Application of PCR for the detection of bovine tuberculosis in cattle

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Abstract

The present study was carried out for the detection and identification of bovine tuberculosis (bTB) using polymerase chain reaction (PCR). A total of 10 suspected cattle of Savar and BAU dairy farm were examined. Lymphnode biopsy, nasal swabs and blood were collected. Smears from lymphnode biopsy and nasal swabs were made onto clear slides and stain with acid fast staining. Portion of lymphnodes were preserved at -20°C and extracted DNA for PCR analysis. Portion of lymphnodes and other tissues were also collected in 10% neutral buffered formalin for routine Hematoxilin and Eosin staining and acid fast staining. In this study, acid fast staining of lymphnodes and nasal smears failed to detect acid fast Mycobacterium. The genome of bovine Mycobacterium in the extracted DNA of lymphnodes which used in PCR reaction was amplified and yielded 600 bp amplicon. This study suggests that, the PCR technique is a useful and rapid diagnostic tool for the identification of bovine TB in dairy cattle. Amplification technology offers the potential for the diagnosis of TB in a few hours with a high degree of sensitivity and specificity.

Keywords: Bovine TB, acid fast, PCR

Introduction

Tuberculosis (TB) is an important zoonotic disease caused by an intracellular acid-fast organism Mycobacterium sp. It has been recognized from 176 countries as one of the important bovine diseases causing great economic loss (Hines et al., 1995; Martin et al., 1994; Samad, 2000). TB is a contagious disease, which can affect most warm-blooded animals, including human being (Radostits et al., 2000). Cattle, goats and pigs are the domestic species most susceptible to infection, while horses are relatively resistant to infection. In cattle, exposure to this organism can result in a chronic disease that jeopardizes animal welfare and productivity and in some countries leads to significant economic losses by causing ill health and mortality. Moreover, human TB of animal origin caused by M. bovis is becoming increasingly evident in developing countries Prasad et al., (2005).

The diagnosis of Mycobacterium-induced disease is most commonly made by direct sample examination and culture. Acid-fast bacilli detection has low sensitivity and specificity, and, at best, can only provide a preliminary diagnosis. Long time is necessary for M. bovis and M. tuberculosis growth and culture, nonfastidious microorganisms will eventually contaminate a small percentage of cultures (Kekkaku, 1998). Following preliminary screening of suspected samples using acid fast staining, isolation can be carried out in a bacteriological medium. However, cross-contamination among bovine carcasses, improper decontamination procedure and duration of isolation procedure (often 3 weeks and up to 8-10 weeks in liquid medium) jeopardizes the isolation of M. bovis. The lengthy duration of isolation procedure imposes an unavoidable delay in important decisions about outbreaks and of suspected herds put under restriction. Shorter time-span diagnostic procedures are required for quicker decision. Therefore, there is an urgent need for a rapid, safe, and reliable method to diagnose of bovine TB. The most promising technique for approaching this diagnostic dilemma is polymerase chain reaction (PCR). PCR has been used to amplify different regions of the mycobacterial genome, making it a good candidate for assisting with species identification in a variety of specimens. Indeed, several research groups have described different PCR protocols for Mycobacterium genome amplification. Based on these results, the US Food and Drug Administration (FDA) have approved the use of PCR as an aid for the diagnosis of TB in clinical samples. The rapid and accurate detection of Mycobacterium sp is of paramount importance in the effective management of TB in man and animal.
Materials and Methods

The samples were collected from suspected cattle of Savar and Bangladesh Agricultural University (BAU) dairy farms (previously PPD positive). The tuberculin test positive cattle of BAU dairy farm (n = 3) were euthanized with saturated MgSO$_4$ and a thorough postmortem examination was carried out to investigate the gross lesions. The samples (liver, heart, lung, lymphnode and spleen) for histopathologic examination were collected and preserved in 10% neutral buffered formalin. Lungs, lymphnodes and spleen were snap freeze and preserved at -20°C for PCR detection of TB. The formalin fixed tissues were processed, sectioned and stained with hematoxylin and eosin (H&E) stain and acid-fast stain (Luna, 1968).

Extraction of DNA: DNA was extracted by locally adopted conventional method (Buckingham and Flaws, 2007). Briefly, a total of 200mg grained sample (lymphnodes) was taken in the microcentrifuge tubes containing 200 µl cell lysis buffer and vortexed. The solution was centrifuged at 5000 rpm for 10 min to collect the supernatant. Equal volume of phenol chloroform isooamyl alcohol was added to the supernatant and vortexed. Then the solution was centrifuged at 15000 rpm for 3 min and the supernatant was collected. Thereafter 10 µl of 5M NaCl was added on the 90µl solution. Ice cool absolute alcohol of 250 µl was added, centrifuged at 15000 rpm for 10 min and the supernatant solution was collected. The solution was desalted twice by using 80% ethanol. The tube was allowed to air dry for 15 minutes and 25µl of nuclease free water was added and stored at 4°C. Thus the DNA samples were evaluated both quantitatively and qualitatively using spectrophotometer and agarose gel electrophoresis, respectively (Spectronic R GeneticsTM New York, USA).

Primer selection for PCR: The oligonucleotide primer used to detect the *Mycobacterium bovis* was listed in Table 1.

**Table 1. Primers and their sequences used in the study**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
<th>Reference</th>
<th>PCR products size (bp)</th>
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<tbody>
<tr>
<td>Forward</td>
<td>5'-CAGGGATCCACCATGTTTACTGCGGTTG-3'</td>
<td>Xiu-yun et al., 2006</td>
<td>600bp</td>
</tr>
<tr>
<td>Reverse</td>
<td>5'-TGGCGAATTCTTACTGCGCCGGGG -3'</td>
<td></td>
<td></td>
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Polymerase chain reaction (PCR): 50 µl PCR mix consisted of 25 µl of 2X Master Mix (GeNei™ PCR Master Mix Kit), 1 µl (10 pmol) of each primer, 3µl template DNA and 20µl nuclease free H$_2$O and 3µl water was added instead of DNA to the water control tube. DNA amplification was carried out in a thermal cycler (Master Cycler Gradient, Eppendorf, Germany) using the thermal profile: 98°C for 5 min, one cycle; 95°C for 1 min, 56°C for 1 min, 72°C for 1 min, 30 cycles; 72°C for 10 min, one cycle. After completion of PCR reaction the tubes were held at 4°C.

Electrophoresis: The amplified PCR products were separated electrophoresed in 1% agarose gel, stained with ethidium bromide and examined under UV light using an image documentation system (Spectronic R GeneticsTM New York, USA).

Results and Discussion

Necropsy findings of suspected cattle

The necropsy examination included enlarged and consolidated lungs with fibrin deposition (Fig. 1) and enlarged spleen (Fig. 3). Pipe steam liver was found in liver and the liver was severely necrosed (Fig. 2). Pipe steam liver indicates infection with fascioliosis which was appeared as risk factor for tuberculosis (Flynn et al., 2007).
Acid-fast staining of impression smears: Impression smears stained with acid fast staining did not reveal acid fast organism (Fig. 4). The undetectable levels of Mycobacterium sp in impression smear of upper respiratory tract suggests that diseased cattle seldom shed the organism at detection levels in nasal discharges (Michel et al. 2007). Lymphnode aspirates were also subjected for smear preparation and acid fast staining. The acid fast organism was not detected in the lymph node aspirates smear (Fig. 5). It was suggested that although the cattle was tuberculin positive, the lymph node was enlarged but the organism left undetected as well (Vitale et al., 1998). The results of this study indicated that detection of acid fast organism in smears preparation is not a valid technique.

Histopathological examination

Severe congestion and accumulation of fibrin in the lung parenchyma was seen. Mononuclear cellular infiltration, proliferation of fibrous connective tissue was seen in spleen (Fig. 6) and lungs (Fig. 7). The liver was cirrhotic and also showed granulomatous reaction. Acid fast organism was not seen in acid fast staining of lung, liver, spleen and lymphnode (Fig. 8 and 9) of a tuberculin +ve cow. The detection of acid fast organism in cattle positive to tuberculin test is not always convincing (Kekkaku, 2003).
Polymerase chain reaction (PCR): The DNA of *Mycobacterium bovis* was amplified using commercial GeNei™ PCR Master Mix Kit and genus specific primer. The DNA of bovine *Mycobacterium* in the extracted DNA in PCR reaction was amplified and gave product of 600 bp, as expected (Fig. 10).
This study provides evidence that PCR is a sensitive screening assay for the detection of *Mycobacterium bovis* DNA in lymph nodes of cattle (Taylor et al., 2007; Sreedevi and Krishnappa, 2003; Romero et al., 1995; Liébana et al., 1995). PCR can generally be used to diagnose bovine TB in field conditions.

**Conclusion**

The present study was conducted principally to develop an effective method for the rapid and sensitive diagnosis of bovine TB in dairy cattle. There are various methods for diagnosis of bovine TB. The definitive diagnosis relies on time-consuming, highly specialized and laborious biochemical tests. For eradication of bovine tuberculosis, a definitive diagnosis depends on the isolation of *Mycobacterium bovis*. PCR method is rapid, sensitive, and specific tool for diagnosis of bovine TB. In this study all the cattle were positive to tuberculin test but in acid fast staining acid fast organism were not seen in any cases of bovine TB. The result of PCR technique revealed that out of nine samples, seven (88%) gave amplified band indicating positive and higher sensitivity of the technique. Rest of the samples failed to yield amplicon specific for *M. bovis* infection and this could due to fact that the cattle may be infected with *Mycobacterium sp* other than *M. bovis*, require further investigation.

**References**


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