Pulmonary Mucormycosis in an Elderly Patient with Multiple Comorbidities

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Abstract

Pulmonary Mucormycosis, a rare opportunistic invasive pulmonary fungal infection.¹ Mucormycosis is less common than other opportunistic fungal infections, such as aspergillus and candida infections, but mortality rate is much higher. We report a case of 79 year old male patient with multiple comorbidities who presented to Manik Hospital and Research Centre with 1 month history of cough, fever and breathing difficulty and diagnosed to have pulmonary mucormycosis and treated successfully.

Keywords: Pulmonary mucormycosis with multiple comorbidities, diagnosis and treatment.

Introduction

Mucormycosis is an opportunistic fungal infection caused by fungi of order mucorales in the class zygomycetes. Also known as zygomycosis, it is commonly caused by fungal species of genera Rhizopus, Lichtheimia and mucor.² Mucormycosis is manifested by a variety of different syndromes, particularly in immunocompromised patients and those with diabetes mellitus. Diabetes remains the most common risk factor; however zygomycosis has increased among transplant recipients and patients with hematological malignancy.³ Although most fungal infections, including mucormycosis, are acquired via inhalation, endobronchial manifestation is rare, and the usual presentation is either pulmonary or systemic disease. Diabetes is common predisposing condition for mucormycosis.⁴ Rhino-orbital-cerebral and pulmonary mucormycosis are acquired by inhalation of spores. In healthy individuals, cilia transport these spores to the pharynx and they are cleared through gastrointestinal tract. In susceptible individuals, infection usually begins in nasal turbinates or the alveoli.⁵ The agents of mucormycosis are angioinvasive; thus, infarction of infected tissues is a hallmark of invasive disease.⁶

Amphotericin B is polyene antifungal drug and is the selected drug for the treatment of mucormycosis.⁷ Drug concentration of Amphotericin B in the lung is lower than other tissues. Therefore pulmonary mucormycosis infection requires a higher dose of Amphotericin B. Intravenous (IV) amphotericin B (lipid formulation) is drug of choice for initial therapy.⁸ Liposomal Amphotericin B is safer, with an improved effect and smaller dose, compared with amphotericin B alone.⁹ Posaconazole used as a stepdown therapy for those patients who have responded to amphotericin B. Posaconazole is
broad spectrum azole that is active in against the agents of mucormycosis and that are available in both parenteral and oral formulations.\textsuperscript{10,11} Treatment of mucormycosis involves a combination of surgical debridement of involved tissues and antifungal therapy.\textsuperscript{12} Aggressive surgical debridement of involved tissues should be undertaken as soon as diagnosis of any form of mucormycosis is suspected. There are reports of patients with early pulmonary infection who were cured with lobectomies.\textsuperscript{13,14}

Herein we report a case of isolated pulmonary mucormycosis presented as non resolving pneumonia in patient with diabetes mellitus, chronic kidney disease, coronary artery disease (post CABG) who was successfully managed with medical therapy.

**Case Report**

A 79 yrs old patient presented in OPD with history of cough with expectoration of purulent yellowish sputum since 1 month, intermittent fever since 3 weeks and breathing difficulty on exertion since 2 weeks. His past history reveals diabetes mellitus and hypertension since 10 yrs, ischemic heart disease (post coronary artery bypass surgery 6 yrs ago) and he was diagnosed to have diabetic nephropathy since 2 yrs. He was on regular medications & follow up for the same. He was nonsmoker and no history of any allergy in past. His X ray chest was done which showed inhomogeneous patchy opacities in right upper & midzone. CBC revealed leukocytosis. His probable diagnosis was community acquired pneumonia. He was treated with oral antibiotics (cefpodoxime 200 mg BD + Azithromycin 500 mg OD) for seven days. However patient returned to OPD reporting no relief of cough and fever and increased breathlessness. Hence he was admitted in Hospital.

Clinical Examination on admission revealed:

- **Temp**: 37.6°C
- **Tachycardia**: rate 114bpm
- **Tachypnea**: rate 30/min
- **BP**: 140/90 mm of Hg
- **Spo2**: 95% on room air

RS- chest auscultation revealed decreased breath sounds and scanty Crackels in right supraclavicular and mammary area.

Cardiovascular, central nervous system and abdominal examination was within normal.

Lab investigation on admission:

- CBC - (Hb)-8.9 gm% (TLC)- 13000 platelet -526000
- ESR- 68mm/l
- Blood urea- 33.9 Sr creatinine-1.59
- Urine- Alb+
- RBSL- 456 mg%
- Na-135 K-4.6
- ABG- PH-7.42, PO2- 78 mm hg, PCO2- 44

Sputum exam- Moderate number of gram-positive cocci as well as occasional gram-negative bacilli,
- AFB : -ve
- Gram positive cocci ++
- Gram negative bacilli ++

Patient was started with intravenous cephalosporins -
- Control of Blood sugar with short acting insulin , and other supportive measures.

However, over next 48 hour, fever not reduced and expectoration and cough persisted. Patient reported minimal increase in dyspnea.

**Chest X Ray**

![Figure 1 Chest X ray showing right upper lobe haziness](image-url)
Blood reports after 2 days
Hb - 9.2 gm%
TLC-30,500
Platelet-4,37,000
Urea-60.9
Creatinine-2.13

Hence we decided to do HRCT chest and Bronchoscopy. HRCT chest showed large hypodence lesion in right upper lobe with surrounding collapse and foci of air.

![HRCT chest image](image1)

**Figure 2.** Chest CT showing large hypodence lesion in right upper lobe with surrounding collapse and foci of air within. consolidation in right upper lobe and middle lobe.

Bronchoscopic biopsy showed- both showing fungal hyphae, anthracotic pigment.

Stain- H E 40x×10x

![Bronchoscopy image](image2)

**Fig. 3-** Bronchoscopy showing fungal hyphae, anthracotic pigment.

His ENT examination and cerebral imaging was normal.

Hence a diagnosis of invasive pulmonary fungal infection-Mucormycosis was established. This was confirmed by the fungal staining & histopathology report of BAL.

Now we started patient on Inj. Liposomal Amphotericin -B with dose adjustments done according to his renal status. His creatinine clearance was 30.23 ml/min

Over next 14 days he showed gradual clinical improvement. His fever reduced, cough and expectoration reduced. Blood sugars were normal on 3 subcutaneous short acting insulin with meals.

His lab after 14 days
Hb-9.8 gm%
TLC-10800
Platelets-515000
Urea- 45.7
Creatinine- 1.78
RBSL- 184

However his chest X ray & repeat HRCT revealed same picture after 14 days. At these pt was given option of surgical intervention along with antifungal agents. However he refused for surgery because of multiple comorbidities and age.

Hence we decided to add second line antifungal agent POSICANAZOLE- mg/kg/day. i.e. 5 ml BID in syrup form. We continued POSICANAZOLE for one month. During this patient showed good clinical improvement. After
one month. His HRCT showed complete resolution of lung lesions. His blood reports at the time of discharge -
Hb-9.9%
TLC-9800
Platelet-490000
FBSL-128
PPBS-194
Urea-43.2
Creatinine-1.56
Thus after one & half month of antifungal therapy (1st + 2nd line antifungal agents) patient showed complete clinical and radiological clearance of pulmonary mucormycosis. He is on regular follow up till date and asymptomatic.

3) Patient had shown only partial clinical response in 14 days of 1st line drug i.e. inj Liposomal Amphotericin-B. However complete response is achieved when treated with 2nd line antifungal agent. Hence we would like to mention use of 2nd line drugs if 1st line drugs fail to achieve recovery.

References
7. Li WF, He C, Liu XF, Wang SY, Qu JL, Lin ZF. A diagnosis neglected for 6 years; Report of a misdiagnosed case of pulmonary mucormycosis and review of

We are reporting this case for following reasons:

1) Incidence of Isolated Pulmonary Mucormycosis is rare.
2) Mortality rate in invasive pulmonary fungal infection is high when associated with comorbidities like diabetes, ischaemic heart disease and Nephropathy. Hence we were successful in complete recovery of patient.


