INTRODUCTION

Mucormycosis is an invasive fungal infection caused by opportunistic fungi of the phylum Glomeromycota. It is frequent in poorly controlled diabetic patients and individuals with immunosuppression. It is usually acquired by direct inoculation through trauma. The clinical presentation is nonspecific, but an indurated plaque that rapidly evolves to necrosis is a common finding. Diagnosis should be confirmed by demonstration of the etiological agent and new molecular diagnostic tools have recently been described. It is an invasive life-threatening disease and in order to improve survival, a prompt diagnosis and multidisciplinary management should be provided. The treatment of choice is amphotericin B, but new azoles, such as posaconazole and isavuconazole, must be considered.

Keywords: Mucormycosis; Mucorales; Mucor; Zygomyces; Rhizopus

EPIDEMOLOGY

In developed countries, patients with allogeneic hematopoietic stem cell transplantations and other hematologic malignancies frequently present mucormycosis.\(^1\)\(^-\)\(^3\) In contrast, in developing countries, uncontrolled diabetes mellitus is the main associated disease.\(^4\)

Its global incidence is unknown; however, in Spain the incidence was 0.43 cases per million per year in 2005. In France, between 1997 and 2006, an increase from 0.7 to 1.2 cases per million was found, mainly in patients with hematologic malignancies and bone marrow transplantation.\(^1\) In the United States of America, there are no extensive prospective reports. In reports from the state of California, incidence was 1.7 cases per million per year.\(^5\)

Zygomycetes are ubiquitous in nature and are usually isolated from decaying organic matter, soil, wood, cotton, bread, fruits, vegetables, and animal excreta.\(^6\),\(^7\) Primary cutaneous mucormycosis is often acquired by direct inoculation, contaminated dressings, surgery, burns, motor vehicle accidents, and insulin injection sites.\(^1\),\(^3\),\(^8\)

There are reports of outbreaks of mucormycosis in healthcare centers due to contaminated adhesive tape, wooden tongue depressors, ostomy bags, and building construction contamination. Other reports have associated this fungus with vascular devices and nitroglycerine patches.\(^7\),\(^9\),\(^11\) In a review of 196 cases of healthcare-associated mucormycosis, 57% involved the skin. Predominant populations were premature infants, surgical patients, and immunocompromised hosts.\(^9\)

According to the largest review of reported cases, the age of presentation ranges from months to 87 years with a mean of 38 years. It is more frequent in men with a ratio of 1.1:1.\(^1\),\(^2\),\(^12\)

The classic major risk factors are uncontrolled diabetes and ketoacidosis, but more recently, hematologic malignancies and allogeneic hematopoietic stem cell transplantation have become more frequent. Other underlying conditions are solid organ transplantation, deferoxamine therapy, drug injections, renal failure, infant low birth weight, malnutrition, HIV infection, systemic lupus erythematosus, burns, trauma, aplastic anemia, and steroid use.\(^1\),\(^2\),\(^7\)

Mucormycosis can also occur in patients with no predisposing factors. According to Roden et al, in 50% of cases of cutaneous mucormycosis, there is no underlying condition; in another review, 40% of patients were immunocompetent.\(^7\)
ETIOPATHOGENESIS

The fungi responsible for mucormycosis, also called zygomycosis, were previously classified in the class Zygomycetes, order Mucorales. Zygomycetes also include the order Entomophthorales, which are responsible for entomophthoramycosis. Recently, molecular phylogenetic analyses have found that Zygomyctota are polyphyletic, and these fungi have been placed in the new monophyletic phylum called Glomeromycota, with the subphyla Mucormycotina and Entomophthoromycotina. In this context, the names “mucormycosis” and “entomophthoramycosis” should be preferred, and zygomycosis should be dismissed.13

Most clinical isolates belong to the genera Rhizopus. In reviews of cutaneous mucormycosis, the most frequent isolated strains are Rhizopus oryzae, Lichtheimia corymbifera and Apophysomyces elegans. Other isolates reported are Mucor sp, Saksenaea vasiformis, Cunninghamella bertholletiae, Rhizomucor spp, and Rhizopus microsporus.6,7-14 In Mexico, there is a case report of a new species called Apophysomyces mexicanus.15

CLINICAL ASPECTS

Cutaneous mucormycosis can be classified as primary and secondary. In primary disease, the skin is infected by direct inoculation and in secondary form, by dissemination from other locations, more commonly from a rhinocerebral infection.3 It can be subcategorized according to the pattern of infection as localized, deep, or disseminated. In a review of 929 cases, 176 patients presented with skin involvement. Most of these cases remained localized to the total depth of the skin, with 24% showing extension to bone or muscle, and 20% developing hematogenous dissemination to other noncontiguous organs.1

The most affected areas of the skin are the arms and legs. Other locations include the scalp, face, thorax, back, abdomen, perineum, breast, neck and gluteal area.7,10,16,17

Primary cutaneous mucormycosis may be gradual in onset or fulminant. The clinical presentation varies. Initially, lesions are indurated plaques that are erythematous to purple. These become necrotic with an erythematous halo that can develop into an eschar.1,14 Other presentations include targetoid lesions, tender nodules, ulcers, purpuric lesions, and swollen and scaly plaques (Figures 1 and 2).3,10-24 In patients with surgical wound infections and burns it often appears as cellulitis and necrosis.15,25

In nosocomial infections, erythema and tenderness that rapidly evolves to necrosis is often found. In some cases, cutaneous lesions resemble contact dermatitis, which progresses to ulceration with necrosis or necrotic exophytic tissue.15 The presence of a fistulous tract from skin to liver allograft has been reported.26

Primary cutaneous mucormycosis by R. variabilis in immunocompetent hosts was reported in China and Japan. Unlike other mucorales, these presented as chronic infections.27 These skin lesions were infiltrated plaques, ulcers and nodules, which usually remained localized and gradually expanded over months and years.21,23

Secondary cutaneous mucormycosis usually results from a rhinocerebral or disseminated infection and is more frequent than primary disease. It has an acute onset and a high mortality. The disease usually starts as sinusitis and the most common cutaneous finding is a necrotic eschar (Figures 3 and 4).13-28 The patients can also have oral involvement with necrotic, black or white ulcers (Figure 5). Other signs are fever, periorbital cellulitis, periorbital edema, ophthalmoplegia, proptosis, loss of vision, and other neurological deficits.29,30 The disease can be divided into three clinical stages: stage I, with signs and symptoms limited to the sino-nasal area; stage II, which is characterized by a sino-orbital infection; and stage III, which has intracranial involvement.30

Figure 1: Cutaneous lesion of mucormycosis

Figure 2: Cutaneous lesion of mucormycosis

Figure 3: Ulceration with necrotic tissue in the eyelid with involvement of the eye

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DIFFERENTIAL DIAGNOSIS
Primary cutaneous mucormycosis should be differentiated from aspergillosis and synergistic gangrene caused by bacterial infections. When targetoid lesions appear, the differential diagnoses include autoimmune disorders, drug reactions, infections, infiltrative diseases, and neoplastic disorders. In some cases, it can mimic tinea corporis or pyoderma gangrenosum.3,18,31

In secondary mucormycosis with rhinocerebral involvement, the differential diagnoses include centrofacial lymphomas, rhinoscleromas, sinusitis, anaerobic infection, and aspergillosis.3

DIAGNOSIS
The clinical findings of cutaneous mucormycosis are nonspecific. Early identification of the fungus is essential to establish prompt antifungal treatment. Early detection can be achieved by direct KOH microscopic examination, observing the presence of non-septated, hyaline, hyphae, 5μm wide and 20 to 50μm long, with irregular branching at right angles, mainly at the periphery of the lesion. Impression smears from the wound edges may also be useful.5

Fungal cultures are positive in 50% of cases, but in recent reviews there has been a clear increase in culture positivity from 72% to 89% in cutaneous locations.1,6 Cultures must be performed in Sabouraud and potato dextrose agar media, avoiding media with antibiotics that inhibit fungus growth.3

A biopsy and molecular diagnostic tests should be performed. The biopsy should be taken from the center of the lesion, including subcutaneous fat. Histology is more useful in primary cutaneous mucormycosis. Common findings are edema, thrombosis, infarctions, necrosis and an inflammatory reaction that includes polymorphonuclear cells, plasma cells, and eosinophils. Thick, hyaline, nonseptated and bifurcated hyphae may be seen with hematoxylin and eosin stain but are best visualized with periodic acid-Schiff and Grocott (Figure 6). Most of the microscopic features are nonspecific and a differential diagnosis with other filamentous fungi must be entertained.3,6

New molecular diagnostic tools have been developed to provide precise identification of the fungus. Still, this technology is not available for most patients. These tests target the 18S ribosomal DNA and are highly specific with no cross-reactivity with other filamentous fungi.32-35

Real-time PCR provides identification of Mucorales in tissue samples and clinical isolates with high specificity. Nevertheless, more sensitivity has been reported in fresh frozen specimens than in formalin-fixed paraffin-embedded tissues.36,37 Bernal-Martinez et al.38 developed a single tube multiplex real-time PCR to detect the genus Mucor, R. oryzae and R. microsporus from clinical and culture isolates. This method has 100% specificity, and provides results in 2 to 3 hours. Other techniques described are real-time PCR followed by high resolution meta-analysis and rolling circle amplification for the ITS region of ribosomal DNA.34,39 However, most of the studies are retrospective and with small samples.

TREATMENT
A multidisciplinary approach is necessary to improve survival in cutaneous mucormycosis. This should include extensive surgical debridement, antifungal therapy, correction of the underlying metabolic or impaired immunological status, and control of other concomitant infections (Table 1).

The antifungal of choice is deoxycolate amphotericin B (d-AmB); however, it is often substituted by lipid formulations be-
cause of their better safety profile. They are also less nephrotoxic allowing longer treatment periods with higher doses. Some studies have shown that amphotericin B (AmB) is the most active drug against mucorales clinical isolates, revealing MICs of ≤1 µg/mL in the majority of the tested strains. In immunocompromised patients, the recommended doses for d-AmB are 1-1.5mg/kg/day, for liposomal amphotericin B (L-AmB), 5-10mg/kg/day, and for amphotericin B lipid complex 5 mg/kg/day. Increasing the dose does not achieve higher plasma levels. In order to increase the survival rate, treatment must be started in the first 5 days after clinical diagnosis. Improvement of comorbidities is imperative. The duration of therapy is unclear. Some authors recommend continuing AmB until clinical and radiological resolution, others recommend 6 to 8 weeks. Sometimes this is not possible because of intolerance, often secondary to renal failure.

Azole derivatives have exhibited variable activity against Mucorales in susceptibility in vitro essays. Posaconazole, followed by isavuconazole, is the most active; itraconazole has limited activity, and voriconazole is not active. Posaconazole has been described as second-line therapy in mucormycosis. Several retrospective studies with this antifungal drug have reported good results. Mostly, this triazole was used in patients with failure or intolerance to AmB. The advantages of the oral formulation is that it allows earlier patient discharge and decreases relapse with prolonged administration. In a single prospective study of 21 patients refractory or intolerant to AmB, a good response was reported with posaconazole, with the exception of patients with disseminated disease. Posaconazole is recommended as second-line treatment for patients with refractory disease or intolerance to AmB or for those who need prolonged treatment maintenance. The suggested dose is 400mg bid, and in most reported cases therapy is provided for several months.

Isavuconazole is a new azole recently approved for treatment of invasive mucormycosis in the United States. It has a safety profile similar to fluconazole with IV and oral formulations available. There are three reported cases of immunocompromised patients with pulmonary, rhinocerebral, and disseminated mucormycosis refractory to AmB and posaconazole but treated successfully with isavuconazole. Recently a multicenter, clinical, single arm, open-label trial was conducted, including 34 patients with invasive mucormycosis. These patients were treated with isavuconazole for a median of 84 days. At the end of treatment, 5 patients presented complete response, 6, partial response, 10, stable disease, and 15, disease progression. There was an analysis using matched historical controls treated with AmB that showed no difference in mortality. The most frequent adverse event were gastrointestinal symptoms, but less than 10% of the patients presented liver enzyme disturbances.

Monotherapy with amphotericin is first-line treatment. There are reports and preclinical data on the use of combinations of different antifungals but there is a lack of clinical studies to support this claim. If necessary, a lipid formulation of AmB should be the primary antifungal, combined with caspofungin or posaconazole.

There is a report on the use of frozen sections to help the pathologist and surgeon define the final surgical margins during surgical debridement of the affected tissue.

PROGNOSIS

The overall mortality of all mucormycosis variants has improved from 84% in the 1950s to 47% in the 1990s, mostly because of treatment with AmB. The overall mortality of cutaneous mucormycosis is less than other presentations of mucormycosis, and in three case series it ranged from 25% to 31%. In localized cutaneous mucormycosis, mortality ranged from 4% to 10%, and in deep extension presentations, it ranged from 26% to 43%. In the largest case series, the mortality rate for disseminated disease was 83%; however, in recent reports it ranges from 26% to 50%.

Roden et al. performed a multivariate regression analysis of risk factors for mortality in mucormycosis and found some significant mortality risk factors that included disseminated disease, renal failure, and infection due to Cunninghamamella species. The use of antifungal therapy and surgical treatment, with no underlying condition or type 1 diabetes mellitus was associated with a significant decrease in mortality. In the experience of a Mexican center, where most cases are associated to uncontrolled diabetes mellitus and metabolic acidosis, the mortality rate was 85% in patients with extensive palpebral disease and with a Glasgow Coma Scale lower than 6.

CONCLUSIONS

Mucormycosis is an emerging invasive fungal disease that requires a high level of clinical skill for a prompt clinical diagnosis and surgical debridement. There is a lack of clinical studies to support the use of combination therapy, especially because of the lack of clinical studies on the use of antifungal combinations.

| Table 1: Treatment of mucormycosis |
|------------------------------------|
| **Antifungal therapy** | **Non pharmacologic treatments** |
| AmB deoxycholate | IV | Standard dose 0.25 - 0.75mg/kg per day |
| Liposomal AmB | IV | 5 - 10mg/kg per day |
| AmB Lipid complex | IV | 5 - 7.5mg/kg per day |
| Posaconazole | PO, IV | 400mg bid |
| Isavuconazole | PO, IV | 200mg tid for 6 doses, then 200mg qd |

AmB, Amphotericin B; PO, by mouth; IV, intravenous
in order to improve survival. Where available, new molecular tests should be used to expedite diagnosis. First-line therapy is amphotericin B combined with surgery; second generation azoles derivatives can also be used. Controlling underlying conditions is also an important aspect of therapy.

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1- In primary cutaneous mucormycosis the disease is most frequently acquired by:
   a. Direct inoculation.
   b. Trauma.
   c. Contaminated dressings.
   d. Surgery.

2- Recently, in developed countries, an emerging risk factor for cutaneous mucormycosis is:
   a. Allogenic hematopoietic stem cell transplantation.
   b. Diabetes Mellitus.
   c. HIV infection.
   d. Aplastic anemia.

3- Which is the most common species isolated in cutaneous mucormycosis?
   a. Rhizopus oryzae.
   b. Lichtheimia corymbifera.
   c. Apophysomyces elegans.
   d. Rhizopus microsporus.

4- In uncontrolled diabetic patients with secondary cutaneous mucormycosis it is frequent to find:
   a. Pulmonary mucormycosis.
   b. Rhinocerebral infection.
   c. Gastrointestinal disease.
   d. Miscellaneous mucormycosis.

5- In China, an emerging form of chronic infection with an indolent course has been reported. The etiological agent related to this variety is:
   a. Rhizopus oryzae.
   b. Lichtheimia corymbifera.
   c. Apophysomyces elegans.
   d. Rhizomucor variabilis.

6- The targeted region of DNA used in molecular diagnostic tools is:
   a. Mitochondrial DNA.
   b. 30S ribosomal DNA.
   c. 18S ribosomal DNA.
   d. Gene encoding a ferulic acid esterase.

7- The \textit{in vitro} tests have showed that the most active antifungal is:
   a. Isavuconazole.
   b. Posaconazole.
   c. Caspofungin.
   d. Amphotericin B.

8- Which of the following factors is not associated with a higher rate of mortality?
   a. Disseminated disease.
   b. Stem cell receptor.
   c. Renal failure.
   d. Infection due to \textit{Cunninghamella} species.

9- Which of the following factors is not associated with a significant increase in survival?
   a. Use of antifungal therapy.
   b. Surgical treatment.
   c. Having no underlying condition.
   d. Use of filgastrim.

10- Which of the following antifungals has good \textit{in vitro} activity?
   a. Posaconazole.
   b. Echinocandins.
   c. Terbinafine.
   d. Itraconazole.

11- In which group of patients is it uncommon to find mucormycosis?
   a. Premature infants.
   b. Surgical patients.
   c. Atopic patients.
   d. HIV patients.

12- In cutaneous primary mucormycosis, extension of the infection usually involves:
   a. Only superficial skin.
   b. Total depth of the skin.
   c. Muscle extension.
   d. Bone extension.

13- What is the first clinical manifestation of rhinocerebral mucormycosis?
   a. Sinusitis.
   b. Periorbital edema.
   c. Ophthalmoplegia.
   d. Proptosis.

14- Which is the initial lesion of primary cutaneous mucormycosis?
   a. Targetoid lesions.
   b. Erythematous plaque.
   c. Tender nodules.
   d. Ulcers.

15- Which is the main differential diagnosis of primary cutaneous mucormycosis?
   a. Lymphoma.
   b. Rhinoscleroma.
   c. Drug reaction.
d. Aspergillosis.

16- Which is the clinical stage of a patient with mucormycosis affecting the sino-orbital area?
   a. Stage I.
   b. Stage II.
   c. Stage III.
   d. Stage IV.

17- What would you expect to observe in a direct KOH microscopic examination of the skin lesions?
   a. Septated hyphae with acute angle or dichotomous branches.
   b. Blastocidal cells forming branched chains of pseudo-hyphae.
   c. Thick-walled spherules filled with endospores.
   d. Non-septated hyphae with right-angle branches.

18- Which of the following antifungals can be used as a strategy for early discharge of the patient?
   a. Posaconazole.
   b. Voriconazole.
   c. Terbinafine.
   d. Itraconazole.

19- Which phylum does mucormycosis fungus belong to?
   a. Chytridiomycota.
   b. Glomeromycota.
   c. Neocllimastigomycota.
   d. Blastocladiomycota.

20- Which is the gold standard for diagnosing mucormycosis?
   a. PCR.
   b. Skin biopsy.
   c. Culture.
   d. KOH microscopic examination.

Answer key
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|   | 1.B | 2.D | 3.B | 4.C | 5.D | 6.D | 7.B | 8.D | 9.A | 10.D | 11.D | 12.C | 13.C | 14.B | 15.A | 16.C | 17.B | 18.C | 19.C | 20.D |
|---|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| 1.B |   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 2.D |   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 3.B |   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 4.C |   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
| 5.D |   |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |

Papers
Information for all members: The EMC-D questionnaire is now available at the homepage of the Brazilian Annals of Dermatology: www.anaisdermatologia.org.br. The deadline for completing the questionnaire is 30 days from the date of online publication.