Otitis, Sinusitis, and Mastoiditis

Ear or Facial Pain Following a Common Cold

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4.1 Otitis – 39
4.1.1 Introduction – 39

4.2 Definitions – 39

4.3 Basic Concepts – 39
4.3.1 Otitis Externa – 39
4.3.2 Acute Otitis Media – 40
4.3.3 Risk Factors – 40
4.3.4 Microbiologic Causes of AOM – 40
4.3.5 Approach to the Diagnosis of AOM – 41
4.3.6 Differential Diagnosis of AOM – 41
4.3.7 Treatment of AOM – 41
4.3.8 Complications of AOM – 43
4.3.9 Follow-Up – 43

4.4 Sinusitis – 44
4.4.1 Introduction – 44

4.5 Definitions – 44
4.5.1 Sinusitis: Inflammation of the Paranasal Sinus Cavity Mucosa – 44

4.6 Basic Concepts – 45
4.6.1 Anatomy and Pathophysiology – 45
4.6.2 Risk Factors – 45
4.6.3 Microbiologic Causes of Acute Bacterial Rhinosinusitis – 45
4.6.4 Approach to the Diagnosis – 45
4.6.5 Differential Diagnosis of Bacterial Rhinosinusitis – 47
4.6.6 Treatment of Bacterial Rhinosinusitis – 47
4.6.7 Complications of Bacterial Sinusitis – 47

4.7 Mastoiditis – 48
4.7.1 Introduction – 48

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J. Domachowske (ed.), Introduction to Clinical Infectious Diseases,
https://doi.org/10.1007/978-3-319-91080-2_4
Learning Objectives
- Know the common clinical presentations for otitis, sinusitis, and mastoiditis.
- Identify common and uncommon microbiologic causes of otitis, sinusitis, and mastoiditis.
- Understand the distinguishing characteristics for acute, recurrent, and chronic clinical courses of each disease.
- List the important risk factors for developing severe infections of the paranasal sinuses.
- Outline the approach to diagnosis, including signs and symptoms that warrant laboratory or imaging evaluations.
- Describe the indications for medical and surgical treatment of otitis, sinusitis, and mastoiditis.

4.1 Otitis

4.1.1 Introduction

The term “otitis” encompasses pathology of both the middle and outer ear. It is generally divided into two categories – otitis media and otitis externa. Otitis media can present either as an acute infectious process of the middle ear (acute otitis media) or as a serous noninfectious process (otitis media with effusion). Otitis externa is an infectious inflammatory condition of the external auditory canal (EAC). Approaches to the diagnosis and treatment of acute otitis media have evolved over the last several decades as new immunizations, and more antibiotic choices have become available.

4.2 Definitions

Otitis media with effusion (OME) – A collection of serous fluid in the middle ear space without signs of acute inflammation. OME is not an infectious process.

Chronic otitis media with effusion (COME) – A collection of serous fluid in the middle ear space that persists for more than 3 months

Acute otitis media (AOM) – An acute infection of the middle ear with signs and symptoms of acute inflammation. An effusion is also present.

Recurrent AOM – Three or more episodes of AOM in a 6-month period or 4 or more episodes of AOM in a 12-month period

Otorrhea – The presence of a discharge from the ear

Chronic suppurative otitis media (CSOM) – The presence of a purulent middle ear effusion associated with otorrhea, secondary to chronic tympanic membrane perforation, for more than 6 weeks in the setting of antibiotic treatment

Otitis externa (OE) – An infection of the external auditory canal

4.3 Basic Concepts

4.3.1 Otitis Externa

Acute otitis externa develops following disruption of the epithelial cell layer of the EAC. Epithelial breakdown can be caused by excessive moisture that leads to maceration, trauma during insertion of a foreign body, occlusion by a device such as a hearing aid or earplug, or dermatologic conditions involving the EAC. Excessive moisture as a cause for breakdown or maceration of the skin that lines the EAC is very common. The frequency of otitis externa among swimmers underscores the importance of keeping the EAC dry and explains why otitis externa is commonly known as “swimmer’s ear.”

The presenting symptoms of otitis externa include otalgia, decreased hearing, sensation of fullness in the ear canal, pruritus, tenderness on palpation, and movement of the EAC or the pinna. Otorrhea, adjacent cervical lymph node enlargement, and local cellulitis may develop later in the course of more severe cases. The most common bacterial causes of OE include Pseudomonas aeruginosa and Staphylococcus aureus [1]. Typical physical examination findings include erythema and edema of the EAC with debris, cerumen, and purulent material filling the canal. Visualization of the tympanic membrane (TM) is often obstructed by the otorrhea and the swelling associated with the inflammation. An unimpaired view of a normal TM, with visible landmarks, is shown in Fig. 4.1.

The diagnosis of OE is made based on the history and the clinical findings. Typical otoscopic findings of OE are shown in Fig. 4.2. When the EAC is cultured, one must be cognizant that culture results may reflect EAC flora rather than a causative organism. First-line antibiotic treatments include topical otic drops of a fluoroquinolone, such as ciprofloxacin with or without topical glucocorticoid drops. If edema of the EAC is severe, placement of a wick inside the EAC may be necessary to ensure delivery of medication to the more proximal areas of infection. Risk of recurrence of OE is increased in individuals with atopic dermatitis, seborrhea, immune compromise, and repeated local trauma to the area when

![Fig. 4.1 Normal tympanic membrane. (Courtesy of Dr. Charles Woods, SUNY Upstate Medical University)](image-url)
40

4.3.2 Acute Otitis Media

The development of AOM begins when there is dysfunction of the eustachian tube. Under normal conditions, the eustachian tube allows the middle ear to drain to the pharynx and to equalize pressure between the middle ear and the environment. Impaired drainage may be present for several reasons. The anatomic position of the shorter eustachian tube in young children maintains a relatively horizontal orientation allowing the drainage to defy gravity. Adenoidal hypertrophy and anatomic anomalies of the palate can also block or impair normal drainage. The presence of gastroesophageal reflux disease, allergic rhinitis, and viral upper respiratory infections can all lead to inflammation of the eustachian tube and surrounding tissues resulting in the presence of increased secretions in the middle ear that accumulates because the eustachian tube is not fully patent [2–4]. In each of these clinical scenarios, negative pressure develops in the eustachian tube and middle ear space. The small number of bacteria normally present in those secretions replicates in the now closed space resulting in acute infection (AOM). Retained nasopharyngeal secretions are very common in individuals with otherwise uncomplicated viral respiratory infections. A subgroup of these individuals experience secondary bacterial infection because the initial viral infection and its associated inflammation allow for the closed space conditions where the bacteria can thrive.

4.3.3 Risk Factors

The risk factors for AOM include any condition that promotes eustachian tube dysfunction. The relative horizontal position of the eustachian tube during early childhood explains why the peak incidence of AOM is between 6 and 18 months of age. Additional risk factors include children who develop their first episode of AOM prior to 6 months of age, children in daycare, and the presence of atopy, adenoidal hypertrophy, chronic sinusitis, ciliary dysfunction, immune-compromising conditions, and craniofacial anomalies. Individuals with trisomy 21 are especially prone to otitis media because their eustachian tubes are short and their pharyngeal muscle tone is weak [2, 3].

4.3.4 Microbiologic Causes of AOM

An accurate description of the infectious etiologies of AOM requires that middle ear fluid is removed during acute infection and submitted for microbiologic testing. The procedure to remove the fluid, tympanocentesis, is no longer performed on a routine basis. Most of the literature that describes the microbiologic causes of AOM was published before the routine introduction of \textit{Haemophilus influenzae} type B (late 1980s) and heptavalent (2000) and 13-valent (2010) conjugate pneumococcal vaccines. Rates of viral, bacterial, and mixed culture results from middle ear effusion vary significantly across those studies with rates of bacterial infection ranging between 50% and 90% of all AOM. The most common viral causes include respiratory syncytial virus; parainfluenza viruses, types 1, 2, and 3; influenza A and B viruses; adenovirus; coronaviruses; parechoviruses; and human metapneumovirus.

The most common bacterial causes of AOM include \textit{Streptococcus pneumoniae}, non-typeable \textit{Haemophilus influenzae}, and \textit{Moraxella catarrhalis}. \textit{S. pneumoniae} was unequivocally the most common bacterial agent of AOM before conjugate pneumococcal vaccine was added to the universal pediatric immunization schedule in 2000. Following vaccine introduction, non-typeable \textit{H. influenzae} became more predominant, and “replacement” pneumococcal serotypes that were not included in the 7- or 13-valent vaccines emerged [2–4].

Non-typeable \textit{H. influenzae} should be suspected in the clinical setting of AOM when purulent conjunctivitis is also present. Suspicions that \textit{H. influenzae} could be the underlying cause of the patient’s condition is important, because unlike \textit{S. pneumoniae}, \textit{H. influenzae} may produce a beta-lactamase, an enzyme that inactivates some of the most common antibiotics used empirically to treat AOM, such as amoxicillin. Less common bacterial etiologies of AOM include...
4.3.5 Approach to the Diagnosis of AOM

AOM is a closed space infection. As the infection progresses, symptoms change from a feeling of fullness to general aches. The pain intensifies as the pressure in the infected space increases until that pressure is relieved. Relief may come spontaneously as serous fluid is reabsorbed or when the pressure exceeds the capacity of the TM and the eardrum ruptures. Tympanocentesis can also be used as a controlled technique to remove middle ear fluid and thereby reduce the pressure. AOM causes ear pain. The pain in young children can manifest itself as fussiness, sleep disruption, or ear tugging. Fever is expected in young children with AOM, but uncommon among older children and adults with AOM. In cases where the TM ruptures, the pain is relieved, but the patient or parent will note a purulent or bloody ear discharge.

A diagnosis of AOM is often suspected while obtaining the history of the illness. The diagnosis is confirmed by finding on the physical examination. A thorough otoscopic exam, including pneumatic otoscopy and clear visualization of the TM and EAC, is essential. Presuming a diagnosis of AOM in the absence of a thorough examination leads to overdiagnosis and subsequent overuse of antibiotics. Pneumatic otoscopy should always be included to assess the mobility of the TM. Efficient pneumatic otoscopy is aided by using a speculum with a tight seal of the EAC [4]. Classic otoscopy findings of AOM include the presence of fluid in middle ear space, outward bulging of the TM secondary to the increased pressure in the middle ear space, loss of visualization of ossicle bony landmarks behind the TM, and erythema of the TM. The middle ear effusion is purulent in AOM but must be carefully distinguished from opaque noninfectious effusions secondary to OME. The absence of erythema of the TM suggests a diagnosis of OME, not AOM. A bulging TM with decreased mobility on pneumatic otoscopy is the most specific exam finding for bacterial AOM [4]. The presence of purulent otorrhea in the EAC is seen if a TM perforation has occurred before the time of examination. This finding must be distinguished from the edema and erythema that occur along the EAC during otitis externa. Bullae may be present on the TM in conjunction with other signs of acute inflammation. The findings are consistent with a form of AOM termed bullous myringitis. The presence of bullae does not suggest a specific microbiologic cause nor does it change the clinical approach to the condition.

In summary, the clinical diagnosis of AOM is made when there is an acute onset of symptoms, the presence of a middle ear effusion, and objective signs of acute middle ear inflammation such as a red bulging eardrum that does not move during pneumatic otoscopy [2–4].

The diagnosis of AOM does not involve any laboratory evaluation or radiographic imaging unless there is evidence for a significant complication or severe or persistent disease. Tympanocentesis with culture of the middle ear effusion is the gold standard for diagnosing the etiologic agent of AOM, but is only necessary in a small subset of cases. When necessary, referral to an otolaryngologist is considered the best practice for those who are not certified and experienced in performing the procedure. The inclination to collect samples from the EAC for microbiologic cultures, even in cases where there is frank otorrhea, is generally discouraged since the laboratory results often reflect EAC flora rather than identifying the true causative organism(s).

4.3.6 Differential Diagnosis of AOM

The differential diagnosis for AOM includes otalgia secondary to OME, referred pain from dental or pharyngeal disease, local herpes zoster infection, otitis externa, adjacent soft tissue infection, or eustachian tube dysfunction secondary to alternative causes.

4.3.7 Treatment of AOM

Recommendations for the treatment of AOM have evolved over the past several decades. Depending on the details of the clinical circumstance, current recommendations now permit a clinical observation period as an alternative to immediate antibiotic therapy for many cases. This approach is supported by the knowledge that many cases are caused by respiratory viruses and by evidence demonstrating that a reasonable proportion resolve without intervention. Ultimately, the approach used should carefully weigh patient-specific factors such as age, severity of presenting symptoms, risk factors for severe infection, history of AOM, and the patient’s availability to follow up. Safety-net antibiotic prescriptions (SNAP) and wait-and-see prescription (WASP) are prescriptions provided at the time of diagnosis allowing patients and families to employ watchful waiting, with the option to fill an antibiotic prescription if the symptoms persist or worsen over the
next 48–72 h [2–4]. Treatment approach algorithms that account for age, examination findings, and illness severity at the time of presentation are shown in Figs. 4.3 and 4.4. Analgesic should be used regularly early in treatment course to provide symptomatic relief of otalgia [2, 4].

The first-line antibiotic choices for the treatment of AOM are shown in Call Out Box 4.1 [4]. High-dose amoxicillin (80–90 mg/kg/day) is recommended to overcome concerns about adequate coverage against penicillin nonsusceptible *S. pneumoniae*. Recommendations regarding duration of antibiotic therapy are age dependent. Except for intramuscular ceftriaxone, a 10-day course is recommended for children less than 2 years of age, whereas a 7-day course can be considered for children between 2 and 5 years of age. A 5- to 7-day course is usually adequate starting at 6 years of age through adulthood [4].

Antibiotic therapy for the treatment of CSOM requires administration of topical fluoroquinolone antibiotic drops with or without glucocorticoid drops. The medications are able to reach the middle ear space by crossing through the perforation in the TM. Fluoroquinolones are used because their spectrum of activity includes the most common etiologic agents including *P. aeruginosa*. 

![Fig. 4.3 Treatment of AOM in patients <24 months of age](image-url)
4.3.8 Complications of AOM

Complications of AOM encompass spread of infection to structures adjacent to the middle ear and include mastoiditis, petrositis, venous sinus thrombosis, and central nervous system infection including brain abscess.

4.3.9 Follow-Up

Patients under treatment or observation for AOM should be reevaluated for the persistence or worsening of symptoms 48–72 h after the initial assessment. If symptoms are improving at 48–72 h, short-term follow-up is not usually necessary. TM abnormalities seen on physical examination can persist for up to 12 weeks.
4.4 Sinusitis

4.4.1 Introduction

Sinusitis refers to an infection of one or more of the paranasal sinus cavities. Acute upper respiratory viral infections that cause the common cold all involve the paranasal sinuses. The vast majority of these infections are self-limiting and do not require treatment with antibiotics. Acute bacterial rhinosinusitis is a more precise name for the condition caused by bacterial pathogens, and like AOM, it typically occurs when drainage is impaired secondary to the inflammation associated with a recent viral infection. Fortunately, only a small number of “colds” become complicated by acute bacterial sinusitis. While viruses and bacteria account for most infections of the paranasal sinuses, molds can cause some of the more severe disease. Immune-compromised patients are particularly vulnerable to these disfiguring, often fatal mold infections.

4.5 Definitions

4.5.1 Sinusitis: Inflammation of the Paranasal Sinus Cavity Mucosa

- **Acute bacterial rhinosinusitis (ABRS)** – A bacterial infection leading to inflammation of the paranasal sinuses which resolves, with treatment, within 30 days
- **Subacute bacterial rhinosinusitis (SBRS)** – A bacterial infection leading to inflammation of the paranasal sinuses which resolves in 30–90 days
- **Recurrent acute bacterial rhinosinusitis (RABRS)** – Bacterial infections leading to inflammation of the paranasal sinuses lasting fewer than 30 days but recurring 3 or more times in a 6-month period or 4 or more times during a 12-month period. Each infection responds well to treatment with antibiotics, and the patient experiences at least 10 symptom-free days in between episodes.
- **Chronic sinusitis** – Inflammation of the paranasal sinuses for more than 90 days with persistent symptoms. This condition is most commonly associated with noninfectious processes such as environmental
acidity, reactions to environmental pollutants, and gastroesophageal reflux disease. Patients with cystic fibrosis and ciliary dyskinesia are especially prone to chronic sinusitis secondary to noninfectious and infectious triggers.

### 4.6 Basic Concepts

#### 4.6.1 Anatomy and Pathophysiology

The anatomy of the paranasal sinuses changes from birth, not becoming fully aerated until adulthood [5, 6]. The ethmoid sinuses, present at birth, grow proportionally with the child. Maxillary sinuses, also present at birth, reach adult size by 4 years of age. The sphenoid sinuses begin to develop around 2 years of age, are functional by age 5, and fully mature by age 12. Finally, the frontal sinuses are first evident by age 6–8 years, not fully maturing until early adulthood.

Acute bacterial rhinosinusitis is incited by predisposing factors such as viral upper respiratory infections, allergic rhinitis, environmental pollutants, nasal septum anomalies, other craniofacial anomalies, adenoidal hypertrophy, masses, polyps, or the presence of a foreign body. Risk factors include smoking, second-hand smoke exposure, and extensive dental disease. All of these conditions impede mucociliary clearance of sinus secretions. Since the sinus cavities are not sterile sites, stasis of secretions increases the likelihood that resident bacterial flora will overgrow resulting in acute bacterial rhinosinusitis [5, 6].

#### 4.6.2 Risk Factors

Acute bacterial rhinosinusitis is most common between 4 and 7 years of age. Children in this age group are old enough to have relatively developed sinuses, yet young enough to experience frequent predisposing viral upper respiratory infections. These upper respiratory infections impede usual sinus clearance. Respiratory viral infections are even more common among children less than 2 years of age, but these younger children have less developed sinuses and wider ostia that facilitate drainage. In addition, younger children will often present earlier with AOM. Treatment of the AOM effectively treats any emerging sinus disease, therefore eliminating the opportunity for the development of acute bacterial sinusitis. A second peak of ABRS occurs during adulthood between 45 and 64 years of age. When the infections become recurrent, it is important to maintain a level of suspicion for the presence of contributing factors such as an anatomic blockage (polyps or a foreign body), gastroesophageal reflux disease, allergic disease, a humoral immune deficiency, cystic fibrosis, ciliary dysmotility, or any other condition that might lead to ongoing sinus inflammation and impaired sinus drainage [5, 6].

#### 4.6.3 Microbiologic Causes of Acute Bacterial Rhinosinusitis

The optimal manner in which to determine the precise microbiology of acute bacterial rhinosinusitis is to obtain sinus fluid during the acute infection. Existing data, obtained decades ago, are based largely on studies where needle aspirates were performed on infected maxillary sinuses and the fluid sent to the microbiology laboratory for culture. The technique is modestly invasive and now seldom performed outside of academic settings, clinical trials, and severe or particularly enigmatic cases [7]. Not unexpectedly, the common causes of sinus infections are nearly identical to the list of pathogens that cause AOM since both processes arise from bacteria that normally inhabit the human upper respiratory tract at low concentrations.

The most common bacterial agents of ABRS include *S. pneumoniae*, non-typeable *H. influenzae*, and *M. catarrhalis*. *S. pneumoniae* and non-typeable *H. influenzae* are equally common in the post-pneumococcal vaccine era with each accounting for approximately 30% of cases. Infection with *M. catarrhalis* is less frequent at an estimated 10% of total cases [5, 6]. Formal sinus aspirate studies have reported that approximately 30% of nasal sinus aspirates are culture-negative for bacteria [7]. Less common causes of ABRS include *S. aureus* and *S. pyogenes*. Anaerobic bacteria have also been implicated either as a primary pathogen or as co-pathogens in several adult studies including *Prevotella*, *Bacteroides*, and *Peptostreptococcus* species. Anaerobic infection should be suspected if there is evidence of extension from dental disease into the sinuses.

#### 4.6.4 Approach to the Diagnosis

The initial presentation of sinusitis varies. A careful history of the clinical course of the symptoms over time, the severity of symptoms, and the presence of fever are key details needed to help distinguish ABRS from the more common viral upper respiratory infections. Symptoms more suggestive of ABRS include fever, headache, facial pain and swelling, and halitosis, while cough, nasal congestion, and sore throat are typically seen during bacterial and viral infections [6] [Call Out Box 4.2].

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**Call Out Box 4.2**

A good general rule of thumb: If more than two mucous membranes are involved during a respiratory infection, the cause is viral, not bacterial. Bacterial infections of the upper respiratory tract tend to be localized. Viral infections can involve the eyes, ears, nose, sinuses, mouth, and pharynx simultaneously.
The color of the nasal discharge is sometimes used to justify treatment with antibiotics. Viral upper respiratory tract infections usually begin with symptoms of clear, watery, nasal discharge. As the infection and associated inflammation proceed, the discharge becomes thicker and often takes on a yellow-green color. This change is expected during a viral infection and does not herald the presence of a bacterial process. The change in color and consistency is secondary to innate immune responses to the virus, with recruitment of inflammatory cells, including neutrophils. Neutrophil myeloperoxidase allows for some of the oxygen-free radicals to be converted to hypochlorite. It is the presence of this halogenated compound that renders the color change, not the presence of bacteria. Acute bacterial infections trigger the same innate immune inflammatory pathways, explaining why the discharge seen during a bacterial infection also has a yellow-green color. While the color and consistency of the nasal discharge is not helpful in distinguishing viral from bacterial sinusitis, several characteristics of the infection are. Table 4.1 lists some of the ways in which the viral and bacterial sinus infections can be differentiated from one another on clinical grounds.

Physical examination findings associated with ABRS include erythema and edema of the nasal turbinates, mucopurulent rhinorrhea, and the presence of postnasal drip. Facial pain and sinus tenderness to palpation are less reliable finding in children, but are quite useful indicators in adolescents and adults. None of the physical examination findings are completely specific to either a bacterial or viral infection. The clues obtained during the history of present illness are often the most important key.

Taking the history of present illness and physical examination findings into account, reliable diagnostic criteria for ABRS have been developed [5, 6] [Call Out Box 4.3].

Imaging is rarely useful in the evaluation of uncomplicated bacterial rhinosinusitis. Plain radiographs or computerized tomography imaging of the sinuses undertaken in the absence of compelling history and severe clinical course is likely to demonstrate misleading results. Mucosal thickening, sinus opacification, and air fluid levels may be present, but none of these findings are specific for bacterial underlying etiology. Sinus abnormalities are typically present on imaging in those who have simple viral upper respiratory infections [5]. In contrast, if bacterial rhinosinusitis is present AND complications or spread to adjacent structures is suspected, imaging is always indicated. Computerized tomography imaging of the sinuses and surrounding structures is first line to define the sinus anatomy and bony structures. If there is suspicion that the process has spread intracranially, magnetic resonance imaging of the brain should be considered.

Laboratory evaluation, as with imaging, is not warranted during the evaluation of uncomplicated ABRS. However, if the patient is ill appearing, is immunocompromised, and fails antibiotic treatment, or local extension of the infection is suspected, a sinus aspirate may be warranted. When obtained, the sample is typically sent for Gram stain, aerobic and anaerobic cultures with susceptibility testing. The presence of more than 10,000 colony forming units of a bacterium is considered clinically significant [5].
The differential diagnosis of bacterial rhinosinusitis includes viral upper respiratory infection(s), allergic rhinitis, a local reaction to environmental pollutants, the presence of a nasal foreign body, and adenoidal hypertrophy or infection. If the chief complaint is a prolonged cough illness, infection with *Bordetella pertussis* should also be considered.

Treatment of bacterial rhinosinusitis is dependent on the clinical course of the illness. Subacute, recurrent, and chronic bacterial rhinosinusitis all warrant prompt antibiotic treatment. In contrast, “persistent” ABRS may be followed with clinical observation for up to 3 days. If symptoms continue or worsen during the observation period, antibiotic therapy is initiated. An observation course must be discussed in detail with patient and indications for returning to care must be explained [5, 6].

First-line antibiotic treatment options for ABRS include either amoxicillin or amoxicillin with clavulanic acid. The use of amoxicillin alone raises concerns regarding coverage of *beta-lactamase*-producing organisms such as non-typeable *H. influenza*, *Moraxella catarrhalis*, *S. aureus*, and most oral anaerobes. However, rates of clinical failure with amoxicillin alone are low, and amoxicillin boasts a long history of safety and tolerability. Amoxicillin-clavulanate provides broader antimicrobial coverage but can cause gastrointestinal side effects and is less palatable in suspension formulations used for young children. Therefore, either medication can be considered first line in the treatment of uncomplicated bacterial rhinosinusitis.

Certain conditions increase the risk for treatment failure on amoxicillin alone and warrant amoxicillin-clavulanate as the preferred first-line treatment. Included in this group are patients with immune-compromising conditions, those who are incompletely immunized, patients who have received antibiotics during the last 4 weeks or recently required hospitalization, and those with chronic bacterial rhinosinusitis [6]. If multidrug-resistant *S. pneumoniae* is suspected or confirmed as the etiologic pathogen, a third-generation cephalosporin (cefdinir, cefixime, or ceftriaxone) or quinolone class antibiotic, such as levofloxacin, may be necessary [6].

Patients with non-type 1 penicillin hypersensitivity will most likely tolerate advanced-generation cephalosporins without allergic manifestations. Those with type 1 penicillin allergy warrant treatment with a non-penicillin alternative such as levofloxacin, clindamycin, or linezolid [5].

If the illness severity warrants inpatient treatment, intravenous antibiotic regimens with antimicrobial spectra of activity that encompass all of the usual suspects include ampicillin-sulbactam, cefotaxime, ceftriaxone, or levofloxacin. In circumstances where intravenous treatment failure is suspected, multidrug-resistant *S. pneumoniae* should be considered as a potential cause while also referring the patient for surgical consultation to determine if a drainage procedure might facilitate improvement. While awaiting surgical advice, vancomycin can be added to the antibiotic regimen to broaden coverage that includes highly resistant *S. pneumoniae* and methicillin-resistant *S. aureus*. Intravenous metronidazole can also be considered to include coverage for the most resistant anaerobes including *B. fragilis* [5].

Adjunctive therapies to treat nasal symptoms, such as intranasal steroids, saline irrigation or lavage, mucolytics, decongestants, and antihistamines, lack a robust evidence basis, but remain common practices by patients and providers.

Complications of bacterial sinusitis occur when the infection extends to adjacent structures. Localized spread of sinus disease may result in periorbital or orbital cellulitis. Bony involvement may include the formation of a subperiosteal abscess. Orbital extension should be suspected when there is local tissue swelling and edema in the periorbital area. Intracranial extension can include septic cavernous sinus thrombosis, meningitis, brain abscess, or osteomyelitis of the frontal bone. Intracranial disease should be considered if the patient develops persistent and severe headache, mental status changes, focal neurologic examination findings, or persistent emesis.

### Case Study

#### Practical Example

**Case 1**

A 7-year-old boy presents to the office for a 14-day history of illness that began with cough, rhinorrhea, and complaints of a sore throat. Symptoms gradually worsened for the first few days, but began to improve on day 6. His parents assumed this had been “a typical cold.” On day 7 of illness, the boy’s symptoms worsened. His cough and rhinorrhea persisted, but secretions became thick and purulent again. He developed new facial pain, fevers to 39 °C, and general malaise. The day prior to the current office visit, the patient began to complain of right eye pain. On physical examination, he appears tired. The right eyelid and surrounding skin is red and swollen. His extraocular movements are intact, but he complaints of discomfort when moving his right eye. He also complains of pain during gentle retropulsion. The conjunctivae are clear and there is no chemosis [Call Out Box 4.4]. His nares show purulent rhinorrhea. The TM’s are unremarkable. The oropharynx is erythematous with postnasal drainage present. The nasal turbinates are erythematous and edematous.
4.7 Mastoiditis

4.7.1 Introduction

Mastoiditis is a suppurative infection of the mastoid air cells that most often presents as a secondary complication of otitis media. Its presentation and progression can be acute and require hospitalization and urgent surgical intervention. Routine vaccination against *S. pneumoniae* has reduced the incidence of AOM and led to a decrease in the frequency of mastoiditis in the past decade.

4.8 Definitions

**Mastoiditis** – A suppurative bacterial infection of the mastoid air cells

**Acute mastoiditis** – Mastoiditis with symptom duration of less than 1 month

**Coalescent mastoiditis** – Infectious destruction of thin bony septae between mastoid air cells

**Subacute or masked mastoiditis** – A low-grade persistent middle ear and mastoid infection with destruction of the bony septae between the air cells

**Chronic mastoiditis** – Mastoiditis with symptoms exceeding 1 month in duration

4.9 Basic Concepts

4.9.1 Pathophysiology

The mastoid sinuses are networks of air cells divided by bony septae located in the posterior portion of both temporal bones. The space connects to the middle ear by a bony passage that is present at birth and increases in size until approximately 2 years of age. As the bony passages grow, they become lined with epithelium contiguous to the middle ear space. This continuity of epithelium allows for the possibility for infections of the middle ear to spread to the mastoid. The mastoids are bordered anteromedially by the middle ear and ossicles, the facial nerve, the bony portion of the EAC, the jugular vein, and the internal carotid artery and medially by the inner ear. The sigmoid sinuses are immediately posterior to the mastoids. The cranial fossa sits above, and the soft tissue and muscles of the lateral neck sit below the mastoids on both sides [8, 9].

Mastoiditis is invariably preceded by inflammation and effusion in the middle ear space. The presence of the effusion creates increased pressure between the middle ear and the mastoid air cells. Untreated spread of infection from the middle ear to the mastoid air cells results in resorption of the thin dividing bony septae and osteomyelitis of surrounding temporal bone.

4.9.2 Risk Factors for the Development of Mastoiditis

As mastoiditis is a secondary complication of AOM, the two infectious processes share the same set of risk factors. Those at risk for severe mastoiditis include patients with immunodeficiencies, functional or anatomic asplenia, and chronic heart or lung disease. Patients who are underimmunized or unimmunized against *S. pneumoniae* are also at increased risk. Individuals with cochlear implants are at risk for hardware-based infections [8].

4.9.3 Microbiologic Causes of Mastoiditis

The most common causes of acute mastoiditis include *S. pneumoniae*, *S. pyogenes*, and *S. aureus* (including methicillin-resistant strains). Less common etiologies include *P. aeruginosa* and other Gram-negative bacteria, non-typeable *H. influenzae*, and resident anaerobes of the oropharynx. *Mycobacterium tuberculosis* is an unusual cause
of mastoiditis but should be considered as a possibility when
the illness is diagnosed in individuals who live or have lived
in TB endemic areas of the world.

Chronic mastoiditis is most frequently caused by P. aeru-
ginosa. Other pathogens to consider when the infection pres-
ents as a chronic process include other Gram-negative bacilli,
S. aureus, and anaerobes [8].

4.9.4 Approach to the Diagnosis
of Mastoiditis

Acute mastoiditis presents with abrupt onset of fever, otalgia,
and a red, swollen postauricular area. The pathophysiology
dictates that these symptoms follow a recent middle ear infec-
tion. If the initial AOM was treated with antibiotics, there may
be an interval of improvement followed by an abrupt worsen-
ing of symptoms. The physical examination reveals an abnor-
mal middle ear consistent with AOM. Otorrhea is present in
approximately half of the cases because the TM has perforated
under the pressure of the infected space. The affected ear will
show edema and erythema of the posterior auricular area. The
pinna begins to protrude due to edematous displacement. As
the infection progresses, fluctuance may develop in the post-
auricular area over the mastoid air cells.

Chronic mastoiditis is preceded by long-standing middle
ear disease. Fevers and postauricular erythema and swelling
become evident as the infection progresses. Chronic mas-
toiditis should be suspected when a patient experiences long-
standing TM perforation with chronic otorrhea. Due to the
chronicity of the process, the patient may also complain of
hearing loss [8, 9].

The diagnosis of mastoiditis is based on the classic clini-
cal signs and symptoms. Once the diagnosis is established,
obtaining fluid for microbiologic culture is important to help
guide definitive treatment. Samples can be obtained via tym-
panocentesis or, more commonly, from a mastoid sample
collected during surgical debridement. Biologic samples
should be sent for Gram stain and aerobic and anaerobic cul-
tures with susceptibility testing. Acid-fast cultures should be
requested in cases where tuberculosis is a possibility [8].

Imaging should be used in cases where mastoiditis has
been diagnosed clinically as the findings will help to guide
decisions regarding surgical care. In the absence of objective
physical examination findings of mastoiditis, the presence of
opacifications in the mastoid air cells is nonspecific since the
finding is also common in uncomplicated serous otitis media
and during uncomplicated AOM [10]. When mastoiditis is
clinically suspected, computerized tomography scanning (CT)
is preferred to evaluate the bony structures. Classic CT find-
ings of mastoiditis include mastoid air cell opacifications,
resorption of the bony septae, and coalescence of air cells. CT
may also be indicated in patients with AOM that has been
unresponsive to antibiotic therapy to rule out the development
of secondary mastoiditis. If intracranial or soft tissue
complications are suspected, magnetic resonance imaging
(MRI) is preferred [8]. Symptoms suggestive of an intracranial
complication include focal deficits on neurologic examination,
hearing changes, vertigo, meningeal signs, or altered mental
status.

4.9.5 Differential Diagnosis of Mastoiditis

Scalp infection, periauricular cellulitis, extension of otitis
externa, and perichondritis of the auricle may all present
with posterior auricular erythema and edema. Parotid swell-
ing secondary to mumps may displace the pinna, but the
direction of deviation is superiorly rather than inferiorly seen
with mastoiditis [8].

The differential diagnosis of abnormal findings of the
mastoid air cells by CT in the absence of objective physical
examination findings of mastoiditis includes underaerated or
sclerotic air cells due to prior AOM and OME.

4.9.6 Treatment of Mastoiditis

All cases of mastoiditis are potential candidates for surgical
intervention, and all cases require treatment with antibiotics.
Antibiotic alone may be appropriate if the illness presents
with focal erythema and edema, but without fluctuance or
signs of adjacent spread. Antibiotic therapy in combination
with surgical intervention is necessary if fluctuance is pres-
ent, there is a history of chronic otorrhea, or the patient has
developed focal neurologic signs and symptoms such as
vomiting, nystagmus, vertigo, or other signs of intracranial
disease. Surgical intervention may be fairly straightforward
with tympanocentesis or placement of tympanostomy tubes
or quite extensive with open debridement of the infected
mastoid tissue and adjacent structures.

The initial antibiotic therapy should be administered par-
terally. The typical duration of therapy is a minimum of 3
weeks. Depending on the clinical response, the antimicro-
bial susceptibilities, and the likelihood of adherence, some
patients may complete therapy with oral antibiotics. The first-
line empiric choice antibiotic for the treatment of acute mas-
toiditis is typically a medication in the penicillin or
cephalosporin class. If the initial presentation is severe, vanco-
mycin is also utilized to ensure coverage against methicillin-
resistant S. aureus and highly resistant S. pneumoniae strains.

Chronic mastoiditis is treated empirically with a modified
penicillin in combination with a β-lactamase inhibitor, such as
ampicillin plus sulbactam or piperacillin plus tazobactam in
combination with gentamicin. This combination provides cov-
verage against, P. aeruginosa, and other Gram-negative bacilli
and anaerobes. In patients with recurrent AOM and a strong
suspicion of P. aeruginosa, piperacillin-tazobactam, ceftazi-
dime, or cefepime can be used along with clindamycin [8].

The prognosis of mastoiditis is good if the disease is diag-
nosed and treated before a more serious complication devel-
ops. Increasing rates of morbidity and mortality are seen
when the infection results in septic thrombosis of the cavern-
ous sinus and/or spread to the temporal lobe of the brain [11].
4.9.7 Complications of Mastoiditis

As many as 20% of mastoiditis cases are associated with complications. Infections caused by *S. pyogenes* and *S. aureus* can be particularly aggressive. Interestingly, patients who have recently been treated with antibiotics for AOM have some of the highest rates of complications. This suggests that causative bacteria are particularly pathogenic or have become resistant to more conservative treatments [11]. Known complications involve contiguous spread of the infection to adjacent structures and include subperiosteal abscesses, Bezold's abscesses (infection of the sternocleidomastoid and trapezius muscle attachments), facial nerve paralysis, meningitis, subdural empyema, brain abscess, venous sinus thrombosis, labyrinthitis, temporal bone osteomyelitis, cerebrospinal fluid otorrhea, or conductive hearing loss secondary to destruction of the bony ossicles. Progressive disease can lead to systemic infection with bacteremia and distal septic emboli. A rare but classic complication known as Gradenigo’s syndrome is diagnosed when there is petrositis with associated otitis media, ipsilateral medial rectus palsy, eye pain, and possible additional cranial nerve abnormalities [8].

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### Case Study

**Practical Example**

**Case**

A 2-year-old boy presents with redness around his ear. Two weeks prior, he had a “cold” with mild symptoms of rhinorrhea and cough. Seven days into his illness, he developed fevers and began tugging on his right ear. His provider diagnosed him with right AOM and treated him with a 10-day course of amoxicillin. Two days into his course of amoxicillin, he seemed to improve and his fevers resolved. Two days after he completed his antibiotic course, his fevers returned and his mother noticed redness and swelling behind his right ear. On physical examination, the boy appeared fussy and slightly ill. He had clear rhinorrhea. His left EAC and TM were unremarkable. His right TM showed dullness and erythema with visible pus behind it. His right EAC was mildly erythematous and edematous. Manipulation of the right pinna and palpation of the posterior auricular area were quite painful. The postauricular area was erythematous and edematous, but without fluctuance. The right pinna was slightly displaced anteriorly.

- What is the boy’s diagnosis?
- Is imaging warranted?
- Should his management include antibiotics, surgery, or both?
- Is inpatient or outpatient treatment more appropriate in this clinical setting?

**Discussion**

The patient’s age, preceding AOM, and physical examination findings are consistent with acute mastoiditis. His presentation was relatively recent in onset. He does not have fluctuance on physical examination or signs and symptoms of adjacent spread. A case could be made for imaging to confirm mastoid findings and rule out any progressive bony disease. His initial management should include treatment with intravenous antibiotics in the inpatient setting.

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### 4.10 Exercises

Please refer to the supplementary information section for answers to these exercises.

Match the clinical scenario with the most likely pathogen. Each pathogen may be used once, more than once, or not at all.

| Pathogen | Characteristic finding |
|----------|------------------------|
| 1. Most common bacterial cause of acute otitis media | A. *Mycobacterium tuberculosis* |
| 2. Classic bacterial cause of concurrent otitis and conjunctivitis | B. *Streptococcus pyogenes* |
| 3. Most common vaccine preventable cause of otitis, sinusitis, and mastoiditis | C. *Moraxella catarrhalis* |
| 4. Gram-positive cause of mastoiditis that produces a β-lactamase | D. *Haemophilus influenzae*, nontypeable |
| 5. An uncommon cause of mastoiditis associated with ossicle damage and hearing loss that should be considered in refugees from Africa and Asia | E. *Streptococcus pneumoniae*  
F. *Staphylococcus aureus* |
4.11 Summary

Otitis, sinusitis, and mastoiditis represent a spectrum of otolaryngologic infections that cause signs and symptoms overlapping with the ubiquitous viral upper respiratory infection or “common cold.” Clinical guidelines for each stress careful consideration of diagnostic criteria, vigilance for the development of complications, the most appropriate antibiotic options, and when necessary, appropriate surgical interventions.

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Further Reading
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