Transudative pleural effusion as an initial presentation of a disseminated cryptococcosis infection in a HIV-negative patient with cirrhosis

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1. Introduction

_Cryptococcus_ is a fungal spore that causes infections, particularly in HIV positive and immunocompromised patients. _Cryptococcus_ is an encapsulated yeast found in the environment. The fungus commonly infects the lungs resulting in pneumonia or the central nervous system, causing meningitis [1]. However, the fungus can disseminate to many different organs. Infrequently, individuals with diabetes mellitus, liver disease, kidney disease, and even no known medical history can have disseminated infection [1–3].

Individuals who are HIV-negative with ESLD have significantly higher mortality than HIV-negative patients without ESLD or HIV-positive patients. This higher mortality is likely related to the liver’s important role in generating an effective immune response, clearing cytokines and bacteria from the vasculature [4]. In patients with ESLD, this response is compromised and results in an increased risk of Cryptococcal meningitis and disseminated disease [3].

_Cryptococcal_ meningitis is a disease with significant morbidity and mortality, particularly in cases with disseminated infection. The presence of either of these conditions demands an aggressive treatment regimen with amphotericin B and flucytosine, while milder infections, such as pulmonary _Cryptococcus_, can be adequately managed with fluconazole [5]. _Cryptococcal_ infection in the lungs usually causes an exudative effusion [6–8]. In our literature review, we have not found a case of an individual with a transudative effusion who had _Cryptococcal_ pneumonia or _Cryptococcal_ pleural infection.

We present a fatal case of disseminated _Cryptococcus_ in a young adult with end-stage liver disease. His course was complicated by many factors: a long-standing ventriculoperitoneal shunt, hypopituitarism, and _Cryptococcal_ invasion of a persistent pleural effusion and his ascitic fluid. His case was particularly unique because despite having disseminated infection, his initial thoracentesis showed a transudative effusion.

2. Case presentation

We present a 32-year-old male transferred to our hospital to manage _Cryptococcal_ pleural infection and spontaneous bacterial peritonitis. The patient experienced abdominal pain and recurrent ascites, requiring multiple paracentesis procedures and albumin therapy for the past six months. He initially presented to an outside hospital with difficulty ambulating, nausea, dizziness, and a sodium level of 120 mEq/L. Paracentesis was done at an outside hospital, and the patient was started on ceftriaxone for possible spontaneous bacterial peritonitis. He developed dyspnea and increased oxygen requirements. Pleural fluid analysis from the outside hospital was positive for Cryptococcal antigen (Ag) with titer 1:256. The patient had no neurological or meningeal symptoms. He was started on IV fluconazole (400 mg) and transferred to our hospital for a higher level of care.

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Our patient’s medical history was significant for cryptogenic cirrhosis, right hypothalamic juvenile pilocytic astrocytoma status-post resection, and radiation (20 years prior) with bilateral occipital ventriculoperitoneal (VP) shunt placement, and panhypopituitarism. He was taking oral hydrocortisone (30 mg/daily) and desmopressin (DDVAP) and was frequently hypernatremic.

On admission, the patient was afebrile with a normal white blood cell (WBC) count of $7.5 \times 10^9$/L (Day 0). Blood cultures were initially negative. HIV test was negative. ALT was 28 units/L, and AST was 35 units/L. The viral hepatitis panel was negative. Given the patient’s history of panhypopituitarism, he was started on stress-dose steroids (IV hydrocortisone 50 mg $+1$). On day +1, our patient had symptomatic relief of his dyspnea after a thoracentesis removed 1.5 L of transudative fluid (Table 1). A paracentesis was performed the following day, which showed 206 nucleated cells/$\mu$L, and ceftriaxone was discontinued since the cell count differential was not consistent with spontaneous bacterial peritonitis. CT thorax after the thoracentesis revealed a massive left-sided pleural effusion and atypical pneumonia on the right with reactive lymphadenopathy (Fig. 1). On day +3, the serum Cryptococcal Ag was positive. Infectious Disease (ID) recommended switching fluconazole to liposomal amphotericin B (3 mg/kg q24h) and flucytosine (25 mg/kg q6h) to treat a disseminated C. neoformans infection. Flucytosine toxicities were monitored with daily complete blood count to ensure there was no agranulocytosis. On day +4, the transudative pleural fluid Cryptococcal Ag was positive. On day +5, the blood cultures from admission grew C. neoformans. In the following days, the pleural and abdominal fluid cultures also grew C. neoformans (Fig. 2). Given fungemia, a transthoracic echo was performed, which showed no vegetation. CT of the head did not reveal any ventriculitis. A lumbar puncture was completed, and the CSF was positive for Cryptococcal Ag on day +4. The CSF culture was positive, resulting six days later (+6). The patient was transferred to the neurologic intensive care unit for the removal of his VP shunt and placement of an external ventricular drain (EVD).

The patient had the left VP shunt removed; however, the right could not be removed due to adhesions. The CSF was monitored frequently and remained positive for Cryptococcal Ag until the tenth day of antifungal therapy. On day +11, a therapeutic thoracentesis revealed an exudative, serosanguinous fluid (Table 1), which grew three Cryptococcus colonies. The patient became lethargic with altered mental status. An EEG was consistent with non-convulsive status epilepticus, and our patient was placed on levetiracetam and lacosamide. He was intubated and put on a midazolam drip with ketamine to achieve burst suppression. The patient required pressor support at this time. Nephrology was consulted for anuric AKI, which was likely due to acute tubular necrosis. The patient had stopped responding to furosemide diuresis and was started on continuous veno-venous hemofiltration (CVVH). His flucytosine was renally adjusted to 25 mg/kg q24 rather than q6. The patient further deteriorated with worsening septic shock despite negative repeat blood cultures drawn on day +13. The patient was started on piperacillin/tazobactam and vancomycin, which were later changed to meropenem and daptomycin. The patient required five pressors, sodium bicarbonate infusion, and an increase in stress steroids. He developed shock liver with disseminated intravascular coagulopathy (DIC). On day +14, the patient started to bleed from his nasogastric tube. He was given multiple transfusions of blood, platelets, fresh frozen plasma, and cryoprecipitate. After a discussion with the family the following day, the patient was placed on comfort care measures and died later that day.

Table 1
Comparison of pleural fluid and ascitic fluid at different time points. Notice that the initial pleural fluid on 9/5 is transudative according to Light’s criteria whereas 9/15 pleural fluid is exudative.

| Date   | Fluid Type | 9/5  | 9/6  | 9/15 | 9/18 |
|--------|------------|------|------|------|------|
| Clarity|            | Asitic| Asitic| Pleural| Pleural|
| Color  |            | Orange| Orange| Orange| Orange|
| Lymphocyte (%) |          | 30   | 56   | 51   | 76   |
| Monocyte (%)    |          | 44   | 9    | 39   | 4    |
| Neutrophil (%)  |          | 23   | 33   | 10   | 20   |
| RBC (/μL)       |          | 12,995| 13,800| 92,571| 12,237|
| Total Nucleated Cells (/μL) | | 789  | 206  | 536  | 357  |
|Amylase (U/L)  |          | 75   | 83   | 39   | –    |
|Glucose (mg/dL) |         | 104  | 120  | 136  | –    |
|Pleural Lactate Dehydrogenase (U/L) | | 94   | 88   | 314  | –    |
|Serum Lactate Dehydrogenase (U/L)    |        | 220  | –    | 490  | –    |
|Pleural Total Protein (g/dL)         |        | 1.7  | 1.9  | 2.0  | 1.1  |
|Serum Total Protein (g/dL)           |        | 4    | 4    | 3.7  | 3.5  |

Fig. 1. CT thorax showing right sided Cryptococcus pneumonia with mediastinal and hilar lymphadenopathy. There is a massive left sided pleural effusion that was present despite a therapeutic thoracentesis performed the previous day removing 1.5 L of transudative fluid.

Fig. 2. Gram stain of the culture from pleural fluid. The insert depicts a budding cryptococcal yeast cell.
afternoon.

3. Discussion

*C. neoformans* is an opportunistic and pathogenic fungus usually acquired by inhalation of yeast or basidiospores from environmental sources such as bird droppings and soil [9]. The infection may remain in various sources such as bird droppings and soil [9]. The infection may remain in the lungs, but in immunocompromised states such as HIV, the fungus can disseminate to the central nervous system, skin, and other parts of the body [10]. An immunodeficient state associated with Cryptococcus is liver cirrhosis, which was the likely risk factor of our patient. There are various *Cryptococcus* presentations in cirrhotic patients, including meningitis, pneumonia, peritonitis, pleuritis, and disseminated infection [7, 8, 11-15]. We present a case of disseminated *Cryptococcus* infection who initially presented with a transudative pleural effusion that was antigen-positive and culture-positive for *C. neoformans*. To our knowledge, this is the first case report of a transudative pleural effusion with a positive *Cryptococcal* Ag and culture.

Disseminated *Cryptococcus* is based on positive cultures from any two organ sites or positive blood culture [12,13,16]. Our patient’s pleural fluid, peritoneal fluid, cerebral spinal fluid (CSF), and blood grew *C. neoformans* at various points in time (Fig. 2). Although the infection source was not identified in our patient, there are two possibilities: lungs or gastrointestinal (GI) tract. Cirrhotic patients are susceptible to increased gut permeability of microorganisms, including fungi, into the bloodstream due to damage to the gut-associated lymphoid tissue [12]. The immunocompromised state that cirrhotic patients experience is called cirrhosis-associated immune dysfunction (CAIDS). CAIDS denotes systemic failure of innate and adaptive immunity [4,12,13,17,18]. This immune dysfunction could explain how subclinical *Cryptococcal* pneumonia and pleural effusion seen in our patient proceeded quickly to disseminate infection.

Cirrhotic patients with *Cryptococcus* have a high mortality rate. In one study, all patients with liver cirrhosis died within the first month after the diagnosis of *C. Neoformans* [19]. A delay in the diagnosis of disseminated disease may be a contributing factor to increased mortality rates. The presentation of *Cryptococal* infection in individuals with liver disease is highly variable. Some patients are asymptomatic, while others rapidly progress to respiratory failure. Presentation variability may contribute to a delay in diagnosing this patient population and ultimately impact their mortality rates [20]. Despite their increased risk, these immunocompromised individuals can still be adequately treated with 400 mg of fluconazole daily for 6–12 months for mild to moderate Cryptococcal pneumonia [5]. At the first sign of *Cryptococcal* meningoitis or disseminated disease, the patient should be transitioned to amphotericin (IV 0.7–1 mg/kg/day) +/- flucytosine (100 mg/kg/day) for two weeks followed by fluconazole (400–800 mg/d) for 8 weeks and then fluconazole (200 mg/d) for 6–8 months [5].

In this case, the initial positive *Cryptococcal* Ag’s significance in the transudative effusion was unclear. Transudative effusions are not typically associated with pleural space infections. We considered this a misclassification of the pleural effusion. Although it is more common for Light’s criteria to misclassify transudative effusions as exudative, it is also possible for exudative effusions to be misclassified as transudative [20]. In our case, we speculate the cause of the misclassification to be concomitant hepatic hydrothorax. A less possible possibility is the protein extravasation from the capillaries took several days before the effusion reached exudative values. We also believed meningococcalis was unlikely because the patient had no fevers or headaches, and the initial confusion improved after less than a day of treatment for hepatic encephalopathy. Because of the above reasons, our initial suspicion of disseminated disease was low. Given the patients initial improvement on fluconazole and his AKI, we were hesitant to transition the patient to amphotericin. We determined his condition warranted more aggressive treatment with amphotericin when his positive serum *Cryptococcal* Ag and blood cultures revealed the disease was disseminated. This transition occurred three days after he arrived at our hospital but twelve days since he first presented to the outside hospital.

As seen with previous cases, there is frequently a rapid progression of disseminated *Cryptococcal* disease to multi-organ failure leading to death in cirrhotic patients [12,13]. Our patient also had other comorbidities, including an AKI and a VP shunt, which further complicated his medical management. Ultimately, he passed away after thirteen days of amphotericin B treatment. An aggressive approach to therapy by starting amphotericin sooner may have changed the outcome. This case highlights the prevalence of *Cryptococcal* infection in cirrhotic patients and the importance of recognizing the ailment despite variable presentations such as transudative pleural effusion.

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Consent

Written informed consent was obtained from the patient or legal guardian(s) for publication of this case report and accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal on request.

Written informed consent was obtained from the next of kin since the patient is deceased.

Declaration of competing interest

Please declare any financial or personal interests that might be potentially viewed to influence the work presented. Interests could include consultancies, honoraria, patent ownership or other. If there are none state ‘there are none’.

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