Eosinophilic pleural effusion: A case and a review

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Abstract
A previously healthy middle-aged woman presented with evanescent skin lesions and bilateral pleural effusions with an eosinophilic predominance. Following this case summary, a description of eosinophilic pleuritis, the epidemiology, etiologic considerations, and selected therapies for this syndrome are discussed. Eosinophilic pleural effusion is caused by myriad etiologies and is a therapeutic challenge.

Keywords: pleural effusion, eosinophilic effusion, dermatitis

Introduction
Pleural effusions are a common clinical problem in adult medicine, but an effusion with a cellular component comprised of many eosinophils is not. Eosinophilic pleural effusions (EPE) have been attributed to a wide variety of etiologic agents, including infections, adverse drug reactions, malignancy, and chest trauma. Even after investigating these causes, the cause of an EPE can still be uncertain. After presentation of a case, an overview of the causes of EPE is discussed followed by comments on therapy.

Case
A 37-year-old Hispanic homemaker developed left sided pleuritic chest pain over a few days in the fall. This was associated with malaise, fatigue, and some sweats but no fever or chills. She was admitted to a hospital and diagnosed with pneumonia for which she received quinolone and macrolide antibiotics. She didn’t respond, and dyspnea developed. Echocardiography was normal. She was transferred to University Medical Center in Lubbock, Texas, when computed tomography of the thorax showed pleural effusions. She had no cough or other focal symptoms except for evanescent skin lesions, occurring over the prior weeks, primarily on the trunk and proximal extremities. These were not pruritic or triggered by trauma or sun exposure. She had no past medical history, medication use, or social history relevant to this presentation. She had traveled to California several months prior to presentation.

She had BMI of 33.6 kg/m², a normal temperature, and normal vital signs except for a respiratory rate of 20 breaths per minute with 91% oxygen saturation on room air. Reduced breath sounds were present at both bases. There were several scattered annular erythematous plaques on the abdomen and proximal thighs (Figure 1). The remainder of her examination was within normal limits showed no adenopathy.

Her leukocyte count was 6,170/µL (75% neutrophils, 20% lymphocytes, 4% monocytes, 0% eosinophils), hemoglobin was 9.4 gm/dL (MCV 76 fl), and the platelet count was normal. C-reactive protein was 12.1 mg/dL. Serum chemistry profile was normal. Both the chest radiograph (Figure 2) and chest computed tomography (Figure 3) showed bilateral pleural effusions and possible lower lung consolidations. Transthoracic echocardiography showed a small pericardial effusion but no other abnormalities. Thoracentesis recovered pleural
fluid with 3200 red cells/mm$^3$ and 2558 leukocytes/mm$^3$ (84% eosinophils, 9% lymphocytes, 7% monocytes). Pleural fluid LDH was 663 IU/L (corresponding serum value: 245 IU/L), and pleural fluid protein was 4.9 g/dL. Gram stain was negative, and cultures for bacterial and fungi had no growth. Investigations conducted over the subsequent week for infectious, autoimmune, and neoplastic conditions are summarized in Table 1. The skin lesions lasted a few days, and then similar lesions appeared at various other sites over the next week.

During this hospitalization, the effusions increased in size. She was started on prednisone with rapid reduction in symptoms and no further increase in the pleural fluid. Her skin lesions completely disappeared. Following hospitalization, the prednisone was tapered over several weeks, but at a lower dose, symptoms and effusions returned. This occurred over the next month at which point she had right video-assisted thoracoscopy which showed “murky fluid” but normal lung adherent to the pleura. Pleural biopsy and mechanical pleurodesis followed without complications. Pathology showed fibrinoid eosinophilic inflammatory infiltrates in the pleura. Lung tissue had patchy alveolar wall fibrosis at subpleural areas, but no eosinophilic alveolar process was apparent beyond the pleural-lung junction. No granulomas, vasculitic features, or malignant cells were seen (Figure 4).

High dose prednisone therapy reduced her symptoms, including the rash, and reduced her pleural effusions. However, whenever the corticosteroid dose was reduced, her conditioned worsened. Two months after the start of her therapy she developed a pericardial effusion, without eosinophilia in the pericardial fluid, which resolved with a pericardial window. She remained without cough, fever, arthralgias, or lymphadenopathy. Ultimately immunosuppression with steroid-sparing regimen consisting of colchicine and hydroxyurea was required to control her disease. Two years later she has fatigue but is asymptomatic with no recurrent effusion or skin lesions.
**DISCUSSION**

Eosinophilic pleural effusions can present both diagnostic challenges and difficult therapeutic decisions. The eosinophil is activated by T lymphocytes and mast cells, particularly upon release of interleukin (IL)-3 and IL-5; the latter prolongs the survival of eosinophils. Eosinophilia exists when 3% or more of peripheral blood leukocytes are eosinophils or when the absolute number is over 350 cells per cubic millimeter. An eosinophilic pleural effusion (EPE) is present when over 10% of the nucleated cells are eosinophils.\(^1\) Eosinophilic pleural effusion results with increased marrow production of eosinophils which migrate into and persist in the pleura. The migration and residence in a tissue is related to expression of vascular cell adhesion molecule-1.\(^1\)

Eosinophilic pleural effusions are a relatively uncommon problem. In a retrospective review of over 2,200 European patients with pleural effusions, from 1995 to 2007, only 7.2% had EPE.\(^1\) The etiology of pleural effusions is unknown in 10–20% of cases. With EPE the proportion of cases due to an unknown etiology may be even higher. In most studies in which an etiology is identified for EPE, malignancies are ranked near the top with lung and breast cancers among the most common followed by lymphomas and a variety of other solid and gastrointestinal neoplasms. An EPE has developed in a patient after mediastinal radiation therapy for Hodgkin's disease. The degree of eosinophilia in the pleural fluid tends to be less in the neoplastic causes than other etiologies.\(^1\) A summary of some of the common causes of EPE is shown in Table 2.

Infectious disease etiologies for EPE typically include parapneumonic effusions from pneumonia and *Mycobacterium tuberculosis* (MTB). The proportion of pleural effusions caused by MTB reflects the prevalence of tuberculosis in the patient population. In Tunisia, where tuberculosis is prevalent and pleural TB relatively common, nearly 40% of pleural effusions were attributed to MTB, whereas a series of...
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The Southwest Respiratory and Critical Care Chronicles 2020;8(33):40–46

patients in Poland included just over 2% caused by MTB. Pleural effusions associated with pulmonary tuberculosis or pleuritis typically have a lymphocytic predominance with <10% eosinophils. If the proportion is greater, tuberculous effusion is unlikely. In patients with “Valley Fever,” caused by the mycosis Coccidioides immitis or C. posadasii endemic to the southwestern deserts of the United States, northern Mexico, and portions of Central and South America, a mild peripheral eosinophilia may occur in 25% of cases with pulmonary infection. Even in cases in which the total blood leukocyte count is normal, mild (>5%) eosinophilia has been documented in 27% of patients by retrospective review. Coccidioidomycosis with pleural effusion occur in 15% of patients with pneumonia. In one small series, 12 cases of EPE were evaluated, and the degree of eosinophils in the pleural fluid ranged from 1–44% of the nucleated cells. Other endemic mycoses have also been associated with EPE but not as frequently as with coccidioidomycosis.

Eliciting a travel history in patients is important since a variety of tissue-migrating parasites have been associated with eosinophilia. For example, paragonimiasis can cause a pulmonary syndrome with hemoptysis, mimicking pulmonary tuberculosis. Paragonimus sp. can infect patients following ingestion of fresh water crustaceans from Asia, South America, and Africa. Eating crawfish in the southern United States

Table 1. Diagnostic tests in the patient in this case report

| Study                              | Result/Interpretation |
|------------------------------------|-----------------------|
| ANA, c-ANCA antibody               | Negative              |
| Antimyeloperoxidase , Antiproteinase 3 antibody | Negative         |
| Strongyloides antibody             | Negative              |
| Interferon gamma-release assay M. tuberculosis | Negative          |
| HIV antibody                       | Negative              |
| Syphilis screen                    | Negative              |
| HCV antibody, HBs antigen          | Negative              |
| Coccidioidomycosis serology        | Negative              |
| Histoplasma urine antigen          | Negative              |
| Beta D glucan antigen (serum)      | Negative              |
| Procalcitonin                      | 0.3 ng/ml             |
| Urine drug screen                  | Negative              |
| Bone marrow biopsy                 | Red cell hypochromia, 1% eosinophils; normal |
| Skin biopsy                        | Lichenoid dermatitis with eosinophils |
| C3, C4                             | Normal                |
| CH50                               | >60 (elevated)        |
| JAK2                               | Negative              |
| CT chest angiography               | Normal                |
| Interleukin 5                      | <5 pg/ml              |
| FISH, HES/4q12                      | Negative for FIP1L1- PDGFRA rearrangement |
Pleural biopsy reveals fibrinoid eosinophilic inflammatory infiltrates, H & E stain.

has also been identified as a source of infection. The lung fluke often stimulates peripheral eosinophilia and EPE. Eosinophilic pleural effusion also occurs in filariasis, ascariasis, sparganosis, strongyloidiasis, and infections with other trematodes and cestodes.7,8

Evanescent eosinophilic skin lesions with EPE, as in this case, suggests the possibility of Strongyloides or a generalized autoimmune disorder. Besides larva currens, strongyloidiasis may be accompanied by peripheral eosinophilia and a form of urticaria around the buttocks and waist. These lesions last 1–2 days and recur.9 The patient presented above had no travel outside the United States. There were no pulmonary, dermal histology, or serologic findings to attribute this case to strongyloidiasis. Biopsy of skin lesions related to strongyloidiasis typically shows filariform fragments, and the enzyme immunoassay for detection of the helminth exceeds 85% sensitivity.10,11

The eosinophilic dermatoses, like EPE, may occur with or without blood eosinophilia. Most cases are related to allergic conditions, such as eczema and atopic dermatitis. Besides parasitic infestations, arthropod bites, vasculitic disorders, and blistering skin conditions such as bullous pemphigoid may exhibit similar histopathology. Eosinophilic dermatosis evolves by poorly understood mechanisms.12 Coccidioidomycosis can be associated with EPE and skin lesions such as erythema nodosum and diffuse evanescent maculopapular rashes. This patient had no pulmonary disease and no evidence of fungus on microbiologic, pathologic, or serologic analysis.

Chest trauma, pneumothorax, thoracotomy, splenectomy, and pulmonary embolism have been associated with EPE. In most of these conditions, eosinophils represent only a minority of the nucleated cells present

| Cause                        | Relative Frequency* (%) | Comment                                      |
|------------------------------|-------------------------|----------------------------------------------|
| Malignancy                   | + + +                   | Half are from lung cancer                    |
| Parapneumonic Effusion       | + +                     | Eosinophils rarely comprise >10%             |
| M. tuberculosis              | +                       | Eosinophils comprise <10% NBC                |
| Coccidioidomycosis           | + +                     | Lymphocytes predominate                      |
| Post chest surgery           | +                       | Including Dressler’s syndrome                |
| Transudative effusions**     | +                       | Almost never seen with CHF                   |
| Autoimmune disorders         | –                       | Includes eosinophilic pneumonia              |
| Post chest trauma            | + +                     | Probably similar mechanism to surgery        |
| Idiopathic                   | + +                     | Often higher % eosinophils in fluid          |

*+ + +, + +, +, – correlates with >25%, >10%, >5%, or <5% of cases, respectively.

**Due to CHF, cirrhosis, nephrotic syndrome.
in the pleural fluid. Trauma and stimulation of pleural mesothelial cells may release cytokines which recruit eosinophils to the affected zone, perhaps in some individuals due to HLA variations and other pre-determined immunologic phenotypes. Rheumatoid arthritis, Churg-Strauss, and Crohn’s disease have rarely been associated with EPE.\textsuperscript{2,13}

Uncommonly EPE has been reported with warfarin, propylthiouracil, sulfasalazine, and nitrofurantoin. Rarely simvastatin, isotretinoin, fluoxetine, valproate, infliximab, and crack cocaine have been implicated in EPE.\textsuperscript{4} There was no recent history of exposure to these drugs in our patient, but a remote drug use history was not obtained. Asbestososis has a notorious association with pleural disease. In a small retrospective evaluation of men exposed to asbestos with pleural disease, 7 of 10 with pleural effusions had “large numbers” of eosinophils. Pathologically, thickened chronic pleuritis was observed with fibrosis, and, in one subject, asbestos bodies were seen.\textsuperscript{15} In our patient there was no known exposure to asbestos. When asbestos exposure has caused disease, the interval between asbestos exposure to presentation is usually several decades. In this instance, there was no history of exposure nor was the plural histopathology suggestive of this disease.

In larger retrospective reviews of EPE, nearly 20% of patients had no discernable cause.\textsuperscript{2} Even after extensive investigations, the etiologic diagnosis for EPE may be difficult. In this case, despite a fairly extensive evaluation of the etiology in this patient, the cause of her EPE was not determined. Empiric therapy with hydroxyurea and colchicine allowed for de-escalation of high dose prednisone. Hydroxyurea has been used successfully in the management of a variety of hypereosinophilic syndromes with less short term morbidity than corticosteroids. There are case reports documenting successful treatment of eosinophilic skin disorders with colchicine.\textsuperscript{16}

Since EPE is not a common syndrome no widescale therapeutic information is available. Thus, anecdotal reports comprise the major reference for our therapeutic experience of EPE.

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