Black molds or dematiaceous fungi are rare etiologic agents of intracerebral abscesses and such infections carry a high mortality of up to 70% despite combined surgical and antifungal therapy. While the growing use of immunosuppressive therapies and organ transplantation have caused an increase in the incidence of rare fungal cerebral infections, occurrence in immunocompetent hosts is also possible. We describe a 60-year-old female patient with a cerebral abscess caused by *Cladophialophora bantiana*. The case illustrates the clinical and radiological similarities between glioblastomas and brain abscesses and emphasizes the need to perform histological and microbiological studies prior to the initiation of any form of therapy. Long-term survival from cerebral black mold abscesses has been reported only when complete surgical resection was possible. The recommended antifungal treatment involves the use of amphotericin B combined with a triazole and, if possible, flucytosine. Highly-active new generation triazole antifungal compounds (voriconazole or posaconazole) are likely to offer improved survival rates for patients with rare mold infections. In particular, posaconazole could be a new therapeutic option given its better tolerance, lower toxicity and fewer drug-drug interactions. We discuss clinical, microbiological and practical pharmacological aspects and review current and evolving treatment options.

**Keywords** brain abscess, *Cladophialophora bantiana*, dematiaceous fungus, posaconazole, voriconazole

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**Introduction**

*Cladophialophora bantiana* is a highly neurotropic dematiaceous fungus and a rare cause of cerebral abscesses. Such infections carry a high mortality of up to 70% and neurosurgical radical resection associated with powerful antifungal treatment is the most successful therapeutic strategy reported to date. We describe a case involving a 60-year-old woman with an unresectable *C. bantiana* brain abscess treated by a combined approach of CT-guided aspiration and aggressive antifungal therapy. Sequential imaging suggests that the fungal infection was contained. This case illustrates the absence of a pathognomonic image of glioblastoma and the absolute need to obtain histopathological confirmation when possible before initiation of any treatment. Highly active new-generation antifungal compounds will probably improve the prognosis of rare mold infections of the central nervous system (CNS) and we review the microbiological and practical pharmacological aspects of these evolving therapies.

**Case Report**

A 60-year-old woman was referred to our institution because of fever and frequent falls ten days after her arrival in the country. Upon admittance, she was sleepy...
and confused. Her medical history included long-term corticoid therapy (10 mg/d) and cholicicine 1 mg/d for systemic sclerosis, pulmonary fibrosis and Raynaud’s phenomenon which was in remission after six months of cyclophosphamide treatment she received three years earlier. She had chronic hepatitis C-related cirrhosis, chronic thrombocytopenia linked to secondary hypersplenism (60 G/L, normal range 150–350 G/L) and arterial hypertension.

Neurological examination was normal apart from drowsiness and confusion, as was the rest of the clinical examination. Laboratory investigations revealed the lack of an inflammatory syndrome (normal leucocytes, C-reactive-protein <1 mg/l). Liver enzymes were known to be chronically elevated because of active hepatitis C. A cerebral CT-scan showed a 2 cm diameter ring-enhancing lesion in the left posterior arm of the internal capsule compressing the left lateral ventricle (Fig. 1A & B). Magnetic resonance imaging (MRI) of the brain showed a ring-enhancing lesion in the body of the caudate infiltrating the corpus callosum and the posterior arm of the internal capsule, strongly suggesting the diagnosis of high grade glioma (Fig. 1C & D). Due to the history of chronic immunosuppressive therapy with prednisone and previous cyclophosphamide treatment for systemic sclerosis, the probability of a secondary cerebral lymphoma or an opportunistic infectious process was considered. Antibiotic treatment with imipenem-cilastatin 500mg i.v. every 6 h associated with dexamethasone 100 mg was started. Serology for toxoplasma was negative (IgM and IgG) eliminating the possible presence of a cerebral toxoplasmosis abscess.

The patient deteriorated during the first week with the appearance of progressive right hemiparesis,

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**Fig. 1** Initial imaging of lesion. (A) Axial section of native CT-Scan demonstrating a discrete hypodensity in the left body of the caudate and mass effect on the lateral ventricle. (B) Contrast-enhanced CT-Scan showing enhancement of the lesion. (C) Axial section of contrast-enhanced T1-weighted MRI. (D) Coronal reconstruction of contrast-enhanced T1-weighted MRI showing extension within corpus callosum.
aphasia and a worsening in sensorium. Notwithstanding both the risk of hemorrhage increased by thrombocytopenia and radiological features advocating glioblastoma diagnosis, a neuronavigation-guided frameless needle biopsy was performed under platelet transfusion. Approximately 5 ml of a total lesion volume of approximately 6–7 ml was aspirated. Microscopic analysis of Gram, Calcofluor white and Grocott silver stained biopsy material revealed a brain abscess suspected to be caused by a black mold.

The presumptive diagnosis was confirmed by further analysis. The aspirated liquid was cultured onto CHROMagar Candida (Beckton Dickinson, New Jersey, USA). After 12 days, 20 mm size colonies were noted developing in culture. They had a velvety texture and were olive gray to black on the obverse and black on the reverse. Microscopic examination of a lactophenol stained slide culture preparation revealed brown septate hyphae with long, sparsely branched conidiophores bearing wavy chains of smooth oval conidia. The latter did not display dark attachment scars as has been described for other Cladosporium spp. Aspirated liquid was also inoculated onto Sabouraud dextrose agar. The isolate recovered could grow at 42°C, a feature that differentiates C. bantiana from other morphologically similar saprophytic fungi (Fig. 2). To corroborate the identification of the species, samples were subjected to a polymerase chain reaction (PCR) amplification with ITS (internal transcribed spacer) 1 and ITS4 primers as previously described [1]. The amplicon was sequenced on an ABI PRISM 310 Genetic Analyser sequencer (Applied Biosystems, Rotkreuz, Switzerland); the resulting sequence was analyzed with the FASTA program and compared with other sequences of the Cladophialophora genus published in GenBank (NCBI) and SeqWeb2.1.0. [1]. The results confirmed the microbiological diagnosis.

C. bantiana was susceptible to all antifungal agents apart from fluconazole. The minimal inhibitory concentrations (MIC; YeastOne™ Trek Diagnostic, Pennsylvania, USA) obtained were (µg/ml): amphotericin B, 1.0 µg/ml (susceptible = S); fluconazole, 64 µg/ml (resistant); itraconazole, 0.023 µg/ml (S); ketoconazole, 0.032 µg/ml (S); flucytosine, 1.0 µg/ml (S); voriconazole 0.064 µg/ml (S); posaconazole, 0.02 µg/ml (S); caspofungin, 0.50 µg/ml (S). Since no breakpoints have been well established for the different antifungal drugs, it is difficult to establish a standardized antifungal therapy. In addition, the efficacy of various antifungal agents against black molds is not clearly defined in cases involving humans. However, there has been limited clinical experience with the newer antifungal compounds like third-generation antifungal triazoles posaconazole and voriconazole.

Voriconazole (400 mg p.o.s b.i.d) and liposomal amphotericin B (5 mg/kg body weight daily) were started and gradually increased to 7 mg/kg/day. Voriconazole was preferred over itraconazole because of better bioavailability (96% vs 55%) and cerebrospinal fluid penetration (90% vs. 50%). Flucytosine could not be added to this regimen because of the risk of myelotoxicity in a patient with chronic thrombocytopenia. Aphasia remained unchanged and right hemiplegia worsened. Brain MRI performed 5 days after starting treatment showed an increase in the initial lesion and new lesions following the biopsy tract (Fig. 3A). The increase in the levels of liver enzymes (ASAT and ALAT) from 59 and 105 U/l at admission to 176 and 310 U/l, respectively (normal range, 11–42 U/l)) confirmed hepatotoxicity and voriconazole was replaced by posaconazole (400 mg b.i.d p.o.s). Ten days after the change to posaconazole, liver enzymes improved rapidly (ASAT 115 U/l and ALAT 189 U/l).

A third MRI performed 11 days after the introduction of posaconazole showed control of the growing brain abscesses. The volume of the lesions was significantly reduced and less midline shift was evidenced (Fig. 3B). Neurological deficits remained stable. Considering the dynamic of abscess growth before the introduction of posaconazole, the findings were interpreted as evidence of the efficacy of the treatment with posaconazole. Unfortunately, the patient developed aspiration nosocomial pneumonia with severe sepsis and died.

**Fig. 2** Histological and cytological (inset) analysis (Grocott silver stain coloration) revealed invasion of brain structures by septed low branching hyphae.
Discussion

Infection of the central nervous system caused by a dermatiaceous mold is referred to as cerebral phaeohyphomycosis, which may be literally defined as ‘infection due to dark-walled fungi’, i.e., those with the dark pigment melanin. While rare, C. bantiana (formerly Cladosporium bantianum, Cladosporium trichoides and Xylohypha bantiana) is the most common agent of this disease.

Although infections caused by dematiaceous fungi are not common, they are being increasingly recognized as being involved in human disease, particularly in the immunocompromised host [2,3]. They can cause soft-tissue infection, sinusitis, mycetoma and CNS abscesses. In particular, C. bantiana has a high specificity for the CNS as evidenced in a recent extensive review of 101 cases of cerebral phaeohyphomycosis in which 48 cases were associated with C. bantiana [2]. The infection usually results from hematogenous dissemination from a primary site of invasion, commonly the lung. However, the absence of a primary focus is not unusual.

Brain abscesses caused by C. bantiana have been reported in both immunocompetent and immunocompromised patients, predominantly in transplant recipients [4–13]. Immunosuppression due to corticoid therapy, neutropenia or diabetes mellitus has also been associated with infections caused by this mold [14,15]. Direct inoculation, eye trauma and intravenous drug use are other known risk factors. Given the increasing use of immunosuppressive therapies and organ transplantation, the incidence of rare fungal diseases, including cerebral mold infections, will certainly be observed more frequently.

Clinically, patients may present with insidious headaches and slow evolving neurological signs. It is important to stress that fever is not always present and infection parameters can be normal on admission of the patient. The abscess can be single or multiple and is usually easy to identify through CT or MRI studies [16]. Radiologically, fungal abscesses caused by rare fungi such as C. bantiana cannot be differentiated with certitude from bacterial abscesses, primary CNS neoplasia or cerebral metastasis. In particular, differentiating between brain abscesses and cystic brain tumors such as high-grade gliomas and metastasis is often difficult, if not impossible [17]. In the immunocompromised host, the differential diagnosis is broader and opportunistic infections (toxoplasmosis, nocardiosis, and listeriosis among others) and specific malignancy like lymphoma should also be considered. Cerebral biopsy with histological studies and exhaustive microbiological cultures for bacteria, mycobacteria and fungi are considered the gold diagnostic standard and should always be performed.

The mortality rate of cerebral C. bantiana infection is high. A recent series confirmed a death rate of 70% despite surgical resection and systemic antifungal therapy [1]. Several studies have shown that radical surgical resection followed by targeted pharmacological treatment enabled good recovery in some cases [18]. One study of 26 cases of cerebral C. bantiana infections concluded that radical surgical resection of the CNS lesions was the best outcome predictor [16]. However, given that CNS disease by C. bantiana is a rare, life-threatening condition, recommendations for systemic antifungal therapy can only be based upon the experience of isolated cases. Based on a large case series [16], systemic antifungal treatment does, however,
appear to affect outcomes. Antifungal therapy is evolving rapidly and more efficient and better tolerated new drugs are now available.

Fluconazole is not active against *C. bantiana* and the classical medical treatment is amphotericin B i.v. alone or associated with flucytosine and itraconazole. In the largest published review of 48 cases, mortality was as high as 70% despite the use of these antifungal agents [2]. The major factor causing such high mortality rates is the poor penetration of amphotericin B (either as deoxycholate or liposomal form) and itraconazole across the blood-brain barrier [19,20]. Furthermore, major adverse effects associated with the use of these drugs often limit and force an interruption of treatment. Amphotericin B is known for its renal toxicity and electrolyte disturbances. Flucytosine use is limited by its high bone marrow toxicity. Itraconazole is an azole used as standard treatment but its oral formulation presents large inter-individual differences in gastrointestinal absorption which is a major limitation and necessitates monitoring of drug levels.

Voriconazole or posaconazole are new second-generation triazole compounds and represent very attractive options to replace itraconazole in *C. bantiana* cerebral infections. Both show broad antifungal spectrum *in vitro*, including *C. bantiana*, with a very good oral bioavailability, high volume distribution and high tissue concentration including the CNS [21]. Successful treatment of *C. bantiana* cerebral infections with both compounds has been reported recently [22,23]. The most common side effects reported with voriconazole are transient visual disturbances and liver toxicity [24]. Posaconazole is also very active, both *in vitro* and *in vivo*, in other human fungal infections such as aspergillosis or zygomycosis [15,21,25]. Efficacy of posaconazole was shown to be superior to itraconazole or amphotericin B in a recent mouse model of *C. bantiana* infection [26]. Posaconazole could be considered as an alternative to voriconazole in cases of pre-existing liver disease or side effects during treatment as occurred in our patient. The case reported herein showed a rapid reduction in liver enzymes after a switch from voriconazole to posaconazole treatment and signs of radiological improvement following posaconazole introduction that were not observed during voriconazole treatment. Other new antifungals are the echinocandins, of which three are currently available, i.e., caspofungin, micafungin, and anidulafungin. They have limited toxicity profiles and minimal drug–drug interactions. Unfortunately, black molds are less susceptible to them.

New antifungal drugs such as voriconazole or posaconazole will probably lead to an improvement of the prognosis of cerebral fungal abscesses but it should be kept in mind that, whenever possible, neurosurgical radical removal of the abscess is currently the treatment of choice and its role will probably not change despite new compounds.

**Conclusions**

Although rare, *C. bantiana* is frequently associated with CNS abscesses. However, other rare fungal etiologic agents should not be excluded from the diagnosis even in the immunocompetent host. Biopsy and microscopic observations, along with microbiological studies for bacteria, mycobacteria and fungi should always be performed to confirm the diagnosis in cases of evolving intracerebral mass lesions. A multidisciplinary approach among neurosurgeons, infectious disease specialists and microbiologists is mandatory in all cases of rare cerebral fungal disease in order to correctly interpret test results and optimize antifungal therapy. The latter is rendered difficult because of the rarity of such diseases, difficulties in extrapolating drug efficacy from *in vitro* susceptibility testing, availability of several new potent drugs, experience in adverse events and drug-drug interactions. Promising new therapies are currently available and additional studies will confirm soon if their introduction into clinical practice will really translate into improved survival. Despite the often fatal outcome, we encourage clinicians to consider newer therapeutic approaches for these life-threatening infections when resection is not possible.

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**References**


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