NOTES

Phaeohyphomycotic Soft Tissue Infections Caused by the Coelomycetous Fungus *Microsphaeropsis arundinis*

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*M. arundinis* is an anamorphic fungal plant inhabitant belonging to the form class Coelomycetes. We describe two cases of *M. arundinis* soft tissue infections in immunosuppressed patients. This organism has not previously been described as causing disease in humans. It was identified on the basis of its typical ostiolate pycnidial conidiomata, ampulliform conidiogenous cells, and small, smooth-walled, brown, cylindrical conidia.

Members of the class Coelomycetes are emerging as an important group causing soft tissue infections in immunosuppressed patients (13, 16). Infections usually are localized, following traumatic implantation, but may progress to invasive subcutaneous disease (2, 4, 5, 7, 9, 11, 17). Airborne transmission of these fungi is unlikely since they typically produce their conidia within enclosed structures known as conidiomata (16). In this report we describe two subcutaneous infections caused by *Microsphaeropsis arundinis* in diabetic patients.

Species of the genus *Microsphaeropsis* Höhnel are coelomycetous anamorphic fungi that have probable affinities to the ascomycete genus *Paraphaeosphaeria* O. Eriksson (3). *Microsphaeropsis* species are typically found as saprobes and parasites of terrestrial plants. They inhabit branches and leaves of various plant hosts and are ubiquitous in soil and freshwater environments.

**Case 1.** An 80-year-old non-insulin-dependent diabetic man presented with a 3-cm-diameter painless, deep granulomatous plaque on the dorsum of the right hand. The lesion had been slowly enlarging over a period of months. Past medical history included ischemic heart disease, chronic renal impairment, and a chronic inflammatory demyelinating polyneuropathy for which he was receiving long-term corticosteroid therapy. His past history was notable for recurrent wound infections of his feet and amputation of two toes 4 years earlier. The ulcers on the feet persisted, and cultures obtained 3 months prior to the present admission yielded bacterial skin flora and an unidentified fungal isolate. Repeat culturing from a different site again yielded the unidentified fungus. Since the significance of this isolate was unknown, it was not investigated any further. At the present admission, tissue obtained from a necrotic lesion on May 17, 2021 by guest
foot, requiring amputation of that foot as well. He was maintained on itraconazole treatment for the *Phialophora* osteomyelitis for 10 months following amputation. There was no subsequent relapse of infection in the legs.

**Histologic examination.** Histopathological examination of material from the first patient showed surface ulceration and an underlying mixed acute and chronic inflammatory response with a prominent granulomatous component. Special stains for fungi (methenamine silver and periodic acid-Schiff [PAS]) highlighted fungal elements as yeast forms and septate pseudohyphae. Many of these organisms were within multinucleate giant cells, and fungal elements were seen extending into the deeper layers of the dermis (Fig. 1A and C). Examination of material from the second patient showed similar features with widespread ulceration and extensive acute inflammation. Chronic inflammation was less prominent, and only a few multinucleate giant cells were discerned. Several of these giant cells contained fungal elements. Methenamine silver and PAS stains again highlighted the budding-yeast forms and septate pseudohyphae which extended into the underlying subcutaneous fat and connective tissue (Fig. 1B and D).

**Mycology.** Specimens collected from both patients were cultured onto BBL Sabouraud dextrose agar Emmons (SDA) containing chloramphenicol (0.16 g/liter) and gentamicin (0.16 g/liter). They were incubated at both 28 and 37°C. BBL Mycosel agar with cycloheximide (0.4 g/liter) (Becton, Dickinson & Co.) and BBL brain heart infusion agar (BHIA) with chloramphenicol (0.16 g/liter) and gentamicin (0.16 g/liter) were also set up at 28 and 37°C.

Growth was apparent within 3 days on SDA and BHIA at both 28 and 37°C, with optimal growth occurring at 28°C. The fungus failed to grow at temperatures above 42°C. Less abundant growth, which occurred more slowly, was detected on the Mycosel agar at 28°C. After 7 days of incubation, subcultures of the colonies measured 15 mm in diameter. They were initially pale grey, fluffy, folded, and spreading. The colonies filled the petri dish within 3 weeks, becoming dark brown to grey with a lighter-colored periphery. Microscopy of the initial cul-

![FIG. 1. (A and B) Histological sections of tissue from case 1 (A) and case 2 (B) demonstrate invasive growth of fungal elements into connective tissue with an associated inflammatory response (PAS stain). (C and D) Methenamine silver stains from case 1 (C) and case 2 (D) further highlight the fungal elements as budding yeast forms and septate pseudohyphae. Magnifications, ×400.](http://jcm.asm.org/)

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tures revealed septate, pigmented, and irregularly shaped hyphae, with swollen segments up to 4 μm in diameter. No conidia or asexual fruiting bodies (pycnidia) were evident after 6 weeks of subculture onto specialty agars including potato dextrose agar (PDA) (Oxoid, Basingstoke, England), Czapek solution agar (Difco Laboratories, Detroit, Mich.), cornmeal agar (Amyl Media Pty Ltd., Dandenong, Victoria, Australia), and 2% water agar.

Subcultures of the isolates were submitted to the Orange Agricultural Institute for specific identification. They were inoculated onto a variety of media to induce sporulation. Colonies on Difco PDA produced dense aerial mycelium, initially greenish-grey in the centre of the colony and whitish at the margin, later becoming dark brown to grey-brown. On BBL malt agar and BBL cornmeal agar, the aerial mycelium was brownish in the center, becoming pale grey-brown at the margins. Colonies were slow growing, reaching only 15 to 20 mm in diameter after 7 days at 25°C. Pycnidium production was slow on PDA, malt agar, and Difco oatmeal agar, occurring only after 8 to 10 weeks. Colonies on carnation leaf agar (tap water agar plus irradiated carnation leaves) produced pycnidia after 4 to 5 weeks under black light and a 12-h/12-h alternate dark-light regimen.

**Identification of isolates.** In vitro, all of our isolates produced subglobose pycnidial conidiomata with dark brown parenchymatous walls (textura angularis) and a single apical ostiole and measured 250 to 350 μm in diameter (Fig. 2A to C). Histological sections of the pycnidia of *M. arundinis* from case 2 were prepared by fixing portions of the colony with visible pycnidia (black dots) in 10% buffered formalin. Following overnight processing, 4-μm thick paraffin sections were prepared and stained with PAS stain. The conidiogenous cells were ampulliform and up to 5 μm long and produced conidia enteroblastically without proliferation (Fig. 2D). The conidia were smooth walled, brown, and cylindrical and measured 3.5 to 4.5 by 1 to 1.5 μm (Fig. 3).
The pycnidial nature of the conidiomata and the enteroblastic nature of the conidiogenesis clearly place them in the genus Microsphaeropsis as currently defined. Of the presently known species, the cylindrical shape and size of the conidia are consistent with an identification of *M. arundinis* (15). These isolates are also identical to a previous isolate from Australia (DAR 35822), which has been identified as *M. arundinis* at the International Mycological Institute.

*M. arundinis* was originally described as a species of *Alveophoma* Bausa Alcade by Ahmad (1). In the original description (in vivo), the conidiomata were described as being scattered or gregarious with one or two ostioles, and the walls were parenchymatous with thickened walls in the outer layers, becoming paler to hyaline in the innermost layer. The conidiomata measured 130 to 165 by 165 to 230 μm. The conidia were described as hyaline and oblong with rounded ends, measuring 6.5 to 7.8 by 1 to 2.5 μm. This fungus was transferred to the genus *Microsphaeropsis* by Sutton (15), who examined the type collection. In contrast to the original description, Sutton described the conidia as brown, cylindrical, and measuring only 4 to 4.5 by 1.5 μm.

*M. arundinis* is distinguished from the species *M. olivacea* and *M. callista* by its smooth, thin-walled, cylindrical, guttulate conidia measuring 4 to 4.5 by 1.5 μm. *M. olivacea* has oval to ellipsoidal, pale brown, smooth, thin-walled conidia measuring 4 to 8 by 2.3 to 5 μm. Conidia of *M. callista* measure 7 to 8 by 4.5 to 5.5 μm, are ellipsoidal to fusiform and thick walled, have a central guttule, and are darker at each end (8). The isolates have been deposited in the culture collections of the Australian Medical Mycology Reference Laboratory (AMMRL); NSW Agriculture Plant Pathology Herbarium, Orange (DAR); and the University of Alberta Microfungus Collection and Herbarium (UAMH). The isolate from case 1 is deposited as AMMRL 159.00 and DAR 75043. Isolates from case 2 are deposited as AMMRL 159.01, AMMRL 159.02, and AMMRL 159.03, DAR 76499, DAR 76031, and DAR 76500, and UAMH 10394, UAMH 10393, and UAMH 10392.

**Bacteriology.** Multiresistant *Staphylococcus aureus* was isolated from three separate tissue samples all collected from lesions on the right leg of the patient in case 2. The fourth tissue sample yielded mixed growth of multiresistant *S. aureus* and *Stenotrophomonas maltophilia*.

**Antifungal susceptibility testing.** The *M. arundinis* isolates from these two patients were tested using the Sensititre Yeast One (Trek Diagnostic Systems Ltd.) colorimetric microbroth dilution method. The sample was incubated at 28°C for 96 h. This longer incubation time was required to allow adequate growth of the fungus, and 28°C rather than 35°C was selected as the better temperature for growth. The MIC of itraconazole was 0.25 mg/liter in both cases. The amphotericin B MIC was 0.25 mg/liter for the organism isolated from the patient in case 1, while two separate MIC tests were performed on isolates from the left calf and the right foot of the patient in case 2. The amphotericin B MICs obtained for these two isolates were 0.125 and 1 mg/liter, respectively.

**Molecular analysis.** Subcultures of the isolates were submitted to the Department of Microbiology at Westmead Hospital for molecular analysis. DNA amplification of the D1 variable region of the 28S gene (large subunit) and the internal transcribed spacer region (ITS1 and ITS2 regions) of the gene coding for rRNA (rDNA) was performed on the three isolates from case 2 and one isolate from case 1. Sequence alignments of the two PCR products (LSU and ITS) using CLUSTAL W showed greater than 99% homology among the four isolates. The identity of the isolates could not be confirmed by DNA sequence analysis using both a BLAST search and a FASTA search, since *Microsphaeropsis* is not yet listed in the GenBank database. The closest match was to a *Paraphaeosphaeria* sp.

**Discussion.** *Microsphaeropsis arundinis* is a plant inhabitant first described from the grass *Arundo donax* L. in West Pakistan (now Pakistan) (1). This grass has a worldwide distribution and is often used for musical instrument and handicraft production. In Australia, *A. donax* is a garden escape weed known as giant reed or elephant grass. It is typically found growing along watercourses, roadsides, and wetlands. It is a robust perennial that grows to a height of 6 m. Although the prevalence of *M. arundinis* on *A. donax* or any other plant species has not been determined in Australia, several other *Microsphaeropsis* species have been described from or reported to occur in Australia. *Microsphaeropsis callista* (H. Syd.) Sutton was isolated from *Eucalyptus haemastoma* leaves (14), while *M. conielloides* Sutton was isolated from leaves of *Eucalyptus pauciflora* and from air (14). *M. olivacea* (Bonord) Höhnel is a ubiquitous species, has been isolated from many plant genera, and has a worldwide distribution, including Australia (15).

*M. arundinis* is confirmed here as a cause of soft tissue infection in two patients, both of whom had diabetes. In the first case, the organism was isolated from a lesion on the dorsum of the hand. Fungal elements were observed by direct microscopy, and their presence was confirmed by histological examination. Treatment with terbinaine appeared to be curative.

The second case was more complicated. The lesions were long standing, and the ulcers became colonized with resistant nosocomial organisms following prolonged hospitalization of the patient. However, despite the mixed infections, we consider *M. arundinis* to be significant, since it was detected by...
histopathological examination, microscopy, and culture from four different lesions. Active inflammation including multinucleate giant cells engulfing fungal elements was present in deep tissue. In addition, a coelomycete was known to be present in the lesions before the patient became colonized with nosocomial organisms during a prolonged hospital admission. Both patients were diabetic and were receiving immunosuppressive medications for intercurrent illness. This would have predisposed them to opportunistic fungal infections. How these patients acquired their infections is not known. The lesion on the dorsum of the hand may well have been acquired by traumatic implantation. The patient never volunteered this information, but he may not have noticed a minor injury. The second patient had chronic nonhealing ulcers on his legs and would have been at risk of infection with multiple organisms. This patient lived in a rural property in Central New South Wales, Australia, an area in which *A. donax* grass is known to occur. However, he did not recall traumatic contact with plant material. He also had a persistent osteomyelitis of his finger as a result of infection with another fungus, *Phialophora verrucosa*, which is associated with traumatic implantation from plant material.

A third isolate of *M. arundinis* identified by this laboratory was obtained from the infected forelimb of a cat. The site of the lesion on the fourth digit was amputated, and the cat was treated with ketoconazole and topical terbinafine without subsequent relapse. This case has been reported elsewhere (8).

A fourth isolate was obtained in 1981 in Sydney, Australia, from an ankle nodule of a patient being treated for acute myeloid leukemia. Histopathology of the nodule showed a leukemic infiltrate and the presence of fungal elements. The fungus was grown on three separate occasions from samples of this site. Isolates have been deposited in the DAR (DAR 35822) and UAMH (UAMH 10391) culture collections (J. Walker, unpublished data). The significance of this isolate could not be established at the time.

Other *Microsphaeropsis* spp. have been implicated in soft tissue infections. *M. olivacea* was reported as causing a skin lesion in an otherwise healthy woman (6), and it was also isolated from the eye of a man with keratitis following traumatic injury (12). *Coniothyrium-Microsphaeropsis* complex was isolated from the ulcerated knee of a diabetic woman who had undergone a renal transplant (10).

Although the identity of our isolates could not be confirmed by molecular analysis, they were shown to have greater than 99% homology to each other, indicating that they are of the same species. Since the taxonomy of the coelomycetes is complex, detailed molecular analysis would be helpful in the definitive characterization of these isolates in particular, as well as in determining the relationships between the various species within the group in general.

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REFERENCES