Subcutaneous phaeohyphomycosis: A rare case

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Abstract
Phaeohyphomycosis is a rare fungal infection caused by melanized fungi which usually involve the skin and subcutaneous tissue. It is an infrequent cause of infection in human beings. Common clinical manifestations are subcutaneous cystic swellings, which is localized, characterized by subcutaneous asymptomatic nodular lesions that develop after traumatic implantation of fungus from contaminated soil, thorns or wood splinters, especially on extremities. We report a case of subcutaneous phaeohyphomycosis presenting as multiple cystic swellings on the left lower leg within a week. The diagnosis of this case is based on histopathology examination of clinical specimens and identification of cultures obtained through biopsy material. We report a case of subcutaneous phaeohyphomycosis in immunocompetent male patient, presenting as multiple cystic swellings over the left lower leg with unknown predisposing factors. Further investigation revealed granulomatous inflammation on histopathology examination and black-coloured colonies identified as Exophiala dermatitidis on fungal culture. This case of phaeohyphomycosis was successfully treated by itraconazole monotherapy without surgical excision and still under follow-up.

Introduction
Phaeohyphomycosis is the general term to describe fungal infection caused by various heterogenous groups of dematiaceous or melanized fungi that contain melanin in their cell walls. It shows melanized filaments in the tissues; therefore, it is defined histopathologically rather than clinically by the presence of brownish septate hyphae, pseudohyphae, yeasts or combination of these forms. It is an infrequent cause of infection in human beings and covers a wide variety of clinical forms, depends on the immune status of the host: superficial (tinea nigra and black piedra); cutaneous (scytalidiosis) and corneal; subcutaneous (mycotic cyst); and systemic in the immunocompromised host. It can be responsible for life-threatening infections in both immunocompromised and immunocompetent individuals. Etiologic agents include Exophiala, Phoma, Bipolaris, Phialophora, Colletotrichum, Curvularia, Alternaria, Exserohilum, and Phialoconium sp. The most common are Exophiala jeaneselmi and Exophiala dermatitidis (formerly Wandiella dermatitidis). These organisms are widespread in the environment, being found in soil, wood, and decomposing plant debris. They are distributed worldwide and are more common in tropical and subtropical climates. The involvement of the skin and subcutaneous tissue is more frequent than any other organs and termed as subcutaneous phaeohyphomycosis. Common clinical manifestations are subcutaneous abscesses or cystic swellings. The mycotic cyst is a localized form, characterized by subcutaneous asymptomatic nodular lesions that develop after traumatic implantation of fungus from contaminated soil, thorns or wood splinters, especially on the extremities. The diagnosis of this case is based on histopathology examination of clinical specimens and identification of cultures obtained through biopsy material. We report a case of subcutaneous phaeohyphomycosis in immunocompetent male patient, presenting as multiple cystic swellings over the left lower leg with unknown predisposing factors that was firstly misdiagnosed as soft tissue tumor and subcutaneous tuberculosis.

Case Report
A 25-year-old male patient presented with multiple large painful swellings on the left lower leg, which was present for about 8 months (Figure 1B, 1C). The patient works as a teacher and farmer. There was no history of trauma, no underlying disease and similar family history, and no suspicion of immunocompromised status. At first, the swelling was quite small in size, slightly itchy, and not painful that lead to misdiagnosing as soft tissue tumor (Figure 1A). The patient provided a history of two predisposing factors that was firstly misdiagnosed as soft tissue tumor (Figure 1B, 1C). The diagnosis of this case is based on histopathology examination of clinical specimens and identification of cultures obtained through biopsy material. The anti-tuberculosis therapy was discontinued and the patient was referred to tertiary hospital (Dr. Soetomo General Hospital) for further investigation.

On physical examination, the patient appeared to be in good health, was afebrile, and had no symptoms other than multiple dark brown, painful, firm-to-soft, cystic swellings on the left lower leg region. The lesions were coalesced to form a large fluctuant mass measuring approximately 20x15x5 cm in size, presented with ulceration on the surface (Figure 1B, 1C). A differential diagnosis of benign lipomatous neoplasm, cutaneous tuberculosis, and deep fungal infection were considered. Routine laboratory investigation values were within normal limits. X-ray for left lower leg discover soft tissue mass appearance without signs of osteomyelitis. The Potassium hydroxide (KOH) preparation and Gram stain from smear showed no fungal appearance and no bacteria respectively. An excisional biopsy sample was taken from the lateral side of the lesion and sent for histopathology examination and fungal culture. Histopathology analysis showed granulomatous inflammation composed of histocytes, lymphocytes, and fibroblasts.
but could not identify the causative fungal elements in the sections stained with haematoxylin and eosin (HE) and periodic acid Schiff (PAS) (Figure 2D, 2E, 2F). Additional Ziehl-Neelsen (ZN) stain showed no acid-fast bacilli (Figure 2G). Fungal culture from skin biopsy specimens grown on Sabouraud’s dextrose agar (SDA) yielded black-coloured colonies identified as *E. dermatitidis* using lactophenol cotton blue preparation (Figure 2A, 2B, 2C).

Based on those findings, the patient was finally diagnosed as subcutaneous phaeohyphomycosis and treated with oral daily itraconazole 400 mg as monotherapy without surgical excision. The patient was followed monthly in mycology division. The progression of the lesions was dramatic and established successful result. At the end of the first month of treatment, edema began to decrease, oozing stopped, and ulcerated lesions healed (Figure 1D, 1E). At the sixth month of itraconazole treatment, the lesions almost completely regressed but the patient still continue the treatment in order to achieve complete clinical cure (Figure 1F, 1G, 1H).

**Discussion**

Dematiaceous or melanized or darkly pigmented fungi contain a large, heterogeneous group of saprobe microorganisms that present in guanos, plants, and wood in decomposition. They are widely distributed around the world but more frequent in subtropical and tropical climates. They have been associated with a variety of clinical syndromes that grouped into three types based on the histopathologic findings i.e., eumycetoma, chromoblastomycosis and phaeohyphomycosis. In histopathology examination, eumycetoma shows the presence of grains, chromoblastomycosis shows sclerotic bodies consisting of thick walled muriform cells, while phaeohyphomycosis is characterized by the presence of dark brown mycelial structures in the involved tissue. These are uncommon causes of human disease, but can be responsible for life-threatening infections in both immunocompromised and immunocompetent individuals.

Phaeohyphomycosis covers a wide variety of clinical forms, depends on the immune status of the host. The involvement of the skin and subcutaneous tissue is more frequent than any other organs and termed as subcutaneous phaeohyphomycosis. Subcutaneous infections mostly occur on the limbs, fingers, wrist, knees, or ankles as a result of traumatic implantation with contaminated matter. The lesion occurs as a single, discrete, asymptomatic small nodule, which is palpable as a smooth swelling. This case had shown such features at the first appearance of the lesion (Figure 1A). In this case, there was no history of trauma, but the trauma could have been so trivial and not noticed by the patient as it was in the left lower leg region and there was a risk factor from outdoor occupation.

Subcutaneous phaeohyphomycosis is a rare fungal infection and is even rarer in immunocompetent individuals because it usually occurs in immunocompromised patients. Some cases have been reported within immunocompetent patients without pathologic antecedents as observed in this case. It shows that immunocompromised...
state is not a necessary prerequisite for phaeohyphomycosis. There are putative mechanisms by which these fungi can cause disease, particularly in immunocompetent individuals. Melanin is a known virulence factor in these fungi, and it acts by scavenging free radicals and hypochlorite that are produced by phagocytic cells in the oxidative process and also prevents their action on the plasma membrane by binding to hydrolytic enzymes. The establishment of fungal appressorium, which helps fungi to penetrate into the host cell is also influenced by the melanin. All these multiple functions may explain the dematiaceous fungi pathogenic potential even in an immunocompetent host.

The diagnosis of phaeohyphomycosis is based on clinical presentation and must rely on pathological examination of clinical specimens and careful gross and microscopical examination of cultures, occasionally requiring the expertise of a mycology reference lab. Without those modalities, the diagnosis and the treatment could be false as observed in this case that the patient was misdiagnosed as soft tissue tumor and subcutaneous tuberculosis and got surgical and anti-tuberculosis therapy respectively, but there was no improvement and on the contrary the lesion became worse. Therefore, referral to a tertiary hospital must be considered in this case for further investigation. In this case, diagnosis of subcutaneous phaeohyphomycosis had been established based on fungal culture colonization identified as *E. dermatitidis* as one of the most common etiological agents of subcutaneous phaeohyphomycosis along with the discovery of granulomatous inflammation in tissue samples.

There are no standard treatment for subcutaneous phaeohyphomycosis. There are no trials comparing different strategies for the treatment of infections caused by dematiaceous fungi. Surgical excision alone of subcutaneous cysts has been successful in a number of cases, but oral systemic antifungals agent is frequently used in combination with surgery when possible.

The patient was treated with itraconazole for antifungal systemic therapy as in vitro susceptibility of most strains of dematiaceous fungi to this drug are high. Successful response without surgical excision was achieved with a six-month itraconazole treatment and the patient still continue the treatment in order to achieve complete clinical cure.

Conclusions

This case highlights successful treatment that can be achieved by itraconazole monotherapy and the importance of considering any subcutaneous soft swellings, especially on the extremities, under differential diagnosis for fungal infections as they may be misdiagnosed as soft tissue tumor and cutaneous tuberculosis, although in immunocompetent individual.

Figure 2. A. Black Coloured Colonies On Fungal Culture; B. The Reverse Of The Colonies Was Black; C. Fungal Morphology On Slide Culture Identified As *E. Dermatitidis*; D. Granuloma In The Dermis Composed Of Histocytes, Lymphocytes, And Fibroblasts (Hex40); E. (Hex400) Closer View; F. No Fungal Elements (Pass400); G. No Acid-Fast Bacilli (Znx400).
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