# Eosinophilic pneumonias

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Eosinophilic Pneumonias

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INTRODUCTION

The terms “eosinophilic pneumonia” and “pulmonary eosinophilia” loosely encompass a broad range of infectious and noninfectious pulmonary conditions that involve infiltration of eosinophils into the lungs, often with accompanying peripheral blood eosinophilia. Conditions in which airway and peripheral eosinophilias are present in the absence of parenchymal infiltration and radiographic changes, such as allergic asthma, are not included in this broad categorization. This review starts with discussions of several infectious causes of eosinophilic pneumonia, which are almost exclusively parasitic in nature. Inclusion of these parasitic etiologies in the differential diagnosis of pulmonary eosinophilia depends highly on epidemiologic considerations that make exposure to the offending organisms more likely. Pulmonary infections due specifically to Ascaris, hookworms, Strongyloides, Paragonimus, filarial nematodes, and Toxocara are considered in detail. The discussion then moves to noninfectious causes of eosinophilic pulmonary infiltration, including allergic sensitization to Aspergillus, acute and chronic eosinophilic pneumonias (AEP and CEP, respectively), Churg-Strauss syndrome (CSS), hypereosinophilic syndromes (HES), and pulmonary eosinophilia due to exposure to specific medications or toxins.

Eosinophil immunobiology is complex and has been reviewed elsewhere (2, 155). However, certain considerations warrant particular mention in a discussion of pulmonary eosinophilia. Eosinophils are integral components of Th2 inflammation (155), a prominent feature of most of the conditions discussed in this review. The presence of the cytokine interleukin-5 (IL-5) in particular is central to eosinophilia, promoting the differentiation, survival, and migration of eosinophils (31, 162, 200, 216). Also important as a common pathophysiologic mechanism of pulmonary tissue damage in the eosinophilic pneumonias is the release of cytotoxic, granule-stored cationic proteins by eosinophils (155). Eosinophils have intricate immunoregulatory functions that are likely important in orchestrating the perpetuation of Th2 inflammation in the eosinophilic pneumonias.

INFECTION CAUSES

Ascaris

Pulmonary eosinophilia due to the passage of helminth larvae through the lungs is referred to as Löffler’s syndrome, though the term is often used more generally to refer to simple pulmonary eosinophilia from any fungal, parasitic, or drug-induced cause. In the original case series by Löffler, most of the cases described were due to Ascaris infections (98). Human pulmonary eosinophilia caused by Ascaris has been described for Ascaris lumbricoides (98) and Ascaris suum (144).

Eosinophilic pulmonary infiltrates and respiratory symptoms due to Ascaris are generally part of a self-limited process due to the transient nature of larval passage through the lungs in the Ascaris life cycle (38). Though many patients remain asymptomatic, 8 to 15% of infected individuals display morbidity, with respiratory symptoms occurring 9 to 12 days after the ingestion of eggs and lasting 5 to 10 days (39, 211). The severity of the clinical syndrome has been noted to correlate with the extent of worm burden (39). Ingestion of A. lumbricoides eggs may occur due to the use of...
human feces contaminated with *Ascaris* eggs as fertilizer, as described in Löfler’s case series (38, 98, 211). Infection with *A. suum* may take place when there is close domestic proximity between pigs and humans (38), though a prominent case series of Löfler’s syndrome due to *A. suum* occurred in patients whose food was maliciously contaminated (144). Symptoms may include cough, dyspnea, wheeze, and hemoptysis and may progress to frank respiratory distress (98, 144). Peripheral eosinophilia (i.e., absolute eosinophil count of ≥500/µl) is often present at the onset of symptoms, but the peak level of eosinophilia will have a delay of several days from presentation (144, 211). If necessary, definitive diagnosis can be made by recovery of *Ascaris* larvae from either respiratory secretions or gastric lavage fluid (144, 211). Eggs will not be detectable in the stool at this early stage in the nematode life cycle. The radiographic pattern found at the time of respiratory symptoms is characterized by patchy areas of consolidation that may coalesce in more severe cases (45). Because pulmonary eosinophilia due to *Ascaris* is self-limiting, specific therapy at the time of symptoms is generally not necessary (38). Corticosteroids may be administered in particularly severe cases and have been observed to result in favorable symptomatic responses (48, 144). Therapy aimed at eradication of adult worms with albendazole or mebendazole should be delayed until after pulmonary symptoms have resolved and organisms are expected to have entered the adult phase (38).

*A. suum* infection has been reported to result in a visceral larval migrans (VLM) pattern of pulmonary eosinophilia (107, 133, 160). However, whether these cases are truly representative of *Ascaris*-associated VLM, with liver and lung involvement, rather than either VLM due to *Toxocara* or misclassified Löfler’s syndrome, remains an unresolved issue (142).

**Hookworm**

Though such cases are uncommon, Löfler’s syndrome may occur after infection with *Ancylostoma duodenale* or *Necator americanus* as larvae migrate through the lungs (10, 45, 61, 96, 163). Initial infection occurs exclusively by the percutaneous route in the case of *N. americanus* and by either the oral or percutaneous route in the case of *A. duodenale* (61). Respiratory symptoms are usually mild and occur several days after the onset of infection (207). As with Löfler’s syndrome due to *Ascaris*, pulmonary symptoms, which include wheeze, cough, and hemoptysis, are accompanied by transient infiltrates on chest imaging and peripheral blood eosinophilia (45). In experimental low-dose *N. americanus* infection of human volunteers, bronchoscopy performed at the time that larvae were expected to be transiting through the lung demonstrated erythema of the airways (111). Also in common with *Ascaris* is the necessity of isolation of hookworm larvae in respiratory or upper gastrointestinal (GI) secretions in order to definitively make the diagnosis, as isolation of hookworm eggs from the stool will not occur for several months after the transient pulmonary component of the syndrome (207). Treatment with a benzimidazole agent is necessary for eradication of adult worms but will not affect the potential pulmonary component of hookworm infection. Corticosteroids may be used if Löfler’s syndrome results in severe respiratory symptoms (207). Löfler’s syndrome associated with cutaneous larval migrans secondary to animal hookworms, such as *Ancylostoma braziliense*, has also been reported but is rare (35, 141, 183, 214).

Notably, experimental hookworm infection in rodents has proven to be a valuable tool in the study of Th2 inflammation and eosinophil recruitment to the lungs, taking advantage of the pulmonary transmigration component of the hookworm life cycle (2, 32). Specific to eosinophilic pneumonias is the observation of increased eosinophil chemoattractant production in the lungs coinciding with the presence of larvae in mice infected with *N. americanus* (32).

**Strongyloides**

While initial *Strongyloides stercoralis* infection by filariform larvae through a transcutaneous route may result in Löfler’s syndrome, this is not a common pulmonary manifestation of strongyloidiasis (175, 207, 211). However, *Strongyloides* may cause pulmonary symptoms and infiltrates as a manifestation of chronic infection or as a result of hyperinfection in immunocompromised hosts (207). The unique life cycle of *S. stercoralis* allows a chronic infection of extended duration to occur due to the ability of new filariform larvae to continuously autoinfect the human host through the perianal skin or bowel mucosa (175). Patients with chronic infection may have repeated episodes of fever and pneumonitis that may be mistaken for recurrent bacterial pneumonias (207). Eosinophilia, though often absent during the acute episodes, may occur during the intervening period between episodes (207). Pulmonary involvement of strongyloidiasis has been reported as an asthma mimic (128).

Serious, potentially fatal pulmonary infections may occur in the context of the hyperinfection syndrome, resulting from heavy parasite burdens, most often in immunocompromised hosts with deficiencies in cell-mediated immunity (49, 79, 97, 116, 148). Symptoms may include dyspnea, cough, pleuritic chest pain, and hemoptysis, accompanied by infiltrates of varying character on chest imaging and often accompanied by a mild blood eosinophilia, unless this is suppressed by concomitant use of corticosteroids or bacterial coinfection (79, 207, 211, 212). Respiratory failure may occur in severe cases (79). Lung abscesses and pleural effusions have been reported as manifestations of pulmonary strongyloidiasis (44, 212). Use of corticosteroids, even for short durations, for underlying chronic lung disease or other indications is a risk factor for the development of hyperinfection syndrome with pulmonary involvement (25, 44, 49, 69, 194). Though rare, pulmonary strongyloidiasis is a potential complication of HIV/AIDS (43, 51, 97, 101). Several other factors, such as hematologic malignancy, immunosuppressive drug therapy, solid organ transplantation, hematopoietic cell transplantation, human T-cell leukemia virus type 1 (HTLV-1) infection, hypogammaglobulinemia, and malnutrition, have all also been associated with *Strongyloides* hyperinfection syndrome (79, 164, 197).

The diagnosis of strongyloidiasis, when suspected in patients with a compatible pulmonary syndrome, can be made by detection of larvae from stool (207). However, because stool studies are often negative due to the small number of larvae usually passed in the stool, aspiration of duodenojejunal fluid and enzyme-linked immunosorbent assay (ELISA) serologic testing may be used as adjunct diagnostic tools (79, 175). Rhabditiform larvae may also be detected in bronchoalveolar lavage (BAL) fluid samples or in sputum examinations, establishing the diagnosis (58, 210). Ivermectin therapy is the treatment of choice, and therapy may need to be extended beyond the usual 2 days to at least 7 days in cases of hyperinfection syndrome (175).
Eosinophilic pulmonary infections are caused by several species of *Paragonimus*, the lung fluke (207). *Paragonimus* infections are generally acquired by the ingestion of raw or undercooked seafood, in particular crabs and crayfish, which serve as secondary hosts in the life cycle of the organism (100). *Paragonimus westermani*, which is endemic to East and Southeast Asia, is the best characterized of the eight species of lung fluke that are thought to lead to human disease (100, 207). It should be noted that *Paragonimus* infections are not limited to East and Southeast Asia but occur throughout the world, in particular in Central and South America (*Paragonimus mexicanus*) and West Africa (*Paragonimus africanus*) (100). While *P. westermani* has been reported for rare infections contracted in the United States (14), most reports of paragonimiasis in the United States have been due to *Paragonimus kellicotti* (91, 106, 140).

Once metacercaria-stage parasites are ingested, they form exocysts in the duodenum that migrate into the peritoneum and through the diaphragm into the lungs (13). As larvae migrate through the upper GI tract and peritoneum, they may initially cause symptoms of abdominal pain and diarrhea after an incubation period of 2 to 15 days (100, 207). As organisms penetrate the diaphragm, pleuritic chest pain may develop, with the potential for pneumothorax or pleural effusion (13, 70). Pleural effusions caused by *Paragonimus* are commonly eosinophilic in nature (185). Though the lungs are usually the destination of migrating larvae, paragonimiasis may involve the brain or skin (100). As larvae continue to migrate within the lung parenchyma, symptoms of cough, low-grade fever, and blood-streaked sputum may occur, accompanied by infiltrates on chest imaging and prominent peripheral blood eosinophilia (10 to 30% of circulating leukocytes) (13, 207). The infection then progresses to a chronic phase within the lung, which may last up to several years, characterized by chronic cough and recurrent hemoptysis with resolution of the previously seen blood eosinophilia (70, 207). During this phase, larvae mature into mature flukes surrounded by a fibrous capsule that forms after the initial period of neutrophilic and eosinophilic inflammation (100). Eosinophil degranulation caused by secreted parasitic proteins may contribute to local inflammation and subsequent fibrous cyst formation (26). The mature flukes produce eggs that are coughed and expectorated or swallowed after rupture of cysts into communicating airways (45).

Chest X-rays may show no abnormality for approximately 20% of infected patients (45, 130). When abnormalities are noted on chest X-ray or computed tomography (CT) scan, they are varied in nature, often consisting of parenchymal opacities that may be hazy and ill-defined in nature or more nodular (72, 130, 217). Thin-walled cysts may be noted, as well as linear opacities corresponding to burrowing organisms (63, 64, 83, 176).

The diagnosis of *Paragonimus* infection is made by identification of eggs in sputum, stool, or gastric aspirates (100). Praziquantel is generally the preferred drug for treatment (71).

**TFPE**

Infections with the filarial nematodes *Wuchereria bancrofti*, *Brugia malayi*, and *Brugia timori* lead to lymphatic filariasis, a manifestation of which is a distinct eosinophilic pulmonary process known as tropical filarial pulmonary eosinophilia (TFPE) (138). True to its name, TFPE occurs in individuals from tropical regions, including but not limited to the Indian subcontinent, Southeast Asia, parts of Africa, and parts of China, or in individuals who have traveled to these regions (126). Humans are infected by third-stage larvae after a bite from a mosquito taking a blood meal. Larvae then mature to male and female adult worms, which mate to produce microfilariae. These are then ingested by mosquitoes taking blood meals, followed by subsequent maturation to third-stage larvae, completing the life cycle of the organism (126). The pathogenesis of TFPE has not been clarified fully, but a strong eosinophilic immunologic response to microfilariae trapped in the lungs likely plays a pivotal role (129, 145, 202). Interestingly, only a small percentage of patients affected with lymphatic filariasis develop pulmonary manifestations, and there is a strong male predominance in developing TFPE (126, 207). The clinical presentation of TFPE is characterized by the gradual onset of paroxysmal asthma-like symptoms and nonproductive cough. Constitutional symptoms of weight loss and fatigue are commonly present as well (126). Chest imaging in TFPE demonstrates reticulonodular opacities with nodules of 1 to 3 mm (Fig. 1) (45, 211). Pulmonary function testing can show both restrictive and obstructive patterns of abnormality (193).

In addition to the clinical presentation and radiographic findings, diagnosis is aided by the presence of a peripheral eosinophilia, which may be high grade and is generally >3,000/microliter. Eosinophils are also present in BAL fluid specimens (145). Serum IgE is elevated in TFPE (127). While not specific for the type of filarial infection, serologic antibody testing can confirm the diagnosis (90). ELISA testing for antigen from *W. bancrofti* is also available (117).

Though not needed for diagnosis, pathology specimens of the lung during TFPE are characterized initially by eosinophilic infiltration that progresses to eosinophilic abscesses and eosinophilic granulomas. As the disease progresses unchecked, fibrosis may develop. Microfilarial fragments are occasionally identifiable on lung specimens (193).

Patients with TFPE are treated with diethylcarbamazine, with the option of using doxycycline as an adjunct therapy against adult worms. Relapses are also treated with the same regimen (137, 152, 187).

**VL M**

VL M is a syndrome due to infection by the dog ascarid, *Toxocara canis*, or the cat ascarid, *Toxocara cati*, that commonly includes pulmonary eosinophilia (9). Humans are accidental hosts for *Toxo-
TABLE 1 Infectious causes of eosinophilic pneumonia

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cara; larvae cannot mature into adults in humans and migrate through tissues, causing disease (36, 207). VLM is predominately a disease of children between the ages of 1 and 5 years, as the infection is usually acquired through the ingestion of embryonated eggs in soil contaminated with dog feces in areas such as public playgrounds (62). Infections are not generally transmitted directly from dogs or cats, as eggs shed from these animals require approximately 3 weeks in the soil before they become infectious (207).

The clinical syndrome observed from *Toxocara* infection commonly affects the liver and the lungs, though other tissues, such as the central nervous system and the eyes (ocular larval migrans), may be affected as well (16, 36, 113). The pulmonary manifestations of VLM are a result of damage from the penetration of larvae into the lungs and from the immunological response to the larvae (78). While some individuals manifest only a peripheral blood eosinophilia, pulmonary symptoms include dyspnea, wheeze, and cough (9, 16, 36). Symptoms may rarely be severe in nature (8). Chest imaging, when abnormal, demonstrates ill-defined airspace opacities that on CT scan are subpleural in location, with a ground-glass or nodular appearance (45, 160). Anti-A or anti-B isohemagglutinin titers are elevated in 50% of patients with VLM (207). Liver involvement may lead to liver function test abnormalities. ELISA testing available from the U.S. Centers for Disease Control and Prevention can assist in establishing the diagnosis. ELISA testing available from the U.S. Centers for Disease Control and Prevention can assist in establishing the diagnosis. ELISA testing available from the U.S. Centers for Disease Control and Prevention can assist in establishing the diagnosis.

**NONINFECTIOUS CAUSES**

**ABPA**

Allergic sensitization to *Aspergillus* species may lead to allergic bronchopulmonary aspergillosis (ABPA), a well-characterized eosinophilic pulmonary condition, with the development of asthmatic symptoms, pulmonary infiltrates, central bronchiectasis, and elevated IgE levels (1, 54, 191, 220). Though ABPA is recognized in general to be due to allergic mechanisms in response to the inhalation of *Aspergillus* rather than due to frank infection, the pathophysiology of the condition is not completely understood. However, Th2 inflammation is a key component of the inflammatory response to *Aspergillus* spores, with *Aspergillus* antigen-responsive circulating Th2 cells being identifiable in ABPA patients (22, 88). T cell clones specific to the *Aspergillus fumigatus* Asp f 1 antigen established from patients with ABPA are predominately of the Th2 phenotype (22). The increase in Th2 inflammation accounts for the observed eosinophilia and elevated IgE levels. The costimulatory molecule OX40 ligand has been found to be important in Th2 inflammatory responses in patients with ABPA associated with cystic fibrosis, with increased Th2 activity inversely correlating with circulating vitamin D levels (88). Proteolytic enzymes and mycotoxins from *Aspergillus* in colonized airways, along with eosinophilic inflammation and neutrophilic inflammation, may cause the characteristic central bronchiectasis seen in many cases of ABPA (50).

The clinical features of ABPA are dominated by asthma symptoms, often with a significant component of cough and expectoration of brownish mucous plugs (1). Peripheral eosinophilia is common, a characteristic that makes it a prominent component of the differential diagnosis for pulmonary eosinophilia in the correct clinical context (18, 21, 154). Chest X-rays and CT scans of patients with ABPA may classically show patchy infiltrates, bronchiectasis, and evidence of mucous impaction (1, 17, 122). Central bronchiectasis is associated with ABPA, though its presence on CT scan is not highly sensitive for a diagnosis of ABPA (149). Pulmonary function testing usually shows an obstructive pattern for patients with ABPA (102, 124). Though pathology is not necessary for diagnosis, findings are variable, with eosinophilic inflammation, mucoid impaction, and bronchocentric granulomatosis (20, 151, 178). Noninvasive, septated hyphae may be observed in the distal airways as well.

The diagnosis of ABPA is based not on a single test result but on a constellation of clinical, radiographic, immunologic, and serologic findings based on the known pathophysiology and presentation of the disease (1, 54). A diagnosis of ABPA may be made if four of the following major features are present: a history of asthma, positive *Aspergillus* skin test reactivity, precipitating antibodies to *A. fumigatus*, elevated total IgE level (>417 IU/ml), elevated circulating levels of specific IgE and IgG, peripheral eosinophilia, central bronchiectasis on chest imaging, and pulmonary...
infiltrates (54). An evaluation for ABPA is often undertaken for patients with difficult-to-treat asthma to find an underlying cause for resistance to therapy. ABPA may also occur as a complicating pulmonary condition in a substantial subset of patients with cystic fibrosis and is suspected in those patients whose exacerbations of respiratory symptoms respond poorly to antibiotic therapy (181).

Treatment for acute or recurrent flares of ABPA consists of systemic glucocorticoids as a first-line therapy, with tapering over a 3- to 6-month period (198). Antifungal therapy with agents effective against Aspergillus species, such as itraconazole or voriconazole, may be used as adjunct therapy targeted at reducing the antigenic stimulus driving ABPA activity (42, 182, 199). The use of antifungal agents has proven effective in improving symptoms and lowering the dose of glucocorticoids in ABPA patients. Omalizumab, a monoclonal antibody against IgE, has also been described in case reports and series as an effective agent in treating ABPA, particularly in those patients with ABPA associated with cystic fibrosis (94, 190, 196).

Several other fungi have been reported to cause a syndrome similar to ABPA (1). However, these are rare compared to the more common ABPA, which is present in a significant subset of patients with asthma (55, 165).

Drugs/Toxins

A host of medications and toxins, both inhaled and ingested, have been associated with the development of pulmonary eosinophilia (179). The most commonly implicated classes of medications are the nonsteroidal anti-inflammatory medications and antibiotics. While several classes of antibiotics are associated with the development of pulmonary eosinophilia, nitrofurantoin in particular has frequently been described as a culprit. Though these medications are the most commonly associated triggers, the list of medications causing pulmonary eosinophilia is lengthy, and all medications should be reviewed carefully whenever eosinophilia with pulmonary infiltrates occurs. Furthermore, environmental exposures, such as particulate metal exposure and scorpion stings, as well as inhalational drugs of abuse, such as crack cocaine, may lead to eosinophilic pulmonary infiltrates (119, 166, 167). The presentation of affected patients may vary, from no symptoms to mild respiratory symptoms and cough to a presentation resembling acute eosinophilic pneumonia in severe cases. Treatment consists of removal of the offending medication or toxin, along with glucocorticoid therapy if warranted by the degree of symptoms (179).

CEP

Chronic eosinophilic pneumonia (CEP) is an idiopathic condition not due to a known infectious or toxic etiology in which eosinophils infiltrate the pulmonary parenchyma and cause symptoms of dyspnea, cough, and hypoxemia of varying severity. Carrington and colleagues first reported CEP in 1969 for a series of 9 female patients presenting with dyspnea, fevers, and weight loss, along with peripheral opacities on chest X-rays and eosinophilic airspace infiltration on lung biopsy specimens (19). These patients had a robust, favorable clinical response to corticosteroid therapy. This first description of CEP remains quite consistent with the current understanding of this syndrome.

The pathophysiology of CEP remains incompletely delineated, though elevations of several cytokine, chemokine, and immuno-modulatory products in studies of BAL fluid obtained from patients with CEP suggest that recruitment of eosinophils into the lung airspaces is a multifactorial process. BAL fluid in CEP contains the cytokines IL-4, IL-5, IL-6, IL-10, IL-13, and IL-18, which is largely consistent with local Th2 inflammation, though the Th1 cytokines IL-2 and IL-12 are present as well (76, 85, 114). Several chemokines are also present in BAL fluid samples, including CXCL17 and CCL22, the chemokine ligands of the receptor CCR4, suggesting that T cell recruitment may be an important factor in the pathophysiology of CEP (74, 114). In addition, CXCL9, CXCL10, CCL5 (RANTES), CCL11 (eotaxin-1), CCL2, and CCL4 have also been observed to be elevated in CEP (73, 75, 89, 186). Other immune mediators, such as tryptase, prostataglandins, and soluble ADAM8, have been detected to be elevated in the BAL fluid of CEP patients as well (7, 109, 132). BAL fluid eosinophils in CEP are activated, with expression of HLA-DR, and have longer survival than their counterparts in the peripheral blood, with decreased apoptosis (11, 159). There is evidence that eosinophils in CEP have locally augmented secretion of their cytotoxic, cationic granule-stored contents, with detection of increased major basic protein (MBP) and eosinophil cationic protein (ECP) in lung-derived specimens (15, 53, 169). Eosinophil derived neurotoxin (EDN) has been detected in high levels in the urine of patients with CEP (37, 136). Electron microscopy studies have demonstrated lysed eosinophils, degranulation, and the uptake of granule-proteins by alveolar macrophages, as well as the presence of cell-free granules (52, 66). T cells also have a likely role in CEP, with increases in activated CD4+ T helper cells present in the alveoli, as well as observations of oligoclonal T cell populations in the BAL fluid (118, 170).

CEP is characterized by respiratory and systemic symptoms in the presence of both lung and blood eosinophilia (203). There is a 2:1 female-to-male predominance of CEP, and the disease often occurs in patients with asthma (104, 105). The onset of symptoms is indolent and most commonly features cough, dyspnea, malaise, fever, and weight loss (105). While a stringent set of diagnostic criteria have not been established for CEP, peripheral blood eosinophilia is usually present, though not always. The presence of eosinophils in BAL fluid is necessary to establish the diagnosis, with eosinophilia exceeding 40% of the total cell count observed in more than 80% of cases (105). Elevations in IgE levels, erythrocyte sedimentation rate, and C-reactive protein may be present as well (105). The original case series of CEP, along with a follow-up radiographic case series, described the chest X-rays of CEP patients as displaying peripheral opacities that are the “photographic negative” of pulmonary edema, which has endured as an identifying characteristic of the disease (19, 121), though the lack of this pattern does not exclude CEP (68, 105). CT imaging of the chest most often shows patchy airspace consolidations with a peripheral distribution, though patterns may vary (46, 105, 112, 115). A representative chest X-ray and CT scan image from a patient with CEP are displayed in Fig. 2. Pleural effusions have been reported as a rare manifestation of CEP, as has radiographic cavititation (93, 161). Pulmonary function testing is often impaired on disease onset, but the pattern of impairment may be either obstructive or restrictive (105).

As with many idiopathic eosinophilic syndromes, the mainstay of treatment is corticosteroid therapy. While the improvement in symptoms and infiltrates with corticosteroids is rapid, relapses are common with attempts to taper the steroid dosage (121). Inter-
estingly, patients with asthma in addition to CEP have been ob-
erved to have lower rates of relapse, perhaps due to high doses of
inhaled corticosteroids (104). Pulmonary function testing may
remain abnormal with an obstructive pattern, despite symptom-
atic and radiographic improvement (41). CEP may rarely progress
to lung fibrosis (41, 218). It should also be noted that CEP shares
many features with Churg-Strauss syndrome (discussed below),
with CEP in some cases possibly representing a prodrome for the
development of Churg-Strauss syndrome (105).

**AEP**

As the name implies, acute eosinophilic pneumonia (AEP) is char-
acterized by a rapidly progressive infiltration of eosinophils into
the lungs, leading to respiratory failure, distinguishing it from the
more indolent and less severe presentation of chronic eosinophilic
pneumonia. AEP is a relatively recently described phenomenon,
with the initial cases reported in 1989 (4, 6). Allen et al. described
a series of four patients who presented acutely with fever, severe
hypoxemia, bilateral pulmonary infiltrates, and BAL fluid eosino-
philia in the absence of identifiable infectious causes and whose
symptoms resolved rapidly with corticosteroid therapy (4). As
with CEP, the early reports of AEP remain representative of the
contemporary understanding of the syndrome.

The underlying mechanisms of AEP are even more poorly un-
derstood than those of CEP, though there is significant overlap
with CEP in the presence of specific inflammatory mediators in
the BAL fluid of patients with AEP. As in CEP, the chemokine
CCL17 has been detected in high levels in AEP, potentially consist-
tent with a role for recruited T cells in the pathogenesis of AEP
(114). An array of both Th2 and Th1 cytokines are elevated in the
BAL fluid in AEP, including IL-1ra, IL-2, IL-5, IL-10, IL-12, IL-13,
and IL-18, perhaps reflecting the known pluripotent secretory po-
tential of eosinophils (3, 76, 180). In particular, elevated BAL fluid
levels of IL-5, the key cytokine promoting differentiation, migra-
tion, and survival of eosinophils, are associated with AEP (120,
134, 155, 184, 215). Cationic granule proteins and inflammatory
lipid mediators are detectable in the urine of subjects with AEP
(136). Vascular endothelial growth factor (VEGF) is elevated in
the lungs in AEP, consistent with increased vascular permeability
contributing to the pathophysiology of the disease (125). BAL
fluid eosinophils in AEP have been noted to have an activated
phenotype, with upregulation of HLA-DR and CD69 (135). Se-
rum IgG levels, particularly those of the IgG2 and IgG4 subclasses,
are lower in patients with AEP than in healthy subjects (110).
Though no specific fungal etiologies have been found to underpin
idiopathic AEP, high concentrations of (1 → 3)-β-D-glucan have
been observed in the BAL fluid of AEP patients (77).

While the etiology of AEP remains incompletely delineated and
no specific inciting agents have been identified, a strong associa-
tion exists between cigarette smoking, particularly of recent onset,
and the development of AEP (171–173, 201, 219). In one AEP case
series, 32 of 33 patients were current smokers at disease onset, with
21 patients having started smoking within 1 month of disease
onset and 6 others having had recent substantial increases in the
quantity of cigarettes smoked (192). Rechallenge with cigarette
smoke has been reported to be associated with recurrence of dis-
ease (201). Another case series of AEP in U.S. military personnel
serving in Iraq featured a history of current smoking in all 18
patients, with recent onset in 14 patients (174). Secondhand
smoke exposure, smoking of flavored cigars, and exposure to
World Trade Center dust have all also been associated with devel-
opment of AEP (5, 87, 153).

The rapid onset of dyspnea and fever of less than 1 month in
duration is characteristic of AEP (67, 203). The condition affects
men more commonly than women (67). Hypoxemia and respira-
tory distress may be severe enough to necessitate initiation of me-
chanical ventilation (67, 143, 203). Chest X-rays demonstrate air-
space and reticular opacities, usually bilaterally, and CT scans
often have a significant component of ground-glass opacification
with accompanying septal thickening and thickening of the bron-
chovascular bundles (23, 33, 84, 115). Obtaining BAL fluid by
bronchoscopy is necessary to make the diagnosis, using an eosin-
ophil percentage of >25% as the criterion (67). While lung biopsy
is not needed diagnostically, pathology may show diffuse alveolar
damage along with eosinophil infiltration (188) (Fig. 3). Periph-
ernal blood eosinophilia is not a necessary diagnostic feature of AEP
(203). AEP is exquisitely responsive to corticosteroid therapy, and
though reported treatment regimens are variable in dosage and
duration, patients do not generally relapse after completion of
therapy (67, 147).

**CSS**

Churg-Strauss syndrome (CSS), also referred to as allergic angitis
granulomatosis, is a condition in which eosinophilic vasculitis and
infiltration potentially involve multiple organs, including the si-
nuses, lungs, peripheral nerves, heart, skin, gastrointestinal tract,
and kidneys (28, 80, 203). In many patients, sinusitis and asthma-
lke symptoms accompanied by peripheral eosinophilia are the
presenting constellation of signs and symptoms (56, 92). Like

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**FIG 2** Representative chest X-ray (A) and CT scan (B) of chronic eosinophilic pneumonia.

**FIG 3** Transbronchial lung biopsy specimen showing acute eosinophilic pneumonia. Numerous infiltrating eosinophils can be seen.
the other noninfectious pulmonary eosinophilic syndromes, the pathophysiology of CSS is not fully understood. While antineutrophil cytoplasmic autoantibodies (ANCA) against myeloperoxidase are often seen in CSS, it is unclear if CSS is truly an autoimmune disease (158, 177). There is likely an important role of T cells in the pathogenesis of CSS, with Th2 inflammation representing an important component of the disease (34, 65, 81, 195). However, the factors that differentiate CSS from other Th2-driven eosinophilic processes remain incompletely delineated. Eosinophil depletion with anti-IL-5 monoclonal antibodies has shown promising preliminary results in treating CSS, emphasizing the mechanistic importance of IL-5 in the disorder (82). Circulating biomarkers of CSS include IL-25 and the chemokines eotaxin-3 (CCL26) and TARC (thymus and activation-regulated chemokine; also called CCL17), suggesting a potential role for these proteins in CSS pathogenesis (34, 146, 189, 221). Several medications used in the treatment of asthma, including leukotriene modifiers, inhaled corticosteroids, and omalizumab, have been associated with the development of CSS (95, 204–206). However, it is likely that these medications do not have a direct causative role in CSS; rather, their initiation accompanies withdrawal or reduction of corticosteroids, which in turn unmask active CSS, or simply reflects their use in treating the asthma of incipient CSS (203).

Respiratory manifestations are an integral component of CSS, with asthma as a major criterion for all systems that have been implemented for the diagnosis of CSS (203). While presentations are variable, CSS has been described as having a prodromal phase in which asthma and rhinosinusitis are the predominant features. The disease may then progress to an infiltrative phase involving the lungs and other organs, followed by a vasculitic phase (56, 92). Regardless of classification scheme, asthma, eosinophilia, pulmonary infiltrates, and eosinophilic vasculitis are components of establishing the diagnosis. As mentioned above, patients with CSS often have a positive ANCA test result. Inflammatory markers, including the erythrocyte sedimentation rate, C-reactive protein, and rheumatoid factor, may be elevated, along with circulating IgE levels (203). Notable extrapulmonary manifestations include mononeuritis multiplex and eosinophilic infiltrative cardiac disease (36, 92). CSS has the potential to be a fatal condition, in particular secondary to its cardiac manifestations (123).

Corticosteroids are the mainstay of therapy for CSS, though remissions may be seen with tapering of steroid therapy (150). Cyclophosphamide and several other immunosuppressive agents have been used to treat CSS as adjunctive, potentially steroid-sparing therapies (47, 57, 59). The use of a monoclonal antibody against IL-5 is an experimental therapy that is not currently commercially available and awaits definitive clinical trials (82).

**HES**

The term “hypereosinophilic syndromes” (HES) refers to a range of disorders characterized by elevated numbers of circulating blood eosinophils accompanied by various end-organ infiltration leading to clinical disease (157, 168, 208). The major subcategories of HES are the myeloproliferative variants and the lymphocytic variants of the disease. While many cases of HES are truly idiopathic, specific etiologies have been identified for some, such as myeloproliferative variant HES due to platelet-derived growth factor α gene rearrangement.

Pulmonary manifestations may occur in up to 40% of patients with hypereosinophilic syndromes, though only a subset of these patients can truly be grouped in the category of pulmonary eosinophilia with distinct radiographic abnormalities, given that most individuals with HES and cough have clear chest X-rays (40, 208). When they are present, infiltrates may occur in any region of the lungs. Biopsy of affected areas may show parenchymal infiltration of eosinophils. Because HES commonly affects the heart, radiographic abnormalities, including interstitial changes and pleural effusions, can often be attributed to congestive heart failure (208). As opposed to many of the other noninfectious causes of eosinophilic pneumonia discussed in this article, corticosteroid therapy may be ineffective in this case, and other immunomodulatory therapies, such as imatinib, are often employed (86, 131). A randomized, placebo-controlled trial has demonstrated that specifically targeting eosinophils with a monoclonal antibody against IL-5 is effective as an adjunct therapy for HES, though not yet commercially available (156).

**CONCLUSIONS**

Eosinophilic pneumonia can be caused by several infectious etiologies of parasitic origin. The noninfectious etiologies can be categorized into allergic, exposure-related, and idiopathic etiologies (Table 2). The clinical history and careful evaluation of patients allow clinicians to first narrow the potential causes to these broad categories and then to a specific diagnosis. Corticosteroids are an
important component of the therapy for many of these conditions, both infectious and noninfectious, though they may precipitate hyperinfection syndrome in cases of strongyloidiasis. While anthelminthic therapy is needed for several of the infections discussed, others are self-limited in nature and likely do not require any therapy.

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