Rapidly progressive fatal orbito-cerebral mucormycosis in an immunocompetent patient following entry of an insect into his eye; a case report and literature review

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Abstract
Mucormycosis, an uncommon and rapidly progressive fungal infection, is mainly seen in immunocompromised patients. However, immunocompetent individuals can also be infected. Here, we present a rapidly progressive orbito-cerebral mucormycosis in an immunocompetent patient, who had a history of entry of an insect into his eye. We report an immunocompetent patient, who had a history of entry of an insect into his eye, presented with left eye panophthalmitis, cellulitis and loss of vision. He underwent orbital exenteration. Then, mucormycosis was diagnosed and amphotericin B was initiated. Leptomeningeal involvement was found in brain magnetic resonance imaging (MRI) and cerebrovascular accident (CVA) was diagnosed. Surgical debridement of necrotic tissue was performed, and postoperatively, our patient died following CVA. Mucormycosis, as a fatal fungal infection, requires early diagnosis, appropriate early surgical and rapid antifungal management so that successful outcome can be achieved.

Introduction
Mucormycosis, an uncommon and rapidly progressive fungal infection, from the Mucorales order, is mainly seen in immunocompromised patients (1). It should be noticed that immunocompetent individuals can also be infected (2). Rhino-orbito-cerebral, cutaneous, pulmonary, gastrointestinal, and disseminated infections are considered to be the clinical presentations of mucormycosis (3). Rhino-orbito-cerebral mucormycosis begins when the fungus affects the nasal mucosa and paranasal sinuses (4). Orbital involvement develops when the orbital wall is invaded through the paranasal sinuses (5), while intracranial invasion is found to occur following progressive ocular involvement (4). We present a rapidly progressive orbito-cerebral mucormycosis in an immunocompetent patient, who had a history of entry of an insect into his eye.

Case Presentation
A 34-year-old white male patient, who had no evidence of immunodeficiency, and no considerable past medical history, presented with left eye panophthalmitis, cellulitis and loss of vision. He had an episode of entry of an insect to his left eye while riding on a motorcycle 25 days earlier. In emergency setting chloramphenicol eye drops BP 0.5% was administered for him, since no remission was observed after a week. Then he was hospitalized and received high corticosteroid doses accompanied by intravenous (IV) antibiotic, while we do not have any documentation of drug and dosage. He underwent ethmoidectomy and exploration of the roof of the orbital cavity in that setup however, they reported no evidence of pus or inflammation. Lateral canthotomy and inferomedial decompression of the orbit was performed following the first surgical procedure. Bloody discharge was seen after the operation.

Then, he was referred to our clinic with left eye swelling, proptosis and chemosis (Figure 1). Laboratory investigations revealed; white blood cell (WBC) $12 \times 10^9$ per liter, and erythrocyte sedimentation rate (ESR)
of 30 mm/1st hour. Vancomycin 1000 mg/kg/dd IV and meropenem 1500 mg/kg/d IV were initiated. Exenteration of the left eye was planned due to panophthalmitis, cellulitis and loss of vision. Color change of floor of the orbit and frontal recess was found after exenteration (Figure 2). Color-changed material was resected and sent to pathology lab. The results of pathology showed mucormycosis (Figure 3). Axial and coronal computed tomography (CT) scan of the orbits showed maxillary antrum involvement, erosion of adjacent bones with spread to lamina papyracea, nasal cavity, and inferior orbital fissure (Figure 4). Amphotericin B 50 mg/kg/d and caspofungin 50 mg/m² daily were added to the antibiotics. Three days after the surgery, the patient was taken to an intensive care unit (ICU) following gradually loss of consciousness. Brain magnetic resonance imaging (MRI) showed left cavernous sinus and internal carotid artery involvement and then cerebrovascular accident (CVA) was diagnosed. After a while, surgical debridement of necrotic tissue was performed. Postoperatively, our patient died following CVA.

Discussion
Mucormycosis, a fungal disease caused by zygomycetes, is appeared to have an acute and fulminant course (5). Although it is an uncommon infection, its incidence is rising (6, 7). Mucormycosis usually presents as pulmonary, gastrointestinal, disseminated or rhinocerebral form (8). Earlier publications reported that the pulmonary mucormycosis is the most common form (2). However, a recent study reported the rhino-orbital form as the most frequent (4). Orbital involvement includes infectious spread into the orbital cavity through the paranasal sinuses. This infection can result in vascular thrombosis, including cavernous sinus thrombosis (1, 4, 9), and ischemia of near tissues, following attachment to the endothelial cells of the vessels (10).

Immunocompromised individuals like patients with malignancies of hematological diseases, long-term corticosteroid consumption, immunosuppressive therapy and HIV infection, are mostly infected. However, there are reports in which immunocompetent individuals are also infected (2,11-13). Several reports have documented the orbital infection in immunocompetent patients (2, 13,14). Rahman et al reported a healthy patient of rhino-orbital mucormycosis who was presented with massive necrosis of the maxilla with retro-bulbar and infra-bulbar involvement. The patient underwent extensive surgical debridement and received medical treatment. However, he
did not survive and died after ten days (2). Another report of an orbital mucormycosis in a healthy patient, with a history of entry of dust into his left eye and presentation with swelling and redness of the eye, showed that orbital exenteration and intravenous amphotericin B led to the patient’s survival (13). However, up to our knowledge, there are few reports of immunocompetent patients who presented with orbital-cerebral mucormycosis. Angali et al reported a case of rhino-orbito-cerebral mucormycosis, who presented with a non-healing ulcer on the face and orbital involvement. Their patient died following severe seizures and status epilepticus (15).

There are two cases of orbital mucormycosis which developed following a traumatic laceration from a tree branch to the temporo-parietal scalp and entry of dust particle to the eye (3, 13). However, up to our knowledge, there is probably no report of orbito-cerebral mucormycosis following entry of insect into the eye. Our patient had a history of entry of an insect to his left eye and was referred with left eye swelling, proptosis and chemosis. He underwent orbital exenteration. Then, mucormycosis was diagnosed and amphotericin B was initiated. Left cavernous sinus and internal carotid artery involvement were found in brain MRI and CVA was initiated. Surgical debridement was performed, and postoperatively, our patient died following CVA. Patient’s survival is reported to exceed 80% when early medical and surgical treatment is performed (16). Badiee et al concluded that effective management of mucormycosis consists of early diagnosis of the disease (13). Similarly, some other reports indicated that early diagnosis and treatment is considered to be important for good prognosis (3,4,15).

Conclusion
Mucormycosis, as a fatal fungal infection with high mortality rate, requires early diagnosis, appropriate early surgical and rapid antifungal management so that successful outcome is possible. Intracranial involvement is still poor prognosis.

Conflicts of interest
The authors report no conflicts of interest.

Ethical considerations
Ethical issues (including plagiarism, data fabrication, double publication) have been completely observed by the authors. The patents of the patient gave her informed consent regarding the publication of this case report.

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References