Mycobacterium tuberculosis
Infection of Domesticated Asian Elephants, Thailand

Taweepoke Angkawanish, Worawidh Wajjwalku, Anucha Sirimalaisuwan, Sittidet Mahasawangkul, Thattawan Kaewsakhorn, Kittikorn Boonsri, and Victor P.M.G. Rutten

Four Asian elephants were confirmed to be infected with Mycobacterium tuberculosis by bacterial culture, other diagnostic procedures, and sequencing of 16S–23S rDNA internal transcribed spacer region, 16S rRNA, and gyrase B gene sequences. Genotyping showed that the infectious agents originated from 4 sources in Thailand. To identify infections, a combination of diagnostic assays is essential.

During the past 2 decades, infections of captive African and Asian elephants with Mycobacterium bovis and M. tuberculosis have been diagnosed worldwide (1–4). Transmission of these infections to other mammals and veterinary personnel has also been observed (5). To date, M. tuberculosis infection has not been reported in elephants in Thailand. Four elephants referred to the National Elephant Institute (NEI) Hospital during 2005–2008, three of which were referred to the Guidelines for the Control of Tuberculosis in Elephants, were further identified by PCR reactions by using 16S rRNA, 16S–23S-rDNA internal transcribed spacers (ITS) (7,8), and gyrase B (gyrB) primers (Table 1). The subsequent sequencing was conducted by using an ABI 3070 system (Applied Biosystems, Foster City, CA, USA). Unambiguous sequences were compared with data available in GenBank (www.ncbi.nlm.gov/BLAST) and analyzed by using ClustalW version 1.4 (www.ebi.ac.uk/Tools/clustalw/). The 16S rRNA and ITS sequencing confirmed that bacteria from lesion tissues of elephants 1, 3, and 4 were identical to those of M. tuberculosis strain American Type Culture Collection elephants 1, 3, and 4 was performed at 21 months, 7 days, and 33 months after admission, respectively, and lesion tissues were collected for bacterial culture, Ziehl-Neelsen (ZN) staining, and histopathologic examination.

A serum sample from elephant 1 was negative for M. tuberculosis at admission, but a sample obtained 10 months later was positive. Bacteria could not be grown from trunk wash samples. Necropsy showed that elephant 1 had tuberculous lesions in the respiratory tract, mediastinal lymph nodes, liver, kidney, and spleen. Histopathologic examination showed caseous necrosis; infiltration of lymphocytes; and accumulation of macrophages and giant cells in lung tissue, lymph nodes, and liver. ZN staining identified acid-fast bacilli. Mycobacteria were cultured from lesion tissue.

A serum specimen from elephant 2 was negative for mycobacteria at admission, but a second sample was positive 23 months later. Bacteria that were positive by ZN staining were cultured from a trunk wash sample. This elephant is still alive and being kept in a restricted area.

Serum samples from elephant 3 were negative at days 1 and 7 after admission, and the elephant died a few hours after the second sample was tested. A stored serum sample from elephant 3, obtained 4 months earlier was also negative. The animal was severely ill and in lateral recumbency. Necropsy showed tuberculous lesions in the lungs, upper trachea, and mediastinal lymph nodes. Histopathologic examination showed caseous necrosis and accumulation of macrophages and giant cells in the lung and lymph nodes. ZN staining showed acid-fast bacilli. Mycobacteria were cultured from lesion tissues.

A serum specimen from elephant 4 was positive at admission. Initially, M. avium bacteria were grown from cultures of trunk wash samples. At necropsy, tuberculous lesions were found in the respiratory organs and mediastinal lymph nodes. Histopathologic examination showed accumulation of macrophages and edema in the lung tissues. ZN staining did not show acid-fast bacilli. However, mycobacteria were cultured from lesion tissues.

Bacteria cultured from trunk wash and tissue samples were further identified by PCR reactions by using 16S rRNA, 16S–23S-rDNA internal transcribed spacers (ITS) (7,8), and gyrase B (gyrB) primers (Table 1). The subsequent sequencing was conducted by using an ABI 3070 system (Applied Biosystems, Foster City, CA, USA). Unambiguous sequences were compared with data available in GenBank (www.ncbi.nlm.gov/BLAST) and analyzed by using ClustalW version 1.4 (www.ebi.ac.uk/Tools/clustalw/). The 16S rRNA and ITS sequencing confirmed that bacteria from lesion tissues of elephants 1, 3, and 4 and from a trunk wash sample of elephant 2 belong to the M. tuberculosis complex. The gyrB sequences of isolates from elephants 2, 3, and 4 were identical to those of M. tuberculosis strain American Type Culture Collection.
(ATCC) 27294 and others (Table 2); the gyrB sequence of the isolate from elephant 1 differed at position 482, which is similar to the M. tuberculosis strain KPM KY679, the ancient TbD-positive strain (9–11). The mycobacterial interspersed, repetitive-unit variable number tandem repeat typing of the exact tandem repeat-A (ETR-A) locus was performed according to protocols of Fleche et al. (12). The sequence of the ETR-A locus showed that different types of M. tuberculosis were present in elephants 2, 3, and 4 because the sequence had 3, 2, and 4 repeats of the typical 75-bp sequence, respectively.

**Conclusions**

We report M. tuberculosis infection in elephants in Thailand. Clinical signs shown by these 4 elephants varied considerably. Elephant 2 showed nasal discharge only; in contrast, elephant 3, showed severe clinical signs and lateral recumbency. Elephant 3 had no antibodies, which may indicate an anergic status of the mycobacteria-specific immune response (13). Histopathologic examination showed that this elephant was severely affected by the infection. Elephant 2 is still alive; cultures of trunk wash samples contain mycobacteria. The elephant was seropositive for M. tuberculosis antigens as defined by the StatPak assay. The other 2 elephants (1 and 4) showed anorexia, chronic weight loss, and comparable lesions at necropsy, but diagnostic assays showed variable results. Trunk wash culture, considered to be the standard for confirmation of M. tuberculosis complex infection in elephants, has its limitations, as described elsewhere (14). This study, which included 3 elephants positive for mycobacteria in tissue culture at necropsy, showed that bacterial cultures of only 2 of 60 trunk wash samples were positive for mycobacteria. The study indicates that serologic tests or other diagnostic procedures could not unequivocally identify infected animals, perhaps because of differences in specific immune responsiveness among species and length of time after infection (13). However, the combination of the different diagnostic observations after infection holds promise for improving the likelihood of confirmed M. tuberculosis infection.

Sequence analysis of 16S and ITS indicated M. tuberculosis complex bacteria in each elephant. Nucleotide sequence polymorphism in the gyrB gene of the mycobacteria isolates (9–11) confirmed the identity of M. tuberculosis for all 4 elephants. M. tuberculosis may be classified into ancestral and modern strains based on M. tuberculosis–specific deletion (TbD1) (15). M. tuberculosis isolated from elephant 1 had a gyrB gene sequence identical to strains of the ancient TbD1-positive strain (Table 2). The other 3 elephants were infected with strains identical to M. tuberculosis ATCC 27294, the modern type, potentially related to major epidemics like the Beijing, Haarlem, and African M. tuberculosis clusters (15).

On the basis of these molecular studies, we believe that M. tuberculosis was probably transmitted to these 4 elephants from humans. In addition, mycobacterial inter-

### Table 1. Primers used to identify bacteria cultured from trunk wash and tissue samples from domesticated Asian elephants, Thailand, 2003–2008*

<table>
<thead>
<tr>
<th>Primer</th>
<th>Forward</th>
<th>Reverse</th>
</tr>
</thead>
<tbody>
<tr>
<td>16s rRNA</td>
<td>5’-AgA gTT gTg gTC CTC TgG CTC Ag-3’</td>
<td>5’-AgAgCT ACC TgT TTA CgA CTT-3’</td>
</tr>
<tr>
<td>ITS</td>
<td>5’-TgT gAC gAC CgG CgG gTg a-3’</td>
<td>5’-TgCT CgA TgA CAA gGg ATC CAC C-3’</td>
</tr>
<tr>
<td>gyrB</td>
<td>5’-TgC gAC gGC ggT gAT gAT gTC C-3’</td>
<td>5’-ACA TAC AGT gTC gTT gCG C-3’</td>
</tr>
</tbody>
</table>

*ITS modified from (7,8); gyrB modified from (9); ITS, internal transcribed spacer; gyrB, gyrase B.

![Table 2. gyrB gene sequence comparisons of 4 Mycobacterium tuberculosis isolates from domesticated Asian elephants, Thailand, 2003–2008](image-url)
spersed, repetitive-unit variable-number tandem-repeat typing of the ETR-A gene *M. tuberculosis* strains in elephants 2, 3, and 4 showed different numbers of the typical 75-bp repeat. Therefore, we conclude that the sources of infection were of different origins. Annual health checks of mahouts and veterinarians who were in contact with the infected animals for >4 years at the NEI did not identify any persons with positive results by chest radiograph when tested as part of the tuberculosis control program in Thailand. To control *M. tuberculosis* complex transmission from humans and other species to wild animals, including elephants, or from wild animals to humans, assays that enable early diagnosis of infection are necessary. Because no assay unequivocally defines the infectious status, a combination of diagnostic approaches is essential.

Further investigation of tuberculosis transmission and surveillance and monitoring of this disease in Thailand will enhance the understanding of its epidemiology. Increased epidemiologic knowledge is essential to control and prevent tuberculosis in elephants.

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Mr Angkawanish is a PhD candidate in the Department of Infectious Disease and Immunology, Faculty of Veterinary Medicine, at Utrecht University, the Netherlands. He is also employed by the National Elephant Institute, Forest Industry Organization, Lampang, Thailand. His research interests include the prevalence of elephant tuberculosis, diagnostic methods, and epidemiology of disease.

References


Address for correspondence: Worawidh Wajjwalku, Faculty of Veterinary Medicine, Kasetsart University, Nakhonpathom, Thailand; email: fvetwww@yahoo.com