Disseminated Trichosporonosis Caused by *Trichosporon loubieri*

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Received 1 March 2003/Returned for modification 16 July 2003/Accepted 19 August 2003

**CASE REPORT**

A 56-year-old female was in good health until 2 months prior to admission when she developed progressive fatigue and bruised easily. Initial evaluation revealed marked pancytopenia. A bone marrow biopsy performed in November 2001 was diagnostic of pre-B-cell acute lymphoblastic leukemia. The patient had no significant past medical or family history. She was a resident of Connecticut and worked in an office setting. She was an avid gardener and swimmer. She denied international travel and gave no history of tobacco, alcohol, or drug use.

She was admitted to another institution for induction chemotherapy with cyclophosphamide, daunorubicin, vincristine, l-asparaginase, and prednisone. She became neutropenic and subsequently developed fevers to 39.4°C with no clinical response to empirical use of imipenem and vancomycin. A yeast grew on blood cultures 2 weeks after initiation of chemotherapy, and the patient was started on amphotericin B lipid complex therapy at 3 mg/kg of body weight/day. Blood cultures (BacT/Alert; bioMérieux, Durham, N.C.) remained positive for 5 days after initiation of antifungal therapy. She received granulocyte-macrophage colony-stimulating factor, and her indwelling catheter was removed. The catheter-tip culture showed no growth, and a cryptococcal antigen test was negative. Other diagnostic evaluations performed at the time were unremarkable and included transesophageal echocardiogram, lumbar puncture, and computed tomography of head, chest, abdomen, and pelvis. The patient developed bilateral upper abdominal pain and elevated liver function tests (LFTs), with the following peak values: alkaline phosphatase, 376 U/liter; aspartate aminotransferase, 27 U/liter; alanine aminotransferase, 89 U/liter; total bilirubin, 1.0 mg/dl; and direct bilirubin, 0.3 mg/dl. Blood cultures showed no growth. Initial chest radiograph demonstrated a left pleural effusion, and a contrast-enhanced abdominal computed tomography revealed multiple hypodense liver lesions and extensive coalescence of hypodensities in an enlarged spleen (Fig. 1). A bone marrow biopsy demonstrated remission of the leukemia. On admission, she was initially continued on liposomal amphotericin B therapy at a dose of 7.5 mg/kg/day.

Given the need for further chemotherapy and the above radiological findings, the patient underwent splenectomy and a liver biopsy. Histopathological sections revealed extensive tissue necrosis with innumerable fungal elements (septate hyphae, pseudohyphae, and pleomorphic yeasts were delineated); angioinvasion was also noted (Fig. 2). There was no growth of the organism from the tissue specimens.

The blood isolate was obtained from the referring institution. Growth in potato-dextrose agar revealed white, dry colonies with a central depression and a radially furrowed outer zone; true and pseudohyphae, arthroconidia, and blastoconidia were observed microscopically. Identification was attempted with the use of the API 20C Aux yeast identification system (bioMérieux). The isolate was classified as *Trichosporon mucoides* with a 67% certainty (API 20C Aux 6745776). Antifungal susceptibilities were determined by using Sensititre YeastOne (TREK Diagnostic Systems, Cleveland, Ohio), and the following MICs were found: amphotericin B, 2 μg/ml; fluconazole, 0.5 μg/ml; itraconazole, 0.5 μg/ml; ketoconazole, 0.5 μg/ml; and flucytosine, 8 μg/ml.

The patient remained febrile with abdominal pain. A follow-up computed tomography revealed progression of the hepatic lesions and no other source of fever. Six days after admission, fluconazole at a dose of 800 mg/day was added to her therapy. She continued to be symptomatic, although abdominal computed tomography 2 weeks later demonstrated stabilization of the hepatic lesions. Given the severity of her illness and her decreased performance status, it was decided to not proceed with consolidation chemotherapy, and she was discharged and went home in January 2002. She continued to receive antifungal therapy at home and died in March 2002.

Given the uncertainty of identification of the isolate, PCR amplification and sequencing of gene segments of the small

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subunit of ribosomal DNA (18S rDNA), large subunit of rDNA (26S rDNA), and the internal transcribed spacer (ITS) region between these two genes were performed using well-described universal fungal primers (6, 22, 29). Briefly, fungal colonies were suspended in sterile water to a density of 0.5 McFarland standard. An aliquot was then flash-frozen by immersion into liquid nitrogen. DNA was isolated from 200 μl of the suspension utilizing the guanidium-based method common to Roche Amplicor system applications (Roche Diagnostics, Indianapolis, Ind.). Five microliters of the isolated DNA was amplified by using an AmpliTaq Gold PCR kit (Applied Biosystems, Foster City, Calif.), using 2.5 U of enzyme and 3 mM MgCl₂. All PCR amplifications were done for 35 cycles, with 1 cycle consisting of denaturation at 94°C for 45 s, annealing at 61°C for 60 s, and extension at 72°C for 90 s. The following primer pairs were used (Table 1). For the 18S rDNA amplicon, the primers described by Einsele (6) were used. For the ITS region amplicon, ITS1 and ITS4 primers were used (29). For the 26S rDNA amplicons, the P1 and P2 primers described by Sandhu (22) were used. PCR amplification was confirmed by electrophoresis on a 1% agarose gel containing ethidium bromide. PCR product was purified by use of Microcon YM-100 filters (Millipore, Billerica, Mass.), and sequencing was performed bidirectionally on an ABI 377 DNA sequencer (Applied Biosystems). Sequences were analyzed by using Lasergene software (DNastar, Madison, Wis.).

Sequences obtained from the three PCRs were compared to the nucleotide sequences in GenBank by using the on-line BLAST (1) program (National Center for Biotechnology Information, Bethesda, Md.) (Table 1). The isolate sequences for the ITS region and 18S rDNA amplicons were identical to those published for the CBS 7065 reference strain of *Trichosporon loubieri* (GenBank accession number AB001759 and AB018027, respectively) (24, 25). The 26S rDNA amplicon sequence differed in only one base pair to the published sequence of the same *T. loubieri* strain (C→T at position 490 of GenBank accession number AF075522) (9).

Further testing of the isolate demonstrated growth at 42°C. The carbohydrate assimilation profile provided by API 20C Aux was compatible with the *T. loubieri* profile published (13, 14), including growth with raffinose and with melibiose and lack of growth with melitose. The ID 32C yeast identification system (bioMérieux) did not demonstrate growth with rhamnose or lactose (ID 32C 757764773). The isolate has been deposited in the American Type Culture Collection (Manassas, Va.) under catalog number MYA-2615.

Since the reclassification of the genus *Trichosporon* in 1992 (14), there had been evidence that only six species were associated with human disease (5, 13). *Trichosporon ovoides* and *Trichosporon inkin* are agents typically associated with capital skin infections. Occasionally, these agents have been reported as causes of systemic infection, usually catheter related (13, 16, 17). On the other hand, trichosporonosis has been recognized as an emerging infection in immunosuppressed patients (2, 10). *Trichosporon asahii* is the most common isolate in these cases, followed by *T. mucoides* (10, 13). *Trichosporon beigeli* was commonly used in the literature prior to 1992, but that designation was rejected after reclassification of the genus (14).

*T. loubieri* type strain (CBS 7065) was isolated from a cow with mastitis (14), but *T. loubieri* had not been isolated in humans until recently (20), when infected cysts in a Nepalese patient with polycystic kidney disease were found after nephrectomy. No tissue invasion was seen on histological examination, no fungemia was documented, but the patient's fever subsided after nephrectomy. The patient reported here developed high-grade fungemia and went on to develop a severe hepatosplenic syndrome, described recently for *T. asahii* (19), after her chemotherapy-induced neutropenia resolved. The infection was finally controlled after a splenectomy and with the addition of high-dose fluconazole to amphotericin B. Histology demonstrated invasive disease and evidence of angioinvasion. Although trichosporonosis is known to cause false-positive cryptococcal antigen test results (18), this was not found in the

FIG. 1. Abdominal contrast-enhanced computed tomography demonstrates multiple hypodense lesions in the liver (left) and necrosis of the spleen (arrow).

FIG. 2. Spleen tissue slide. Multiple hyphal elements are observed in an intravascular space. Gomori’s methenamine silver stain was used. Magnification, ×100. (Photomicrograph copyright Danny Milner.)
We thank Akhtari Alam and Esperanza Albano, clinical mycology lab, Brigham & Women’s Hospital, for valuable assistance and Takashi Sugita, Department of Microbiology, Meiji Pharmaceutical University, Tokyo, Japan, for insightful comments and for running the ID 32C carbohydrate assimilation test on the isolate.

REFERENCES

# Table 1. T. loubieri isolate rDNA PCR products

| rDNA segment | Primers* | PCR product (bp) | GenBank accession no. | % Homology to:
|--------------|----------|-----------------|-----------------------|------------------------
| 18S (6)      | ATT GGA GGG CAA GTC TGG TG CCG ATC CCT AGT CCG CAT AG | 462 | AY101606 | T. loubieri CBS 7065 100 (462/462) AB001759 97.4 (450/462) AB001763 |
| ITS region (29) | TCC GTA GGT GAA CTC GCG G TCC TCC GCT TAT TGA TAT GC | 491 | AY101607 | T. loubieri CBS 7065 100 (491/491) AB018027 84.9 (417/491) AB018030 |
| 26S (22)     | ATC AAT AAG CGG AGG AAA AG CTC TGG CCT CAC CCT ATT C | 727 | AY101608 | T. loubieri CBS 7065 99.8 (605/606) AF075522 93.7 (568/606) AF075515 |

* The forward primer is shown on the first line, and the reverse primer is shown on the second line.

% Homology to the T. loubieri CBS 7065 or T. mucoides CBS 7625 sequence (CBS, Centraalbureau voor Schimmelcultures, Utrecht, The Netherlands). The number of homologous base pairs to the total number of base pairs is shown in the parentheses. The GenBank accession number of the T. loubieri CBS 7065 or T. mucoides CBS 7625 strain sequence is shown on the second line.

present case. These findings confirm that T. loubieri is a human pathogen and that it is capable of causing invasive infection.

Gueho and colleagues mentioned the ability of T. loubieri to grow with rhamnose (14), but this is not an exclusive characteristic and only T. loubieri’s ability to grow at 42°C is used in the physiological key for Trichosporon identification (14). In addition, the carbohydrate assimilation profiles of the two variants of T. loubieri from the patient with infected renal cysts (20) suggest that intraspecies variations exist. This underscores the relevance of molecular confirmation of Trichosporon clinical isolates for definitive identification (9, 25). Analysis of rDNA intergenic spacer regions may further aid in the classification of this genus and of species-specific strains (23).

The relative resistance of these organisms to amphotericin B is important to note. Clinical failures and the microbiologic resistance of Trichosporon spp. have been well-documented (26–28), and it has been shown that amphotericin B is fungistatic. This resistance exists in part the recalcitrant course of the patient presented despite the use of high doses of lipid formulations of amphotericin B. Interestingly, there was no growth of the organism from the surgical specimens obtained after several weeks of therapy. There is growing evidence that azole drugs have good activity against Trichosporon spp. (3, 8, 11, 21) and that combined administration of fluconazole and amphotericin B may be superior to either drug used alone against invasive infection (4, 15). Echinocandins have no activity against Trichosporon species (7), and a breakthrough infection in a patient receiving caspofungin was recently reported (12).

Given the robustness of the current classification of the genus Trichosporon (9, 14, 25), species-specific classification should be attempted on all relevant clinical isolates. Caution should be used when interpreting identification based solely on sugar assimilation profiles. Azole-based treatment regimens should be considered the first-line treatment of trichosporonosis.

(A preliminary report of this work was presented at the 40th Annual Meeting of the Infectious Diseases Society of America, Chicago, Ill., 25 October 2002.)

Nucleotide sequence accession numbers. The rDNA sequences of the isolate have been deposited in GenBank (accession numbers AY101606, AY101607, and AY101608 for the 18S, ITS, and 26S rDNA gene segments, respectively).
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