Apply traditional and molecular protocols for the detection of carrier state of visceral leishmaniasis in black Bengal goat

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Abstract: Visceral leishmaniasis (VL) is the second largest parasitic killer of human in the world after malaria which is responsible for an estimated 500,000 new cases of VL in each year with 8-10% mortality. The parasite is transmissible to humans and animals by the bite of phlebotomine sand fly. The clinical manifestations are highly diverse, humans and dogs are naturally infected, and the diseases are associated with several risk factors, yet to understand. The aims of this study were to apply traditional and molecular detection tools and more emphasis was given to identify goat as a carrier of visceral leishmaniasis. To demonstrate the promastigote and amastigote phases of Leishmania in tissues, traditional impression smear staining technique was used. For the confirmation of the species of Leishmania involved specific technique like polymerase chain reaction (PCR) was applied. A total of twenty goats were investigated and samples were tested using impression smear staining, histopathology and PCR. Blood smear and impression smears were prepared from spleen, liver, bone marrow and stained with Giemsa’s stain. Using Giemsa’s staining out of twenty goats investigated, six (6/20) were found to contain promastigote and amastigote stages of Leishmania in their visceral organs. Histopathological examination from the liver section of suspected goats showed degeneration, necrosis and non specific fibrous connective tissue proliferation compared to non-reactive goats. There was accumulation of macrophages in lymphoid follicles of spleen in five suspected goats. A highly sensitive and specific primers were used in PCR amplification with the extracted DNA from liver and spleen of suspected six goats. Results of PCR showed that two of them were generated 145bp amplicon selective for L. donovani in their liver and spleen.

Leishmaniasis has a great public health significance and the protozoa found in goats of Fulbaria Upazilla may possess threat for transmission in human and other animals, require further investigation.

Key words: Black Bengal Goats, Visceral leishmaniasis, Giemsa’s staining, Polymerase Chain Reaction (PCR)

1. Introduction

Leishmaniasis is a worldwide vector borne zoonotic disease caused by several species of the intracellular protozoan parasite. There are four forms of leishmaniasis including, visceral, cutaneous, diffuse cutaneous and mucocutaneous leishmaniasis [1]. Among them, visceral leishmaniasis remain in the top important vector borne zoonotic disease in many parts of the world and is caused by more than 20 species of the protozoan genus, Leishmania [2]. It is the second-largest parasitic killer in the world after malaria and responsible for an estimated 500,000 cases per year [3]. Visceral leishmaniasis (VL) is the most severe form of leishmaniasis and if left untreated, is usually fatal. Even with treatment, case fatality rates often exceed 10% in VL-endemic areas of Asia and Africa [4]. VL is a serious international public health problem affecting about 88 countries of the world, of which 66 being in the old world (Asia, Africa and Europe) and the remaining 22 in the new world (South & Central America). Leishmaniasis is considered as one of the most neglected disease in Bangladesh based on the limited resources invested in diagnosis, treatment, control and its strong association with the poverty [5]. In Bangladesh, sporadic kala-azar cases were reported in 1970 and an outbreak occurred in Pabna district in 1980 [6]. This is one of the major health problem because 34 out of 64 districts are at risk, where 80% patients died within five years of affliction due to lack of diagnostic and treatment facilities in rural areas [7]. The true incidence of kala-azar in Bangladesh is not well documented but assumed to be close to 40,000–45, 000 cases per year [8]. Ninety percent of total cases of VL in Bangladesh is reported in 10 districts during last 11 years (1994-2004) and of which more than 50% of cases were reported in Mymensingh District. Out of 12 Upazilla of Mymensingh five were confirmedly reported to have kala-azar patients. Among these five, Fulbaria Upazilla showed the highest incidence and becoming epidemic in this area. Using the population of the respective Upazilla as the denominator, the incidence of kala-azar in Fulbaria Upazilla ranged from 30 to 33/10,000/year [8].

Also VL is an endemic disease of great public health importance in some rural communities of Bangladesh. Dog, cat, jackle [9], rodent, cattle, goat may act as a silent carrier and could play role in disease
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transmission in human. Detection of normal parasites in a clinical and subclinical sample is necessary to confirm a suspected case of leishmaniasis. Most commonly used methods to diagnose VL is to examine organs (liver and spleen) stained with modified Giemsa’s [10]. PCR-based assays need to apply as highly sensitive and specific (upto 100%) detection tool to diagnose Leishmania parasite irrespective of species or genus [11]. DNA from liver, spleen, lymphnode, blood and bone marrow smears stained with Giemsa stain are usually the most effortable primitive diagnosis tool [12]. It needs to adopt and correlate findings of traditional and molecular detection tool to understand the accuracy of detection techniques as well as the epidemiology of VL.

Goat may act as a carrier of VL but the possible role of goat for VL transmission has not been studied in Bangladesh. The asymptomatic goats in VL endemic area could carry organism in the liver and spleen but the authenticity of this findings has not confirmed yet. Therefore, this study was designed to standardize traditional and molecular tools for the detection of leishmanial protozoa in black bengal goat and identify goat as a carrier of VL in Fulbaria Upazilla, Mymensingh, Bangladesh.

II. Materials And Methods

A total of twenty goats of ill health and both sexes were randomly selected and collected from Fulbaria Upazilla, Mymensingh during the period from January to May, 2013. Among them, fifteen goat samples were collected from slaughter house of Fulbaria Upazilla and another five goat samples were collected from suspected cases. Systemic dissection and investigation were carried out and liver, spleen, blood and bone marrow were collected.

2.1 Impression smears examination

Thin smears of blood and impression smears from liver, spleen and bone marrow were made on to clean grease free glass slides, dried in air and fixed in ice cold absolute methanol (acetone free) for 15 minutes. The slides were dried in air and placed in Coplin jar containing working Giemsa’s solution (Luna, 1968) and allowed to stain for 50 minutes. Slides were washed in running tape water for 30sec, dried in air and examined at low (10x) and high power (40x and 100x) microscopic field. Extensive investigation was carried out for the presence of promastigote and amastigote stages of leishmanial protozoa in blood, in tissue spaces and in macrophages of various organs were employed and protozoal morphology was investigated and images were captured.

2.2 Histopathological investigation

During necropsy, portion of liver and spleen were collected and fixed in 10% buffered neutral formalin for histopathological studies. Formalin fixed tissue samples were trimmed, processed, sectioned and stained with H&E (Luna, 1968). A low (10x) and high power (40x, 100x) microscopic investigation were carried out to observe the changes in the internal organs specific to VL.

2.3 Detection of species of Leishmania by PCR

2.3.1 DNA extraction

For DNA extraction from the liver and spleen of goats Wizard Genomic DNA Purification Kit (Promega) was used according to the manufacturer’s protocol. Briefly 100µl of 20% liver samples suspension was mixed with 600 µl of nuclei lysis solution, incubated for 15 min at 65 °C temperature and 3µl of RNase Solution was added to the nuclear lysate and mixed by inversion. The mixture was incubated for 30minutes at 37°C. 200µl of Protein Precipitation solution was added and vortexed vigorously for 20 s and centrifuged at 13000 rpm for 4 min and the precipitated protein was formed a tight white pellet. The supernatant was transferred to a new tube containing 600µl of room temperature isopropanol and centrifuged at 13000 rpm for 1 min. The supernatant was discharged and 600µl of 70% ethanol was added. After the centrifugation at 13000 rpm for 1 min, the ethanol was aspirated and the pellet air dried. The pellet was suspended in 100µl of the DNA rehydration solution. Then incubating at 65°C for 1 hour and finally stored the DNA at 4°C. The DNA samples were evaluated quantitatively and qualitatively using spectrophotometry (A260 and A280) and agarose gel electrophoresis. One set of primers (Table 1) were used to identify the species of leishmanial protozoa.

2.3.2 PCR

PCR reactions were performed onto each DNA sample in 25µl reaction volume containing 7 µl of Nuclease free water, 2x PCR master mix (Promega Corporation, USA), .25µl forward mix and .25µl reverse primers and 5 µl of DNA template. A total of 45 cycles of DNA amplification reaction for L. donovani was carried out. The thermal profile used for L. donovani comprised an initial denaturation for two minutes at 94°C followed by 45 cycles of DNA amplification reaction in a Master Cycler (Master Cycler Gradient, Eppendorf, Germany). The condition of PCR amplifications were denaturation for 60sec at 94°C, primer annealing for 90sec at 62°C and extension for 30sec at 70°C followed by a final extension for 10min at 70°C. The PCR reactions were

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finally terminated by adding 3μl 50mM EDTA and PCR products were analyzed by electrophoresed in 2% agarose gel, stained with ethidium bromide and examined under UV light using an image documentation system (Cell Biosciences, Alphalmager HP, USA).

Table 1: Primers and their sequences for Leishmania donovani used in this study

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
<th>Product size</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDFl(RV1) sense</td>
<td>5CTTTCTGTCCGCCGGTAGG-3</td>
<td>145bp</td>
<td>[13]</td>
</tr>
<tr>
<td>LDRI(RV2) antisense</td>
<td>5CCACCTGGCCTATTTTACCA-3</td>
<td></td>
<td></td>
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III. Results And Discussion

A total of twenty goats were examined in this study, where special fascination was focused to the detection of Leishmania protozoa in goats. For this, consequent studies on systemic necropsy of goats, blood smear and impression smears were performed from liver, spleen and bone marrow aspiration, systemic histopathology and PCR amplification was done using species (L. donovani) specific primers. In this study, Physical examination of the goats investigated revealed depression, loss of condition particularly decreased muscle mass over shoulders, hips, spine, serosanguineous nasal discharge, dull hair coat, splenomegaly and generalized lymphadenopathy. The goats were ill health, rough hair coat, alopecic but major gross changes characteristics of VL were not observed either in spleen, liver and bone marrow. However, in common the spleen appeared relatively larger. Blood smear and impression smears taken from the cut surface of liver, spleen and bone marrow stained with Giemsa’s stain and were examined under light microscope. From this smear morphologically two forms of leishmanial stage was found, one is promastigote form and another is amastigote. Promastigotes are slightly elongated and contain a single nucleus with an anterior flagellum originating from a kinetoplast while amastigotes are slightly round to oval, still contain a single nucleus and kinetoplast, but retain only a rudimentary flagellum. In this study, pear or sperm head shaped leishmanial promastigote was seen in impression smear of liver (Figure 2a). Impression smear of spleen revealed highest number of promastigote (Figure 2c). Dark colored amastigote inside the ruptured macrophage was seen in bone marrow aspirates (Figure 1a). In blood (buffy coat) smear promastigote stage of Leishmania was also observed (Figure 1b). The concentration of leishmanial promastigotes (per 10x microscopic field) appeared higher in the impression smears prepared from spleen (N=5-15), followed by liver (N=5-7), bone marrow (N=3). Using smear microscopy, the species of Leishmania involved whether pathogenic or non pathogenic remain undetected. However, comparative data on the to the morphologic analysis of the developmental stages of leishmanial protozoa in man and animals is very limited. [14] investigated clinically suspected cases of visceral leishmaniasis (83.4%) in canids and identified positive cases by examining bone marrow. However studies showed that bone marrow examination was the least sensitive method for the detection of visceral leishmaniasis [15,16,17]. This study showed that bone marrow examination was the least sensitive method for the detection of visceral leishmaniasis [15,16,17].

Histopathological examination was done in liver and spleen of six suspected goats. The results of histopathological examination revealed numerous rod shaped structure similar to promastigote stages of Leishmania in the liver. Numerous dark color bodies in Vonkupffer cell of liver was seen (Figure 3a) and these could be the amastigote stages of the protozoa. Histopathological examination of spleen revealed nodule like structure (Figure 3b) and containing infiltration of macrophage in lymphoid follicle. Notable changes in the bone marrow was not seen in this study. Histopathological study did not reveal much amastigote stage of protozoa in the macrophages of liver, spleen, blood and bone marrow. The presence of large number of promastigote in liver and spleen without tissue reaction and lacking of inflammation indicate the role of goats as a symptomatic carrier of VL. Scatteredly distributed granulomatous reaction were seen in liver parenchyma. The nodule of hepatic parenchyma contain infiltration of macrophage. There was lacking of caseous necrotic centre. Fibrotic patches were well documented along the hepatic parenchyma could provoke by dispatching of leishmanial promastigotes. There were numerous darkly stained promastigotes dispatched congested central vein indicating stage of disseminating of the protozoa. [18] reported the reason of the granuloma formation and presence of necrosis or proliferation of connective tissues and giant cells in chronically infected liver. There was a infiltration of macrophage in lymphoid follicle of spleen and nodule like structure was also seen. Early lesions with high parasitic load can show granuloma at the edge of lesions and deeper sections of the biopsy [19]. [20] described granulomatous leishmanial lesions in monkey, consisted of macrophages containing parasites, lymphocytes, plasma cells, and occasionally eosinophils. Differences were not observed between promastigote or amastigote derived nodular lesion or strains of leishmanial protozoa involved, course of infection and variation in either the sex and ages of the hosts and geography.

Many techniques such as isoenzyme electrophoresis, monoclonal antibodies, in situ hybridization, molecular karyotyping and restriction enzyme are currently used to detect Leishmania compared to PCR. These
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techniques require a larger quantity of sample material, involve a lengthy technical process and may present difficulties in the interpretation of results [21], [22], [23]. PCR assays have greatly improved the sensitivity and specificity to diagnose leishmanial infection in canids [24]. However, to make this technique achievable, the sampling methods need to be noninvasive, easy and painless. Using PCR on to the DNA from Conjunctival swab diagnosed symptomatic dogs successfully and observed 92% sensitivity [25]. However, [26] found 91.7% and [27] obtained between 73.9 and 95.6% sensitivity depending upon the PCR method used to diagnose VL. In this study as the animals were examined at necropsy and from slaughter house, PCR amplification (Table 1) with the DNA template from the liver and spleen of goats were carried out (Figure 4) using primers specific for L. donovani (Figure 4). PCR amplification with the DNA from these organs showed 145bp amplicon indicating infectivity of goats with L. donovani. This study provide evidence that the goats (N=20) at Fulbaria Upazilla were infected with L. donovani. Molecular detection using PCR to investigate the carrier state of VL in goats was reported by [28] where spleenomegali and fibrotic condition of liver were well demarcation of leishmaniasis. As there was lacking data regarding infectivity of goats with Leishmania, to the best of our knowledge may be this is the first study in Bangladesh describing occurrence of VL in domestic small ruminants. The result may come to an assumption that goats can be considered as an important carrier for leishmanial infection similar to other wild canids [29].

Fig.1 Impression smears prepared from bone marrow (a) and buffy coat (b) of blood of a goat and stained with Giemsa’s. Dark colored numerous amastigotes (red arrow, 100 X, left) was seen inside the ruptured macrophage. Promastigote (white arrow, 10 X, right) phase of Leishmania protozoa was seen in the smear prepared from blood sample.

Fig.2 Morphological investigation of promastigote stages of leishmanial protozoa (black arrow) in different organs impression smears prepared from goat and stained with Giemsa’s. The promastigote stage of the parasite was seen in the intercellular spaces and found to contain a tail (white arrow) and a head (black arrow) with variable morphology. Under oil immersion microscopy ballon shaped (a, black arrow), sperm head shaped (b, black arrow) and sometimes tadpole shaped (c, black arrow) appearance of the protozoa were seen.

Fig.3 Section of the liver and spleen of a goats (a & b) and stained with H&E. Dark color amastigote was seen (black arrow, 100X, left) in Vonkupffer cell of liver section (a). Scatteredly distributed granulomatus reaction were seen in liver parenchyma. The nodule of hepatic parenchyma contain infiltration of macrophage and closely packed collection of macrophagess replacing hepatic parenchyma and lacking caseous necrotic centre.

Nodule like structure (black arrow, 10X, right) in lymphoid follicle of spleen was seen.
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Fig. 4 Agarose gel electrophoresis of PCR products with the genomic DNA obtained from the visceral organs of goats. *Leishmania donovani* specific primers were found to amplify genomic DNA of *Leishmania* in PCR settings. Lane - 1, 2, 3, 4, 5, 6 containing DNA template from the liver of goats and Lane – 7, 8, 9, 10, 11, 12 containing DNA template from the spleen of goats. Lane – M containing 100bp DNA ladder, Lane – NC is for Negative control, Lane – PC is for Positive control. PCR amplification showed the presence of *L. donovani* in goats 6 and 8 by generating 145 bp amplicon.

IV. Conclusions

This study concludes that probably this is the first study in Bangladesh detecting *Leishmania sp.* in goats by using PCR. However further studies are needed by increases sample size & to identify genomic DNA in sand flies by using PCR which transmitting VL to other animals and human beings.

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References


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