Rhino orbital mucormycosis – A case series

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Abstract
Mucormycosis is a fulminating, opportunistic and often a fatal mycotic infection. Its incidence is commonly observed among immune suppressed patients. We are reporting 3 cases of Rhino-orbital mucormycosis with central retinal artery occlusion. All 3 were males in the age range of 50-70 years. All were diabetic with similar clinical presentation; they were treated with I.V Amphotericin B along with control of blood sugars. Although the general outcome was good in all the three patients the visual prognosis remained poor.

Keywords: Central retinal artery occlusion, Diabetes, Immunocompromised Mucormycosis, Rhino-orbital.

Introduction
Mucormycosis is a relatively rare and fatal infective condition. The causative agents belong to the Mucoraceae family, Absidia, Mucor, and Rhizopus which may often be seen in decaying organic matters like food soil and excreta of animals.¹ They have ability to multiply rapidly and release numerous spores that become airborne and which can gain access to humans either by inhalation or ingestion. The spores which enter the person with normal immune system are usually killed by phagocytic action, and rarely immunocompromised people become a victim of this infection.² In India the incidence of mucormycosis was observed to be 0.14 cases per 1000 population.³ In US the incidence was reported to be 20 per 100,000 cases with 0.7% among autopsied patients.⁴ Among US population the cumulative 1 year incidence of Mucormycosis was reported to be 4 per 1000 stem cell and 0.6 per 1000 solid organ transplantation cases respectively.⁵

Rhino-Orbito-cerebral mucormycosis is often observed with immunocompromised state, like blood dyscrasia, malignancy, diabetes mellitus with poor glycemic control, prolonged corticosteroid therapy, post organ transplantation and in severe debilitating conditions like burns and trauma⁶ As rhino- orbito-cerebral mucormycosis occurs rarely, it may pose a challenge for clinicians to diagnose and treat this fatal condition. Early identification of clinical presentation followed by aggressive antifungal therapy, debridement and correction of immunocompromised state is very important to reduce the mortality rate in this condition⁷. Ocular morbidity commonly seen in mucormycosis is central retinal artery occlusion when there is orbital involvement, which may often lead to irreversible blindness. The disease presents often with clinical symptoms resembling facial or orbital cellulitis with rapid progressive clinical course which may end in fatal outcome if not managed properly. Hence the authors decided to write up these cases to resolve the diagnostic dilemma among clinicians treating these patients.

Case Report 1
A 70 year old male with diabetes mellitus for five years and coronary artery disease presented with complaints of fever, right side facial swelling and numbness for 1 month. He had history of sudden painless loss of vision in the right eye. On ocular examination of right eye there was no perception of light, lid edema, ptosis, raised tear film height, grade four relative afferent pupillary defect and restriction of extraocular movements in all quadrants. Fundus showed Mild disc edema, cattley tracking of vessels suggestive of a Central retinal artery occlusion (CRAO), macula showed cherry red spot, background retina was edematous. Based on the presenting features and clinical findings a clinical diagnosis of rhino-orbital-cerebral mucormycosis was made with differential diagnosis of orbital cellulitis, maxillary sinusitis and superior orbital fissure syndrome. Paranasal sinus debridement with orbital decompression was done. Histopathological investigation of debrided tissue sample revealed angioinvasive fungal filaments confirming the diagnosis of Mucormycosis. (Fig. 1 & Fig. 2) Treatment was initiated with IV Liposomal Amphotericin B 0.5mg/kg in view of right maxillary sinus fungal involvement with orbital invasion. In addition, patient’s cardiac status was evaluated, echocardiography was done which identified vegetation of 1x6mm attached to basal chordae near the left ventricular apex suggestive of infective fungal endocarditis which could have led to CRAO. The general status of the patient improved although the visual prognosis remained poor.

Fig. 1: Histopathological staining showing broad aseptate fungi
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Case Report 2
A 54 year old male with diabetes mellitus for fifteen years and chronic kidney disease, presented with complaints of right side facial swelling and pain with sudden loss of vision right eye. History of brownish nasal discharge occasionally blood stained for one week. History of pulmonary tuberculosis 10 years back, which was treated. On examination, there was no perception of light in right eye with periorbital edema, proptosis and conjunctival chemosis (Fig. 3) with grade four relative afferent pupillary defect and restriction of extraocular movements in all quadrants. Fundus showed pale disc with well defined margins, cattle tracking of vessels suggestive of a Central retinal artery occlusion (CRAO) (Fig. 4), macula showed cherry red spot, background retina was pale. Oral examination revealed ulcer in the hard palate (Fig. 5). Computed tomography of Paranasal Sinus showed involvement of nasal cavity, maxillary, anterior and posterior ethmoid sinus (Fig. 6). Potassium hydroxide mount done showed Broad Aseptatic Hyphae, Zygomycetes with branches confirming the diagnosis of mucormycosis. Medical management involved initiation of Intravenous liposomal Amphotericin B 0.5mg/kg followed by Endoscopic debridement of mucormycosis with palatal fenestration. The general status of the patient improved although the visual prognosis remained poor.

Case Report 3
A 71 year old male with history of diabetes mellitus for 10 years and hypertensive for the past 3 years presented with the complaints of intractable pain in and around the orbit with sudden loss of vision in the right eye. Ocular examination of right eye revealed no perception of light, periorbital edema, proptosis, conjunctival chemosis with grade 3 relative afferent pupillary defect and restriction of extraocular movements in all direction of gaze. Fundus examination revealed pale disc with well-defined margins, cattle tracking of vessels suggestive of a Central retinal artery occlusion (CRAO), macula showed cherry red spot, background retina was pale. Oral examination revealed a 2 x 4mm black necrotic ulcer over hard palate. MRI revealed Right sided

Fig. 2: Gomori silver staining

Fig. 5: Hard palate lesion

Fig. 6: CT of Paranasal Sinus showed involvement of nasal cavity, maxillary, anterior and posterior ethmoid sinus

Fig. 4: Central Retinal artery occlusion

Fig. 3: Right eye with conjunctival chemosis

Fig. 3: Right eye with conjunctival chemosis

Fig. 5: Hard palate lesion

Fig. 6: CT of Paranasal Sinus showed involvement of nasal cavity, maxillary, anterior and posterior ethmoid sinus

Fig. 4: Central Retinal artery occlusion
pansinusitis with no direct extension to orbit and an 18x7mm defect in the hard palate. Subsequent treatment involved Intravenous liposomal Amphotericin B 0.5mg/kg followed by endoscopic debridement of mucormycosis with palatal fenestration. Tissue culture of debrided sample showed Rhizopus Arrhizus growth confirming the diagnosis of mucormycosis. The general status of the patient improved although the visual acuity remained poor.

Discussion

Mucormycosis is an uncommon fungal infection which is invasive in nature with potential fatal outcome. Because mucormycosis is not a notifiable disease and the incidence is relatively rare, the real data on incidence is difficult to estimate. There are no clear demographic factors which predispose individuals to this condition. Reviews from previous studies show a similar male to female distribution; however a review retrospectively for the past 30 years of all published cases of pulmonary mucormycosis by Lee et al had a male-to-female ratio of 3:1, which is similar to our series where all three patients were male.

Mucormycosis is commonly reported among patients with poor glycemic control, immunosuppressive conditions like HIV, Malignancy and post organ donation, because of their immunocompromised state. In our series all the three patients were chronic diabetics with poor compliance to treatment which may have predisposed to the development of the disease. The possible mode of transmission of this condition was by inhalation of fungal spores into paranasal sinus and extending into adjacent tissues. Upon germination the fungus invade the adjacent structures like palate, sphenoid sinus, orbit and finally the brain. The initial presentation is usually nonspecific in rhino-orbital-cerebral mucormycosis. It usually presents with eye or facial pain, numbness followed by conjunctival chemosis and eventually loss of vision. Elevated white blood count is seen in patient with functional bone marrow. Fever may not be a typical symptom since in half of the patients it is not seen. If untreated, the infection may spread to the orbit, which can result in compromised extraocular muscle function, proptosis and chemosis.

Similar results were seen in case series published by Bavikar et al and Kim JG et al where patients presented with facial swelling associated with pain, numbness and loss of vision. Prompt diagnosis of rhino orbital mucormycosis along with aggressive treatment are vital for good prognosis of the patient. Symptoms correlating with mucormycosis in a susceptible individual should call for immediate initiation of treatment while necessary steps are taken towards confirmation of mucormycosis. The presence of branching aseptate hyphae in debrided tissue in histopathological examination may confirm the clinical diagnosis, along with fungal cultures for further confirmation.

Imaging studies in the early stages of the disease may not be helpful. In correlation with the clinical finding CT scan may also be used to detect the progression of the disease. Soft tissue involvement of fungal infection can be accurately identified by MRI. For determining the progression of the disease and surgical intervention both CT and MRI scans should be frequently obtained. Management of rhino-orbital-cerebral mucormycosis involves addressing three key issues: correction of underlying predisposing factors, early initiation of antifungal therapy and appropriate surgical intervention.

Although amphotericin B deoxycholate is the drug of choice for mucormycosis, lipid formulations of amphotericin B are considered to be very effective. Recommended initial doses for the lipid formulation of amphotericin are 5-7.5 mg/kg/day, up to 10 mg/kg/day for possible involvement of CNS.

Debridement of necrotic tissue along with Antifungal therapy is important for patient survival. In rhino orbital mucormycosis, surgical care includes drainage of the sinuses and even removal of the orbital contents. Repeated surgery may be required, especially for rhino-orbital mucormycosis.

Conclusion

Rhino orbital mucormycosis is a rare disease with fatal outcome. Patients presenting with symptoms like pain over face with facial swelling, and orbital involvement along with predisposing conditions should routinely be examined for mucormycosis. Early diagnosis, prompt antifungal therapy, surgical debridement and reversal of underlying predisposing conditions is the most important criterion for conservation of orbit and thereby vision of the patient.

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References


