BLACKWELL’S
UNDERGROUND CLINICAL VIGNETTES

MICROBIOLOGY
VOL. I, 3E

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Notice
The authors of this volume have taken care that the information contained herein is accurate and compatible with the standards generally accepted at the time of publication. Nevertheless, it is difficult to ensure that all the information given is entirely accurate for all circumstances. The publisher and authors do not guarantee the contents of this book and disclaim any liability, loss, or damage incurred as a consequence, directly or indirectly, of the use and application of any of the contents of this volume.
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<tr>
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<td>Anemia—Aplastic Crisis (Parvovirus B19)</td>
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<td>Atypical Mycobacterial Infection</td>
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<td>Cat-Scratch Disease</td>
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<td>Colorado Tick Fever</td>
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<td>Cryptosporidiosis</td>
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ACKNOWLEDGMENTS

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For submitting comments, corrections, editing, proofreading, and assistance across all of the vignette titles in all editions, we collectively thank:

For generously contributing images to the entire *Underground Clinical Vignette* Step 1 series, we collectively thank the staff at Blackwell Science in Oxford, Boston, and Berlin as well as:

- **Axford, J. Medicine.** Osney Mead: Blackwell Science Ltd., 1996. Figures 2.14, 2.15, 2.16, 2.27, 2.28, 2.31, 2.35, 2.36, 2.38, 2.43, 2.65a, 2.65b, 2.65c, 2.103b, 2.105b, 3.20b, 3.21, 8.27, 8.27b, 8.77b, 8.77c, 10.81b, 10.96a, 12.28a, 14.6, 14.16, 14.50.


Please let us know if your name has been missed or misspelled and we will be happy to make the update in the next edition.
We were very pleased with the overwhelmingly positive student feedback for the 2nd edition of our Undergraduate Clinical Vignettes series. Well over 100,000 copies of the UCV books are in print and have been used by students all over the world.

Over the last two years we have accumulated and incorporated over a thousand “updates” and improvements suggested by you, our readers, including:

- many additions of specific boards and wards testable content
- deletions of redundant and overlapping cases
- reordering and reorganization of all cases in both series
- a new master index by case name in each Atlas
- correction of a few factual errors
- diagnosis and treatment updates
- addition of 5–20 new cases in every book
- and the addition of clinical exam photographs within UCV—Anatomy

And most important of all, the third edition sets now include two brand new COLOR ATLAS supplements, one for each Clinical Vignette series.

- The UCV–Basic Science Color Atlas (Step 1) includes over 250 color plates, divided into gross pathology, microscopic pathology (histology), hematology, and microbiology (smears).
- The UCV–Clinical Science Color Atlas (Step 2) has over 125 color plates, including patient images, dermatology, and funduscopy.

Each atlas image is descriptively captioned and linked to its corresponding Step 1 case, Step 2 case, and/or Step 2 MiniCase.
How Atlas Links Work:

Step 1 Book Codes are:  
A = Anatomy  
B = Behavioral Science  
BC = Biochemistry  
M1 = Microbiology, Vol. I  
M2 = Microbiology, Vol. II  
P1 = Pathophysiology, Vol. I  
P2 = Pathophysiology, Vol. II  
P3 = Pathophysiology, Vol. III  
PH = Pharmacology

Step 2 Book Codes are:  
ER = Emergency Medicine  
IM1 = Internal Medicine, Vol. I  
IM2 = Internal Medicine, Vol. II  
NEU = Neurology  
OB = OB/GYN  
PED = Pediatrics  
SUR = Surgery  
PSY = Psychiatry  
MC = MiniCase

Case Number  
M-P3-032A  
ER-035A, ER-035B

Indicates Type of Image:  
H = Hematology  
M = Microbiology  
PG = Gross Pathology  
PM = Microscopic Pathology

- If the Case number (032, 035, etc.) is not followed by a letter, then there is only one image. Otherwise A, B, C, D indicate up to 4 images.

**Bold Faced Links:** In order to give you access to the largest number of images possible, we have chosen to cross link the Step 1 and 2 series.

- If the link is bold-faced this indicates that the link is direct (i.e., Step 1 Case with the Basic Science Step 1 Atlas link).
- If the link is not bold-faced this indicates that the link is indirect (Step 1 case with Clinical Science Step 2 Atlas link or vice versa).

We have also implemented a few structural changes upon your request:

- Each current and future edition of our popular *First Aid for the USMLE Step 1* (Appleton & Lange/McGraw-Hill) and *First Aid for the USMLE Step 2* (Appleton & Lange/McGraw-Hill) book will be linked to the corresponding UCV case.
- We eliminated UCV → First Aid links as they frequently become out of date, as the *First Aid* books are revised yearly.
• The Color Atlas is also specially designed for quizzing—captions are descriptive and do not give away the case name directly.

We hope the updated UCV series will remain a unique and well-integrated study tool that provides compact clinical correlations to basic science information. They are designed to be easy and fun (comparatively) to read, and helpful for both licensing exams and the wards.

We invite your corrections and suggestions for the fourth edition of these books. For the first submission of each factual correction or new vignette that is selected for inclusion in the fourth edition, you will receive a personal acknowledgement in the revised book. If you submit over 20 high-quality corrections, additions or new vignettes we will also consider inviting you to become a “Contributor” on the book of your choice. If you are interested in becoming a potential “Contributor” or “Author” on a future UCV book, or working with our team in developing additional books, please also e-mail us your CV/resume.

We prefer that you submit corrections or suggestions via electronic mail to UCVteam@yahoo.com. Please include “Underground Vignettes” as the subject of your message. If you do not have access to e-mail, use the following mailing address: Blackwell Publishing, Attn: UCV Editors, 350 Main Street, Malden, MA 02148, USA.

Vikas Bhushan
Vishal falling
Tao Le
October 2001
HOW TO USE THIS BOOK

This series was originally developed to address the increasing number of clinical vignette questions on medical examinations, including the USMLE Step 1 and Step 2. It is also designed to supplement and complement the popular First Aid for the USMLE Step 1 (Appleton & Lange/McGraw Hill) and First Aid for the USMLE Step 2 (Appleton & Lange/McGraw Hill).

Each UCV 1 book uses a series of approximately 100 “super-potent” cases as a way to condense testable facts and associations. The clinical vignettes in this series are designed to incorporate as many testable facts as possible into a cohesive and memorable clinical picture. The vignettes represent composites drawn from general and specialty textbooks, reference books, thousands of USMLE style questions and the personal experience of the authors and reviewers.

Although each case tends to present all the signs, symptoms, and diagnostic findings for a particular illness, patients generally will not present with such a “complete” picture either clinically or on a medical examination. Cases are not meant to simulate a potential real patient or an exam vignette. All the boldfaced “buzzwords” are for learning purposes and are not necessarily expected to be found in any one patient with the disease.

Definitions of selected important terms are placed within the vignettes in (small caps) in parentheses. Other parenthetical remarks often refer to the pathophysiology or mechanism of disease. The format should also help students learn to present cases succinctly during oral “bullet” presentations on clinical rotations. The cases are meant to serve as a condensed review, not as a primary reference. The information provided in this book has been prepared with a great deal of thought and careful research. This book should not, however, be considered as your sole source of information. Corrections, suggestions and submissions of new cases are encouraged and will be acknowledged and incorporated when appropriate in future editions.
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<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>5-ASA</td>
<td>5-aminosalicylic acid</td>
</tr>
<tr>
<td>ABGs</td>
<td>arterial blood gases</td>
</tr>
<tr>
<td>ABVD</td>
<td>Adriamycin/bleomycin/vincristine/dacarbazine</td>
</tr>
<tr>
<td>ACE</td>
<td>angiotensin-converting enzyme</td>
</tr>
<tr>
<td>ACTH</td>
<td>adrenocorticotropic hormone</td>
</tr>
<tr>
<td>ADH</td>
<td>antidiuretic hormone</td>
</tr>
<tr>
<td>AFP</td>
<td>alpha fetal protein</td>
</tr>
<tr>
<td>AI</td>
<td>aortic insufficiency</td>
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<tr>
<td>AIDS</td>
<td>acquired immunodeficiency syndrome</td>
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<tr>
<td>ALL</td>
<td>acute lymphocytic leukemia</td>
</tr>
<tr>
<td>ALT</td>
<td>alanine transaminase</td>
</tr>
<tr>
<td>AML</td>
<td>acute myelogenous leukemia</td>
</tr>
<tr>
<td>ANA</td>
<td>antinuclear antibody</td>
</tr>
<tr>
<td>ARDS</td>
<td>adult respiratory distress syndrome</td>
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<tr>
<td>ASD</td>
<td>atrial septal defect</td>
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<tr>
<td>ASO</td>
<td>anti-streptolysin O</td>
</tr>
<tr>
<td>AST</td>
<td>aspartate transaminase</td>
</tr>
<tr>
<td>AV</td>
<td>arteriovenous</td>
</tr>
<tr>
<td>BE</td>
<td>barium enema</td>
</tr>
<tr>
<td>BP</td>
<td>blood pressure</td>
</tr>
<tr>
<td>BUN</td>
<td>blood urea nitrogen</td>
</tr>
<tr>
<td>CAD</td>
<td>coronary artery disease</td>
</tr>
<tr>
<td>CALLA</td>
<td>common acute lymphoblastic leukemia antigen</td>
</tr>
<tr>
<td>CBC</td>
<td>complete blood count</td>
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<tr>
<td>CHF</td>
<td>congestive heart failure</td>
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<tr>
<td>CK</td>
<td>creatine kinase</td>
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<tr>
<td>CLL</td>
<td>chronic lymphocytic leukemia</td>
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<tr>
<td>CML</td>
<td>chronic myelogenous leukemia</td>
</tr>
<tr>
<td>CMV</td>
<td>cytomegalovirus</td>
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<tr>
<td>CNS</td>
<td>central nervous system</td>
</tr>
<tr>
<td>COPD</td>
<td>chronic obstructive pulmonary disease</td>
</tr>
<tr>
<td>CPK</td>
<td>creatine phosphokinase</td>
</tr>
<tr>
<td>CSF</td>
<td>cerebrospinal fluid</td>
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<tr>
<td>CT</td>
<td>computed tomography</td>
</tr>
<tr>
<td>CVA</td>
<td>cerebrovascular accident</td>
</tr>
<tr>
<td>CXR</td>
<td>chest x-ray</td>
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<tr>
<td>DIC</td>
<td>disseminated intravascular coagulation</td>
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<tr>
<td>DIP</td>
<td>distal interphalangeal</td>
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<tr>
<td>DKA</td>
<td>diabetic ketoacidosis</td>
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<td>DM</td>
<td>diabetes mellitus</td>
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<td>DTRs</td>
<td>deep tendon reflexes</td>
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<td>DVT</td>
<td>deep venous thrombosis</td>
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<td>Abbreviation</td>
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<td>EBV</td>
<td>Epstein-Barr virus</td>
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<td>electrocardiography</td>
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<td>Echo</td>
<td>echocardiography</td>
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<td>ejection fraction</td>
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<td>esophagogastroduodenoscopy</td>
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<td>EMG</td>
<td>electromyography</td>
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<td>ERCP</td>
<td>endoscopic retrograde cholangiopancreatography</td>
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<td>ESR</td>
<td>erythrocyte sedimentation rate</td>
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<td>FEV</td>
<td>forced expiratory volume</td>
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<tr>
<td>FNA</td>
<td>fine needle aspiration</td>
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<tr>
<td>FTA-ABS</td>
<td>fluorescent treponemal antibody absorption</td>
</tr>
<tr>
<td>FVC</td>
<td>forced vital capacity</td>
</tr>
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<td>GFR</td>
<td>glomerular filtration rate</td>
</tr>
<tr>
<td>GH</td>
<td>growth hormone</td>
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<tr>
<td>GI</td>
<td>gastrointestinal</td>
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<td>GM-CSF</td>
<td>granulocyte macrophage colony stimulating factor</td>
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<td>genitourinary</td>
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<td>HAV</td>
<td>hepatitis A virus</td>
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<tr>
<td>hCG</td>
<td>human chorionic gonadotrophin</td>
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<td>HEENT</td>
<td>head, eyes, ears, nose, and throat</td>
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<td>human immunodeficiency virus</td>
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<td>HLA</td>
<td>human leukocyte antigen</td>
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<td>HPI</td>
<td>history of present illness</td>
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<td>heart rate</td>
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<td>HRIG</td>
<td>human rabies immune globulin</td>
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<td>ID/CC</td>
<td>identification and chief complaint</td>
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<td>IDDM</td>
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<tr>
<td>Ig</td>
<td>immunoglobulin</td>
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<td>IGF</td>
<td>insulin-like growth factor</td>
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<td>intramuscular</td>
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<td>JVP</td>
<td>jugular venous pressure</td>
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<td>KUB</td>
<td>kidneys/ureter/bladder</td>
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<td>LDH</td>
<td>lactate dehydrogenase</td>
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<tr>
<td>LES</td>
<td>lower esophageal sphincter</td>
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<tr>
<td>LFTs</td>
<td>liver function tests</td>
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<tr>
<td>LP</td>
<td>lumbar puncture</td>
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<tr>
<td>LV</td>
<td>left ventricular</td>
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<td>LVH</td>
<td>left ventricular hypertrophy</td>
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<tr>
<td>Lytes</td>
<td>electrolytes</td>
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<tr>
<td>MCHC</td>
<td>mean corpuscular hemoglobin concentration</td>
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<tr>
<td>MCV</td>
<td>mean corpuscular volume</td>
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<tr>
<td>MEN</td>
<td>multiple endocrine neoplasia</td>
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<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>MGUS</td>
<td>monoclonal gammopathy of undetermined significance</td>
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<tr>
<td>MHC</td>
<td>major histocompatibility complex</td>
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<tr>
<td>MI</td>
<td>myocardial infarction</td>
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<tr>
<td>MOPP</td>
<td>mechlorethamine/vincristine (Oncovorin)/procarbazine/prednisone</td>
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<tr>
<td>MR</td>
<td>magnetic resonance (imaging)</td>
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<tr>
<td>NHL</td>
<td>non-Hodgkin's lymphoma</td>
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<td>NIDDM</td>
<td>non-insulin-dependent diabetes mellitus</td>
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<tr>
<td>NPO</td>
<td>nil per os (nothing by mouth)</td>
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<tr>
<td>NSAID</td>
<td>nonsteroidal anti-inflammatory drug</td>
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<td>PA</td>
<td>posteroanterior</td>
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<td>PIP</td>
<td>proximal interphalangeal</td>
</tr>
<tr>
<td>PBS</td>
<td>peripheral blood smear</td>
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<tr>
<td>PE</td>
<td>physical exam</td>
</tr>
<tr>
<td>PFTs</td>
<td>pulmonary function tests</td>
</tr>
<tr>
<td>PMI</td>
<td>point of maximal intensity</td>
</tr>
<tr>
<td>PMN</td>
<td>polymorphonuclear leukocyte</td>
</tr>
<tr>
<td>PT</td>
<td>prothrombin time</td>
</tr>
<tr>
<td>PTCA</td>
<td>percutaneous transluminal angioplasty</td>
</tr>
<tr>
<td>PTH</td>
<td>parathyroid hormone</td>
</tr>
<tr>
<td>PTT</td>
<td>partial thromboplastin time</td>
</tr>
<tr>
<td>PUD</td>
<td>peptic ulcer disease</td>
</tr>
<tr>
<td>RBC</td>
<td>red blood cell</td>
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<td>RPR</td>
<td>rapid plasma reagin</td>
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<tr>
<td>RR</td>
<td>respiratory rate</td>
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<tr>
<td>RS</td>
<td>Reed-Sternberg (cell)</td>
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<tr>
<td>RV</td>
<td>right ventricular</td>
</tr>
<tr>
<td>RVH</td>
<td>right ventricular hypertrophy</td>
</tr>
<tr>
<td>SBFT</td>
<td>small bowel follow-through</td>
</tr>
<tr>
<td>SIADH</td>
<td>syndrome of inappropriate secretion of ADH</td>
</tr>
<tr>
<td>SLE</td>
<td>systemic lupus erythematosus</td>
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<tr>
<td>STD</td>
<td>sexually transmitted disease</td>
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<tr>
<td>TFTs</td>
<td>thyroid function tests</td>
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<tr>
<td>tPA</td>
<td>tissue plasminogen activator</td>
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<td>TSH</td>
<td>thyroid-stimulating hormone</td>
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<tr>
<td>TIBC</td>
<td>total iron-binding capacity</td>
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<tr>
<td>TIPS</td>
<td>transjugular intrahepatic portosystemic shunt</td>
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<tr>
<td>TPO</td>
<td>thyroid peroxidase</td>
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<tr>
<td>TSH</td>
<td>thyroid-stimulating hormone</td>
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<td>TTP</td>
<td>thrombotic thrombocytopenic purpura</td>
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<tr>
<td>UA</td>
<td>urinalysis</td>
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<tr>
<td>UGI</td>
<td>upper GI</td>
</tr>
<tr>
<td>US</td>
<td>ultrasound</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<tr>
<td>VDRL</td>
<td>Venereal Disease Research Laboratory</td>
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<td>WPW</td>
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A 25-year-old IV drug abuser presents with a high fever with chills, malaise, a productive cough, hemoptysis, and right-sided pleuritic chest pain.

HPI He also reports multiple skin infections at injection sites.

PE VS: fever. PE: stigmata of intravenous drug abuse at multiple injection sites; skin infections; thrombosed peripheral veins; splenomegaly and pulsatile hepatomegaly; ejection systolic murmur, increasing with inspiration, heard in tricuspid area.

Labs CBC: normochromic, normocytic anemia. UA: microscopic hematuria. Blood culture yields *Staphylococcus aureus*.

Imaging Echo: presence of vegetations on tricuspid valve and tricuspid incompetence. CXR: consolidation.

Treatment High-dose intravenous penicillinase-resistant penicillin in combination with an aminoglycoside. If the isolated *S. aureus* strain is methicillin resistant, vancomycin is the drug of choice.

Discussion In drug addicts, the tricuspid valve is the site of infection more frequently (55%) than the aortic valve (35%) or the mitral valve (30%); these findings contrast markedly with the rarity of right-sided involvement in cases of infective endocarditis that are not associated with drug abuse. *Staphylococcus aureus* is responsible for the majority of cases. Certain organisms have a predilection for particular valves in cases of addict-associated endocarditis; for example, enterococci, other streptococcal species, and non-albicans *Candida* organisms predominantly affect the valves of the left side of the heart, while *S. aureus* infects valves on both the right and the left side of the heart. *Pseudomonas* organisms are associated with biventricular and multiple-valve infection in addicts. Complications of endocarditis include congestive heart failure, ruptured valve cusp, myocardial infarction, and myocardial abscess.
A 25-year-old male complains of increasing shortness of breath and ankle edema that have progressively worsened over the past 2 weeks.

He also complains of fatigue, palpitations, and low-grade fever. His symptoms followed a severe URI. He denies any history of joint pain or skin rash (vs. rheumatic fever).

JVP elevated; pitting pedal edema; fine inspiratory crepitations heard at both lung bases; mild hepatosplenomegaly.

ASO titers not elevated. CBC: lymphocytosis. ECG: first-degree AV block. ESR elevated; increased titers of antibodies to coxsackievirus demonstrated in serum.


Dilated heart with foci of epicardial, myocardial, and endocardial petechial hemorrhages.

Endomyocardial biopsy reveals diffuse infiltration by mononuclear cells, predominantly lymphocytes; focal fibrosis.

Manage congestive heart failure and arrhythmias; cardiac transplant in intractable cases.

Coxsackie B is most often implicated in viral myocarditis. Nonviral causes of myocarditis include bacteria such as Borrelia burgdorferi (Lyme disease), parasites such as Trypanosoma cruzi (Chagas' disease), hypersensitivity reaction (systemic lupus erythematosus, drug reaction), radiation, and sarcoidosis; may also be idiopathic (giant cell myocarditis).
ID/CC A 35-year-old male complains of fever, nonproductive cough, and chest pain.

HPI He states that the chest pain developed after he had a severe cold for 1 week. He describes the pain as severe, crushing, and constant over the anterior chest and adds that it worsens with inspiration and is relieved by sitting up and bending forward.

PE VS: low-grade fever; sinus tachycardia. PE: triphasic pericardial friction rub (systolic and diastolic components followed by a third component in late diastole associated with atrial contraction); elevated JVP; inappropriate increase in JVP with inspiration (Kussmaul's sign); pulsus paradoxus may also be seen.

Labs Moderately elevated transaminases and LDH; elevated ESR; serum CPK-MB normal. CBC: neutrophilic leukocytosis. ECG: diffuse ST-segment elevation (vs. myocardial infarction); PR-segment depression.

Imaging Echo: pericardial effusion. CXR: apparent cardiomegaly (due to effusion).

Gross Pathology In long-standing cases, pericardium may become fibrotic, scarred, and calcified.

Micro Pathology Pericardial biopsy reveals signs of acute inflammation with increased leukocytes, vascularity, and deposition of fibrin.

Treatment Analgesics for pain; steroids in resistant cases; indomethacin; surgical stripping of scarring in severe cases.

Discussion Acute pericarditis is commonly idiopathic. Known infectious causes include coxsackievirus A and B, tuberculosis, staphylococcal or pneumococcal infection, amebiasis, or actinomycosis; noninfectious causes include chronic renal failure, collagen-vascular disease (systemic lupus erythematosus, scleroderma, and rheumatoid arthritis), neoplasms, myocardial infarction, and trauma. Long-term sequelae include chronic constrictive pericarditis.

Atlas Link [UCC] PG-M1-003

3 PERICARDITIS—ACUTE
ID/CC A 64-year-old male presents with rapidly progressive dyspnea and fever.

HPI He has a history of orthopnea and paroxysmal nocturnal dyspnea and also reports pink, frothy sputum (HEMOPHTYSIS). One month ago he underwent a bioprosthetic valve replacement for calcific aortic stenosis. He is not hypertensive and has never had overt cardiac failure in the past.

PE VS: fever; hypotension. PE: bilateral basal inspiratory crackles heard; cardiac auscultation suggestive of aortic incompetence (early diastolic murmur heard radiating down left sternal edge).

Labs CBC: normochromic, normocytic anemia. Three consecutive blood cultures yield coagulase-negative Staphylococcus epidermidis; strain found to be methicillin resistant.

Imaging CXR (PA view): suggestive of pulmonary edema. Echo: confirms presence of prosthetic aortic valve dehiscence leading to incompetence and poor left ventricular function.

Treatment High-dose parenteral antibiotics—vancomycin (drug of choice for methicillin-resistant S. aureus), gentamicin, and oral rifampicin; surgical replacement of damaged prosthetic valve; prophylactic antibiotics (amoxicillin) for patients receiving oral/dental treatments to prevent transient bacteremia.

Discussion Prosthetic valve endocarditis is subdivided into two categories: early prosthetic valve endocarditis (EPVE), which becomes clinically manifest within 60 days after valve replacement (most commonly caused by Staphylococcus epidermidis, followed by gram-negative bacilli and Candida), and late prosthetic valve endocarditis (LPVE), which is manifested clinically more than 60 days after valve replacement (most commonly caused by viridans streptococci).
ID/CC A 25-year-old female complains of low-grade fever and myalgia of 3 weeks' duration.

HPI She has a history of rheumatic heart disease (RHD). One month ago, she underwent a dental extraction and did not take the antibiotics that were prescribed for her.

PE VS: fever. PE: pallor; small peripheral hemorrhages with slight nodular character (Janeway lesions); small, tender nodules on finger and toe pads (Osler's nodes); subungual linear streaks (splinter hemorrhages); petechial hemorrhages on conjunctiva, oral mucosa, and upper extremities; mild splenomegaly; apical diastolic murmur on cardiovascular exam; fundus exam shows oval retinal hemorrhages (Roth's spots).

Labs CBC/PBS: normocytic, normochromic anemia. UA: microscopic hematuria. Growth of penicillin-sensitive Streptococcus viridans on five of six blood cultures.

Imaging Echo: vegetations along atrial surface of mitral valve.

Gross Pathology Embolism from vegetative growths on valves may embolize peripherally (left-sided) or to the lung (right-sided).

Micro Pathology Bacteria form nidus of infection in previously scarred or damaged valves; bacteria divide unimpeded once infection takes hold with further deposition of fibrin and platelets; peripheral symptoms such as Osler's nodes are believed to result from deposition of immune complexes.

Treatment IV β-lactamase-resistant penicillin and gentamicin; bacteriostatic treatments ineffective.

Discussion S. viridans is the most common cause of subacute infective endocarditis, while Staphylococcus aureus is the most common cause of acute bacterial endocarditis. Prophylactic antibiotics should be given to all RHD patients before any dental procedure. The disease continues to be associated with a high mortality rate.
ID/CC A 54-year-old female who underwent a left mastectomy with axillary lymph node dissection a year ago presents with pain together with rapidly spreading redness and swelling of the left arm.

HPI One year ago, she was diagnosed and operated on for stage 1 carcinoma of the left breast.

PE Left forearm swollen, indurated, pink, and markedly tender; overlying temperature raised; margins and borders of skin lesion ill defined and not elevated (vs. erysipelas).

Labs Needle aspiration from advancing border of the lesion, when stained and cultured, isolated β-hemolytic group A streptococcus.

Treatment Penicillinase-resistant penicillin (nafcillin/oxacillin).

Discussion Cellulitis is an acute spreading infection of the skin that predominantly affects deeper subcutaneous tissue. Group A streptococci and Staphylococcus aureus are the most common etiologic agents in adults; Haemophilus influenzae infection is common in children. Patients with chronic venous stasis and lymphedema of any cause (lymphoma, filariasis, post-regional lymph node dissection, radiation therapy) are predisposed; recently, recurrent saphenous-vein donor-site cellulitis was found to be attributable to group A, C, or G streptococci.
ID/CC  A 16-year-old teenager presents to the outpatient clinic with a painful facial rash and fever.

HPI  One week ago, the patient went on a camping trip and scratched his face on some low-lying tree branches. There is no medical history of diabetes, cancer, or other chronic conditions.

PE  VS: fever (39.0°C); tachycardia (HR 110); BP normal. PE: erythematous, warm, plaque-like rash extending across cheeks and face bilaterally with sharp, distinct borders and facial swelling.

Labs  CBC: leukocytosis with neutrophilia. ESR elevated.

Treatment  Antibiotics with sufficient coverage for penicillinase-producing Streptococcus and Staphylococcus spp. (e.g., cephalaxin); analgesics/antipyretics; elevate the affected part to reduce swelling.

Discussion  Erysipelas is an acute inflammation of the superficial layers of the connective tissues of the skin, usually on the face, almost always caused by infection with Group A Streptococcus which is part of normal bacterial skin flora. Risk factors include any breaks in the skin or lymphedema.

Atlas Link  [Image] MC-136
ID/CC  A 30-year-old slaughterhouse worker presents with a painful red swelling of the index finger of his right hand.

HPI  The swelling developed 4 days after he was injured with a knife while slaughtering a pig.

PE  Well-defined, exquisitely tender, slightly elevated violaceous lesion seen on right index finger; no suppuration noted; right epitrochlear and right axillary lymphadenopathy noted.

Labs  Biopsy from edge of lesion yields *Erysipelothrix rhusiopathiae*, a thin, pleomorphic, nonsporulating, microaerophilic gram-positive rod.

Treatment  Penicillin G or ciprofloxacin in penicillin-allergic patients.

Discussion  Erysipeloid refers to localized cellulitis, usually of the fingers and hands, caused by *Erysipelothrix rhusiopathiae*; infection in humans is usually the result of contact with infected animals or their products (often fish). Organisms gain entry via cuts and abrasions on the skin.
ID/CC A 10-year-old male complains of a spreading skin rash and painful swelling of both wrists.

HPI The patient’s mother states that the rash began with erythema of the cheeks ("slapped-cheek appearance") and subsequently progressed to involve the trunk and limbs.

PE Erythematous lacy/reticular skin rash involving face, trunk, and limbs; bilateral swelling and painful restriction of movement at both wrist joints.

Labs Serology detects presence of specific IgM antibody to parvovirus; ASO titer (to rule out acute rheumatic fever) normal; rheumatoid factor (to rule out rheumatoid arthritis) negative.

Treatment Self-limiting disease.

Discussion A small (20- to 26-nm), single-stranded DNA virus, parvovirus B19 causes erythema infectiosum (fifth disease) in schoolchildren, aplastic crises in persons with underlying hemolytic disorders (e.g., sickle cell anemia), chronic anemia in immunocompromised hosts, arthralgia/arthritis in normal individuals, and fetal loss in pregnant women.

Atlas Link UCV2 MC-299
ID/CC  A 5-year-old white male presents with golden-yellow, crusted lesions around his mouth and behind his ears.

HPI  He has a history of intermittent low-grade fever, frequent "nose picking," and purulent discharge from his lesions. He has no history of hematuria (due to increased risk of poststreptococcal glomerulonephritis).

PE  Characteristic "honey-colored" crusted lesions seen at angle of mouth, around nasal orifices, and behind ears.

Labs  Gram-positive cocci in chains (Streptococci) in addition to pus cells on Gram stain of discharge; β-hemolytic streptococci (group A streptococci) on blood agar culture; ASO titer negative.

Gross Pathology  Erythematous lesions surrounding natural orifices with whitish or yellowish purulent exudate and crust formation.

Micro Pathology  Inflammatory infiltrate of PMNs with varying degrees of necrosis.

Treatment  Cephalosporin, penicillin, or erythromycin if allergic.

Discussion  Impetigo is a highly communicable infectious disease that is most often caused by group A streptococci, occurs primarily in preschoolers, and may predispose to glomerulonephritis. It occurs most commonly on the face (periocular area), hands, and arms. Staphylococcus aureus may coexist or cause bullous impetigo; group B streptococcal impetigo may be seen in newborns.
ID/CC  A 30-year-old male homosexual visits his family doctor complaining of a nonpruritic skin eruption on his upper limbs, trunk, and anogenital area.

HPI  He has been HIV positive for about 3 years and admits to having continued unprotected intercourse.

PE  Multiple painless, pearly-white, dome-shaped, waxy, umbilicated nodules 2 to 5 mm in diameter on arms, trunk, and anogenital area; palms and soles spared.

Gross Pathology  Firm, umbilicated nodules containing thick yellowish material.

Micro Pathology  Stained histologic sections confirm diagnosis with large cytoplasmic inclusions (molluscum bodies) in material expressed from lesions.

Treatment  Lesions may resolve spontaneously or be removed by curettage, cryotherapy, or podophyllin; no antiviral drug or vaccine available.

Discussion  Molluscum contagiosum is a benign, autoinoculable skin disease of children and young adults; it is caused by a poxvirus (DNA virus) and is transmitted through sexual contact, close bodily contact, clothing, or towels. It is one of many opportunistic infections seen in AIDS patients (difficult to eradicate).

Atlas Link  UCM2 MC-143

MOLLUSCUM CONTAGIOSUM
ID/CC A 30-year-old black male presents with a nonpruritic skin rash on the trunk, upper arm, and neck.

HPI The patient is otherwise in excellent health.

PE Multiple hypopigmented, scaling, confluent macules seen on trunk, upper arms, and neck; no sensory loss demonstrated over areas of hypopigmentation; Wood's lamp examination of skin macules displays a pale yellow to blue-white fluorescence.

Labs Examination of KOH mounting of scales from lesions demonstrates the presence of short, thick, tangled hyphae with clusters of large, spherical budding yeast cells with characteristic "spaghetti-and-meatballs" appearance.

Treatment Topical selenium sulfide; antifungal agents such as miconazole and clotrimazole; oral itraconazole in recalcitrant cases.

Discussion Pityriasis versicolor, which is common in young adults, is a relatively asymptomatic superficial skin infection caused by the lipophilic fungal organism *Pityrosporum orbiculare* (also termed *Malassezia furfur*). The lesions, which usually have a follicular origin, are small, hypopigmented-to-tan macules with a branlike scale; the macules are distributed predominantly on areas of the upper trunk, neck, and shoulders.
ID/CC An 18-month-old male is brought to the pediatrician following the appearance of an extensive skin rash.

HPI Four days ago he suddenly developed a very high fever (40°C) with no other symptoms or signs. The fever continued for 4 days until the day of his admission, when it abruptly disappeared, coinciding with the onset of the rash.

PE Child looks well; in no acute distress; generalized rash apparent as discrete 2- to 5-mm rose-pink macules and papules on trunk, neck, and extremities (face is spared); lesions blanch on pressure; no lymphadenopathy; splenomegaly may also be present.

Labs CBC/PBS: WBCs variable; relative lymphocytosis with atypical lymphocytes.

Treatment Supportive; foscarnet.

Discussion Roseola infantum, also called exanthem subitum, is caused by human herpesvirus 6. It is the most common exanthematosus disease in infants 2 years of age or younger and is a frequent cause of febrile convulsions.
ID/CC: A 2-month-old female infant presents with extensive bullae and large areas of denuded skin.

HPI: Her mother had suffered from staphylococcal mastitis 1 week ago.

PE: VS: fever. PE: large areas of red, painful, denuded skin on periorbital and peribuccal areas; flaccid bullae with easy dislodgment of epidermis under pressure (Nikolsky's sign); mucosal surfaces largely uninvolved.

Labs: Vesicle fluid sterile; Staphylococcus aureus on blood culture.

Treatment: IV penicillinase-resistant penicillin (e.g., nafcillin, oxacillin). Treat with erythromycin if patient is allergic to penicillin.

Discussion: Scalded skin syndrome is caused by the exfoliating effect of staphylococcal exotoxin. The action of the exotoxin is to degrade desmoglein in desmosomes in the skin.
A 30-year-old man presents with a bilateral red pruritic skin eruption in the groin area.

Bilateral, circular papulosquamous skin eruption on erythematous base with active, advancing peripheral (serpiginous) border over scrotum and perineum.

Microscopic examination reveals long septate hyphae on KOH skin scrapings.

Topical antifungal agents (Whitfield’s ointment, clotrimazole, miconazole); systemic therapy with oral griseofulvin, ketoconazole, or itraconazole in resistant cases.

Tinea cruris and tinea corporis (common ringworm) occur sporadically; *Trichophyton rubrum* is the most common cause. The inflammatory form, which is usually localized to the limbs, chest, or back, is commonly caused by *Microsporum canis* or *Trichophyton mentagrophytes*. Ringworm of the scalp, known as tinea capitis, is commonly seen in children and is caused by *Trichophyton tonsurans*.

**Tinea Corporis (Ringworm)**
ID/CC A 28-year-old male presents with a red, pruritic skin eruption on his trunk and his upper and lower limbs of a few hours’ duration.

HPI One day earlier, he was prescribed cotrimoxazole for a UTI. He has not experienced any dyspnea.

PE Erythematous, warm, urticarial wheals (hives) seen over trunk, legs, and arms; no angioedema or respiratory distress.

Labs CBC: leukocytosis with eosinophilia. No parasites revealed on stool exam.

Gross Pathology Linear or oval, raised papules or plaque-like wheals up to several centimeters in diameter.

Micro Pathology Wide separation of dermal collagen fibers with dilatation of lymphatics and venules.

Treatment Topical agents to reduce itching; avoidance of causative agent (in this case, cotrimoxazole); antihistamines (primarily H₁ blockers but also H₂ blockers); consider glucocorticoids.

Discussion Mast cells and basophils are focal to urticarial reaction. When stimulated by certain immunologic or nonimmunologic mechanisms, storage granules in these cells release histamine and other mediators, such as kinins and leukotrienes. These agents produce the localized vasodilatation and tranudation of fluid that characterize urticaria.

Atlas Link UCMC MC-021
A 7-year-old male is brought to his family physician complaining of a thick yellowish discharge in his eyes that prevents him from opening his eyes in the morning; for the past few days, his eyes have been blood-red, painful, and watery. His eye pain is exacerbated by exposure to light (phosphobia).

HPI Three of his classmates and a neighbor had a similar episode about 7 days ago (suggesting a local epidemic of such cases).

PE VS: no fever. PE: normal visual acuity; erythematous palpebral conjunctiva; watery eyes; remains of thick mucus found on inner canthal area; no corneal infiltrate on slit-lamp exam; normal anterior chamber; mild preauricular lymphadenopathy.

Labs Stained conjunctival smears reveal lymphocytes, giant cells, neutrophils, and bacteria.

Treatment Topical antimicrobial eye drops; cool compresses; minimize contact with others to avoid spread; avoid use of topical steroid preparations, as these can exacerbate bacterial and viral eye infections.

Discussion Conjunctivitis is a common disease of childhood that is mostly viral (adenovirus) and self-limiting; it occurs in epidemics, and secondary bacterial infections (staphylococci and streptococci) may result. Visual acuity is not affected.
ID/CC  A 35-year-old woman complains of fever and pain in the face and upper teeth (maxillary sinus), especially while leaning forward.

HPI  She has had a chronic cough, nasal congestion, and discharge for the past few months.

PE  VS: fever. PE: halitosis; greenish-yellow postnasal discharge; bilateral boggy nasal mucosa; bilateral percussion tenderness and erythema over zygomatic arch; clouding of sinuses by transillumination; dental and cranial nerve exams normal.

Labs  Nasal cultures reveal *Streptococcus pneumoniae*.

Imaging  CT, sinus: partial opacification of maxillary sinus with air-fluid level.

Gross Pathology  Erythematous and edematous nasal mucosa.

Micro Pathology  Presence of organisms and leukocytes in mucosa.

Treatment  Oral decongestants; amoxicillin, Bactrim, or fluoroquinolone.

Discussion  Other pathogens include other streptococci, *Haemophilus influenzae*, and *Moraxella*. The obstruction of ostia in the anterior ethmoid and middle meatal complex by retained secretions, mucosal edema, or polyps promotes sinusitis. *Staphylococcus aureus* and gram-negative species may cause chronic sinusitis. Fungal sinusitis may mimic chronic bacterial sinusitis. Complications include orbital cellulitis and abscesses.
ID/CC A 17-year-old boy presents with itchy eyes, nasal stuffiness, increased lacrimation, sneezing, and a watery nasal discharge.

HPI He has had similar episodes in the past that have corresponded with changing of the seasons. His mother is known to have bronchial asthma.

PE VS: no fever. PE: pallor; boggy nasal mucosa; nasal polyps present; conjunctiva congested; no exudate.

Labs Conjunctival and nasal smear demonstrates presence of eosinophils; no bacteria on Gram stain; no neutrophils. Allergen skin tests (sensitized cutaneous mast cells) show positive sensitivity.

Gross Pathology Nasal mucosa hyperemic and swollen with fluid transudation.

Micro Pathology Local tissue inflammation and dysfunction of upper airway because of type I, IgE-mediated hypersensitivity response.

Treatment Oral decongestants with intranasal corticosteroids; antihistamines; intranasal cromolyn sodium, especially before anticipated contact with allergen.

Discussion Allergic rhinitis is commonly caused by exposure to pollens, dust content, and insect matter; symptoms are mediated by the release of vasoactive and chemotactic mediators from mast cells and basophils (e.g., histamine and leukotrienes) with IgE surface receptors.
ID/CC  A 20-year-old male presents with a runny nose, nasal congestion, sore throat, headache, and sneezing.

HPI  He notes that his wife currently has similar symptoms.

PE  VS: mild fever. PE: rhinorrhea; congested and inflamed posterior pharyngeal wall; no lymphadenopathy.

Labs  Routine tests normal; routine throat swab staining and culture negative for bacteria.

Gross Pathology  Nasal membranes edematous and erythematous with watery discharge.

Micro Pathology  Mononuclear inflammation of mucosa; focal desquamation.

Treatment  Symptomatic.

Discussion  Colds occur 2 to 3 times a year in the average person in the United States; the peak incidence is in the winter months. Rhinoviruses account for the majority of viral URIs, followed by coronaviruses. Spread occurs by direct contact and respiratory droplets.
ID/CC  A 60-year-old male presents with swelling and a vesicular skin eruption on the left side of his face.

HPI  The patient reports that before the rash developed, he had severe radiating pain on the left side of his face. He also recalls having suffered an attack of chickenpox during his childhood.

PE  Unilateral vesicular rash over left forehead and nasal bridge, including the tip of the nose, indicating involvement of the nasociliary branch of the trigeminal nerve (Hutchinson’s sign); skin of lids red and edematous; slit-lamp examination reveals numerous rounded spots composed of minute white dots involving epithelium and stroma, producing a coarse subepithelial punctate keratitis; cornea is insensitive.

Micro Pathology  Vesicular skin lesions with herpesvirus inclusions that are intranuclear and acidophilic with a clear halo around them (Cowdry type A inclusion bodies); syncytial giant cells also seen.

Treatment  Acyclovir; steroids; cycloplegics. Triflurouromidine for HSV keratitis.

Discussion  Herpes zoster ophthalmicus is caused by the varicella zoster virus, which causes chickenpox as a primary infection. Zoster is believed to be a reactivation of the latent viral infection. In zoster ophthalmicus, the chief focus of reactivation is the trigeminal ganglion, from which the virus travels down one or more branches of the ophthalmic division such that its area of distribution is marked out by rows of vesicles or scars left by the vesicles. Ocular complications arise during subsidence of the rash and are generally associated with involvement of the nasociliary branch of the trigeminal nerve.
**ID/CC**  An 18-year-old male complains of severe irritation in the left eye, blurred vision, excessive lacrimation, and photophobia.

**HPI**  He reports that he has had similar episodes in the past that were treated with an antiviral drug. His records indicate that he suffered the first attack at the age of 7, at which time his condition was diagnosed and treated as a severe follicular keratoconjunctivitis; his records also indicate a history of recurrent episodes of herpes labialis.

**PE**  Examination of left eye reveals circumcorneal congestion; fluorescein staining of cornea reveals infiltrates spreading in all directions, coalescing with each other and forming a large, shallow ulcer with crenated edges ("DENDRITIC ULCER"); cornea is insensitive.

**Labs**  HSV-1 demonstrated on immunofluorescent staining of epithelial scrapings as well as in the aqueous humor.

**Treatment**  Trifluridine eye drops; acyclovir has been shown to decrease recurrences.

**Discussion**  Most ocular herpetic infections are caused by HSV-1. It is also the primary cause of corneal blindness in the United States. Primary infections present as unilateral follicular conjunctivitis, blepharitis, or corneal epithelial opacities; recurrences may take the form of keratitis (> 90% of cases are unilateral), blepharitis, or keratoconjunctivitis. Branching dendritic ulcers, usually detected by fluorescein staining, are virtually diagnostic; deep stromal involvement may result in scarring, corneal thinning, and abnormal vascularization with resulting blindness or rupture of the globe.
A 20-year-old male swimmer complains of severe pain and itching in the right ear that is associated with a slight amount of yellowish (purulent) discharge.

The patient has no previous history of discharge from the ear and no history of associated deafness or tinnitus.

Red, swollen area seen in right external auditory meatus that is partially obliterating the lumen; movement of tragus is exquisitely painful (tragal sign).

Gram stain of aural swab reveals presence of gram-negative rods; culture isolates Pseudomonas aeruginosa.

Red, swollen area seen in cartilaginous part of external auditory meatus; when visualized, tympanic membrane is erythematous and moves normally with pneumatic otoscopy (vs. acute otitis media).

Eardrops (either a combination of polymyxin, neomycin, and hydrocortisone or ofloxacin); gentle removal of debris in ear.

Otitis externa is most common in summer months and is thought to arise from a change in the milieu of the external auditory meatus by increased alkalization and excessive moisture; this leads to bacterial overgrowth, most commonly with gram-negative rods such as Pseudomonas (also causes malignant otitis externa) and Proteus or fungi such as Aspergillus.
ID/CC  An 18-month-old white female presents with irritability together with a bilateral, profuse, and foul-smelling ear discharge of 2 months' duration.

HPI  The patient had recurrent URIs last year, but her mother did not administer the complete course of antibiotics. The patient's mother has a history of feeding her child while lying down.

PE  Bilateral greenish-white ear discharge; perforated tympanic membranes in anteroinferior quadrant of both ears; diminished mobility of tympanic membrane on pneumatic otoscopy.

Labs  Gram-negative coccobacilli on Gram stain of discharge from tympanocentesis; Haemophilus influenzae seen on culture.

Gross Pathology  Possible complications include ingrowth of squamous epithelium on upper middle ear (cholesteatoma) if long-standing; conductive hearing loss; mastoiditis; and brain abscess.

Micro Pathology  Hyperemia and edema of inner ear and throat mucosa; hyperemia of tympanic membrane; deposition of cholesterol crystals in keratinized epidermoid cells in cholesteatoma.

Treatment  Keep ear dry; amoxicillin-clavulanic acid; surgical drainage for severe otalgia; myringoplasty.

Discussion  Otitis media is the most common pediatric bacterial infection and is caused by Escherichia coli, Staphylococcus aureus, and Klebsiella pneumoniae in neonates; in older children it is usually caused by pneumococcus (Streptococcus pneumoniae), H. influenzae, Moraxella catarrhalis, and group A streptococcus. Resistant strains are becoming increasingly common.
ID/CC  A 6-year-old male presents with complaints of a mild sore throat and eye irritation.

HPI  His mother says that he has spent hours at the community swimming pool this summer.

PE  Mild rhinopharyngitis; bilateral conjunctival congestion with scanty mucoid discharge.

Labs  Viral culture of conjunctival and nasopharyngeal swab yields adenovirus.

Treatment  No specific treatment; self-limiting illness.

Discussion  Adenovirus infections occur most often in infants and young children, who acquire the virus by the respiratory or fecal-oral route. The most common respiratory tract syndrome in this age group is mild coryza with pharyngitis; in older children, these symptoms may be accompanied by conjunctivitis. May also cause hemorrhagic cystitis in children. On electron microscopy it is seen as a double-stranded nonenveloped DNA virus surrounded by a 20-faced icosahedral protein capsid from which 12 antenna-like fibers or pentons extend radially.
ID/CC A 9-year-old male complains of pain during swallowing (odynophagia) for 2 days, accompanied by muscle aches, headache, and fever.

HPI He has otherwise been in good health.

PE VS: fever. PE: moderate erythema of pharynx; enlarged, erythematous tonsils covered with white exudate; tender cervical adenopathy.

Labs CBC: neutrophilic leukocytosis. *Streptococcus pyogenes* isolated on throat swab and culture.

Gross Pathology Hyperemia and swelling of upper respiratory tract mucosa; cryptic enlargement of tonsils with purulent exudate; enlargement of regional lymph nodes.

Micro Pathology Acute inflammatory response with polymorphonuclear infiltrate, hyperemia and edema with pus formation; hyperplasia of regional lymph nodes; dilatation of sinusoids.

Treatment Oral penicillin V.

Discussion Streptococcal pharyngitis is an acute bacterial infection produced by gram-positive cocci in chains (*Streptococcus*); pharyngitis is most commonly caused by group A streptococcus. Complications due to immune-mediated cross-reactivity and molecular mimicking may include glomerulonephritis and rheumatic fever.

Atlas Link M-M1-026
**ID/CC**  A 30-year-old female presents to the surgical ER complaining of a stabbing right upper quadrant abdominal pain.

**HPI**  She is a prostitute who has been receiving treatment for gonococcal pelvic inflammatory disease.

**PE**  Right upper quadrant tenderness; cervical motion tenderness and mucopurulent cervicitis found on pelvic exam.

**Labs**  Cervical swab staining and culture identifies *Neisseria gonorrhoeae*.

**Imaging**  US: no evidence of cholecystitis. Peritoneoscopy: presence of “violin string” adhesions between liver capsule and peritoneum.

**Gross Pathology**  Adhesions noted between liver capsule and peritoneum.

**Treatment**  Antibiotic therapy (ceftriaxone and doxycycline) for patient (and for partner if warranted).

**Discussion**  Acute fibrinous perihepatitis (Fitz–Hugh–Curtis syndrome) occurs as a complication of gonococcal and chlamydial pelvic inflammatory disease and clinically mimics cholecystitis.
ID/CC  A 25-year-old male presents with sudden-onset, severe vomiting, nausea, abdominal cramps, and diarrhea.

HPI  He had returned home about 2 hours after attending a birthday party at which meat and milk were served in various forms. The friend who was celebrating his birthday reported similar symptoms.

PE  VS: no fever. PE: mild dehydration; diffuse abdominal tenderness; increased bowel sounds.

Labs  Toxigenic staphylococcus recovered from culturing food. Coagulase-positive staphylococcus cultured from nose of one of the cooks at party.

Micro Pathology  No mucosal lesions.

Treatment  Fluid and electrolyte balance; antibiotics not indicated.

Discussion  Staphylococcus aureus food poisoning results from the ingestion of food containing preformed heat-stable enterotoxin B. Outbreaks of staphylococcal food poisoning occur when food handlers who have contaminated superficial wounds or who are shedding infected nasal droplets inoculate foods such as meat, dairy products, salad dressings, cream sauces, and custard-filled pastries. The incubation period ranges from 2 to 8 hours; the disease is self-limited.
ID/CC An 11-year-old white male presents with jaundice and dark yellow urine that has been present for the last several days.

HPI He also complains of nausea, vomiting, and malaise. For the past 2 weeks, he has had a low-grade fever and mild abdominal pain. He recently returned from a vacation in Mexico, where he said he consumed a lot of shellfish.

PE Icterus; tender, firm hepatomegaly; no evidence of splenomegaly or free fluid in the peritoneal cavity.

Labs Direct hyperbilirubinemia; elevated serum transaminases (ALT > AST); moderately elevated alkaline phosphatase; prolonged PT; increased urinary urobilinogen and bilirubin; positive IgM antibody to hepatitis A (HAV) indicative of active HAV infection.

Gross Pathology May often appear normal.

Micro Pathology Multifocal hepatocellular necrosis with Councilman bodies; lymphocytic infiltrates around necrotic foci; loss of lobular architecture.

Treatment Supportive management; passive vaccination available.

Discussion In hepatitis A infection, virus is shed 14 to 21 days before the onset of jaundice; patients are no longer infectious 7 days after the onset of jaundice. It is spread by fecal-oral transmission and is endemic in areas where there are contaminated water sources. There is no chronic carrier state; recovery takes place in 6 to 12 months. HAV is a naked, single-stranded RNA virus of the picornavirus family. A killed vaccine is available; passive immunization in the form of immune serum globulins is also available.
A 25-year-old male medical student presents with jaundice and dark yellow urine.

He admits to having experienced an accidental needle stick 2 months ago, which he did not report. He also complains of nausea, low-grade fever, and loss of appetite.

Icterus; tender, firm hepatomegaly; no evidence of ascites or splenomegaly.

Direct hyperbilirubinemia; elevated serum transaminases (ALT > AST); mildly elevated alkaline phosphatase; HBsAg positive; IgM anti-HBc positive (present during window period).

US, abdomen: hepatomegaly; increased echogenicity.

Liver may be enlarged, congested, or jaundiced; in fulminant cases of massive hepatic necrosis, liver becomes small, shrunken, and soft (acute yellow atrophy).

Liver biopsy reveals hepatocellular necrosis with Councilman bodies and ballooning degeneration; inflammation of portal areas with infiltration of mononuclear cells (small lymphocytes, plasma cells, eosinophils); prominence of Kupfer cells and bile ducts; cholestasis with bile plugs.

Supportive care; follow up to determine continued presence of HBsAg for at least 6 months as sign of chronic hepatitis; vaccine available for prevention.

Hepatitis B immune globulin plus hepatitis B vaccine are recommended for parenteral or mucosal exposure to blood and for newborns of HBsAg-positive mothers. The infection is divided into the prodromal, icteric, and convalescent phases; 5% proceed to chronic hepatitis with increased risk for cirrhosis and hepatocellular carcinoma. Unlike hepatitis A, hepatitis B has a long incubation period (3 months). Hepatitis B virus is an enveloped, partially circular DNA virus of the hepadna family that contains a DNA-dependent DNA polymerase. The continued presence of HBsAg after infection has clinically resolved indicates a chronic carrier state.

Hepatitis B—Acute
ID/CC  A 30-year-old male is referred for an evaluation of intermittent jaundice over the past 2 years.

HPI  He also complains of diarrhea, skin rash, and weight loss. He received a blood transfusion 3 years ago, when he was injured in a motorcycle accident. He denies any IV drug use or any history of neuropsychiatric disorders in his family.

PE  Icterus; firm, tender hepatomegaly; splenomegaly; no evidence of ascites; no Kayser-Fleischer rings found on slit-lamp examination (vs. Wilson’s disease).

Labs  Direct hyperbilirubinemia; markedly raised serum transaminase levels; hepatitis B (HBV) serology negative; enzyme immunoassay of antibodies to structural and nonstructural enzyme proteins of hepatitis C (C200, C33c, C22-3) positive.

Micro Pathology  On liver biopsy, presence of ballooning degeneration; fatty changes; portal inflammation with necrosis of hepatocytes within parenchyma or immediately adjacent to portal areas ("piecemeal necrosis").

Treatment  Ribavirin and α interferon; supportive management.

Discussion  Hepatitis C belongs to the flavivirus family and is currently the most important cause of post-transfusion viral hepatitis; 90% of cases involve percutaneous transmission. Greater than 50% of cases progress to chronic hepatitis, leading to cirrhosis in 20%.

Atlas Link  UG-30 M-M1-031
ID/CC A 10-year-old male complains of generalized weakness, faintness on exertion, and occasional epigastric pain.

HPI His mother has noticed that he often eats soil and other inedible things (PICA).

PE Pallor; puffy face and dependent edema.

Labs CBC: microcytic, hypochromic anemia; eosinophilia. Low serum iron and ferritin; elevated serum transferrin; reduced bone marrow hemosiderin; hypoproteinemia; stool exam revealed eggs of *Ancylostoma duodenale* (ovoid eggs with thin transparent shell that reveal the segmented embryo within).

Treatment Albendazole or mebendazole; iron supplementation to treat iron deficiency anemia.

Discussion Infection with hookworms, either *Ancylostoma duodenale* or *Necator americanus*, is more likely where insanitary conditions exist; individuals at risk include children, gardeners, plumbers or electricians who are in contact with soil, and armed-forces personnel. Hookworm eggs excreted in the feces hatch in the soil, releasing larvae that develop into infective larvae. Percutaneous larval penetration is the principal mode of human infection. From the skin, hookworm larva travel via the bloodstream to the lungs, enter the alveoli, ascend the bronchotraheal tree to the pharynx, and are swallowed. Although transpulmonary larval passage may elicit a transient eosinophilic pneumonitis (LOFFLER'S PNEUMONITIS), this phenomenon is much less common with hookworm infections than with roundworm infections. The major health impact of hookworm infection, however, is iron loss resulting from the 0.1 to 0.4 mL of blood ingested daily by each adult worm. In malnourished hosts, such blood loss can lead to severe iron deficiency anemia.
A 14-year-old malnourished child died soon after hospitalization due to an extensive small bowel rupture and shock.

He had presented to the emergency room with massive bloody diarrhea. His history at admission revealed the presence of abdominal pain, fever, and diarrhea of a few days' duration; his symptoms had developed after he ate leftover meat at a fast-food restaurant.

He was dehydrated, pale, and hypotensive at time of admission and developed signs of peritonitis and shock shortly before his death.

Culture and exam of necrotizing intestinal lesions isolated Clostridium perfringens type C producing beta toxin.

Autopsy revealed ruptured small intestine, mucosal ulcerations, and gas production in the wall.

Microscopic exam revealed necrosis and acute inflammation in the ileum.

Patient died despite aggressive fluid and electrolyte replacement, bowel decompression, and antibiotic therapy (penicillin, clindamycin, or doxycycline); surgery had been planned in view of rupture of the small bowel.

Necrotizing enterocolitis is a condition affecting poorly nourished persons who suddenly feast on meat (pigs). It is associated with Clostridium perfringens type C and beta enterotoxin; beta toxin paralyzes the villi and causes friability and necrosis of the bowel wall. Immunization of children in New Guinea with beta-toxoid vaccine has dramatically decreased the incidence of the disease.
ID/CC  A 7-year-old male who has been hospitalized for treatment of acute lymphocytic leukemia complains of copious watery diarrhea, right lower quadrant abdominal pain, and fever.

HPI  He was diagnosed as neutropenic (due to aggressive cytotoxic chemotherapy) a few days ago.

PE  VS: fever; tachycardia; tachypnea. PE: pallor; sternal tenderness; axillary lymphadenopathy; hepatosplenomegaly; abdominal distention; moderate dehydration.

Labs  CBC: severe neutropenia; anemia; thrombocytopenia. PBS and bone marrow studies suggest he is in remission; blood culture grows Clostridium septicum.

Imaging  CT, abdomen: thickening of cecal wall.

Gross Pathology  Mucosal ulcers and inflammation in ileocecal region of small intestine.

Treatment  Aggressive supportive measures; surgical intervention; appropriate antibiotics (penicillin G, ampicillin, or clindamycin).

Discussion  Neutropenic enterocolitis is a fulminant form of necrotizing enteritis that occurs in neutropenic patients; neutropenia is often related to cyclic neutropenia, leukemia, aplastic anemia, or chemotherapy. In postmortem exams of patients who have died of leukemia, infections of the cecal area (typhilitis) are frequently found; Clostridium septicum is the most common organism isolated from the blood of such patients.
A 25-year-old male complains of midepigastric pain that usually begins 1 to 2 hours after eating and occasionally awakens him at night.

The patient has been diagnosed with duodenal ulcers several times in the past, but his symptoms have consistently recurred even after therapy with H$_2$ blockers, antacids, and sucralfate.

VS: stable. PE: pallor; epigastric tenderness on deep palpation.

CBC: normocytic, normochromic anemia. Stool positive for occult blood.

UGI: ulcerations in antrum of stomach and duodenum; antral biopsy specimens yield positive urease test.

Gross Pathology
Grossly round ulcer (may also be oval) seen as sharply punched-out defect with relatively straight walls and slight overhanging of mucosal margin (heaped-up margin is characteristic of a malignant lesion); smooth and clean ulcer base.

Micro Pathology
No evidence of malignancy; antral biopsies reveal presence of chronic mucosal inflammation.

Triple therapy with amoxicillin, metronidazole, and bismuth subsalicylate; triple therapy with clarithromycin, omeprazole, and tinidazole is now considered effective and relatively free of side effects.

Helicobacter pylori grows overlying the antral gastric mucosal cells; 40% of healthy individuals and approximately 50% of patients with peptic disease harbor this organism. Although H. pylori does not breach the epithelial barrier, colonization of the antral mucosal layer by this organism is associated with structural alterations of the gastric mucosa and hence with a high prevalence of antral gastritis. Despite the fact that H. pylori does not grow on duodenal mucosa, it is strongly associated with duodenal ulcer, and eradication of the organism in patients with refractory peptic ulcer disease decreases the risk of recurrence.

ID/CC A 4-year-old male is brought to the physician by his parents, who complain that the child has had intense perianal itching, especially during the night.

HPI The child is otherwise healthy, and his developmental progress is normal.

PE Perianal excoriation noted.

Labs Cellulose adhesive tape secured to perianal area during the night reveals presence of *Enterobius vermicularis* eggs that were flattened on one side, were embryonated, and had a thick shell; no parasites found on stool exam.

Treatment Strict personal hygiene; drugs used include albendazole, mebendazole, piperazine, and pyrantel pamoate.

Discussion Infection is caused by *Enterobius vermicularis*. Adult worms are located primarily in the cecal region; female adult worms migrate to the perianal area during the night and deposit their eggs. Direct person-to-person infection occurs by ingestion and swallowing of eggs; autoinoculation occurs by contamination of fingers. The life cycle is completed in about 6 weeks.
A 10-month-old male presents with fever and severe vomiting followed by watery diarrhea.

His stools are loose and watery without blood or mucus.

VS: fever; tachycardia. PE: child is irritable; moderate dehydration.

Absence of leukocytes on fecal stain; rotavirus detected with ELISA; electron microscopy with negative staining identifies rotavirus on stool ultrafiltrates.

Major histopathologic lesions are characterized by reversible involvement of the proximal small intestine; mucosa remains intact with shortening of villi, a mixed inflammatory infiltration of lamina propria, and hyperplasia of the mucosal crypt cells; electron microscopy reveals distended cisterns of endoplasmic reticulum, mitochondrial swelling, and sparse, irregular microvilli.

Fluid replacement therapy.

Rotavirus group A is the single most important cause of endemic, severe diarrheal illness in infants and young children worldwide; it occurs with greater frequency during winter months in temperate climates and during the dry season in tropical climates. In the United States, rotavirus accounts for 50% of all childhood diarrheas, has an incubation period of 48 hours, is transmitted by the fecal-oral route, and lasts only a few days. Some children subsequently develop lactose intolerance, which lasts for a few weeks.
A 30-year-old male presents with sudden-onset, crampy abdominal pain and diarrhea.

The diarrhea is watery and contains mucus. The patient also complains of low-grade fever with chills, malaise, nausea, and vomiting. Careful history reveals that he had ingested partially cooked eggs at a poultry farm 24 hours before his symptoms began.

VS: fever; tachycardia. PE: mild diffuse abdominal tenderness; mild dehydration.

Stool culture yields Salmonella typhimurium; stained stool demonstrates PMNs.

Intestinal mucosal erythema (limited to the colon) and some superficial ulcers.

Mixed inflammatory infiltrate in mucosa; superficial epithelial erosions.

Fluid and electrolyte replacement therapy; antibiotics withheld, as they prolong carrier state. Antibiotic therapy only for malnourished, severely ill, bacteremic, and sickle cell disease patients.

Salmonella infection is acquired through the ingestion of food (eggs, meat, poultry) or water contaminated with animal or human feces; individuals with low gastric acidity are also susceptible.
A 50-year-old alcoholic white male presents with fever, abdominal pain, and rapidly progressive distention of the abdomen.

**HPI**  
He was diagnosed with alcoholic cirrhosis 1 month ago, when he was admitted to the hospital with jaundice and hematemesis.

**PE**  
VS: fever. PE: icterus; on palpation, abdominal tenderness with guarding; fluid thrill and shifting dullness to percussion (due to ascites); splenomegaly; decreased bowel sounds.

**Labs**  
CBC: leukocytosis. Ascitic fluid leukocyte count > 500/cc; PMNs (350/cc) elevated; ascitic proteins and glucose depressed; gram-negative bacilli in ascitic fluid; Escherichia coli isolated in culture; elevated AST and ALT (AST > ALT).

**Imaging**  
KUB: ground-glass haziness (due to ascites); no evidence of free air. US, abdomen: cirrhotic shrunken liver; ascites; splenomegaly; increased portal vein diameter and flow. EGD: esophageal varices.

**Gross Pathology**  
Fibrinopurulent exudate covering surface of peritoneum; fibrosis may lead to formation of adhesions.

**Micro Pathology**  
PMNs and fibrin on serosal surfaces in various stages with presence of granulation tissue and fibrosis.

**Treatment**  
Specific organism-sensitive antibiotics or empiric therapy (such as cefotaxime or β-lactamase-resistant penicillin) for gram-negative aerobic bacilli and gram-positive cocci; supportive treatment for cirrhosis.

**Discussion**  
The spontaneous or primary form of peritonitis occurs in patients with advanced chronic liver disease and concomitant ascites; E. coli is the most common cause of secondary peritonitis.
A 25-year-old male U.S. citizen on vacation in Mexico presents with abrupt-onset explosive watery diarrhea, abdominal cramps, and a low-grade fever and chills.

The patient does not complain of tenesmus or passage of blood or mucus in his stools, but he does complain of a feeling of urgency to defecate.

VS: low-grade fever. PE: unremarkable.

No erythrocytes, WBCs, or parasites seen in stained stool; bioassays for enterotoxigenic *Escherichia coli* (ETEC) reveal presence of the labile enterotoxin (LT) (tests available only for research purposes).

Fluid replacement; antibiotics (fluoroquinolone or TMP-SMX) with loperamide; prevention with careful hygienic practices and prophylactic fluoroquinolone or bismuth subsalicylate with loperamide.

Traveler’s diarrhea is a self-limited condition that develops within 1 to 2 days of ingestion of contaminated food or drinks. Over three-fourths of cases of traveler’s diarrhea are caused by bacteria, with enterotoxigenic *E. coli* the most frequent cause (may also be caused by enteropathogenic *E. coli* and, in Mexico, by an enteroadherent *E. coli*). Other common pathogens include *Shigella* species, *Campylobacter jejuni*, *Aeromonas* species, *Plesiomonas shigelloides*, *Salmonella* species, and noncholera vibrios. Rotavirus and Norwalk agent are the most common viral causes; *Giardia*, *Cryptosporidium*, and, rarely, *Entamoeba histolytica* are parasitic pathogens. Enterotoxigenic *E. coli* produce enterotoxins that bind to intestinal receptors and activate adenyl cyclase in the intestinal cell to produce an increase in the level of the cyclic nucleotides cAMP (LT, labile toxin) and cGMP (ST, stable toxin), which markedly augments sodium, chloride, and water loss, thereby producing a secretory diarrhea.
ID/CC A 30-year-old male presents with sudden-onset fever, colicky abdominal pain, and watery diarrhea.

HPI He had eaten raw oysters at a friend's party the day before (incubation period 4 hours to 4 days).

PE VS: fever; tachycardia. PE: no dehydration; diffuse abdominal tenderness; increased bowel sounds.

Labs *Vibrio parahaemolyticus* isolated from stool in a high-salt content (halophilic vibrio) culture medium; PMNs in stool; Kanagawa phenomenon (beta-hemolysis on medium containing human blood; done as an indicator for pathogenicity) positive.

Treatment Fluid and electrolyte balance; antibiotics not required (since they do not shorten course of infection).

Discussion Seafood is the main source of the organism. After ingestion, *Vibrio parahaemolyticus* multiplies in the gut and produces a diarrheal enterotoxin.
ID/CC  A 35-year-old male presents to the emergency room with high-grade fever, marked weakness, and a hemorrhagic vesiculobullous skin eruption.

HPI  He had just returned from deep-sea fishing in the Gulf of Mexico, where he had consumed large quantities of seafood. He has been diagnosed with chronic liver disease (due to hemochromatosis).

PE  VS: fever; hypotension; tachycardia. PE: icterus; vesiculobullous skin lesions seen on an otherwise-bronzed complexion.

Labs  Blood culture on high-salt medium (halophilic bacteria) reveals growth of *Vibrio vulnificus*; evidence of hemochromatosis (hyperglycemia, hyperbilirubinemia, increased serum iron).

Treatment  Ceftazidime and doxycycline, ciprofloxacin; supportive.

Discussion  Halophilic *Vibrio vulnificus* should be suspected and treated in any individual with chronic liver disease who presents with septicemia and skin lesions 1 to 3 days following seafood ingestion.
ID/CC A 56-year-old white male complains of diarrhea and bloating for several months along with ankle swelling.

HPI He also complains of memory loss, fever, arthritis in the knees and hands, and weight loss.

PE VS: fever. PE: thin, gaunt male; muscle wasting; swollen, tender right wrist and ankle; axillary and femoral lymphadenopathy; ecchymoses of chest and arms.

Labs CBC/PBS: macrocytic, hypochromic anemia; hypoalbuminemia; increased fecal fat (steatorrhea).

Imaging UGI/SBFT: nonspecific dilatation of small bowel.

Gross Pathology Atrophy of intestinal mucosa; inflammatory infiltrate in synovia of joints.

Micro Pathology Small bowel biopsy reveals characteristic macrophages containing bacilli with PAS reagent staining; characteristic gram-negative actinomyete bacilli in macrophages, PMNs, and epithelial cells of lamina propria; dilated lymphatics; flattening of intestinal villi.

Treatment Bactrim (TMP-SMX) or ceftriaxone for 1 year.

Discussion Caused by infection with Tropheryma whippelii; produces malabsorption of fat-soluble vitamins, protein, iron, folic acid, and vitamin B₁₂.
ID/CC  A 28-year-old female complains of painful swelling of both knees and tender skin eruptions on both shins.

HPI  For the past 2 weeks she has also had watery diarrhea that developed after she consumed some raw pork. She also complains of low-grade fever and mild abdominal pain.

PE  VS: low-grade fever; tachycardia. PE: mild dehydration; swollen and warm knee joints with painful restriction of all movements (arthritis); multiple tender, erythematous plaques and nodules (erythema nodosum) seen over both shins.

Labs  CBC: leukocytosis. *Yersinia enterocolitica* isolated from stool; patient is HLA-B27 positive.

Micro Pathology  Oval ulcers with long axis in the direction of bowel flow, similar to ulcers caused by typhoid fever (intestinal tubercular ulcers are transverse).

Treatment  Supportive; antibiotics (aminoglycosides, fluoroquinolones) indicated in severe infections.

Discussion  *Yersinia enterocolitica* is an invasive gram-negative intracellular pathogen that causes gastroenteritis, most frequently involving the distal ileum and colon (enterotoxin mediated), mesenteric adenitis (due to necrotizing and suppurative gut lesions) and ileitis (pseudappendicitis), and sepsis; infection may trigger a variety of autoimmune phenomena, including erythema nodosum, reactive arthritis, and possibly Graves' disease, especially in HLA-B27-positive individuals. Spread is by the fecal-oral route and occurs via contaminated milk products or water, swine, or household pet feces.
A 3-year-old albino male is referred to a specialist for an evaluation of a suspected immune deficiency.

His parents report recurrent respiratory, skin, and oral infections with gram-negative and gram-positive organisms. He also has a history of bruising easily.

Partial albinism; light-brown hair with silvery tint; nystagmus; photophobia on eye reflex exam; chronic gingivitis and periodontitis; purpuric patches over areas of repeated minimal trauma; mild hepatomegaly; no lymphadenopathy.

CBC/PBS: decreased neutrophil count with normal platelet count; large cytoplasmic granules (giant lysosomes) in WBCs on Wright-stained peripheral blood smears. Prolonged bleeding time; impaired platelet aggregation; normal clotting time and PTT; normal nitroblue tetrazolium test.

Largely supportive; ascorbic acid, prophylactic antibiotics, acyclovir.

Chédiak–Higashi syndrome is an autosomal-recessive disorder that is due to a defect in polymerization of microtubules in leukocytes that causes impairment of chemotaxis, phagocytosis, and formation of phagolysosomes. Patients with this disorder usually present with recurrent pyogenic staphylococcal and streptococcal infections.
An 8-year-old child with sickle cell anemia is seen with complaints of sudden-onset pallor of the skin and mucous membranes, fatigue, and malaise.

The child suffered a mild prodromal illness before developing severe pallor.

VS: no fever; tachycardia; tachypnea; BP normal. PE: severe pallor; mild icterus; no lymphadenopathy, splenomegaly, or hepatomegaly noted.

CBC: severe anemia (Hb 2 g/dL); reduced leukocyte and platelet counts; mild hyperbilirubinemia; absent reticulocytes and sickled RBCs on peripheral blood smear.

Bone marrow biopsy reveals increased numbers of giant pronormoblasts (diagnostic of parvovirus infection).

Blood transfusions to tide over the crises. Spontaneous recovery in 1 to 2 weeks.

Parvovirus infection is the cause of transient aplastic crises (may also be due to folic acid deficiency) that occur in patients who have severe hemolytic disorders; cessation of erythropoiesis for about 10 days in a normal adult as a result of parvovirus infection would produce a 10% drop in hemoglobin concentration (i.e., a fall of 1% daily would lead to a decline in hemoglobin concentration of 1 to 2 g/dL after 10 days). A patient with severe hemolysis in whom the bone marrow is turning over at a rate seven times normal would experience a 70% decrease in hemoglobin concentration (i.e., a drop from 10 g/dL to 3 g/dL) as a result of a 10-day cessation of erythropoiesis. Although parvovirus can affect all precursor cells, the red cell precursors are most profoundly affected.

Atlas Link: HCV1 H-M1-046
ID/CC  A 35-year-old Finnish man complains of easy fatigability and shortness of breath.

HPI  He often eats undercooked or raw freshwater fish. He also reports vague digestive disturbances such as anorexia, heartburn, and nausea.

PE  PE: pallor.

Labs  CBC/PBS: megaloblastic anemia. Blood vitamin B$_{12}$ levels low; stool exam reveals presence of operculated eggs and proglottids of *Diphyllobothrium latum*.

Treatment  Niclosamide or praziquantel.

Discussion  *Diphyllobothrium latum* (fish tapeworm) infection is found in cold climates where raw or undercooked fish are eaten. The adult worm attaches to the human jejunum and competes for absorption of vitamin B$_{12}$, producing a deficiency that resembles pernicious anemia. Prevention includes proper preparation of fish.
A 45-year-old male with refractory acute myeloid leukemia who underwent a bone marrow transplant from a nonidentical donor presents with an extensive skin rash, severe diarrhea, and jaundice.

Prior to the transplant, which occurred 2 months ago, he received preparative chemotherapy and radiotherapy along with broad-spectrum antibiotics. Engraftment was confirmed within 4 weeks by rising leukocyte counts.

VS: BP normal. PE: patient is cachectic and moderately dehydrated; icterus noted; violaceous, scaly macules and erythematous papules resembling lichen planus seen over extremities.

CBC: falling blood counts; relative eosinophilia. Elevated direct serum bilirubin and transaminases; stool exam reveals no infectious etiology; skin biopsy taken.

Skin biopsy specimens reveal vacuolar changes of basal cell layer with perivenular lymphocytic infiltrates (CD8+ T cells).

High-dose cyclosporine therapy, rabbit anti-thymocyte globulin, methylprednisolone or anti-T-cell monoclonal antibodies.

Approximately 30% of bone marrow transplant recipients develop graft-versus-host disease (GVHD). This attack is primarily launched by immunocompetent T lymphocytes derived from the donor’s marrow against the cells and tissues of the recipient, which it recognizes as foreign. Cyclosporin A is effective for prevention of GVHD.
A 20-year-old male presents with an extensive purpuric skin rash, oliguria, and marked weakness; he also complains of bloody diarrhea of 1 week's duration.

The patient ate a hamburger at a fast-food restaurant 2 to 3 days prior to the onset of his diarrhea. He has no associated fever.

VS: no fever. PE: dehydration; pallor; extensive purpuric skin rash.

Stool examination reveals presence of RBCs but no inflammatory cells or parasites; culture isolates sorbitol-negative *Escherichia coli*; serotyping studies and effect on HeLa cell culture reveal presence of *enterohemorrhagic E. coli* (EHEC) serotype O157:H7; elevated BUN and creatinine. CBC/PBS: microangiopathic anemia and thrombocytopenia. PT, PTT normal.

Sigmoidoscopy: moderately hyperemic mucosa with no evidence of any ulceration.

Pathology localized to kidney, where hyaline thrombi were seen in afferent arterioles and glomerular capillaries.

Dialysis and blood transfusion for management of HUS; fluid and electrolyte maintenance; antimicrobial therapy. Most patients who develop HUS as a complication of *E. coli* hemorrhagic colitis die as a result of hemorrhagic complications.

Hemorrhagic colitis associated with a Shiga-like toxin producing *EHEC* O157:H7 is characterized by grossly bloody diarrhea with remarkably little fever or inflammatory exudate in stool; a significant number of patients develop potentially fatal HUS. EHEC infections can be largely prevented through adequate cooking of beef, especially hamburgers.

**Atlas Link** UCMR B-M1-049

**HEMOLYTIC-UREMIC SYNDROME (HUS)**
ID/CC  A 34-year-old male presents to his primary care physician with a hard, red, painless swelling on his left mandible that has slowly been growing over the past few weeks and has now begun to drain pus.

HPI  The patient recently had a tooth extraction.

PE  No acute distress; no other significant findings.

Labs  Gram stain of exudate reveals branching gram-positive filaments and characteristic “sulfur granules”; non-acid-fast and anaerobic (distinguishes actinomyces from Nocardia).

Imaging  XR: no bony destruction.

Gross Pathology  Sinus tracts from region of infection to surface with granular exudate.

Micro Pathology  Granulation tissue and fibrosis surrounding a central suppurative necrosis; granulation tissue may also enclose foamy histiocytes and plasma cells.

Treatment  Amoxicillin followed by amoxicillin or penicillin G followed by oral penicillin V and, if necessary, surgical drainage and removal of necrotic tissue.

Discussion  Actinomyces israelii is a part of the normal flora of the mouth (crypts of tonsils and tartar of teeth), so most patients have a history of surgery or trauma. There is no person-to-person spread. Actinomycosis is a chronic suppurative infection and can also involve the abdomen or lungs, especially following a penetrating trauma such as a bullet wound or an intestinal perforation. Pelvic disease is associated with IUD use. Spread occurs contiguously, not hematogenously.

Atlas Link  [Image] M-MI-050
ID/CC  A 7-month-old girl is brought to the pediatric clinic with wheezing, respiratory difficulty, and nasal congestion of 3 hours' duration.

HPI  She has had rhinorrhea, fever, and cough and had been sneezing for 2 days prior to her visit to the clinic.

PE  VS: tachypnea. PE: nasal flaring; mild central cyanosis; accessory muscle use during respiration; hyperexpansion of chest; expiratory and inspiratory wheezes; rhonchi over both lung fields.

Labs  CBC/PBS: relative lymphocytosis. ABGs: hypoxemia with mild hypercapnia. Respiratory syncytial virus (RSV) demonstrated on viral culture of throat swab.

Imaging  CXR: hyperinflation; segmental atelectasis; interstitial infiltrates.

Micro Pathology  Severe bronchiolitis produces bronchiolar epithelial necrosis, lymphocytic infiltrate, and alveolar exudates.

Treatment  Humidified oxygen, bronchodilators, aerosolized ribavirin.

Discussion  RSV is the most common cause of bronchiolitis in infants under 2 years of age; other viral causes include parainfluenza, influenza, and adenovirus. RSV shedding may last 2 or more weeks in children.
An **8-year-old** female presents with pain and swelling of her knee joints, elbows, and lower limbs along with **fever** for the past 2 weeks; she also complains of shortness of breath (dyspnea) on exertion.

**HPI**  
The patient had a **sore throat 2 weeks ago**.

**PE**  
VS: fever. PE: **blanching, ring-shaped erythematous rash over trunk and proximal extremities** (**erythema marginatum**); **subcutaneous nodules** at occiput and below extensor tendons in elbow; **swelling with redness of both knee joints and elbows** (**polyarthritis**); painfully restricted movement; pedal edema; increased JVP; high-frequency apical systolic murmur with radiation to axillae (**mitral valve insufficiency due to carditis**); bilateral fine inspiratory basal crepitant rales; mild, tender hepatomegaly.

**Labs**  
CBC: leukocytosis. *Streptococcus pyogenes* on throat swab; markedly elevated ASO titers; elevated ESR; elevated C-reactive protein (CRP); negative blood culture. ECG: prolonged P-R interval.

**Imaging**  
CXR: cardiomegaly; increased pulmonary vascular markings. Echo: vegetations over mitral valve with regurgitation.

**Gross Pathology**  
Acute form characterized by **endo-, myo-, and pericarditis** (**pancecarditis**); chronic form characterized by fibrous scarring with calcification and mitral stenosis with verrucous fibrin deposits.

**Micro Pathology**  
Myocardial muscle fiber necrosis enmeshed in collagen; characteristic finding is fibrinoid necrosis surrounded by **perivascular accumulation of mononuclear inflammatory cells** (**Aschoff cells**).

**Treatment**  
Aspirin, corticosteroids, and diuretics; penicillin or erythromycin.

**Discussion**  
Acute rheumatic fever is a sequela of upper respiratory infection with group A, β-hemolytic streptococcus; it causes **autoimmune** damage to several organs, primarily the heart. The systemic effects of acute rheumatic fever are immune mediated and are secondary to cross-reactivity of host antistreptococcal antibodies.

**Atlas Link**  
[UCV1 M-M1-052]
ID/CC A 48-year-old missionary who has lived in Cameroon, West Africa, for 20 years is airlifted home because of lethargy, nuchal rigidity, persistent headache, and drowsiness that have not responded to antibiotics and supportive treatment.

HPI He states that over the years he has been bitten in the neck several times by a mutumutu, or tsetse fly (Glossina palpalis). He has also had intermittent, generalized erythematous rashes accompanied by fever.

PE Alert but somewhat incoherent and confused; sometimes delusional; nuchal rigidity and tremors of face and lips; splenomegaly; generalized rubbery, painless lymphadenopathy, predominantly in posterior neck and supraclavicular areas (Winterbottom's sign).

Labs PBS/LP: hypercellular, trypanosomal forms present; lymphocytes in CSF. Elevated IgM.

Gross Pathology Chancre with erythema and induration at bite site; chancre resolves spontaneously; spleen and lymph nodes enlarged during systemic stage; leptomeninges enlarged during CNS involvement.

Micro Pathology Skin: edema, mononuclear cell inflammation, organisms, and endothelial proliferation; spleen and lymph nodes: histiocytic hyperplasia; CNS: mononuclear cell meningoencephalitis.

Treatment Suramin; pentamidine or eflornithine.

Discussion Also called sleeping sickness. African trypanosomiasis is a systemic febrile disease endemic to Africa whose chronic form causes a meningoencephalitis. It is caused by the flagellated protozoans Trypanosoma brucei gambiense (West African) and Trypanosoma brucei rhodesiense (East African), which are transmitted by the tsetse fly.
A 28-year-old male homosexual complains of continuous low-grade fever, weight loss, and diarrhea of 1 month’s duration.

HPI He also complains of an extensive skin rash, mucous membrane eruptions, recurrent herpes zoster infection, and oral ulcerations. He reports practicing receptive anal intercourse.

PE VS: low-grade fever. PS: cachectic; generalized lymphadenopathy; maculopapular rash; severe seborrheic dermatitis; aphthous ulcers; white confluent patch with corrugated surface (oral hairy leukoplakia) along lateral borders of tongue; penile warts (condylomata acuminata); extensive multiple pruritic, pink, umbilicated papules 2 to 5 mm in diameter (molluscum contagiosum).

Labs CBC: anemia; leukopenia with lymphopenia; thrombocytopenia. Low CD4+ count; elevated CD8+ T-cell count; ELISA for HIV-1 positive; Western blot confirmatory; PCR for viral RNA (investigation of choice in window period) positive.

Micro Pathology Oral hairy leukoplakia; lesions show keratin projections resembling hairs, koilocytosis, and little atypia; hybridization techniques reveal Epstein–Barr virus in lesions.

Treatment Prophylactic antibiotics for prevention of opportunistic infections while monitoring CD4+ T-cell counts; antiretroviral drugs (zidovudine, didanosine, zalcitabine, and protease inhibitors); counseling and rehabilitative measures.

Discussion AIDS-related complex (ARC) consists of symptomatic conditions in an HIV-infected patient that are not included in the AIDS surveillance case definition and that meet at least one of the following criteria: (1) the conditions are indicative of a defect in cell-mediated immunity; or (2) the conditions have a clinical course or management that is complicated by HIV infection.

Atlas Link 2C172 Z-M1-054
A 28-year-old male from India complains of gradual-onset, intermittent, crampy abdominal pain with one to four foul-smelling, frothy loose stools daily.

His stools sometimes contain blood and mucus. He also complains of flatulence, tenesmus, and, at times, alternating diarrhea and constipation.

Slight tenderness during palpation of cecum and ascending colon; no hepatomegaly.

CBC: mild leukocytosis; no eosinophilia. Fresh stool examination reveals presence of Entamoeba histolytica cysts and motile hematophagous trophozoites; serology for antiamebic antibodies is positive.

Colonoscopy: multiple colonic mucosal ulcers that are slightly raised and covered with shaggy exudate; mucosa between ulcers normal.

Biopsy specimens reveal lesions extending under adjacent intact mucosa to produce classical “flask-shaped” ulcers; amebic trophozoites demonstrated at base of ulcer.

Metronidazole (drug of choice) followed by paromomycin or iodoquinol.

Entamoeba histolytica cysts are infective and are transmitted through contaminated water, raw vegetables, food handlers, and fecal-oral or oral-anal contact. The sites of involvement, in order of frequency, are the cecum and ascending colon, rectum, sigmoid colon, appendix, and terminal ileum. Trophozoites are the invasive form of the organism, causing colitis or distant infection by hematogenous spread. Complications include perforation of the bowel; liver abscess with pleural, pericardial, or peritoneal rupture; bowel obstruction by ameboma; and skin ulcers around the perineum and genitalia.

[Atlas Link: M-M1-055]
A 45-year-old male Peace Corps volunteer who recently spent 2 years in rural Mexico complains of a spiking fever, malaise, headache, and right upper quadrant abdominal pain.

He admits to having had bloody diarrhea with mucus (dysentery) and tenesmus that disappeared with some pills that he took several months ago.

VS: fever (39.6°C). PE: pallor; slight jaundice; tender 3+ hepatomegaly with no rebound tenderness; pain on fist percussion of right lower ribs.

CBC: leukocytosis with neutrophilia. Amebic cysts in stool specimen (not concurrent with abscess); positive serology for antibodies to Entamoeba histolytica.

CXR: elevation of right hemidiaphragm; small right pleural effusion. CT/US: cavitating lesion in right lobe of liver (due to abscess).

Multiple colonic mucosal ulcers, slightly raised and covered with shaggy exudate; enlarged liver with one large abscess on right lobe containing chocolate-colored pus; abscess may rupture and spread to lungs, brain, or other organs.

Sterile pus; ameba may be obtained from periphery of lesion.

Metronidazole; needle evacuation; surgery in case of treatment failure or rupture.

Prior travel to endemic areas plus a triad of fever, hepatomegaly, and right upper quadrant pain are hallmarks of hepatic liver abscess. Colitis preceeds the liver abscess; amebas then invade the gut wall and enter portal circulation.
A 15-year-old male who resides in Florida presents with nausea and vomiting, fever, and marked neck stiffness.

He also complains of a severe bifrontal headache. Careful history reveals that he swam for several hours in brackish water approximately a week ago.

VS: fever; tachycardia. PE: signs of meningeal irritation (neck rigidity, positive Kernig's sign and Brudzinski's sign); funduscoppy reveals mild papilledema.

Labs LP: bloody CSF (raised RBC count may also be due to examiner's inability to recognize proliferating amebas) shows intense neutrophilia, pleocytosis, high protein, and low sugar; no organism seen on Gram, ZN, or India ink staining of CSF; wet preparation of CSF reveals viable Naegleria trophozoites; diagnosis confirmed using direct fluorescent antibody staining.

Gross Pathology Lesions are mostly present in the olfactory nerves and brain. Focal hemorrhages, extensive fibrinoid necrosis, and blood vessel thrombosis with nerve tissue necrosis.

Micro Pathology Naegleria fowleri trophozoites seen as 10- to 20-μm-diameter organisms with large nucleus, small granular cytoplasm, distinct ectoplasm, and bulbous pseudopodia.

Treatment Intracisternal and IV amphotericin B, miconazole, rifampin; prognosis is very poor.

Discussion Primary amebic meningoencephalitis is caused by amebas of the genus Naegleria or Acanthamoeba. The former most often affects children and young adults, appears to be acquired by swimming in warm, fresh/brackish water, and is almost always fatal, with the ameba gaining entry into the arachnoid space through the nasal cribriform plate. Acanthamoeba infections involve older, immunocompromised individuals and are sometimes characterized by spontaneous recovery.
ID/CC A 30-year-old male goes to the emergency room because of dyspnea, cyanosis, hemoptysis, and chest pain.

HPI He has had a high fever, malaise, and a nonproductive cough for 1 week. The patient is a sheep farmer who remembers having been treated for dark black skin lesions in the past.

PE VS: fever. PE: dyspnea; cyanosis; bilateral rales heard over lungs.

Labs CBC: normal. Negative blood and sputum cultures; diagnosis of anthrax confirmed by fourfold increase in indirect microhemagglutination titer.

Imaging CXR: mediastinal widening. CT, chest: evidence of "hemorrhagic mediastinitis."

Gross Pathology Patchy consolidation; vesicular papules covered by black eschar.

Micro Pathology Lungs show fibrinous exudate with many organisms but few PMNs.

Treatment Isolate and treat with IV penicillin G or ciprofloxacin.

Discussion Anthrax is caused by infection with Bacillus anthracis. A cell-free anthrax vaccine is available to protect those employed in industries associated with a high risk of anthrax transmission (farmers, veterinarians, tannery or wool workers).
ID/CC  A 38-year-old male receiving cytotoxic chemotherapy (immunosuppressed) for acute leukemia presents with pleuritic chest pain, hemoptysis, fever, and chills.

HPI  He also complains of dyspnea, tachypnea, and a productive cough.

PE  VS: fever. PE: severe respiratory distress; bilateral rales heard over lungs.

Labs  CBC: severe neutropenia. Negative blood and sputum culture for bacteria.

Imaging  CXR: necrotizing bronchopneumonia.

Gross Pathology  Necrotizing bronchopneumonia; abscesses.

Micro Pathology  Lung biopsy identifies Aspergillus with septate, acutely branching hyphae (visualized by silver stains); necrotizing inflammation; vascular thrombi with hyphae (due to blood vessel invasion).

Treatment  IV amphotericin B or itraconazole.

Discussion  The most lethal form of infection, invasive aspergillosis, is seen primarily in severely immunocompromised individuals, i.e., patients with AIDS; patients with prolonged, severe neutropenia following cytotoxic chemotherapy; patients with chronic granulomatous disease; and patients receiving glucocorticoids and other immunosuppressive drugs (e.g., transplant recipients).

Atlas Links  UCM # M-M1-059A, M-M1-059B, M-M1-059C
A 50-year-old male presents to the ER with complaints of recurrent, sudden-onset, severe breathlessness, wheezing, fever, chills, and a productive cough (sometimes producing brown bronchial casts).

The patient has had steroid-dependent chronic bronchial asthma for many years and has no history of foreign travel or contact with a TB patient. He has a history of occasional hemoptyis.

VS: fever; marked tachycardia; severe tachypnea. PE: respiratory distress; central cyanosis; wheezing; rhonchi and coarse rales over both lung fields.

CBC: eosinophilia. Oxygen saturation low. Very high titers of specific IgE antibodies against Aspergillus present (specific marker for the disease); sputum cultures positive for Aspergillus; skin tests to Aspergillus antigens positive. PFTs: obstructive picture (due to underlying asthma).

CXR: segmental infiltrate in upper lobes (these infiltrates are segmental because they correspond directly to the affected bronchi); branching, fingerlike shadows from mucoid impaction of dilated central bronchi (virtually pathognomonic of allergic bronchopulmonary aspergillosis). CT, chest: evidence of proximal bronchiectasis.

Oral corticosteroids or beclomethasone.

Allergic bronchopulmonary aspergillosis (ABPA) is a hypersensitivity disorder that primarily affects the central bronchi; immediate and Arthus-type hypersensitivity reactions are involved in its pathogenesis. The onset of the disease occurs most often in the fourth and fifth decades, and virtually all patients have long-standing atopic asthma. Untreated ABPA leads to proximal bronchiectasis.
A 50-year-old alcoholic male presents with a high-grade fever, cough, copious, foul-smelling sputum, and pleuritic right-sided chest pain.

His wife reports that he was brought home in a semiconscious state a few days ago, when he was found lying on the roadside heavily under the influence of alcohol.

VS: fever. PE: signs of consolidation elicited over right middle and lower pulmonary lobes.

Sputum reveals abundant PMN leukocytes and mixed oral flora; culture yields Bacteroides melaninogenicus (Prevotella melaninogenica) and other Bacteroides species, Fusobacterium, microaerophilic streptococci, and Peptostreptococcus.

CXR: consolidation involving apical segment of right lower lobe and posterior segments of middle lobe; large cavity with air-fluid level (ABSCESS) also seen.

Clindamycin.

Alcoholism, drug abuse, administration of sedatives or anesthesia, head trauma, and seizures or other neurologic disorders are most often responsible for the development of aspiration pneumonia. Because anaerobes are the dominant flora of the upper GI tract (outnumbering aerobic or facultative bacteria by 10 to 1), they are the dominant organisms in aspiration pneumonia; of particular importance are Bacteroides melaninogenicus (Prevotella melaninogenica) and other Bacteroides species (slender, pleomorphic, pale gram-negative rods), Fusobacterium nucleatum (slender gram-negative rods with pointed ends), and anaerobic or microaerophilic streptococci and Peptostreptococcus (small gram-positive cocci in chains or clumps).
A 38-year-old HIV-positive male is admitted to the hospital with fever, rigors, night sweats, and diarrhea.

HPI He reports excessive weight loss over the past few weeks. He was treated for *Pneumocystis pneumonia* a few weeks ago and still reports a persistent productive cough.

PE VS: fever. PE: patient is extremely emaciated; hepatosplenomegaly and lymphadenopathy noted.

Labs CD4+ count < 50/μl; *Mycobacterium avium-intracellulare* isolated on blood culture; smears of tissues obtained from lymph nodes, bone marrow, spleen, liver, and lungs reveal evidence of acid-fast bacilli, and cultures yield *M. avium* intestinal infection with *M. avium* proven by culture of stools and colonic biopsy specimens.

Imaging CT, abdomen: hepatosplenomegaly; retroperitoneal lymphadenopathy; bowel mucosal fold thickening.

Micro Pathology Despite the presence of many mycobacteria and macrophages, well-formed granulomas were typically absent due to profound impairment of cell-mediated immunity.

Treatment The primary treatment regimen includes clarithromycin and ethambutol with or without rifabutin. The failure rate of therapy is high.

Discussion *Mycobacterium avium* complex is now the most frequent opportunistic bacterial infection in patients with AIDS; it typically occurs late in the course of the syndrome, when other opportunistic infections and neoplasia have already occurred. Prophylaxis against *M. avium-intracellulare* is recommended in AIDS patients with a CD4+ count of < 100/μl (administer azithromycin, clarithromycin, or rifabutin).
A 20-year-old male from India presents to the ER with severe nausea and vomiting.

Careful history reveals that 2 hours ago he ate some unrefrigerated fried rice that his wife had cooked the night before. He does not complain of any fever or diarrhea (may or may not be present).

VS: no fever. PE: mild dehydration; diffuse mild abdominal tenderness.

Fecal staining reveals no RBCs, WBCs, or parasites; Bacillus cereus, a gram-positive rod, isolated from vomitus and stool and shown to produce the emetogenic enterotoxin.

Supportive.

Bacillus cereus causes two distinct syndromes: a diarrheal form (mediated by an Escherichia coli LT-type enterotoxin with an incubation period of 8 to 16 hours; caused by meats and vegetables) and an emetic form (mediated by a Staphylococcus aureus-type enterotoxin with an incubation period of 1 to 8 hours; caused by fried rice). Proper food handling and refrigeration of boiled rice are largely preventive.
ID/CC  A 30-year-old male who recently emigrated from Peru presents with an extensive nodular skin eruption, mild arthralgias, and occasional fever.

HPI  One month ago, the patient had a high-grade fever that was accompanied by excessive weakness, dyspnea, and passage of cola-colored urine; the fever subsided after 2 weeks, but his weakness has progressed since that time.

PE  Pallor; mild icterus; extensive skin rash comprising purplish nodular lesions of varying sizes seen on face, trunk, and limbs; mild hepatosplenomegaly; funduscropy reveals retinal hemorrhages.

Labs  Intraerythrocytic coccobacillary-form bacteria visible in thick and thin films stained with Giemsa; bacteria seen and isolated from skin lesions; indirect serum bilirubin elevated. PBS: macrocytic, hypochromic anemia with polychromasia; marked reticulocytosis (due to hemolytic anemia); Coombs’ test negative.

Micro Pathology  Skin biopsy of vascular skin lesions reveals endothelial proliferation and histiocytic hyperplasia; electron microscopy of verrucous tissue shows *Bartonella bacilliformis* in interstitial tissue.

Treatment  Chloramphenicol, penicillin, erythromycin, norfloxacin, and tetracycline are effective; rifampicin is indicated for treatment of verrucous forms.

Discussion  Bartonellosis is a sandfly-borne bacterial disease occurring only on the western coast of South America at high altitudes; the causative agent is a motile, pleomorphic bacillus, *Bartonella bacilliformis*. Two stages of the disease are recognized: an initial febrile stage associated with a hemolytic anemia (Orova fever) and a later cutaneous stage characterized by hemangiomatous nodules ( verruga peruana ).
A 32-year-old male is referred to a tertiary care center with chronic pneumonia and warty lesions on his left upper limb.

The patient is from the southeastern United States. His skin lesions are nonpruritic and painless. He also complains of malaise, weight loss, night sweats, chest pain, breathlessness, and hoarseness.

VS: fever; tachycardia; mild tachypnea. PE: bilateral rales and rhonchi; raised, verrucous, and crusted lesions with serpiginous border located on left upper extremity; small abscesses demonstrable when superficial crust was removed.

Sputum and pus from cutaneous lesions demonstrate spherical cells (8 to 15 mm in diameter) that have a thick-walled, refractile double contour and show unipolar (broad-based) budding; culture of pus and sputum on Sabouraud's agar yields growth of Blastomyces; no evidence of acid-fast bacilli found either on staining or on culture; Gomori's methenamine silver staining of lung tissue does not reveal Pneumocystis.

CXR: bilateral alveolar consolidations with air bronchograms.

Epithelioid macrophages and giant cells surrounding a suppurative center; skin lesions show pseudoepitheliomatous hyperplasia very similar to squamous cell carcinoma.

Itraconazole is treatment of choice in most patients; amphotericin B, fluconazole, and ketoconazole are alternative drugs.

Blastomycosis is a systemic mycotic infection of humans and dogs that is characterized by suppuration and granulomatous lesions and is caused by the dimorphic fungus Blastomyces dermatitidis; the disease is endemic in the southeastern and south-central portions of the United States, and several pockets of infection extend north along the Mississippi and Ohio rivers into central Canada. Clinical disease most commonly involves the lungs (acquired by spore inhalation) and then, by hematogenous dissemination, the skin, the skeletal system, and the male genitourinary tract. Infection cannot be passed from person to person.

Atlas Link M-M1-065
ID/CC A 25-year-old male presents with sudden-onset **double vision** (**diplopia**), **dry mouth**, **weakness**, **dysarthria**, and **dysphagia**.

HPI He has no previous history of episodic weakness or of dog or tick bites (vs. myasthenia gravis, rabies, or Lyme disease). Last night, he ate some **home-canned food**.

PE VS: no fever. PE: patient alert; ptosis; bilateral **third and tenth cranial nerve palsy**; symmetric **flaccid paralysis** of all four limbs; deep tendon reflexes reduced; no sensory loss seen; decreased bowel sounds.

Labs Botulinum toxin detected in patient’s serum and canned-food sample with specific antiserum.

Treatment Antitoxin; close monitoring of respiratory status; intubation for respiratory failure.

Discussion The disease is characterized by gradual return of muscle strength in most cases. Botulinum toxin is a zinc metalloprotease that cleaves specific components of synaptic vesicle docking and fusion complexes, thus **inhibiting the release of acetylcholine at the neuromuscular junction**. The disease in adults is due to **ingestion of the toxin** rather than to bacterial infection. Botulism is also seen in infants secondary to the ingestion of *Clostridium botulinum* spores in **honey**.
ID/CC A 28-year-old white male visits his family doctor complaining of acute pain in both hip joints together with weakness, backache, myalgias, arthralgias, and undulating fever of 2 months' duration; this morning he woke up with pain in his right testicle.

HPI For the past 3 years he has worked at the largest dairy farm in his state. He enjoys drinking “crude” milk.

PE VS: fever. PE: pallor; marked pain on palpation of sacroiliac joints; mild splenomegaly; generalized lymphadenopathy.

Labs CBC: relative lymphocytosis with normal WBC count. Positive agglutination titer (> 1:160); rising serologic titer over time; small gram-negative rod Brucella abortus on blood culture.

Imaging XR, hips: joint effusion and soft tissue swelling without destruction. MR, spine: evidence of spondylitis.

Gross Pathology Lymphadenopathy and splenomegaly; hepatomegaly rare.

Micro Pathology Granulomatous foci in spleen, liver, and lymph nodes, with proliferation of macrophages; epithelioid and giant cells may be seen.

Treatment Combination therapy with doxycycline or TMP-SMX and rifampin or streptomycin.

Discussion Also called Malta fever, a microbial disease of animals, brucellosis is caused by several species of Brucella, a gram-negative, aerobic coccobacillus. It is transmitted to humans through the drinking of contaminated milk or through direct contact with products or tissues from animals such as goats, sheep, camels, cows, hogs, and dogs. The clinical picture is often vague; thus, a high index of suspicion may be necessary for diagnosis.
ID/CC  A 26-year-old female presents to the ER with intense, acute-onset left lower quadrant **crampy abdominal pain**, foul-smelling stools with streaks of blood, urgency, tenesmus, and fever.

HPI For the past 2 days, the patient has also had headaches and myalgias. She frequently drinks **unpasteurized** ("raw") **milk** that she buys at a health-food store.

PE VS: fever (39°C); tachycardia; normal RR and BP. PE: no dehydration; diffuse abdominal tenderness more marked in left lower quadrant.

Labs Stool smear shows leukocytes (due to invasive tissue damage in the colon) and **gram-negative, curved bacilli**, often in pairs, in "gull-wing"-shaped pattern; dark-field exam shows motility; culture in microaerophilic, 42°C conditions on special agar yields **Campylobacter jejuni**, indicated by oxidase and catalase positivity.

**Gross Pathology** Friable colonic mucosa.

**Micro Pathology** Nonspecific inflammatory reaction consisting of neutrophils, lymphocytes and plasma cells with hyperemia, edema and damage to epithelium, glandular degeneration, ulcerations, and crypt abscesses caused by colonic tissue invasion of the organism.

**Treatment** Self-limiting disease. Severe cases (i.e., high fever, severe diarrhea) can be treated with **fluoroquinolones**.

**Discussion** One of the primary causes of "traveler’s diarrhea." Sources of infection include **undercooked food** and contact with **infected animals** and their excreta. Prevent by improving public sanitation, pasteurizing milk, and proper cooking.
A 49-year-old morbidly obese, diabetic woman presents with pruritus in the skin folds beneath her breasts.

She admits to having this problem chronically, especially in the warm summer months, when she perspires more heavily.

Superficially denuded, beefy-red areas beneath breasts with satellite vesicopustules and whitish curd-like concretions on surface.

Clusters of budding cells with short hyphae seen under high-power lens after skin scales have been put in 10% KOH; Candida albicans isolated in Sabouraud’s medium.

Rash has whitish-creamy pseudomembrane that covers an erythematous surface.

Yeast invades superficial layers of epithelium.

Keep affected areas dry; clotrimazole or other antifungal agents locally.

Other superficial areas of infection include the oral mucosa (thrush), vaginal mucosa (vaginitis), and esophagus (GI candidiasis). Systemic invasive candidiasis may be seen with immunosuppression, in patients receiving chronic broad-spectrum antibiotics, in AIDS patients, or in those receiving hyperalimentation.
ID/CC  A 25-year-old female presents with painful lumps in her right axilla and neck together with low-grade fever.

HPI  Three weeks ago she was scratched on her right forearm by her pet cat; an erythematous pustule initially developed at the site but resolved spontaneously within 10 days.

PE  VS: fever. PE: tender right axillary and cervical lymphadenopathy.

Labs  Lymph node biopsy diagnostic; serologic indirect immunofluorescent antibody test for Bartonella henselae is positive.

Micro Pathology  Hematoxylin and eosin staining reveals granulomatous pathology with stellate necrosis and surrounding palisades of histiocytes; Warthin–Starry silver stain reveals clumps of pleomorphic, strongly argyrophilic bacilli.

Treatment  Symptomatic; fluctuant node may need aspiration; azithromycin given to immunocompromised patients.

Discussion  Bartonella henselae is the agent that causes cat-scratch disease. Lymphadenopathy can persist for months and can sometimes be mistaken for a malignancy. Individuals who are immunocompromised may present with seizures, coma, and meningitis.
ID/CC  An 8-year-old white female enters the emergency room complaining of headache, malaise, and bilateral swelling of the right eye.

HPI  She recently returned from a year-long stay in Brazil, where her father works as a logger in the Amazon forest. Over the past week she had a high fever, which was treated at home as malaria.

PE  VS: fever (39°C); tachycardia. PE: right eyelid swollen shut (Romaña's sign); markedly hyperemic conjunctiva; ipsilateral retroauricular and cervical lymph nodes; hepatosplenomegaly.

Labs  PBS: trypanosomes on thick blood smear. ECG: right bundle-branch block; ventricular extrasystoles.

Gross Pathology  Encapsulated, nodular area (chagoma) or Romaña's sign may be seen at point of entry, commonly the face.

Micro Pathology  Intense neutrophilic infiltrate with abundant macrophages at site of entry; myocardial necrosis with mononuclear cell infiltration; pseudocysts in infected tissues contain parasites that multiply within cells; denervation of myenteric gut plexus.

Treatment  Nifurtimox for acute disease.

Discussion  Chagas' disease is a parasitic disease that is restricted to the Americas (endemic in South and Central America) and is produced by Trypanosoma cruzi, a thin, undulating flagellated protozoan; it is transmitted by contamination of a reduviid bug bite with injection of its feces. Also known as American trypanosomiasis. Long-standing cases show myocardial involvement with dilated cardiomyopathy, life-threatening conduction defects, and apical aneurysm formation and may also show megaesophagus or megacolon.

Atlas Link  UCMID M-M1-071
ID/CC  A 35-year-old male complains of cough productive of mucopurulent sputum and breathlessness.

HPI  Before the onset of these symptoms, he had a sore throat with hoarseness. He has no history of hemoptysis, sharp chest pain, or high-grade fever.

PE  Crepitations heard over left lung base.

Labs  CBC: normal leukocyte count. Sputum exam revealed no bacterial organism; microimmunofluorescence detected species-specific antibodies directed against Chlamydia outer-membrane proteins; cultivation of C. pneumoniae demonstrated on HEP-2 and HL cell lines.

Imaging  CXR: left lower lobe subsegmental infiltrate with interstitial pattern.

Treatment  Doxycycline is the drug of choice; erythromycin and fluoroquinolones may also be used.

Discussion  The peak incidence of chlamydia pneumonia is in young adults. The mode of transmission would appear to be from person to person.
ID/CC An 8-year-old male who recently emigrated from India presents with bilateral eye irritation and photophobia.

HPI He reports recurrent episodes of similar eye irritation and redness in the past.

PE Conjunctival congestion; multiple (> 5) follicles, each at least 0.5 mm in diameter, seen in upper tarsal conjunctiva; inflammatory thickening of tarsal conjunctiva; new vessels (PANUS) seen in cornea at superior limbus; punctate keratitis.

Labs Diagnosis confirmed by demonstration of characteristic cytoplasmic inclusion bodies (HALBERSTEDER-PROWZEK BODIES) in Giemsa staining of conjunctival scrapings.

Micro Pathology Chlamydia trachomatis is typically seen in conjunctival scrapings in colony form in the epithelial cells as H-P inclusion bodies. Histologically there is lymphocytic infiltration involving the whole adenoid layer of parts of the conjunctiva; special aggregations of lymphocytes form follicles that tend to show necrosis and certain large multinucleated cells (LEBER’S CELLS).

Treatment Topical tetracycline with systemic tetracycline/doxycycline/erythromycin/azithromycin; prophylaxis of family contacts with topical tetracycline.

Discussion Chlamydia trachomatis causes a variety of ocular diseases, including neonatal inclusion conjunctivitis, sporadic inclusion conjunctivitis in adults, and sporadic as well as endemic trachoma; trachoma is endemic in North Africa, in the Middle East, and among the Native American population of the southwestern United States. In endemic areas, trachoma is transmitted from eye to hand to eye, especially among young children in regions where standards of cleanliness are poor. Sporadic trachoma infection in nonendemic areas as well as sporadic inclusion conjunctivitis in adults results from transmission of the agent from the genital tract to the eye. Trachoma is a major cause of blindness in endemic areas.

Atlas Link UCBID M-M1-073
ID/CC A 30-year-old man has sudden severe, profuse (several liters per day) watery diarrhea, protracted vomiting, and abdominal pain.

HPI He has just returned from a trip to rural India.

PE Severe dehydration; low urine output; generalized mild abdominal tenderness with no signs of peritoneal irritation; stools have characteristic “rice-water” appearance; (gray, slightly cloudy fluid with flecks of mucus), with no blood.

Labs Stool culture reveals gram-negative rods with “darting motility”; O1 antigen detected; Vibrio cholerae isolated on culture media; serum chloride levels decreased; serum sodium levels increased.

Treatment Vigorous rehydration therapy with oral and/or IV fluids; tetracycline, ciprofloxacin, or doxycycline.

Discussion A heat-labile exotoxin produced by Vibrio cholerae that acts by permanently stimulating Gs protein via ADP ribosylation, resulting in activation of intracellular adenylate cyclase, which in turn increases cAMP levels and produces secretory diarrhea.
ID/CC  A newborn baby is referred to the pediatrician for further evaluation of an unusually small head, low birth weight, and an extensive erythematous rash.

HPI  Intrauterine growth retardation was prenatally diagnosed on ultrasound. The child’s mother had a flulike episode during the first trimester of her pregnancy.

PE  Small for gestational age; generalized hypotonia with sluggish neonatal reflexes; extensive “pinpoint” petechial skin rash (mulberry muffin rash); microcephaly; chorioretinitis, mild icterus; hepatosplenomegaly; sensorineural hearing loss in right ear.

Labs  CBC/PBS: mild thrombocytopenia; atypical lymphocytosis. Moderately elevated direct serum bilirubin and transaminases. UA: cells in urine found to have large intranuclear inclusions (owl’s eye inclusions); cytomegalovirus isolated on tissue culture.

Imaging  XR/CT, head: periventricular calcifications; microcephaly.

Treatment  Ganciclovir (only for immunocompromised patients).

Discussion  A congenital herpesvirus infection involving the CNS with eye and ear damage, congenital cytomegalovirus is a common cause of mental retardation.
A 13-year-old white female visits her pediatrician complaining of fever, severe dyspnea, and a dry cough.

She was recently diagnosed with acute lymphocytic leukemia, for which she received a bone marrow transplant. She is currently on immunosuppressive therapy.

VS: fever; tachypnea. PE: pallor; crepitant rales over both lung fields; mild cyanosis; no hepatosplenomegaly.

CBC/PBS: anemia; leukopenia. ABGs: hypoxemia. No organism in induced sputum stained with Gram, Giemsa, ZN, and methenamine silver.

CXR: diffuse, bilateral interstitial infiltrates.

Interstitial pneumonitis; hepatitis.

Characteristic intranuclear inclusions with surrounding halo (owl's- or bull's-eye cells) on transbronchial lung biopsy.

Ganciclovir (CMV is resistant to acyclovir).

An enveloped, double-stranded DNA virus belonging to the herpesvirus group; the most common cause of pneumonia and death in bone marrow transplant patients. It is also common in AIDS patients.

Atlas Link [UCM1 M-M1-076]
ID/CC A 30-year-old homosexual white male presents to his family physician with a rapidly progressive diminution of vision.

HPI He is known to be HIV positive and periodically comes in for checkups.

PE Cotton-wool exudates, necrotizing retinitis, and perivascular hemorrhages on funduscopic exam.

Treatment Ganciclovir; foscarnet (CMV is resistant to acyclovir).

Discussion CMV retinitis is an important treatable cause of blindness that occurs in 20% of AIDS patients; 50% to 60% of patients develop retinal detachment within 1 year. Toxoplasmosis and progressive multifocal leukoencephalopathy (PML) are other important causes of blindness in AIDS patients.
ID/CC A 19-year-old migrant worker from the southwestern United States is brought to the family doctor complaining of cough, pleuritic chest pain, fever, and malaise.

HPI He also complains of a backache and headache along with an erythematous skin rash (due to hypersensitivity reaction) in his lower limbs.

PE VS: fever; tachypnea. PE: central trachea; coarse, crepitant rales over both lung bases; tender, erythematous nodules over shins (erythema nodosum); periarticular swelling of knees and ankles.

Labs Positive skin test with coccidioidin; dimorphic fungi (hyphae in soil; spherules in body tissue); Coccidioides immitis on silver stain and sputum culture; positive latex agglutination test. CBC/PBS: eosinophilia.

Imaging CXR: nodular infiltrates and thin-walled cavities in both lower lungs.

Gross Pathology Caseating granulomas; often subpleural and in lower lobes; necrosis and cavitation may also be present.

Micro Pathology Silver-stained tissue sections show spherules filled with endospores.

Treatment Amphotericin B or itraconazole.

Discussion Endemic in the southwestern United States, coccidioidomycosis is produced by C. immitis and is transmitted by inhalation of arthrospores. Systemic dissemination is frequent in blacks as well as in immunosuppressed and pregnant patients. Meningitis or granulomatous lung disease may result, which may lead to death.
**ID/CC** A 28-year-old male who lives in the *northwestern United States* complains of a high-grade *fever with rigors*, generalized aches, myalgias, headache, and backache.

**HPI** Four days ago he returned from a hiking trip during which he was *bitten by a tick*; he took amoxicillin as prophylaxis against Lyme disease.

**PE** VS: fever.

**Labs** CBC: leukopenia; relative lymphocytosis. Viral antigen detected in RBCs by immunofluorescence; *Colorado tick virus cultured* in suckling mice by intracerebral inoculation of blood clot; indirect fluorescent Ab test positive.

**Treatment** Symptomatic.

**Discussion** Colorado tick fever virus is an 80-nm double-shelled *reovirus* that is covered with capsomeres; its icosahedral core contains 12 *segments of dsRNA*. The disease is a zoonosis that is transmitted by a wood tick, *Dermacentor andersoni*. It occurs primarily in the Rocky Mountain region, primarily affecting hikers. Since no specific therapy exists, prevention is key (wear clothing that covers the body).
A 2-year-old male is brought to the ER by his parents with sore throat, inspiratory stridor, and a barking cough of 1 day’s duration.

The patient has no significant past medical history.

VS: fever (38.6°C); tachypnea. PE: respiratory distress; nasopharyngeal discharge; diffuse rhonchi and wheezes; examination of extremities reveals some cyanosis.

Throat and nasal swabs isolate parainfluenza virus; serodiagnosis and hemagglutinin inhibition tests reveal type 1 (most common cause).

CXR: air trapping. XR, neck: subglottic narrowing.

Inflammation and edema of larynx, trachea, and bronchi.

Most cases respond to supportive therapy such as humidified air, removal of secretions, and bed rest. Severe cases may require humidified oxygen, racemic epinephrine, or high-dose corticosteroids.

Differentiate croup from Haemophilus influenzae type B and influenza A virus. Modes of transmission include respiratory droplets and person-to-person contact; tends to peak in the fall and winter. Most cases of croup are due to parainfluenza virus type 1; type 3 is a prominent cause of bronchiolitis in babies.
ID/CC  A 30-year-old man with AIDS presents with chronic, recurrent profuse, nonbloody, watery diarrhea.

HPI  The diarrhea has recurred over the past 2 months with intermittent cramping, and previous treatments have not been effective.

PE  VS: no fever. PS: moderate dehydration; thin; generalized lymphadenopathy.

Labs  Acid-fast staining demonstrates oocysts of Cryptosporidium in fresh stool.

Gross Pathology  Intestinal mucosa appears normal.

Micro Pathology  Blunting of intestinal villi; mixed inflammatory cell infiltrates with eosinophils in lamina propria; organisms visible on brush borders.

Treatment  No treatment found effective; supportive management with maintenance of fluids and nutrition.

Discussion  Cryptosporidium parvum infection presents as acute diarrhea in malnourished children and as severe diarrhea in immunocompromised patients (part of HIV wasting syndrome); the disease is mild and self-limiting in immune-competent patients. The disease is acquired through the ingestion of oocysts (fecal-oral transmission) that may be killed by chlorination.

Atlas Link  [Link to image] M-M1-081
ID/CC  A 5-year-old white male presents with malaise, anorexia, low-grade fever, sore throat of 3 days’ duration, and dyspnea on exertion.

HPI  The child was raised abroad. His immunization status cannot be determined.

PE  VS: fever; tachycardia with occasional dropped beats. PE: cervical lymphadenopathy (bull’s-neck appearance); smooth, whitish-gray, adherent membrane over tonsils and pharynx; no hepatosplenomegaly; diminished intensity of S1.

Labs  Metachromatic granules in bacilli arranged in “Chinese character” pattern on Albert stain of throat culture; Corynebacterium diphtheriae confirmed by growth observed on Löeffler’s blood agar; erythema and necrosis following intradermal injection of C. diphtheriae toxin (positive Schick’s test); immunodiffusion studies (Elek’s) confirm toxigenic strains of C. diphtheriae. ECG: ST-segment elevation; second-degree heart block.

Imaging  Echo: evidence of myocarditis.

Gross Pathology  Pharyngeal membranes not restricted to anatomic landmarks; pale and enlarged heart.

Micro Pathology  Polymorphonuclear exudate with bacteria; precipitated fibrin and cell debris forming a pseudomembrane; marked hyperemia, edema, and necrosis of upper respiratory tract mucosa; exotoxin-induced myofibrillar hyaline degeneration; lysis of myelin sheath.

Treatment  Begin treatment on presumptive diagnosis; specific antitoxin and penicillin or erythromycin; respiratory and cardiac support; confirm eradication by repeating throat culture.

Discussion  A bacterial infection of the throat, diphtheria is preventable by vaccine and is caused by toxigenic Corynebacterium diphtheriae, a club-shaped, gram-positive aerobic bacillus. Diphtheria toxin is produced by β-prophage-infected corynebacteria; it blocks EF-2 via ADP ribosylation and hence ribosomal function in protein synthesis. The toxin enters the bloodstream, causing fever, myocarditis (within the first 2 weeks), and polyneuritis (many weeks later).

Atlas Links  UCM1 M-M1-082 UCM2 MC-324

82 DIPHTHERIA
A 56-year-old male professor of veterinary medicine from New Zealand experiences sudden high fever with chills, jaundice, and right upper quadrant pain while attending a conference in the United States.

His past history is unremarkable. He has been healthy and has been physically active working in the field with sheep and breeding dogs.

VS: fever; hypotension (BP 90/50). PE: hepatomegaly; jaundiced sclera; on palpation of epigastrum and right hypochondrium, abdomen is tender with no rebound tenderness.

CBC: leukocytosis with neutrophilia; slight eosinophilia. Strongly positive immunoblot test for antibodies to echinococcal antigens; elevated direct bilirubin and alkaline phosphatase.

CT/US, abdomen: multiple large septated liver cysts impinging on bile ducts, producing biliary dilatation (due to obstruction).

Liver is most common site of invasion, but cysts may also form in lungs, kidney, bone, and brain; each cyst contains millions of scoleces and consists of two layers: an inner germinal layer and an outer laminated layer; usually surrounded by fibrotic reaction.

Giant cell reaction surrounding cyst with eosinophilic infiltration.

Surgically remove cysts if possible; albendazole may be effective.

Echinococcosis is a zoonosis produced by Echinococcus granulosus. It is acquired through the ingestion of food or drink contaminated with the feces of dogs or other carnivores that have eaten contaminated meat; humans are the intermediate host of parasitic larvae. Accidental spilling of cyst fluid, either spontaneously or during surgery, may result in secondary seeding or anaphylaxis and even death. Also known as hydatid disease.
ID/CC  A 28-year-old male who is a resident of the southeastern United States presents with a high fever with chills, headache, and myalgias.

HPI  He remembers having been bitten by a tick a week before developing his symptoms; however, he reports no skin rash.

PE  VS: fever. PE: no skin rash noted.

Labs  CBC: leukopenia and mild thrombocytopenia. Characteristic intraleukocytic inclusion bodies and serologic response to Ehrlichia antigens demonstrated; E. chaffeensis cultured from blood and detected by PCR.

Treatment  Doxycycline.

Discussion  Ehrlichicac are gram-negative, obligately intracellular bacteria. The two types of Ehrlichia species that affect humans are E. chaffeensis (which attacks macrophages and monocytes) and an E. equi-like organism (which attacks granulocytes). Preventive measures include wearing clothing that covers the body and using insect repellants.
A 30-year-old male from Texas presents with fever and a skin rash that began about 2 weeks ago.

The onset was gradual, with prodromal symptoms of headache, malaise, backache, and chills. These symptoms were followed by shaking chills, fever, and a more severe headache accompanied by nausea and vomiting. A remittent pattern of fever accompanied by tachycardia continued for 10 to 12 days, with the rash appearing around the fifth day of fever. The patient worked at a rat-infested food-storage depot this summer.

VS: fever. PE: discrete, irregular pink maculopapular rash seen in axillae and on trunk, thighs, and upper arms; face, palms, and soles only sparsely involved; mild splenomegaly noted.

The Weil–Felix agglutination reaction for Proteus strain OX-19 was positive; complement-fixing antibodies to the typhus group antigen were demonstrated; endemic typhus (due to Rickettsia typhi) was confirmed serologically by using specific washed rickettsial antigens in IFA tests.

Antibiotic treatment with doxycycline (chloramphenicol is used as an alternative).

Murine typhus is a natural infection of rats and mice by Rickettsia typhi; spread of infection to humans by the rat flea is incidental and occurs when feces from infected fleas are scratched into the lesion. Cases can occur year-round; however, most occur during the summer months, primarily in southern Texas and California.
ID/CC  A 28-year-old Guatemalan male is brought to the hospital complaining of severe headache, photophobia, and fever over the past 2 weeks.

HPI  As a political dissident, he spent 4 months in a refugee camp in southern Mexico before entering the United States.

PE  VS: fever (40°C). PE: papilledema and delirium; bilateral swelling of parotid glands 1 week later; toxic facies; maculopapular rash on trunk and extremities; face, palms, and soles spared; mild splenomegaly.

Labs  Positive Weil-Felix reaction to OX-19 strains of Proteus; rise in complement fixation titer for Rickettsia prowazekii; specific antibodies. UA: proteinuria; microscopic hematuria.

Gross Pathology  Myocarditis and pneumonia may be present; cerebral edema; maculopapular rash.

Micro Pathology  Zenker’s degeneration of striated muscle; thrombosis and endothelial proliferation of capillaries with abundant rickettsiae and perivascular cuffing; accumulation of lymphocytes; microglia and macrophages (typhus nodules) in brain.

Treatment  Doxycycline; chloramphenicol.

Discussion  Epidemic typhus is a febrile illness caused by Rickettsia prowazekii, a gram-negative, nonmotile, obligate intracellular parasite; it is transmitted via body lice and is associated with war, famine, and crowded living conditions. The rash should be differentiated from Rocky Mountain spotted fever, which starts peripherally on the wrists and ankles and also includes the palms and soles.

Atlas Link  UCMZ MC-169
ID/CC A 4-year-old male presents with fever, hoarseness, and respiratory distress because of partial airway obstruction.

HPI The child is also unable to speak clearly and has pain while swallowing (odynophagia).

PE VS: fever; tachypnea. PE: patient is leaning forward with neck hyperextended and chin protruding; drooling; marked suprasternal and infrasternal retraction of chest; inspiratory stridor on auscultation.

Labs Culture of throat swab (no role in management of acute disease) reveals penicillinase-resistant Hemophilus influenzae; blood cultures also positive.

Imaging XR, neck: marked edema of epiglottis and aryepiglottic folds (“thumbs-up” sign).

Gross Pathology Epiglottis is cherry-red, swollen, and “angry-looking.” Rapid cellulitis of epiglottis and surrounding tissue leads to progressive blockage of airway.

Treatment Preservation of airway; IV cefuroxime.

Discussion The principal cause of acute epiglottitis in children and adults is H. influenzae type b; other pathogens include H. parainfluenzae and group A streptococcus. Characterized by rapid onset.

Atlas Link UPML PG-M1-087
ID/CC  A 30-year-old soldier who had been admitted for a gunshot wound in the right thigh presents with severe pain and swelling at the site of his injury.

HPI  The patient’s right lower limb had become discolored, and several bullae had appeared on the skin. He has passed very little urine over the past day, and the urine he has passed has been dark (“cola-colored”).

PE  VS: low-grade fever; marked tachycardia. PE: diaphoresis; skin of right thigh discolored (bronze to purple red); site of injury exquisitely tender and tense and oozing a thin, dark, and foul-smelling fluid; crepitus while palpating thigh.

Labs  CBC: low hematocrit. Gram stain of exudate and necrotic material at wound site reveals presence of large gram-positive rods; anaerobic culture of exudate and blood yields Clostridium perfringens type A; culture isolate demonstrates positive Nagler reaction (due to presence of alpha toxin lecithinase); further labs confirm presence of intravascular hemolysis, myo- and hemoglobinuria, and acute tubular necrosis.

Imaging  XR, right thigh: presence of gas in soft tissues.

Gross Pathology  Overlying skin purple-bronze, markedly edematous with vesiculobulbous changes with little suppurative reaction.

Micro Pathology  Coagulative necrosis, edema, gas formation, and many large gram-positive bacilli found in affected muscle tissue; relatively sparse infiltration of PMNs noted in the bordering muscle tissue.

Treatment  Surgical debridement; antibiotics (penicillin, clindamycin, tetracycline, metronidazole); hyperbaric oxygen therapy and polyvalent antitoxin; supportive management of associated multiorgan failure.

Discussion  A rapidly progressive myonecrosis caused by Clostridium perfringens type A, traumatic gas gangrene develops in a wound with low oxygen tension (embedded foreign bodies containing calcium or silicates cause lowering of oxygen tension, leading to germination of the spores). The most important toxin is the alpha toxin lecithinase, which produces hemolysis and myonecrosis.

Atlas Link  UCM  PG-M1-088

88 GAS GANGRENE—TRAUMATIC
ID/CC  A 4-year-old female is brought to the pediatrician because of lack of appetite; nausea and vomiting; chronic, foul-smelling diarrhea without blood or mucus; and a bloated sensation.

HPI She has been in several day-care centers over the past 3 years.

PE Low weight and height for age; mild epigastric tenderness.

Labs Binucleate, pear-shaped, flagellated trophozoites (GIARDIA LAMBLIA) on freshly passed stool; cysts found on stool exam.

Treatment Metronidazole.

Discussion The most common protozoal infection in children in the United States, giardiasis is transmitted mainly through contaminated food or water and causes malabsorption.

Atlas Links UCM1 M-M1-089A, M-M1-089B
A 3-day-old female neonate presents with a thick eye discharge. The mother admits to having multiple sexual partners and complains of a vaginal discharge. She did not receive adequate antenatal care.

Exam of both eyes reveals a thick purulent discharge and marked conjunctival congestion and edema; conjunctival chemosis is so marked that cornea is seen at bottom of a crater-like pit; corneal ulceration noted.

Conjunctival swabs on Gram staining reveal presence of gram-negative diplococci both intra- and extracellularly in addition to many PMNs; conjunctival swab and maternal cervical culture yield Neisseria gonorrhoeae.

Aqueous penicillin G or ceftriaxone for a total of 7 days. Also treat mother and her sexual contacts. Educate the mother regarding the importance of safe sex.

Caused by Neisseria gonorrhoeae, gonococcal ophthalmia neonatorum is contracted from a mother with gonorrhea as the fetus passes down the birth canal; infection does not occur in utero. Corneal inflammation is the major clinical sign that may produce complications such as corneal opacities, perforation, anterior synechiae, anterior staphyloma, and panophthalmitis. It is now common practice to prevent this disease by treating the eyes of the newborn with an antibacterial compound such as erythromycin ointment or 1% silver nitrate; however, home childbirth bypasses this prophylactic procedure, and thus some cases are still occurring in the United States.
ID/CC  A 19-year-old white male presents with burning urination; profuse, greenish-yellow, purulent urethral discharge; staining of his underwear; and urethral pain.

HPI  Four days ago, he had unprotected sexual contact with a prostitute.

PE  Mucopurulent and slightly blood-tinged urethral discharge; normal testes and epididymis; no urinary retention.

Labs  Smear of urethral discharge reveals intracellular gram-negative diplococci in WBCs; gonococcal infection confirmed by inoculation into Thayer-Martin medium.

Gross Pathology  Abundant, purulent urethral exudate.

Treatment  Ceftriaxone plus doxycycline or erythromycin for possible coinfection with Chlamydia.

Discussion  A common STD caused by Neisseria gonorrhoeae, gonorrhea may involve the throat, anus, rectum, epididymis, cervix, fallopian tubes, prostate, and joints; conjunctivitis is also found in neonates. Neonatal conjunctivitis may be prevented through the instillation of silver nitrate or erythromycin eye drops at birth.

Atlas Links  UCV1 M-M1-091  UCV2 IM2-018
ID/CC A 28-year-old male immigrant presents with inguinal swelling and a painless penile ulcer.

HPI He admits to unprotected intercourse with multiple sexual partners, many of whom were prostitutes. He first noticed a papule on his penis several weeks ago.

PE Soft, painless, raised, raw, beef-colored, smooth granulating ulcer noted on distal penis; multiple subcutaneous swellings (pseudorubores) noted in inguinal region, some of which have ulcerated.

Labs Giemsa-stained smear from penile and inguinal regions demonstrate characteristic "closed safety pin" appearance of encapsulated organisms within a large histiocyte (Donovan bodies).

Micro Pathology Characteristic histologic picture of donovanosis comprises some degree of epithelial hyperplasia at margins of lesions; dense plasma cell infiltrate scatters histiocyte-containing Donovan bodies.

Treatment Treat with doxycycline or double-strength TMP-SMX.

Discussion Granuloma inguinale, a slowly progressive, ulcerative, granulomatous STD involving the genitalia, is caused by the gram-negative bacillus Calymmatobacterium granulomatis (formerly Donovania granulomatis); it is seen in Giemsa-stained sections as a dark-staining, encapsulated, intracellular rod-shaped inclusion in macrophages, the so-called Donovan body. The disease is endemic in tropical areas such as New Guinea, southern India, and southern Africa.
A 60-year-old male presents with cough productive of mucopurulent sputum together with mild fever and worsening breathlessness.

He is a chronic smoker who has been diagnosed with COPD.

VS: fever. PE: in moderate respiratory distress; emphysematous chest with obliterated cardiac and liver dullness; wheezing and crackles heard over both lung fields.

Haemophilus influenzae organisms seen as small, pleomorphic gram-negative bacilli on Gram stain of sputum; nontypable H. influenzae isolated on sputum culture (to grow in culture, H. influenzae requires both factor X–hematin and factor V–nicotinamide nucleoside present in erythrocytes).

Amoxicillin/ampicillin therapy; TMP-SMX, azithromycin, and clarithromycin are also excellent drugs for the treatment of clinically mild to moderate H. influenzae infections of the upper respiratory tract.

Infections caused by nontypable, or unencapsulated, Haemophilus influenzae strains have been increasingly recognized in pediatric and adult populations. Nontypable H. influenzae strains are frequent respiratory tract colonizers in patients with COPD and commonly exacerbate chronic bronchitis in these patients; nontypable strains are also the most common cause of acute otitis media in children.
A 25-year-old male presented with sudden-onset breathlessness, cough, cyanosis, and high-grade fever.

The patient failed to improve on 100% oxygen, became hypotensive, and died of type 2 respiratory failure a few hours after admission. He had been in perfect health and had been hiking in several rodent-infested areas before falling ill.

On admission he was found to have fever, tachycardia, cyanosis, hypotension, and rales on auscultation over both lung fields; no meningeal signs or localizing CNS signs could be demonstrated.

ABGs: respiratory acidosis with hypoxia and hypercapnia. CBC: leukocytosis; hemoconcentration; thrombocytopenia; atypical lymphocytosis. Increased LDH and ALT levels; prolonged PT index; sputum exam and blood culture did not yield any organism; IgM antibody to hantavirus and immunohistochemical stains for hantavirus antigen in tissues confirmed infection with the virus.

CXR: noncardiogenic pulmonary edema (bat-wing edema pattern).

Histopathologic exam of lung tissues was suggestive of acute respiratory distress syndrome (adult hyaline membrane disease).

Patient died despite intensive ventilatory support (Sin Nombre virus most frequently causes hantavirus pulmonary syndrome in the United States).

A virus closely related to the Hantaan virus (which produces Korean hemorrhagic fever and hemorrhagic fever with renal syndrome) has been recovered from mice in various regions of the United States; rodents are the natural host for this group of viruses. Infected rodents shed the virus in saliva, urine, and feces for many weeks, and humans are believed to acquire the infection via exposure to rodent excrement or saliva, either by the aerosol route or by direct inoculation.
**ID/CC** A 35-year-old male who works as a U.N. health worker presents with a high-grade fever and massive hematemesis.

**HPI** He recently returned from Zaire, where he worked in a tick-infested forest.

**PE** VS: fever. PE: extensive ecchymosis.

**Labs** CBC: leukopenia; severe thrombocytopenia. LFTs: elevated AST. Crimean-Congo virus isolated.

**Treatment** Treatment involves a 10-day course of ribavirin; platelet transfusions; avoid salicylates; barrier nursing and containment of infected secretions, since airborne infection may occur in hospital environment.

**Discussion** The agent responsible for Crimean-Congo hemorrhagic fever is a bunyavirus; reservoirs include wild and domesticated sheep, cattle, goats, and hares. The disease is transmitted by a tick vector, usually an ixodid of the genus *Hyalomma*; endemic areas include the Middle East and western China. The disease targets individuals of all ages and affects males and females equally.
A 10-year-old male is brought to the ER in a state of shock accompanied by massive hematemesis.

The family had just returned from a vacation in Thailand. His parents say that he had a high-grade fever for 5 to 6 days, for which he was receiving presumptive treatment for malaria.

VS: hypotension; tachycardia. VS: cool, clammy extremities; petechial skin rash over extremities, axillae, trunk, and face; bleeding from venipuncture sites.

CBC: thrombocytopenia; hematocrit increased by > 20%. Abnormal clotting profile suggestive of disseminated intravascular coagulation (DIC); paired sera reveal significant rise in titer of hemagglutination inhibition antibodies against Dengue virus serotypes 1 and 2.

US: bilateral pleural effusion and ascites.

Symptomatic; manage shock with fluids and hemodynamic monitoring; fresh blood/platelet-rich plasma; avoid salicylates.

Dengue hemorrhagic fever is caused by a mosquito-borne (Aedes aegypti) flavivirus and is characterized by four distinct dengue serotypes (type 2 is considered the most dangerous). A. aegypti has a domestic habitat (stagnant water in flower pots, old jars, tin cans, and old tires) and bites during the day. Dengue fever has shown an increase in incidence in Southeast Asia, Central and South America, and the Caribbean. Since no specific therapy exists, prevent by avoiding contact with infected A. aegypti.
A 58-year-old man who was hitchhiking through central and southern Africa was admitted to a hospital in Zaire in a state of shock following massive hemorrhage from the GI tract (hematemesis and melena); he died within 6 hours of admission. Ten days later, a male doctor who had attended this patient and had attempted resuscitation became ill with a similar disease syndrome.

HPI At admission, he gave an 8-day history of progressive fever, severe headaches, myalgias, and watery diarrhea. He also reported an erythematous, measles-like skin rash that had begun to desquamate.

PE VS: fever. PE: splenomegaly; hepatomegaly.

Labs CBC: leukopenia; Pelger-Huët anomaly of neutrophils with atypical mononuclear cells; thrombocytopenia with abnormal platelet aggregation. Markedly elevated AST and ALT; blood was inoculated intraperitoneally into young guinea pigs and into various tissue culture cell lines, and Ebola virus was detected by indirect immunofluorescent staining techniques.

Gross Pathology At autopsy, lymph nodes, liver, and spleen found to be most conspicuously involved (replication of Ebola virus can occur in virtually all organs); stomach and intestines filled with blood; petechiae seen over bowel mucosa.

Micro Pathology Severe congestion and stasis of spleen; widespread necrosis of liver cells; electron microscopy of liver revealed pleomorphic virus particles appearing in contrast preparations as long, filamentous forms, U-shaped forms, and some circular forms resembling a doughnut.

Treatment Supportive care, since no specific treatment exists; a prior outbreak was brought under control by isolating patients and instituting strict barrier nursing.

Discussion A hemorrhagic, febrile infection of humans due to infection with the Ebola and Marburg viruses, both of which are filoviruses that are structurally indistinguishable but antigenically distinct. This disease is a zoonosis but the reservoir is unknown. Individuals can become infected through person-to-person or nosocomial contact.
ID/CC A 25-year-old male woodcutter who lives in South Korea is admitted to the ER in a state of shock and massive epistaxis.

HPI The patient had been complaining of fever, malaise, headache, myalgias, back pain, abdominal pain, nausea, and vomiting for the past week; he also complained of extremely reduced urine output. Careful history revealed that before he fell ill, he and his friend were cutting wood in the forest when they accidentally disturbed a rodent-infested area.

PE VS: hypotension. PE: epistaxis; facial flushing; petechiae and subconjunctival hemorrhages.

Labs Deranged RFTs suggestive of acute renal failure. CBC: thrombocytopenia. Serology and culture identify hantavirus, Hantaan serotype.

Treatment Supportive management in the form of dialysis (for renal failure); management of shock and hemorrhage; IV ribavirin (must start within first 4 days of manifestation of disease).

Discussion Korean hemorrhagic fever with renal syndrome is caused by the Hantaan serotype of hantavirus. Its reservoirs are various rodents that are found distributed over Europe and Asia; humans acquire the disease mainly by inhaling aerosols of rodent virus.
ID/CC A 7-year-old male complains of a high fever and a very sore throat.

HPI The pain is so severe that the child refuses to swallow. He is adequately immunized and achieved normal developmental milestones.

PE VS: fever. PE: characteristic grayish-white vesicular lesions, some of which have ulcerated, noted over soft palate and tonsils.

Labs Coxsackievirus A isolated from mucosal lesions.

Treatment Self-limiting condition.

Discussion In hand, foot, and mouth disease (HFMD), patients complain of fever, weakness, and decreased appetite along with similar lesions noted in the oral cavity, palms, soles, and buttocks. Herpangina may be caused by coxsackievirus A1–A10, A16, A22, and B1–B5. Outbreaks of HFMD are usually caused by coxsackievirus A16.
ID/CC A 25-year-old homosexual male visits a health clinic complaining of headache, low-grade fever, and a **painful skin rash in the perianal area**.

HPI He has no history of penile ulcerations and admits to **unprotected anal sex** with **multiple partners**.

PE Perianal **vesicular rash** in clusters on **erythematous base**; no penile ulceration; painful inguinal lymphadenopathy.

Labs **Multinucleated giant cells with intranuclear inclusions** surrounded by clear halo on Pap-stained section or Tzanck preparation of scrapings from base of vesicles.

Gross Pathology Clear liquid in vesicles; secondary bacterial infection may result; painful ulcerations when vesicles rupture.

Micro Pathology Inflammatory infiltrate with abundant lymphocytes.

Treatment **Acyclovir**.

Discussion An enveloped, double-stranded DNA virus transmitted by sexual contact, HSV 2 has a **tendency to recur** and can be **transmitted to the fetus through the birth canal**. Condom use appears to be one of the most effective means of preventing transmission.

Atlas Links **UCM1 M-M1-100**  **UCM2 IM2-019A, IM2-019B**
A 45-year-old HIV-positive male is seen by his family doctor following the appearance of a painful, burning skin rash on the left side of his chest that is accompanied by a headache and low-grade fever.

The patient had chickenpox as a child. He had been well until 1 year ago, when he was diagnosed with non-Hodgkin’s lymphoma, for which he is currently undergoing chemotherapy.

Vesicular rash on erythematous base; in dermatomal distribution (left T6–T8); exquisitely tender to touch.

Acantholytic cells on Tzanck smear from base of vesicles.

Intranuclear eosinophilic inclusions surrounded by clear halo (Cowdry A inclusions).

Acyclovir.

Shingles represents a reactivation of a latent infection with varicella-zoster virus; the rash typically follows the distribution of a nerve root. It is commonly seen in immunosuppressed patients and is also associated with trauma, ultraviolet radiation, hypothermia, and emotional stress. Postherpetic neuralgia is a common complication in the elderly.

IM2-020A, IM2-020B

HERPES ZOSTER (SHINGLES)