ATS Core Curriculum 2021. Pediatric Pulmonary Medicine:
Pulmonary Infections

Jane E. Gross
National Jewish Health

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The following is a concise review of the Pediatric Pulmonary Medicine Core reviewing pediatric pulmonary infections, diagnostic assays, and imaging techniques presented at the 2021 American Thoracic Society Core Curriculum. Molecular methods have revolutionized microbiology. We highlight the need to collect appropriate samples for detection of specific pathogens or for panels and understand the limitations of the assays. Considerable progress has been made in imaging modalities for detecting pediatric pulmonary infections. Specifically, lung ultrasound and lung magnetic resonance imaging are promising radiation-free diagnostic tools, with results comparable with their radiation-exposing counterparts, for the evaluation and management of pulmonary infections. Clinicians caring for children with pulmonary disease should ensure that patients at risk for nontuberculous mycobacteria disease are identified and receive appropriate nontuberculous mycobacteria screening, monitoring, and treatment. Children with coronavirus disease (COVID-19) typically present with mild symptoms, but some may develop severe disease. Treatment is mainly supportive care, and most patients make a full recovery. Anticipatory guidance and appropriate counseling from pediatricians on social distancing and diagnostic testing remain vital to curbing the pandemic. The pediatric immunocompromised patient is at risk for invasive and opportunistic pulmonary infections. Prompt recognition of predisposing risk factors, combined with knowledge of clinical characteristics of microbial pathogens, can assist in the diagnosis and treatment of specific bacterial, viral, or fungal diseases.

Keywords: molecular diagnostics; imaging; COVID-19; nontuberculous mycobacteria; immune compromise
DIAGNOSTIC ASSAYS
Marianne S. Muhlebach and Sara Abu-Nassar

A variety of organisms infect the airways and lung parenchyma, including bacteria, mycobacteria, viruses, and fungi. Diagnostic methods have improved greatly in recent years, yet culture remains the clinical gold standard for bacteria, mycobacteria, and fungi. Disadvantages of culture include a limited spectrum of culturable organisms, the time until results, and personnel efforts. Advantages include quantitative results and the ability to further test the organism. Besides conventional identification of organisms by growth characteristics and biochemical testing, 16S rRNA sequencing and mass spectrometry (matrix-assisted laser desorption ionization–time-of-flight mass spectrometry) have enhanced the sensitivity and accuracy of species-level identification.

Culture-independent methods typically rely on amplification of bacterial or viral nucleic acids and subsequent identification (1). Polymerase chain reaction (PCR) with or without quantitation has been available the longest, and newer, refined methods of nucleic acid extraction and amplification have streamlined the process, including isothermal nucleic acid extraction and amplification for high-throughput and point-of-care testing. Several syndromic panels for upper and lower respiratory tract infections are commercially available (2). These multiplex panels include bacterial, atypical bacterial, and viral co-detection, often with concomitant detection of bacterial resistance genes. Sensitivity and specificity vary among panels and specimen types. Sensitivity for different targets may differ within a sample, and detection of bacterial resistance in a mixed infection may not be pathogen specific (3). Increasingly, panels are semiquantitative to decrease uncertainty about contamination versus a clinically relevant bacterial load. Figure 1 demonstrates standard pathways for respiratory organism identification.

Clinicians should understand the appropriate specimen type (swab, induced or expectorated sputum, bronchoalveolar lavage fluid) relevant to the clinical question and recognize that sample quality and contamination are a few of the issues that may affect interpretation, especially for molecular diagnostics (4).

Viral Diagnostics

Viral detection is largely PCR-based, with many point-of-care methods for single or multiplex detection being available. Separate panels are optimized and Food and Drug Administration–approved for upper and lower respiratory tract samples. Antigen-based viral detection is widely used (e.g., influenza, severe acute respiratory syndrome coronavirus 2 [SARS-CoV-2]) and has rapid results but lower performance.
PCR

Gram Stain

AFB Smear

Culture

Bacterial culture up to 2 weeks
AFB culture up to 6 weeks
Fungal culture up to 8 weeks

Drug Sensitivities
1-2 days
2-3 days
4-5 days

1-4 hours

1-4 hours

NAA Test to identify Tb

If positive 1-4 hours

Results in Gram + or – and the shape

DNA and dNTPs

Antigen Antibody

Detection Area

Single Strip Multiple Strip Microarray

Sample Pad Conjugate Pad Absorption Pad

Dengue HIV Influenza Malaria

Figure 1. Pathways for respiratory organism identification. AFB = acid-fast bacillus; dNTPs = deoxyribonucleotide triphosphates; HIV = human immunodeficiency virus; NAA = nucleic acid extraction and amplification; PCR = polymerase chain reaction; Tb = tuberculosis.

Multiplex Panels test for:
1. Viral, bacterial and atypical bacteria (Chlamydia pneumoniae and Mycoplasma)
2. Antituberculosis medications

Options for Microbiology

Obtaining a Sample

Induced or Expectorate Sputum

Bronchial Sample

Nasal or Oropharyngeal Swab

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Fungal Diagnostics

Fungal diagnostics are often used in immunocompromised patients (ICPs). Culture and histopathology of secretions or tissues with fungus-specific staining remain valuable. Molecular methods include pan-fungal PCR and pathogen-specific PCR. Some syndromic lower respiratory infection panels include targets for *Aspergillus fumigatus*, *Pneumocystis jirovecii*, and *Cryptococcus neoformans* (5). To date, there are few published evaluations of those tests. Antigen detection assays are available for *Cryptococcus*, *Aspergillus* (galactomannan), *Candida* (enolase), and β-glycan as nonspecific markers of fungal infection. Given the concern of contamination from upper airway secretions, evaluation of the host’s immune response to fungus can provide specificity. Assays include complement-fixation, immunodiffusion, and enzyme immune assays.

Mycobacterial Diagnostics

Diagnosis of mycobacterial infections remains challenging, given the fastidious nature of the organisms. Culture remains the gold standard, followed by *Mycobacterium tuberculosis*–specific PCR (6). Skin and serologic tests are routinely used for tuberculosis, with cautious interpretation being used for ICPs. For nontuberculous mycobacteria (NTM), culture remains the standard in most settings; however, molecular assays and panmycobacterial PCR with gene sequencing for subtype identification are becoming available (7).

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NEW AND EVOLVING IMAGING MODALITIES

Diana Y. Chen and Nazia Hossain

In addition to clinical assessment and microbial tests, imaging studies can aid in the diagnosis and management of pulmonary infections. Although plain chest radiography (CXR) is often the preferred initial modality, findings are often normal or nonspecific. Chest computed tomography (CT) remains the gold standard for characterization of pulmonary infections but increases the total-dose radiation exposure. Lung ultrasound (LUS) and lung magnetic resonance imaging (MRI) have gained attention as radiation-free alternatives for pulmonary imaging. Video 1 shows common radiologic findings for pulmonary infections from the imaging modalities described in this review.

LUS

Although the diagnosis of pneumonia is made in a clinical context, CXR studies are often obtained, although appreciable false-negative rates and wide inter- and intraobserver interpretation variability are reported. LUS is an advantageous alternative diagnostic modality that can detect the presence of consolidations, focal B-lines, pleural line abnormalities, and effusions with significantly better sensitivity than (95.5% vs. 86.8%) and specificity similar to (95.3% vs. 98.2%) CXR alone or in combination with clinical findings, and the use of LUS results in less interpretation variability (1, 2). LUS is better for diagnosing and characterizing pleural effusions and identifying mediastinal lymphadenopathy in suspected pulmonary tuberculosis than CXR (3, 4).

LUS techniques yield quality imaging results in children because of their unique anatomical features, including a thinner chest wall and smaller thoracic width. Advantages to LUS include the lack of radiation exposure, lower relative cost, potential for expanded access in low-resource settings, and rapid availability of results (5).

Chest CT versus MRI

CT is the gold standard for detecting pulmonary infection and for diagnosing bronchiectasis and air trapping. Despite routine use of pediatric low-dose CT protocols, minimizing the cumulative dose of radiation exposure in children remains crucial, particularly in patients with DNA-repairing deficiencies and oncologic conditions. With advances in imaging quality, emerging studies have aimed to validate MRI as a radiation-free alternative.

The advancements and development of MRI with fast-imaging sequences and higher field strengths have enhanced image resolution while reducing total scan times and the need for prolonged sedation (6). In detection of consolidations, cavitary lesions, nodules, ground-glass opacities (GGOs), pleural effusions, and lymph nodes, MRI has
demonstrated high sensitivity and specificity compared with CT (6, 7). Bronchiectasis scoring by using MRI correlates significantly with pulmonary exacerbations, respiratory symptoms, and the forced expiratory volume in 1 second in patients with cystic fibrosis (CF) (8). The availability of fast MRI sequences may vary among institutions because of the need for specific knowledge, training, and expertise. Lung MRI techniques continue to develop rapidly, and multicenter standardization trials are ongoing (9).

Coronavirus disease

Coronavirus disease (COVID-19), caused by SARS-CoV-2, has rapidly resulted in over 2.5 million deaths worldwide as of February 24, 2021. CT patterns of COVID-19 in adults are well described, with bilateral GGOs appearing in up to 88% of cases (10). In contrast, children with COVID-19 are three times more likely to have normal CT findings (35.7% of children vs. 8.4–10.2% of adults). CT manifestations are also less severe and show less bilateral involvement (27.7% of children vs. 73.8–78.8% of adults). GGO remains the most common feature (37.2%), followed by consolidations or infiltrates (22.3%). As GGOs are an atypical radiologic feature of other viral infections in children, detection can aid in early identification and risk stratification (11).

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NTM Diagnosis and Management

Stacey L. Martiniano and Patricia Lenhart-Pendergrass

NTM are ubiquitous in the environment, particularly in soil and water sources. They can cause uncommon, but significant, pulmonary disease in children (1, 2). These atypical pathogens are slow growing compared with typical bacteria and require special culture conditions for proper isolation and identification (3). NTM species associated with pulmonary disease in children include the M. abscessus group (MAB) (which includes the subspecies abscessus, massiliense, and bolletii), which typically grow within 1 week in culture, as well as slower growing species such as M. avium complex (MAC) (which includes M. avium, M. intracellulare, and M. chimaera, among others) and M. kansasii, which can take weeks to grow in culture (4).

In the pediatric population, NTM infections most commonly occur in children with CF. Pulmonary NTM can also be diagnosed in children with non-CF bronchiectasis, including primary ciliary dyskinesia, autosomal dominant hyper-IgE syndrome, and primary or secondary immunodeficiencies (5). MAC and MAB are the most common species associated with pulmonary disease in the United States.

Screening and Diagnosis

Annual screening for NTM should be performed in older children with CF via an expectorated or induced sputum culture, even if they are asymptomatic. Routine screening should also be considered in children with non-CF bronchiectasis. In addition, the presence of NTM should be considered in pulmonary infections or exacerbations that are unresponsive to treatment of typical pathogens or that present with an unexpected clinical decline. Because NTM can cause transient or indolent infection, or be a contaminant, it is imperative that careful consideration of the cause and a formal diagnosis of NTM pulmonary disease is made before starting treatment. Diagnostic criteria are shown in Figure 2. If a patient is on chronic

| Clinical and Radiographic Criteria |
|-----------------------------------|
| 1. Pulmonary or systemic symptoms, and |
| 2. Nodular or cavitary opacities on chest radiograph, or a high-resolution computed tomography scan that shows bronchiectasis with small nodules, and |
| 3. Appropriate exclusion of other diagnoses |

| Microbiologic Criteria |
|------------------------|
| 1. Positive culture from at least 2 separate expectorated sputum samples, or |
| 2. Positive culture from at least one bronchial wash or lavage, or |
| 3. Transbronchial or lung biopsy with mycobacterial histopathologic features (granulomatous inflammation or acid-fast bacilli) and positive culture for NTM from biopsy, sputum, or bronchial washing |

Figure 2. Clinical and microbiologic criteria for diagnosis of nontuberculous mycobacteria pulmonary disease. Adapted from Reference 4.
azithromycin therapy for immunomodulation, it should be stopped upon isolating NTM to avoid partial treatment and induced resistance.

Management

Management of NTM disease in children requires a prolonged course of multiple antibiotics, with the treatment choice being based on the species or subspecies and guided by susceptibilities. Expert consultation is recommended, especially for drug-resistant MAC or MAB (4). Standard treatment approaches are shown in Table 1 for the most common NTM (4, 6). Newer antibiotics that show in vitro activity against NTM are included on the basis of updated treatment guidelines (6). Typically, MAC and M. kansasii can be treated with an oral antibiotic regimen. MAB requires an intensive phase of intravenous and oral antibiotics, which is followed by a continuation phase of oral and inhaled antibiotics. Daily treatment is recommended in children (vs. thrice-weekly treatment in adults) because of drug metabolism. Any patient started on NTM treatment requires frequent follow-up with intensive drug-toxicity monitoring (e.g., audiograms, vision testing, electrocardiography, and laboratory testing). The goal of treatment is to achieve 12 months of consecutive negative culture results. If microbiologic conversion cannot be achieved, drug concentrations should be measured and alternate antibiotics can be trialed.

| Organism          | Antimicrobial Regimens                                                                 |
|-------------------|----------------------------------------------------------------------------------------|
| **M. avium complex** | • Oral azithromycin or clarithromycin, rifampin or rifabutin,* and ethambutol          |
|                   | • For severe disease or if resistance is suspected, addition of IV or inhaled amikacin or clofazimine should be considered. |
| **M. abscessus group** |                                                                                     |
| Intensive phase   | • IV amikacin plus IV imipenem, cefoxitin, or tigecycline                               |
|                   | • Oral azithromycin or clarithromycin*                                                |
|                   | • 1 or 2 of the following: oral clofazimine, omadacycline, tedizolid, linezolid, bedaquiline, or moxifloxacin |
| Continuation phase| • Inhaled amikacin                                                                     |
|                   | • Oral azithromycin or clarithromycin†                                                 |
|                   | • 1 or 2 of the following: oral clofazimine, omadacycline, tedizolid, linezolid, bedaquiline, or moxifloxacin |
| **M. kansasii**   | Oral rifampin, isoniazid or macrolide, and ethambutol                                   |

*Additional antimycobacterial agent needed if acquired or inducible resistance is present.

Definition of abbreviations: IV = intravenous; M. = Mycobacterium.

Data are from References 4 and 6.

If a patient cannot receive rifamycin because of a drug–drug interaction (e.g., ivacaftor), then an alternate antibiotic (e.g., clofazimine) should be used.

*Additional antimycobacterial agent needed if acquired or inducible resistance is present.
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PEDIATRIC COVID-19
Ruobing Wang and Timothy Klouda

Since its identification in December 2019, SARS-CoV-2 has infected 20 million people in the United States, with an estimated 12% of cases occurring in children (1). The incidence of pediatric COVID-19 is likely underestimated because the majority of children with COVID-19 have mild or no symptoms, resulting in reduced disease screening and reporting. Despite children having better outcomes, they may contribute to transmission and are at risk for severe infection. Moreover, the pandemic has affected the lives of children because of disrupted education, decreased access to medical care, and increased mental health disorders (2).

Pathogenesis
The SARS-CoV-2 virus gains entry into cells by binding to the ACE2 receptor (3). The reason children experience less severe symptoms is unknown, and hypotheses include children having lower levels of ACE2, differential immune responses, and fewer comorbidities (4). To date, only age (less than 1 yr old) and having a preexisting medical condition have been identified as risk factors for severe disease in children (5).

Clinical Presentation
Children typically have mild symptoms and a fast recovery, with 17.4–19.3% of children with COVID-19 being asymptomatic (6, 7). The incubation period is estimated to be 2–14 days, and symptomatic patients typically recover in 2–4 weeks (8). Common symptoms include fever (51.2–59.1%), cough (37.0–55.9%), rhinorrhea (9.9–20.0%), and pharyngitis (8.3–18.2%) (6, 7).

Diagnostic Testing
Diagnosis of COVID-19 is typically through detection of viral mRNA from a respiratory sample with reverse transcription–PCR. The accuracy of testing results depends on numerous factors, such as the sample source and the viral load present. Consultation with an infectious disease expert is recommended in certain clinical scenarios (e.g., congenital or acquired immunodeficiency) to ensure proper test interpretation and determine the need for possible retesting (8). Although laboratory findings have limited diagnostic and prognostic value, lymphopenia as well as elevated creatinine kinase, liver enzymes, and procalcitonin have been documented (7). CT abnormalities can present before clinical symptoms and may include GGOs and consolidations (6, 7).

Treatment and Management
Currently, there are no specific therapies for children with COVID-19. Stable patients are advised to quarantine at home (8). Treatment strategies for hospitalized patients include supplemental oxygen, fluid resuscitation, and empiric antibiotics when indicated (8). Systemic corticosteroids are not recommended. Patients in whom supplemental oxygen fails should be escalated to noninvasive positive pressure ventilation before mechanical ventilation with lung-protective strategies (9). Monoclonal antibodies to SARS-CoV-2 can be administered to high-risk pediatric patients 12 years or older who test positive for SARS-CoV-2 (10). To date, the Pfizer–BioNTech COVID-19 mRNA vaccine is the only vaccine available to patients 12 years or older. Clinical trials for patients under 12 years of age are currently ongoing.

Complications and Prognosis
An estimated 13.3% of pediatric patients with COVID-19 are admitted to the hospital, with 3.5% requiring intensive care unit (ICU) care. Mortality is low and is estimated to be less than 1% (5). Multisystemic inflammatory syndrome in children is a rare complication defined by fever, systemic inflammation, and
multiorgan dysfunction seen after COVID-19 infection in patients 3–12 years old. There are no established guidelines, and treatment includes intravenous immunoglobulin and glucocorticoids. With multisystemic inflammatory syndrome in children, 80% of patients require ICU care, and mortality is estimated at 2% (11).

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INFECTIONS IN THE IMMUNOCOMPROMISED HOST

Stephen Kirkby and Robin Ortenberg

The pediatric ICP is at risk for invasive and opportunistic pulmonary infections. Prompt recognition of predisposing risk factors, combined with knowledge of clinical characteristics of microbial pathogens, can assist in the diagnosis and treatment of specific bacterial, viral, or fungal diseases.

Classification of Patients at Risk for Infection

There are hundreds of primary immunodeficiencies, further characterized by specific cellular defects (1). Disorders affecting T cells are likely to present with recurrent or severe viral and fungal infections. B-cell abnormalities lead to decreased antibody production and recurrent bacterial pneumonias, specifically caused by encapsulated organisms (2). Secondary immunodeficiencies vary in severity depending on the underlying cause of disease and include diabetes, malignancy, malnutrition, sickle-cell disease, human immunodeficiency virus, and immunomodulatory medication use. In addition, significant immunosuppressive states exist related to solid organ or bone marrow transplantation (3).

Diagnostic Evaluation

Evaluation of the ICP with respiratory symptoms should aim to quickly identify the source organism and rule out noninfectious etiologies. Imaging with CXR or CT can assist in identifying pathognomonic patterns of disease, in conjunction with specific laboratory studies. Bronchoscopy with bronchoalveolar lavage remains a key tool for diagnosis through cultures, cytology, and molecular diagnostic testing (such as PCR) (4). There may also be a limited role for lung biopsy (transbronchial or radiology-guided biopsy) or sampling of other tissues, such as lymph nodes or skin (Table 2). Infectious disease consultants can offer valuable assistance in specific testing, particularly in complicated cases.

Bacterial Infections

Bacterial infections are commonly encountered among ICPs and may be acquired in the community or healthcare setting. Clinicians should have high clinical suspicion and quickly institute appropriate broad antibiotic coverage for gram-positive and gram-negative species, as the typical treatment for community-acquired pneumonia in immunocompetent hosts may be insufficient (5). Slow-growing bacterial species including mycobacteria and Nocardia species may also be identified in ICPs. Practitioners can tailor treatment to a specific pathogen once cultures and drug susceptibility patterns are identified.

Fungal infections

Although many fungi represent commensal or nonpathologic organisms, others can cause true infection in the ICP. P. jirovecii is a common infection among organ transplant recipients as well as patients with chronic systemic glucocorticoid therapy or prolonged neutropenia or lymphopenia. Prophylactic strategies are often used in these high-risk patients. Aspergillus species, including A. fumigatus, may cause invasive pulmonary aspergillosis with sequelae such as necrotizing pneumonia, vascular invasion, and hematologic spread. These patients may present with hemoptysis or pulmonary infarction. The classic radiologic finding in invasive pulmonary aspergillosis is a cavitary lesion with surrounding GGO, the so-called “halo sign.” The highest-risk patients include those with hematologic malignancies and bone marrow transplant recipients. Other important fungal pathogens to consider in ICPs include mucormycosis, cryptococcosis, and Candida.
Table 2. Characteristics of selected pulmonary infections in immunocompromised patients

| Organism                                      | Associated Immune Disorders and Risk Factors | Clinical Presentation | Characteristics of Diagnosis                                                                 |
|-----------------------------------------------|---------------------------------------------|-----------------------|-------------------------------------------------------------------------------------------------|
| *Aspergillus* species, specifically *A. fumigatus* (fungus) | • Neutropenia  
• Hypo- or agammaglobulinemia  
• Immunosuppressive therapy  
• BMT | • Fever  
• Cough  
• Chest pain  
• Throat pain | • Imaging: ground-glass opacities; cavitary nodules present with invasive disease  
• Serum testing: galactomannan; *Aspergillus* PCR  
• Bronchoscopy: tracheobronchitis with mucosal plaques and/or ulceration  
• BAL: fungal culture*; galactomannan: some studies show low sensitivity and possibility of cross-reactivity with other fungal infections; *Aspergillus* PCR: does not distinguish colonization from active infection  
• Biopsy of nodule/tissue: septate hyphae with acute branching pattern  
• Elevated β-1,3-D-glucan in serum and/or BAL  
• Serum antigen testing  

| *C. neoformans* (fungus) | • HIV  
• Congenital T-cell disorders  
• Immunosuppressive therapy | • Cough  
• Fever  
• Lymphadenopathy  
• Neurologic disease: headaches; meningismus | • Imaging: diffuse interstitial findings; pulmonary nodules  
• BAL: India-ink stain reveals halo sign of polysaccharide capsule*; antigen testing  
• Low β-1,3-D-glucan in serum and/or BAL |
Table 2. Continued.

| Organism | Associated Immune Disorders and Risk Factors | Clinical Presentation | Characteristics of Diagnosis |
|----------|---------------------------------------------|-----------------------|------------------------------|
| P. jirovecii (fungus) | hypo- or agammaglobulinemia | Dyspnea | Imaging: bilateral infiltrates; ground-glass or cystic lesions; nodules, with or without cavitation |
| | congenital T-cell disorders | Fever | BAL or sputum: direct identification of organism*; tinctorial staining; fluorescent antibody staining; PCR |
| | HIV | Cough | Elevated β-1,3-β-D-glucan levels in serum and/or BAL |
| | post-BMT | cyanosis (late) | |
| | immunosuppressive therapy | hypoxic respiratory failure (late) | |
| Candida species (fungus) | neutropenia | commonly presents with systemic findings due to hematogenous spread | Imaging: microabscesses |
| | congenital T-cell disorders | white plaques on buccal mucosa | Bronchoscopy: evidence of oropharyngeal plaques |
| | post-BMT | | Blood culture |
| | chronic illness | | BAL: Gram stain or potassium hydroxide preparation; budding yeasts with or without pseudohyphae |
| | | | Elevated β-1,3-β-D-glucan (Fungitell) levels in serum and/or BAL |
| CMV (virus) | post-BMT | fever | BAL: histopathology "owl-eye" inclusions or positive CMV-specific immunohistochemistry*; quantitative PCR; positive culture result |
| | HIV | malaise | Serum: quantitative viral load; whole-blood PCR |
| | neonates | cough | |
| Organism                  | Associated Immune Disorders and Risk Factors | Clinical Presentation                                                                 | Characteristics of Diagnosis                      |
|--------------------------|---------------------------------------------|----------------------------------------------------------------------------------------|----------------------------------------------------|
| *L. monocytogenes* (bacteria) | • Neonates                                  | • Fevers                                                                               | • Radiology: ground-glass opacities                |
|                          | • HIV                                       | • Chills                                                                                | • Serum and CSF: Gram stain and culture*; PCR*     |
|                          | • Post-BMT status                           | • Can present initially with septicemia                                                 | • BAL usually low-yield                            |
|                          | • Chronic gastrointestinal disease          | • Neurologic symptoms: meningismus; seizures; cranial nerve abnormalities; encephalitis |                                                    |

**Table 2. Continued.**

Definition of abbreviations: *A. fumigatus* = *Aspergillus fumigatus*; *BAL* = bronchoalveolar lavage; *BMT* = bone marrow transplantation; *C. neoformans* = *Cryptococcus neoformans*; *CMV* = cytomegalovirus; *CSF* = cerebrospinal fluid; *HIV* = human immunodeficiency virus; *L. monocytogenes* = *Listeria monocytogenes*; *P. jiroveci* = *Pneumocystis jiroveci*; *PCR* = polymerase chain reaction.

*Gold standard.
infections. Published guidelines detail the diagnosis of pulmonary fungal infections (6).

**Viral Infections**

Seasonal community-acquired viral infections, including with influenza, respiratory syncytial virus, adenovirus, and other common upper respiratory tract viruses, are more likely to cause clinical pneumonia in pediatric ICPs. A high level of suspicion and prompt diagnostic testing via PCR panels are indicated. Solid organ and bone marrow transplant recipients are at particularly high risk for cytomegalovirus, which can cause pneumonitis and other organ disease. Risk is based in part on the seropositivity of the donor and recipient, and prophylactic strategies are often employed (7).

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