Molecular detection of *Leishmania infantum* DNA according to clinical stages of leishmaniasis in dog

Deteção molecular de DNA de *Leishmania infantum* em diferentes estágios clínicos da leishmaniose em cães

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Abstract

The aim of this study was to compare molecular tests used to diagnose *Leishmania* spp. in dogs with different stages of infection. Blood and conjunctival swab (CS) samples from dogs classified in four clinical stages were subjected to different PCR protocols (13A/13B, MC1/MC2, LITSR/L5.8S and LEISH-1/LEISH-2 primers). To the study, 22.3% (48/215) of dogs were classified as without clinical signs, 67.5% (145/215) stage I (mild disease), 7.0% (15/215) stage II (moderate disease) and 3.2% (7/215) stage III (severe disease). The results showed that in blood samples, 13A/13B detected a significant higher number of positive dogs in stage I (25/145) and in total (42/215) (p≤0.05). However, when CS samples were tested, no difference was observed (p>0.05). On the other hand, in blood samples, MC1/MC2 detected significantly fewer positive dogs classified as without clinical signs (0/48), in stage I (0/145) and in total (1/215) (p≤0.05). Likewise, in CS samples, this primers showed also lower detection (1/215) (p≤0.05). So than, we can conclude that PCR on blood samples with 13A/13B primers has greater capacity to detect positive dogs, mainly at the initial of clinical disease than do other primers and MC1/MC2 are not a good choice to detect *Leishmania infantum* infection in dogs.

Keywords: Canine visceral leishmaniasis, dog, *Leishmania* spp., PCR, stages of disease.

Resumo

O objetivo deste estudo foi comparar testes moleculares usados para diagnosticar *Leishmania* spp., em cães apresentando diferentes estágios de infecção. Amostras de sangue e suabe conjuntival (SC) de cães classificados em quatro estágios clínicos foram submetidas a diferentes PCRs (primers 13A/13B, MC1/MC2, LITSR/L5.8S e LEISH-1/LEISH-2 primers). Para o estudo, 22,3% (48/215) dos cães foram classificados como sem sinais clínicos, 67,5% (145/215) estágio I (doença leve), 7,0% (15/215) estágio II (doença moderada) e 3,2% (7/215) estágio III (doença grave). Os resultados mostraram que, em amostras de sangue, 13A/13B detectou número significativamente maior de cães positivos no estágio I (25/145) e no total (42/215) (p<0,05). No entanto, quando as amostras de SC foram testadas, nenhuma diferença foi observada (p>0,05). Por outro lado, no sangue, MC1/MC2 detectou significativamente menos cães positivos sem sinais clínicos (0/48), em estágio I (0/145) e no total (1/215) (p<0,05). Da mesma forma, em amostras de SC, MC1/MC2 também apresentou menor detecção (1/215) (p<0,05). Assim, a PCR em amostras de sangue com 13A/13B tem maior capacidade de detectar cães positivos, principalmente no início da doença do que outros primers, e o par de primers MC1/MC2 não é uma boa escolha para detectar infecção por *Leishmania infantum* em cães.

Introduction

The leishmaniases are a group of zoonotic infectious diseases that affect humans and domestic and wild animals. They are caused by protozoa of the genus *Leishmania* (DESJEUX, 2004; WHO, 2016). Today, they are considered to be among the six most important types of infectious and parasitic diseases in the world (LANGONI et al., 2015). One hundred and two countries in East and Southeast Asia, the Middle East, North and East Africa, Southern Europe (Mediterranean) and Central and South America are considered endemic for leishmaniases (ALVAR et al., 2012; WHO, 2016). In these countries, it has been estimated that approximately 0.2 to 0.4 million new cases of visceral leishmaniasis (VL) and 0.7 to 1.2 million new cases of cutaneous leishmaniasis (CL) occur every year (WHO, 2016). These cases form a challenge for public health (LOPES et al., 2016).

The leishmaniases has been described in several domestic and wild animals, including in carnivores, didelphimorphs, rodents, equids and primates (ASHFORD, 1996; DUARTE et al., 2010; SOARES et al., 2013; BENASSI et al., 2017). In domestic environments, dogs are considered to be the main epidemiological reservoir, with great importance for maintenance of the zoonotic VL cycle (QUINNELL & COURTENAY, 2009; MAIA & CAMPINO 2011). The concern regarding canine visceral leishmaniasis (CVL) is that it is more prevalent than is VL in humans. Great numbers of parasites are harbored in the skin of dogs, which favors vector infection (ARIAS et al., 1996).

Despite the importance of making accurate diagnoses among infected dogs for effective disease control, this remains a challenge (FERREIRA et al., 2014). Currently, immunoenzymatic (ELISA) and immunochromatographic (Dual Path Platform, Bio-Manguinhos®) serological tests are recommended for diagnosing *Leishmania* spp. infection in dogs from Brazil (BRASIL, 2011). However, serological tests have limitations such as low diagnostic sensitivity in dogs without clinical sings and cross-reactions with other trypanosomatids (TRONCARELLI et al., 2009; GRIMALDI et al., 2012). As an alternative to indirect tests, molecular methods play an important role in confirmation of diagnoses, identification of causative *Leishmania* spp. and detection of carriers (GRAMICCIA, 2011).

The polymerase chain reaction (PCR) is the molecular method most used in these studies, and it enables selective amplification of target sequences in the parasite’s DNA (DEGRAVE et al., 1994). The technique has been shown to be an important tool for detecting and diagnosing *Leishmania* spp. infection (MAIA & CAMPINO, 2008; ASSIS et al., 2010; COURA-VITAL et al., 2011; SOLANO-GALLEGO et al., 2011; MOHAMMADIHA et al., 2013). Several molecular techniques, biological samples and target genes are used in detecting *Leishmania* spp. DNA (MAIA & CAMPINO, 2008). However, the results obtained by several research groups are quite different. Thus, PCR use is far from having been standardized for *Leishmania* spp. infection (LACHAUD et al., 2002). In Brazil, there is no consensus among researchers regarding the primers that would be best for making the diagnosis of *Leishmania* spp. infection (LOPES et al., 2016). According to Courtenay et al. (2002) there is a positive correlation between parasite DNA detection by PCR and clinical disease. Thus, with the aim to elucidated differences among molecular tests used to diagnose *Leishmania* spp., related to different clinical phases of infection, this study was performed.

Materials and Methods

Study area

This study was carried out on the canine population of four cities in the state of São Paulo, Brazil. In this state, 10,934 cases of cutaneous leishmaniasis (CL) and 2,546 cases of visceral leishmaniasis (VL) were registered over the last 26 years (BRASIL, 2018a, b). The cities of Pirassununga (21° 59’ 46” S and 47° 25’ 33” W) and Santa Cruz das Palmeiras (21° 48’ 49” S and 47° 15’ 23” W) are areas where CL is endemic with 117 and 6 human cases, respectively, registered over the last 17 years, but no VL cases were registered (BRASIL, 2018c, d, e, f). On the other hand, Ilha Solteira (20° 38’ 44” S and 51° 06’ 35” W) and Andradina (20° 53’ 46” S and 51° 22’ 46” W) are endemic regions for both CL and VL. Between 2001 and 2017, 3 and 18 human CL cases were registered in Ilha Solteira and Andradina, respectively (BRASIL, 2018c, d) and 12 and 98 cases of human VL were registered in the same cities (BRASIL, 2018e, f).

Samples

Between 2014 and 2015, biological samples were collected from 215 dogs. All the animals came from municipal kennels or from animal protection shelters belonging to animal protection organizations. Blood samples were collected directly from the cephalic or external jugular veins. Samples were collected in tubes containing EDTA (at a proportion of 1 mg/mL) for hemograms and DNA analysis, and in tubes with no anticoagulant for serological and biochemical tests. Hemograms and biochemical tests were done on the same day after sampling. Blood and serum samples were stored at -20 ºC until serological analysis and DNA extraction. Exfoliative epithelial cells were collected from both eyes using sterile swabs manufactured for bacteriological isolation. The conjunctival swab samples were stored at 4 ºC until DNA extraction.

This study was approved by the Ethics Committee for Animal Use, under registration number 7839110215.

Animal ranking

The clinical signs observed and the quantitative results from the biochemical and serological tests were used to rank the animals into four clinical stages: without clinical signs, mild disease (I), moderate disease (II) and severe disease (III), in accordance with an adaptation of the disease classification of Solano-Gallego et al. (2011), as described in Table 1.

Clinical examinations

At the time of the sample collection, the dogs were also subjected to clinical examinations. Thus, each animal was carefully examined regarding mucosal color, nutritional status, abdominal
and lymph node alterations (lymphadenopathy) and ocular and dermatological alterations: alopecia, dermatitis, onychogryphosis and conjunctivitis. All signs observed were recorded on an individual clinical sheet for each dog.

**Hematological tests**

Hemograms were performed using automated equipment (BC-2800 Vet®; Mindray) to obtain total counts of leukocytes, red blood cells, hemoglobin, hematocrit and platelets. Differential leukocyte counts and cell morphology evaluations were performed using Rosenfeld-stained blood smears. The hepatic and renal function of these dogs was assessed through measurements of alanine aminotransferase (ALT/GOT Liquiform®; LabTest, ref. 109-4/30), aspartate aminotransferase (AST/GOT Liquiform®; LabTest, ref. 109-4/50) and creatinine (CREATININA K®; LabTest, ref. 109-4/30), urea (UREA UV Liquiform®; LabTest, ref. 1008-4/30), aspartate aminotransferase (ASAT/GOT Liquiform®; LabTest, ref. 109-4/30), 2% agarose gel (CORTES et al., 2004; Amresco). sterile deionized water was used as a negative control. DNA samples extracted from Leishmania amazonensis (IFLA/BR/1967/ph8) and L. infantum (MCAN/BR/1984/CCC-17.481), which were provided by the Leishmaniasis Laboratory of the Oswaldo Cruz Institute (FIOCRUZ), Rio de Janeiro, were used as a positive control. Twelve microliters of sample buffer (10 mM Tris, 10 mM EDTA, 0.005% m/v bromophenol blue and 10% v/v glycerol) and subjected to electrophoresis on 1.5% agarose gel (CORTES et al., 2004) stained with SYBR Safe (Invitrogen). The run was performed in a thermocycler Veriti® (Applied Biosystems). Sterile deionized water was used as a negative control. DNA samples extracted from Leishmania amazonensis (IFLA/BR/1967/ph8) and L. infantum (MCAN/BR/1984/CCC-17.481), which were provided by the Leishmaniasis Laboratory of the Oswaldo Cruz Institute (FIOCRUZ), Rio de Janeiro, were used as a positive control. Twelve microliters of sample buffer (10 mM Tris, 10 mM EDTA, 0.005% m/v bromophenol blue and 10% v/v glycerol) and subjected to electrophoresis on 1.5% agarose gel (EL TAI et al. 2000) and 2% agarose gel (CORTES et al., 2004; RODGERS et al., 1991) stained with SYBR® Safe (Invitrogen). The run was performed in a 1x TBE buffer at 100 V (volts) for 60 min with DNA MW Marker 100-bp Ladder® (Amresco). A UV Photo Doc-I® (UVP) transilluminator was used to view the amplified products.

**Quantitative PCR (qPCR)**

To exclude false negatives stemming from PCR errors or sample degradation, a real-time PCR for the endogenous β-actin gene was performed as described by Manna et al. (2006) (Table 2).

<table>
<thead>
<tr>
<th>Clinical stages</th>
<th>Serology</th>
<th>Clinical signs</th>
<th>Laboratory findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without clinical signs</td>
<td>Negative</td>
<td>Absent</td>
<td>No clinicopathological abnormalities</td>
</tr>
<tr>
<td>Stage I</td>
<td>Negative to low positive antibody levels</td>
<td>Dogs with moderate clinical signs such as peripheral lymph adenomegaly or papular dermatitis</td>
<td>Usually without observed clinical and pathological abnormalities</td>
</tr>
<tr>
<td>Stage II</td>
<td>Low to medium positive antibody levels</td>
<td>Dogs, which apart from the signs listed in stage I, may present: diffuse or symmetrical cutaneous lesions such as exfoliative dermatitis, onychogryphosis, ulcerations, anorexia, weight loss, fever, and epistaxis</td>
<td>Creatinine &lt; 1.4 mg/dl</td>
</tr>
<tr>
<td>Stage III</td>
<td>Medium to high positive antibody levels</td>
<td>Dogs, in addition to the signs listed in stages I and II, may show signs originating from lesions of the immune complex: vasculitis, arthritis, uveitis, and glomerulonephritis. At very severe cases: pulmonary thromboembolism or nephrotic syndrome</td>
<td>Clinopathological abnormalities such as mild non-regenerative anemia, hyperglobulinemia, hypoaalbuminemia, serum hyperviscosity syndrome Creatinine &lt; 1.4 mg/dl</td>
</tr>
</tbody>
</table>

*EL = ELISA level.
Table 2. Name, target gene, nucleotide sequence and the references of the primers and probe used to the study.

<table>
<thead>
<tr>
<th>PRC Target</th>
<th>Primer</th>
<th>PCR Concentration</th>
<th>PCR Thermal Condition</th>
<th>Size</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>kDNA</td>
<td>13A (5’-GTGGGGGAGGGGCCTTTCT-3’)</td>
<td>200mM Tris-HCl; 500mM KCl; 1.5mM MgCl2; 0.31mM DNTPs; 0.26µM primer; 1U Taq Polimerase</td>
<td>94°C 3 min, 35 cycles (94°C 40s, 56°C 30s, 72°C 30s), 72°C 5 min</td>
<td>120bp</td>
<td>Rodgers et al. (1990)</td>
</tr>
<tr>
<td>Leishmania</td>
<td>13B (5’-ATTTTACACCAACCCCCAGT-3’)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ssu-rDNA</td>
<td>LITSR (5’-CTGGATCATTTTCGGATGATG-3’)</td>
<td>50mM Tris-HCl; 10mM KCl; 1.5mM MgCl2; 0.2mM DNTPs; 0.5µM primer; 2U Taq Polimerase</td>
<td>95°C 4 min, 35 cycles (95°C 30s, 53°C 30s, 72°C 1 min), 72°C 5 min</td>
<td>Sequencing</td>
<td>El Tai et al. (2000)</td>
</tr>
<tr>
<td>Leishmania</td>
<td>L5-8S (5’-TGATACCATTTTCGGATGATG-3’)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mammalian</td>
<td>β-actin_S (5’-CTGGGACACCATCTTCTA-CAA-3’)</td>
<td>500mM Tris-HCl; 10mM KCl; 1.5mM MgCl2; 0.31mM DNTPs; 0.26µM primer; 1U Taq Polimerase</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>β-actin gene</td>
<td>β-actin_AS (5’-GCCTCGGTCAGCA-3’)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fluoregenic probe (5’-CCACCGCGATGGTCCT-3’)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Amplification for *L. infantum* was performed in accordance with Francino et al. (2006) (Table 2). Both qPCRs were performed using a LightCycler® 480 II thermocycler (Roche). The standard reaction curve was obtained using canine DNA (for the endogenous β-actin gene) and a DNA sample extracted from *L. infantum* (MCAN/BR/1984/CCC-17.481) in a ten-fold serial dilution (for the *L. infantum* gene). The DNA concentration was estimated by measuring the absorbance at 260 and 280 nm in a DS-11 spectrophotometer (DeNovix). The analyses were performed in accordance with the standards established through MIQE (Minimum Information for Publication of Quantitative real-time PCR Experiments) (BUSTIN et al., 2009). Sterilized ultrapure water was used as a negative control.

DNA sequencing

After electrophoresis on 1.5% agarose gel, qPCR products for ITS-1 were removed from the gel and purified using the Illustrated GFX PCR DNA & Gel Band® purification kit (GE Healthcare), in accordance with the manufacturer's instructions. DNA sequencing was performed using 20 ng/µL of purified PCR product and 5 µM of each primer. The samples were sent to the DNA Sequencing Service of the Human Genome and Stem Cell Research Center, at the Biological Institute (IB) of the University of São Paulo (USP). Chromatograms obtained using forward and reverse primers were assembled through the Sequence Scanner 2 software v2.2 and compared using the Clustal W software (available in BioEdit Sequence Alignment Editor, version 7.1.11, Ibis Biosciences, Carlsbad, CA, USA). The BLAST program (ALTSCHUL et al., 1990) was used to analyze the nucleotide sequences (BLASTN), with the aim of searching for similar genes in international databases (GenBank) and comparing these with the sequences obtained here.

Data analysis

The chi-square test with a significance level of 5% was calculated using the R software, version 3.1.1 (R Development Core Team, 2014) in order to assess the associations between the positivity of dogs classified as without clinical signs, stage I (mild disease), stage II (moderate disease) and stage III (severe disease) in each PCR protocol (13A/13B, MC1/MC2, LITSR/L5.8S and LEISH-1/LEISH-2) for *Leishmania* spp. detection.

Results

In this study, all the dogs were ranked in accordance with the disease classification described above. Thus, 22.3% (48/215) were classified as without clinical signs, 67.5% (145/215) in stage I (mild disease), 7.0% (15/215) in stage II (moderate disease) and 3.2% (7/215) in stage III (severe disease).

All the DNA samples were positive for the endogenous β-actin gene. This confirmed the quality of the process of DNA extraction and proved that there were no inhibitors and/or false negative results in the molecular analysis. Regarding DNA extracted from blood samples, tests using primers 13A/13B detected a significantly
higher number of positive dogs in stage I (25/145) and in total (42/215) than did the other tests (p<0.05). On the other hand, the primers MC1/MC2 detected significantly fewer positive dogs classified as without clinical signs (0/48) or in stage I (0/145) and in total (1/215) (p<0.05). Although LITSR/L5.8S showed a higher number of positive animals (21/215), there was no significant difference between the results from use of these primers and those from qPCR with LEISH-1/LEISH-2 (15/215), independent of clinical stage (p>0.05) (Table 3).

Regarding DNA extracted from conjunctival swab samples from the same dogs. The 13A/13B set of primers presented greater detection of positive animals in total (28/215) and also of dogs classified as without clinical signs (5/48) or in stage I (16/145), but no significant difference was observed (p>0.05). Like in the blood samples, the MC1/MC2 primers detected a statistically lower number of positive animals (1/215) than did the other tests on the swab samples (p≤0.05) (Table 4).

### Discussion

Here, we tested four different molecular tests for detecting *Leishmania* spp. infection, on 215 dogs that had been classified into four different clinical stages. Our results showed that the primers 13A and 13B (RODGERS et al., 1990) detected a greater number of positive dogs at the onset of the clinical signs of disease (stage I) and in total among the samples, with no significantly greater detection among without clinical signs dogs than was seen among the other primers tested.

The primers 13A/13B, MC-1/MC-2 and LEISH-1/LEISH-2 are sequences that target kinetoplast minicircle DNA (kDNA), while the primers LITSR/L5.8S are designed to target the genomic region of ribosomal (rDNA) and, more specifically, the internal transcribed spacer 1 (ITS-1). In a comparison of six primers for different gene targets, in relation to diagnosing leishmaniases, Kolta and others (2016) reported that PCR using the kDNA gene target presented good sensitivity and specificity. Their results have also been confirmed by other authors (REAL et al., 1999; LACHAUD et al., 2002; BENSOUSSAN et al., 2006; MOREIRA et al., 2007; ESMAEIL et al., 2011; LOPES et al., 2016).

In our analysis, kDNA-PCR was able to detect positive dogs in all types of samples. This was seen especially when the primers 13A/13B were used. These detected a significantly larger number of positive dogs in blood samples (p<0.05) and a larger but non-significant number, in conjunctival swabs samples (p>0.05). In several molecular surveys targeting *Leishmania* spp. kDNA, the diagnosis was reported to be sensitive (REAL et al., 1999; MIRO et al., 2008; SOLANO-GALLEGO et al., 2011). According to Miró et al. (2008), the sensitivity of PCR protocols can be correlated with the number of copies of the amplified DNA region that are present in the parasite. Each parasite has around 10,000 minicircles with a conserved region of approximately 200 bp and a variable region of 600 bp (MOHAMMADIHA et al., 2013). Specifically, the gene locus of the primers 13A/13B is present in thousands of copies in the cellular genome of the parasite (LIU et al., 2005). Thus, this makes the kDNA the most sensitive target for making the molecular diagnosis of the parasite (MOHAMMADIHA et al., 2013), for both clinically and subclinically infected dogs.

### Table 3. Molecular tests for *Leishmania* spp. detection in blood samples from dogs classified in different clinical stages.

<table>
<thead>
<tr>
<th>Clinical Classification</th>
<th>13A/13B</th>
<th>Conventional PCR</th>
<th>LITSR/L5-8S</th>
<th>Real-Time PCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without clinical signs (N= 48)</td>
<td>9 a</td>
<td>0 b</td>
<td>6 a</td>
<td>4 ab</td>
</tr>
<tr>
<td>Stage I (N= 145)</td>
<td>25 a</td>
<td>0 c</td>
<td>10 a</td>
<td>7 b</td>
</tr>
<tr>
<td>Stage II (N= 15)</td>
<td>4 a</td>
<td>0 a</td>
<td>1 a</td>
<td>0 a</td>
</tr>
<tr>
<td>Stage III (N= 7)</td>
<td>4 a</td>
<td>1 a</td>
<td>4 a</td>
<td>4 c</td>
</tr>
<tr>
<td>Total (N= 215)</td>
<td>42 a</td>
<td>1 ab</td>
<td>21 b</td>
<td>15 b</td>
</tr>
</tbody>
</table>

Proportion test (chi-square). Different lowercase letters (a, b, c) indicate statistical significance between columns at the 5% probability level (p ≤ 0.05). 13A/13B: *Leishmania* spp.; MC1/MC2: kDNA *Leishmania infantum*; LITSR/L5-8S: SSU-rDNA *Leishmania* spp.; LEISH-1/LEISH-2: kDNA *Leishmania infantum*.

### Table 4. Molecular tests for *Leishmania* spp. detection in conjunctival swab samples from dogs classified in different clinical stages.

<table>
<thead>
<tr>
<th>Clinical Classification</th>
<th>13A/13B</th>
<th>Conventional PCR</th>
<th>LITSR/L5-8S</th>
<th>Real-Time PCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Without clinical signs (N= 48)</td>
<td>5 a</td>
<td>0 a</td>
<td>4 a</td>
<td>3 a</td>
</tr>
<tr>
<td>Stage I (N= 145)</td>
<td>16 a</td>
<td>0 b</td>
<td>10 a</td>
<td>14 a</td>
</tr>
<tr>
<td>Stage II (N= 09)</td>
<td>4 a</td>
<td>0 a</td>
<td>1 a</td>
<td>3 a</td>
</tr>
<tr>
<td>Stage III (N= 13)</td>
<td>3 a</td>
<td>1 a</td>
<td>2 a</td>
<td>4 a</td>
</tr>
<tr>
<td>Total (N= 215)</td>
<td>28 a</td>
<td>1 b</td>
<td>17 a</td>
<td>24 a</td>
</tr>
</tbody>
</table>

Proportion test (chi-square). Different lowercase letters (a, b, c) indicate statistical significance between columns at the 5% probability level (p ≤ 0.05). 13A/13B: *Leishmania* spp.; MC1/MC2: kDNA *Leishmania infantum*; LITSR/L5-8S: SSU-rDNA *Leishmania* spp.; LEISH-1/LEISH-2: kDNA *Leishmania infantum*.
affected dogs (LACHAUD et al., 2002; MOREIRA et al., 2007). In our study, the primers 13A/13B showed higher sensitivity for detecting *Leishmania* spp. kDNA from dogs with clinical signs (stages I, II and III) and without clinical signs. However, there was a significant difference (p<0.05) only in relation to those classified in stage I, thus showing that this technique is useful for detecting the early stages of CVL.

Although also targeting the kDNA, the primers MC-1/MC-2 were less able to detect positive dogs through blood and CS samples, in comparison with 13A/13B (p<0.05). This may demonstrate that MC-1/MC-2, which is a set of primers that is considered specific for *L. infantum* kDNA, has lower sensitivity, such that it can only detect animals with higher parasite loads (RAMOS et al., 2012).

In addition, the primers LEISH-1/LEISH-2 are considered efficient for use in qPCR by many authors (FRANCINO et al., 2006; RAMOS et al., 2012; MOHAMMADIHA et al., 2013) and are similar to the primers 13A/13B. However, the TaqMan-MGB probe that was used determines specificity for *L. infantum* DNA (FRANCINO et al., 2006). For this reason, our results showed lower detection of positive dogs through blood samples, compared with what was shown by 13A/13B (p<0.05). In relation to conjunctival swab samples, these primers sets did not differ statistically. Interestingly, qPCR detected a higher number of positive dogs through CS (24/215) than through blood samples (15/215). This may have been due to differences in the distribution of the parasites, among tissues and the local immune response (MAIA et al., 2009). In addition, the presence of PCR inhibitors in blood and low parasite loads could explain these results (REALE et al., 1999; LACHAUD et al., 2002).

In fact, conjunctival swabs have been shown to be a sensitive and practical method for collecting samples. This method provides consistent diagnoses through cPCR (LEITE et al., 2010; OLIVEIRA et al., 2015; PEREIRA et al., 2016) and through real-time PCR (SOLANO-GALLEGO et al., 2011; FERREIRA et al., 2014; LEITE et al., 2015; BENASSI et al., 2017), in both symptomatic dogs (STRAUSS-AYALI et al., 2004; FERREIRA et al., 2008; PILATTI et al., 2009) and asymptomatic dogs (LEITE et al., 2010).

Comparison of diagnoses made using specific primers for *L. infantum* kDNA (MC-1/MC-2 and LEISH-1/LEISH-2) showed that the MC set presented significant lower detection of positive animals from both the blood and the CS samples. Ramos et al. (2012) also observed the same result using bone marrow, spleen and lymph node tissue to compare the diagnoses with these same primers and protocols. In addition, the MC set detected positive dogs only at the most severe stage of disease (stage III) (p<0.05). On the other hand, the LEISH set was able to diagnose *L. infantum* kDNA at all stages (without clinical signs, stages I, II and III). These divergent results may have been due to the lower sensitivity of the primers MC-1/MC-2, such that high parasite loads in blood and tissues would be needed (RAMOS et al., 2012). In turn, qPCR using LEISH-1/LEISH-2 was able to detect parasite DNA even in samples with small amounts of genetic material (RAMOS et al., 2012). Although Mohammadiha et al. (2013) did not find any significant relationship between clinical signs and amounts of *Leishmania* spp. kDNA in the blood, other authors showed that there was a positive relationship (MANNA et al., 2006; REIS et al., 2006; RODRIGUEZ-CORTES et al., 2010).

Regarding the rDNA target region, the primers LITSR/L.5.8S were designed to target the ITS-1 genomic region between the 18S rRNA and 5.8S rRNA genes of the parasite (SALLOUM et al., 2016). Our results showed that 21 dogs were cPCR-positive in blood samples and, although this was a notable detection rate, it was lower than the number of dogs that were found to be positive through cPCR using the 13A/13B set (p<0.05). Bensousan et al. (2006) compared use of ITS-1 PCR with use of PCR on the mini-exon region (ME-PCR) and found that ITS-1 was more sensitive for diagnosing cutaneous leishmaniasis. Subsequently, Koltas et al. (2016) observed that ITS-1 had a better detection rate than that of ME-PCR, but that both of these had lower sensitivity than small subunit rRNA (SSU rRNA). This was supported by other authors, who found that SSU rRNA had better performance in diagnosing leishmaniasis (SCHÔNIAN et al., 2003; ALBUQUERQUE et al., 2017). Schônian et al. (2003) reported that ITS-1 PCR was able to detect 0.2 parasites in samples.

However, the main limitation of use of ITS-1 is that it detects *Leishmania* only at the genus level, such that additional tools are needed for species identification (ALBUQUERQUE et al., 2017). In fact, to distinguish between species, the ITS-1 PCR products need to be subjected to sequencing analysis (TENÓRIO et al., 2014; SALLOUM et al., 2016) or to the restriction fragment length polymorphism (RFLP) technique (SCHÔNIAN et al., 2003; SALLOUM et al., 2016). Although the ITS-1 region is a conserved region of DNA, it has several polymorphisms that allow identification of species (ROELFSEMA et al., 2011). Identification at species level is very important in areas in which several *Leishmania* species are implicated in the leishmaniasis cycle (TSOKANA et al., 2014; ALBUQUERQUE et al., 2017). Thus, several studies have used the L.5.8SR/LITS set to distinguish between *Leishmania* species (SCHÔNIAN et al., 2003; TENÓRIO et al., 2014; BABUADZE et al., 2016; SALLOUM et al., 2016). ITS-1 PCR is considered to be the next most sensitive discrimination method after RFLP (KOLTAS et al., 2016) or sequencing (SALLOUM et al., 2016). In our study, we used sequencing followed by cPCR, and it was possible to identify some sequences with 99% or 100% similarity to *L. infantum*.

Lastly, it should be remembered that despite the efforts of many research groups, the continuing lack of a gold-standard test for diagnosing *Leishmania* spp. infection is a major impediment (RODRIGUEZ-CORTES et al., 2010). Molecular tools such as PCR have improved epidemiological studies on leishmaniasis around the world (GRAMICCIA, 2011) and can help in research on disease control (SCHÔNIAN et al., 2008). We performed four different PCRs to detect *Leishmania* spp. infection in dogs at several clinical stages and our results showed variation in sensitivity between the primer sets used. The sensitivity of each technique may vary over the course of evolution of the disease and between individuals; therefore, a combination of serological and molecular tests may be necessary to correct the diagnosis (ASSIS et al., 2010; NUNES et al., 2007; QUEIROZ et al., 2010; SILVA et al., 2014). In addition, it has been recommended that the diagnosis of this disease should be based on clinical signs and on the epidemiological characteristics of the region, which may contribute towards correctly identifying truly positive animals (SILVA et al., 2014).
Conclusion

Our results emphasize that using associations of more than one technique and/or type of sample are necessary in order to better detect *Leishmania* spp. infection. Moreover, cPCR using the primers MCI/MC2 does not readily detect infected dogs; while cPCR using the primers 13A/13B is better for detecting *Leishmania* spp. infection in blood, mainly at the onset of dog’s disease.

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