Mucormycosis and chromoblastomycosis occurring in a patient with leprosy type 2 reaction under prolonged corticosteroid and thalidomide therapy

Flávia Machado Alves Basílio¹  Maira Mitsue Mukai²  Rosângela Lameira Pinheiro⁴  Mariana Hammerschmidt¹  Betina Werner³  Sandra Moritz⁵

Abstract: Mucormycosis is an uncommon fungal infection caused by Mucorales. It frequently occurs in patients with neutropenia, diabetes, malignancy and on corticoid therapy. However, it is rare in patients with AIDS. Clinical disease can be manifested in several forms. The case reported illustrates the rare occurrence of chromoblastomycosis and mucormycosis in an immunosuppressed patient with multibacillary leprosy, under prolonged corticosteroid and thalidomide therapy to control leprosy type 2 reaction. Neutrophil dysfunction, thalidomide therapy and work activities are some of the risk factors in this case. Chromoblastomycosis was treated by surgical excision and mucormycosis with amphotericin B. Although the prognosis of mucormycosis is generally poor, in the reported case the patient recovered successfully. This case should alert dermatologists to possible opportunistic infections in immunosuppressed patients.

Keywords: Chromoblastomycosis; Immunocompromised host; Leprosy; Mucormycosis

INTRODUCTION

Mucormycosis is an emerging life-threatening infection caused by Mucorales (Rhizopus, Rhizomucor, Absidia, Mucor, Cunninghamamella). Given the ubiquitous nature of these fungi (found in soil and in decaying matter), most humans are exposed to these organisms on a daily or weekly basis. Nonetheless, they rarely cause disease because of their low virulence, mainly affecting individuals with immunosuppressed conditions.¹

Chromoblastomycosis is a chronic disease cau-
sed by traumatic inoculation of saprophagous dematiaceous fungi, when they enter through an open wound and infect both skin and subcutaneous tissue. This disease is mostly reported in tropical and subtropical areas (higher in rural populations) and is often caused by *Fonsecaea (F.) pedrosoi*. Countries with the highest numbers of cases are Madagascar and Brazil.

The case reported illustrates the rare occurrence of chromoblastomycosis and mucormycosis in an immunosuppressed patient with multibacillary leprosy, under prolonged corticosteroid and thalidomide therapy to control leprosy type 2 reaction.

**CASE REPORT**

A 28 year-old white male, rebar setter, born in Itaperuçu-PR and residing in Curitiba-PR/Brazil, was diagnosed with leprosy and neuritis in 2001, confirmed by skin biopsy (Figure 1). He was being treated with multibacillary chemotherapy and prednisone 1 mg/kg since May 2001. Three months later, he developed flu-like symptoms. Rifampin was discontinued and dapsone and clofazimine were maintained. Erythema nodosum appeared in May 2002, and thalidomide was added to the treatment. In April 2003, at the end of the multibacillary chemotherapy, dapsone and clofazimine were suspended. During this time the patient was kept on prednisone and thalidomide to control neuritis and leprosy type 2 reaction (both drugs were suspended only in 2007).

In August 2003, he had a single erythematous, hyperkeratotic plaque with black dots on his right hand (Figure 2). A skin biopsy demonstrated large pigmented muriform cells inside multinucleated giant cells and *Fonsecaea pedrosoi* was isolated from the fungi culture (Figure 3). Chromoblastomycosis was diagnosed and treated by surgical excision, with total resolution. Erythema nodosum and neuritis persisted, and increased doses of both prednisone (80 mg/d) and thalidomide (300 mg/d) were necessary.

In December 2003, six nodular, non-tense, warm and erythematous lesions were noticed on his thighs and trunk (anterior and posterior). Systemic complaints were absent and blood analysis was normal, except for an elevation of serum glucose (229 mg/dl). HIV test (ELISA) was negative. Seven days later, a yellow-greenish fistulization replaced the first lesions and new ones appeared. Skin biopsy of a left thigh lesion showed acute and chronic inflammation with broad dermal necrosis; secondary vasculitis was seen focally and fungi were not detected. A PAS-stained pus smear with diastase highlighted broad, pleomorphic, non-septated hyphae with irregular, non-parallel contours (Figure 4). *Mucor sp* was isolated from the culture.

Treatment was started with parenteral amphotericin B 1 mg/kg/d (total dose=1880 mg) with concomitant serum glucose control. Surgical drainage was performed in the larger lesions. Prednisone was tapered during this period. The patient was discharged from the hospital with only skin scars in the previously involved areas (Figure 5).

In February, 2004, the patient returned with papular lesions, plaques and fever. A new biopsy of a thigh lesion revealed virchowian leprosy and Lucio’s phenomenon (necrotizing erythema). Modified multibacillary chemotherapy was started, with dapsone, clofazimine and ofloxacin (due to rifampin intolerance), and was finished after 24 months of treatment, in March, 2006.

During follow-up, patient had been free from lesions or reactional states since July, 2006, and last negative bacterioscopy was in December, 2008. Outpatient treatment was finished in January, 2010.

**DISCUSSION**

Mucormycosis is described in several clinical forms, with rhinocerebral disease affecting the paranasal sinus orbits and the brain (80% to 90% of cases). The pulmonary, disseminated, cutaneous, and gastrointestinal forms are less common. The major route of infection is via inhalation of fungal spores; other routes include ingestion and traumatic inoculation.

The cutaneous form (shown in this case) is less common than other clinical forms, but potentially lethal if early treatment is not started. It is usually related to local trauma, with several cases described after the use of adhesive bandages and invasive catheters contaminated by spores. Two different presentations of primary cutaneous mucormycosis have been described. The superficial form appears as vesicles or pustules that ulcerate with scar formation, usually in health-
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be obtained in order to identify the hyphae and culture can identify the causative organism. Vascular invasion is a hallmark of zygomycosis, commonly associated with thrombosis of vessels and necrosis of surrounding tissue.

The mucormycosis treatment approach should begin with primary prevention, including careful use of immunosuppressive therapy in patients whose condition demand the use of such agents. Correction of predisposing factors such as hypoalbuminemia, hypogammaglobulinemia, neutropenia, and hyperglycemia are also recommended.

Amphotericin B is the antifungal agent of choice against mucormycosis. The recommended dose is 1mg/kg/d and the duration of treatment is between 3 and 6 weeks. Liposomal amphotericin presents reduced nephrotoxic effect and improved cell penetration. Besides, recent data suggest that posaconazole may be useful as salvage therapy, and adjunctive immune therapy with recombinant granulocyte colony-stimulating factor (G-CSF) and GM-CSF, or with recombinant IFN-\(\gamma\), has been used successfully in conjunction with lipid formulations of amphotericin B in treatment of mucormycosis.

An iron chelator, deferasirox, does not allow iron utilization by the fungus and has been used as an...
agent for infection treatment, unlike deferoxamine that is a risk factor, since it allows the fungus to utilize deferoxamine-bound iron by recognizing it as a siderophore. 

Furthermore, early and repeated surgical debridement of the involved tissues associated with an antifungal agent is an important part of the treatment and hyperbaric oxygen may be useful as an auxiliary treatment in cutaneous and rhinocerebral mucormycosis.

Chromoblastomycosis is primarily a disease of tropical or subtropical regions, mainly in people working outdoors in agricultural occupations. The characteristic lesion is the warty papule or plaque eventually leaving a fibrotic scar.

The disease spreads in the adjacent skin with characteristic, satellite lesions surrounding the primary source. The lesions develop slowly at the site of implantation. Over the years, the nodule grows centrifugally. In many instances, the central parts of the lesion heals, leaving ivory-colored scars. Mortality due to chromoblastomycosis is a rare event.

The diagnosis is based on KOH examination, identification of organism in histological sections and culture of the organism, which reveal slow growth of green to black colonies, and microscopic analysis of the conidia may identify the agent. A skin biopsy usually shows hyperkeratotic pseudoepeitheliomatous hyperplasia with neutrophilic granulomatous reaction in the dermis. Pigmented large fungi are easily seen when stained with H & E.

Chromoblastomycosis lesions are recalcitrant and extremely difficult to eradicate. Chemotherapy (especially itraconazole), surgical excision and/or cryosurgery have been used throughout the years, but an effective treatment has not yet been established. Recent studies provide evidence that the HIV PIs (nel-finavir and saquinavir) have multiple effects on crucial biological processes of F. pedrosoi, showing the possibility of exploiting HIV PIs as a potential therapeutic agent, alone or in combination with currently used antymycotic drugs, in chromoblastomycosis.

Reported cases of mucormycosis in literature include several clinical conditions, but none was described in association with leprosy.

In this case, the patient had some risk factors to development of mucormycosis, such as: prolonged corticosteroid therapy, which has negative effects on neutrophilic granulocytes (although the number of these cells is enhanced), causing impaired margination, reduced chemotactic activity, affecting phagocytosis and intracellular killing by neutrophils; thalidomide therapy, leading to diminished TNF alfa production and suppressed Th1 response; his profession, rebar setter, includes activities for steel reinforcement for concrete structures in which there is contact with sharp edges, iron and wood materials in a damp environment, the perfect place for fungi reproduction. This way, probably traumatic inoculation of fungal spores was the route of infection for both mucormycosis (multiple lesions) and chromoblastomycosis (single lesion).

Patients with leprosy are not susceptible to infections caused by Mucorales, as the neutrophil functions remain intact. However, the immunosuppressive effects of the therapy used to control reactional states may increase the prevalence of such infections. Probably the patient would not have the two diseases if he were not under prolonged corticosteroid and thalidomide therapy. In addition, work activities may have contributed significantly to mucormycosis and chromoblastomycosis infections in this case.

Mucormycosis has a high mortality rate and the survival outcome depends on early diagnosis, followed by an effective and aggressive treatment.

Although the prognosis of mucormycosis is generally poor, in the reported case the patient recovered successfully.

This case alerts dermatologists to possible infections occurring in immunosuppressed patients. A high index of suspicion is warranted in order to perform an early diagnosis and treatment, and better prognosis.
REFERENCES


MAILING ADDRESS:
Flávia Machado Alves Basilio
Rua General Carneiro, 181
Alto da Glória
80060-900 Curitiba, PR
E-mail: flavia_mab@yahoo.com.br