Serratia pneumonia presenting as hemoptysis in a patient with sarcoidosis: a case report

Paul Zarogoulidis1
Konstantinos Porpodis1
Maria Konoglou2
Maria Saroglou2
Alexandros Mitrakas1
Dimitrios Matthaios3
Panagiotis Tzouopoulos1
Konstantinos Archontogeorgis4
Andrew Koulelidis4
Konstantinos Zarogoulidis1
Stavros Tryfon2

1Pulmonary Department, “G. Papanikolaou” General Hospital, Aristotle University of Thessaloniki, Greece; 21st Pulmonary Clinic, “G. Papanikolaou” General Hospital, Thessaloniki, Greece; 31st Internal Medicine Department, University General Hospital of Alexandroupolis, Greece; 4Pulmonary Department, University General Hospital of Alexandroupolis, Democritus University of Thrace, Greece

Introduction: Serratia marcescens is a Gram-negative bacillus which belongs to the family Enterobacteriaceae. It is a facultative anaerobe and produces red pigment at room temperature. It naturally occurs in soil and water as well as the intestines, and it is responsible for nosocomial infections. There have been few reports about community acquired pneumonia of Serratia.

Case presentation: This report presents a 37-year-old man with hemoptysis, fever, and shortness of breath. The clinical and laboratory examinations revealed that the patient had pseudo-hemoptysis due to S. marcescens pneumonia, on an immunocompromised pattern, because of the coexistence of sarcoidosis (stage 1).

Conclusion: Appropriate antibiotic therapy for Serratia was administered, and the patient’s symptoms regressed. The patient is healthy and asymptomatic after 1-year follow-up. To the best of the authors’ knowledge, this is the first reported case of a pseudohemoptysis in a patient with pulmonary sarcoidosis.

Keywords: Serratia marcescens, pseudohemoptysis, pulmonary sarcoidosis

Introduction
Serratia, is an opportunistic pathogen, mainly responsible for infecting the urinary system, respiratory tract, central nervous system, and bloodstream.1 Infections of colonized individuals are associated with invasive devices, surgery, and immunocompromised states. Serratia ubiquitously thrives in moist environments and frequently contaminates liquids and equipment. In the outpatient setting, it has been associated with community-acquired pneumonias, but only limited case reports exist, and usually are associated with some degree of immunocompromise. This paper reports a case where the poor living environment and coexistence of sarcoidosis were likely the predisposing factors that developed pneumonia from Serratia.

Case report
A 37-year-old man presented in the emergency department complaining of productive cough with blood-tinged sputum, intermittent fever, and shortness of breath for the last 20 days. He was a lifelong nonsmoker, currently unemployed, and occasionally homeless. His past medical history was remarkable only for arterial hypertension treated with atenolol.

On examination, he was ill-appearing but in no acute distress. His axillary temperature was 37.8°C, blood pressure was 140/80 mm Hg with a normal heart rate of 90 bpm, and oxygen saturation SpO2: 94%. Lung auscultation revealed crackles over the lower right lung.
On laboratory testing, the patient had a mildly elevated white blood cell count of 10,900 cells/μL, with normal differential. The hemoglobin level was 10.6 g/dL, with a hematocrit of 32.5%; the platelet count was 455,000/L. The erythrocyte sedimentation rate was 55 mm after the first hour. A basic metabolic panel was notable for an elevated serum creatinine of 2.52 mg/dL, with urea of 57 mg/dL, and normal electrolytes. Serum angiotensin converting enzyme was slightly elevated at 57 IU/mL (normal range 8–52 IU/mL). The acute renal failure was attributed to the Serratia infection, since the patient did not previously have this disorder and Serratia is known to induce renal failure.1

There was no lymphadenopathy observed in general examination, but chest x-radiography demonstrated bilateral hilar prominence, lower-right lung lobe opacification, and right hemidiaphragm elevation (Figure 1). Chest high-resolution computed tomography scan was performed according to the department’s protocol for investigating the thorax. The scan revealed consolidation of the anterior segment of the right lower lobe, and bilateral paratracheal, subcarinal, and hilar lymphadenopathy (Figure 2). In addition, transbronchial needle biopsy aspiration was conducted as a diagnostic approach.

Pulmonary function testing after remission of the productive cough (2 days) showed a restrictive pattern with a forced vital capacity (FVC) of 49% predicted, a forced expiratory volume in one second (FEV1) of 47% predicted, and an FEV1/FVC ratio of 71%. The total lung capacity was 66% of predicted, and the residual volume was 79% of predicted. There was reduction of the diffusion lung capacity for carbon monoxide to 68% of predicted.

Serologic studies for human immunodeficiency, hepatitis B virus, and a tuberculin skin test were negative. Urine antigen for Streptococcus pneumoniae and Legionella were given upon admission, but they were also negative.

Fiberoptic bronchoscopy revealed a partial extraluminal compression of the trachea without other endobronchial lesions. No evidence of inflammation, granulomas, malignancy, vasculitis, or alveolar wall necrosis were found on biopsy. The cytological results from the bronchoalveolar lavage fluid (BALF) showed no malignancy. The cell count on the BALF was 10.5 × 10^10 cell/L, with 78% alveolar macrophages, 12% lymphocytes, and 10% neutrophils. No red blood cells or hemosiderin-laden macrophages were seen. BALF flow cytometry showed that 69% of the lymphocytes were T-lymphocytes, 7% were B-lymphocytes, and the T4/T8 ratio was 1.9, while the blood T4/T8 ratio was 1.4. Microbiology cultures of the BALF grew Serratia marcescens, which was sensitive to ciprofloxacin, tobramycin, and amikacin. The smear for acid-fast bacilli was negative, and cultures for mycobacteria remained negative after 6 weeks.

The patient was treated with ciprofloxacin 500 mg twice daily for 3 weeks. After the third day of treatment, the cough improved, and the reddish appearance of the sputum completely resolved. On follow-up after 1 month, chest x-radiography presented complete remission on right lower lobe infiltrate, while the perihilar lymphadenopathy persisted. A surgical mediastinal biopsy showed nonnecrotizing granulomas, consistent with sarcoidosis. The final diagnosis was pneumonia due to S. marcescens and stage I pulmonary sarcoidosis.

After 1-year follow-up, the patient remains asymptomatic, and on chest x-radiography the infiltrate has completely resolved, while enlargement of the perihilar lymph nodes persists (Figures 3 and 4).

**Discussion**

This is a case of pseudohemoptysis due to pulmonary infection with S. marcescens and concomitant sarcoidosis.
Serratia pneumonia presenting as hemoptysis has previously been documented. An infection can be the presenting problem subsequently leading to the diagnosis of sarcoidosis, as in this case.

In sarcoidosis immunopathogenesis, the immune system undergoes a reactivity change. Primary features are a reduction in circulating CD4+ lymphocytes (T-helper cells), while their numbers in tissue increase, which is associated with a marked increase in tissue cytokine production, particularly interferon-γ, granulocyte-macrophage colony-stimulating factor, and interleukin-2. This T-helper 1 (Th-1) cytokine profile recruits macrophages, eventually forming a granulomatous reaction.

In contrast to these highly active tissue-based responses, the immunity related to the circulatory cells expression is depressed by cutaneous anergy and poor responses to antigen recall testing. In addition, the production of inhibitory cytokines by macrophages appears to interfere with the normal immune response. Girard et al reported five cases and reviewed 65 additional cases of opportunistic infections associated with sarcoidosis. While the majority of these occurred in patients receiving corticosteroids and were accompanied by CD4 lymphocytopenia, they described opportunistic infections in four untreated individuals. Observations have suggested a pathogenetic role of Gram-negative infections in sarcoidosis by driving an interleukin-18 response as another feature of the Th-1 pathway. This has been postulated for infections with Hemophilus influenzae and Moraxella catarrhalis.

Conclusion
In summary, this report describes a rare case of an opportunistic infection in an outpatient setting. In this case, the sarcoidosis was the underlying condition that induced an infection by Gram-negative bacilli.

Consent
Written informed consent was obtained from the patient upon discharge for publication of this case report and all accompanying images.

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Disclosure
The authors declare that they have no conflicts of interest in this work.
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