Canine Babesiosis in Northwestern India: Molecular Detection and Assessment of Risk Factors


1. Introduction

Amongst the various prevalent canine vector-borne diseases, canine babesiosis is very common and clinically significant disease caused by intraerythrocytic apicomplexan protozoa belonging to genus Babesia, distributed worldwide, including India. Babesia species often referred to as piroplasms comprise two main species, B. canis and B. gibsoni, based on their size. B. canis is a large piroplasm (4-5 μm), which usually occurs as a single pear-shaped piroplasm or in pairs of merozoites divided by binary fission within the erythrocyte.

Previous studies, on the basis of differences in the geographical distribution, vector specificity, and antigenic properties [1, 2], recognized that large canine piroplasms are subdivided into three species, namely, B. canis transmitted by Dermacentor reticulatus (in Europe), B. vogeli transmitted by Rhipicephalus sanguineus (in tropical and subtropical regions), and B. rossi transmitted by Haemaphysalis elliptica (in South Africa). B. gibsoni has been found to be associated with infection of dogs in Asia, North America, northern and eastern Africa, and Europe [3-5]. It is a small parasite that commonly appears as individual ring forms or pyriform bodies ranging between 1.0 and 2.5 μm in size [3].

Clinically canine babesiosis has been found to result in a wide range of presentations from subclinical disease to serious illness characterised by fever, pallor, jaundice, splenomegaly, weakness and collapse associated with intra- and extravascular haemolysis, hypoxic injury, systemic inflammation, thrombocytopenia, and pigmenturia [6].

As far as the diagnosis of canine babesiosis is concerned, direct microscopic examination of the stained blood smear is the most commonly used method as it is conclusive, feasible, and cost effective diagnostic method but not necessarily detects parasites in dogs with unapparent or chronic infections since the level of parasitemia is very low [7]. As regards, the serological methods, indirect fluorescent antibody test (IFAT) and enzyme linked immunosorbent assay (ELISA) for B. gibsoni parasites, are considered to be highly sensitive, but...
only moderately specific because of antigenic cross-reactions to *B. canis* [8] and normal dog erythrocytes [8, 9]. Therefore, the development of highly specific and sensitive system for the diagnosis of canine babesiosis is still awaited. In this regard, recent advances in molecular biology techniques like polymerase chain reaction (PCR) have made it possible to detect and identify piroplasms with greater sensitivity and specificity than traditional methods [10, 11].

Regarding Indian scenario, though there are sporadic reports of canine babesiosis based on conventional diagnostic methods [12–15], the true status of canine babesiosis is still not clear barring few reports [16, 17] employing the PCR based assays. Furthermore, molecular detection of canine babesiosis has not yet been explored from Punjab, north state of India, so the present work was carried out to know the status of canine babesiosis in this part of the country through PCR based assays.

2. Materials and Methods

2.1. Geographical Area. The study was conducted from Ludhiana district of Punjab state, in the northwestern region of India. The climate of the region under study is excessively hot and dry during summers. Winters are cool with some frosts, and the average annual rainfall is 565.9 mm. These environmental conditions provide favourable and conducive conditions for the survival and propagation of ticks and *Rhipicephalus sanguineus* is the major tick infesting canines [18].

2.2. Samples. A total of 214 blood samples were collected aseptically from cephalic vein of the selected dogs in EDTA coated vials from the dogs presented to Small Animal Clinics, Teaching Veterinary Clinical Complex, GADVASU, Ludhiana, as well as local private veterinary clinics from a period of one year (April 2012 to March 2013). Dogs were selected on the basis of presence of naturally acquired tick infestation at the time of presentation and/or showing clinical signs in accordance with the haemoproteozoan infection, namely, fever, haemoglobinuria, anemia, and so forth. The collected blood samples were utilized immediately for the preparation of thin blood smears and were then kept at −20°C until DNA extraction.

Microscopic examination of blood samples was done after staining the prepared thin blood smears with Giemsa as per standard protocol [19] and examined under oil immersion objective of the microscope to detect the piroplasms and the results obtained were compared to that of PCR assay.

2.3. Genomic DNA Isolation. For conducting the PCR assay, genomic DNA was isolated from whole blood using QIAamp DNA blood mini kit (QIAGEN, GmbH, Germany) following the manufacturer’s recommendations with minor modifications and stored at −20°C till use. Genomic DNA of *B. gibsoni* was isolated and utilized as a positive control from infected blood sample showing parasitemia in blood smear examination. Genomic DNA was also isolated from the whole blood of infection-free puppy and used as a negative control along with nuclease-free water.

Table 1: Evaluation of diagnostic/screening PCR assays over blood smear examination.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>PCR (95% CI)</th>
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<tbody>
<tr>
<td>Sensitivity*</td>
<td>100% (78.47, 100)</td>
</tr>
<tr>
<td>Specificity*</td>
<td>90.5% (85.64, 93.83)</td>
</tr>
<tr>
<td>Diagnostic accuracy*</td>
<td>91.12% (86.55, 94.24)</td>
</tr>
</tbody>
</table>


2.4. PCR Protocol. The PCR assay was optimized targeting a portion of the 18S rRNA gene to amplify *B. gibsoni* as described by Inokuma et al. [20]. The sequences of the primers were as follows:

Gib599 Forward: 5’CTCGGCTACTTGCCTTGTC3’;

Gib1270 Reverse: 5’GCCGAAACTGAATAACGGC3’.

PCR assay in a final volume of 25 μL was carried out in a PCR thermal cycler (Applied Biosystems, USA). The master mix consisted of 2.5 μL of 10X PCR buffer (MBI Fermentas), 0.5 μL of 10 mM dNTP mix (MBI Fermentas), 1.5 μL of 25 mM MgCl2 (MBI Fermentas), 1.0 U of recombinant Taq DNA polymerase (MBI Fermentas), 1 μL each (20 pmol) of the primers, and 5 μL of template DNA isolated from field samples. The volume was made up to 25 μL with nuclease-free water. The PCR cycling conditions were initial denaturation at 95°C for 5 min, 40 cycles of denaturation at 95°C for 30 sec, annealing at 56°C for 30 sec, and extension at 72°C for 30 sec, and the final extension was performed at 72°C for 5 min. The PCR products obtained were checked for amplification by electrophoresis on a 1.5% agarose gel and visualized using gel documentation system (Syngene, UK). In order to check the specificity of the assays, isolated genomic DNA of large *Babesia, Ehrlichia canis, Hepatozoon canis*, and *Trypanosoma evansi* isolated from the microscopically positive cases were also employed in the PCR to see the amplification, if any.

2.5. Statistical Analysis. All data analyses were performed by using statistical software program (SPSS for Windows, Version 19.0, USA). Association between the prevalence of *B. gibsoni* by PCR and various risk factors, namely, sex, age, breed of the host, and season, was carried out by Chi square (χ²-test). Variables with significant association at *P* < 0.05 (two-sided) were subjected to the multivariate logistic regression model. The results were each expressed as *P* value and odds ratio (OR) with a 95% confidence interval (CI 95%).

3. Results

3.1. Blood Smear Examination. In the present study, examination of Giemsa-stained peripheral thin blood smears of 214 canines revealed an overall prevalence of canine babesiosis as 7.47% with 0.93% (2/214) positivity for the piroplasms of large
Table 2: Final logistic regression model for factors associated with prevalence of *B. gibsoni* by PCR on animal levels.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Regression coefficient (β)</th>
<th>Standard error (SE)</th>
<th>P value</th>
<th>Odds</th>
<th>CI (95%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>−1.330</td>
<td>0.199</td>
<td>0.000</td>
<td>0.398</td>
<td>0.080–1.799</td>
</tr>
<tr>
<td>Sex</td>
<td>0.346</td>
<td>0.151</td>
<td>0.022</td>
<td>0.849</td>
<td>0.403–1.791</td>
</tr>
<tr>
<td>Breed</td>
<td>0.179</td>
<td>0.200</td>
<td>0.371</td>
<td>3.345</td>
<td>1.045–10.710</td>
</tr>
<tr>
<td>Season</td>
<td>−0.223</td>
<td>0.186</td>
<td>0.230</td>
<td>2.143</td>
<td>0.788–5.830</td>
</tr>
</tbody>
</table>

*Confidence interval.

4. Discussion

In the present study by using conventional parasitological techniques, a statistically higher percent positivity was recorded for *B. gibsoni* infection than large *Babesia* (P = 0.0051) with the overall prevalence of canine babesiosis as 7.47% (16/214). Previously, from the same region, Eljadar et al. [21] examined a total of 951 suspected dog samples from Small Animal Clinics, GADVASU, Ludhiana, and three local private veterinary hospitals for haemoproteozoon infections and reported 1.26% samples to be positive for *B. canis* and 3.17% to be positive for *B. gibsoni*. The comparative higher prevalence of *B. gibsoni* over *B. canis* recorded by him is in congruence with that of the present study. Similar findings were recorded in earlier studies by Singh et al. [15, 22] from this region revealing the prevalence of *B. gibsoni* and *B. canis* in the range of 0.65%–8.26% and 1.43%–4.51%, respectively.

Microscopic detection of *B. gibsoni*, though smaller in size than large *Babesia*, was easier because of its frequent appearance in the circulating host blood. This might also be due to a low level parasitaemia in case of large *Babesia* infection especially during very early or carrier stage which is beyond the level of microscopic detection [6, 10, 23]. The prevalence of canine babesiosis from various parts of northern India has been reported to be ranging from 0.66 to 8.9% [14, 22, 24, 25] while from Southern India Senthil Kumar et al. [13] recorded 3.9% and 84.9% prevalence of *B. canis* and *B. gibsoni*, respectively. Wide variation in climatic conditions prevailing in different parts of India might be responsible for varying percentage of these tick borne infections.

On the basis of present findings PCR based assay was able to detect 15.42% prevalence of *B. gibsoni*. Higher detection of canine babesiosis by PCR based assays as compared to microscopy as observed in the present study has also been reported by several authors worldwide indicating the higher sensitivity levels of PCR [10, 17, 26–30]. As far as the detection of *B. gibsoni* with PCR based assays is concerned, many studies have been carried out worldwide and the prevalence has been recorded to be ranging from 3.3 to 55% [17, 20, 31–34].

As far as evaluation of various risk factors is concerned for canine babesiosis, several authors have observed the prevalence of the haemoproteozoon infections to be highest in young dogs [35, 36]. In terms of sex of the host, from the data obtained in the current study, it can be concluded that the assays recorded no statistical significance difference in the prevalence of the disease among males and female dogs. These results are incongruous with Amuta et al. [28] and Singh et al. [14].
Regarding breed of the host, the results revealed that blood smear examination and PCR detected a statistically nonsignificant difference in the prevalence of the *B. gibsoni* among the various breeds and nondescript dogs. In seasonal prevalence of the disease, the disease was most prevalent in warm seasons as compared to winters. The probable reason behind this trend may be correlated to the seasonal activity of the brown dog tick, *Rhipicephalus sanguineus* which is in its abundance in hot and humid period of the year, thus resulting in the higher incidence of haemoprotozoan infections in warm months during warmer seasons [37].

**Conflict of Interests**

The authors declare that there is no conflict of interests regarding the publication of this paper.

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**References**


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