**CASE PRESENTATION**

A 19-year-old woman with a medical history significant for poorly controlled diabetes mellitus (DM) (hemoglobin A1C 11.2%) presented with a 12-day history of right-sided pleuritic chest pain. One month earlier, she had developed sudden-onset fever, dry cough and shortness of breath. At the time, she did not have any gastrointestinal symptoms including epigastric pain, heartburn, vomiting or retching. She denied sick contacts and had no preceding symptoms of an upper respiratory tract infection. She was seen at a local emergency department and was released home on a 10-day course of oral doxycycline (100 mg every 12 h) to treat a community-acquired pneumonia (CAP). After an initial two-week improvement, her complaints relapsed, with fevers, dry cough, poor appetite and the above-described chest pain. The patient did not smoke and she denied alcohol abuse or illicit drug use.

On presentation to the hospital, she was febrile (temperature 38.2°C), her oxygen saturation was 86% on room air and she was thin. She was awake and oriented but in obvious pain. She had no jugular venous distention. Her heart sounds were regular without murmurs or gallops. The lung examination revealed reduced air entry over the right lung base with bronchial breath sounds and egophony. The abdomen was soft without tenderness or distension and she had no evidence of subcutaneous emphysema or lymphadenopathy. There were no lower limb edema and no joint swelling or deformities.

The laboratory tests revealed a white blood cell count of 26.0×10⁹/L and a hemoglobin level of 119 g/L. Blood chemistry showed normal kidney and liver function. Tests for HIV and pregnancy were negative. Chest imaging included a chest x-ray and a contrasted chest computed tomography scan (Figures 1 and 2). The patient was admitted to hospital with severe sepsis due to CAP complicated by an empyema and was started on intravenous broad-spectrum antibiotics, vancomycin (15 mg/kg every 12 h) and piperacillin/tazobactam (3.375 g every 6 h). She underwent computed tomography-guided drainage 48 h afterwards. An interventional radiologist performed the procedure under strict sterile techniques. Cytopathology (Figure 3) and cultures were ordered on the drained fluid.

What is your diagnosis?

**DIAGNOSIS**

The fluid Gram stain was negative and the culture showed pure growth of *Candida albicans*. Hematoxylin and eosin-stained section of the pleural fluid showed numerous neutrophils and cellular debris consistent with empyema with occasional pseudohyphae. No squamous cells or ingested material was identified.

Based on the cytopathology findings and the aforementioned culture results, the patient was diagnosed with candida empyema thoracis (CET) that complicated an inadequately treated CAP with parapneumonic effusion. CET has been reported to be a part of a mixed infection in up to 46% of cases (1). A second pleural fluid sample was sent from the drain and it, once again, grew *C albicans*. She was started on intravenous micafungin (100 mg daily) with significant subsequent clinical improvement.

In the present case, we speculate that doxycycline disrupted the normal flora and allowed for candidal overgrowth. The patient’s poorly controlled DM facilitated translocation from mucus surfaces into the bloodstream and, subsequently, into the parapneumonic effusion, worsening the patient’s symptoms (2,3).

Before discharge, the intravenous antifungal was changed to oral fluconazole (200 mg daily) and her antibacterial therapy was de-escalated.
to four-week course of the above therapy. She improved markedly and continues to be followed in the outpatient clinic.  

**DISCUSSION**

CET is a rare entity. Most cases are acquired in health care settings (4). Community-acquired CET comprises <15% of reported cases (4). Risk factors for this infection include malignancies, especially of the esophagus and stomach; DM; cirrhosis; and immunocompromised states including the use of long-term steroid therapy, HIV infection or having undergone organ transplantation (1,4). Only a minority of individuals with this ailment have no underlying morbidity. The infection results from direct inoculation into the pleural space following esophageal rupture, or after thoracentesis or thoracotomy. The former mechanism of infection is the most common (1,4). On the other hand, it can spread from contiguous foci such as a fungal lung infection or a subdiaphragmatic abscess that complicated visceral perforation. On occasion, CET can result from hematogenous seeding. Fungemia was documented in 5% to 20% of patients with this entity in the two largest case series in the literature (1,4).

C. albicans is the most commonly implicated culprit causing fungal empyema, followed by Candida tropicalis and Candida glabrata (1,4). On rare occasions, other fungi, such as Aspergillus and Cryptococcus, have also been implicated in causing this infection.

Patients with CET complain of fever, shortness of breath, cough and chest pain. They may also complain of the manifestations of the underlying condition that has led to the CET. For example, patients with esophageal rupture may complain of vomiting, severe chest pain and subcutaneous emphysema (5). Furthermore, patients presentation may be complicated by florid sepsis with significant leukocytosis, organ failure and shock.

Diagnostic criteria for candida/fungal empyema thoracis include (4):

- Fungal isolation from exudative pleural effusion;
- Signs of active infection such as fever and leukocytosis;
- Isolating the same fungus from pleural effusion in more than one occasion, or from pleural effusion and another specimen such as blood or tissue cultures.

The latter point is not needed to confirm the diagnosis if the fungus has been cultured from a specimen that was appropriately collected from a sterile site (6). Furthermore, new markers of invasive fungal infection (ie, beta-D-glucan and galactomannan) can be used to corroborate the diagnosis. Candida isolated from previously placed chest tubes or drains is considered a to be colonizer, unless infectious symptoms and signs persisted until antifungal therapy was initiated (1).

Early aggressive therapy is imperative to optimize outcome (1,4). It consists of empyema drainage and antifungal therapy. The initiation of broad-spectrum antibiotic therapy pending culture results is indicated, given that approximately one-half of fungal empyema thoracis cases have a bacterial coinfection (1). It is important to note that the concentration of antifungal agents in pleural fluid is variable (1). Voriconazole and micafungin have been reported to provide good pleural penetration (7-9). On the other hand, anidulafungin and liposomal amphotericin B are reported to have poor pleural concentrations (10-12). Data regarding fluconazole pleural concentration are lacking; however, it has been used successfully alongside empyema drainage in the treatment of the majority of patients in the above-mentioned case series (1,4).

The prognosis of CET is guarded, with reported mortality rates as high as 60 to 70% (1,4). Being immunocompromised, developing acute respiratory failure or developing septic shock are risk factors for poor outcome.

**CONCLUSION**

CET is an uncommon invasive candidiasis. Predisposing risk factors include malignancies, DM, cirrhosis and immunocompromised states. The prognosis of this infection is grim unless prompt drainage and effective antimicrobial therapy are initiated in a timely manner.

**DISCLOSURES:** The authors have no conflicts of interest to declare.

**REFERENCES**
