Acute Pulmonary Infections

LEARNING OBJECTIVES

- Appreciate the epidemiology of acute pulmonary infections that require pediatric intensive care.
- Review the signs and symptoms of bronchiolitis and pneumonia.
- Review host defense mechanisms during acute pulmonary infections.
- Review the common etiologies of bronchiolitis.
- Review the common etiologies of pneumonia.
- Understand the pathophysiology of bronchiolitis and pneumonia in children.
- Understand the treatment options, including modes of ventilation, for bronchiolitis and pneumonia.
- Understand an effective management strategy for parapneumonic effusions and empyemas.

INTRODUCTION

Acute lower respiratory infection is a common cause of morbidity in infants and children, and at times, requires intensive care and mechanical ventilation. Viral bronchiolitis and bacterial pneumonia account for the majority of lower respiratory tract infections that lead to respiratory insufficiency and pediatric intensive care admission. Twenty-seven percent of children who require mechanical ventilation for at least 24 h in pediatric intensive care units are diagnosed with bronchiolitis and 16% have the diagnosis of pneumonia. The median length of time intubated for an acute pulmonary infection leading to respiratory failure is approximately 7 days.

Viral bronchiolitis remains the leading cause for hospital admission in infancy and the most frequent cause of acute respiratory failure in children admitted to pediatric intensive care units in North America. Pneumonia in children younger than 5 years of age has an annual incidence
of 34–40 cases per 1,000. Community acquired pneumonia can also lead to severe respiratory compromise especially in children with pre-existing disease. A detailed understanding of the diverse etiologies and distinct clinical courses of acute pulmonary infections is essential for the pediatric critical care practitioner. This chapter will focus on bronchiolitis and pneumonia as the two leading causes of pulmonary infections leading to PICU admission.

BRONCHIOLOITIS

Epidemiology

Approximately one third of children develop bronchiolitis during the first 2 years of life. Of these, only 1 in 10 (3% of all infants in the United States) will require hospitalization. Although hospitalization rates have increased over the last three decades, mortality remains low. Overall mortality rate is 1–2%, but as high as 5% in high risk infants. Most deaths occur in infants younger than 6 months of age with co-morbidities such as prematurity, congenital heart disease, congenital or acquired lung disease or immunodeficiency.

Etiology of Viral Bronchiolitis

Respiratory syncytial virus (RSV) was first isolated in 1957 and still represents the major cause of bronchiolitis. Other causative viruses include parainfluenza, adenovirus, enterovirus, influenza and most recently human metapneumovirus and human bocavirus (HBoV). In the northern hemisphere, RSV outbreaks occur from October to June. Human metapneumovirus (hMPV) recently has been identified as the causative agent in 3–19% of bronchiolitis cases, possibly surpassing parainfluenza as the second most common etiology. Its prevalence is slightly higher in the late winter and spring. Parainfluenza infections peak at 10 months of age, representing approximately 7–10% of cases of bronchiolitis. Parainfluenza (PIV-3) is endemic throughout the year, but especially common in the late spring.

Males are 1.5–2 times more likely to require hospitalization for bronchiolitis and are likely to have more severe disease. An X-linked genetic trait that results in a reduced tolerance to hypoxia has been postulated and would be consistent with the observation of increased mortality in newborn males with infant respiratory distress syndrome. Virtually all children by the age of two will have been infected with RSV, all children by the age of five will have been infected with hMPV, and all children by the age of nine will have been infected with HBoV.

The remainder of the discussion on bronchiolitis will be divided into RSV and non-RSV bronchiolitis. Although etiologic agents may differ, clinical courses are often similar.

Respiratory Syncytial Virus (RSV)

Respiratory syncytial virus (RSV) accounts for 50–80% of bronchiolitis, infecting one-half of all infants within the first year of life and hospitalizing approximately 120,000 infants yearly (about 3% of affected infants). Approximately 10% of these infants require mechanical ventilation. Co-infection with either hMPV or rhinovirus occurs in 10–30% of young children.

Two types of RSV exist – types A and B. Type A is more common and is believed to cause more severe disease, although data is not conclusive. Both types may exist simultaneously in the community. Infants less than 1 year will typically shed the virus for about 9 days. Children with immunodeficiencies may shed the virus for months. The immune response varies with age and contributes to both termination of the disease and its pathologic features.

The virus is transmitted from respiratory secretions by close contact with infected persons or by contact with contaminated objects or surfaces. There is a 45% RSV transmission rate within families and about one-half of hospital workers will acquire RSV. Therefore, hand washing and the wearing of gowns and gloves is of primary importance to attenuate transmission.
Pathophysiology

Antibody-Mediated Immunity

RSV introduced onto the nasal or conjunctival mucosal surface causes profuse rhinorrhea within a few days. During the first 2 months of life, passively acquired maternal antibodies are protective. However, as maternal antibody titers gradually decrease, infants become susceptible to severe disease. Cell-bound IgA may develop to help clear the virus. Circulating IgG directed against the glycoprotein (G) and fusion (F) proteins (operative in syncytia formation) on the viral surface will develop several days later. Infants less than 3 months of age appear to induce a weaker antibody response likely due to the presence of maternal antibodies. Virus-specific IgE in the respiratory tract is associated with disease severity. Often, complete and effective immune responses are not induced, thus re-infections are possible even during the same season.

Cell-Mediated Immunity

Epithelial cells and alveolar macrophages are key activators of cellular immunity. Although these cells enhance viral clearing, they also contribute to airway inflammation through the release of cytokines and chemokines. These include interleukin (IL)-1, tumor necrosis factor-alpha, IL-6, IL-8, macrophage-inflammatory protein (MIP)-1-alpha and RANTES (regulated upon activation, normal T cell expressed and secreted). Release of these cytokines and chemokines are believed to be partially responsible for airway inflammation and hyperreactivity. The effects of these mediators persist beyond the acute infection and contribute to prolonged pulmonary dysfunction.

Children who require mechanical ventilation have lower peripheral T cell counts compared to hospitalized infants not requiring mechanical ventilation. These infants demonstrate low T cell proliferative responses and interferon (IFN-γ) production. IL-12 is required for the initiation of cellular immunity. The length of time requiring mechanical ventilation has been found to be inversely related to IL-12 production. The role of Th1/Th2-like cytokine profiles, expressed as IFN-γ/IL-4 ratios, is controversial. In some studies, these ratios decreased after polyclonal stimulation in hospitalized infants with RSV. However, more recent studies have shown normal ratios following polyclonal stimulation.

Neutrophils are the predominant cell found in the airways of infants with RSV bronchiolitis. Elevated levels of IL-8 are found in high concentrations in the nasal secretions of infected children and act as a neutrophil chemoattractant. Further evidence of cellular induced injury is seen in post-mortem examination where peribronchial lymphocyte infiltration with bronchial epithelial necrosis is typically present.

Clinical Presentation and Course

Infants typically present with tachypnea, rhinorrhea, cough, low-grade fever, irritability, poor feeding and vomiting. Respiratory rates greater than 60 breaths per minute are often associated with room air saturations of less than 96%. Infants may also have tachycardia, mild conjunctivitis, otitis media, or pharyngitis. Low-grade fever usually persists for 1–3 days. In addition, infants may develop a metabolic acidosis from poor caloric and fluid intake.

Apnea often is the first presenting symptom of RSV bronchiolitis in small infants. The etiology of apnea remains unknown; however, is likely related to the immaturity of the respiratory control center in the brainstem. The incidence of apnea in infants with bronchiolitis is approximately 16–20%.

The heterogeneous nature of RSV induced lung disease can cause atelectasis in some areas and overdistension in others. Chest roentgenograms often show hyperinflation with flattening of the diaphragms and patchy or peribronchial infiltrates. Atelectasis, especially of the right upper lobe, is often seen. Infants may have high lung volumes with the functional residual capacity often being twice normal. The decrease in dynamic compliance and increase
in airway resistance leads to marked increase in work of breathing, often worse during expiration from lower airway obstruction. Alterations in gas exchange and hypoxemia are secondary to a ventilation-perfusion mismatch.

The anatomical differences between young infants and older children contribute to the severity of the disease in the young. Due to the highly compliant cartilaginous chest wall and poor thoracic musculature, the infant’s chest wall has difficulty countering the lung’s inherent tendency towards collapse. This leads to a greater propensity of small infants towards atelectasis compared with older children. The absence of effective collateral ventilation in infants also contributes to the development of atelectasis and impaired gas exchange. Cellular debris in small airways and peribronchial edema increase airways resistance leading to wheezing as the predominant symptom in some infants.

Despite the potential for severe impairment in lung function, most hospitalized infants improve within 3–5 days. Typically, by 2 weeks, they have normal respiratory rates, oxygenation, and ventilation. Chest radiographs usually normalize by day 9. However, about 20% of infants will have a protracted course, with some mild respiratory symptoms persisting for months.

Viral respiratory infections have been linked to the development of asthma later in childhood. The Tucson Children’s Respiratory Study group prospectively followed for 13 years, 880 infants who had bronchiolitis and found an increased risk for subsequent wheezing episodes.

**High Risk Populations**

Some infants are at an increased risk for severe RSV disease such as those with chronic lung disease due to prematurity (bronchopulmonary dysplasia), cystic fibrosis, congenital heart disease, and immunodeficiencies. In children with cystic fibrosis, RSV accounted for 18% of symptomatic infections, 33% of hospitalizations for infants less than 1 year, and 43% of infants requiring mechanical ventilation. In a study of hospitalized infants with congenital heart disease infected with RSV, 33% required intensive care, 19% received mechanical ventilation, and 3.4% died. Children having undergone hematopoietic stem cell transplants who develop RSV infections have an extremely high mortality of 60–80% despite mechanical ventilation and antiviral therapy. Environmental factors such as crowding, passive exposure to tobacco smoke, and lack of breast-feeding are associated with the development of severe disease. Compared to national averages, Native American and Alaskan children younger than 1 year of age have higher rates of infections.

**Non-RSV Bronchiolitis**

**Parainfluenza**

There are three subtypes of human parainfluenza viruses. HPIV-3 is most frequently isolated from children with bronchiolitis, while PIV-1 and PIV-2 most commonly cause croup. Similar to RSV, both cell-mediated hyper-responsiveness to viral antigen and virus-specific IgE responses are observed in children with parainfluenza bronchiolitis. Upper airway edema with concomitant obstructive symptoms may be present. Children that are infected with parainfluenza have a significant likelihood of developing asthma later in life.

**Metapneumoviruses**

The human metapneumoviruses (hMPV) are a group of RNA viruses of the Paramyxoviridae family identified in humans in 2001. hMPV appears to be the second most common cause of bronchiolitis in children throughout the world. The majority of children are born with maternal hMPV specific IgG which wanes to around 25% by 6–12 months of age. By age five, essentially 100% of children have been exposed to hMPV and will have neutralizing antibody to hMPV. There are two subgroups, A and B, with group A having more severe clinical symptoms. Clinical presentation of children with this virus is similar to RSV. The pulmonary inflammation generally peaks on day 5 which includes interstitial edema and inflammatory cell infiltrates of the bronchioles and alveoli. These inflammatory changes can persist for up
to 21 days. About half of infected children are 0–12 months of age, and infection is primarily in the winter months.

**Human Bocavirus**

Human bocavirus (HBoV) was recently discovered in 2003. With amino acid sequencing, this new member of the Parvoviridae family was found to be closely related to the bovine parvovirus and the canine minute virus, hence the name bocavirus (BO for Bovine and CA for Canine). Detection of the HBoV from the respiratory tract in symptomatic children and its absence of detection in non-symptomatic controls strongly suggest the virus to have a role in respiratory infections in children. Co-infection is commonly described in up to 60% of samples. It remains unclear if HBoV is a primary pathogen or acts to exacerbate other viral illnesses. The pathogenesis of HBoV has not been well described, but with the high occurrence of wheezing and lower respiratory tract symptoms in children infected with the virus, it is speculated that this virus may be a significant contributor to asthma exacerbations. The majority of infected children have rhinorrhea, cough, and wheezing, however, diarrhea has been reported in up to 25% of these children. In children with high viral loads, HBoV has been detected in the serum suggesting the potential for disease beyond the respiratory tract.

**Influenza A and B**

Both influenza A, including novel influenza strains such as H1N1, and influenza B can cause a clinical picture consistent with bronchiolitis in the small infant. These viruses may cause severe multisystem disease and are discussed in greater detail in the viral pneumonia section.

**Diagnosis**

Rapid diagnostic assays are available for early detection of many viruses. The older assays are antigen-based and include indirect immunofluorescence/direct immunofluorescence (IFA/DFA), enzyme immunoassay (EIA), optical immunoassay (OIA), and neuraminidase activity assays. Although still widely used because they are inexpensive and technically simple, they have a low specificity and sensitivity. Molecular assays are becoming the new “gold standard” for respiratory virus detection – replacing tissue culture that may take days. The published sensitivities and specificities approach 100% when compared to tissue culture or antigen assay. These assays generally use polymerase chain reaction (PCR) amplification. Significant advancements in these assays are being made to simplify the performance of the assay and decrease the required time. The most important cause of false negative test results remains poor specimen handling or inadequate sample collection. Other than aiding with cohorting of hospitalized patients, serologic detection of respiratory viruses is rarely clinically useful.

**Treatment**

Regardless of the viral etiology of bronchiolitis, supportive care remains the mainstay of treatment. Supplemental humidified oxygen is frequently needed. Due to many infants being obligate nasal breathers, frequent nasal suctioning may be beneficial to maintain an unobstructed upper airway. The affected infant or child is often unable to take adequate fluids complicated by increased insensible losses from the respiratory tract; hence, intravenous fluids may be required. Infants and children with severe respiratory distress should be kept NPO in the event respiratory failure ensues and endotracheal intubation is required. Antibiotics are not routinely indicated in previously healthy children infected with RSV. Progressive disease, leukocytosis, persistent fever, consolidation on radiograph or systemic toxicity should prompt an evaluation of bacterial co-infection and the use of empiric antibiotics.

High risk patients often require close monitoring and care in an intensive care unit. These include infants less than 6 weeks of age or infants with a history of prematurity, congenital heart disease, bronchopulmonary dysplasia, immunodeficiency or neurologic disease. Infants with RSV bronchiolitis typically have a combination of hyperinflation, pulmonary infiltrates,
and atelectasis. Therefore, no one mode of ventilation can be recommended for all infants. Non-invasive positive pressure (NIIPP) modes (CPAP or BiPAP) may be attempted in infants where their primary respiratory embarrassment is secondary to atelectasis. However, this may not be suitable if the disease process appears severe or protracted as prolonged use of NIPP may make feeding difficult, cause breakdown of facial tissue, or be difficult to maintain without significant sedation that further compromises ventilation. If an infant requires endotracheal intubation, the mode of mechanical ventilation should be tailored to the predominant lung pathology present (i.e. atelectasis versus hyperinflation). Children with significant air trapping may need mechanical ventilation similar to a child with asthma, providing low respiratory rates and longer inspiration and exhalation times. The more typical infant will lose functional residual capacity (FRC) because of atelectasis and alveolar infiltrates. Therefore, despite having some air trapping, these infants often need PEEP to be adjusted to recruit alveoli and return FRC to normal. In the setting of elevated pulmonary vascular resistance (PVR) which may occur in infants with congenital heart disease or bronchopulmonary dysplasia, lowering PVR by traditional methods such as maintaining oxygenation, deep sedation, muscle relaxation and even nitric oxide may be indicated.

Ribavirin is the only FDA-approved antiviral drug for RSV. Ribavirin inhibits viral replication and is active against RSV, influenza A and B, adenoviruses, and hepatitis viruses. For lower respiratory tract diseases, ribavirin is typically administered via aerosolization. In 1996, a meta-analysis of studies involving ribavirin was discouraging and was consistent with the common clinical experience that ribavirin did not improve clinical outcomes. Therapy targeted at attenuating the virus-induced inflammatory cascade has also been disappointing. Corticosteroid administration was not associated with reduction in clinical scores, the need for hospitalization, or the length of hospitalization. Routine use of any corticosteroid given via any route (intravenous, enteral or aerosolized) is not indicated, except in patients with pre-existing chronic lung disease.

Bronchodilators have not shown a clear benefit in patients with acute RSV bronchiolitis. In 12 randomized control trials, involving 843 infants, evaluating the effect of salbutamol or albuterol on bronchiolitis, 9 (75%) showed no effect. The remaining three studies demonstrated only a small transient improvement in the acute clinical score. Although the routine use of bronchodilator therapy cannot be recommended, it has become acceptable practice to attempt to see if individual infants are beta agonist responsive or not. If no clinical response is seen after a trial of a beta agonist, its use should be discontinued.

In the 1990s, five randomized trials involving 225 infants, evaluating the effect of nebulized adrenaline (epinephrine) on bronchiolitis showed clinical improvement, with reductions in oxygen requirement, respiratory rate, wheezing, and decrease in pulmonary vascular resistance. Two of these studies showed lower hospital admission rates and earlier discharge. A 2004 Cochrane systematic review suggested a potential benefit with epinephrine administration. However, subsequent studies have not supported its routine use. As with albuterol, a clinical trial in selected infants seems reasonable.

Nebulized hypertonic saline has been used for treating hospitalized, as well as ambulatory, children with viral bronchiolitis with variable success. A recent Cochrane meta-analysis of nebulized hypertonic saline has shown an improvement in clinical scores and decrease in hospital duration.

Several studies have evaluated the benefit of surfactant and nitric oxide for severe respiratory distress. The results have been inconclusive and do not currently support their routine use. Heliox, a mixture of oxygen (20–30%) and helium (70–80%) with lower viscosity than air has been used successfully in cases of airway obstruction, croup, airway surgery, and asthma to reduce respiratory effort during the period of airway compromise. Several studies have shown improved respiratory distress scores in patients on heliox with continuous positive airway pressure obviating the need for intubation and mechanical ventilation.

Prevention

Palivizumab is a neutralizing humanized mouse monoclonal antibody directed against the RSV-F glycoprotein. It was licensed by the Food and Drug Administration (FDA) in 1998
Palivizumab should be used as preventive therapy in infants with chronic lung disease and congenital heart disease. Cardiopulmonary bypass significantly lowers the serum level of palivizumab, so it should be redosed following surgery if continued protection is desired.

Palivizumab should be administered intramuscularly as 15 mg/kg every 30 days for a total of five doses during RSV season, which is generally from November through March, to high risk infants. Infants or children that develop an RSV infection should continue to receive prophylaxis following recovery because the naturally acquired antibodies are not fully protective. Motavizumab, a new, enhanced potency, humanized RSV monoclonal antibody has demonstrated 50–100 times greater neutralizing activity against RSV. In completion of a phase III trial, motavizumab was found equal to palivizumab for the prevention of RSV hospitalization and superior to palivizumab for reduction of RSV-specific outpatient medically attended lower respiratory tract infections (MALRIs).

PNEUMONIA

Pneumonia describes any inflammatory condition of the lung in which the alveoli are compromised by aspirated foreign matter, inflammatory fluid, or cellular debris. Infection is the primary cause of parenchymal injury to the lung. Pathogens include viruses, bacteria and fungi.

Clinical Presentation

Signs and symptoms of pneumonia are non-specific and may be occult in the young infant. Children often have fever, chills, headache, malaise, restlessness, and irritability. Gastrointestinal complaints such as abdominal pain, distention, or emesis may also be present in young children. The symptoms are often preceded by minor upper respiratory tract infections characterized by low-grade fever and rhinorrhea. With more significant involvement of the lower respiratory tract, tachypnea, dyspnea, cough, nasal flaring, grunting, or retractions may be seen. The older child may demonstrate productive sputum and complain of pleuritic chest pain. On auscultation of the chest, rales and/or decreased breath sounds might be heard over areas of consolidation or pleural effusions. However, due the short path for transmission of breath sounds and the small chest size in infants, breath sounds may not be decreased, even in the presence of effusions. Children with pleural irritation might prefer to lie on the affected side with legs flexed and may complain of radiating pain to the neck and shoulder or into the abdomen.

Epidemiology

Community acquired pneumonia (CAP) is a common, and at times, a serious infection in children. The incidence of CAP is 35–40 cases per 1,000 children less than 5 years of age and 11–16 cases per 1,000 children 5–14 years of age. The exact prevalence of the etiologic agents causing pediatric pneumonia is difficult to ascertain. It is often difficult to differentiate viral from bacterial pneumonia based solely on clinical examination. Specific pathogens causing CAP can be determined in only approximately one-third of children using commonly available cultures, antigen detection, or serologic techniques. Blood cultures yield pathogens in only about 10–15% of infants and children with bacterial CAP and many children do not undergo viral testing as it is often unnecessary. With these inherent limitations, it is generally thought that viruses account for approximately 80% of CAP in children under the age of 2 years and approximately 50% of CAP in preschool children ages 2–5 years.
Viral causes decline in the school age and adolescent child and bacterial causes such as *Streptococcus pneumoniae* and *Mycoplasma* become important pathogens (Fig. 25-1).

Overall, bacteria account for 20–30% of community-acquired pneumonias. The likelihood of infection with different bacteria varies by age. In the newborn period, organisms from the maternal genital tract are likely causes and include *Group B Streptococcus*, *Escherichia coli*, enteric Gram-negative bacilli, *Listeria*, and *Chlamydia*. In older infants, *Streptococcus pneumoniae* becomes a significant cause and remains so until 6 years of age. *Group A Streptococcus* and *Staphylococcus aureus* are uncommon causes. *Moraxella catarrhalis* is a common cause of upper respiratory tract disease, but rarely causes pneumonia. About 20% of infants with pertussis will have bacterial co-infection. In children older than 6 years of age, *Streptococcus pneumoniae* remains the most common cause. *Hemophilus influenzae* type B (HIB), and most recently *Streptococcus pneumoniae*, have decreased significantly as causes of CAP due to the widespread use of effective vaccines.

In the older child and young adolescent, the atypical pneumonias, *Mycoplasma* and *Chlamydia*, become more prevalent and viral causes less common. Rare bacterial pneumonias can occur with animal contact and include: *Francisella tularensis* (rabbits); *Chlamydia psittaci* (parrots and birds); *Coxiella burnetii* (sheep); and *Salmonella choleraesuis* (pigs). Children with congenital anatomical defects, immunodeficiencies, and genetic disorders are at increased risk for bacterial, viral and fungal pneumonia.

**Normal Host Defense Mechanisms**

The airways are normally sterile below the sublaryngeal area to the lung parenchyma. There are several protective mechanisms that include anatomic and mechanical factors, local immune defenses, and the systemic immune response. Microbes are filtered by nasal hairs or are expelled from the airways by the epiglottic reflex, cough reflex, and mucociliary apparatus. Immunoglobulin A (IgA) is the predominant immunoglobulin present in the upper respiratory tract. IgA is able to bind two antigens simultaneously, forming large antigen-antibody complexes. In this manner, the microbes are neutralized and removed by ciliary clearance, thus preventing microbial binding to the epithelium. In the lower tract, immunoglobulin G (IgG) provides humoral protection by opsonizing microbes for phagocytosis by neutrophils and macrophages, activating the complement cascade, and by neutralizing bacterial.
endotoxin. Alveolar macrophages produce superoxide anions, hydrogen peroxide, and hydroxyl radicals that serve an important role in the host defense; however, uncontrolled production can lead to lung injury. In addition to oxygen radicals, a number of cytokines are produced by the alveolar macrophages. These include IL-1, IL-6, TNF, transforming growth factor-β (TGF-β), chemotactic factors, platelet derived growth factor, and M-CSF. These cytokines play a central role in phagocytic recruitment and activation.

Infection occurs when one or more of the defense mechanisms is altered or if the inoculum is too large. Pathogens typically gain entry through inhalation of aerosolized material or through aspiration of resistant organisms inhabiting the upper airways. Less frequently, pneumonia can occur via hematogenous spread.

In children with bacterial pneumonia, a significant portion will have a concurrent or preceding viral infection. Viral infection may predispose to bacterial superinfection by reducing clearance mechanisms and by weakening the host immune response.

**Pathophysiology**

Pathogens entering the lower airways evoke an exudative consolidation of pulmonary tissues. Initially, there is hyperemia of lung parenchyma due to vascular engorgement and capillary leak causing exudation and intra-alveolar fluid accumulation. Fibrin is then deposited and the airways are infiltrated with neutrophils. Consolidation causes a decrease in lung compliance and vital capacity and a total reduction in the surface area available for gas exchange. A physiologic shunt (V/Q mismatch) occurs as there is increased blood flow through poorly ventilated segments of lung, resulting in hypoxia. Compensatory hypoxic vasoconstriction may occur in an attempt to reduce V/Q mismatch and hypoxia, especially in localized areas of consolidation.

With treatment, resolution of consolidation will occur in 8–10 days. The exudate undergoes enzymatic digestion and is either reabsorbed or removed by coughing. If the bacterial infection extends into the pleural cavity, an empyema may result.

**SPECIFIC ETIOLOGIES**

**Bacterial Pneumonia**

**Streptococcus Pneumoniae**

*Streptococcus pneumoniae* is a Gram-positive diplococcus that is frequently found in the upper respiratory tract. There are over 80 capsular serotypes with 80% of infections caused by 14 serotypes. It is the most common bacterial cause for pneumonia occurring at a peak age of 13–18 months. Typically, it causes a lobar or segmental consolidation, but it may manifest as patchy infiltrates in infants. Pleural effusions occur in up to 20% of children that require hospitalization (Fig. 25-2). Pneumatocoele formation is rare. Hemolytic uremic syndrome is associated with neuraminidase-producing strains.

Treatment is typically with a penicillin or cephalosporin. Emerging resistance may require initial therapy with vancomycin. In hospitalized patients, parenteral therapy is generally needed for 48–72 h after fever resolves, followed by completion of 7–10 days of enteral therapy.

Pneumococcal conjugate vaccines (PCV) have been developed that confer immunity against 7 and 13 serotypes. The 7-valent PCV (Prevnar) was licensed for use in the United States in 2000. A 13-valent PCV has been recently introduced and will replace the 7-valent PCV. The PCVs have been highly effective at reducing hospitalizations among children younger than 2 years for pneumococcal pneumonia.

PCV is now recommended universally for children younger than 24 months of age and older children at high risk due to underlying diseases. High risk children include those with sickle cell disease and other types of functional asplenia, human immunodeficiency syndrome, primary immunodeficiency, children receiving immunosuppressive therapy, and children with chronic pulmonary or cardiac disease. A 23-valent PCV is available for
high risk children who need expanded serotype coverage. Children with sickle cell disease or functional asplenia should continue to receive antibiotic prophylaxis regardless of whether or not they have received pneumococcal vaccines.

**Chlamydia Trachomatis**

Approximately 50–75% of infants born to *Chlamydia trachomatis*-infected mothers will become infected at one or more anatomical site, including conjunctiva, nasopharynx, rectum, and vagina. About 30% of infants with nasopharyngeal infections will develop pneumonia. The infants usually present at about 4–12 weeks of age with cough and congestion, but an absence of fever. The cough often interferes with the ability to feed. Infants generally have tachypnea and rales on examination and chest x-ray frequently shows hyperinflation. A peripheral eosinophilia may be present. *C. trachomatis* is susceptible to macrolides, tetracyclines, quinolones, and sulfonamides. Erythromycin for 2–3 weeks is the treatment of choice for neonatal pneumonia.

**Chlamydia pneumoniae and Mycoplasma Pneumoniae**

*Mycoplasma pneumoniae* and *Chlamydia pneumoniae* play a greater role in causing respiratory tract disease in children than previously thought. An indolent course that develops over 5–7 days manifested by low-grade fever, scratchy sore throat, aches, and headaches characterizes both pathogens. After a few days, rales may be heard, particularly in the bases where the infiltrates tend to occur. These organisms have been associated with the initiation, promotion, and exacerbation of asthma in children. In addition, a pertussis-like illness with acute bronchitis has been described. A recent study has shown that nearly half of the cases of community-acquired pneumonia in children aged 2–14 years were associated with *M. pneumoniae* or *C. pneumoniae*. Classic atypical pneumonias caused by these organisms are usually mild and self-limited. However, a number of studies have suggested that severe pulmonary infection may occur in otherwise healthy children. Pleural effusions, pneumatoceles, lung abscesses, pneumothoraces, bronchiectasis, chronic interstitial fibrosis, and acute respiratory distress syndrome although rare complications, have all been reported. Serological testing is the most common means of diagnosis, but this is often retrospective. Cultures obtained from swabbing the nasopharynx may take several days to grow. PCR techniques are currently being refined and standardized. Treatment with antibiotics reduces the rate of recurrent wheezing episodes, decreases morbidity, and shortens the duration of symptoms. The organisms are susceptible to tetracyclines, macrolides, and quinolones. The optimal doses and duration of treatment is unclear; however, some data suggest that prolonged treatment for greater than 2 weeks may be more desirable to decrease symptoms and eradicate the organism from the nasopharynx.
**Staphylococcus Aureus**

*Staphylococcus aureus* is a Gram-positive organism that can be found on the skin, nasal mucosa, and other mucus membranes. About 20–30% of children are carriers. It is generally spread by direct contact or by respiratory particles. *S. aureus* is an unusual cause of lower airway disease in otherwise healthy children. It is more typically isolated from infants and young children with debilitating conditions. Primary *S. aureus* pneumonia presents in the winter or early spring with a short febrile prodrome and a rapid onset of pulmonary symptoms. Blood cultures are positive in 20–30% of patients. Secondary staphylococcal pneumonia will have a more prolonged prodrome with no seasonal predilection, but is often seen after influenza infections. As this secondary pneumonia is usually a result of hematogenous spread, blood cultures are positive in about 90% of patients. Unilateral lobar disease is more typical with primary disease, while diffuse bilateral infiltrates are more frequent with secondary pneumonia. Effusions can be diagnosed in about 15% of children at presentation, but ultimately will develop in about 75% of cases. Pneumatoceles occur in up to 45–65% of children. Treatment is with nafcillin or oxacillin, but more organisms are becoming resistant and require therapy for serious or invasive disease with vancomycin, linezolid, daptomycin, or quinupristin-dalfopristin.

Methicillin resistant *Staphylococcus aureus* (MRSA) was once considered to be restricted to hospitals and long-term care facilities. However, community acquired MRSA (CA-MRSA) is now a significant cause of a variety of infections (including pneumonia) in children without prior health care facility exposure. The majority of community acquired MRSA infections involve minor skin and soft tissue infections, but invasive and sometimes fatal infections can occur in otherwise healthy individuals. CA-MRSA and healthcare-associated MRSA (HA-MRSA) can be distinguished by several important features. Patients with CA-MRSA by definition have not had recent hospitalization (acute or chronic care), prolonged antibiotic use or chronic underlying disease. Toxin production also distinguishes CA-MRSA from HA-MRSA. Panton valentine leukocidin (PVL) is a toxin which is present in most CA-MRSA isolates, but rarely in HA-MRSA isolates. PVL toxin lyses white blood cells leading to leukopenia and a decreased ability to kill *S. aureus*. Its production has been implicated as a contributor to the development of CA-MRSA necrotizing pneumonia. CA-MRSA isolates, unlike HA-MRSA, lack multi-drug resistance. CA-MRSA is generally more susceptible to clindamycin, trimethoprim-sulfamethoxazole and doxycycline than HA-MRSA, probably because HA-MRSA has developed resistance to survive in the healthcare setting.

**Group A Beta-Hemolytic Streptococcus**

*Group A beta-hemolytic Streptococcus* (GABHS) is a Gram-positive organism responsible for about 15% of pharyngitis and tonsillitis in children. It is rare as a primary cause of pneumonia. When it does occur, the children generally have high fever and appear toxic. The pneumonia is typically lobar. Associated empyemas are common and pneumatoceles may develop. There are several virulent toxin-producing GABHS M-serotypes that are associated with toxic shock syndrome. Pre-existing varicella disease with disruption of skin and soft tissue as the port of entry is reported approximately 40–50% of the time. An associated pneumonia occurs in 10–20% of children with toxic shock syndrome. GABHS are highly susceptible to penicillins and cephalosporins. In cases of toxic shock, clindamycin is often added to inhibit the production of streptococcal pyrogenic exotoxins A (SPE-A) and B (SPE-B).

**Group B Streptococcus**

About 30–40% of infants with perinatally acquired *Group B Streptococcus* (GBS) infections will have pneumonia. The infant usually has systemic disease and blood cultures are frequently positive. Late-onset GBS is predominantly caused by the type III serotype. In these infants, the infection is usually manifest as bacteremia without a focus or with meningitis. Pneumonia is rare in late-onset disease. GBS is uniformly sensitive to penicillin.

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**While Staphylococcus aureus pneumonia is uncommon, effusions ultimately develop in about 75% of cases and pneumatoceles occur in 45–60%**.
Pertussis

Pertussis, or “whooping cough” is a highly contagious respiratory tract infection caused by the Gram-negative pleomorphic bacillus *Bordetella pertussis* and less commonly *Bordetella parapertussis*. With the development and widespread use of a vaccine in the 1940s, a significant and sustained decrease in incidence has occurred. However, despite immunization rates greater than 80%, cyclical recurrences of the disease have occurred every 3–4 years since the 1980s. This is likely secondary to the waning of immunity in adolescents and young adults. Under-immunized or unimmunized infants are the most vulnerable. Nearly all deaths reported from pertussis occur in infants younger than 3 months of age.

Pertussis is often divided into catarhal (fever, rhinorrhea and initiation of cough), paroxysmal (severe coughing episodes, lymphocytosis, potential for complications) and convalescent stages (slow waning of cough over weeks to months). Complications include secondary bacterial or viral pneumonia, apnea, malnutrition, pulmonary hypertension and neurologic involvement including seizures and encephalopathy. Infants less than 6 months of age are at highest risk for complications and mortality. Characteristic paroxysms of cough with an end inspiratory whoop occur in children. Infants may present with a nonspecific cough with associated apnea and cyanosis, without a whoop. Adolescents may be asymptomatic or have only a mild prolonged cough. An increased white blood count up to 100,000 with a lymphocytosis is characteristic early in the course of the disease. The preferred test for laboratory confirmation is the detection of *B. pertussis* DNA by PCR assay. Bacteriologic culture provides a definitive diagnosis.

If administered during the early stages of the disease (first 7–10 days of illness), erythromycin for 14 days may decrease symptoms and reduce the risk of spread. A 5 day course of azithromycin or a 7–10 day course of clarithromycin has been found to be as effective with less gastrointestinal symptoms. Corticosteroids, bronchodilators, or intravenous immunoglobulins have not demonstrated efficacy. Supportive care with supplemental oxygen, mechanical ventilation, intravenous fluids, maintenance of adequate caloric intake, and treatment of secondary bacterial infections are the mainstay of therapy. The use of extracorporeal membrane oxygenation in infants with hypoxemia, pulmonary hypertension and right heart failure refractory to conventional mechanical ventilation has resulted in poorer outcomes than expected. Vaccination in infancy with booster doses in adolescence is preventative.

Viral Pneumonia

About 80–85% of pneumonias in children are caused by viruses. There is considerable evidence that viral infections often precede bacterial pneumonias and cause weakening of the host defenses. Viral pneumonias with RSV and parainfluenza are discussed in more detail in the bronchiolitis section.

Influenza

Influenza is the main viral cause of pneumonia in school-aged children requiring hospitalization. There are three serotypes, A, B, and C which are further divided into subtypes based on the hemagglutinin and neuraminidase genes. Hemagglutinin 1, 2, and 3 and neuraminidase 1 and 2 typically infect humans. The gene segments for the surface glycoproteins are unstable, so mutations, called antigenic shift, occur regularly. Epidemics occur annually during the winter months with a short, 1–3 day incubation period. The virus causes destruction of the ciliated respiratory epithelium within 1 day of symptoms. Airway edema and infiltration with inflammatory cells into the airway mucosa and epithelium follows. Slow repair occurs over 2–4 weeks. A severe fulminating pneumonia may result in hemorrhagic exudates that contain many polymorphonuclear and mononuclear cells. Destruction of the respiratory epithelium often leads to secondary bacterial infections.

During the 2003–2004 influenza season, 143 influenza-related deaths occurred in children; of these, 41% were less than 2 years of age. Forty-five percent of the older children (2–17 years of age) did not have an underlying medical condition. Rare complications of...
influenza include acute myositis, rhabdomyolysis, myocarditis, pericarditis, Reye syndrome, encephalitis, transverse myelitis, and Guillain-Barré syndrome.

Children may present with an abrupt clinical course manifested by high fever, myalgias, headaches, scratchy sore throats, and dry cough. Peripheral white blood counts are usually less than 5,000. Pulmonary infiltrates often involve multiple lobes. Bacterial co-infection, especially with MRSA, increases morbidity and mortality significantly.

Rimantidine and amantadine can shorten the course for influenza type A disease by limiting viral replication, but only if given within the first 48 h of the disease. Prophylactic dosing is 70–90% effective and does not interfere with antibody production from the vaccine. Both drugs have central nervous system and gastrointestinal side effects, including an increase in the incidence of seizures. Oseltamivir and zanamivir have recently been approved for the treatment of influenza infections in children. They inhibit neuraminidase, an enzyme produced by influenza A and B. The course of disease in healthy adults can be reduced by 1–2 days, if started within 48 h of the onset of symptoms. Zanamivir is a dry powder aerosol that must be delivered by a special breath-activated device. Bronchospasm in patients with asthma has been reported. Aspirin or aspirin-containing products should be avoided due to the risk of Reye syndrome.

Immunoprophylaxis is the most effective strategy for the prevention of influenza infection. Inactivated vaccines have efficacy rates from 70% to 90%. Currently, the inactivated vaccine is recommended for all children older than 6 months of age with high risk conditions including chronic pulmonary or cardiac disease, immunosuppressive disorders, sickle cell disease and other hemoglobinopathies, diseases requiring long-term aspirin therapy, chronic metabolic and renal diseases; healthy children aged 6–23 months; and household contacts over the age of 6 months of high risk persons. A live, attenuated influenza vaccine was licensed in 2003. It is administered by the intranasal route and is approved for healthy children aged 5–17 years.

Avian Influenza

Avian influenza viruses do not normally infect species other than birds and pigs. However, in 1997, the first human death from avian influenza occurred in Hong Kong in a 3 year old with Reye syndrome. Subsequently, an epidemic occurred among humans in Hong Kong with close contact to live, infected poultry. The subtype H5N1 appears to be the most ominous due to its ability to rapidly mutate and infect new species. The overall mortality rate is greater than 70%. The avian viruses are not believed to be transmissible from person-to-person, but some recent cases are being investigated for this possibility. Children uniformly present with fever and cough. Symptoms range from typical influenza-like symptoms to conjunctivitis to respiratory disease and failure. Significant laboratory data include leukopenia and thrombocytopenia. All children who developed pneumonia and progressed to ARDS died. Diagnosis remains difficult, as no tests are widely available. Of the antiviral drugs available for influenza A, the most recent H5N1 strains in Southeast Asia are resistant to rimantadine and amantadine. Therefore, treatment is mainly supportive. A prototype H5N1 vaccine was made available to manufacturers in April 2004, but production is difficult because the standard means of producing influenza vaccines from specially grown chicken eggs is not feasible. H5N1 kills the embryo before enough viruses can be harvested for vaccine production.

Novel H1N1 Influenza A

In April, 2009, The Centers for Disease Control confirmed the emergence of a novel influenza A (H1N1) virus with genes from swine viruses of the Eurasian lineage and genes from avian influenza viruses. By June, 2009, the first influenza pandemic since 1968 was declared, affecting over 191 countries and territories. In comparison to illnesses with seasonal influenza, the majority of cases occurred in individuals younger than 65 years of age, with nearly half of the cases occurring in children under 18 years of age.

The clinical symptoms can be typical for influenza; fever, sore throat, cough, and muscle aches with the addition of vomiting and diarrhea in children. A wide range of complications
have been reported that include mild-to-moderate (otitis media, sinusitis, myositis, and febrile seizures) to more severe complications such as myocarditis, rhabdomyolysis or encephalitis. Severe complications may frequently involve invasive bacterial co-infection (i.e. MRSA) and/or exacerbation of underlying medical conditions in particular asthma. Children who present initially with uncomplicated influenza may have rapidly progressive hypoxemic respiratory failure and multiorgan system dysfunction that is refractory to all therapies (Fig. 25-3).

Of reported H1N1 deaths, approximately 20% were in children. The majority of these children had comorbid asthma, neuro-developmental conditions, or obesity. An American Academy of Pediatrics Work Group identified children at greatest risk for life-threatening H1N1 influenza disease (Table 25-1).

The Centers for Disease Control has recommended prompt empiric antiviral therapy for infants, children, and adolescents of any age presenting with suspected or confirmed H1N1 influenza and any of the following conditions:

- Illness requiring hospitalization
- Progressive, severe, or complicated illness, regardless of previous health
- Presence of significant risk factors (see Table 25-1)

The H1N1 strain has been found to be resistant to amantadine and rimantadine, but is usually sensitive to neuraminidase inhibitors, specifically oseltamivir or zanamir. In 2009, oseltamivir was emergently approved for treatment in children less than 12 months of age. Resistance to oseltamivir has been reported and is thought due to the H275Y mutation. Interestingly, the mutation confers resistance to oseltamivir, but not to zanamir. Peramivir, a neuraminidase inhibitor, an unapproved (investigational) antiviral available in an intravenous formulation received an emergency use authorization permit from the FDA for use in children with confirmed severe refractory H1N1 influenza. Its use should be restricted to children that are not responding to either oral or inhaled antiviral drugs or if the parenteral route is the only dependable method of drug delivery.

A vaccine was manufactured and licensed using the same standards as seasonal influenza by late 2009. A single dose was found to provide adequate protection in children older than 10 years of age, younger children requiring two doses separated by at least 21 days.
Adenovirus

Adenoviruses have been implicated in 4–10% of pneumonias in children. Adenoviruses are classified into 49 serotypes with types 3, 7, 7a, 11, and 21 being the most common etiologic agents of lower respiratory disease and causing a severe necrotizing pneumonitis. These serotypes are associated with serious pulmonary sequelae, such as bronchiectasis, bronchiolitis obliterans, unilateral hyperlucent lung, and persistently abnormal pulmonary function tests. Adenovirus infections peak between 6 months and 5 years of age. Mortality from severe respiratory infections can be high, because the disease often involves multiple organ systems. Survivors may have permanent lung injury often in the form of bronchiolitis obliterans. In the immunocompromised host, mortality rates are as high as 50–80%. Cidofovir has in vitro activity against adenovirus, but proof of efficacy is limited. Therapy is supportive.

Severe Acute Respiratory Syndrome–Associated Coronavirus (SARS-CoV)

Severe acute respiratory syndrome is a newly described pulmonary infection caused by a novel SARS-associated coronavirus. SARS-CoV is highly contagious and was coined “the first plague of the twenty-first century”. The disease rapidly spreads among household contacts and healthcare personnel. Children less than 18 years of age account for only approximately 5% of those affected, with a mean age of 12 years. No deaths were reported among children in the 2003 outbreak. Children and adults present with fever, malaise, cough, coryza, chills or rigor, sputum production, headache, myalgia, leukopenia, lymphopenia, thrombocytopenia, mildly elevated activated partial thromboplastin times, and elevated levels of lactate dehydrogenase. Radiographs of the chest show non-specific infiltrates. Apart from diarrhea, patients have minimal extrapulmonary symptoms. Early diagnosis by reverse transcription-polymerase chain reaction (RT-PCR) can be made with 80% sensitivity on nasopharyngeal aspirates within the first 3 days of the illness. The clinical course follows a triphasic pattern. There is an incubation period of 2–10 days with a prodrome of high fever, chills, malaise, headache, and myalgias. Diarrhea occurs in up to 20% of adults. After 2–7 days, the disease progresses to involve the lower airways with a dry non-productive cough and dyspnea. In 10–20% of cases, acute respiratory distress syndrome (ARDS) follows and often patients require mechanical ventilation. Deaths occur from respiratory failure. Young children run a milder and shorter biphasic clinical course. Cough is found in approximately half the children, and crackles are rarely heard despite radiographic evidence of infiltrates. A regimen of antibiotics, ribavirin, and corticosteroids was proposed based on initial anecdotal success. However, ribavirin has demonstrated minimal activity against SARS-CoV isolates in vitro. Non-randomized studies of corticosteroids have reported favorable outcomes. A pediatric series of 44 children with confirmed SARS treated with ribavirin and corticosteroids showed no adverse effects and all survived.

TABLE 25-1

| HIGH RISK CONDITIONS ASSOCIATED WITH LIFE-THREATENING H1N1 INFECTION |
|---------------------------------------------------------------|
| 1. Neurological disorders, such as epilepsy, cerebral palsy, developmental delay and neuromuscular disorders |
| 2. Chronic respiratory diseases associated with impaired pulmonary function and/or difficulty handling lung secretions, moderate and especially severe persistent asthma, technology-dependent children (e.g., those requiring oxygen, tracheostomy, or a ventilator) |
| 3. Primary immunodeficiencies or conditions that require medications or treatments that result in secondary immunodeficiencies |
| 4. Congenital heart disease |
| 5. Metabolic (e.g., mitochondrial) or endocrine disorders, especially if cardiopulmonary function is impaired |

Adapted from http://www.aap.org/new/swineflu.htm
Hantavirus Cardiopulmonary Syndrome (HCPS)

Hantavirus Cardiopulmonary Syndrome is a viral zoonotic disease that affects healthy children and adolescents who are exposed to aerosols of rodent excreta. The deer mouse is the main rodent reservoir. Most cases occur in the southwestern United States, but cases have been confirmed in 30 states. HCPS presents with a prodrome of fever, chills, myalgia, headache, and gastrointestinal symptoms. Respiratory compromise requiring supplemental oxygen generally occurs within 72 h. The disease can progress to respiratory distress and ARDS. The majority of deaths result from hypoxemia and cardiac dysfunction with marked hypotension and ventricular arrhythmias. In adults, the case fatality rate is approximately 38%. A recent case series of 13 children aged 10–16 years, revealed that 92% of infected children developed HCPS, 33% died, and 67% were critically ill and required mechanical ventilation. Treatment is supportive as ribavirin has not been proven to reduce mortality. Extracorporeal membrane oxygenation was used on two patients, one of which survived. Laboratory evaluation reveals thrombocytopenia, leukocytosis, and circulating immunoblasts. An elevated prothrombin time of $\geq 14$ s is predictive of severe disease. No deaths were reported in children younger than 14 years of age. Diagnosis can be made by detection of hantavirus-specific immunoglobulin M, hantavirus-specific RNA by polymerase chain reaction, or hantavirus antigen by immunohistochemistry.

Pneumonia in the Immunocompromised Host

Respiratory infections in children with primary or acquired immunodeficiencies requiring intensive care are not uncommon. These infants and children are susceptible to many organisms that are rarely pathogenic in a normal host. Primary immunodeficiencies include abnormalities or deficiencies in immunoglobulins and antibodies, T and B cells, phagocytes, natural killer cells, and complement. Acquired immunodeficiencies include asplenia, human immunodeficiency virus (HIV), corticosteroid therapy, and immunosuppression used for marrow or solid organ transplants.

Immunocompromised children can present with attenuated signs and symptoms of respiratory infections. In addition to physical examination and chest roentgenograms, these children often require chest computed tomography to better delineate the extent of disease. Bronchoalveolar lavage, needle aspiration, or lung biopsies might be required to make a definitive diagnosis. Pulmonary specimens should be tested for common bacteria as well as for Pneumocystis Carinii, acid-fast bacilli, Nocardia, Legionella, Crytococcus, Aspergillus, Candida, Histoplasma, Coccidioides, and Blastomyces. Viruses such as cytomegalovirus, varicella, herpes virus, and measles should be considered.

Pneumocystis carinii Pneumonia (PCP)

*Pneumocystis carinii* (now known *Pneumocystis jiroveci*) is an opportunistic pulmonary pathogen in infants and children with human immunodeficiency virus (HIV) and other primary immunodeficiencies, malnutrition, hematological malignancies, solid organ and bone marrow transplant recipients, and patients on high dose corticosteroid therapy for inflammatory and collagen-vascular diseases. It is a unicellular organism that exists as a cyst (the diagnostic form). The organism attaches to the type I alveolar cells resulting in an alveolitis characterized by ventilation-perfusion mismatch and decreased pulmonary compliance. If untreated, PCP carries a mortality rate of 25–50%, and nearly 100% in the HIV-seropositive child. Fortunately, the incidence has markedly decreased with the administration of chemoprophylactic agents to high risk patients. Children typically present with fever, tachypnea, non-productive cough, and hypoxia with an absence of rales on auscultation of the chest. Initially, they may have an elevated pH and low carbon dioxide levels. Lactate dehydrogenase levels are generally elevated. Bilateral diffuse alveolar infiltrates are seen with initial hilar involvement subsequently spreading to the periphery (Fig. 25-4). Diagnosis is made by demonstrating the organism with the methenamine silver nitrate stain on pulmonary tissue, respiratory secretions, or lung fluid. Bronchoalveolar lavage is the most widely used technique to obtain lung fluid for diagnosis. Treatment consists of supportive therapy.
with supplemental oxygen; ultimately continuous positive airway pressure or mechanical ventilation may be necessary if respiratory failure occurs. Trimethoprim-sulfamethoxazole (TMP-SMX) is the recommended initial treatment. In patients that cannot tolerate TMP-SMX, then pentamidine isethionate should be used. Corticosteroids in anti-inflammatory doses as an adjunct to antimicrobial therapy have improved clinical outcomes. Concurrent pulmonary infections were found in 35% of patients, most frequently bacterial or cytomegalovirus pneumonia.

**DIAGNOSIS OF PNEUMONIA**

Determination of the etiologic agent in pneumonia is difficult. Fortunately, in most community-acquired pneumonias, identification of the specific causative organism is not critical. However, in children with a complicated course that fails to respond to standard therapies, definitive diagnosis of the etiologic agent is essential. Complete blood counts, inflammatory markers, and chest radiographs do not differentiate the causative agents for pneumonia. Blood cultures are rarely positive outside of the neonatal period. Rapid antigen tests are available for RSV, parainfluenza, influenza, and adenovirus. Nasopharyngeal swabs for viral cultures generally take 7–8 days to become positive, and in one study, 86% of the patients had been discharged prior to the positive results. Older children and adolescents might be able to produce sputum for Gram stain and culture. An adequate specimen should contain more than 25 leukocytes and fewer than 25 squamous epithelial cells per low-power field. In the intubated patient, sputum can be more easily acquired. However, interpretation of the results of Gram stains and cultures is at times difficult in differentiating colonizing from pathologic organisms. Colonization of the endotracheal tube may occur as early as 12 h, but most frequently between 60 and 96 h. The oropharynx becomes colonized within 36 h, the stomach at 36–60 h, and the lower respiratory tract between 60 and 84 h. In addition, a comparison of infectious agents isolated by both tracheal aspirates and bronchoalveolar lavage found only 36% concordance.

Bronchoalveolar lavage (BAL) can be safely used to obtain secretions from the lower airways for Gram stain and culture. It is especially useful in the diagnosis of pneumonia in the immunocompromised child. However, BAL performed directly through the bronchoscope carries a risk of contamination. The smallest bronchoscope that can accommodate a protected specimen brush is 4.8 mm and requires a 6.5 mm endotracheal tube for passage. The smallest flexible fiberoptic bronchoscope with a suction channel has an external

**FIGURE 25-4**

Chest radiograph of severe *Pneumocystis carinii* pneumonia in a 13 month old male with combined immunodeficiency. Note the diffuse alveolar involvement and air bronchograms. (Image provided courtesy of FA Maffei)
diameter of 2.8 mm and is too small to admit a double-sheathed brush. Non-bronchoscopic double-lumen plugged catheters can be inserted blindly through the endotracheal tube to obtain a non-contaminated specimen. The sensitivity and specificity of these samples are similar to those obtained by a bronchoscopic guided protected specimen. Transthoracic needle aspirations are performed in some centers with good results. One study reported a diagnostic success rate in 59% of patients. The incidence of pneumothorax was approximately 20%, but none required subsequent placement of a pleural drainage catheter. A lung biopsy is rarely needed to make a definitive diagnosis.

**TREATMENT**

Supportive treatment with oxygen and intravenous fluids are often standard therapies. As both pneumonia and mechanical ventilation can cause an elevation in anti-diuretic hormone levels, careful fluid monitoring is essential to avoid overhydration, excessive lung water and hyponatremia. Initial antibiotic choices should be empiric and based upon the likely organisms for each age group, because of the difficulty in identifying the causative agent.

The child’s respiratory status including respiratory rate, work of breathing, pulse oximetry, and central nervous system response should be closely monitored. Non-invasive bi-level positive airway pressure (BiPAP) has been effective for use in children with mild to moderate respiratory insufficiency, defined as an A-a gradient >100 and <250 or PaO$_2$/FiO$_2$ ratio <200 but >100 mm Hg. Serial evaluation of mask-face contact areas is essential to avoid skin breakdown.

Children with moderate or severe respiratory insufficiency often require intubation and mechanical ventilation. Children with respiratory failure secondary to pneumonia often require increased positive end expiratory pressure (PEEP), increased inspiratory time, and aggressive pulmonary toilet to recruit alveoli. For patients requiring high levels of PEEP, adequate sedation is often required to prevent patient/ventilator asynchrony and barotrauma. Spontaneous respirations should be encouraged while on mechanical ventilation. Rarely, the use of neuromuscular blockade is required to allow mechanical ventilation. Prone positioning may improve ventilation/perfusion (V/Q) mismatching in dependent lung regions. Lung protective strategies allowing permissive hypercapnea with small lung volumes to ventilate and appropriate PEEP to maintain alveolar recruitment is recommended for children with pneumonia. High frequency oscillatory ventilation can also be utilized to maintain mean airway pressure and alveolar recruitment. Airway pressure release ventilation (APRV) provides recruitment of alveoli while allowing spontaneous respirations. In children with severe respiratory distress syndrome, treatment with bovine surfactant may improve oxygenation. Extracorporeal life support continues to have a role in children with reversible severe acute hypoxic respiratory failure refractory to mechanical ventilation.

Pneumonias can often be complicated by the development of pleural effusions and empyemas. These occur when the fluid production by the interstitial lung tissue exceeds the maximum pleural lymphatic flow. Parapneumonic effusions often occur from pneumonia as white blood cells and other debris of infection block the lymphatics resulting in elevation of protein in the pleural space, increase in colloid osmotic pressure, and consequent failure of fluid reabsorption. On physical exam, the child will have decreased breath sounds over the effusion. In older children, auscultatory percussion changes might be appreciated. Plain chest radiographs can reveal most clinically significant effusions. Ultrasound and chest computed tomograms are useful in determining the volume and quality of the fluid and the presence of loculations. Simple parapneumonic effusions or transudates can also be differentiated from exudates by using the criteria of Light et al. (Table 25-2). A pleural fluid pH less than 7.2 indicates a complicated effusion that is likely exudative and requires drainage whereas a pleural fluid pH more than 7.3 suggests that the effusion may be managed with systemic antibiotics alone.

Complicated parapneumonic effusions or empyemas occur when the fluid becomes purulent. During this stage, the effusions undergo a fibrinopurulent stage with many polymorphonuclear leukocytes, bacteria, and cellular debris entering the fluid. Fibrin is deposited over the pleural surfaces and loculations begin to form. The pH and glucose levels fall as the LDH levels rise. If untreated, they often progress to a third organizing stage in which the exudate
develops into an inelastic, fibrotic peel that restricts the lung. Simple parapneumonic effusions usually resolve with thoracentesis or tube thoracostomy and antibiotic treatment of the pneumonia. More complicated parapneumonic effusions have been successfully treated with thoracotomy tubes and fibrinolytics. However, although risks for bleeding are reportedly low, this therapy requires close monitoring of chest tube drainage and instillation of expensive medications with intermittent clamping of the chest tube. No single recommendation for the choice of fibrinolytic agent or dosage has been established. Also, if tried late in the organizing phase, this is often unsuccessful due to loculations and the high viscosity of the purulent fluid. Surgical debridement either by open procedure or by video-assisted thorascopic surgery (VATS) is often needed for organizing, complicated parapneumonic effusions. Multiple studies have reported that early VATS or thoracotomy for empyema leads to a shorter hospital stay. The treatment modality is best determined by the temporal stage and nature of the effusion.

**CONCLUSION**

Acute pulmonary infections are common diagnoses that require admission to the pediatric intensive care units. Understanding the pathophysiology of lower respiratory infections enables the intensivist to tailor therapy to the individual child and pathogen. Early establishment of a specific etiology and the selection of the correct treatment plan directly impacts clinical outcome.

### REVIEW QUESTIONS

1. **A 3 month old, former 27 week premature infant with bronchopulmonary dysplasia presents with clinical signs of bronchiolitis. Analysis of nasopharyngeal secretions by polymerase chain reaction testing identifies respiratory syncitial virus.** Which of the following therapies have been proven to be a consistent benefit for RSV bronchiolitis?
   - A. Aminophylline
   - B. Bronchodilators
   - C. Corticosteroids
   - D. Ribavirin
   - E. Supportive care

2. **Palivizumab is indicated for which of the following children?**
   - A. A 5 month old, former 27 week premature infant who just underwent surgical repair of a large ventricular septal defect who received palivizumab 2 weeks ago
   - B. A 9 month old, former 28 week premature infant with mild bronchopulmonary dysplasia who received palivizumab 2 weeks ago
   - C. A 1 month old, former 36 week premature infant with peripheral pulmonic stenosis who has never received palivizumab
   - D. A 2 month old full term infant with a urea cycle defect who has never received palivizumab
   - E. An 8 month old, former 25 week premature infant with bronchopulmonary dysplasia who received his fifth dose of palivizumab a month ago

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**TABLE 25-2**

| LIGHT CRITERIA WITH INDIVIDUAL SENSITIVITY AND SPECIFICITY OF TESTS TO DISTINGUISH EXUDATIVE FROM TRANSUDATIVE EFFUSIONS |
| --- |
| Pleural fluid may be classified as exudative, if one or more of the following criteria are met: |
| ▪ Pleural fluid protein divided by serum protein > 0.5 (Sensitivity 98%, Specificity 83%) |
| ▪ Pleural fluid lactate dehydrogenase (LDH) divided by serum LDH > 0.6 (Sensitivity 86%, Specificity 84%) |
| ▪ Pleural fluid LDH is more than two-thirds of the upper limit of normal for serum LDH (Sensitivity 82%, Specificity 89%) |

Adapted from Light (2002)
3. A 5 year old, unimmunized male with moderately severe asthma requires hospital admission with a 12 h history of fever, cough and myalgias in the middle of an H1N1 influenza outbreak. The most appropriate initial management of this child includes which of the following?
   A. Intravenous peramivir administered after confirming the diagnosis with rapid testing
   B. Intravenous zanamivir administered as soon as possible
   C. Oral oseltamivir administered as soon as possible
   D. Oral oseltamivir administered as soon as possible
   E. Orally inhaled zanamivir administered after confirming the diagnosis with rapid testing

4. A 7 year old presents with a high fever, respiratory distress, and a parapneumonic effusion on chest radiograph. Which of the following findings would MOST likely suggest the need for video-assisted thoroscopic surgical drainage of this effusion?
   A. A mediastinal shift away from the effusion
   B. A pleural fluid pH > 7.3 and glucose >200 mg/dL
   C. Persistent drainage for more than 5 days from a percutaneously placed thoracostentesis catheter
   D. The persistence of fever following 48 h of parenteral antibiotics
   E. The presence of loculations on ultrasound or computer tomography images

5. A 4 year old male presents with acute hypoxemic respiratory failure (PaO2/FiO2 ratio = 150), disseminated intravascular coagulation, and renal insufficiency secondary to catecholamine-resistant shock. Rapid antigen testing identifies the H1N1 virus. In addition to oral oseltamivir, the initial antimicrobial coverage should include which of the following?
   A. Cefepime
   B. Intravenous immunoglobulin
   C. Intravenous zanamivir
   D. Trimethoprim-sulfamethoxazole
   E. Vancomycin

6. A 16 year old male presents with a 3 day history of fever, chills, myalgia, headache, and gastrointestinal symptoms. On clinical exam, he is febrile, tachypneic with scattered rales, and hypotensive. There is no rash or evidence of animal bite on exam. His initial laboratory results are remarkable for thrombocytopenia, leukocytosis with an increased percentage of circulating immunoblasts, and elevated levels of lactate dehydrogenase, aspartate aminotransferase, and alanine aminotransferase. His prothrombin time is 16 s. He is admitted and his respiratory status continues to deteriorate ultimately requiring mechanical ventilation. He remains in refractory shock for several days. After an extensive diagnostic work-up, he is diagnosed with hantavirus cardiopulmonary syndrome based on the detection of hantavirus-specific immunoglobulin M. Of the following, which is most likely to be part of his medical history?
   A. An underlying immunodeficiency
   B. Being a member of the high school wrestling team
   C. Exposure to rodent excrement
   D. Intravenous drug use
   E. Residence in the Southeastern United States

7. Corticosteroids have the MOST established benefit in which of the following clinical scenarios?
   A. A 7 week old infant with severe bronchiolitis secondary to respiratory syncitial virus
   B. A 6 month old, unimmunized infant with severe hypoxia and respiratory failure secondary to pertussis
   C. A 14 year old female with necrotizing pneumonia secondary to community acquired methicillin resistant Staphylococcus aureus
   D. A 14 month old with a history of acquired immune deficiency syndrome and currently in hypoxic respiratory failure secondary to Pneumocystis jiroveci pneumonia (Pneumocystis carinii pneumonia)
   E. A 16 year old native American female with severe cardiopulmonary dysfunction secondary to a Hantavirus infection

**ANSWERS**

1. E
2. A
3. D
4. E
5. E
6. C
7. D

**SUGGESTED READINGS**

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