Mycobacterium heckeshornense Infection in HIV-infected Patient

To the Editor: Mycobacterium heckeshornense is a slow-growing scotochromogen phenotypically and phylogenetically related to M. xenopi. It was first identified as a cause of lung infection on the basis of unique 16S rRNA and 16S–23S spacer sequencing (1). Published data are limited to the original description and 5 case reports in English-language literature (2–5). We report disseminated M. heckeshornense infection in an HIV-infected patient and document its role as an emerging pathogen.

A man 40 years of age with diffuse large B-cell lymphoma had advanced HIV infection and a CD4 count <10 cells/mm³. Antiretroviral therapy (ART) comprising abacavir/lamivudine and lopinavir/ritonavir was initiated. Ongoing night sweats and weight loss after chemotherapy prompted submission of blood cultures, which were positive for M. heckeshornense after 41 days’ incubation.

Progressive wasting after 2 months’ ART prompted treatment for M. heckeshornense with isoniazid 300 mg 1×/d, clarithromycin 500 mg 2×/d, moxifloxacin 400 mg 1×/d, vitamin B6 25 mg 1×/d, and rifabutin 150 mg 3×/wk. Pretreatment blood and urine cultures grew M. heckeshornense after 41 and 30 days’ incubation, respectively. After 18 months of ART and antimycobacterial therapy, the patient’s condition improved, and his mycobacterial blood culture remained negative.

Blood for cultures was collected in Vacutainers (Becton Dickinson, Sparks, MD, USA), injected into Myco-F lytic media (Becton Dickinson), and incubated at 35–37°C in a fully automated BACTEC 9000 MB (Becton Dickinson) blood culture instrument for 42 days. Ziehl-Neelsen stain confirmed acid-fast bacilli (AFB) after 41 days. Gen-Probe AccuProbe DNA probes (Gen-Probe Incorporated, San Diego, CA, USA) for M. avium and M. tuberculosis complexes were negative. The initial blood culture was subcultured to Middlebrook 7H11 media and incubated at 35–37°C, demonstrating a growth range of 37°–45°C (optimal growth at 42°–45°C). The isolate was forwarded to the National Reference Centre for Mycobacteriology (Winnipeg, Manitoba, Canada) for partial 16S ribosomal RNA gene sequence identification. It corresponded with 100% sequence identity to the type strain of M. heckeshornense. The 16S rRNA gene sequence analyzed for this isolate was 1,314 bp long and presented a divergence of 2.6% from its closest species, M. xenopi.

Optimal growth temperature and extended time to isolation of the initial isolate prompted a second incubation period of 42 days (total of 84 days) for all subsequent cultures. A subsequent blood culture was positive at 53 days of incubation. If negative after this reincubation period, a terminal Ziehl-Neelsen smear was performed to confirm absence of AFB.

Susceptibility testing of the initial blood and urine isolates was performed by using the radiometric
broth MIC method (Becton Dickinson BACTEC 460 radiometric system; Becton Dickinson Microbiology Systems) and by microbroth dilution method (TREK Diagnostics, Cleveland, OH, USA) for isoniazid and streptomycin. Susceptibility results were as follows: amikacin MIC \( \leq 2.0 \) \( \mu g/mL \), ciprofloxacin MIC \( \leq 1.0 \) \( \mu g/mL \), clarithromycin MIC \( \leq 16.0 \) \( \mu g/mL \), rifabutin MIC \( \leq 0.12 \) \( \mu g/mL \), ethambutol MIC \( \leq 8.0 \) \( \mu g/mL \), isoniazid MIC \( 1.0 \) \( \mu g/mL \), and streptomycin MIC \( 8.0 \) \( \mu g/mL \) (6).

Disseminated disease caused by nontuberculous mycobacteria (NTM) has been described for HIV-infected patients with CD4 counts \(<50\) cells/mm\(^3\). More than 90% of NTM cases are caused by \( M. avium–intracellulare \) complex (MAC) (7). Reports of \( M. hecketshornense \) infection are limited to immunocompetent persons with lung disease (1–3), tenosynovitis (4), and axillary lymphadenitis (5), attesting to the organism’s virulence.

No criteria exist for diagnosing disseminated NTM infection other than MAC, which is based on clinical signs and isolation from cultures of blood, lymph node, bone marrow, or other sterile sites. The most common diagnostic method is blood culture, positive for \( >90\% \) of cases (7). \( M. hecketshornense \) was considered the etiologic agent for the patient reported here on the basis of repeated blood culture isolation, clinical signs, and improvement with treatment. Concomitant antiretroviral therapy and immune restoration also likely contributed to the patient’s improvement.

The incidence of \( M. hecketshornense \) infection may be underestimated because of incubation time allowed by the BACTEC 9000 series instruments at 42 days. The positive cultures for this patient were obtained close to or after the traditional 42-day period and without reincubation may have been missed. Piersimoni et al. (8) found that most blood cultures for \( M. xenopi \) were detected with terminal AFB and visual growth inspection performed after the isolates had been determined as negative by conventional means at 42 days, which suggests a need for prolonged incubation.

The optimal treatment for \( M. hecketshornense \) infection has not been established. Its phenotypic and genotypic resemblance to \( M. xenopi \) suggests that similar treatment may be reasonable. Current recommendations for treatment for \( M. xenopi \) infection include isoniazid, rifampin or rifabutin, and ethambutol, with or without an initial phase of streptomycin (9). For HIV-infected patients, consideration should be given to prolonged treatment similar to that for disseminated MAC infection, generally \( \geq 12 \) months and accompanied by a sustained (\( \geq 6 \) months) increase in CD4 counts to \( >100 \) cells/mm\(^3\) (9).

We report disseminated \( M. hecketshornense \) in an HIV-infected patient, documenting the pathogen’s increasing clinical spectrum. Its isolation requires prolonged incubation and may be missed by standard mycobacterial isolation instruments.

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References


We document expansion of *B. procyonis* roundworms into northwestern and southeastern Florida.

In 2006 and 2007, nine ascarids (>3 inches) were collected from the feces of an unrecorded number of raccoons admitted to a rehabilitation center in northern Florida. In September 2008, December 2009, and June 2010, one ascarid each was found in the feces of a 4- and a 6-month-old raccoon from Leon County, Florida, and a 6-month-old raccoon from Wakulla County, Florida, after routine treatment with pyrantel pamoate (20 mg/kg). In July 2010, a juvenile (6-month-old) raccoon from Broward County, Florida, which had been admitted to a rehabilitation center, passed several ascarids (2 collected for testing) in its feces after ivermectin treatment (0.2 mg/mL) for mange. The 14 ascarids were preserved in 70% ethanol, and adult males were identified as *Baylisascaris* spp. on the basis of their morphologic characteristics (perianal rough patches). The ascarids were subsequently confirmed as *B. procyonis* by sequence analysis of the 5.8S rRNA gene or the internal transcribed spacer (ITS-1) and ITS-2 regions (7,8). The complete sequences of the 5.8S rRNA gene and ITS-2 region from 2 ascarids from northern Florida and 1 from southern Florida were identical to *B. procyonis* sequences (GenBank accession nos. AJ001501 and AB051231, respectively). ITS-1 sequences from the 2 ascarids from northern and southern Florida were 99.1% (424/428; AB053230) to 100% identical (AJ007445 and ascarids from Georgia, Kentucky, and Texas [6]), respectively, to *B. procyonis* sequences.

Several previous studies did not detect *B. procyonis* roundworms in raccoons or latrine sites in central Florida (n = 51 from Glades, Highlands, Hillsborough, and Orange counties), southern Florida (n = 90 from around Miami and n = 64 fecal samples on Key Largo), and numerous counties throughout Florida (n = 177) (1,3,9). Historically, *B. procyonis* roundworms have been absent throughout most of the Southeast, but the parasite was recently detected in north-central Georgia (5,6). How the species became established in Florida remains unclear. Establishment could have resulted from natural dispersal of infected raccoons from *B. procyonis*-endemic areas; however, recent examination of several raccoon populations in southern Georgia failed to detect such infections (6). Alternatively, the parasites could have been introduced from the movement of infected raccoons, exotic pets (e.g., kinkajou [*Potos flavus*]), or natural wildlife intermediate hosts (1).

Additionally, because domestic dogs can serve as definitive hosts, an infected dog from a *B. procyonis*-endemic area may have passed eggs into the environment (1). Veterinarians in Florida should be aware of this possible zoonosis and carefully examine ascarid eggs detected in fecal specimens because *B. procyonis*-infected dogs often have mixed infections with *Toxocara canis, Toxascaris leonina*, or both, which have morphologically similar eggs (1). Physicians, veterinarians, and wildlife biologists in Florida should be aware of this serious pathogen and the likelihood its range will increase, as highlighted by the recent detection of *B. procyonis* roundworms in a kinkajou from southern Florida (K.P. Kazacos et al., unpub. data).

This study also highlights the importance of wildlife rehabilitation centers as resources for the study of wildlife/zoonotic diseases. Animals admitted to rehabilitation centers are often ill or injured, which may increase pathogen shedding or transmission. Additionally, young raccoons are likely to be infected with *B. procyonis* roundworms, and kits as young as 3 months of age can be patent. Numerous fatal *B. procyonis* larva migrans infections have occurred among animals in rehabilitation centers and zoological parks. These infections were