Rapid detection of *Mycobacterium tuberculosis* complex in cattle and leche (Kobus leche kafuensis) at the slaughter house

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Abstract

The detection and diagnosis of tuberculosis (TB) in food-producing animals is critical to human health. In this study we applied the loop-mediated isothermal amplification (LAMP) system to detect *Mycobacterium tuberculosis* complex (MTC) directly in 57 cattle and six leche (Kobus leche kafuensis) carcasses exhibiting lesions characteristic of TB. The samples were first subjected to Ziehl-Neelsen microscopy, followed by culture and LAMP assay. In addition, multiplex-PCR was used to determine the species involved. Of the samples from the cattle, 84.2% (95% confidence interval: 71.6-92.1) were found positive with Ziehl-Neelsen microscopy, 93.0% (95% confidence interval: 82.2-97.7) with culture, and 94.7% (95% confidence interval: 84.5-98.6) with the LAMP system while the *Kobus leche kafuensis* samples were all positive for all techniques showing a high amplification efficiency and has been used to diagnose several other diseases.10 The objective of our study was to evaluate the applicability of the LAMP system, in the veterinary field, to detect MTC directly from suspected TB lesions of cattle and wildlife being slaughtered for food.

Materials and Methods

**Sampling**

Our study was conducted on samples collected from the slaughtered animals along the examination line. The animals were examined for gross lesions according to the standard postmortem procedures as described previously.18 Organs and tissues with suspected TB lesions were collected after detailed postmortem examination of the entire carcasses. Following collection, the specimens were placed into a cooler box with ice packs before transportation to the laboratory for analysis.

**Preparation of samples for evaluation**

To prepare the suspected TB samples for analysis, the suspected tissues with lesions were trimmed of fat and then a 500-mg sample was collected. The sample was then minced with sterile scissors and homogenized in a sterilized glass homogenizer, after which 1 mL of phosphate buffer (pH 6.8) was added. After thorough mixing, 1 mL of 5% sodium hydroxide was added. This was mixed thoroughly and then incubated for 15 min at room temperature. To this mixture 10 mL of phosphate buffer was added and then centrifuged at 1500 g for 20 min. The pellet was collected and then resuspended in a final volume of 0.5 mL of phosphate buffer. This was used for inoculation into 2% Ogawa medium and preparation of slides for Ziehl-Neelsen (ZN) microscopy. Cultures were monitored for growth up to 8 wk at 37°C. Another 100 μL of the suspension was used to prepare DNA directly by using DNAzol reagent (Invitrogen, Carlsbad, CA, USA). The suspension was mixed with 1.0 mL of DNAzol reagent and mechanical disruption was used as previously described.19 DNA was extracted according to the manufacturer’s instructions. Genomic DNA from *Mycobacterium* bacterial cultures was also prepared from colonies using DNAzol and mechanical disruption as described earlier. The extracted DNA was then dissolved in 50 μL TE buffer consisting of 10 mM Tris/HCL (pH 8.0) and 1 mM EDTA.

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LAMP
LAMP reactions were performed in a total volume of 25 µL consisting of 30 pmol each of inner primers FIP and BIP, 5 pmol each of outer primers F3 and B3, 20 pmol each of loop primers FLP and BLP, 1.4 mM deoxynucleotide triphosphate, 0.8 M betaine, 20 mM Tris/HCl (pH 8.8), 10 mM KCl, 10 Mm (NH4)2SO4, 8 mM MgSO4 and 8 U Bst DNA polymerase (New England Biolabs, USA) with 2 µL sample DNA. The sequences of the primers used are shown in Table 1.

The primers used in our study have also been used by other workers to detect MTC in human sputum samples. The mixture was then incubated at 64°C for 1 hr in a thermal dry heat block (ALB 221; Iwaki, Tokyo, Japan). A negative control (buffer) and positive control were included in each run. Results were visualized with the fluorescence detection reagent (Eiken Chemical Co., Tochigi, Japan) according to the manufacturer's instructions.

Multiplex-PCR primers and conditions
Genomic DNA from Mycobacterium bacteri- al cultures was used as a template. Primer pairs for cfp32 (a specific gene for MTC), RD9 (region of difference 9 seen only in M. tubercu- losis and M. canettii), and RD12 (region of dif- ference 12 deleted in M. bovis, M. caprae, and M. canettii) were obtained from an earlier publi- cation. The general PCR recipe contained 7.4 µL H2O, 2 µL 10 x Taq buffer, 2 µL dNTPs (2.5 mM each), 0.2 µL Taq (Takara), 1 µL target DNA, 2.2 µL of 10 µM cfp32 primers, 0.7 µL of 5 µM RD9 primers, and 0.8 µL of 5 µM RD12 primers. Appropriate negative controls consisting of PCR mix without target DNA were included. The PCR was performed using the following program: denaturation for 1 min at 98°C followed by 35 cycles of 5 sec at 98°C, 20 sec at 58°C, and 1 min at 68°C with final extension for 5 min at 72°C in a thermalcycler (iCycler, Bio-Rad Laboratories Inc., CA, USA). All PCR products were identified by gel электро- phoresis in a 2.0% agarose gel and were visualized by ethidium bromide staining.

Results
A total of 388 carcasses were examined, comprising 358 cattle and 30 lechwe. Of these animals examined, 57 (15.9% CI: 12.4-20.2) cattle carcasses had lesions characteristic of TB in the tissues and organs while 6 (20% CI: 8.40-39.1) lechwe carcasses had such lesion exhibits as well. When these samples were subjected to ZN microscopy, 48 (84.2% CI: 71.6-92.1) and 6 (100% CI: 51.7-100) samples from the cattle and lechwe, respectively, were found positive. In the case of the cattle, 9 samples were found to be negative, from those observed with characteristic TB lesions. On culture of postmortem TB-positive samples, 53 (93.0% CI: 82.2-97.7) cattle and all lechwe observed at meat inspection were found to be positive for acid-fast bacilli. The positive sample specimens increased in number with the LAMP system. Furthermore, the positive reactions on LAMP were easy to determine with the naked eye, as shown in Figure 1.

The overall results for the detection of TB- positive samples are shown in Table 2. All of the culture positive samples were positive on the LAMP assay. The culture isolates were also subjected to the LAMP system and were found to be MTC. In terms of sensitivity, the LAMP system was 100% positive in culture positive samples. An additional sample not detected with culture was also found to be positive. The sensitivity of ZN microscopy was found to be 85.7%, while culture was 93.7%, and LAMP 95.2%, respectively. In all these diagnostic procedures, sensitivity was taken as the proportion of samples testing positive for Mycobacterium tuberculosis.

The multiplex-PCR was successfully used to identify the MTC species involved in cattle and lechwe. The amplicon of 786bp of the cfp32 region was observed as indicated in Figure 2, showing that the MTC species involved in these cases was M. bovis. All of the cultures were positive for this amplicon indicating the MTC strain in the domestic and wildlife animals to be M. bovis.

Discussion
We successfully applied the LAMP assay technique for the rapid detection of the MTC group directly from clinical samples. It is important to note that the diagnosis of TB in

Table 1. Primers for the specific detection of the Mycobacterium tuberculosis complex used in the study.

<table>
<thead>
<tr>
<th>Name of primer</th>
<th>Sequence</th>
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<tbody>
<tr>
<td>FIP</td>
<td>CACCAACGTGGTACTCATGCAAGCGGAAGGTCT</td>
</tr>
<tr>
<td>BIP</td>
<td>TCGGGATACGGCTGGACACACAGCATGCACCCGT</td>
</tr>
<tr>
<td>F3</td>
<td>CTGCTCATGGAGCGCAGG</td>
</tr>
<tr>
<td>B3</td>
<td>GCTCATCCACACCCGC</td>
</tr>
<tr>
<td>FLP</td>
<td>GTTGCCACTCAGAGTATCTCCG</td>
</tr>
<tr>
<td>BLP</td>
<td>GAAACTGGGTCTAATACCGG</td>
</tr>
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Table 2. Number and percentage of samples testing positive for Mycobacterium tuberculosis under different procedures.

<table>
<thead>
<tr>
<th>Sample source</th>
<th>Ziehl-Neelsen microscopy</th>
<th>Culture</th>
<th>LAMP</th>
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<tbody>
<tr>
<td>Cattle</td>
<td>48 (84.2%)</td>
<td>53 (93.0%)</td>
<td>54 (94.7%)</td>
</tr>
<tr>
<td>Lechwe</td>
<td>6 (100%)</td>
<td>6 (100%)</td>
<td>6 (100%)</td>
</tr>
<tr>
<td>Total</td>
<td>54</td>
<td>59</td>
<td>60</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>85.7%</td>
<td>93.7%</td>
<td>95.2%</td>
</tr>
<tr>
<td>95% CI</td>
<td>74.1-92.9</td>
<td>83.7-97.9</td>
<td>85.8-98.8</td>
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animals largely depends on the tuberculin skin testing and slaughterhouse surveillance for undetected infections. In addition, it would be useful to develop a rapid and sensitive method to apply to doubtful TB-suspected samples. Slaughterhouse surveillance of TB is one of the key tools for detecting and confirming suspected TB cases. This is aided by microscopic analysis of sample smears, which is a rapid method of detecting acid-fast bacilli. However, this method is limited in sensitivity and the ability to identify infecting species. The culture method can be used efficiently to diagnose TB, but it is time consuming as it requires a minimum of four weeks. Therefore, a test that combines the rapidity of microscopy and sensitivity of bacterial culture methods would facilitate easy diagnosis at slaughter houses and establishments requiring definite results at the shortest possible time. The test can be a valuable tool in veterinary diagnostics. In our study, there was efficient amplification of the LAMP system to detect MTC from suspected specimens from cattle and lechwe. These results clearly demonstrate the high sensitivity of the LAMP system in detecting TB as observed by others who have been working on human specimens. This result also suggests that LAMP may be superior to all procedures being used at the moment to detect TB in animals. One sample was detected as positive on LAMP although negative on culture. This could be attributed to low numbers of MTC viable cells in the section of the specimens under investigation. M. bovis was identified from the results of the multiplex-PCR, which targets three genetic regions that are cfp32, RD9 and RD12. In summary, our study clearly provides evidence that the LAMP assay is a method that allows direct detection of MTC from processed clinical/abattoir specimens. This means that the technique can also be applied in the field of veterinary medicine for rapid confirmation of suspected cases. The ease of use and high sensitivity of this assay may facilitate the diagnosis and confirmation of MTC at slaughter houses, hence improving surveillance and control of TB at herd levels and in veterinary medicine in general.

References