Case of early-disseminated *Rhizopus microsporus* var. *microsporus* mucormycosis in a renal transplant patient

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**Abstract:** Mucormycosis is a rare infection caused by the ubiquitous filamentous fungi of the order Mucorales and class Zygomycetes. These species are vasotropic, causing rapid onset of tissue infarctions and necrosis and subsequent thrombosis by invading vascular bed. The disease spectrum ranges from involvement of skin, sinuses, lung, and brain to disseminated and mostly fatal infections, especially in immunocompromised hosts. Here, we present a case of a fatal disseminated mucormycosis in a 56-year-old female who had deceased donor renal allograft transplantation ~2 weeks prior to presentation. She presented with shortness of breath and dry cough. Despite being on broad-spectrum antibiotics/antifungals and proper management by transplant, infectious disease, and primary team, she died within 3 weeks of admission. Autopsy showed disseminated mucormycosis of lungs and thyroid. Disseminated infection within 2 weeks of solid organ transplantation in this patient was one of the rare features of mucormycosis.

**Keywords:** Zygomycetes, immunocompromised, transplant

**Case**

A 56-year-old black woman from South Carolina presented with complaints of shortness of breath and dry cough for 7 days. She had past medical history of end-stage renal disease status post deceased donor renal transplant nearly 2 weeks ago, diet-controlled diabetes mellitus, hypertension, hypothyroidism, and anemia of chronic disease. Institutional approval by Northshore- LIJ Health and written informed consent from patient was obtained.

She was on maintenance immunosuppressant and prophylactic antibiotics (tacrolimus 1 mg PO Q12h, mycophenolate 500 mg PO Q12h, prednisone 20 mg PO Q24h, valganciclovir 450 mg every other day, and TMP-SMX 1/2 tab Q24h).

On initial physical examination, patient was pale looking but afebrile and on no acute distress. Her vitals on admission were blood pressure 124/74 mmHg, pulse rate 85/min, respiratory rate 22/min, and saturation of 96% on 2 L/min of oxygen via nasal cannula. Her admission labs were significant for white blood cell (WBC) count of 14,000/mm³ with neutrophilic pleocytosis, hemoglobin 7.7 g/dL, and platelet count of 1,55,000/mm³. She had normal electrolytes and liver enzymes. She was found to have acute or chronic kidney injury with serum creatinine of 2.6 mg/dL (baseline creatinine 1.5 mg/dL upon discharge 2 weeks ago). Serum levels of her immunosuppressive drugs were within normal limits. Chest X-ray on admission showed left upper lobe round infiltrate (Figure 1). The patient was treated with oxygen and multiple antimicrobials (vancomycin, aztreonam, and azithromycin) for several days.
All initial cultures including blood and sputum were negative for any bacteria or fungi. On day 5 of antimicrobial therapy, her WBC count decreased to 9,000/mm$^3$. However, she persistently felt short of breath and continued having cough. A repeat chest X-ray showed expanding left upper lobe lesion, while computed tomography scan of chest (Figure 2) done at that time showed a left upper lobe lung lesion with cavitations, along with diffuse bilateral ground glass opacification. Due to continued cough and new left-sided chest pain which she had developed, she went for bronchoscopy with bronchoalveolar lavage and transbronchial biopsy (TBBx) nearly 10 days after admission which revealed septal fibrosis with mild chronic inflammation and focal hemosiderin-laden macrophages.

Primary transplant team added IV micafungin on her empiric antibiotic regimen after the TBBx because of no significant clinical improvement. Her doses of immunosuppressive medications were decreased as well. Samples from bronchoalveolar lavage and TBBx were negative on culture and special stains for any bacterial, mycobacterial, or fungal pathogens.

Post bronchoscopy, she started spiking temperature (101.4°F) and clinically looked more lethargic and confused. Her WBC count increased to 23,000/mm$^3$ (88% neutrophils). The infectious disease team was consulted nearly 2 weeks after initial admission.

The patient underwent thoracotomy with a wedge biopsy at that time. She was started on empirical IV voriconazole. Wedge biopsy results revealed diffuse alveolar hemorrhage, with hemosiderin-laden macrophages and possible capillary fibrin-platelet thrombi. However, silver stain, Gram stain, and acid-fast bacilli stain did not reveal any organisms. Again, multiple cultures for bacterial, mycobacterial, and fungal pathogens were negative. She was transferred to medical ICU after thoracotomy and remained intubated. Serum fungal markers including cryptococcal antigen, beta-D-glucan, and galactomannan antigen were negative.

Despite empiric broad-spectrum antibiotics and antifungals, she remained febrile with persistent leukocytosis, deteriorating renal function, and no improvement in oxygenation. She required five units of red blood cells transfusion post open lung biopsy for persistent anemia. Multiple attempts to wean ventilation failed. On day 20, she had massive hemoptysis followed by hypotension and bradycardia requiring cardiopulmonary resuscitation. However, she died. On autopsy, both of her lungs were heavy, boggy, and congested. There were white lines in the interlobular septa, with the right more prominent than the left, suggestive of lymphatic congestion. Left lung upper and lower lobes had a hemorrhagic appearance throughout, and blood oozed from the cut surfaces. There was a 8×5×2.5 cm well-circumscribed hemorrhagic cavity filled with blood clot in the left upper lobe. The tracheal mucosa had bilateral ulcerative lesions (0.5 cm) directly beneath the vocal cords 1 cm from the midline. Microscopically, the fungi displayed broad-based pauci-septate hyphae with dichotomous wide-angle branching in the left upper lobe hemorrhagic lesion and surrounding lung parenchyma, as well as in the pleura (Figure 3). A focus of thyroid tissue also contained these fungal hyphae (Figure 4). Associated with the fungus was necrosis and vascular thrombosis. Microbiological diagnosis of *Rhizopus microsporus* var. *microsporus* was made.

**Literature review**

Mucormycosis refers to a spectrum of disease presentations caused by ubiquitous saprophytic fungi of the

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**Figure 1** Left upper lobe round infiltrate in chest X-ray.

**Note:** The arrow indicate the cavitary lesion in the lung.

**Figure 2** Computed tomography scan of chest showing left upper lobe lung lesion with cavitations, along with diffuse bilateral ground glass opacification.

**Notes:** The arrow indicate the cavitary lesion in the lung. The inset shows that the cavitary lesion in lung is on that level on coronal view.
class Zygomycetes. It is a severe invasive opportunistic fungal infection that usually occurs in patients who are in immunocompromised states as HIV, diabetes, neutropenia, organ transplant, malignancy, or prolonged use of steroids. Concomitant diabetes mellitus in organ transplant recipients is the most important risk factor in developing this infection. Incidence of mucormycosis has been increasing over several years, due to increasing use of immunosuppressives and antifungals lacking anti-Mucorales activity such as caspofungin or voriconazole. Invasive fungal infections in stem cell transplant recipients account for 7.2%–8% and ∼2% in recipients of solid organ transplantation (SOT).

It is an extremely rare condition in renal allograft recipients with an incidence of 0.2%–1.2%. In the multicenter TRANSNET study, it was found that the incidence of mucormycosis was 0.07% in SOT recipients in 1 year. These patients are more prone to develop opportunistic infections due to their immunosuppressed state. One study reported that 100% of patients received immunosuppressive, and 78.9% received a cumulative steroid dose (≥600 mg of prednisone) therapy before diagnosis of zygomycosis. Our patient had several important risk factors as being an organ transplant recipient on immunosuppressive regimen, concomitant diabetes, and glucocorticoid treatment as part of her antirejection medication regimen. However, it is not clear why she had invasive fungal infection so early after transplant. Main cause of early-onset nosocomial zygomycosis can be due to donor-derived infection. In a recent review on 169 cases of health care-associated mucormycosis, 14% had graft-transmitted infection, representing 60% of SOT recipients. Environmental contamination or nosocomial source has been occasionally found to be transmitting filamentous fungi in SOT recipients. This usually occurs when there is breach in aseptic barrier while obtaining, transplanting, or implanting organ. Contamination of preservation fluid is rarely shown or suspected to be the mode of transmission of aspergillosis and mucormycosis associated with invasive fungal infections in the recipients. So far, 23 cases of donor-derived opportunistic filamentous fungal infections have been reported in SOT patients (71% Aspergillus sp., 21% Mucorales). A majority (91%) of them were kidney transplant recipients. In our particular case, none of the recipient’s organs from the deceased donor were reported to have invasive fungal infections or unexplained death. Preservative fluid culture was also negative for any fungal pathogens. On the other hand, non-graft-transmitted mucormycosis develops after a median of 5 months (range: 1.5–12 months) in SOT, with a significant earlier occurrence in liver transplant recipients.

In our patient, nosocomial mucormycosis occurred after 3 weeks of transplant, which has never been reported before to best of our knowledge.

Roden et al found that the most common sites of mucormycosis in 61 SOT recipients were the lungs (37%), skin lesions (16%), rhinocerebral infection (16%), sinus disease (15%), gastrointestinal infection (7%), disseminated disease (2%), and other less common sites (7%). Disseminated mucormycosis is defined as an involvement of two or more noncontiguous organs and usually presents with nonspecific manifestation. In a study done on ten SOT recipients with zygomycosis, 12.9% developed disseminated mucormycosis. The majority had lungs involved which was identified on autopsy or with a cavitary radiologic appearance highly suggestive of invasive fungal infection. In our case, involvement of lung and thyroid tissue classifies it as a disseminated infection. The case described is a unique case of
disseminated mucormycosis because only three documented human infections owing to this organism (R. microsporus var. microsporus) have been described so far.14-16 None of the published cases were associated with SOT recipients, however.

In one of the largest study done on mucormycosis in a renal transplant patient, it was found that there is no statistically significant difference in mortality rate between sex, different age, and time from transplant. Conclusion from this 2 decades of study was that mucormycosis after renal transplantation has a poor prognosis, particularly in patients with pulmonary involvement.17

The mortality rate among SOT recipients with zygomycosis has typically ranged from 49% to 71% and up to 96% in disseminated form of infection.18,19 Successful treatment requires a combination of early diagnosis, reversal of underlying risk factors, reduction of immunosuppression, prompt administration of antifungal therapy (amphotericin B, the drug of choice) if possible, and aggressive surgical debridement when applicable. A high index of suspicion is required for rapid and timely diagnosis of invasive zygomycosis, preferably by culture and histological confirmation. Diagnosis of mucormycosis is very difficult due to the rarity of the problem and significant limitations of diagnostic tests. In our particular patient, definitive tests included fungal cultures and lung biopsy. However, both test missed the diagnosis most likely due to sampling bias or inadequate tissue. Many missed cases of these deep-seated fungal infections are diagnosed on autopsy. In fact, autopsy series have demonstrated that up to 50% of cases of mucormycosis are diagnosed postmortem.20 Another study reviewed autopsies with SOT in UCLA Medical Center (Santa Monica, CA, USA), especially focusing on the deep-seated fungal infection, and the incidence of fungal infections, causative fungi, and organs involved were evaluated. Deep-stated fungal infections were demonstrated in 21.0% of the patients with SOT. The incidence of fungal infections was 26.1% in kidney transplantation, which was the highest rate in SOT autopsies. The most common causative fungi were Aspergillus sp., seen in 70.6% of SOT autopsies.21

From our case report, we showed that invasive fungal infection can occur very early after transplant and should be regarded as differential when these patients present with nonspecific clinical feature with no identified source of infection.

Disclosure
The authors report no conflicts of interest in this work.

References