contrast, emergency procedures to reverse an imminent life-threatening condition should never be withheld and emergency reversal of anticoagulation may be required. This chapter reviews current literature and recommendations for select ED procedures such as lumbar puncture, central venous catheterization, arthrocentesis, paracentesis, thoracentesis, and tube thoracostomy in the anticoagulated patient.

ASSESSING RISK OF BLEEDING AND THROMBOSIS

All anticoagulants inhibit stable clot formation and increase the risk of bleeding. Routine coagulation testing, that is prothrombin time (PT), partial thromboplastin time (PTT), international normalized ratio (INR), and platelet count, in anticoagulated patients, is recommended prior to performing an invasive procedure. Although laboratory tests to determine drug presence, drug concentration, and level of anticoagulant effect can be useful in the assessment of bleeding risk, standard coagulation assays accurately monitor the degree of anticoagulation for only a few agents. PT and INR conveniently assay the extrinsic pathway to monitor the quantitative effect of warfarin therapy. PTT or activated partial thromboplastin time (aPTT) accurately monitors the anticoagulant effect of heparin. Newer nonvitamin K antagonist oral anticoagulants (NOACs) were developed to have predictable pharmacokinetics with minimal interactions to avoid the need for routine laboratory testing. Coagulation assays to quantify the effect or presence of a NOAC would be useful in many clinical circumstances such as in active bleeding, suspected overdose, and the perioperative period, but do not exist for many. The direct measurement of drug concentration is not suitable in clinical practice because of the time required to perform the laboratory analysis. Standard coagulation assays can be useful in extreme scenarios, but are generally insufficient in determining drug concentration or degree of anticoagulant effect of NOACs (Table 72.1).

Normal thrombin time is suitable for excluding significant dabigatran levels but too sensitive for determining the degree of anticoagulant effect. Dilute thrombin time, ecarin chromogenic assay, and ecarin clotting time may be useful to determine the degree of anticoagulant effect, but is not standardized across laboratories. Although using standard coagulation assays to quantify dabigatran effect is not recommended, an elevated PT/INR suggests a supratherapeutic dabigatran effect, and a normal PTT excludes it. Unfortunately, none of these tests are accurate predictors of bleeding risk with dabigatran and patients may have bleeding when these tests are within the normal range.

Anti-factor Xa activity is valuable to determine the effect of factor Xa inhibitors such as rivaroxaban and apixaban, but this test is not widely available. Although it is not recommended to use standard coagulation assays to quantify drug effect, a normal PT and INR level virtually excludes supratherapeutic rivaroxaban effect.

For patients on warfarin, understanding how INR correlates to the risk of bleeding is important. Spontaneous bleeding events are uncommon when the INR is normal or within a therapeutic range (i.e., 2 to 3). The relative risk for bleeding with an INR between 3 and 5 is 2.7 (95% confidence interval [CI] 1.8–3.9). However, when the INR is above 5, the relative risk of a spontaneous hemorrhagic event increases dramatically to 21.8 (95% CI 12.1–39.4). These findings are similar to a previous retrospective review in 1995 that also showed adverse events occurred in 75 per 100 patient-years for an INR greater than or equal to 6.5.4 Patients with severe coagulopathy (INR greater than 9) have a poor prognosis, with 67% experiencing spontaneous hemorrhagic events and a 74% mortality rate.

Patients without a history of bleeding or anticoagulant use do not require routine coagulation studies prior to performing a procedure unless history or physical examination suggest a bleeding disorder or use of anticoagulation. A 2005 systematic review failed to demonstrate the utility of routine coagulation studies on nonanticoagulated patients prior to the performance of a procedure. Abnormal laboratory findings such as thrombocytopenia, an elevated INR, or other coagulation abnormalities are not necessarily absolute contraindications to performing an invasive procedure. Numerous studies have demonstrated that select procedures can safely be performed even in the setting of anticoagulation with a vitamin K antagonist (VKA). Determining a patient’s risk for bleeding requires examination of both the procedure to be performed, the anticoagulant medication, and the level of anticoagulation. Unfortunately, there are no validated systems available to quantify risk of bleeding.

The risk of thrombosis after anticoagulant reversal is significant for a subset of ED patients. These patients include those with a recent diagnosis of pulmonary embolism, significant clot burden, or those with mechanical hardware such as a prosthetic cardiac valve. When clinically feasible, it is best to allow the anticoagulant effect of a medication to wane rather than emergently reversing the medication in these patients. This approach is rarely possible in the ED, so timing of the procedure should be coordinated with the inpatient provider if the patient requires admission.

Many procedures report contraindications with severe coagulopathy (INR greater than 9) and disseminated intravascular coagulation (DIC). Proper coagulation is significantly
which anticoagulated patients will bleed from a procedure is difficult. Furthermore, the simple act of controlling periprocedural bleeding may not entirely end the risk of serious harm. For instance, periprocedural bleeding with percutaneous coronary intervention is associated with an increased short- and long-term morbidity and mortality, including major adverse cardiovascular events and readmission rates well after the bleeding is controlled.13,14

When managing spontaneous, periprocedural, or postprocedural bleeding associated with anticoagulation, there are a few principles to consider. Most importantly, never withhold emergency life-saving procedures such as endotracheal intubation, tube thoracostomy, cardiac defibrillation, pericardiocentesis, or vascular access when necessary.

Stabilization with supportive treatments such as oxygenation, intravascular volume resuscitation, and repletion of blood products via transfusion are the initial steps of assessment and management of bleeding in the anticoagulated patient. Patients with prolonged or severe bleeding may have presenting symptoms in various stages of circulatory shock. Poor blood supply and intravascular volume leads to poor tissue perfusion. Cellular hypoxia, damage, and resulting inflammatory response from hypoperfusion can exacerbate the patient's clinical status. Adequate resuscitation and stabilization is key to the initial phase of management.

### MANAGEMENT OF ANTICOAGULANT ASSOCIATED BLEEDING

Although periprocedural bleeding is typically not life threatening and can often be controlled with direct pressure, predicting disrupted in these clinical settings and invasive procedures may lead to severe bleeding. However, life-saving measures such as thoracostomy, central venous catheterization, and endotracheal intubation are sometimes required even in these severe settings. Additional life-saving procedures that may carry a high risk of bleeding in the setting of anticoagulation, but should not be withheld, include defibrillation and pericardiocentesis. The gravity of the clinical scenario is important to keep in mind when weighing the risk of performing a procedure in the setting of anticoagulation.

Limited data are available on the safety of performing procedures on patients taking newer NOACs. These NOACs may confer the same, less, or increased risk of bleeding as traditional VKA agents. The lack of an effective reversal agent for many of the NOACs is another factor that complicates the decision to perform a procedure in the setting of anticoagulation. Fortunately, many of the NOACs have a duration of action that is less than the traditional VKAs.

### TABLE 72.1 Summary of Anticoagulant Testing and Management

<table>
<thead>
<tr>
<th>ANTICOAGULANT</th>
<th>LABORATORY TESTING TO CONSIDER</th>
<th>PHARMACOLOGY</th>
<th>AGENTS TO REVERSAL OF REMOVAL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitamin K Antagonists</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low-molecular-weight heparin (LMWH)</td>
<td>Anti-Factor Xa</td>
<td>3–6 hours</td>
<td>Consider protamine sulfate or rVIIa in life-threatening bleeding</td>
</tr>
<tr>
<td>Unfractionated heparin</td>
<td>PTT, aPTT</td>
<td>60–90 minutes</td>
<td>Protamine sulfate</td>
</tr>
<tr>
<td>Warfarin</td>
<td>PT/INR</td>
<td>20–60 hours (duration of action 2–5 days)</td>
<td>Vitamin K, fresh frozen plasma (FFP), prothrombin complex concentrate (PCC), recombinant Factor VIIa</td>
</tr>
<tr>
<td>Direct Thrombin Inhibitors</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Argatroban</td>
<td>Activated clotting time (ACT), aPTT</td>
<td>39–51 minutes</td>
<td>Consider hemodialysis, rFVIIa or FFP but generally not indicated due to short half-life</td>
</tr>
<tr>
<td>Dabigatran</td>
<td>Thrombin time (TT), dilute TT, ecarin chromogenic assay (ECA), and ecarin clotting time (ECT), PT/INR, PTT</td>
<td>12–17 hours</td>
<td>Idarucizumab, PCC, hemodialysis</td>
</tr>
<tr>
<td>Factor Xa Inhibitors</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Apixaban</td>
<td>Anti-Factor Xa, PT/INR</td>
<td>12 hours</td>
<td>PCC</td>
</tr>
<tr>
<td>Rivaroxaban</td>
<td></td>
<td>5–9 hours</td>
<td></td>
</tr>
<tr>
<td>Anti-Platelet Agents</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aspirin</td>
<td>Platelet function testing</td>
<td>Duration of action 7–10 days, irreversibly inhibits platelet</td>
<td>Platelet transfusion</td>
</tr>
<tr>
<td>Clopidogrel</td>
<td></td>
<td>7–10 hours</td>
<td></td>
</tr>
<tr>
<td>Prasugrel</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ticagrelor</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*aPTT, Activated partial thromboplastin time; INR, international normalized ratio; PT, prothrombin time; PTT, partial thromboplastin time.*
Hemostatic measures such as direct compression of a bleeding or oozing vessel, wound, or region should be performed concurrently with stabilization to prevent further blood loss. Additional measures may be needed to control bleeding when compression fails or when the bleeding originates from a noncompressible site. These measures include the use of topical hemostatic agents, systemic hemostatic agents (i.e., tranexamic acid), procedural intervention, operative management, or intraarterial embolization.

The clinician can assess the severity of coagulopathy with laboratory analysis. Patients may experience bleeding at subtherapeutic, therapeutic, or supratherapeutic levels of anticoagulation. Determining qualitative level of anticoagulation may help guide therapy. However, many anticoagulants do not have an associated diagnostic test that is accurate, timely, and clinically relevant to determine severity of coagulopathy like warfarin or heparin. Obtain routine coagulation assays (i.e., PT, PTT, and INR) when no relevant study exists. Routine coagulation assays may sometimes be helpful in determining the lack of a supratherapeutic anticoagulant effect but are often difficult to interpret.

When bleeding is severe, life-threatening, refractory to hemostatic efforts, or the coagulopathy is determined to be severe, restoring the ability to generate an effective clot by administration of an antidote, coagulation factor, blood product, or removal of the offending anticoagulant (i.e., hemodialysis) may be necessary. Consideration for the anticoagulant’s duration of action is also important when deciding whether to reverse anticoagulation.

Withholding further doses of anticoagulation may be necessary after stabilization and disposition. When the clinical decision to withhold anticoagulant is not clear, consultation with the prescribing physician (e.g., cardiologist prescribing aspirin and clopidogrel for cardiac stent) or hematologist may be warranted.

### Reversal of Anticoagulation

The decision to discontinue or reverse anticoagulation prior to a procedure is difficult. Although there are no validated data that can accurately stratify risk for peri- or postoperative thromboembolism, the American College of Chest Physicians (ACCP) has published guidelines that stratify patients into low-, moderate-, and high-risk groups. These guidelines are listed in Table 72.2.

Risk factors that increase the likelihood of a thromboembolic event with anticoagulant disruption include a prosthetic heart valve, atrial fibrillation, recent cerebrovascular accident (CVA) or transient ischemic attack, recent venous thromboembolism (VTE), the presence of a VTE risk factor (i.e., protein C/S deficiency, antiphospholipid antibody), and a high CHADS2 score (congestive heart failure, hypertension, age, diabetes mellitus, CVA). Patients with mechanical valves are at higher risk for thromboembolic events when anticoagulation is disrupted than patients with bioprosthetic valves. Mitral valves present a higher risk of thromboembolic events than aortic prosthetic valves. The postoperative period just after (less than 3 months) mechanical valve placement is also a high-risk time for thromboembolic events.

Procedures on patients with prosthetic valves typically require a delayed approach to operative management, except when there is an emergency. In general, discontinuing anticoagulation in a patient with a prosthetic valve is safer than reversal of anticoagulation to perform a procedure. Reversal of anticoagulation puts patients at higher risk for thromboembolic events than does simply discontinuing the anticoagulant. When possible, discontinue anticoagulation and delay invasive procedures on patients with prosthetic valves until coagulation status returns to near normal. For patients with low risk bioprosthetic valves that require discontinuation of anticoagulation, warfarin should be discontinued 48 to 72 hours prior to the procedure or until the

### TABLE 72.2 Suggested Patient Risk Stratification for Perioperative Arterial or Venous Thromboembolism

<table>
<thead>
<tr>
<th>RISK STRATUM</th>
<th>MECHANICAL HEART VALVE</th>
<th>ATRIAL FIBRILLATION</th>
<th>VTE</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Any mitral valve prosthesis</td>
<td>CHADS2 score of 5 or 6</td>
<td>Recent (within 3 mo) VTE. Severe thrombophilia (e.g., deficiency of protein C, protein S or antithrombin, antiphospholipid antibodies, or multiple abnormalities)</td>
</tr>
<tr>
<td></td>
<td>Older (caged-ball or tilting disc) aortic valve prosthesis</td>
<td>Recent (within 3 mo) stroke or transient ischemic attack, Rheumatic valvular heart disease</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Recent (within 6 mo) stroke or transient ischemic attack</td>
<td>VTE within the past 3 to 12 mo; Nonsevere thrombophilic conditions (e.g., heterozygous factor V Leiden mutation, heterozygous factor II mutation); Recurrent VTE; Active cancer (treated within 6 mo or palliative)</td>
<td></td>
</tr>
<tr>
<td>Moderate</td>
<td>Bileaflet aortic valve prosthesis and one of the following: atrial fibrillation, prior stroke or transient ischemic attack, hypertension, diabetes, congestive heart failure, age &gt; 75 yr</td>
<td>CHADS2 score of 3 or 4</td>
<td></td>
</tr>
<tr>
<td>Low</td>
<td>Bileaflet aortic valve prosthesis without atrial fibrillation and no other risk factors for stroke</td>
<td>CHADS2 score of 0 to 2 (and no prior stroke or transient ischemic attack)</td>
<td>Single VTE occurred 12 mo ago and no other risk factors</td>
</tr>
</tbody>
</table>

CHADS2, Congestive heart failure-hypertension-age-diabetes-stroke; VTE, venous thromboembolism.

INR drops below 1.5. Anticoagulation may then be restarted 24 hours following the procedure. High-risk patients with prosthetic valves (e.g., mitral valve) require bridging therapy to reduce the time off anticoagulation.

For patients taking VKAs that require reversal of anticoagulation for an urgent procedure, administration of oral vitamin K is recommended. If more immediate reversal is required for a procedure, administration of fresh-frozen plasma or prothrombin complex concentrate in addition to oral vitamin K is recommended. For patients taking antiplatelet agents such as aspirin or clopidogrel, platelet transfusion is recommended for reversal of the effect (Fig. 72.1).

**Figure 72.1** Anticoagulant and coagulopathy reversal agents. A, Oral vitamin K reverses the effect of vitamin K antagonists, such as warfarin. Onset of action ranges from hours to days. B, Intravenous vitamin K can be used in patients who require emergency procedures; however, onset of action is measured in hours, as the liver must regenerate coagulation factors. There is a risk of anaphylaxis with intravenous vitamin K, and it should be administered via slow infusion (over 20 to 60 minutes.) C, Fresh frozen plasma (FFP) can be used to reverse the effect of vitamin K antagonists in emergency situations. D, Prothrombin complex concentrates (such as the four factor PCC Kcentra [CSL Behring LLC., Kankakee, IL], pictured) provides rapid international normalized ratio correction and may be safer than FFP. PCCs can also be used to reverse the effects of direct thrombin inhibitors (e.g., dabigatran) and factor Xa inhibitors (e.g., apixaban). E, Platelet transfusions can be used to reverse the effects of antiplatelet medications such as aspirin or clopidogrel, and also to correct thrombocytopenias.
Procedures

Lumbar Puncture

The decision to perform a lumbar puncture (LP) should be individualized and based on an assessment of the risks and benefits of the procedure. When the risks of an LP are high, empiric antibiotic therapy may be appropriate for conditions such as meningitis. When the condition is difficult to diagnose or treat, LP may be necessary. In these cases, consultation with interventional radiology or anesthesiology may be warranted to reduce the risk of hemorrhage.

When an LP is performed in an anticoagulated ED patient, it is critical to monitor for complications. The most important complication associated with LP in the setting of anticoagulation is a spinal epidural hematoma (SEH). This rare, but catastrophic, complication is more likely to occur with a difficult or traumatic LP in anticoagulated patients or those with a platelet disorder. An emergency magnetic resonance image should be performed if there is suspicion for an SEH. If an SEH is diagnosed, immediate decompression with a laminectomy should be performed to avoid irreversible spinal cord ischemia.

Many clinicians feel that spinal procedures such as an LP may be safely performed on patients taking aspirin or a nonsteroidal antiinflammatory drug (NSAID) without discontinuing the medication or transfusing platelets. It is important to consider the specific medication. Aspirin irreversibly inactivates cyclooxygenase-1 (COX-1) and blocks thromboxane production for the life of a platelet (7–10 days) within an hour of ingestion. Therefore platelet function does not return to normal until the permanently inhibited platelets are replaced by enough newly synthesized active platelets. The ability for platelets to aggregate can be seen after 4 days following aspirin cessation, as it requires only a few uninhibited platelets (newly synthesized or transfused) to recruit aspirin treated platelets to action. Horlocker and colleagues studied 391 patients undergoing spinal anesthesia and also on antiplatelet therapy including aspirin, naproxen, piroxicam, ibuprofen, indomethacin, dipyridamole, and sulindac. One hundred and thirteen patients were on multiple antiplatelet agents preoperatively. No SEHs were reported in this study. The authors concluded that preoperative antiplatelet medication was not a contraindication to spinal anesthesia. Importantly, it is not recommended to perform an LP on patients who are using aspirin or an NSAID concurrently with another anticoagulant such as heparin, low-molecular-weight heparin (LMWH), or other antiplatelet agents.

The safety of performing an LP in patients receiving newer antiplatelet agents, including clopidogrel, ticlopidine, abciximab, eptifibatide, or tirofiban is not well studied. The American Society of Regional Anesthesia and Pain Medicine (ASRA) recommends discontinuation of these medications prior to LP. Reversing the effect of these medications and the period of discontinuation prior to the performance of the procedure depends on the medication. Pharmacologically, normal platelet activity is expected 8 hours after discontinuing tirofiban and eptifibatide, 24 to 48 hours after discontinuing abciximab, and 1 to 2 weeks after discontinuing clopidogrel and ticlopidine. The ASRA, ACCP, and the American Heart Association (AHA) recommend discontinuing clopidogrel 7 to 10 days prior to neuraxial anesthesia or surgery. Other organizations recommend a 5-day washout period prior to giving spinal injections in patients receiving these medications.

Heparin increases the risk of SEH. In a study from 1981, 2% (7 of 342) of patients receiving heparin for anticoagulation developed SEH after LP. Risk for SEH increased with a traumatic procedure, starting anticoagulation within an hour of the LP, and concomitant aspirin therapy. Patients with preexisting coagulopathy were excluded from this study. This study suggests that the risk of SEH in patients on heparin prior to the LP is at least 2%. In a study by Tryba and colleagues, the risk of SEH increased tenfold in patients receiving heparin or aspirin who experienced a traumatic LP. As intravenous (IV) heparin has a short duration of action, it should be discontinued for at least 4 hours prior to the LP and the aPTT has normalized. Heparin infusion should not be resumed for at least an hour after an LP is performed. The risk for SEH from LP in patients on twice daily subcutaneous heparin for the prophylaxis of deep venous thrombosis is low. Nonetheless, it is recommended to discontinue for at least 8 hours prior to neuraxial procedures.

LMWH has a half-life that can range between 2 and 6 hours depending on the dose, route of administration, and renal function. The ASRA guidelines recommend discontinuing LMWH 12 hours prior to neuraxial procedures when used at doses intended for deep vein thrombosis prophylaxis and 24 hours when used at doses intended for anticoagulation. Anticoagulation with LMWH should be withheld for 18 to 24 hours after the LP to prevent SEH.

Warfarin is associated with a high risk of SEH following an LP. Warfarin should be stopped for 5 days and the INR normalized prior to performing an LP. The administration of vitamin K with fresh frozen plasma (FFP), or the administration of prothrombin complex concentrates (PCCs) is recommended for complete warfarin reversal.

There is currently no study that has evaluated the safety of LP in the setting of anticoagulation with the new direct thrombin inhibitors or factor Xa inhibitors. Little is known about procedural safety when anticoagulated with these medications, although many clinicians assume that the risk of bleeding is increased similarly to warfarin. Duration of action, reversal strategies, and diagnostic testing are complicated and medication specific.

Performance of an LP in patients with hemophilia is safe following 100% factor replacement. In a study by Silverman and colleagues, 30 of 33 patients (91%) with severe factor deficiency had no serious complications with LP after adequate factor replacement. In a review of six articles that evaluated neuraxial procedures in patients with hemophilia, there were no SEHs in 107 procedures on 85 patients, of which 53 of the procedures were diagnostic LPs in the ED. In 105 of the 107 procedures (98%), factor levels were replaced to normal. One case of an SEH with neurologic impairment occurred in a patient with undiagnosed hemophilia.

There is scant literature on the performance of an LP in patients with select platelet disorders, including von Willebrand disease (vWD) and idiopathic thrombocytopenic purpura (ITP). Choi and colleagues evaluated 74 neuraxial procedures (all for obstetrical anesthesia) performed in 72 patients with vWD. Sixty-four patients (86%) required no treatment secondary to normal vWD indices, whereas 10 patients required treatment with desmopressin (DDAVP), vWF/factor VIII concentrate, or factor VIII alone. No complications were noted. In the same study, there were no complications reported in 326 neuraxial procedures in patients with ITP. Pretreatment with corticosteroids, IV immune globulin, or platelet transfusion...
was variable among the reports. No pretreatment was provided in 103 procedures that included a patient with a platelet count of 2 × 10^9/L. Based on the results of their study, the authors concluded that it is safe to perform a lumbar puncture without providing platelet transfusion when the platelet count is greater than 50 × 10^9/L.

Performing an LP in the setting of thrombocytopenia has been well studied in the pediatric oncology population. In 2000, Howard and colleagues reported on the safety of performing LPs in children with platelet counts greater than 10 × 10^9/L without transfusing platelets. Nine hundred forty-one procedures were performed on patients with platelet counts below 50 × 10^9/L without the development of an SEH. Twenty-nine procedures were performed on patients with platelet counts less than or equal to 10 × 10^9/L. No complications were reported, although the study was not powered to determine patient safety. Nonetheless, the authors recommended platelet transfusion for an LP when the platelet count is less than or equal to 10 × 10^9/L. A 2010 systematic review by van Veen states that a platelet count greater than or equal to 40 × 10^9/L is safe to perform an LP provided that the platelet count is stable, the patient does not have an acquired or congenital coagulopathy, the platelet function is normal, and the patient is not receiving an antiplatelet or anticoagulant medication. The authors also stated that it may be safe to perform an LP at platelet counts 20 to 40 × 10^9/L, however there were insufficient data to recommend safety at this platelet level without transfusion. The American Association of Blood Banks recommends prophylactic platelet transfusion for patients having an elective diagnostic LP for a platelet count less than 50 × 10^9/L. However, this is a weak recommendation based on very low-quality evidence.

**Vascular Access**

Patients frequently require central venous access for resuscitation, medication administration, or hemodynamic monitoring. Critically ill patients are sometimes anticoagulated or develop a coagulopathy secondary to their illness. Providers are oftentimes hesitant to perform central venous cannulation in anticoagulated patients or those with a coagulopathy because of the risk of bleeding.

Bleeding complications during central venous catheterization in patients with abnormal hemostasis is relatively low. Arterial puncture is the most common complication associated with central venous catheter insertion, and occurs in 3% of catheters placed in the internal jugular vein and 0.5% of catheters placed in the subclavian vein. Numerous studies have demonstrated that central venous cannulation is safe in patients with a coagulopathy or disorder of hemostasis. Emergency reversal of anticoagulation prior to the insertion of a central venous catheter is common practice but does not appear to be evidence-based.

In one study, 76 consecutive patients with various disorders of hemostasis (thrombocytopenia, anticoagulation with heparin, anticoagulation with warfarin, and abnormalities in coagulation assays) received central venous access procedures. Of the 104 procedures, only one patient required an intervention beyond 20 minutes of site compression to control bleeding. The authors noted that bleeding problems are uncommon and serious bleeding is rare. Providers should make a case-based clinical decision weighing the risk of hemorrhage versus the risk of thrombosis and fluid overload associated with reversal.

In a 2000 study evaluating the complications of central venous catheter placement in patients with various hemostasis disorders, thrombocytopenia (platelet count less than 50 × 10^9/L) was the only significant predictor for bleeding complications. However, the authors call into question the utility of platelet transfusion because the complications were easily addressed with compression and suture. Other studies have also found that a platelet count less than 50 × 10^9/L is an independent risk factor for bleeding. Further studies are required to assess the benefit of preprocedural platelet transfusion for central venous catheter insertion. Platelet transfusion to maintain counts greater than 50 × 10^9/L is recommended in patients with evidence of postprocedural bleeding at catheter insertion site until hemostasis is achieved.

In 1999, Fisher and colleagues demonstrated a low incidence of major vascular complications in patients with cirrhosis and an elevated INR. Of 658 cannulations, there was one major event (hemothorax) caused by accidental subclavian artery puncture. The mean INR was 2.4 (range 1 to 16) and no attempt was made to reverse coagulopathy prior to the procedure. Similar results were found in two additional studies on coagulopathic patients including patients with liver disease receiving central venous access. Patients with DIC should have vigorous correction of coagulopathy abnormalities in conjunction with hematology consultation prior to central venous catheterization when patient stability allows for it. Targets for correction include PT less than 1.5 times normal and fibrinogen greater than 1.0 g/L.

Factor replacement is recommended for patients with hemophilia prior to central venous cannulation. The goal of management is to achieve a circulating factor level of 100% prior to the procedure followed by a continuous factor infusion for 2 to 3 days afterwards to prevent serious bleeding complications.

In nonurgent situations IV heparin infusion should be discontinued for 3 hours prior to central venous cannulation. In patients receiving therapeutic anticoagulation with subcutaneous LMWH, the recommendation is to wait 18 hours after the last injection before performing nonurgent central venous cannulations. Similarly, nonurgent central venous cannulation in patients on warfarin should be withheld until the INR is less than 1.5.

The use of ultrasound for the placement of central venous catheters is well established and has been shown to increase success rates and decrease complication rates. The use of ultrasound in patients with disorders of hemostasis is safe and preferred when available. There are no data comparing ultrasound-guided versus traditional approach in the setting of anticoagulation.

In emergency situations site selection for central venous cannulation may improve outcomes. The safest technique in an anticoagulated patient appears to be an ultrasound-guided internal jugular approach. The lower risk of malposition, ease of ultrasound-guided technique, and the ability to manually compress a bleeding vessel in the event of a bleeding complication, makes the internal jugular access approach favored over the subclavian approach. Though the subclavian approach is associated with fewer arterial punctures, the provider is unable to manually compress bleeding vessels effectively. The femoral approach is also suitable for emergency central venous access because it is amenable to ultrasound-guided technique, and the provider is able to manually compress a bleeding vessel.
Importantly, femoral venous catheters are less favorable long-term because of an increased risk of infection. The use of intraosseous access is not contraindicated in anticoagulated patients. The manufacturer of the EZ-IO (Vida-Care, San Antonio, TX) device notes that applying pressure to the site for 1 to 2 minutes controls bleeding after needle removal, but more time may be required for patients on anticoagulant therapy.51

Patients requiring temporary placement of a hemodialysis catheter for emergency hemodialysis may have an increased risk of complications with central venous access as a result of comorbid conditions leading to anticoagulation or impaired hemostasis, particularly uremia-induced platelet dysfunction. If a patient requires emergency access for hemodialysis, the preferred approach site is the femoral vein because of the lower complication rate. Whereas complication rates are similar to typical central venous catheter placement, inadvertent dilation of the artery or unrecognized arterial puncture with a hemodialysis catheter can lead to massive bleeding. Although there are no data or guideline specifically targeted for the placement of hemodialysis catheters in the setting of anticoagulation, mechanical errors may be complicated by anticoagulation.

**Arthrocentesis**

Though numerous providers recommend reversing anticoagulation in patients receiving warfarin who require an arthrocentesis, there are little data to support this practice.52,53 In the largest study to date, Ahmed and Gertner evaluated the safety of arthrocentesis in patients receiving a VKA medication.11 Among the 456 procedures performed, there were no statistical differences in overall bleeding complications between patients with an INR greater than 2 and those with an INR less than 2. The authors concluded that arthrocentesis can be performed on patients who are anticoagulated without the need for reversal or discontinuation of anticoagulation prior to the procedure. Of note, 103 of the procedures were on patients with an INR greater than 3 and the highest INR was 7.8. In the latest study by Bashir and colleagues, 2084 knee and shoulder joint injections were performed on patients taking warfarin.12 The mean INR was 2.77 (range 1.7 to 5.5). Eighty-seven percent of the patients had an INR greater than or equal to 2. Nineteen- or 21-gauge needles were used for the arthrocentesis. There were no procedural complications noted in this study, leading the authors to conclude that joint injections are safe in the setting of therapeutic anticoagulation with warfarin. Additional studies have failed to demonstrate a higher risk of bleeding with arthrocentesis in the setting of anticoagulation.9,32 There appears to be no difference between the types of joint requiring arthrocentesis. Most studies have included both shoulder and knee joint aspiration.

There is no available literature evaluating the safety of arthrocentesis in patients receiving an antplatelet agent or a nonvitamin K anticoagulant. Hemarthrosis is a common presentation for patients with bleeding disorders such as hemophilia. It is typically recommended that patients or parents replace factors to achieve circulating factor activity of 40% to 50% at the first sign of acute bleeding episodes. Arthrocentesis is not always required to make the diagnosis of hemarthrosis in this clinical setting. However, if performing an arthrocentesis in the setting of hemophilia is required, the recommendation is to perform the procedure after appropriate factor replacement.

Spontaneous nontraumatic hemarthrosis is an infrequent complication with supratherapeutic anticoagulation and may even occur within the target therapeutic range.14 Routine coagulation studies are recommended if hemarthrosis is suspected in an anticoagulated patient. Temporary discontinuation of anticoagulants is typically recommended as part of treatment of hemarthrosis. If arthrocentesis is required, reversal of anticoagulation is not recommended if the INR is less than 4.5.7 Although arthrocentesis can safely be performed in the setting of supratherapeutic anticoagulation as previously mentioned, there are no safety data for the setting of hemarthrosis.11 Clinical judgement should be used to determine the necessity of this procedure.

**Paracentesis**

Paracentesis is a relatively safe procedure. The frequency of complications such as minor bleeding, or major complications such as abdominal hematoma that require a transfusion is low.55 In one report, abdominal hematomas occurred in only 1% of patients despite cirrhosis being a common comorbidity.62 There are currently no data to support the routine administration of FFP or platelets prior to paracentesis in patients with mild to moderate coagulopathy (PT or PTT up to twice normal) or thrombocytopenia (platelet count 50 to 99 × 109/L).55,57,58 However, caution should be exercised in patients with severe coagulopathy or DIC.55

In a 2009 prospective study evaluating the complication rate of paracentesis in cirrhotic patients, local bleeding occurred in 2.3% of cases and major bleeding requiring a medical or surgical intervention occurred in 1% of cases. Two of the five patients with major bleeding included a major hematoma, whereas three cases involved bleeding into the peritoneal cavity. A nonsignificant trend towards increased risk of complications was seen in patients with a platelet count less than 50 × 109/L. Therefore, the authors advised caution when performing paracentesis on patients with a platelet count greater than 50 × 109/L.59

In a 2011 retrospective study evaluating the safety of paracentesis in patients with cirrhosis, the authors concluded that the procedure was safe.60 Of 209 paracenteses reviewed for the study, 19% were performed on patients with a coagulopathy. The most common complication was local bleeding (3%). One patient experienced an abdominal hematoma, but no further data were given about this patient. The authors concluded that performing a paracentesis in the setting of coagulopathy is safe and the only absolute contraindications are DIC and fibrinolysis.

Although ultrasound-guidance technique has been shown to decrease the risk of bleeding with paracentesis in typical conditions, there are no data demonstrating this benefit in anticoagulated patients.61

**Thoracentesis and Tube Thoracostomy**

The major procedural complication associated with thoracentesis is pneumothorax, not bleeding.62 Numerous studies have failed to demonstrate an increased risk of bleeding with thoracentesis in the setting of anticoagulation.57,63,64

Additionally, the correction of abnormal coagulation laboratory values prior to the procedure is unlikely to have benefit. In a 2013 retrospective study of 1009 ultrasound-guided thoracenteses, Hibbert and colleagues evaluated patients treated with and without preprocedural transfusion of FFP or platelets for abnormal coagulation profiles.64 The authors
found no statistical difference in the bleeding complication rate (overall 0.4%) between those treated with transfusion prior to the procedure and patients not treated with blood products. In this study, the patients who did not receive FFP or platelet transfusions had a mean INR of 1.9 prior to the procedure. Seventy-four patients (14%) who were not treated with a transfusion had a platelet level of less than $50 \times 10^3/\text{L}$. There were no bleeding complications seen in the group that did not receive FFP or platelets compared with four bleeding complications that occurred in the group that received transfusions.

When performed under ultrasound guidance, the risk of bleeding is low even when coagulation studies or the platelet count are abnormal. However, there are no data available to demonstrate a lower bleeding complication rate of ultrasound-guided technique compared to the traditional approach in anticoagulated patients.

Recent studies have raised concerns about the risk of bleeding when thoracentesis is performed on patients taking clopidogrel. One prospective study of 25 consecutive patients noted a low rate of clinically significant hemorrhage in patients receiving thoracentesis while on clopidogrel, but further studies are required to determine safety. In that study, one patient developed a significant hemothorax requiring blood transfusion and tube thoracostomy. A 2013 case report also raised similar concerns as a patient on clopidogrel and aspirin developed a hemothorax after chest tube insertion. However, there are no data to support this recommendation. In fact, two studies failed to demonstrate a high risk for bleeding in patients taking clopidogrel especially when using small bore chest tubes.

In patients with a tension pneumothorax there are no absolute contraindications to tube thoracostomy. In stable patients undergoing chest tube placement, the need for anticoagulation reversal should be considered. Several clinicians have recommended that coagulopathies and platelet defects be corrected prior to tube thoracostomy.

**Dental Procedures**

Managing postoperative bleeding after a dental procedure can be a challenging task for EPs especially in the setting of anticoagulation. Dental procedures such as extractions are commonly performed without the cessation of anticoagulants. Multiple studies and authors have concluded that the risk of cessation of anticoagulation including antiplatelet therapy is much greater than the risk of significant bleeding with continued anticoagulation for dental procedures.

Multiple organizations including the AHA, ACCP, American College of Surgeons, American Dental Association, and The American College of Cardiology all agree that single or dual antiplatelet therapy should not be interrupted for dental procedures. To date there are no clinical trials demonstrating the safety of dental procedures in patients taking NOAC medications. Some have concluded that it appears safe to continue NOAC medications such as dabigatran or rivaroxaban for dental procedures based on management recommendations for warfarin and LMWH.

Bleeding and oozing from a dental procedural site is common after extractions such as wisdom tooth extractions even in the setting of normal coagulation. Patients commonly come to the ED after futile attempts to control the bleeding at home. Simple local therapy such as compression, packing, and vasoconstrictor infiltration is typically sufficient to control bleeding, even in the anticoagulated patient (see Chapter 72). Although none are specific for anticoagulated patients, there are multiple commercially available products that can be utilized to achieve hemostasis. Thrombin soaked gelatin sponges, oxidized cellulose material, chitosan coated gauze dressing, and topical tranexamic acid have all been described for this purpose.

Patients with hemophilia and vWD have an increased risk of bleeding during and after dental procedures. In addition to utilizing standard therapies, IV or local tranexamic acid and epsilon-aminocaproic acid in patients with hemophilia may also aid in achieving hemostasis, as has been described in a limited number of trials. However, a 2015 Cochrane review was unable to definitively conclude its efficacy.

Although no studies exist looking specifically at the safety of performing simple oral and dental procedures in the ED such as incision and drainage of a dental abscess, they are likely benign, considering the safety data for more invasive dental procedures such as extraction.

**Epistaxis**

Epistaxis is a common presenting symptom in the ED and may be complicated by anticoagulant use. Whether the bleeding is the result of excessive anticoagulation, achieving hemostasis may be more difficult than normal clinical scenarios. Evaluation of coagulation parameters (PT, INR, PTT, platelet count, or other anticoagulant-specific laboratory assays) should be routinely ordered for anticoagulated patients presenting with epistaxis. A complete blood count is also suggested for patients with prolonged or severe bleeding with epistaxis.

Anticoagulant reversal is rarely necessary unless laboratory findings reveal a markedly abnormal degree of anticoagulation or the patient has severe, symptomatic, or life-threatening bleeding. Specific factor replacement is necessary for patients with hemophilia and severe bleeding.

A variety of topical hemostatic agents are available and can be applied to the nasal cavity for the management of epistaxis in the anticoagulated patient when standard therapy is not adequate. Cellulose, gelatin, and thrombin compounds can be placed directly on the bleeding site to promote clot formation even in fully anticoagulated patients. Both topical and IV use of tranexamic acid has also been described in the management of epistaxis, but there is a lack of evidence-based data supporting its efficacy. Recently a case report described the successful use of topical tranexamic acid in the management of rivaroxaban associated epistaxis after failure with a thrombin-soaked inflated nasal tampon.

**Nasogastric Tube Insertion**

There is a lack of information available about the safety of placing a nasogastric tube (NGT) in the setting of anticoagulation. Severe coagulopathy is a commonly stated relative contraindication of NGT passage for risk of epistaxis without reference to evidence-based data.

Similarly, the often stated NGT placement contraindication of esophageal varices is unproven. Although mechanical or chemical irritation is thought to cause esophageal varices rupture and bleeding, there are no evidence-based data that stratifies the risk of NGT placement in this clinical setting. In an anesthesia study of patients undergoing hepatic transplantation with esophageal varices, 0 of 73 patients developed bleeding after NGT placement. This further calls into question the link between NGT placement and causation of bleeding esophageal varices. Additional research is required to prove or disprove this link.
Thrombocytopenia is a common finding in cirrhosis and is a poor indicator for risk of future gastrointestinal hemorrhage. Furthermore, routine coagulation assays are not reliable indicators of coagulation status in patients with cirrhosis. Whereas platelet transfusion is commonly recommended prior to endoscopy for a platelet count of less than 30 × 10^9/L, there is no analogous recommendation for NGT insertion.

**Labor and Delivery**

Delivery of a newborn in the ED is a relatively rare event. Postpartum bleeding and hemorrhage is a complication exacerbated by maternal factors such as anticoagulation and bleeding disorders. Pregnant patients diagnosed with venous thromboembolism are typically treated with LMWH. In the final weeks of pregnancy obstetricians occasionally switch patients to unfractionated heparin to reduce time to normalization of coagulation studies when labor begins. However, fully anticoagulated patients may present with precipitous labor and develop postpartum hemorrhage. Reversal of anticoagulation may be necessary in the event of refractory bleeding. There is no recommendation for prophylactic reversal of anticoagulation for delivery.

Patients with vWD are at increased risk for postpartum hemorrhage. These patients have a higher risk of delayed postpartum hemorrhage reported up to 2 to 3 weeks after delivery. According to case reports, postpartum hemorrhage may still occur despite prophylaxis with factor VIII, cryoprecipitate, FFP, and DDAVP. Expert opinion recommends that a von Willebrand factor activity of greater than or equal to 50 IU/dL should be achieved before delivery and maintained for at least 3 to 5 days afterward. Prophylaxis with DDAVP or von Willebrand factor concentrate should be given prior to childbirth if time permits. However, there are no randomized trials that have studied DDAVP’s efficacy to prevent or treat postpartum hemorrhage.

Although a rare occurrence because of its X-linked nature, replacement of factors is recommended for pregnant women with severe hemophilia A or B in labor. Risk for bleeding begins when levels are below 30% of normal. Risk increases with severity of disease especially those with less than 1% normal factor activity.

**Wound Management**

**Control of Hemorrhage**

Hemostasis is an essential step in wound management. Inadequate hemostasis can lead to the formation of a hematoma within a closed wound, which may cause dehiscence of wound edges, impaired healing, and increased risk for wound infection. Anticoagulated patients may come to the ED for prolonged or severe episodes of bleeding from an acute traumatic wound. Most bleeding wounds are effectively managed with routine application of direct pressure and elevation. However, anticoagulants may impair the integral mechanisms of hemostasis and prevent adequate spontaneous hemostasis.

There are multiple techniques that can be utilized to control a bleeding wound including compression, vessel crushing, ligation, electrocautery, or application of a vasoconstrictor or hemostatic agent. These traditional techniques are not specific to anticoagulated patients, but they may be useful in achieving hemostasis. There are a few topical hemostatic agents that have been shown effective in the setting of anticoagulation.

Chitosan is a complex carbohydrate derived from chitin that interacts with platelets and red blood cells to form a gel-like clot independent from coagulation factors. In a heparinized swine model, Millner showed chitosan granules (Omni-Stat, Medtrade Products Ltd., Crewe, United Kingdom) and dressings (Celox Gauze, Medtrade) were both efficacious in providing hemostasis over plain gauze compression. The long-term stability of the clot is not well-established and therefore these products are used as temporary management if definitive therapy such as surgical intervention is required. Chitosan-based dressings (Hemcon, Tricol Biomedical, Portland, OR) have also been used externally for hemostasis in pediatric patients with bleeding tendencies from congenital or acquired bleeding disorders.

Topical thrombin has also shown efficacy as a topical hemostatic agent in anticoagulated patients, in particular heparin and clopidogrel therapy. These commercially available topical thrombin products aid in the production of fibrin in one of the final steps of the coagulation pathway to create a stable clot.

IV tranexamic acid has been shown to be efficacious in trauma patients especially when given early after injury. However, anticoagulated patients were not included in the study. The use of topical tranexamic acid for the control of bleeding has been promising in many clinical scenarios but remains unproven especially in the setting of anticoagulation.

**Incision and Drainage**

There is no literature reporting complications in anticoagulated patients who undergo incision and drainage. Although bleeding is a risk with incision and drainage, local therapy alone is typically sufficient to provide hemostasis. EPs should make a case-based clinical decision weighing the risk of hemorrhage for patients requiring incision and drainage.

**SUMMARY**

Procedures in the setting of anticoagulation present a unique complexity in the ED. The EP should be mindful of potential bleeding complications and should be prepared to identify and treat these events. Reversal of anticoagulation is not always necessary prior to performing a procedure and the EP should be attentive to the risk of thrombosis. In some instances, the risk-benefit ratio may favor delaying a procedure until the patient’s coagulation status returns to normal range with time, or the use of medications or blood products. Lastly, emergency procedures to correct an imminent life-threatening life-saving condition should never be withheld in the setting of anticoagulation.

**REFERENCES ARE AVAILABLE AT www.expertconsult.com**
REFERENCES


63. Patel MD, Joshi SD: Abnormal preprocedural international normalized ratio and platelet counts are not associated with increased bleeding complications after ultrasound-guided thoracentesis. AJR Am J Roentgenol 197:164, 2011.


INTRODUCTION

This appendix presents a list of calculations commonly used by emergency physicians. It is not all-inclusive and purposely does not include decision tools or scales that do not require algebraic calculations. Many of these equations can be found in online calculators on websites such as https://www.mdcalc.com or http://reference.medscape.com/guide/medical-calculators.

ENGLISH-TO-METRIC CONVERSIONS

Patients frequently express common figures such as weight, temperature, and volume in standard measurements. Please see Table A.1 for conversion factors and simple examples.

MEAN ARTERIAL PRESSURE

Calculation of mean arterial pressure (MAP) provides a weighted average of systolic blood pressure (SBP) and diastolic blood pressure (DBP). It is a determination of tissue perfusion pressure and is normally 70 to 100 mm Hg in adults. To determine MAP:

\[
\text{MAP} = \frac{\text{SBP} + (2 \times \text{DBP})}{3}
\]

Example: A hypertensive emergency is diagnosed in an elderly, hypertensive patient. Current recommendations are to reduce MAP by 10% to 20% in the first hour. His blood pressure is 240/120. To calculate the current MAP,

\[
\text{MAP} = \frac{240 + (2 \times 120)}{3} = 160 \text{ mm Hg}
\]

QT AND QTc INTERVALS

The QT interval on the electrocardiogram (ECG) represents the period of ventricular electrical activity from activation to repolarization. The most important determinant of the QT interval is the heart rate. As the heart rate increases, the QT interval shortens. Many ECG machines calculate this; however, the presence of U waves or other ECG abnormalities can result in inappropriate readings. To calculate the rate-corrected QT interval (QTc) using Bazett's formula, divide the QT interval by the square root of the R-R interval (the interval between the R wave on two consecutive QRS complexes). The R-R interval may be measured from the ECG or calculated as 60 divided by the heart rate (in beats/min). The interval is represented in seconds or milliseconds:

\[
\text{QTc} = \frac{\text{QT}}{\sqrt{\text{R-R}}}
\]

The QTc is normally less than 0.46 second in men and 0.44 second in women. This may be calculated online at: http://www.medical-calculator.nl/calculator/QTc/.

The list of drugs that can cause QTc prolongation is quite lengthy. The clinical significance of QTc prolongation is often unclear. The following Internet websites offer up-to-date information on this ever-changing topic:

https://crediblemeds.org/
http://www.ncbi.nlm.nih.gov/pmc/articles/PMC1767957/.

Table A.2 shows the normal range of the QT interval in adults.

Example: A 21-year-old man ingested a large quantity of amitriptyline (tricyclic antidepressant) tablets. His ECG revealed a QT interval of 0.37 second and a heart rate of 120 beats/min. The QTc is calculated as:

\[
\text{R-R interval} = 60/120 = 0.50
\]

\[
\text{QTc} = 0.37/\sqrt{0.5} = 0.523 = 523 \text{ msec}
\]

The patient's QTc is significantly prolonged for his heart rate and indicates significant cardiac effects from overdose of a tricyclic antidepressant.

PREDICTED PEAK EXPIRATORY FLOW RATE

The peak expiratory flow rate (PEFR), measured in liters per minute, is a useful means of assessing airway obstruction. It is measured by having a patient exhale maximally through a peak-flow meter. Normal values range from 350 to 600 L/min. Comparison between the initial and posttreatment PEFR in patients with exacerbations of asthma helps determine the degree of severity. Patients with an initial PEFR of less than 20% of predicted or with a subsequent value of less than 60% of predicted after initial therapy may require further evaluation, treatment, or both. Many patients monitor PEFR on themselves and are able to state a personal best, which is the preferred standard for that individual. Other patients may be monitored with an estimated PEFR. Estimations are based primarily on a patient's gender, age, and height. Although graphs and tables are available to provide values across a range of ages and heights, the PEFR can also be approximated by using the following formulas:

Adults: \[\text{PEFR} = 13 \times (\text{Height (in)} - 40) + 110\]

Female children/adolescents:

\[\text{PEFR} = \text{Height (m)} \times 5.5 - \text{Age} \times 0.03 - 1.11\]

Male children/adolescents:

\[\text{PEFR} = \text{Height (m)} \times 6.14 - \text{Age} \times 10.043 + 0.15\]

These may also be found online in many places including: http://www.mdcalc.com/estimatedexpected-peak-expiratory-flow-peak-flow/.

ENDOTRACHEAL TUBE SIZE

Adults

Select the largest-diameter endotracheal tube (ETT) that can be tolerated for adults. A 7.5-mm cuffed ETT is well tolerated.
Appendix

COMMONLY USED FORMULAS AND CALCULATIONS

**TABLE A.1** Standard-to-Metric Calculations

<table>
<thead>
<tr>
<th>Fahrenheit to Celsius</th>
<th>A patient’s temperature is 100.4°F &lt;br&gt;( (^\circ F - 32)/1.8 = 38^\circ C )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pounds to kilograms</td>
<td>A patient reports his weight to be 154 lb &lt;br&gt;( 154 \text{ lb}/2.2 = 70 \text{ kg} )</td>
</tr>
<tr>
<td>Inches to centimeters</td>
<td>The patient’s height is 72 inches &lt;br&gt;( 72 \text{ inches} \times 2.54 = 182.9 \text{ cm} )</td>
</tr>
<tr>
<td>Fluid ounces to milliliters</td>
<td>The child drinks 4 oz of formula at a sitting &lt;br&gt;( 4 \text{ oz} \times 30 = 120 \text{ mL} )</td>
</tr>
</tbody>
</table>

**TABLE A.2** Normal Range of the QT Interval in Adults

<table>
<thead>
<tr>
<th>HEART RATE (beats/min)</th>
<th>NORMAL QT RANGE (sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>40</td>
<td>0.42–0.53</td>
</tr>
<tr>
<td>50</td>
<td>0.37–0.48</td>
</tr>
<tr>
<td>60</td>
<td>0.34–0.44</td>
</tr>
<tr>
<td>70</td>
<td>0.31–0.41</td>
</tr>
<tr>
<td>80</td>
<td>0.29–0.38</td>
</tr>
<tr>
<td>90</td>
<td>0.28–0.36</td>
</tr>
<tr>
<td>100</td>
<td>0.27–0.34</td>
</tr>
<tr>
<td>110</td>
<td>0.25–0.32</td>
</tr>
<tr>
<td>120</td>
<td>0.24–0.31</td>
</tr>
<tr>
<td>130</td>
<td>0.23–0.30</td>
</tr>
<tr>
<td>140</td>
<td>0.22–0.29</td>
</tr>
<tr>
<td>150</td>
<td>0.21–0.28</td>
</tr>
</tbody>
</table>

by most adult female patients. An 8.0-mm cuffed ETT is well tolerated by most adult male patients.

**Pediatrics**

An uncuffed ETT should be used for children younger than 8 years. A number of techniques are available for estimating the appropriate size of ETT in children. Commonly used formulas for estimating tube size and depth of insertion use age in years and are as follows:

\[
\text{ETT size} = \frac{(\text{Age}+16)}{4} \text{ or } \frac{\text{Age}}{4}+4
\]

To estimate depth of insertion for a child older than 2 years:

\[
\text{Depth of insertion} = 3 \times \text{Internal diameter of the ETT}
\]

**VENTILATOR SETTINGS**

The recommended initial ventilator settings follow. Adjustments in these ventilator settings may be made according to the patient’s clinical situation:

\[
\text{Tidal volume (V}_T\text{) } = 6–12 \text{ mL/kg}
\]

\( V_T \) is often based on ideal body weight, and there is greater recognition that a lower \( V_T \) may be beneficial, such as 5 to 10 mL/kg. (Adjust \( V_T \) to limit inflation or plateau pressures to 30 cm H\text{}_2\text{O} or lower. Lower tidal volumes are recommended to limit ventilator-induced lung injury.)

\[
\text{Rate } = 10 \text{ to } 12 \text{ breaths/min for adults, } 16 \text{ to } 20 \text{ breaths/min for children, and } 20 \text{ to } 30 \text{ breaths/min for infants.}
\]

Fi\text{O}_2\ (\text{fraction of inspired oxygen}) = 50\% \text{ to } 100\% \text{ initially; reduce Fi\text{O}_2\ as quickly as possible to avoid oxygen toxicity.}

Inspiratory-to-expiratory (I/E) ratio = 1:2. To allow complete exhalation, the I/E ratio should be at least 1:2.

\[
\text{Minute ventilation (V}_E\text{) } = \text{Rate} \times V_T
\]

\( V_E \) will vary with the patient’s mass and disease process; the normal adult range is approximately 7 to 10 L/min.

**Example:** A 6-year-old girl with asthma has respiratory distress and altered mental status and requires endotracheal intubation. Her weight is 20 kg. To prepare for intubation and mechanical ventilation, use the following equipment and settings:

\[
\begin{align*}
\text{ETT size (mm): } & (6+16)/4 = 5.5\text{-mm ETT (uncuffed).} \\
\text{Depth of insertion: } & 3 \times 5.5 = 16.5 \text{ cm.} \\
\text{V}_T & = 12 \text{ mL/kg} \times 20 \text{ kg} = 240 \text{ mL.} \\
\text{Respiratory rate} & = 16 \text{ breaths/min.} \\
\text{FiO}_2 & = 100\%. \\
\text{I/E ratio} & = 1:2.
\end{align*}
\]

**IDEAL BODY WEIGHT**

Ideal body weight (IBW) is used in many applications including mechanical ventilator settings, drug dosing, and renal function measurement. Many equations have been described but the Devine formula is the most widely used. This formula, however, cannot be used for patients less than 5 feet (60 in) tall.

\[
\text{IBW (male) } = 50 \text{ kg} + [2.3 \times (\text{height in in}) – 60] \\
\text{IBW (female) } = 45.5 \text{ kg} + [2.3 \times (\text{height in in}) – 60]
\]

Calculate online at: http://www.mdcalc.com/ideal-body-weight/.

**Example:** A 55-year-old female presents with an asthma exacerbation and is intubated. She has a weight of 112 kg and body mass index of 40 and so IBW is used in calculating tidal volume. Her height is 66 inches.

\[
\text{IBW } = 45.5 \text{ kg} + [2.3 \times (66 \text{ in} – 60)] = 59.3 \text{ kg}
\]
**ABSOLUTE NEUTROPHIL COUNT**

The absolute neutrophil count (ANC) is the total number of neutrophils in the blood, both mature and immature. The value represents the actual number of neutrophils as compared to a percentage of white blood cells (WBCs). It can be an important measure in an immunocompromised patient as it can help to assess a patient's risk for developing opportunistic infections and guide treatment. Immunocompromised patients who are neutropenic and present with fever are at high risk of developing sepsis. It is calculated as:

$$\text{ANC} = \frac{\text{(% neutrophils} + \text{% bands})/100}{\text{WBC count}}$$

- Normal: > 1500 cells/µL
- Mild neutropenia: 1000 to 1500 cells/µL
- Moderate neutropenia: 500 to 1000 cells/µL
- Severe neutropenia: < 500 cells/µL

**WHITE BLOOD CELL CORRECTION IN CEREBROSPINAL FLUID**

When performing a lumbar puncture on a patient to evaluate for meningitis, the number of WBCs in the cerebrospinal fluid (CSF) is an important value for diagnosis. However, a traumatic lumbar puncture can falsely elevate this value. A formula using the WBCs and red blood cells (RBCs) in CSF and in blood can correct the WBCs in CSF. The normal value for WBCs and RBCs in CSF is < 5 cells/µL. All values are in cells/µL except RBCs in blood, which is cells × 10^6/µL.

$$\text{Adjusted WBC}_{\text{CSF}} = \frac{(\text{WBC}_{\text{blood}} \times \text{RBC}_{\text{CSF}})}{(\text{RBC}_{\text{blood}} \times 10^6)}$$

**Example:** A 25-year-old male presents with fever, sore throat, and headache. Meningitis is suspected. His blood shows elevated WBC at 15,000 cells/µL. A lumbar puncture (LP) is completed and results show elevated WBCs of 25 cells/µL. The rest of his CSF values are unremarkable. His RBC in blood is 5 × 10^6/µL.

$$\text{Adjusted WBC in CSF} = 25 - [(15,000 \times 7000)/5 \times 10^6]$$

Adjusted WBC in CSF = 4 cells/µL

His WBCs in CSF are actually within normal range and the elevation is likely due to a traumatic LP.

**CREATININE CLEARANCE**

Creatinine clearance (CrCl) occasionally needs to be calculated by emergency physicians to risk-stratify patients undergoing imaging studies requiring intravenous (IV) contrast material, as well as to determine the severity of renal failure in patients without a known baseline. CrCl is best calculated from a collection of urine over a 24-hour period. However, if the patient’s CrCl is in steady state (i.e., without recent change), it is possible to estimate CrCl by using a formula that incorporates serum creatinine (Scr) in mg/dL, lean body weight (LBW) in kg, age in years, and gender:

$$\text{CrCl} = \frac{[(140 - \text{Age}) \times \text{LBW}]}{[72 \times \text{Scr}]}$$

- Normal values: 74 to 160 mL/min.
- Mild renal impairment: 40 to 60 mL/min.
- Moderate renal impairment: 15 to 40 mL/min.
- Severe renal impairment: < 15 mL/min (indication for renal dialysis).

The following examples illustrate the importance of calculating CrCl:

**Example 1:** A 71-year-old woman has upper abdominal tenderness. A computed tomography (CT) scan is planned, but the clinician is concerned about the risk for contrast-induced nephropathy. The patient’s Scr is 1.4 mg/dL and she weighs 55 kg. To calculate her CrCl:

$$\text{CrCl} = \frac{[(140 - 71) \times (55 \times 10^3)]}{[72 \times 1.4]}$$

$$\text{CrCl} = 32.0 \text{ mL/min}$$

The patient has moderate renal impairment.

**Example 2:** A 21-year-old man has right lower quadrant tenderness. A CT scan is planned, but the radiologist is concerned about the risk for contrast-induced nephropathy. The patient’s Scr is 1.4 mg/dL, and he has an LBW of 100 kg. To calculate his CrCl:

$$\text{CrCl} = \frac{[(140 - 21) \times (100 \times 10^3)]}{[72 \times 1.4]} = 118 \text{ mL/min}$$

The patient has adequate CrCl. Note that despite an identical serum creatinine level, the second patient has normal renal function, whereas the first has renal insufficiency.

**FRACTIONAL EXCRETION OF SODIUM**

Acute kidney injury (AKI) can have many etiologies. Calculating the fractional excretion of sodium (FENa) can help to differentiate prerenal disease, intrinsic renal disease, and postrenal disease. The calculation measures the percent of filtered sodium that is excreted in the urine. Low values of FENa, less than 1%, in the setting of AKI suggests a prerenal etiology as the kidneys are appropriately retaining sodium in response to decreased perfusion. Higher values of FENa, greater than 2%, suggests an intrinsic etiology of AKI. The higher values can be caused by either salt wasting by damaged nephrons or an appropriate response by remaining nephrons to volume expansion. FENa greater than 4% can be suggestive of a postrenal cause of AKI. To calculate FENa, a serum creatinine (Scr), serum sodium (SNa), urine creatinine (Ucr), and urine sodium (Una) need to be measured.

$$\text{FENa} = \frac{(\text{Scr} \times \text{Ucr})}{(\text{SNa} \times \text{Una})} \times 100$$

**Example:** A 65-year-old previously healthy male had a syncopal episode while working outside on a hot day. He was found to have AKI with an Scr of 2.0 mg/dL. His SNa is 150 mEq/L, UNa is 18 mEq/L, and UCr is 30 mg/dL.

$$\text{FENa} = \frac{(2.0 \times 18)}{(150 \times 30)} \times 100$$

$$\text{FENa} = 0.8\%$$

The patient’s FENa is less than 1% which suggests a prerenal cause, likely hypovolemia.
HYPERNATREMIA

Elevation of the serum sodium concentration is proportionate to the free water deficit when volume is depleted. Because 60% of the adult body is water, the total-body free water deficit is calculated by using measured Na⁺, desired Na⁺, and body weight in kilograms. To calculate the free water deficit:

\[
\text{Ideal total body water (TBW) = 60\% \times \text{weight (kg)}}
\]

\[
\text{TBW deficit = } \frac{[\text{Measured Na⁺} - \text{Normal Na⁺}]}{\text{Normal Na⁺}} \times \text{TBW}
\]

(Assume normal Na⁺ to be 140 mmol/L.)

Example: An elderly man is brought to the ED in a coma. He has severe dehydration, his IBW is 70 kg, and his serum Na⁺ level is 165 mmol/L. To determine his free water deficit:

\[
\text{Ideal TBW = 0.6} \times 70 \text{ kg = 42 L}
\]

Free water deficit = \[(165 - 140)/140\] \times 42 = 7.5 L

Fluid correction for hypernatremia should take place over a 48- to 72-hour period to avoid the potential for cerebral edema. Calculate free water deficit and fluid rate online: http://www.medcalc.com/sodium.html.

POTASSIUM

Serum potassium (K⁺) levels change with acid-base status. In acidic states, K⁺ moves out of cells as H⁺ moves in, thus raising serum K⁺ levels. In alkalotic states, K⁺ moves into cells as H⁺ moves out, thus lowering serum K⁺ levels. The change in K⁺ varies inversely with pH at the following rates:

The serum K⁺ concentration increases 0.6 mmol/L for each 0.1-unit decrease in pH.

The serum K⁺ concentration decreases 0.6 mmol/L for each 0.1-unit increase in pH.

CALCIUM

Approximately 50% of serum calcium is bound to serum proteins (primarily albumin), 40% is in the free ionized state (the physiologically active form), and 10% is mixed with serum anions (phosphate, bicarbonate, citrate, and lactate). For this reason, serum calcium is lowered approximately 0.8 mg/dL for every decrease in albumin of 1 g/dL. To correct for decreased albumin (at levels < 4 g/dL), the following formula can be used:

\[
\text{Corrected Ca^{2+} (mg/dL) = Serum Ca^{2+} (mg/dL) + (0.8 \times (4.0 - \text{serum albumin} [g/dL]))}
\]

Example: A malnourished man has a serum calcium level of 7.5 mg/dL and a serum albumin level of 2 g/dL. To calculate his corrected calcium level:

\[
\text{Corrected Ca^{2+} (mg/dL) = 7.5 mg/dL + (0.8 \times (4.0 - 2.0 [g/dL]) = 9.1
}\]

MAINTENANCE IV FLUID RATE

To calculate maintenance IV fluids for a pediatric patient, use the following formula:

4 mL/kg per hr for the first 10 kg, plus
2 mL/kg per hr for the second 10 kg, plus
1 mL/kg per hr for each further kg.
The major positive charge in serum. An elevated AG usually means that there is some unmeasured anion, toxin, or organic acid in the blood. **Box A.1** lists many substances that can cause an AG acidosis. The AG is normally 8 to 12 mmol/L:

$$\text{AG} = \text{Na}^+ - (\text{Cl}^- + \text{HCO}_3^-) = 8 - 12 \text{ mmol/L}$$

**Example:** A suicidal young male drank an unknown amount of antifreeze. His electrolyte levels are Na$^+$, 144; K$^+$, 3.1; Cl$^-$, 108; and HCO$_3^-$, 14. The AG is calculated as follows:

$$\text{AG} = 144 - 108 - 14 = 22 \text{ mmol/L}$$

**Example:** An elevated AG presumably because of ingestion of ethylene glycol.

**Fluid Resuscitation of Burned Patients**

Various formulas for IV fluid resuscitation in patients with burns have been recommended. The Parkland formula is commonly used and is calculated as follows:

$$\text{Replacement fluid} = 4 \text{ mL} \times \text{Weight [kg]} \times \frac{(% \text{Body surface area [BSA] burned})}{2}$$

Count only second- and third-degree burns in calculating BSA. The total volume should be administered in the first 24 hours with half the fluid given in the first 8 hours and the remaining half in the next 16 hours. Clinical parameters, including urine output, vital signs, and central venous pressure or pulmonary capillary wedge pressure, should be monitored carefully to assess the adequacy of resuscitation.

**Example:** A 65-kg woman has second- and third-degree burns covering 35% of her BSA. To determine her anticipated 24-hour fluid resuscitation needs:

- Replacement fluid = $4 \times 65 \times 0.35 = 9100 \text{ mL}$
- Half is given in the first 8 hours: $9100 / 2 = 4550 \text{ mL}$
- $4550 \text{ mL/hr} = 569 \text{ mL/hr}$ for the first 8 hours
- Half is given in the second 16 hours: $284 \text{ mL/hr}$

Do not forget to add maintenance fluid rates to the results of the Parkland formula. Most online calculators do not include this!

**Anion Gap**

The anion gap (AG) is an estimate of the amount of unmeasured negatively charged ions in serum that are not bicarbonate (HCO$_3^-$) and chloride (Cl$^-$). The AG is calculated by subtracting the sum of HCO$_3^-$ and Cl$^-$ values from sodium (Na$^+$), which

<table>
<thead>
<tr>
<th>EACH mg/dL OF</th>
<th>INCREASES SERUM mOsm/kg BY</th>
<th>FOR EACH SERUM mOsm/kg INCREASE DUE TO</th>
<th>THE CORRESPONDING mg/dL CHANGE IS (= Mol Wt/10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methanol</td>
<td>0.31</td>
<td>Methanol</td>
<td>3.2</td>
</tr>
<tr>
<td>Ethanol</td>
<td>0.22</td>
<td>Ethanol</td>
<td>4.6</td>
</tr>
<tr>
<td>Acetone</td>
<td>0.17</td>
<td>Acetone</td>
<td>5.8</td>
</tr>
<tr>
<td>Isopropyl alcohol</td>
<td>0.17</td>
<td>Isopropyl alcohol</td>
<td>6.0</td>
</tr>
<tr>
<td>Ethylene glycol</td>
<td>0.16</td>
<td>Ethylene glycol</td>
<td>6.2</td>
</tr>
<tr>
<td>Glycerol</td>
<td>0.11</td>
<td>Glycerol</td>
<td>9.2</td>
</tr>
<tr>
<td>Mannitol</td>
<td>0.05</td>
<td>Mannitol</td>
<td>18.2</td>
</tr>
</tbody>
</table>

ACID-BASE BALANCE

A thorough understanding of blood gas interpretations is crucial for critically ill patients but is beyond the reach of this section. Unfortunately, most online calculators either struggle with mixed acid-base disorders, or cannot handle different units (kPa vs. mm Hg).

By using the combination of an arterial blood gas (ABG) sample and serum electrolyte levels, a patient’s acid-base status can be evaluated. A single acid-base disorder may be present as a primary disorder (respiratory acidosis, respiratory alkalosis, metabolic acidosis, or metabolic alkalosis), or several disorders may be present at the same time. When an acid-base disorder is present, the body will try to compensate to preserve the pH as close to normal as possible. The compensatory responses in relation to the primary disorder are as follows:

<table>
<thead>
<tr>
<th>Primary Disorder</th>
<th>Compensatory Response</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metabolic acidosis</td>
<td>Increased ventilation</td>
</tr>
<tr>
<td>Metabolic alkalosis</td>
<td>Decreased ventilation</td>
</tr>
<tr>
<td>Respiratory acidosis</td>
<td>Increased renal reabsorption of HCO₃⁻ in the proximal tubule</td>
</tr>
<tr>
<td></td>
<td>Increased renal excretion of H⁺ in the distal tubule</td>
</tr>
<tr>
<td>Respiratory alkalosis</td>
<td>Decreased renal reabsorption of HCO₃⁻ in the proximal tubule</td>
</tr>
<tr>
<td></td>
<td>Decreased renal excretion of H⁺ in the distal tubule</td>
</tr>
</tbody>
</table>

When interpreting acid-base disorders, the following basic steps are suggested:

Knowledge of normal values:

- Normal serum pH = 7.40
- Normal partial pressure of carbon dioxide (Pco₂) = 40 mm Hg

First, determine the patient’s pH status. Acid-base changes are either metabolic or respiratory. Simple disturbances are categorized by examining pH, Pco₂, and HCO₃⁻ (Table A.4).

If pH is less than normal (< 7.35), the patient is acidemic. If pH is above normal (> 7.45), the patient is alkalemic. If the patient is acidemic and Pco₂ is decreased (> 45 mm Hg) as a primary disorder, respiratory acidosis is present. If the patient is alkalemic and Pco₂ is decreased (< 35 mm Hg) as a primary disorder, respiratory alkalosis is present. If the patient is acidemic and the arterial HCO₃⁻ level is less than normal (< 22 mEq/L) as a primary disorder, metabolic acidosis is present. If the patient is alkalemic and the HCO₃⁻ level is greater than normal (> 26 mEq/L) as a primary disorder, metabolic alkalosis is present.

Acid-base homeostasis is normally maintained because a change in pH will usually trigger a compensatory change to minimize the change in pH, although the compensation is never complete. The degree and timing of compensation are determined by the primary disturbance itself and by individual physiology. Respiratory compensation for metabolic disorders is more rapid and occurs through a change in the respiratory rate, which in turn adjusts the Pco₂. Metabolic compensation for a respiratory disturbance requires renal adjustment of HCO₃⁻ and can take 3 to 5 days. The predicted compensatory responses for the primary disturbances are shown in Table A.4. It is important to remember that physiologic compensatory mechanisms may themselves be compromised or overwhelmed by the acid-base disorder (Fig. A.1).

Example 1: A 58-year-old woman has had profuse diarrhea for 1 week. Initial laboratory data include the following: Na⁺, 133; K⁺, 2.8; pH 7.26; Cl⁻, 118; Pco₂, 13; HCO₃⁻, 5.

1. Acidity is present (pH < 7.40).
2. The primary process is metabolic (HCO₃⁻ < 22 mEq/L, and Pco₂ is not increased).
3. Compensation. In primary metabolic acidosis, the formula that checks for compensation is:

\[ \Delta Pco₂ \times 1.3 \times \Delta HCO₃⁻ \]

(where \( \Delta HCO₃⁻ \) is normal – current values)

\[ \Delta Pco₂ = 1.3 \times (25 - 5) = 26 \text{ mm Hg} \]

The predicted Pco₂ is the normal Pco₂ – ΔPco₂

\[ = 40 - 26 = 14 \]

The actual Pco₂ is 13 mm Hg.
The respiratory compensation is predicted by the equation
and suggests that no other acid-base disorder is present.

**Example 2:** A 74-year-old nursing home resident is admitted
to the hospital with hypotension (96/70) and fever (39°C). He
has had a positive urine culture for *Escherichia coli* and two
positive blood cultures with the same organism. His laboratory
values are as follows:

- Sodium, 138 mmol/L.
- Potassium, 3.2 mmol/L.
- Chloride, 105 mmol/L.
- pH, 7.49.
- Pco₂, 25 mm Hg.
- HCO₃⁻, 22 mmol/L.

1. Alkalosis is present (pH > 7.44).
2. The primary process is respiratory (Pco₂ < 40 mm Hg and
HCO₃⁻ was not increased).
3. Compensation. The decrease in Pco₂ is 40 - 25, or
15 mm Hg. The formula for an expected decrease in HCO₃⁻
is 2 mmol for every 10–mm Hg decrease in Pco₂. In this instance the expected decrease in HCO₃⁻ for the 15–mm Hg decrease in Pco₂ is 3 mmol/L, which is nearly identical to the actual decrease (i.e., 25 – 22). Therefore only acute respiratory alkalosis is present with normal compensation.

**WINTER’S FORMULA**

In a pure primary metabolic acidosis, the body's normal response is a change (decrease) in P CO₂. The expected compensation can be predicted by Winter's formula:

\[
\text{Expected P CO}_2 = 1.5 \times \text{HCO}_3^- + 8 + 2
\]

**Example:** HCO₃⁻ calculated from ABG analysis is 10 mEq/L. With pure metabolic acidosis the expected P CO₂ is:

\[
\text{Expected P CO}_2 = 1.5 \times 10 + 8
\]

Expected P CO₂ = 23 mm Hg (range, 21 – 25 mm Hg)

If the measured P CO₂ is higher than the expected P CO₂, a concomitant respiratory acidosis is also present. Normocapnia or hypercapnia in the presence of severe metabolic acidosis may be a harbinger of impending respiratory failure and suggests the possible need for mechanical ventilation.

A lower P CO₂ would suggest a concomitant respiratory alkalosis, such as seen with salicylate poisoning.

Another useful tool in estimating P CO₂ in metabolic acidosis is the recognition that P CO₂ is approximately equal to the last two digits of the pH. In the previous example, the expected pH should be 7.23.

**GLASGOW COMA SCALE**

For head-injured patients, the Glasgow Coma Scale (GCS) is a frequently referenced assessment of neurologic function. It is calculated by using the best result of neurologic testing of eye opening, motor function, and verbal function. Each of the three subscores is determined by using verbal and painful stimuli. Point assessments are listed in Table A.5, with adjustments noted for pediatric patients. Points are given for the best response observed for each category. For example, a person with right-sided motor deficits but otherwise normal function would have a normal GCS score because left-sided function is normal. Scores of 13 to 15 correlate with mild brain injury, 9 to 12 with moderate brain injury, and 3 to 8 with severe brain injury.

**Example:** An intoxicated male is involved in a motor vehicle crash and is brought to your ED. He opens his eyes only to painful stimuli, is muttering incomprehensibly, and localizes pain on the left and withdraws from pain on the right. What is his GCS score, and what severity of injury does this score represent?

GCS = Eye opening + Verbal response + Motor response

Eye opening to pain = 2 points
Incomprehensible sounds = 2 points
Localizing pain is 5 points and withdrawal from pain is 4 points. The better score is used.

\[2 + 2 + 5 = 9 \text{ points}\]

**NATIONAL INSTITUTES OF HEALTH STROKE SCORE**

The full National Institutes of Health (NIH) stroke scale is shown in Table A.6. The scale includes directions for patient assessment, as well as recommended adjustments for special situations that might impair the ability of the patient to respond to the assessor. Training scenarios and the NIH stroke scale can be found at [http://www.nihstrokescale.org/](http://www.nihstrokescale.org/).

The scale itself, in unedited format, is presented in Table A.6. Scores range from 0 (normal neurologic function) to 34 (maximal injury). Use of thrombolytics has been suggested for patients with scores between 8 and 26 who have no contraindications and can receive the intervention within a specified time of onset; for many sites the time of onset should be within 3 hours.

**Example:** A 74-year-old, left-handed woman arrives at the ED 90 minutes after the onset of left-sided arm and leg weakness. She has no headache. By 2 hours from symptom onset, she has confirmed normal findings on head CT and does not have contraindications to thrombolytics. Her symptoms have...
TABLE A.6 National Institutes of Health Stroke Scale

Administer stroke scale items in the order listed. Record performance in each category after each subscale examination. Do not go back and change scores. Follow the directions provided for each examination technique. Scores should reflect what the patient does, not what the clinician thinks that the patient can do. The clinician should record answers while administering the examination and work quickly. Except when indicated, the patient should not be coached (i.e., repeated requests to patient to make a special effort).

If any item is left untested, a detailed explanation must be clearly written on the form. All untested items will be reviewed by the medical monitor and discussed with the examiner by telephone.

<table>
<thead>
<tr>
<th>Instructions</th>
<th>Scale Definition</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>1a. Level of Consciousness (LOC): The investigator must choose a response, even if a full evaluation is prevented by obstacles such as an endotracheal tube, language barrier, or orotracheal trauma or bandages. A 3 is scored only if the patient makes no movement (other than reflexive posturing) in response to noxious stimulation.</td>
<td>0 = Alert; keenly responsive 1 = Not alert but arousable by minor stimulation to obey, answer, or respond 2 = Not alert, requires repeated stimulation to attend, or is obtunded and requires strong or painful stimulation to make movements (not stereotyped) 3 = Responds only with reflex motor or autonomic effects or is totally unresponsive, flaccid, areflexic</td>
<td></td>
</tr>
<tr>
<td>1b. LOC Questions: The patient is asked the month and his or her age. The answer must be correct—there is no partial credit for being close. Aphasic and stuporous patients who do not comprehend the questions will score 2. Patients unable to speak because of endotracheal intubation, orotracheal trauma, severe dysarthria from any cause, language barrier, or any other problem not secondary to aphasia are given a 1. It is important that only the initial answer be graded and that the examiner not “help” the patient with verbal or nonverbal cues.</td>
<td>0 = Answers both questions correctly 1 = Answers one question correctly 2 = Answers neither question correctly</td>
<td></td>
</tr>
<tr>
<td>1c. LOC Commands: The patient is asked to open and close the eyes and then to grip and release the nonparetic hand. Substitute another one-step command if the hands cannot be used. Credit is given if an unequivocal attempt is made but not completed because of weakness. If the patient does not respond to command, the task should be demonstrated (pantomime) and the result scored (i.e., follows none, one, or two commands). Patients with trauma, amputation, or other physical impediments should be given suitable one-step commands. Only the first attempt is scored.</td>
<td>0 = Performs both tasks correctly 1 = Performs one task correctly 2 = Performs neither task correctly</td>
<td></td>
</tr>
<tr>
<td>2. Best Gaze: Only horizontal eye movements will be tested. Voluntary or reflexive (oculocephalic) eye movements will be scored, but caloric testing is not done. If the patient has a conjugate deviation of the eyes that can be overcome by voluntary or reflexive activity, the score will be 1. If a patient has an isolated peripheral nerve paresis (cranial nerve III, IV, or VI), score a 1. Gaze is testable in all aphasic patients. Patients with ocular trauma, bandages, preexisting blindness, or other disorders of visual acuity or fields should be tested with reflexive movements and a choice made by the investigator. Establishing eye contact and then moving about the patient from side to side will occasionally clarify the presence of a partial gaze palsy.</td>
<td>0 = Normal 1 = Partial gaze palsy. This score is given when gaze is abnormal in one or both eyes but forced deviation or total gaze paresis is not present 2 = Forced deviation or total gaze paresis not overcome by the oculocephalic maneuver</td>
<td></td>
</tr>
</tbody>
</table>
### 3. Visual:
Visual fields (upper and lower quadrants) are tested by confrontation, with finger counting or by visual threat used as appropriate. Patient must be encouraged, but if they look at the side of the moving fingers appropriately, this can be scored as normal. If there is unilateral blindness or enucleation, visual fields in the remaining eye are scored. Score 1 only if a clear-cut asymmetry, including quadrantanopia, is found. If the patient is blind from any cause, score 3. Double simultaneous stimulation is performed at this point. If there is extinction present, the patient receives a 1 and the results are used to answer question 11.

<table>
<thead>
<tr>
<th>Instructions</th>
<th>Scale Definition</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 = No visual loss</td>
<td>1 = Partial hemianopia</td>
<td>2 = Complete hemianopia</td>
</tr>
</tbody>
</table>

### 4. Facial Palsy:
Ask or use pantomime to encourage the patient to show his or her teeth or raise the eyebrows and close the eyes. Score the symmetry of grimace in response to noxious stimuli in a poorly responsive or noncomprehending patient. If facial trauma, bandages, an orotracheal tube, tape, or other physical barrier obscures the face, these impediments should be removed to the extent possible.

<table>
<thead>
<tr>
<th>Instructions</th>
<th>Scale Definition</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 = Normal symmetric movement</td>
<td>1 = Minor paralysis (flattened nasolabial fold, asymmetry on smiling)</td>
<td>2 = Partial paralysis (total or nearly total paralysis of the lower part of the face)</td>
</tr>
</tbody>
</table>

### 5 and 6. Motor Arm and Leg:
The limb is placed in the appropriate position: extend the arms (palms down) 90 degrees (if sitting) or 45 degrees (if supine) and the leg 30 degrees (always tested supine). Drift is scored if the arm falls before 10 seconds or the leg before 5 seconds. An aphasic patient is encouraged by using urgency in the voice and pantomime, but not noxious stimulation. Each limb is tested in turn, beginning with the nonparetic arm. Only in the case of amputation or joint fusion at the shoulder or hip may the score be “9,” and the examiner must clearly write the explanation for scoring as a “9.”

<table>
<thead>
<tr>
<th>Instructions</th>
<th>Scale Definition</th>
<th>Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 = No drift; the limb holds 90 (or 45) degrees for a full 10 seconds</td>
<td>1 = Drift; the limb holds 90 (or 45) degrees but drifts down before a full 10 seconds; does not hit the bed or other support</td>
<td>2 = Some effort against gravity; the limb cannot get to or maintain (if cued) 90 (or 45) degrees, drifts down to the bed, but has some effort against gravity</td>
</tr>
<tr>
<td>5a. Left Arm</td>
<td>5b. Right Arm</td>
<td></td>
</tr>
<tr>
<td>6a. Left Leg</td>
<td>6b. Right Leg</td>
<td></td>
</tr>
<tr>
<td>Instructions</td>
<td>Scale Definition</td>
<td>Score</td>
</tr>
<tr>
<td>--------------</td>
<td>-----------------</td>
<td>-------</td>
</tr>
<tr>
<td><strong>7. Limb Ataxia:</strong> This item is aimed at finding evidence of a unilateral cerebellar lesion. Test with the eyes open. In case of a visual defect, ensure that testing is done in the intact visual field. The finger-nose-finger and heel-shin tests are performed on both sides, and ataxia is scored only if present out of proportion to weakness. Ataxia is absent in a patient who cannot understand or is paralyzed. Only in the case of amputation or joint fusion may the item be scored “9,” and the examiner must clearly write the explanation for not scoring. In the case of blindness, test by touching the nose from an extended arm position.</td>
<td>0 = Absent 1 = Present in one limb 2 = Present in two limbs. If present, is ataxia in the right arm: 1 = Yes, 2 = No 9 = Amputation or joint fusion; explain</td>
<td></td>
</tr>
<tr>
<td><strong>8. Sensory:</strong> Sensation or grimace in response to a pinprick when tested or withdrawal from a noxious stimulus in an obtunded or aphasic patient. Only sensory loss attributed to stroke is scored as abnormal, and the examiner should test as many body areas (arms [not hands], legs, trunk, face) as needed to accurately check for hemisensory loss. A score of 2, “severe or total,” should be given only when severe or total loss of sensation can be clearly demonstrated. Stuporous and aphasic patients will therefore probably score 1 or 0. A patient with a brain stem stroke who has bilateral loss of sensation is scored 2. If the patient does not respond and is quadriplegic, score 2. Patients in coma (item 1a = 3) are arbitrarily given a 2 on this item.</td>
<td>0 = Normal; no sensory loss 1 = Mild to moderate sensory loss; the patient feels that a pinprick is less sharp or dull on the affected side or there is a loss of superficial pain with pinprick but the patient is aware of being touched 2 = Severe to total sensory loss; the patient is not aware of being touched on the face, arm, and leg</td>
<td></td>
</tr>
<tr>
<td><strong>9. Best Language:</strong> A great deal of information about comprehension will be obtained during the preceding sections of the examination. The patient is asked to describe what is happening in the attached picture, to name the items on the attached naming sheet, and to read from the attached list of sentences. Comprehension is judged from responses here, as well as to all the commands in the preceding general neurologic examination. If visual loss interferes with the tests, ask the patient to identify objects placed in the hand, repeat, and produce speech. An intubated patient should be asked to write. A patient in coma (question 1a = 3) will arbitrarily score 3 on this item. The examiner must choose a score in a patient with stupor or limited cooperation, but a score of 3 should be used only if the patient is mute and follows no one-step commands.</td>
<td>0 = No aphasia; normal 1 = Mild to moderate aphasia; some obvious loss of fluency or facility of comprehension, without significant limitation in ideas expressed or form of expression. Reduction in speech and/or comprehension, however, makes conversation about the material provided difficult or impossible. For example, in conversation about the materials provided, the examiner can identify a picture or naming card from the patient's response. 2 = Severe aphasia; all communication is through fragmentary expression; great need for inference, questioning, and guessing by the listener. The range of information that can be exchanged is limited; the listener carries the burden of communication. The examiner cannot identify materials provided from the patient's response. 3 = Mute, global aphasia; no usable speech or auditory comprehension</td>
<td></td>
</tr>
<tr>
<td><strong>10. Dysartrhia:</strong> If the patient is thought to be normal, an adequate sample of speech must be obtained by asking the patient to read or repeat words from the attached list. If the patient has severe aphasia, the clarity of articulation of spontaneous speech can be rated. Only if the patient is intubated or has another physical barrier to producing speech may the item be scored “9,” and the examiner must clearly write an explanation for not scoring. Do not tell the patient why he or she is being tested.</td>
<td>0 = Normal 1 = Mild to moderate; the patient slurs at least some words and, at worst, can be understood with some difficulty 2 = Severe; the patient's speech is so slurred that it is unintelligible in the absence of or out of proportion to any dysphasia or is mute or anarthric 9 = Intubated or other physical barrier; explain</td>
<td></td>
</tr>
</tbody>
</table>
11. Extinction and Inattention (Formerly Neglect): Sufficient information to identify neglect may be obtained during the prior testing. If the patient has a severe visual loss preventing visual double simultaneous stimulation and the cutaneous stimuli are normal, the score is normal. If the patient has aphasia but does appear to attend to both sides, the score is normal. The presence of visual spatial neglect or anosognosia may also be taken as evidence of abnormality. As the abnormality is scored only if present, the item is never untestable. This is an additional item, not a part of the NIH Stroke Scale score.

\[
\begin{align*}
0 & = \text{No abnormality} \\
1 & = \text{Visual, tactile, auditory, spatial, or personal inattention or extinction to bilateral simultaneous stimulation in one of the sensory modalities} \\
2 & = \text{Profound hemi-inattention or hemi-inattention to more than one modality. Does not recognize own hand or orients to only one side of space}
\end{align*}
\]

A. Distal Motor Function: The patient’s hand is held up at the forearm by the examiner and the patient is asked to extend the fingers as much as possible. If the patient cannot or does not extend the fingers, the examiner places the fingers in full extension and observes for any flexion movement for 5 seconds. The patient’s first attempts only are graded. Repetition of the instructions or of the testing is prohibited.

\[
\begin{align*}
0 & = \text{Normal (no flexion after 5 seconds)} \\
1 & = \text{At least some extension after 5 seconds, but not fully extended. Any movement of the fingers that is not command is not scored} \\
2 & = \text{No voluntary extension after 5 seconds. Movements of the fingers at another time are not scored}
\end{align*}
\]

- Left Arm
- Right Arm

TABLE A.7 Definitions of Commonly Used Epidemiologic Terms

<table>
<thead>
<tr>
<th>Definition</th>
<th>Formula</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prevalence</td>
<td>( \frac{a + c}{a + b + c + d} )</td>
<td>Incidence of disease in the population tested</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>( \frac{a}{a + c} )</td>
<td>Probability of a positive test result, disease present</td>
</tr>
<tr>
<td>Specificity</td>
<td>( \frac{d}{b + d} )</td>
<td>Probability of a negative test result, disease absent</td>
</tr>
<tr>
<td>False-negative rate</td>
<td>( \frac{c}{a + c} )</td>
<td>Probability of a negative test result, disease present</td>
</tr>
<tr>
<td>False-positive rate</td>
<td>( \frac{b}{b + d} )</td>
<td>Probability of a positive test result, disease absent</td>
</tr>
<tr>
<td>Positive predictive value</td>
<td>( \frac{a}{a + b} )</td>
<td>Probability of a disease present, test positive</td>
</tr>
<tr>
<td>Negative predictive value</td>
<td>( \frac{d}{c + d} )</td>
<td>Probability of a disease absent, test negative</td>
</tr>
<tr>
<td>Overall accuracy</td>
<td>( \frac{a + d}{a + b + c + d} )</td>
<td>Probability of a “true” test result</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Disease State</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td>a (True positive)</td>
<td>b (False positive)</td>
</tr>
<tr>
<td>Negative</td>
<td>c (False negative)</td>
<td>d (True negative)</td>
</tr>
<tr>
<td>a + c = All patients with disease</td>
<td>b + d = All patients without disease</td>
<td>a + b + c + d = All patients tested</td>
</tr>
</tbody>
</table>

is decreased, although she is aware of the testing (1 point). She has a mild expressive aphasia but is easily comprehensible (1 point), has some slurring of words but is still comprehensible (1 point), and exhibits no evidence of neglect (0 points). Her total NIH score is 14, and as a result she may be a candidate for thrombolytic therapy.

**DIAGNOSTIC PROBABILITY**

The probability of obtaining a certain test result in the presence or absence of a particular disease entity for a given population with a given disease prevalence is presented in Table A.7.

No medical test is totally accurate. The parameters listed in Table A.7, when available, can help guide a clinician’s test selection. When this information is not available, it may be difficult to identify random laboratory errors or detect failures. Knowledge of disease prevalence, combined with the sensitivity and specificity of the test, yields the positive (or negative) predictive value of that test. For a given sensitivity and specificity, predictive value is directly proportional to prevalence. Hence, even a test with high sensitivity and specificity may not detect a rare disease. This underscores the importance of pretest clinical evaluation.

**ACKNOWLEDGMENT**

The editors and authors wish to acknowledge the contributions of M. John Mendelsohn to this Appendix in previous editions.
Index

A

Abdomen
ascites of, see Ascites; Paracentesis
injury of, see Abdominal injury
lavage of, see Peritoneal lavage
Abdominal hernia, 897–903
bowel obstruction and, 897, 898f
classification of, 897–899
definition of, 899–901
differential diagnosis of, 901, 901f
en masse reduction of, 900, 901f
epigastic, 899, 899f
temoral, 898, 898f
history in, 899
hydrocele, 899, 899f
incarcerated, 900
incisional, 898–899, 899f
inguinal
direct, 897, 898f
indirect, 897, 898f
pantaloons, 898
physical examination in, 899
radiologic imaging of, 899–900, 900f
reduction of, 901–902
complications of, 903
contraindications to, 901–902
frog-leg technique for, 902, 903f
indications for, 901–902
interpretation for, 903
procedure for, 902, 903f–903f
spigelian, 899, 899f
strangulated, 900, 900f
umbilical, 899, 899f
Abdominal injury
penetrating
emergency department thoracotomy for, 339
resuscitative thoracotomy for, 342
peritoneal lavage in, see Peritoneal lavage, diagnostic
Abdominal pain, pelvic examination for, 1216
Abdominal thrusts (Heimlich maneuver), 443–444
Acid citrate dextrose, in autotransfusion, 483–484, 494f
Acid phosphatase, 1234–1235
Acidosis
local anesthesia and, 541
metabolic, 1534, 1534f
acapnography and, 34–37, 37f
in hypothermia, 1417
restraint and, 1488
respiratory, 1534, 1534f, 1535f
rewarming, 1412
Acquired immunodeficiency syndrome (AIDS), transmission of, 503
Acrômioiacular joint

corticosteroid injection of, 1084–1085, 1084f
subluxation and dislocation of, 997–999, 998f
first degree (type I), 997–998
second degree (type II), 998
third degree (type III), 998, 998f
fourth degree (type IV), 998
fifth degree (type V), 998
sixth degree (type VI), 998
radiographic examination of, 998–999
ACTH stimulation test, before etomidate use, 117
Activated charcoal, 864–868, 865f
complications of, 867–868
contraindications to, 866
indications for, 865–866, 866f
multiple doses of, 868–869, 868f
complications of, 869
contraindications to, 868–869
indications for, 868, 868f
technique for, 869
technique for, 865f, 866–867, 867f
Activated partial thromboplastin time
(aPTT), in routine coagulation testing, 1520
Active compression decompression CPR, 334
Acute asthma attacks, evaluation of, 232–233, 232
Acute lung injury (ALI)
mechanical ventilation in, 172–173, 172f
transfusion-related, 504, 511
Acute myocardial infarction, paced cardiac rhythm and, 269, 270r
Acute respiratory distress, severity and response to treatment to, capnography and, 33, 33f
Acute respiratory distress syndrome (ARDS), mechanical ventilation in, 172–173, 172f
Adam’s apple, 127
Adenosine, in supraventricular tachycardia, 232–233, 232f
Adenosine deaminase, in pleural effusion, 486, 487f
Adenosine in, pleural effusion, 194f
Adenosine triphosphate, in compartment syndrome, 1129
Adrenal function, etomidate effects on, 117
β-adrenergic blockers, in supraventricular tachycardia, 233–234

Note: Page numbers followed by “f” refer to illustrations; page numbers followed by “r” refer to tables; page numbers followed by “h” refer to boxes.
Airway management (Continued)

King Laryngeal Tube for, 58–60, 60f
laryngeal mask airways for. see Laryngeal mask airways
manual airway maneuvers for, 39–40, 40f
oral/nasopharyngeal and nasopharyngeal
artificial airways for, 43–45, 43f–44f
oxygen therapy for, 45–49, 46f
patient positioning in, 40–41, 41f
retroglottic airway devices for, 58–61, 59f
Sellick maneuver for, 51–52
suctioning in, 42–43, 42f
techniques in, 39–45
triple airway maneuver for, 40
Airway pressure, in mechanical ventilation,
160–162
Airway pressure release ventilation, 166–167,167f
Akinosi-Vazirani block, 555, 555f
Albumin, in paracentesis, 895
Alcohol
agitation caused by, 1495
hypothermia and, 1410–1411
sexual assault and, 1240
Allentaniol, in procedural sedation and analgesia, 136f
Alkaline burns, 795–796, 796f
Alkaline burns, eye irrigation for, 1307
Alkaline phosphate, in diagnostic peritoneal lavage, 884
Alkalization, urinary, 1467–1468, 1467f
Alkalis
metabolic, 1534, 1535f
Winter’s formula in, 1536
Alveolar bone
Alveolar interdependence, 161f
Alveolar nerve, 547f
Alveolar nerve block
anterior middle superior, 551, 551f
inferior, 552–554, 553f
middle superior, 549–550, 550f
posterior superior, 546–549
Alveolar osteitis, 1397–1398
Alveolar ventilation, 160
Amanita phalloide, 859
Amatoxin, 1461, 1461f
Ambient light, pulse oximetry and, 29
Ambu bag, 146f
American Association of Blood Banks (AABB), 502–503
American Burn Association (ABLA)
grading system for burn severity and disposition of patients, 777f
on incidence of burn injury, 774
American Heart Association (AHA), in
opharyngeal and nasopharyngeal
American Implantable Cardioverter Defibrillator
Amniotic fluid, 1187, 1187f
Amputations
amputated part care in, 951, 952f
complications of, 955
contraindications to, 947f, 948
dermal slice wounds in, 953–954, 953f–955f
ear, 954–955
field, 955
fingertip, 702–706, 705f, 953–954, 953f–955f
general considerations for, 948–949
hand, 951f
indications for, 947f–948f, 948
management of, 949, 949f, 949f, 950f–951f
mechanism of, 948
nose, 954–955
patient assessment of, 949, 949b, 949f, 950f–951f
penis, 954–955
special considerations of, 951–955
hand function, 951
lower extremity, 951
stump care in, 949–951
Amylase
in diagnostic peritoneal lavage, 884
in pleural effusion, 194f
Anal canal
abscess of, 760, 761f
Anatomy
anoscopy, 760, 761f, 905f
Anal fissure, 915–916, 915f
Anastrozole
for burns, 784
capnography and, 33–34, 34f–36f, 35r–36r
general, 956f
rectal administration of, 486
Analgesics, ophthalmic, 1296–1298
Anesthesia
for burns, 784
capnography and, 33–34, 34f–36f, 35r–36r
general, 956f
nasoendotracheal intubation in, 498
ophthalmic, 1296–1298
contraindications of, 1296–1298
complications of, 1298f
indications for, 1296, 1298f
procedure of, 1296–1298
Anchoring device, of peripheral intravenous access, 399
Andexanet alfa, 517
Anesthesia
for burns, 784
nebulized, 125
of peripheral intravenous access, 397
spermatocord, 1143f, 1144–1145
in thoracentesis, 188–190, 191f
Anesthetic agents
hazards and complications of, 1075
malignant hyperthermia and, 122
ophthalmic, 1296–1298
complications of, 1298f
contraindications to, 1296, 1298f
indications for, 1296, 1298f
procedure of, 1296–1298
Anogenital ultrasonography, 1110f
Arthrocentesis of, 111f
arthrocentesis of, 111f, 1112f
articular, 111f
contraindications to, 1109f
arthrocentesis of, 1110f
Arthralgia
Ankle
arthrocentesis of, 1119, 1120f
ultrasound in, 1110f–1114f, 1112f
arthrocentesis of, 1098–1101
arthrocentesis of, 1110f
contraindications to, 1021–1023, 1022f
post-reduction care for, 1023
radiographic examination of, 1022–1023, 1022f
reduction techniques for, 1023, 1024f
nerve blocks at, 562f, 577–581, 578f–580f
soft cast for, 1053

Arthrocentesis
Arthrography
Ascites
Aspirate
Aspiration
Ascitic fluid laboratory data in, 894, 894a
causes of, 884, 885r
classification of, 886b
pathophysiological, 885b
determination of, 884
differentiation of, 884
Asphyxia
Atrial (AV) block, transvenous pacing for, 288
Atrial flutter, 221, 223f, 239f
cardiodesignation in, 238–239
carotid sinus massage in, 225, 227f
diltiazem in, 234–235
verapamil in, 235
Atrioventricular (AV) block, transvenous pacing for, 288
Atrium chest drainage systems, 497
Atropine
for bradycardia, 115
ocular, 1299–1301, 1299r
Atypical antipsychotic agents, for restraint, 1394–1395
Auditory reflex, in brain death testing, 1284
Auricular hematomas, 1358
incision for, 1359, 1360
indications for, 1358
needle aspiration for, 1360, 1360r
Autologous blood, in autotransfusion, 492
Autologous donations, 508
Automated external defibrillator (AED), 249, 249f
in children, 253
Automatic implantable cardioverter defibrillator (AICD)
ACLS interventions and, 265–268
amiodarone and, 266
cardiopulmonary resuscitation and, 265–268
characteristics of, 259–262, 261f–262f
defibrillation and, 243
electromagnetic interference and, 272, 273b
external cardiac defibrillation and, 265–268
inactivation of, 271, 271b
malfunction of, 270–274, 270b
out-of-hospital discharge of, 272, 273b
recalls of, 272
response to magnet displacement, 263–264
ventricular tachycardia and, 266, 267f
Autotransfusion, 491–499
advantages of, 497f
anatomy for, 492
antiocoagulation for, 493–494, 494t
Atrium chest drainage systems, 497f
blood filters for, 493
commercial systems, 494–495, 495f
coagulation for, 494–499
continuous, 493–496, 495f
Pleur-evac system, 495
contraindications, 491f, 493
equipment for, 491f, 493
hematologic complications for, 498, 498b
with heparin, 492, 492f
historical techniques for, 494, 494f
indications for, 493, 493f
in-line blood collection and infusion procedure for, 496–498
nonhematologic complications, 498–499
pathophysiology for, 492
Pleur-evac chest drainage systems for,
procedure for, 496–498
units for, 494–496
vacuum suction for, 493
Avulsion, 624, see also Wounds
injuries, 948
of nail, 697–698, 700f
of teeth, 1391–1393, 1393f
Awake intubation, 124–126, 124f
Axilla, abscess of, 756, 757f
Axillary artery, injuries to, in shoulder dislocation, 984, 985f
Axillary nerve, in shoulder dislocation, 983–984, 985f
Azithromycin, for antibiotic-resistant N. gonorrhoeae, 1235–1236
Babcock clamps, 1152, 1154f
Bag-back (slaps), for foreign body airway obstruction, 41–42
Backboards, 922, 923f
application of, 928f
logroll maneuver, 927, 928f
for standing position, 929
Backward, upward, rightward pressure (BUPR), 75–77, 77f
Baclofen, acute intrathecal, 1501
Bacteremia, prophylactic antibiotics in, 746–747
Bacterial vaginosis (BV), vaginal discharge, 1217
Bacteriuria, 1442, 1448–1449
catheter-associated, 1168
Bag-mask ventilation (BMV), 49–52, 49f, 64–65
complications of, 50–51, 51b
contraindications to, 49, 50f
foreign body airway obstruction and, 41 indications for, 49
pediatric, 50
technique, 49–50, 51f–52f
Bag-valve-mask technique, for nasal foreign body removal, 1377f, 1378
Bag-valve-mask ventilation, 152
Bair Claw, 134, 134f
Baker cyst, 1098
Balloon catheter, for nasal foreign body removal, 1377–1378, 1377f
Balloon tamponade, of gastroesophageal varices, 852–858, 852f–853f
aftercare, 854–857
chest radiography for, 857f
complications of, 852f, 857–858
contraindications to, 852f, 853
endoscopic band ligation in, 853
historical perspective on, 852, 853f
indications for, 852f, 853
procedure for, 854, 855f–857f
sclerotherapy in, 853
vasoactive agents in, 853
Balloon-tipped pacing catheter, 292f
Bandage contact lenses, for corneal abrasion, 1319–1321
complications of, 1321
contraindications to, 1321
equipment for, 1321
indications for, 1321
procedure for, 1321
Bandages, 640
for ankles, 1052–1053
elastic, for splinting, 1042–1043
for foreign body airway obstruction and, 41 indications for, 49
Barotrauma, with percutaneous transluminal intubation, 139–140
Bar factory, local anesthesia and, 541
Baric probe, 1086
Barotrauma
Barotrauma, with percutaneous transluminal intubation, 139–140
Barrier precautions, 1470, 1471f
Bartholin abscess, 1213–1214, 1214f
incision and drainage for, 1213–1214, 1215f–1216f
Word catheter for, 1214, 1216f
Bladder (Continued)

key procedure sequence of, 1172–1173
pathophysiology of, 1174
tapping, 1442–1443
Blanket, heat exchange, 1412
Bleeding, 154–155
after corticosteroid injection therapy, 1076
anticoagulant associated, management of,
1521–1528
with arthrocentesis, 1116
assessing risk of, in setting of
anticoagulation, 1520–1521
in brain, 1278
central venous catheterization and, 1525
with cricothyrotomy, 134
gingival, 1396–1397
International normalized ratio and,
1520
intraperitoneal, 1218
nasal, 1363, 1365f
posttonsillectomy, 1383, 1383f
risk of, in thoracentesis of patients taking
clopidogrel, 1527
with tracheostomy, 154–155, 155f
upper gastrointestinal, nasogastric feeding
tube in, 829
from urethral catheterization, 1168
vaginal, in adult woman, 1214–1217
Blind nasotracheal intubation, 102–105, 105f
anterior to epiglottis to, 103
arytenoid cartilage to, 103–104, 104f
complications of, 104–105
contraindications to, 102–103
esophageal placement of, 104
indications for, 102–103
laryngospasm with, 104
arytenoid cartilage to, 103–104, 104
vocal cords to, 103–104
procedure and technique for, 93
physical difficulties of, 103–104
technical difficulties of, 103–104
vocal cords to, 103–104
Blister
burn-related, 780f, 785f, 789, 789f
frostbite-related, 1418, 1419f
Blood
alkalinization of, 1467–1468, 1467b
cultures, 1452–1457
acidosis vs. anaerobic bottles for,
1455–1456, 1456b, 1456f
contaminants in, 1456, 1456b
fungal, 1457
indicators for, 1452, 1452b
intraoperative specimens for, 1454
obtaining, techniques for, 1453–1454,
1454f
outpatient, 1452–1453, 1453f
phlebotomy for, 1454
sets of, 1453, 1455f, 1456f
timing of, 1454
volumes, 1454–1455
drawing of, from central venous catheter,
408
febral, 1458–1460, 1459f
gastric, testing for, 1460
hemodialysis, 1460, 1460f
ordering of, 518–519
request forms for, 519
amongologic tests on, 1464–1465
in urine, 1126f, 1445f, 1446, 1447f
Blood groups, 500, 501f
Blood levels, high, local anesthesia and,
549–551
Blood patch, epidural, 1270
Blood pressure, 6–11
in children, 1
diastolic, 7–8
Blood pressure (Continued)
measurement of
bilateral, 10
complications of, 8
contraindications to, 6
cuff testing for, 7
Doppler ultrasound in, 11–12, 12f–13f,
14r
equipment for, 6–7
errors in, 9
indications for, 6
Korotkoff sound in, 7
procedure for, 7–8
respiration and, 10
monitoring of, for arterial puncture and
cannulation, 377
normal, 2
physiology of, 6
sphygmomanometer, 7–8
Blood sampling
arterial, 356–359, 358f–359f
for arterial puncture and cannulation, 377
capillary, 353–354, 355f
Blood tests, in sexual assault evaluation, 1234
Blunt trauma, 338–339, 875–876, 876f, 877f
Body cooling unit, 1427–1428
Body surface mapping, in
electrocardiography, 280–281
Body-piercing, removal and, 728–730,
729f–730f
“Boils”, 754–756, 756f
Bone Injection Gun (BIG), 465, 465f,
468–470, 471f
Bone marrow aspiration needle, 464
Bone marrow aspiration needle, 464
Bone, see Tracheal tube introducer
Boutonnière deformity, 967f
Brachial artery
arterial cannulation for, 7–8
pressure, differential, 10
Brachial plexus
anatomy of, 566, 566f
injuries to, in shoulder dislocation, 984,
985f
Brachial vein
catheterization, for transvenous cardiac
catheterization, 295
internal, for transvenous cardiac pacing,
294f
Bradyarrhythmia, 5
atropine for prevention of, in intubation,
115
in cold water immersion, 1420
bradycardia, 5
atropine for prevention of, in intubation,
115
in cold water immersion, 1420
diving, 231
paradoxical, 15–16
transcutaneous pacing in, 288f
Bradykinin
hyperventilation, 34
Brain
abscess of, 1260
bleeding in, 1244
hemiation of, 1243, 1244f, 1270
infarction of, carotid sinus massage and,
1228
nasogastric tube in, 835, 837f
strokes, 1243
volume of, 1243
Brain death
definition of, 1284
testing, 1284–1285
Brainstem reflex testing, 1284–1285
Braxton-Hicks contractions, 1186
Breast abscess, 758, 758f
Breathing
bedside toxicologic test on, 1463–1464
diaphramatic, 4
INDEX

Breathing (Continued)
periodic, 4
rapid, in mechanical ventilation, 174
Breecnh presentation, 1188–1190, 1188f, 1200–1203, 1202f
Brescia–Cimino formula, 452
Broad-spectrum antibiotic ointments, for burns, 787
Bromocriptine, in neuroleptic malignant syndrome, 1425
Bronchospasm
acute, capnogram for, 34
in esophageal foreign bodies, 808
in intubation, 115–116
Bronchus
injury to, 341
nasogastric tube in, 835, 836f
Browiocatheter, 447, 448f
Brudzinski’s sign, 1259
Bullying, 400
Brut, carotid, 228–229, 230f
Buccal mucosa, laceration of, 1394–1395
Buccal nerve, 547f
Buddy tapping, 1042, 1045f
Bundle branch block, transcutaneous pacing in, 290–291, 291f
Bunion, 1057, 1058f
Bunion bursitis, 1078, 1100
Bupivacaine, 525–526f
induced ventricular dysrhythmias, 542
infiltration of, 533f–534r, 534
lidocaine and, 534, 54sf
in regional anesthesia, 560
for toothache, 1387
Burns
774–805
abuse-related, 792–794, 793f
acid, 795–796, 795f
air bag, 796–797, 796f
alkali, 795–796, 795f, 796f
with blisters, 780f, 785f, 789, 789f
cement, 796f
characteristics of, 775f
chemical, 780f, 790f, 795–800, 801f
chronic acid injury with, 800
coneval, 791, 792f, 797
deep, 774, 776f
depth of, 774–777, 776f
dressings, 783–784, 784f–785f, 788f
edema with, 786, 786f
electrical, 800f, 801f
emergere skaratomy for, 803–805, 804f
facial, 791–792, 792f
first-degree, 774, 775f, 776f
flash, 791, 792f
fluid resuscitation in, 1533
foot, 790, 790f
fourth-degree, 774–777, 776f
hand, 780f, 790–791, 791f
histopathology of, 778–779
hot tar, 794–795, 794f–795f
hydrocarbon, 797
hydrofluoric acid injury with, 797–800, 798f–799f
hypothermia with, 780–782
major, 774, 777f
initial care of, 781–782, 781f–782f
minor, 774, 777f, 780f
clinical issues in, 784–787
follow-up care of, 787–789, 788f, 788f
infections in, 789–790
initial care of, 782–783
outpatient care of, 783–787, 784f–785f
moderate, 774, 777f
open, care for, 783
phenol injury with, 797
Burns (Continued)
phosphorus, 800
in pregnancy, 794
pruritus with, 785
radiation, 803
"rule of nines" in, 777, 779f
second-degree, 774, 775f, 776f
size of, 777–778, 777f
treatment of
aloe vera cream for, 787
anesthesia, 786
blister management in, 783, 789, 789f
broadspectrum antibiotic ointments, 787
chemical, 801f
cooling therapy in, 780–782, 781f, 783f
corticosteroids for, 787
emergency, 780–781
emollients for, 788
fluid resuscitation in, 781
honey for, 787
other topical preparations, 787
outpatient physical therapy in, 788
outpatient visit, inpatient care, 779–780
silver sulfadiazine for, 786–787, 787f
with topical antimicrobials, 786
wound evaluation in, 774–778, 775f
wound healing of, 788–789
zones of, 778–779, 779f
Bursae, 1071
Bursitis, 1071, 1072f
anserine, 1078, 1098, 1100f
bunion, 1078, 1100
ischioiogluteal, 1096–1097
prepatellar, 1078, 1097–1098, 1097f, 1099f
radiohumeral, 1078f, 1085–1086
subacromial, 1078f, 1080f, 1083f
suprapatellar, 1098
treatment of, 1071–1104
trochanteric, 1078f, 1095–1096, 1096f–1097f
Butterfly needle
for phlebotomy, 1457, 1457f
for scalp vein intravenuous catheterization, 360, 362f
Button battery ingestion, 824–825, 826f
Caloric testing (Continued)
contraindications to, 1281
equipment for, 1281
indications for, 1281
interpretation of, 1282–1284
physiology for, functional anatomy of, 1281, 1282f
procedure for, 1281
Canalith-repositioning maneuvers, for vertigo, 1286–1288, 1287f–1288f
complications of, 1288
contraindications to, 1286–1287
indications for, 1286–1287
procedure for, 1287–1288
Canaliths, in DIIX-Hallpike test, 1285
Cancer
abscess and, 739, 740f
in fetal development, 1507
pericardial effusion in, 313
Candida, urinary, 1499f
Candidiasis, vaginal discharge, 1217
Cantholysis, 1334, 1335f
Canthotomy, lateral, 1334, 1335f
Capillary blood sampling, 333–334
Capillary refill, 14–16
Capillary refill time (CRT), 16–17
Capnography, 29, 30f–31f
in CPR, 336–337, 337f
indications for
in intubated patients, 31–33
in spontaneously breathing patients, 31–33
limitations of, 37
physiology in, 31
in procedural sedation and analgesia, 598–600
technology for, 30
terminology for, 30
Capnometer, 30
Captain Morgan technique, in hip dislocation, 1014, 1015f
Carbon dioxide (CO₂)
concentration of, and time, 29–30
monitoring of, 29–37
mainstream or sidestream, 30, 30f
quantitative or qualitative, 30, 31f
retention in oxygen therapy, 48–49
in percutaneous translaryngeal ventilation, 139–140
Carbon monoxide poisoning, 781–782
oxygen therapy for, 43
Carboprost tromethamine, for postpartum hemorrhage, 1205f
Carbuncles, 754–756, 756f
Cardboard splints, 931
Cardiac arrest
algorithm, 245f
artificial perfusion during. see Cardiopulmonary resuscitation (CPR)
central venous catheterization in, 430
defibrillation and, 241
hypothermia and, 342, 1410, 1416–1417
identifying cause of, capnography and, 32
in mechanically ventilated patients, 175–177
oxygen therapy during, 45, 46f
pediatric, 250
resuscitation, prognosis after initiation of, capnography and, 32
targeted temperature management for, 1422
transvenous cardiac pacing in, 290
unwitnessed, 249
automated external defibrillator application in, 249, 249f
medication for, 249, 249f
Cardiac pacing

- (Continued)
thrombosis with, 301
in trauma, 290
- ultrasound for, 300f–301h, 300f
US guidance in, 299
venous access for, 295

Cardiac trauma, penetrating, 311–312
Cardiopulmonary bypass

in hyperthermia, 1432
in hypothermia, 1415–1416
Cardiopulmonary resuscitation (CPR), 241–242, 332–337
active compression decompensation, 334
adjuncts to improve quality of, 333–336
capnography in, 32, 336–337, 337f
in central venous catheterization, 408
chest compressions-, 333
compressions in, 332, 333b
feedback devices of, 334, 335f
monitoring of, 334, 335f
duration of, emergency department thoracotomy and, 340
effectiveness of, capnography and, 32, 32f
epinephrine in, 249
extracorporeal, 335–336, 336
goal of, 342
hemopericardium with, 312
high-quality, 240f
in hypothermia, 1410, 1416–1417
impedance threshold device in, 334, 335f
leadership in, 333
mechanical devices for, 335, 335f
monitoring during, 336–337
open-chest, 342
in patients with implantable pacemakers, 299–303
pneumothorax and, 196, 197f
in pulseless ventricular tachycardia, 241–242
pulses during, 6
workup in, 333
ultrasound monitoring in, 337
ventilations in, 332
in ventricular fibrillation, 241–242
Cardiovascular disease, in procedural sedation and analgesia, 596
Cardioversion, 238–257
algorithm for, 254f
in atrial fibrillation, 236, 236b
cardiocerebrovascular, 256
- complications of, 258f
contraindications to, 258f, 257
electrical, 235–237
- electrode position in, 256
energy requirements for, 256–257
equipment for, 238f, 254–255, 255f
indications for, 238f, 254
- pediatric, 257
- physiology of, 253–254
- sedation for, 255–256, 256t
- technique for, 253–257
treatment with, 254
- Carotid bruit, 228–229, 230f
- Carotid pulse, 5
- Carotid sinus, 229f
- Carotid sinus massage (CSM), 5, 227–230
- complications of, 229–230
- contraindications to, 228–229
- diagnostic use of, 223–226, 227f, 228t
- technique of, 229f–230f
- unsuccessful, 229
- Carotid sinus syndrome (CSS), 225–226
- Carpal dislocation/dissociation, 1011–1012, 1011f–1013f
- Carpal tunnel syndrome, 1078r, 1093–1094, 1093f
- Carpometacarpal joint arthritis, 1116, 1116f
dislocation of fingers, 1011
thumb, 1008, 1008f
- Casts
- bivalving of, 1055
- burn injury from, 1053
dermatitis and, 1054
- ischemia from, 1053, 1053f
- pain from, 1055
- pruritus and, 1054, 1054f
- removal of, 1056f
- Cat bite, 648–649, 650f, 652f
- Catecholamine reactions, local anesthesia and, 543
- Catecholamines, in intracranial hypertension, 113
- Catfish sting, 725–726, 726f
- Cathartics, 869
- Catheterization

- central venous. see Central venous catheterization
- umbilical artery, 370–371, 370f–371f
- umbilical vein, 367–369, 368f–369f
- urethral, 1157–1170, 1159f
- aftercare for, 1168
- bleeding after, 1168
- complications of, 1168–1170
- contraindications to, 1162–1164
- difficult, 1163, 1168
- equipment for, 1164, 1164f, 1164t
- female, 1164–1167, 1166f
- anatomy of, 1160, 1161f–1162f
difficult, 1168
- indications for, 1162
- infection from, 1168
- key procedure sequence of, 1159–1160
- long-term catheter use in, 1168
- male, 1164–1167, 1165f, 1167f
- anatomy of, 1160, 1162f
difficult, 1163, 1168
- mechanical complications of, 1168
- nontubing catheter removal after, 1168–1170, 1169f, 1170f
- pathophysiology of, 1160
- traumatic Foley catheter removal after, 1170, 1171f
- undesirable catheter retention in, 1168–1170
- Catheters

- for arterial cannulation, 379, 379f
- for arterial puncture, 379
- balloon, for nasal foreign body removal, 1377–1378, 1377f
- Bronchi, 447, 448f
- central venous, 412–413, 413f, 413r
- complications of, 413
- misplacement of, 429–430
- placement of, 415f–416f, 417–418
- replacement of, 418–419
- for compartment pressure measurement, 1127, 1138f
- in cooling techniques, 1432
coudé, 1164, 1164f
- cuffed, tunneled right atrial, 447, 448f
de Pezzer, in anorectal abscess, 762
in diagnostic peritoneal lavage, 880
- displacement or malposition in, 459
- dual-balloon, for posterior nasal packing, 1369, 1371f
eMBOLIZATION of, with central venous catheterization, 437
- in endotracheal drug administration, 479
Central venous catheterization (Continued)

contraindications to, 406f, 409–410
dressing for, 425, 428f
for emergency venous access, 408
equipment for, 406f, 411–413
external jugular vein for, 430
femoral vein for, 407, 408f, 409t, 410, 420, 424–425, 425f
guidewire for, 411–412, 412f
placement of, 413–417, 415f–416f, 418f
for high-flow fluid bolus, 408
historical perspective on, 405
in hypoperfusion, 409
for inability to achieve peripheral access, 408
indications for, 406f, 407–409
internal jugular vein, 405–407, 407f, 409t, 410, 415f–416f, 420, 422–424, 423f
needle for, 411f, 411r, 414–417, 417f
over-the-needle technique for, 419
pediatric, 364–367, 364f
complications of, 367
equipment and setup for, 364
femoral, 365–366, 365f
indications and contraindications in,
internal jugular vein, 366, 367f
subclavian, 366–367
umbilical, 367–369, 368f–369f
preparation for, 413
problems related to, 301
radiographs for, 429, 429f
Selinger (wire) technique for, 410, 413–417
for serial blood drawing, 408
sheath introducer in, 417–418, 419f
subclavian vein for, 405, 407f, 409t, 410, 420
infracavicular, 420–421, 421f
supracavicular, 421–422, 422f
technique for, 413–419, 415f–416f, 419f
training for, 438
ultrasound for, 411, 413, 414f, 426b–428b,
survey, 414f
Central venous pressure measurement
of, 430
complications of, 431f
contraindications to, 430, 431f
errors in, 433–434, 434b
indications for, 430, 431f
interpretation of, 434
manometry for, 432–433, 432f
transducer for, 432–433, 433f
monitoring of, 407–408, 410–415
in cardiac tamponade, 434–435
in fluid challenge, 434
physiology of, 431
Cephalic vein catheterization of, 430
venous cutdown of, 441–442, 441f
Cerebral blood flow (CBF), 1245
Cerebral infarction, carotid sinus massage and, 228
Cerebral perfusion pressure (CPP), 1245,
1246f
Cerebrospinal fluid (CSF), 1244–1245, 1245f,
1263f
anatomy of, 1258–1259
examination of, 1258–1280
analysis with infections, 1275–1279
appearance in, 1272–1273
in bacterial infections, 1275, 1275r
cells in, 1273
in cryptococcal meningitis, 1279
culture in, 1276
in cytomegalovirus infection, 1280
Cerebrospinal fluid (CSF) (Continued)
glucose in, 1273–1274, 1273b
in immunocompromised patients, 1279–1280
in mycobacterial tuberculosis, 1280
in neurophilis, 1278
pressure in, 1269
in primary CNS lymphoma, 1280
in progressive multifocal leukoencephalopathy, 1280
protein in, 1274
red blood cells in, 1272
in resuscitative thoracotomy, 1279–1280
in intracranial shunt assessment of, 1256–1257, 1257f
excessive drainage of, 1253
physiology of, 1258–1259
Cerumen impaction removal, 1351–1355,
1352f, 1354f
ceruminosities in, 1353, 1354f
complications of, 1353–1355
contraindications to, 1352, 1352f
indications for, 1352
irrigation in, 1353, 1354f
manual instrumentation in, 1353, 1354f
procedure for, 1352–1353
Ceruminolytics, 1353, 1354f
Cervical collars, 921, 921f
application of, 925f
Cervical esophagostomy, 840
Cervical extrication splints, 921–922, 921f
Cervical myofascial pain syndrome, 1103
Cervical pharyngostomy, 840
Cervical spine
immobilization of C-collars for, 921, 921f
indications of, 930
extrication splints for, 921–922, 921f
helmet removal and, 942
indications for, 919
in pediatric patients, 929–930
procedure for, 924–929, 925f
injury to, 918–919
Cervicis, vaginal discharge in, 1217
Cervix
dilation of, 1190
effacement of, 1190, 1190f
Cesarean section, perimortem, 1207, 1208f
Chair technique
in elbow dislocation, 1001, 1001f
in shoulder dislocation, 993
Chandy maneuver, 55–57
Chelation therapy, for chronic acid injury, 800
Chemical burns, 780, 780f, 795–800, 795b, 801f
Chemical cautery, for hemorrhage, in IVADs, 458–459
Chemical dermatitis, 562
Chemical terrorism, assessment and triage of victims of, capnography and, 33, 33et
Chemotherapeutic agents, extravasated, antidotes for, 402f
Cherry-red spot, 1331, 1333f
Chewy undertow, 201–202, 201f–202f
Direct laryngoscopy (Continued)
indications for, 67f, 69
learning, 72, 74f
optimal patient positioning for, 71–72, 71f
procedure and technique of, 72–85
Direct pathway, between vestibular and oculomotor nuclei, 1281
Direct pressure, for hemorrhage, in IVADs, 457–458
“Directed donations”, 508
Disfellowshipping, 519
Dislocations
general principles of, 980–983, 981f
management of, 980–1026
preparation of patient with, 980
timing of reduction for, 982–983, 982f
Displacement, of catheter, in IVADs, 459
Disseminated intravascular coagulation, 506
Disposition, paracentesis and, 896
Displacement, of catheter, in IVADs, 459
Disfellowshipping, 519
Direct pressure, for hemorrhage, in IVADs, 457–458
Direct pathway, between vestibular and oculomotor nuclei, 1281
Diving reflex, 231
DIX–Hallpike test, for vertigo, 1285–1286, 1286f
Dry needling, 1074
Dry tap
ultrasound in, 1110
in culdocentesis, 1221–1222, 1223f
in pericardiotomies, 330
Dundee micropuncture technique, 1154, 1154f
DuoDERM, 641
Dyes, toluidine blue, 1229f
perineal staining with, 1232, 1233b, 1233f
Dyshemoglobinemias, effects of, pulse oximetry and, 28
Dysphagia, in esophageal foreign bodies, 808
Dyspnea
spirometry for, 23
in transtracheal oxygen therapy, 157
Dysrhythmias
drug-induced, 291
mechanism of, 241
with pericardiocentesis, 330
reentry in, 222–225, 224f, 224f
transtracheal suctioning and, 147
ventricular, bupivacaine-induced, 542
E
Ear, 1348–1361
amputations of, 954–955
anatomy of, 1348–1349, 1349f
anesthesia of, 1349–1351, 1350f
pericardiocentesis from, 1351–1355, 1352f, 1354f
lacerations of, 642, 1350f
syringing, 1353, 1354f
temperature in, 18, 20f
wick placement, 1355, 1356f
Eardrops, 1352, 1352f
EasyTube, 60–61
Echocardiography, for pericardial effusion or tamponade, 318–322, 319f
Eclipse sign, 1300f
Ecstasy, overdose, 1425–1426
Ectopic pregnancy, 1218–1220, 1218r, 1219f
Edema
in brain, 1243
with burns, 786, 786f
in foreskin, 1151f
pulmonary
after pericardiocentesis, 331
after thoracentesis, 195
after tube thoracostomy, 220
noninvasive positive pressure ventilation in, 168
Edoxaban, 517
Edrophonium (Tensilon) test, for myasthenia gravis, 1291–1292, 1291f
Effective dose, 1505
Effusion
joint, 1106, 1107f
parapneumonic, 199–200
Einhoven, Willem, 275
Einhoven's triangle, 277f
Eject Helmet Removal System, 946f
ELA-Max, 529–530
Elbow, 956
arthrocentesis of, 1117, 1118f
ultrasound in, 1110–1114b, 1113f
barly, 1086
corticosteroid injection of, 1085–1089
dischonings of, 999–1002
anterior, 1000f, 1002
reduction techniques for, 1002, 1002f
posterior, 1000–1002, 1000f
post-reduction care for, 1002
radiologic examination of, 1000, 1000f
INDEX 1555

Elbow (Continued)
 reduction techniques for, 1000–1002
 chair technique in, 1001, 1001f
 recommended initial approach in, 1000–1001, 1001f
 traction-countertraction method in, 1001, 1001f
 golfer’s, 1085, 1085f
 nerve blocks at, 562r, 566–567, 567f
 nursemaid’s, 1002–1005, 1003f
 clinical assessment of, 1002–1003, 1003f
 radiographic examination for, 1003
 reduction techniques for, 1004–1005, 1004f
 tennis, 1085, 1085f
 Elderly patient
 abuse of, burns and, 792–794, 793f
 agitation in, 1496
 carotid sinus syndrome in, 225–226
 spinal cord injury in, 919
 Electrical alternans, 318, 318f
 Electrical burns, 800, 801f
 Electrical storm, 270
 Electrocardiography (ECG), 275–287
 alternative leads for, 281–284, 283f, 283r
 amplitude, alteration in, 281
 arm electrode reversal in, 284, 284f
 artifacts in, 286–287, 287f
 basic equipment for, 275–278
 body surface mapping in, 280–281
 calibration for, 278, 279f
 central venous catheter intracardiac leads for, 284
 computer provided information, 278
 correlation of, regional action potentials and, 240f
 80-lead, 280–281
 electrode misplacement in, 284–286
 esophageal leads for, 284
 15-lead, 279–280, 279f
 in hypothermia, 1407, 1409f
 indications for, 275
 invasive procedural leads for, 280
 leg electrode reversal in, 284, 285f
 Lewis leads for, 281, 283f, 283r
 limb electrode reversal in, 284, 285f, 286r
 limb-precordinate leads for, 283
 machine, in transvenous cardiac pacing, 293, 293f, 302f
 modified bipolar chest leads for, 283–284, 283f, 283r
 in myocardial infarction, 275
 P wave, 282f
 paper speed in, 278, 281, 282f
 for pericardial effusion or tamponade, 317–318, 318f
 for pericardiocentesis, 280, 326–327, 327f
 posterior leads for, 278–279, 279f
 precordial electrode misplacement in, 285–286, 286f
 prehospital, 281
 in procedural sedation and analgesia, 600
 right-sided leads for, 280, 280f
 12-lead, 275–276, 276f–277f, 276r
 electrode placement for, 277–278
 pediatric, 278
 vertical sternal (Barker) leads for, 283, 283f, 283r
 Electrocautery, pulse oximetry and, 29
 Electrodes
 position of, in cardioversion, 256
 in transvenous cardiac pacing, 292
 Electroencephalography, in increased intracranial pressure, 1248
 Elemental alkali metal burns, 800
 Elevated plasma free hemoglobin, with autotransfusion, 498–499
 Elevation, in wound care, 642–643, 643f
 Embolism
 air, with autotransfusion, 498
 fat with autotransfusion, 498
 with intraosseous infusion, 475
 pulmonary, see Pulmonary embolism
 Embolization catheter, with central venous catheterization, 437
 cerebral, in arterial cannulation, 392
 in IVADs, 456
 Emergence reaction in ketamine, 616
 ketamine-related, 118–119
 Emergency airway algorithm, 68–69, 69f
 Emergency childhood, 1186–1210
 Emergency dental procedures, 1384–1404
 Emergency department (ED)
 airway management in, 39
 pulse oximetry in, 26
 spirometry in, 23
 Emergency department thoracotomy (EDT), 338–339
 Emergency intubation, general approach to, 63
 Emergency technique, ideal vs., 68
 EMLA cream in intact skin, 529–530
 mucosal application of, 527f, 528
 TAC and, 532
 Emollients, for burns, 788
 Emopymas, 190–200f
 in stable patients, 201
 tube thoracostomy for, 206–207
 Encephalopathy syndrome, 1426
 Endocarditis, prophylactic antibiotics in, 1417
 Endoscopically assisted, 1378
 Endoscopic third ventriculostomy (ETV), 1257
 Endoscopy, for esophageal foreign bodies, 81f
 Endotracheal administration, of medication, 476–481, 476f
 Endotracheal intubation, preoxygenation prior to, 48
 Endotracheal tube, 81f
 monitoring position of, during transport, 32
 pediatric, 70f
 placement, verification of, capnography and, 31–32
 replacing, 108f
 size, 1529–1530
 standard, using, 93
 End-tidal CO₂ pressure (PETCO₂), 29–30, 30f
 accuracy of, ET tube placement and, 31–32
 in acute respiratory distress, 33, 33f
 colorimetric detector for, 30
 CPR effectiveness and, 32
 detection, 83–84
 serum bicarbonate and, 34–37, 37f
 English nail anvil, 1067f, 1068–1069
 English-to-metric conversions, 1529, 1530f
 Entrapment, of esophageal foreign bodies, 808f
 Envenomation
 coelenterate, 724
 stingray, 726
 Enzyme-linked immunosorbent assay, in meningitis, 1276
 Enzymes, in diagnostic peritoneal lavage, 884
 Eosinophils, CSF, 1273
 Epicondylitis, 1078f
 lateral, 1085–1086, 1085f–1086f
 medial, 1085–1086, 1085f, 1087f
 Epidermoid tumor, 1271
 Epididymitis, acute, 1142
 Epidural blood patch, 1270
 Epidural hematoma, 1248, 1248f–1249f
 spinal, 1260–1261
 Epiglottitis, in esophageal foreign bodies, 808
 Epiglottoscopy, 72
 Epinephrine
 anxiety and, 543
 in auricular field blocks, 1351
 in CPR, 249
 depot effect of, 477
 endotracheal drug administration of, 477f
 local anesthesia and, 526, 533–534, 535f, 539f
 for priapism, 1148, 1148f
 push-dose, how to make, 1148, 115f
 regional anesthesia and, 561
 for wound, 632
 Epinephrine-cocaine, 532–533
 Epiglottis
 acute, 318, 318f
 Epiphysis, intraosseous infusion-related injury to, 475
 Episiotomy, 1203–1205, 1204f
 Epistaxis
 anatomy of, 1361, 1361f
 anticoagulation and, 1527
 patients with, 1363
 management of, 1363–1373, 1363f–1365f
 anterior nasal packing for, 1366–1368, 1367f
 antibiotics for, 1372
 cautery for, 1365–1366, 1366f
 complications of, 1368, 1369f, 1372
 contraindications to, 1364
 equipment for, 1364, 1364f
 indications for, 1364
 inflatable balloon packs for, 1369, 1371f
 patient disposition in, 1372
 posterior gauze pack for, 1368–1369, 1370f
 posterior nasal packing for, 1368–1372, 1370f
 testing for, 1363–1364
 Epley maneuver procedure, 1287, 1287f
 Epstein-Barr virus (EBV), transfusion transmission of, 503
 Epulis gravidarum, 733–734
 Equipment failure, of tracheostomy, 153
 Equivalent dose (Hₑ), 1505
 Ergonovine maleate, for postpartum hemorrhage, 1205f
 Erythromycin, for antibiotic-resistant N. gonorrhoeae, 1235–1236
 Escharotomy
 complications of, 805
 emergency, 803–805
 indications for, 804
 technique of, 804–805
 for chest, 805
 for limbs, 804–805, 804f
 for neck, 805
 for penis, 805
 Eskimo technique, in shoulder dislocations, 993f–992f, 993
 Esmolol, in supraventricular tachycardia, 214
 Esophageal bougienage, for removal of esophageal foreign bodies, 819–820, 820f, 820r
 Esophageal detector device, of tracheal tube, 81–81
Evaporation, heat loss with, 1407
Evaporative cooling, 1427–1428, 1427r
Excited delirium syndrome (EXDS), 1481–1483
Explosion, defibrillation and, 243
Examination, for intravenous regional anesthesia, 593
Extensor carpi radialis brevis, 956, 957f–958f
Extensor carpi radialis longus, 956, 957f–958f
Extensor carpi ulnaris, 956–958, 957f–958f
Extensor digiti minimi, 956, 957f–958f
Extensor digitorum communis, 956, 958f, 961f, 968f
Extensor digitorum longus, 972, 972f
Extensor hallucis longus, 972, 972f
Extensor indicis proprius, 956, 958f
Extensor pollicis brevis, 956, 957f–958f
Extensor pollicis longus, 956, 957f–958f
Extensor tendon injury, foot, 972–973, 972f
in hand and wrist, 956–972, 957f
antibiotics in, 962
functional anatomy of, 956–958, 957f–958f, 960f
general approach of, 958–962, 961f
repair of aluminum foam splint in, 963f, 970f
care after, 971
complications of, 970–971
preparation for, 961f, 962
zones 1 and 2, 968–970, 969f
zone 3, 967–968, 967f
zone 4, 965f, 966–967
zone 5, 965f–966, 965f
zone 6, 963–965, 964f
zone 7 and 8, 963
knee, 978–979, 979f
patterns and management of, 962–970
postrepair care and rehabilitation of, 971–972
External auditory canal, 1348–1349
anesthesia of, 1350f, 1351
examination of, 1351, 1351f
foreign body removal in, 1355–1358, 1358f
complications of, 1357–1358
contraindications to, 1356
cyanoacrylate for, 1357, 1358f
Fogarty catheters for, 1357, 1358f
Foam, in hand and wrist, 1303
contraindications for, 1305
procedure for, 1305
contraindications to, 1305
duration of, 1306–1307
equipment for, 1305
indications for, 1305
Morgan lens, 1307, 1308f
procedure for, 1305–1307, 1306f–1307f
prolonged, 1307
normal, 1300f
optokinetic nystagmus testing of, 1296, 1300f
patching of, 1316
penlight examination of, 1300f
Pseudomonas infection of, 1318
slit lamp examination of, 1328–1331
stye of, 1336, 1337f
tonometry of, 1322–1328
transillumination of, 1299, 1300f
visual acuity, assessment of, 1295–1296
EZ-IO Device, 465–466, 466f

F
Face
Abscess in, 1399–1400
laceration of, 685–686, 685f–687f
epithelial stitch in, 685, 687f
natural skin lines and, 685, 685f
stellate, 683–685, 685f
tape closure in, 685–686, 687f
natural wrinkles of, 685–687, 687f
topolical wound anesthesia for, 532
Facial burns, 791–792, 792f
Factor II, direct inhibitors, 516–517, 517r
Factor VII, 511–513
for hemophilia, 512
for intracranial hemorrhage, 512–513
for trauma, 513
warfarin reversal in, 515–516, 516r
Factor VIII
concentrates, 513
in cryoprecipitate, 511
Heart (Continued)

direct cardiac compressions of, 345–346, 345f
Foley catheter insertion into, 347, 348f
hemorrhage in, staples for, 346, 346f
partial-oclusion clamps for, 347, 349f
Sauerbruch maneuver for, 346–347, 347f
suture repair of, 346, 347f
trauma to, with central venous catheterization, 437
water-bottle, 317, 318f
wounds of, hemorrhagic, 346–347, 346f
Heart failure, congestive
pericardial effusion in, 313
pleural effusion in, 181, 182f
Heart rate (HR)
fetal, 1187, 1187f, 1191, 1192f
local anesthesia and, 541
normal, 239–240
in pediatrics, 2r
Heart rhythm, 5–6
Heat cramps, 1423
Heat exchange blanket, 1412
Heat exhaustion, 1423, 1423f
Heat illness, 1422
Heat injury, from splinting, 1053–1054
Heat loss, 1407
Heatstroke, 1423–1426
exertional, 1423–1424
Heel
capillary blood sampling from, 353
pain in, 1100–1101, 1100f
Heel pain syndromes, 1057–1058
Heel spur pain, 1057
Heel stick, in neonates, 1454
Heimlich maneuver, 41–42, 42f
Helmet, removal of, 940–946, 942f
contraindication to, 943
indication for, 942–943
motorcycle helmet, 945f, 946
sport helmet, 943–946, 944f
Hemangioma, capillary, lobar, 733–734, 734f–735f
Hemarthrosis, 1108–1109, 1109f
arthrocentesis and, 1526
spontaneous nontraumatic, 1526
Hemocrit, of pericardiocentesis fluid, 329
Hematoma
temporal, 984–985
Hemoglobin
fetal, pulse oximetry and, 28
in transfusion guideline, 505
Hemoglobin-based oxygen carriers (HBOCs), 508
Hemodynamics
in autotransfusion, 498
drug-induced, 504
Hemopericardium
acute, 311–313
atriaumatic, 312–313
blunt trauma causing, 312
focused assessment with sonography in trauma (FAST) examination in, 340–341, 340f
miscellaneous trauma causing, 312
nontraumatic, 312
penetrating trauma causing, 311–312
traumatic, 311–312
Hemoperitoneum, 1220
Hemophilia
central venous cannulation and, factor replacement prior to, 1525
dental procedures and, 1527
factor VII for, 512
lumbar puncture in patients with, 1524
Hemorrhage
amputations and, 949
epidural, spinal, 1272
gastrointestinal, bedside tests for, 1458–1460
in great-vessel wounds, control of, 349, 349f
intracranial
factor VII for, 512–513
traumatic, 510
and IVADs, 456–459, 457f
approach to, 457–459, 457f–458f
chemical cautery for, 458–459
dilysis clamps for, 458, 458f
direct pressure for, 457–458
suture for, 459, 459f
thrombogenic and dermal adhesive agents for, 458
vasoconstrictive agents for, 458
oral, 1396–1397
postpartum, 1205–1207, 1205f, 1206f
poststenoltemecy, 1383, 1383f
retrobulbar, 1333–1334, 1334f
subarachnoid, 1259
subconjunctival, 1337, 1337f
in wounds, control of, 631–632, 633f, 1528
cardiac, 346–347, 346f–347f
Hemorrhagic shock, 1426
infusion rate for, 520
Hemorrhoids
conservative treatment of, 908
external, 907f, 908
surgical excision of, 907f, 908–910, 909f–910f
internal, 904, 907f
management of, 907–910
Hemostasis, wound management and, 1528
Hemostatic agents, for hemorrhage control of, 632
in IVADs, 458
Hemothorax, 199
autotransfusion with, 492, 492f
with central venous catheterization, 436, 436f
symptoms of, 200
thoracentesis and, 195
tube thoracotomy for, 205–206, 206f
under sound in, 492, 492f
Heparin
central venous cannulation and, 1525
low-molecular-weight, 1524
spinal epidural hematoma and, 1524
Heparitin-associated coagulopathy, in IVADs, 459
Hepatitis B
occupational exposure to, 351–352, 1476
prevention of, sexual assault and, 1236
Hepatitis B immune globulin (HBIG), for hepatitis B, 1236
Hepatitis C
occupational exposure to, 351–352, 1477
transfusion transmission of, 502–503
Herniation
after lumbar puncture, 1270–1271, 1271b
of brain, 1243, 1245f, 1270
Heroin, body packing of, 870, 870f
Herpes zoster, dental pain and, 1387f
Herpetic keratitis, 1303
Herpetic whitlow, 765f, 767, 767f
Hexachlorophene, 627, 627f
for skin disinfection, 1453
Hexylresorcinol, 797
Hickman catheter, 447
HI-D Big Stick, 42, 42f
Hidradenitis suppurativa, 756–758, 757f
High-flow nasal cannula oxygen (HFNC), 48
High-flow prapism, 1146
High-frequency ventilation, 166
Hill-Sachs deformity, 984–985
Hindfoot, dislocations of, 1023
HINTS examination, for vertigo, 1288–1290, 1289f
complications of, 1290
contraindications to, 1288
interpretation of, 1289–1290
procedure for, 1288–1289
Hip
arthrocentesis of, ultrasound in, 1110–1114f, 1113f
corticosteroid injection of, 1095–1097
dislocation of, 1012–1016
analgesia and anesthesia for, 1013–1014
anterior, 1015–1016, 1016f
reduction techniques for, 1015–1016, 1017f
fractures with, 1012–1013
posterior, 1014, 1014f
Allen technique for, 1014, 1015f
Captain Morgan technique for, 1014, 1015f
reduction techniques for, 1014
Stimson technique for, 1014, 1015f
Whistler technique for, 1014, 1015f
radiographic examination of, 1013, 1013f
nerve blocks at, 562f
prostheses, dislocation of, 1014–1015, 1016f
Hognose device, 1357, 1357f
Homatropine, oculair, 1299–1301, 1299f
Honey, for burns, 787
Hordeolum, 1336, 1337f
Horizontal head impulse test, for vertigo interpretation of, 1289
procedure for, 1288–1289, 1289f
Horizontal mattress stitch, 679–680, 682f
Horse collar, 926f
Hot tar burns, 794–795, 794f–795f
Housemaid's knee, 1097
Huber needle, 448, 449f, 453
Human anthemophilic factor, 513
Human bites, 650–653, 651f, 733, 733f
tendon injury with, 965–966, 965f
β-human chorionic gonadotropin (β-hCG) testing, in sexual assault evaluation, 1234
in urine, 1450, 1451f, 1451f
Human immunodeficiency virus (HIV)
infection
CSF analysis in, 1279
cytomegalovirus infection and, 1280
Indwelling vascular access devices (IVADs) (Continued)
dialysis clamps for, 458, 458f
 direct pressure for, 457–458
 mechanical, 457
 suture for, 459
 thrombogenic and dermal adhesive agents for, 458
 vasoconstrictive agents for, 458
 historical perspective of, 447
 infection with, 454–456
 long-term venous access catheters in, 457–459
 midline peripheral catheters in, 449–450
 PICC in, 449, 449f–450f
 pinch-off syndrome in, 456
 steal syndrome with, 460
 temporary dialysis catheters in, 450–451, 450f–451f
 thrombus formation with, 456
 totally implantable, 448, 449f
 accessing, 453

Infants
arterial blood sampling in, 356–359, 358f–359f
 arterial catheterization in, percutaneous, 372–373
 bag blows for, 41
 bag-mask ventilation for, 46
 capillary blood sampling in, 353–354, 355f
 central venous catheterization in, 364–367
 chest thrusts for, 41
 direct laryngoscopy for, 72–75, 75f
 heart rate in, 5
 heel stick in, 353, 1454
 hemorrhagic shock and encephalopathy syndrome in, 1426
 intravenous infusion in, 463
 lumbar puncture in, 1266–1267
 with meningitis, 1259
 peripheral venous catheterization in “mini-cutdown”, 364f
 percutaneous, 359–361, 361f
 venous catheterization in, 361–364, 363f
 rehydration techniques in, 373–376
 discharge after, 376
 laboratory tests and, 375
 nasogastric tube, 375
 nasogastric tube rehydration and, 375
 oral, 374–375, 374f
 parenteral, 375
 subcutaneous, 375–376, 376f, 376t
 umbilical artery catheterization in, 370–371, 370f–371f
 umbilical vein catheterization in, 367–369, 368f–369f
 urine collection in, 1442–1443
 venipuncture in, 354–356, 356f

Infection (Continued)
tracheostomy and, 153
 in transfusion therapy, 502–503, 503f
 from urethral catheterization, 1168
 venous cutdown and, 445
 with Trousseau’s, 459, 637f, 653
 Infectious exposure management, 1470–1480
 Infectious keratitis, 1321–1322
 Infiltration anesthesia, 533–536, 533f
 agents for, 534, 534f
 buffering for, 535
 contraindications to, 533
 equipment for, 534–535
 field block anesthesia and, 533
 indications for, 533
 injection in, 535–536, 536f
 temperature manipulation in, 535
 Infraorbital nerve block, 551–552, 552f
 Infraorbital nerve, 551, 552f
 Infraspinatus muscle syndrome, 1073r, 1102f, 1113
 Infuse-A-Port, 448
 Inguinal hernia
 direct, 897, 898f
 indirect, 897a, 898f
 pantaloon, 898
 Inhalation injury, 791, 792f
 Inhibitor-bypassing activity, factor VIII, 513–514
 for anti-factor Xa anticoagulants, 518f
 for direct factor II inhibitors, 516–517, 517f
 for warfarin reversal, 515f–516t
 Injury classification system, 963f
 In-line filtration, in autotransfusion, 493
 Innominate artery, tracheostomy-related bleeding from, 154–155, 155f
 Inspiratory-to-expiratory time ratio, 164, 167f
 Insulin infusion devices, 1499, 1500e, 1500f
 Insulin-dependent diabetics, abscess and, 739, 740f
 Intercostal muscle syndrome, 1073r, 1102f, 1113
 Intercostal nerve block, 562f, 563–566, 564f
 Intercostal space number, 278
 Internal hemorrhoids, 904, 907–908, 907f
 International Consensus Conference on Cardiopulmonary Resuscitation and Emergency Cardiopulmonary Care, 41
 International normalized ratio (INR)
 bleeding and, 1520
 elevated, management of, 512f
 in routine coagulation testing, 1520
 in warfarin reversal, 516f
 Interphalangeal joint arthrodesis of, 1116–1117, 1117f, 1119, 1120f
 fingers, dislocation of
distal, 1005f, 1010, 1011f
 proximal, 1009–1010, 1009f
 dorsal, 1005f, 1009–1010, 1009f
 lateral, 1010
 volar, 1010
 foot, dislocation of, 1025–1026, 1025f
 thumb, dislocation of, 1006, 1006f
 toe, dislocation of, 1025–1026, 1025f
 Interscalene nerve block, 566, 582–587, 584f–585f
 Interscapular hypodermoclysis, 490f
 Intersection syndrome, 1090–1093, 1092f
 Intersphincteric abscess, 761, 761f
 Intervention, immediate, spirometry and, 23
 Intracranial hypertension idiopathic, 1295–1296
 rapid-sequence intubation and, 114–115
 Intracranial pressure (ICP)
 increased
 diuresis in, 1247
 fluid management, 1247
 glucose control in, 1248
 head position, 1247
 hyperventilation in, 1247
 hypothermia and, 1248
 management of, 1243–1257
 medical treatment of, 1246–1251
 operative management in, 1251
 oxygenation in, 1246
 paralytics in, 1246–1247
 pathophysiology of, 1243–1245, 1244f, 1244f
 sedation in, 1246–1247
 seizure prophylaxis in, 1247–1248
 signs and symptoms of, 1245–1246
 skill trephination and, 1248–1251
 1249f–1250f
 steroids in, 1248
 titration of ETCO2 in, 32–33
 in intubation, 113
 in rapid-sequence intubation, 116f
 tracheal suctioning and, 145
 Intracranial shunts, 1251–1257
 assessment of, 1253–1255
 catheter for, 1251, 1251f
 clinical presentation of, 1253
 components of, 1252f
 failure of, 1253
 infection with, 1253
 management of, 1243–1257
 principle of, 1251
 tapping of, 1255–1257, 1257f
 valves for, 1251–1252, 1252f
 Intranasal administration, of medication, 481–485, 482f
 Intravascular ligation, 690–693, 693f
 Intravenous infusion, 461–475
 anatomy for, 462–463, 462f
 blood transfusion through, 520
 Bone Injection Gun (BIG) for, 465, 465f, 468–470, 471f
 comparative trials between devices, 466
 compartment syndrome with, 475
 complications of, 462f, 471–475
 bony, 474–475
 soft tissue, 474–475
 technical difficulties, 471–474, 474f
 contraindications to, 462f, 464
 distal femur for, 467
 distal tibia for, 467, 467f
 duration of use for, 475
 epiphyseal injury with, 475
 equipment for, 462f, 464–467
 EZ-JO Device for, 465–466, 466f
 EZ-JO needle for, 470–471, 472f–473f
 FAST-I Intravenous Infusion System for, 465, 465f, 468
 fat embolism with, 475
 historical perspective on, 461–462
 humerus for, 467–468
 indications for, 462f, 463–464
 in infants, 463
 infection with, 474, 475f
 infusion rate in, 462
 manual needle insertion for, 468, 469f
 medications and fluids for, 463f
 needles for, 464–465, 464f
 NIO device for, 466, 466f, 471, 473f
 pain with, 475
 physiology of, 462–463
 proximal tibia for, 467, 467f
 setup for, 464–467
Intravenous access (Continued)
simulation studies comparing devices, 466–467
sites for, 467–468, 467f
skin sloughing with, 474–475
sternum for, 468
training for, 475
in US military, 463–464
Intrathoracic pressure, increased, spirometry and, 23
Intravaginal testicular torsion, 1142
Intravascular hemolysis, 1447f
Intravascular injection, of anesthetics, 563
Intravenous access
transillumination of, 395
preparation of, 396–400
percutaneous brachial vein cannulation, 399–400
positioning of, 397–399
preparation of, 396–400
safety, 396–397
transillumination of, 395
ultrasound guidance of, 395
Intravenous catheter safety device, 396f
Intravenous contrast with CT, esophageal
Intravenous fluid rate, maintenance, calculation of, 1532–1533
Intravenous regional anesthesia. see Regional anesthesia, intravenous
Intubating laryngeal mask airway, 62, 91, 92f
flexible endoscopic intubation through, 95–97
types of, 92
Intubation
aphneic oxygenation during, 65
tracheal, 62–110
Inverse ratio ventilation, 166–167
Inverting horizontal mattress stitch, 681
IO catheterization, 371–372, 372f
Iodine compounds, for skin disinfection, 563
Iodine solution, 627, 627r
Iontophoresis, 530
IR radiation, capnography and, 30
Iron overdose, deferoxamine for, 468
Irradiation, prenatal, 1505, 1506f
K
Katz Extractor, for nasal foreign body removal, 1357–1378, 137ff
Kendrick Extrication Device (KED), 921f, 922
Keratitis
air bag, 796–797, 796f
herpetic, 1303
infectious, 1321–1322
Kernig’s sign, 1259
Ketamine in cardiodversion, 255–256, 256r
contraindications to, 616
for ear canal foreign body removal, 1356
in ear procedures, 1351
emergence reaction with, 118–119, 616
etomidate or, 117
intranasal, 481
in procedural sedation and analgesia, 608f–609f, 611–612r, 615–617
for rapid-sequence intubation, 117–119, 117r
for restraint, 1490r, 1495
Ketones, in urine, 1444, 1445r
Ketorolac, in heat strokes, 1432
Kilograms, metric conversions of, 1530r
King laryngeal tube, 58–60, 60f
orotracheal intubation with a, 101–102, 102f
Kleihauer-Betke test, 507
Knee
arthrocentesis of, 1118–1119, 1119f
ultrasound in, 1110–1114, 1110f–1111f
corticosteroid injection of, 1097–1098
dislocations of, 1016–1020
anterior, 1018f
clinical assessment of, 1017f, 1018 lateral, 1018f
posterior, 1018f
post-reduction care for, 1020
reduction technique for, 1019–1020, 1019f
vascular injury with, 1018–1019, 1018f
nerve blocks at, 562r
splints for, 1044–1046
immobilizer as, 1044, 1048f
Jones compression dressing as, 1046
posterior, 1044–1046, 1048f
Knee extensor tendon rupture, 978–979
Korokoff sound, 7, 8f
Kussmaul respirations, 3–4, 4f
Labor, 1186–1187
betamethasone in, 1194
cervical dilation in, 1190
cervical effacement in, 1190, 1190f
delivered, anticoagulation and, 1528
evaluation of, 1187–1188, 1188f
fetal distress during, 1191–1193
fetal heart rate in, 1187, 1187f, 1191, 1192f
fetal station in, 1190–1191, 1191f
indomethacin in, 1193, 1193r
ketorolac in, 1194
Laryngectomy, 158
Laryngoscopy, 69–70, 69
Lactating mothers, 1507–1508

Laryngoscopy (Continued)
contraindications to, 1338
direct, 67f, 69–85, 73f
contra indications to, 67f, 69
equipment for, 67f
indications for, 1338
learning, 72, 74f
optimal patient positioning for, 71–72, 71f
procedure and technique of, 72–85
equipment for, 1338
flexible, 1338–1339, 1341f
indications for, 1338
mirror, 1339–1342, 1342f
video, 85–95, 86f, 89f
with angulated blades, 87–90, 87f
with standard Macintosh blades, 85–87, 86f
with tube channel, 90–92, 90f
Laryngospasm
in nasotracheal intubation, 104
in tracheal intubation, 80
Larynx, 1338–1342, 1339f
examination of, 1338
pediatric co. adult, 127, 128f
Lateral neck stabilizers, 924, 924f
Lavage
in gastrointestinal tract bleeding, 830
lung, hyperthermia, 1432
acidity; curettage, 1435
Leakage, gastroenterostomy feeding tube and, 849
Left bundle branch block (LBBB), transvenous cardiac pacing in, 290
Left ventricular assist devices, 262, 263f
blood pressure and, 9
complications of, 270–271
LEMON score, 127–129, 129f
Leonard catheter, 447
Leukemia, lumbar puncture in patients with, 1260
Leukocyte esterase, in urine, 1444, 1445r
Levator scapulae muscle syndrome, 1073r, 1101, 1102f
Levin tube, 828, 836f
Levonorgestrel, 1236–1237
Lewis, Thomas, 275
Lewis leads, for electrocardiography, 281, 283f, 283r
Lidocaine, 525–526r
alkalinize, 353
bupivacaine and, 534, 534r
dorotrahal drug administration of, 477r
epinephrine and, 534
in increased intracranial pressure, 1246
infiltration of, 534, 534r
in injection therapy, 1078
in intact skin, 529–530
in intravenous regional anesthesia, 588–590, 592
mucosal application of, 527, 527b, 527r
for nasal anesthesia, 1361
for phimosis, 1157f
for prapitism, 1148, 1148r
in regional anesthesia, 560
for shoulder dislocation, 986, 989f
spinal puncture and, 1262
for toothache, 1386
viscous, 529
Lorazepam
in increased intracranial pressure, 1247–1248
Lorazepam solution, 512
Life-threatening airway obstruction, 157
Ligamentum flavum, 1264, 1265f
Ligation, of veins, 632, 633f

Light’s criteria, 193, 193b
Limb injury, in regional anesthesia, 563
Limb-precordium leads, for electrocardiography, 283
Lims, escharotomy for, 804–805, 804f
Lingual nerve, 542f
Lip laceration, 690–693, 692f
Lipid emulsion, intravenous, local anesthesia and, 542, 542b
Lisfranc fracture-dislocation, 1023, 1025f
Liver
thoracentesis-related puncture of, 195
ultrasound of, 1434, 1435f, 1440f
Lobar capillary hemangioma, 733–734, 734f–735f

Local anesthesia, 523–544
acid-base status and, 541
agents for, 523
active for, 525
activity profile of, 525–526, 525r
clearance of, 540
dose of, 523, 524r, 540, 540r
inadvertent intravascular injection of, 540–541
mechanism of action of, 525
pK, of, 525, 526f
potency of, 526
protein binding of, 526, 526r, 541
skin testing of, 543
toxicity of, 539
allergic reactions with, 542–543
catecholamine reactions with, 543
duration of, 526, 526f
epinephrine and, 526, 534, 535b
hematoma block in, 536–537, 536f
in hemorrhoid excision, 908, 910f
high blood levels in, 539–541
inadvertent intravascular injection of, 540–541
injuries in, 538
intratracheal, 537
intrapleural, 537
maximum safe dosage for, 540, 540r
mechanism of action of, 525
nerve structure and, 524–525, 524f
onset of action of, 525–526, 525r
pH and, 526, 535
potency of, 526, 526r
in spinal puncture, 1267
systemic toxic reactions of, 538–542, 539r
cardiovascular, 541
CNS, 541
high blood levels and, 539–541
host factors and, 541
lipid emulsion in, 542, 542b
prevention of, 541
Treatment of, 541–542
for toothache, 1387
for tube thoracostomy, 209–210, 210f
vasoergic reactions with, 543–544, 543f–544f
wound healing in, 537–538
wound infection in, 538

Local injection therapy
contraindications to, 1074b
equipment for, 1078, 1078f
Local twitch response, 1071
Locked scalpula, 997
Logroll maneuver, 927, 928f
Long arm splint
anterior, 1037–1038
posterior, 1036–1037, 1037f
Lorazepam
in increased intracranial pressure, 1247–1248
in neuroleptic malignant syndrome, 1425
for restraint, 1490r, 1493–1494
Ophthalmologic procedures (Continued)
eyes
- dilating, 1298–1302
- irrigation of, 1305–1308
- patching, 1316
- fluorescein examination, 1302–1305
globe
- luxation of, 1334–1336, 1336f
- protection, evaluation and, 1309, 1309f
- infectious keratitis, 1321–1322
- Marcus Gunn pupil, APD or, 1336–1337, 1337f
- ocular FB removal, 1334, 1335
- with tube channel, 90–92
- ear canal débridement and wick placement for rapid-sequence intubation, 120
Optic nerve
- cantholysis for, 1334, 1335
- for increased intracranial pressure, 1246
- Oxyhemoglobin for CSF, 1272
- Oxyhemoglobin dissociation curve, 25–26, 26f
Oxytocin, 1199–1200, 1205–1207, 1205f

P
Pacemaker
- characteristics of, 258–262, 259f, 260f–261f
- complications of, 268–269, 268b
defibrillation and, 243
- electrocardiography of, 265
- electromagnetic interference and, 272, 273b
- failure to capture of, 269
- failure to sense of, 269
- indications for, 262–263
- malfunctions of, 268–269
- output failure of, 268f
- radiography of, 260f–261f, 265
- recalls of, 272
- response to magnet placement, 263–264, 264f–265f
tachycardia with, 269
- in twiddler's syndrome, 271–272, 272f
- Pacing bradycardic rhythms, for transcutaneous cardiac pacing, 306–307
- Pacing catheters, in transvenous cardiac pacing, 292
- Pacing generator, in transvenous cardiac pacing, 292, 292f, 293b
Paddling
- for immobilization, 924, 924f
- for splitting, 1029–1030, 1031f, 1034–1035, 1034b
Pain
- in burn injury, 784
- from cast, 1055
- dental, 1386, 1387f
- heel, 1057–1058, 1100–1101, 1100f
- with intraosseous infusion, 475
- management of, 483
- as vital sign, 21–22
- Palm laceration, 976f–977f
- Palmis longus tendon, 569, 571f
- Pancuronium
- in mechanical ventilation, 171
- for rapid-sequence intubation, 121r, 123
- Papain
- for esophageal foreign bodies, 816
Paper speed, 278
in electrocardiography, 281, 282f
Paracetamol, 884–896, 885f
- albumin in, 895
- anticoagulation and, 1526
- cell count in, 895, 895r
- chronic ambulatory peritoneal dialysis and, 896
- complications of, 893–894
- intraperitoneal, 894
- local, 894
- systemic, 893–894
- culture in, 895
- Paracresis (Continued)
- disposition and, 896
- Gram stain in, 895
- indications and contraindications to, 884–886
- anatomic, 886
- systemic, 886
- inspection in, 894–895
- interpretation of, 894–895, 894b
- medical therapy and, 896
- miscellaneous in, 895
- preliminary actions in, 886
- procedure for, 887–889, 887f–889f
- site of entry in, 886–887, 887f
- in spontaneous bacterial peritonitis, 896
- technique of, 886–887
- ultrasound guidance for, 889, 8918–893b, 892f–893f
- volume of fluid removed with, 890, 890f–891f
- Paradichlorobenzene poisoning, 1461
- Paradoxical bradycardia, 15–16
- Paradoxical undressing, 1406
- Paralysis, succinylcholine and, 122
- Paralytics, evaluation of airway before using, 124
- Paraphimosis, 1151–1155, 1151f
- anatomy of, 1152
- dorsal slit procedure for, 1156, 1159f
- iatrogenic, 1151
- key procedure sequence of, 1152
- pathophysiology of, 1152
- reduction, 1153f
- adjunctive techniques to assist, 1152–1155, 1154b, 1154f
- aftercare for, 1155
- complications of, 1155
- foreskin edema, 1154, 1154b
- manual, 1152, 1153f
- Parapneumonic effusion, 181, 195, 195b
- Parasitic infection, transfusion transmission of, 502
- Parasympatholytics, ocular, 1299f
- Parenteral rehydration, in infants and children, 374–375, 374f
- Osteitis, alveolar, 1397–1398
- Otitis externa
- ear canal debridement and wick placement for, 1355, 1356f
- malignant, 1352f, 1355
- Otolaryngologic procedures, 1338–1383
- “Outrigger” finger splint, 1042, 1045f
- Overdrive pacing, in ventricular tachycardia, 1057–1058
- Oversensing, 269
- Over-the-needle catheter cannulation, 383, 384f
- Overwear syndrome, with hard contact lenses, 1317
- Oxygen
- high saturation of, pulse oximetry and, 29
- hyperbaric, for frostbite, 1419
- inspired, fraction of, 163
- supplemental, in procedural sedation and analgesia, 600
- therapy, 45–49
- during cardiac arrest, 45, 46b
- complications of, 48–49
- indications and contraindications, 45
- during neonatal resuscitation, 45, 46b
- oxygen delivery devices for, 45–48
- Oxygen (Continued)
- partial non-rebreather mask for, 47
- preoxygenation prior to endotracheal intubation in, 48
- procedure for, 48
- simple masks for, 46–47
- standard flowmeter for, 47, 47f
- Venturi mask for, 47–48, 47f
- toxicity, 49
- warm humidified, in hypothermia, 1413
- Oxygenation
- assessment of, devices for, 23–37
- in decreased intracranial pressure, 1246
- Oxyhemoglobin for CSF, 1272
- Oxyhemoglobin dissociation curve, 25–26, 26f
Oxytocin, 1199–1200, 1205–1207, 1205f
Pressure sores, from splinting, 1053–1054, 1054f
Pressure ulcers, cervical spine immobilization and, 930
Pressure-cycled ventilation, 165
Pressure(s), 1113, 1269, 1272
Preterm premature rupture of membranes (pPROM), 1186
Priapism, 1145–1150, 1145f
anatomy of, 1146, 1146f
key procedure sequence of, 1146
pathophysiology of, 1146
stuttering, 1147

treatment of, 1147–1150, 1147f
aftercare for, 1150
aspiration/irrigation technique for, 1148–1150, 1148f, 1149f–1150f
complications of, 1150
contraindications to, 1147
indications for, 1146–1147
minimally invasive technique for, 1148, 1148f, 1149f

Prilocaine, 525r–526f
Procainamide
in malignant hyperthermia, 1424–1425
in supraventricular tachycardia, 233

Procare, 525r–526r
infiltration of, 533r–534r, 534

Procedural sedation and analgesia (PSA)

(Continued)
renal evaluation for, 598
respiratory evaluation for, 596
routes of administration for, 602–603
sufentanil in, 615
supplemental oxygen with, 600
terminology for, 594, 595f
thiopental in, 611r–612r, 614

Prochlorperazine, rectal administration of, 487r

Procidentia, 913, 914f
Profound multifocal leukoencephalopathy,
1280

Prolapse
cord, 1191–1193
rectal, management of, 913–915, 914f
Promethazine, rectal administration of, 487r, 488

Pronation method, in nursemaid’s elbow reduction, 1004

Proparacaine, ocular, 1296, 1298r

Prophylactic measures, for infection, in IVADS, 453–456

Propofol in
awake intubation, 125
in cardiovension, 255–256, 256r
in increased intracranial pressure, 1246
in procedural sedation and analgesia, 607b, 611r–612r, 613–614, 613f
in rapid-sequence intubation, 117r, 119

Proport, 448

Propranolol
dentrochael drug administration of, 478
in supraventricular tachycardia, 234

Prosthetic valves, reversal of anticoagulation, 1522–1523

Protective gear, for splinting, 1030

Protein
binding, in local anesthesia, 525r–526r, 526, 541

CSF, 1274
in synovial fluid, 1121
in urine, 1444–1446, 1445r, 1447b

Proteinuria, 1444–1446, 1447b

Prothrombin complex concentrate, 514
for anti-factor Xa anticoagulants, 518r
for warfarin reversal, 515, 515r–516r

Prothrombin time (PT), in routine coagulation testing, 1520

Pruritus in burn injury, 785
with burns, 785
from splinting, 1054, 1054f

Pseudoneurocyrrus, in IVADS, 456–457

Pseudomonas infection of eyes, 1318
of foot, 1066, 1066f
Pseudothrombophlebitis syndrome, 1098
Pseudotumor cerebri, 1243, 1244f, 1259–1260

Psychiatric disorder, agitation caused by, 1496
Psychological support, for sexual assault, 1237

Psychostimulant overdose, 1425–1426
Pterygium, 1332
Puric hair samples, in sexual assault, 249–250
Purpura, 624, 648, 649f, 973, 976f, see also Wounds
Porcine cerebral, 1243, 1244f, 1259–1260

Pulmonary artery catheter, 409
misplacement in, 297–298

Pulmonary edema after thoracentrysis, 195
after tube thoracostomy, 220
noninvasive positive pressure ventilation in, 168

Pulmonary embolism (PE), diagnosis of, during pregnancy, 1509–1510, 1510b–1511a, 1510f

Pulv, disease of, 1398–1399
Pulvitis, 1398–1399

Pulse, 5–6
apical, 5–6
blood pressure and, 10
bounding, 6
during cardiopulmonary resuscitation, 6
checks, in CPR, 332–333

Pulse oximetry, 25–29, 407–408
clinical applications of, 26b
clinical utility of, 26
indications for, 26, 27f
interpretation of, 26–28
limitations of, 27–28
phyiology in, 25–26
probes for, 26, 27f
site of, 29
in procedural sedation and analgesia, 598
procedure for, 26
technology for, 25, 25f

Pulse plethysmograph, 26, 28f
Pulse pressure, 9–10
Pulse rate, 5

Pulseless electrical activity, pericardioenteesis in, 323

Pulseless ventricular tachycardia (VT)
CPR in, 241–242
defibrillation in, 243
Pulselessness, duration of, emergency department thoracotomy and, 340
Pulse-pressure variation, 10
Pulsus paradoxus, 10–11, 11f, 316–317, 317f
Puncture wounds, 624, 648, 649f, 973, 976f, see also Wounds
Corneal, 1316
foot, 639f, 1062–1063
sole of, 717–718, 718f

Pupil. see also Eyes
afferent defect of, 1336–1337, 1337f
dilating, 1298–1302
agenurosis, 1299–1301, 1299t, 1301f
complications of, 1301–1302, 1302b
contraindications to, 1299, 1300f
indications for, 1299, 1300f
procedure for, 1301

Marcus Gunn, 1336–1337, 1337f
Pupillary response, in brain death testing, 1284

Purpura, posttransfusion, 505
Purulent meningitis, empirical therapy for, 1277r
Pyogenic granuloma, 733–734, 734f–735f

Q
QT interval, 1529, 1530r

Qp/c interval, 1529

Quadriceps tendon rupture, 978
Quenchenstedt test, 1272
Quinton catheter, 450–451

R
Raf catheter, 447

Rabies, prophylaxis for, 650, 652r
Radial artery
for arterial cannulation, 387–389, 387f
arterial puncture of, 381f
Radiotherapy

in acromioclavicular joint subluxation and dislocation, 998–999
in ankle dislocations, 1022–1023, 1022f–1023f
in anterior shoulder dislocations, 984–986, 986f–988f
in central venous catheterization, 429, 429f
in dislocation, 980–981
in esophageal foreign bodies, 809–810, 809f–811f
of foot, 1063, 1063f
in foreign body removal, 708–709, 711f, 713f–714f
in hand injuries, 1005, 1005f
in hip dislocation, 1013, 1013f
in intracranial shunt, 1254, 1255f–1256f
in joint injury, 702, 704f
of left ventricular assist devices, 266f
in nursemaid's elbow, 1003
of pacemaker, 260f–261f, 265
in patellar dislocation, 1021
in posterior elbow dislocations, 1000, 1000f
in posterior shoulder dislocations, 994–996, 994f–996f
in sternoclavicular dislocation, 999f
Radiohumeral bursitis, 1078f, 1085–1086
Radiophysics, 1504–1505, 1505f
Raney clamps, in scalp bleeding, 632, 635f
Rape trauma syndrome, 1237
Rapid Rhino device, 1367, 1367f–1368f
Rapid-sequence intubation (RSI), 112f
atracurium for, 121, 123
awake intubation vs., 124
decision to perform, 68, 68h
etomidate for, 116–117, 117r
fentanyl for, 117f, 120
in increased intracranial pressure, 1246
intracranial hypertension with, 114–115
ketamine for, 117f–119, 119f
methohexital in, 117r
midazolam for, 117f, 119–120
mivacurium for, 121f, 124
for neuromuscular blocking agents, 120–123
neuromuscular blocking agents for, 121f
nondepolarizing agents for, 122–123
overview of, 111–113
paracurium for, 121f, 123
preoxygenation in, 64–65, 111, 112f
Rapid-sequence intubation (RSI) (Continued)
pressor response in, 113–114
prophyl for, 117f, 119
protocols in, 111–113, 113b, 116b
rocuronium for, 121f, 123
"second look," airway evaluation before, 124
succinylcholine for, 121–122, 121r
thiopental in, 116b, 117r
vecuronium for, 121f, 123
β1-Receptor agonists, in preterm labor, 1193–1194
Receptor 1507, 1506
Recumbent, 760, 761f
anatomy of, 904
digital examination of, 904
foreign body in, 912f–913f
management of, 910–912
removal of, 910, 913f
prolapse complications of, 915
indication for, 915
management of, 913–915, 914f
procedure for, 915
reduction of, 914f, 915
Rectus abdominis syndrome, 1073f, 1102f, 1103
Red blood cells
absolute number of, 1274
antigens and antibodies of, 500, 501f
clearance from first to last tubes, 1274
count, in diagnostic peritoneal lavage, 883, 883f
in CSF, 1272
preparations of, 500–502, 502f
irradiated, 502
leukocyte-reduced, 502
packed red blood cells in, 501
wash, 501–502
whole blood in, 500
urinary, 1440f
Reese shoe, 1052, 1052f
Regional anesthesia
in burn injury, 784
of head and neck, 545–559, 545f, 546f
Akinoi-Vazirani block for, 555, 555f
anatomy for, 545
anterior middle superior alveolar block for, 551, 551f
anterior superior alveolar nerve block for, 550–551, 550f
complications of, 545f
contraindications to, 545f
equipment for, 545–546, 545f
general recommendations of, 546–558,
Gow-Gates block for, 554–555, 554f
indications for, 545f
inferior alveolar nerve block for, 552–554, 553f
infraorbital nerve block for, 551–552, 552f
mental nerve block for, 555–556, 555f–556f
middle superior alveolar nerve block for, 549–550, 550f
occipital nerve block for, 557, 557f
ophthalmic (V1) nerve block for, 557–558, 558f
posterior superior alveolar nerve block for, 546–549, 549f
scap, 556–557, 556f
supraperiosteal injection for, 546, 549f
Regional anesthesia (Continued)
intravenous, 588–593, 588f
agents for, 592–593
complications of, 593
contraindications to, 588–589, 588f–589f
equipment for, 589, 589f
exanguination for, 591
indications for, 588–589, 588f–589f
mechanism of action of, 592
procedure for, 589–592, 591f, 592b
site of injection for, 593
of lower extremity, 560–587
agents for, 560
at ankle, 562f, 577–581, 578f–580f
complications of, 562–563
deep peroneal nerve block, 562f,
579f–580f, 580–581
digital nerve block, 569–572
equipment for, 560
femoral nerve block, 562f, 575–577, 576f
hematoma with, 563
indications for, 560
infection with, 563
injection for, 561–562
instructions for, 560
intercostal nerve block and, 563–566,
564f–565f
intravascular injection with, 563
lumbar injection with, 563
metatarsal, 562f, 581–587, 581f–582f
nerve injury with, 562–563, 562r
paresthesia for, 561
patient positioning for, 560–561
peroneal nerve block, 562f
posterior tibial nerve block, 562f,
578–580, 579f–580f
precautions for, 562–563
preparation for, 560
saphenous nerve block, 562f, 579f–580f,
581
sites for, 561
superficial peroneal nerve block, 562f,
579f–580f, 580
sural nerve block, 562f, 579f–580f, 580
systemic toxicity with, 563
tibial nerve block, 562f
at toes, 562f, 581–587, 581f
ultrasound for, 561, 582b–587b,
583f–587f
of thorax, 560–587
agents for, 560
complications of, 562–563
equipment for, 560
hematoma with, 563
indications for, 560
infection with, 563
injection for, 561–562
instructions for, 560
intercostal nerve block and, 563–566,
564f–565f
interscalene nerve block and, 582b–587b,
584f–585f
intravascular injection with, 563
lumbar injection with, 563
nerve injury with, 562–563, 562r
paresthesia for, 561
patient positioning for, 560–561
precautions for, 562–563
preparation for, 560
sites for, 561
systemic toxicity with, 563
ultrasound for, 561, 582b–587b,
583f–587f
of upper extremity, 560–587
agents for, 560
complications of, 562–563
Respiratory rate, 3
abnormal, 3–4, 4f
in children, 1, 2f
in mechanical ventilation, 163

Respiratory alkalosis, 1534, 1535
Respiratory acidosis, 1534, 1534
Respiration, 2–4
Resolution, degree of, 1434

Resuscitation, principles of, 239
Resuscitative thoracotomy, 338–352, 338f
airway control in, 343
anesthesia and amnesia and, 343
anterolateral thoracotomy incision in,
343–345, 344f
aortic cross-clamping in, 349–351, 349f
blunt and penetrating abdominal injury
and, 342
and cardiopulmonary resuscitation,
duration of, 340
in children, 351
complications of, 351–352, 352f
Core base expander in, 351–351, 350f
direct cardiac compressions in, 345–346,
345f
equipment for, 342
Foley catheter in, 347, 348f
general technique for, 344f
hemodynamic monitoring in, 351
hemorrhagic cardiac wounds in, control of,
346–347, 346f–347f
hemorrhagic great-vessel wounds in,
control of, 349, 349f
indications and contraindications to,
339–341, 339f
airway analgesia, 341–342
blunt and penetrating abdominal injury
as, 342
blunt cardiac injuries as, 341
nontraumatic arrest as, 342

Resuscitative thoracotomy (Continued)
nontraumatic hypothermic cardiac arrest
as, 342
penetrating cardiac injuries as, 340–341
pulmonary injuries as, 341
internal cardiac defibrillation and, 345
neurologic outcomes and, 340
partial-occlusion clamps in, 347, 349f
pericardiostomy and, 345
procedure for, 342–351
Sauerbruch maneuver in, 346–347, 347f
Retardation, in fetal period, 1507
Retinoblastoma hemorrhage, 1333–1334, 1334f
Retrocaecal abscesses, 1057–1058, 1101
Retroglottic airway devices, 58–61, 59f
Retrograde cystogram, 1178, 1179–1181,
1182f–1183f
Retrograde intubation, 106–109, 107f
comlications of, 106, 108
contraindications to, 106
equipment for, 106
indications for, 106
procedure for, 106–107
technique of, 106–107
Retrograde technique, for fishhook removal,
721, 721f
Retrograde urethrograph aftercare for,
1181
anatomy of, 1177
comlications of, 1181
contraindications to, 1178
indications for, 1177–1178
key procedure sequence of, 1177
pathophysiology of, 1177, 1178f–1179f
procedure for, 1178–1179, 1178s, 1180f
Return of spontaneous circulation (ROSC),
capnography and, 32
Reverse Bigelow technique, for anterior hip
dislocation, 1016
Reverse Wood's screw maneuver, 1200, 1201f
Rewarming methods
active core, 1413–1416, 1413f
arteriovenous anastomoses, 1412
bladder irrigation for, 1414
cardiac bypass for, 1415–1416
experimental, 1416
external
active, 1411–1413
passive, 1411
for frostbites, 1419
gastric irrigation for, 1414
heated humidified oxygen/air for,
1413–1414
heated saline for, 1413, 1413f
hemodialysis for, 1416
peritoneal dialysis for, 1410f, 1414
prehospital care, 1409, 1410f
rate of, 1411, 1411f
thoracic cavity lavage for, 1415
warm air inhaled for, 1411f
Rh compatibility, 507
Rh immune prophylaxis, 507
Rh sensitization, 507
Rhabdomyolysis, from drugs, 1125, 1126f
RhOGAM, 507
Rib fractures, intercostal nerve block for,
563, 564f
Riddell Revolution football helmet, 945f
Right bundle branch block (RBBB),
transvenous cardiac pacing in, 290
Right-sided heart catheterization,
complications of, 301–303, 303f
Rigid splints, 931, 932f, 933
Ring block, 1156, 1158f
Ring cutter, 727, 729f
Ring removal, 727, 728f
Rigten maneuver, 1199f
Sutures (Continued)
techniques for, 667–685, 668f
tension and, 670–671
tie in, 669f
in tongue lacerations, 693–694, 693f
for tube thoracostomy, 212, 215f
undermining and, 670, 670f
Swathe and sling, 1043
Sweat glands, abscess in, 756, 757f
Swimmer's ear, 1355
“Swirl” sign, 1246f
Swiss roll technique, for paronychia, 765
Synchronized intermittent mandatory ventilation, 165–166
Syncope, in carotid sinus syndrome, 225–226
Sympathetic ganglion, 1143–1145
Synthetic dressings, 783–784
T
Tarsal burns, hot, 794–795, 794f
Tape, wound, 655–657, 656f
Tear ducts, removal of, 732, 733f
darts, removal of, 732, 733f
Tattooing, traumatic, 722–723, 723f
Teeth, 1384, 1385f–1386f
avulsion of, 1391–1393, 1393f
deciduous, 1384, 1385f
dry socket of, 1397–1398
emergency materials for, 1403, 1403f
extraction of, bleeding after, 1396
fractures of, 1387–1390
calcium hydroxide for, 1388, 1389f
Ellis class I, 1387, 1388f
Ellis class II, 1387–1389, 1388f
Ellis class III, 1385f–1389f
soft tissue laceration with, 1387, 1387f
intrusion of, 1391–1393, 1391f
luxation of, 1390–1391, 1390f
pain in, 1386
permanent, 1385f
reimplantation of, 1391–1392, 1393f
subluxation of, 1390–1391
supernumerary, 1384
Tegaderm, 641
Temperature, 17–21
contraindications to, 18
conversion scale for, 20f
core... afterdrop of, 1412
measurement of, 1406–1407, 1406f
ear, 20r
indications for, 18
Temperature (Continued)
infiltration anesthesia and, 535
interpretation of, 19–21
measurement sites for, 18–19
monitoring of, 20
normal, 17–18, 20r
oral, 18, 20r
physiology of, 17–18
pulse rate and, 5
rectal, 18, 20r, 1406f, 1407
respiratory rate and, 2–3
Temporal arteries, for arterial cannulation, 392–394
Temporal dural arteriovenous fistula, 940–942
Temporal arteries, for arterial cannulation, 392–394
Temporal arteries, for arterial cannulation, 392–394
Temporal arteries, for arterial cannulation, 392–394
Temporal arthritis, 1100–1101, 1100f
Tachycardia
Systolic arterial blood pressure, 7–8
Systemic toxicity, with local anesthetics, 563
Syphilis, transfusion transmission of, 502
Syncope, in carotid sinus syndrome, 225–226
Synchronized intermittent mandatory ventilation, 165–166
Sweat glands, abscess in, 756, 757f
Swimmer's ear, 1355
“Swirl” sign, 1246f
Swiss roll technique, for paronychia, 765
Synchronized intermittent mandatory ventilation, 165–166
Syncope, in carotid sinus syndrome, 225–226
Sympathetic ganglion, 1143–1145
Synthetic dressings, 783–784
Tarsal burns, hot, 794–795, 794f
Tape, wound, 655–657, 656f
Tear ducts, removal of, 732, 733f
darts, removal of, 732, 733f
Tattooing, traumatic, 722–723, 723f
Teeth, 1384, 1385f–1386f
avulsion of, 1391–1393, 1393f
deciduous, 1384, 1385f
dry socket of, 1397–1398
emergency materials for, 1403, 1403f
extraction of, bleeding after, 1396
fractures of, 1387–1390
calcium hydroxide for, 1388, 1389f
Ellis class I, 1387, 1388f
Ellis class II, 1387–1389, 1388f
Ellis class III, 1385f–1389f
soft tissue laceration with, 1387, 1387f
intrusion of, 1391–1393, 1391f
luxation of, 1390–1391, 1390f
pain in, 1386
permanent, 1385f
reimplantation of, 1391–1392, 1393f
subluxation of, 1390–1391
supernumerary, 1384
Tegaderm, 641
Temperature, 17–21
contraindications to, 18
conversion scale for, 20f
core... afterdrop of, 1412
measurement of, 1406–1407, 1406f
ear, 20r
indications for, 18
Thoracentesis (Continued)
clopidogrel, risk of bleeding in patients taking, 1527
complications of, 187f, 195f
contraindications to, 185–186, 187f
equipment for, 186, 187f
indications for, 185, 187f
insertion site for, 187–188, 190f
needle aspiration technique for, 190
over-the-needle catheter insertion technique for, 191, 192f
in parapneumonic effusion, 195, 195f
patient position for, 187–188, 188f
pediatric, 191
technique for, 186–191
ultrasound for, 187–188, 188f–190f,
189f–1906
Thoracic cavity lavage, in hypothermia, 1415
Thoracic duct laceration, catheter-related, 437
Thoracic spine, injury to, 917
Thoracolumbar spine, immobilization of complications of, 930
full-body spine boards for, 922, 923f, 927–929
full-body splints, 923–924, 923f
procedure for, 926–929
in recumbent position, 926–929
scoop stretchers for, 922–923, 923f,
926–927
in sitting position, 926
in standing position, 929
Thread tourniquet, hair, 734, 735f
Three-bottle system, in autotransfusion, 494–495, 495f
Three-in-one block. see Femoral nerve block
Thrombin, topical
in oral hemorrhage, 1397, 1397f
as topical hemostatic agent, 1528
Thrombocytopenia, lumbar puncture and, 1525
Thrombogenic agents, for hemorrhage, in IVADs, 458
Thrombolysis, for frostbite, 1419
Thoracolumbar spine, immobilization of complications of, 930
full-body spine boards for, 922, 923f, 927–929
full-body splints, 923–924, 923f
procedure for, 926–929
in recumbent position, 926–929
scoop stretchers for, 922–923, 923f,
926–927
in sitting position, 926
in standing position, 929
Thrombosis
anticoagulation and assessment of, 1520–1521
risk factors in, 1522
in arterial cannulation, 392
central venous catheterization and, 435b, 437
transvenous cardiac pacing and, 301
Thrombosed external hemorrhoids, surgical excision of, 908–910, 910f
Thrombosis
anticoagulation and assessment of, 1520–1521
risk factors in, 1522
in arterial cannulation, 392
central venous catheterization and, 435b, 437
transvenous cardiac pacing and, 301
Thrombotic occlusion, after arterial cannulation, 391
Thrombus formation, in IVADs, 459
Femoral nerve block
Thoracic spine, injury to, 917
Thoracic spine, injury to, 917
Thoracic spine, injury to, 917
Thoracic spine, injury to, 917
Thoracic spine, injury to, 917
Tick removal, 730–731, 731f
Ticlopidine, spinal puncture bleeding and, 1261
Tight lens syndrome, 1321
Tilt testing, 15f
Tinel’s test, 1091f
Tissue adhesive (tissue glue), 657–661, 659f–660f
combinations of, 660–661
procedure for, 659–660, 660f
in toothache, 1386
Tocolytic therapy, 1193, 1193f
Tourniquet
in extensor tendon injury, 962
hair-thread, 734, 735f
in hemorrhage, control of, 632–634, 636f
in intravenous regional anesthesia, 589–590, 591f
in toenail removal, 1068–1069, 1069f
for venous cutdown, 442, 443f
Tonic epidermal necrosis (TEN), 800–803, 802f
clinical features of, 801–802
management of, 802–803, 802r
Tonic shock syndrome
after nasal puncture, 1372
splitting and, 1054
Toxins
catfish, 725–726
coeleterates, 724
stingray, 726
Toxoplasmosis, 1279–1280
Trachea
anatomy of, 142, 143f
injury to, 341
physiology of, 142
stent for, 157
Tracheal buttons, 149
Tracheal chondritis, 157
Tracheal intubation, 62–110
airway anatomy for, 62, 63f
changing tubes in, 109–110, 109f
Combitube in place, 101–102, 102f
comlications of, 67f, 84–85
difficult airway, assessing for, 65–67, 67f
digital, 105–106, 106f
direct laryngoscopy in, 67f, 69–85, 73f
contraindications to, 67f, 69
equipment for, 67f
indications for, 67f, 69
optimal patient positioning for, 71–72, 71f
procedure and technique of, 72–85
equipment for, 62, 69–78
flexible endoscopic, 95–101, 96f
complications of, 100
contraindications to, 96, 97f
direct laryngoscopy of, 100, 100f
indications for, 96, 97f
oral approach of, 98–100, 99f–100f
procedure for, 96–100
technique of, 96–100
laryngospasm with, 80
optical and video intubating stylets of, 101
optical laryngoscopes in, 85–99
orotracheal intubation with a king
laryngeal tube, 101–102, 102f
positioning and securing tube, 80, 82f
preoxygenation in, 64–65
preparation for, 62–64, 64f
retrograde, 106–109, 107f
comlications of, 106, 108
contraindications to, 106
equipment for, 106
indications for, 106
procedure for, 106–107
technique of, 106–107
tracheal stricture with, 85
tube introducer for, 78–80, 78f–79f
tube passing, 77–78, 77f
tube placement
clinical assessment of, 80–81
confirmation of, 80–84, 82f
tube tip design in, 71f
tubes for, 70–71, 70f
video laryngoscopes in, 85–95, 86f
with angulated blades, 87–90, 87f
with standard Macintosh, 85–87, 86f
with tube channel, 90–92, 90f
Tracheal stenosis, 153–154
Tracheal tube introducer, 78–80, 78f–79f
Tracheal tube passing, 77–78, 77f
Tracheobronchitis, 153
Tracheoesophageal fistula, 154
Tracheoinnominate artery fistula (TIF), 155
Tracheomalacia, 153–154, 154f
Tracheostomal tube dislodgment, 152
Tracheostomy, 127
bleeding with, 154–157
care, 142–159
changing of, 147–151, 150f–151f
comlications of, 148f
contraindications to, 148f
equipment for, 143, 143f, 148, 148f
false passage with, 153
indications for, 147–148, 148f
procedure for, 149–151, 150f
tube components in, 149
tube sizing for, 148–149, 148r
comlications of, 143, 151–156, 159
in children, 159
cuff, 144, 158–159
comlications with, 153
uffed or unuffed tubes in, 149
dislodgment of, 152
equipment for, 158
failure, 153
evaluation of, 142–143
false passage of, 153
humidifiers for, 159
infection with, 153
mechanical ventilation and, 179
mini-, suctioning procedure, 147
misplacement of, 153
in obese patients, 157–158
obstruction in, 151–152
pediatrics, 158–159
routine maintenance of, 143–144, 144f
stoma, 152
suctioning of, 144–147, 146f, 159
comlications of, 145f, 147
contraindications to, 145, 145f
equipment for, 145–146, 145f
indications for, 145, 145f
procedure for, 146–147, 146f
technique for, 146–147
tracheal buttons with, 149
tracheal stent with, 157
tracheoesophageal fistula with, 154
tracheomalacia with, 153–154
tube, 152
fracture, 153
obstruction, 151–152
sizing, 148–149, 148r, 158, 158r
ventilation with, 144, 144f
Traction splint, 935, 935f–936f
application of, 935f
Traction-countertraction method
in posterior elbow dislocations, 1001, 1001f
in shoulder dislocations, 990–993, 991f–992f
Traditional (open) technique, for surgical cricoidotomy, 128f, 130–133, 133f–135f
Tranexamic acid
for epistaxis, 1372–1373
in anticoagulated patient, 1527
for hemostasis, 1528
Transcutaneous pacemaker, 288
Transducers
for central venous pressure, 432–433, 433f
for ultrasound, 1434, 1436f–1437f
Transesophageal puncture (TEP), for voice restoration, 156
Transfusion reactions, 503–505
Transfusion therapy, 500–522
abbreviated crossmatch in, 507
acute lung injury related to, 504, 511
allergic reactions to, 503
anaphylactic reaction to, 501–504
blood products for, 508–517, 509f
administration of, 519–521
for Jehovah’s Witnesses, 519
caevts in, 522f
Tones
dislocations of
interphalangeal, 1025–1026, 1025f
tomathalgonal and, 1024–1025, 1025f
fracture of, 1060–1061, 1061f
nerv block at, 562r, 581–587, 581f–582f
Toudilue blue dye staining, in sexual assault, 453
Tourniquets
accessing, 453
Traction splint, 935, 935f–936f
application of, 935f
Traction-countertraction method
in posterior elbow dislocations, 1001, 1001f
in shoulder dislocations, 990–993, 991f–992f
Traditional (open) technique, for surgical cricoidotomy, 128f, 130–133, 133f–135f
Tranexamic acid
for epistaxis, 1372–1373
in anticoagulated patient, 1527
for hemostasis, 1528
Transcutaneous pacemaker, 288
Transducers
for central venous pressure, 432–433, 433f
for ultrasound, 1434, 1436f–1437f
Transesophageal puncture (TEP), for voice restoration, 156
Transfusion reactions, 503–505
Transfusion therapy, 500–522
abbreviated crossmatch in, 507
acute lung injury related to, 504, 511
allergic reactions to, 503
anaphylactic reaction to, 501–504
blood products for, 508–517, 509f
administration of, 519–521
for Jehovah’s Witnesses, 519
caevts in, 522f
This page intentionally left blank
This page intentionally left blank
This page intentionally left blank
This page intentionally left blank
Fast answers and trusted evidence

Drive better outcomes with a clinical search engine that makes it easier to find and apply relevant knowledge.

Fast
Anticipates your query, recognizing concepts and offering shortcuts

Convenient
Accessible at the patient’s bedside or on the go, making it easy to discover, share, and apply content anywhere

Complete
Draws relevant answers from a wide range of current, comprehensive content across 30+ medical and surgical specialties

Trusted
Content from Elsevier, the name healthcare professionals worldwide rely on

ckey.co/books

ELSEVIER