Chronic Meningitis and Hydrocephalus due to *Sporothrix brasiliensis* in Immunocompetent Adults: A Challenging Entity

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Chronic meningitis caused by *Sporothrix* sp. is occasionally described in immunosuppressed patients. We report the challenges in diagnosing and managing 2 nonimmunocompromised patients with hydrocephalus and chronic meningitis caused by *Sporothrix brasiliensis*. This more virulent species appears to contribute more atypical and severe cases than other related species.

**Keywords.** *Sporotrichosis; Sporothrix brasiliensis; chronic meningitis; hydrocephalus; virulence; CNS*

Chronic meningitis caused by *Sporothrix* spp. has occasionally been described in patients with immunosuppression from alcohol abuse, cirrhosis, transplantation, diabetes, and Hodgkin’s disease. It has recently been increasingly reported in AIDS patients as part of a cat-associated epidemic of sporotrichosis in Rio de Janeiro State, Brazil [1, 2]. *Sporotrichosis* has therefore been suggested as a differential diagnosis of chronic meningitis in immunosuppressed patients living in endemic or hyperendemic sporotrichosis areas [3]. The clinical outcomes of chronic meningitis due to *Sporothrix* sp. in immunosuppressed patients are poor, with an overall 50% mortality rate [1]. We present 2 cases of hydrocephalus due to unsuspected chronic meningoencephalitis caused by *Sporothrix brasiliensis* in nonimmunocompromised adults and highlight the challenges in diagnosing and managing these cases.

**CASE 1**

A previously healthy 46-year-old male presented in January 2016 with a 1-month history of headaches, retro-orbital pain, episodes of confusion, and gait impairment. His previous medical history was unremarkable, and he denied alcohol use. The patient was from the countryside of Paraná State, where he was a quarry worker. Initial investigation showed a cerebrospinal fluid (CSF) with very mild mononuclear pleocytosis (7 cells/mm³), low glucose (12 mg/dL), and elevated protein (148 mg/dL), but negative microbiological analyses. Magnetic resonance imaging (MRI) revealed basal meningitis (Figure 1A). Because tuberculosis is highly endemic in Brazil, he was empirically treated for tuberculous meningitis with the standard regimen for 4 months, without showing signs of improvement.

On readmission, the patient was wheelchair bound due to motor weakness and lack of balance and was confused and disoriented. MRI showed hydrocephalus, and a ventriculoperitoneal shunt (VPS) placement was indicated. This procedure resulted in partial neurological improvement. The patient was discharged from the hospital but kept on antituberculosis therapy. HIV serology was negative. The patient presented relapses of the central nervous system (CNS) manifestations that were caused by obstruction of the CSF pathway secondary to formation of a pseudocyst around the peritoneal tip of the shunt, which accumulated large volumes of CSF. The CSF pathway was reestablished each time via a surgical procedure; 3 consecutive CSF samples were collected and showed mild mononuclear pleocytosis, low glucose, and elevated protein; all were negative on direct mycological exam (DME), but 2 yielded *Sporothrix* sp.

The isolate was subsequently identified as *S. brasiliensis* by DNA sequencing of the calmodulin gene (GenBank MG869808). The patient was treated with amphotericin (accumulated dose 2 g), with good clinical response. As neither itraconazole nor serum level monitoring was available [4], he was treated with fluconazole (800 mg/d), scheduled for 1 year; no relapses were recorded at the time of writing. However, the patient presented neurological sequelae (ataxia and extrinsic ocular motor paresis related to basal meningitis), which are slowly improving.

**CASE 2**

Case 2 was a previously healthy 40-year-old male with progressive weight loss and lethargy dating from October 2016. He was a farmer in the countryside of Bahia State. He had been treated for “depression,” but his symptoms evolved, with headaches,
vomiting, and confusion. He was HIV-negative and denied alcohol abuse. In April 2017, a cranial computed tomography scan showed hydrocephalus and transepidual edema without anomalous intraparenchymal contrast enhancement or intraparenchymal lesions. C, Upper image: baseline CSF direct mycological exam (August 2017) of patient 2 showing yeast cells suggestive of *Sporothrix* sp. free (thin arrows) or engulfed by macrophages (thick arrows). Lower image: calcofluor white staining of patient 2’s CSF sample collected on August 2017, showing a yeast aggregate surrounded by extracellular matrix suggestive of a biofilm-like structure. D, Fibrin sheath on the tip of the ventriculoperitoneal shunt removed (August 2017) from patient 2: culture of the material from the tip yielded *Sporothrix* sp., subsequently identified by molecular method as *S. brasiliensis*.

As with patient 1, a peritoneal pseudocyst with 1500 mL of CSF was found, and a clinical hypothesis of peritonitis due to secondary bacterial infection (although cultures for bacteria were negative) was made. This led to empirical treatment with vancomycin and ceftazidime; abdominal symptoms improved significantly. CSF analysis at this time showed 12 mononuclear cells, glucose 53 mg/dL, and protein 43 mg/dL; however, DME yielded yeast forms, and culture revealed *Sporothrix* sp. (Figure 1C). The peritoneal tip of the VPS showed a fibrin sheath (Figure 1D), which was also positive for *Sporothrix* sp. on culture. Molecular identification revealed the species *S. brasiliensis* (GenBank MG867724), as in case 1. The catheter was removed, and a new VPS was positioned. A qualitative immunoelectrophoresis test for anti-*Sporothrix* antibodies yielded positive results in serum and CSF.

However, the patient’s follow-up was complicated. Subsequent episodes of obstruction accompanied by recrudescence of hydrocephalus required successive neurosurgical procedures, including
a neuro-endoscopic ventricular septostomy and, ultimately, a ventriculostomy shunt with 2 proximal catheters with a Y-connector. The patient was started on amphotericin deoxycholate, but attempts to move to itraconazole failed because of the development of severe phamacoderm. The patient remains hospitalized, receiving lipo-somal amphotericin and monitoring of hydrocephalus. All subse-
quent CSF analyses were negative for Sporothrix but still presented mild biochemical and cellular abnormalities, with the exception of the most recent (December 2017), which was normal.

DISCUSSION
The presented cases highlight the challenges in diagnosing and managing chronic meningitis caused by Sporothrix brasiliensis. Although currently unusual, this issue will likely be of growing importance due to the continuous expansion of the sporotrichosis epidemic in Brazil, which is causing atypical and more severe cases [5]. The diagnosis was delayed in both patients; they neither presented clini-co-laboratorial evidence of immunosup-
pression nor were from municipalities where cases of human or feline sporotrichosis had been previously reported, according to the local health authorities. However, a case of cutaneous sporotrichosis was documented in patient 1’s municipality several months after the presented case, likely transmitted by a cat with an illness typical of sporotrichosis. Unfortunately, the animal could not be located for confirmation. In neither case was CNS involvement preceded by manifestations of cutaneous-lymphatic sporotrichosis or associated disseminated disease. In immuno-
suppressed patients, CNS involvement is commonly part of a constellation of manifestations suggestive of hematogenous dis-
ssemation [2, 3]. Therefore, case 1 was empirically treated for tuberculosis for more than 4 months, whereas case 2 was diagnosed as idiopathic hydrocephalus 8 months before Sporothrix brasiliensis was isolated from CSF. These 2 cases are similar to those previously described in the literature with regards to both the difficulty in recovering the agent from the CSF [1, 2, 6, 7] and the delay in starting appropriate treatment, likely worsening the progno-
thesis. Published cases refer to several weeks to many months from initial symptom presentation to identification of the fungus [1]. The usual scenario is that of a patient present-
ing the clinical syndrome of chronic meningitis, manifested initially by headache that progressed to lethargy, confusion, or other less frequent manifestations such as vomiting, seizures, gait disturbances, and other neurological deficits. It is note-
worthy, like in the cases reported here, that fever was not always present. Preceding or associated cutaneous-lymphatic sporotrichosis was also not always present, particularly in nonimmuno-
nocompromised patients. Moreover, most patients, like these, did not report a history of unusual environmental exposure to Sporothrix sp. CSF analysis is also indistinguishable from that of chronic meningitis caused by other agents: elevated protein and low glucose levels were generally described, as well as low to moderate pleocytosis, although cases with 0 cells have already been described [1, 6]. The patients in this report followed this pattern, with case 1 presenting more prominent CSF alterations than case 2. Brain imaging, when reported, was either normal or showed meningeal enhancement; in a few patients there were signs of vasculitis and infarcts. Parenchymal lesions were occasion-
ally described, mostly in immunosuppressed, HIV-infected patients [1, 7].

In light of the lack of awareness of Sporothrix sp. as a cause of chronic meningitis and the difficulties in obtaining positive fungal cultures from CSF in meningeal sporotrichosis, Scott et al. [6] recommended, 30 years ago, systematic serological testing of the CSF for antibodies to Sporothrix sp. when investiga-
ting chronic meningitis cases with difficult etiological diag-
oses. In Brazil, serological testing is restricted to a few research laboratories. Our laboratory (LIM-53) developed a qualitative immuno-electrophoresis test for detecting anti-Sporothrix sp. serum antibodies in the 1980s, which will now be included in the screening panel for deep myoses (histoplasmosis, asper-
gillosis paracoccidioidomycosis). This test was performed in patient 2, yielding positive results in serum and CSF. However, despite its high specificity, it still has low sensitivity (≤40%) and needs improvement. The importance of serological diagnosis was recently demonstrated by a case report, which could only be diagnosed through CSF antibody detection [7].

The conditions of both patients were and still are difficult to manage. Although they presented very mild CSF pleocyto-
sis, they evolved with hydrocephalus and required ventricular shunts. Moderate hydrocephalus was also present in several of the patients described in the literature [1], but placement of VPS was not frequently reported, except in Scott et al’s case series, where 5 of the 7 patients underwent VPS; only 1 of these patients died, a patient with a 6-month-delayed diagnosis who could only be treated with a small amount of amphotericin B [6]. More recently, Freitas et al. [2] reported that 2 of the 4 HIV-infected patients from the hyperendemic area of Rio de Janeiro died of hydrocephalus complications; however, it was not mentioned whether these patients underwent VPS. In the patients in this study, the ventricular shunts were very difficult to manage due to frequent obstructions, requiring successive surgical interventions to restore the CSF pathway, indicating the presence of a significant and persistent local inflammatory rea-
tion. Both patients responded to treatment; however, patient 1 developed ataxia and extrinsic ocular motor paresis secondary to basal meningitis, from which he has yet to recover. Patient 2 remains hospitalized on antifungal therapy and in the process of neurological rehabilitation.

It is not known how Sporothrix sp. reaches the CNS. In immuno-
suppressed patients, it is speculated that it is caused by hematogenic spread in the setting of a disseminated disease [1–
3]. However, this would not apply to nonimmunocompromised patients without disseminated disease. An attractive yet specula-
tive hypothesis relies on the recent discovery that the CNS has a
functional lymphatic system connected to the body’s lymphatic system through deep cervical lymph nodes [8]. It is thus possible that our patients acquired the infection through either the respiratory route or an unnoticed traumatic skin inoculation, and cells from the initial local inflammatory response (e.g., macrophages) that phagocytosed yeast cells would have migrated to the lymphatic circulation and eventually reached the CNS through this pathway. Indeed, Figure 1C shows some CSF macrophages engulfing yeast cells in patient 2. In addition, it has been suggested that, at least in experimental models, S. brasiliensis is more virulent in vivo than the other species [9]. In vitro, it has also been shown that Sporothrix brasiliensis has a greater ability to disarm murine macrophages, promoting its survival within macrophages (L. Rossato, PhD, S. A. Almeida, PhD, 2017, unpublished data), and to form biofilms [10]. In fact, biofilm-like structures were observed in patient 2’s CSF (Figure 1C), which may help to explain the recurrent hydrocephalus shunt catheter obstructions.

These cases carry some potentially important messages: that the Brazilian epidemic, due to the more virulent Sporothrix brasiliensis, contributes more to atypical and unexpectedly aggressive cases, even in nonimmunocompromised individuals without a clear-cut epidemiological link, compared with previous outbreaks due to Sporothrix schenckii [11–14]; that sporotrichosis should come to mind in cases of chronic meningitis with complex etiological diagnosis; and that serological tests should urgently be made widely available to enable earlier diagnosis, prevent severe CNS damage, and reduce mortality rates.

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