SUBCUTANEOUS PHYCOMYCOSIS: A CASE REPORT
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ABSTRACT: Subcutaneous phycomycosis is a rare fungal infection of the subcutaneous tissue caused by Zygomatous group of fungi. We report an rare case of an 60 years old farmer with nasal subcutaneous phycomycosis which presented as painless, non-tender, swelling on his nose. Local examination revealed a non-tender, dull, erythematous, woody hard, uniform, smooth, non-pitting swelling on the root of the nose, extending to the upper lips, left and right cheek. The overlying skin was intact, and a finger could be insinuated beneath the swelling. Histopathological analysis of the biopsied tissues revealed chronic granulomatous inflammation containing fungal hyphae with surrounding langhans giant cells and eosinophilia. The swelling decreased in the second month of treatment after the patient was started on oral itraconazole.

KEYWORDS: Phycomycosis; zygomatous fungi; potassium iodide; Azole.

INTRODUCTION: Subcutaneous phycomycosis is a rare, chronic, localized, subcutaneous zygomycosis, characterized by painless, woody swelling of the rhinofacial region. The disease occurs mainly in the tropical rain forests of Africa, South-East Asia. A few cases have been reported from India. We report a case of this rare subcutaneous phycomycosis.

CASE REPORT: A 60 year old farmer presented to opd with a slowly progressive swelling of the nose, forehead, upper lip and nasal block of 2 year duration. He was treated earlier with various antibiotics, with no improvement.

General and systemic examinations were unremarkable, except for a disfigured facial appearance. Local examination revealed a non-tender, dull, erythematous, woody hard, uniform, smooth, non-pitting swelling on the root of the nose, extending to the upper lips, left and right cheek and forehead. The overlying skin was intact, and a finger could be insinuated beneath the swelling. There was no regional lymph node enlargement.

The patient did not have a history of fever, vomiting, and his history was not suggestive of any significant illness

Routine blood examination was normal. X-ray of paranasal sinuses showed features of frontal and maxillary sinuses.

The biopsied tissue was sent for potassium hydroxide preparation and fungal culture. Broad thin walled non-septate mycelia were found in the KOH preparation.

The patient was treated with capsul itraconazole 200 mg twice a day for 3 month with remarkable improvement. After 3 months itraconazole dose tapered to 100 mg twice day. Patient is still on follow up.

DISCUSSION: Subcutaneous phycomycosis is a fungal infection of the subcutaneous fats caused by Zygomatous group of fungi in the family Entomophthoraceae, mainly in the genus Basidiobolus (common in children) and Conidiobolus in adults (Prasad PV, 2002).¹ The causative organisms are
Basidiobolus ranarum and Conidiobolus coronatus respectively. It can cause a variety of clinical manifestations including subcutaneous zygomycosis, gastrointestinal zygomycosis and occasionally an acute systemic illness.

Documentations as early as 1886 describe this fungus as a saprophyte in the gastro-intestinal tract of reptiles and amphibians. Soil and vegetation contaminated by faeces from these animals seems a likely source of the infection in man (JOE HF, 1980). Reports have shown fungal spores to be found on the bristles of mites other insects. When these are eaten by reptiles and amphibians, the saprophytic cycle of the fungus is complete. Such mites and insects may comprise a further possible source of infection in man (HC, 1999).

Granulomatous lesions due to Basidiobolus ranarum have been found in the nostrils and on the legs of horses, but reports on animal infection are scanty. In humans a chronic infection of the subcutaneous tissue is usually produced characterized by the formation of firm and non-tender swellings, generally on the extremities, trunk and rarely other parts of the body. Our patient is among the rare cases as he presented with a nasal swelling. The disease appears to be a primary rather than a secondary infection, no predisposing factors are known (HC, 1999).

Granulomatous inflammatory features may also be present (Manjir RN, 2009).

Most patients with phycomycosis respond very well to oral potassium iodide therapy as also to azoles (Sujatha S, 2003). Our patient also responded well to itraconazole and the swelling reduced dramatically after one month of treatment. Relapse chances are high in patient who undergo excisional biopsy (Manjir RN, 2009). Hence surgery should be avoided as it may hasten the spread of infection.

Treatment of subcutaneous phycomycosis is difficult because the diagnosis is usually established late, but patients often respond to oral potassium iodide, oral itraconazole (200 to 400 mg/day), ketoconazole (200 to 400 mg/day), fluconazole (100-200 mg/day), amphotericin-B, and cotrimoxazole. Of these, itraconazole and fluconazole are both effective and relatively safe. Treatment should be continued for at least 1 month after the lesions have cleared.

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CASE REPORT

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