Natural canine infection by *Leishmania infantum* and *Leishmania amazonensis* and their implications for disease control

Infeção natural por *Leishmania infantum* e *Leishmania amazonensis* e suas implicações para o controle da doença

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Abstract

Leishmaniasis is a major public health problem worldwide. Because *Leishmania* can adapt to new hosts or vectors, knowledge concerning the current etiological agent in dogs is important in endemic areas. This study aimed to identify the *Leishmania* species detected in 103 samples of peripheral blood from dogs that were naturally infected with these protozoa. The diagnosis of leishmaniasis was determined through parasitological examination, the indirect enzyme-linked immunosorbent assay (ELISA) and the polymerase chain reaction (PCR). The *Leishmania* species were identified by means of PCR-restriction fragment length polymorphism (PCR-RFLP). The samples were subjected to PCR using oligonucleotide primers that amplify the intergenic region ITS1 of the rRNA gene in order to identify the species. The amplified DNA was digested using the restriction enzyme HaeIII. A restriction profile identical to *L. amazonensis* was shown in 77/103 samples and the profile was similar to *L. infantum* in 17/103. However, a mixed profile was shown in 9/103 samples, which impeded species identification. In conclusion, the infection in these dogs was predominantly due to *L. amazonensis*, thus indicating that diagnosing of cases of canine leishmaniasis needs to be reexamined, since the causative agent identified is not restricted to *L. infantum*.

Keywords: Canine, *Leishmania* spp., leishmaniasis, zoonosis.

Resumo

Leishmaniose é um grande problema de saúde pública global. Devido à adaptação de *Leishmania* a novos hospedeiros ou vetores, conhecimentos sobre o agente etiológico atual em cães é importante em áreas endêmicas. Este estudo teve como objetivo identificar as espécies de *Leishmania* detectadas em 103 amostras de sangue periférico de cães naturalmente infectados com este protozoário. O diagnóstico de leishmaniose foi determinado por exame parasitológico, ensaio imunoenzimático (ELISA) e a reação em cadeia da polimerase (PCR). A identificação das espécies de *Leishmania* foi realizada por PCR – seguido da análise do polimorfismo no comprimento de fragmentos de restrição (PCR-RFLP). As amostras foram submetidas a PCR utilizando-se iniciadores oligonucleotídicos que amplificam a região intergénica ITS1 do gene de rRNA para identificar as espécies, o DNA amplificado foi digerido com a enzima de restrição HaeIII. Observou-se que 77/103 amostras mostraram um perfil de restrição idênticos a *L. amazonensis*, 17/103 foram semelhantes para *L. infantum*. 09/103 mostraram um perfil misto, o que impediu a identificação da espécie. Em conclusão, a infeção nestes cães era predominantemente devido a *L. amazonensis*, indicando que o diagnóstico de casos de leishmaniose canina precisa ser reexaminada, já que o agente causador não está restrito a *L. infantum*.

Introduction

Leishmaniasis is one of the principal global public health problems. Protozoa of the genus *Leishmania* are distributed worldwide and the epidemiology of the leishmaniasis depends on the characteristics of the species involved (WHO, 2013). Their geographical distribution is limited by vector distribution and, over recent years, the number of cases has continually increased (WHO, 2013).

Leishmaniasis is a disease characterized by damage to the skin, mucosa and viscera, according to the species of *Leishmania* and the host immune response. The disease can be caused by over 30 species of *Leishmania* and transmission occurs through the bites of sandflies of the genera *Phlebotomus* and *Sergentomys* in the Old World and *Lutzomyia* in the New World (BATES, 2007), although other possible vectors (COUTINHO et al., 2005; DANTAS-TORRES et al., 2010; FERREIRA et al., 2009) and transmission via blood transfusion have been reported (OWENS et al., 2001).

Visceral leishmaniasis (VL) can be fatal, mucocutaneous leishmaniasis causes mutilation and diffuse cutaneous leishmaniasis (CL) has a long disease course due to cellular immune deficiency. CL can be debilitating when the patient presents multiple lesions (ALVAR et al., 2004; DESJEUX, 2004).

VL mainly affects domestic dogs, which are the main reservoir of the parasite for humans (DANTAS-TORRES, 2007). There have also been reports of infection in cats (SOBRINO et al., 2008), wild canids (LIMA et al., 2009) and horses (DANTAS-TORRES, 2007). In CL transmission foci, dogs and other animals are not exposed as reservoirs but as accidental hosts, together with humans (DANTAS-TORRES, 2007).

Therefore, knowledge concerning the *Leishmania* species circulating in dogs in endemic areas is important. To diagnose the disease in dogs, the most commonly used methods are parasitological and/or serological tests, but these do not determine the species of the parasite (DEGRAVE et al., 1994).

Visceral leishmaniasis has spread out across the entire Brazil, with autochthonous cases reported in 25% of Brazilian municipalities in 21 states (WERNECK, 2014). The current strategies for the prevention and control of VL applied by the Brazilian Ministry of Health of March 14, 1963, which states pets infected with VL should be put down.

 Immediately following this, 2 ml of peripheral blood were collected by means of jugular vein puncture and were stored in tubes containing EDTA. One milliliter was used for DNA extraction and the remainder was centrifuged at 3000 rpm for 15 min at 4 °C in order to obtain plasma for detection of anti-*Leishmania* antibodies through the enzyme-linked immunosorbent assay (ELISA). The crude *Leishmania infantum* antigen preparation and the ELISA procedure were performed as previously described by (LIMA et al., 2003). This study was approved by the Research Ethics and Animal Experimentation Committee of the Araçatuba School of Veterinary Medicine of São Paulo State University (FMVA/UNESP), under protocol no. 002232.

DNA extraction from blood samples

DNA extraction from peripheral blood samples from the dogs was performed using the DNAeasy® commercial kit (QIAGEN, Valencia, California, 91355, USA), following the manufacturer’s recommendations. The DNA was quantified using a spectrophotometer at 260/280 nm (NanoDrop Technologies ND 1000 UV/VIS, USA), to evaluate the degree of purity, and was then stored at −20 °C.

Polymerase chain reaction

Determination of the *Leishmania* species was performed by means of PCR-RFLP. The initial step involved PCR with primers that amplified the intergenic region ITS1 of the rRNA gene (5′ AGCTGGATCATTTTCCGATG 3′, 5′ TATGTTGACGGTTATCCAGGC 3′).
The PCR reaction was performed with 100 ng of purified DNA, 12.5 µl of JumpStart Taq ReadyMix (20 mM of Tris-HCl at pH 8.3, 100 mM of KCl, 3 mM of MgCl₂, 0.4 mM of each dNTP dATP, dCTP, dGTP, dTTP, 0.1 U/µl of Taq DNA polymerase and JumpStart Taq antibody) (JumpStart™ Taq ReadyMix™, Sigma-Aldrich®, Inc., St. Louis, MI 63103, USA), 10 pmol of each oligonucleotide primer and 6.5 µl of ultrapure water, in a final reaction volume of 25 µl.

PCR was performed under the following conditions: an initial denaturation cycle at 94 °C for 60 s, followed by 35 cycles of denaturation at 94 °C for 30 s, primer annealing at 56 °C for 30 s and extension at 72 °C for 40 s.

The amplified fragments were viewed using a transilluminator on 2% agarose gel stained with ethidium bromide. Samples were considered positive when an amplified fragment of 227-268 bp was detected. For Leishmania species identification, the amplified DNA was then digested with the restriction enzyme HaeIII: 2 U for 1 h at 37 °C (Restriction Endonuclease HaeIII; Sigma-Aldrich®, Inc., USA). The digested product was then viewed using a transilluminator on 2% agarose gel stained with ethidium bromide, and was documented on 10% polyacrylamide gels stained with red gel (Uniscience Ltd., Cambridge, UK). The restriction profile of the sample was compared with the standard PCR restriction product obtained when using the species L. infantum (IOC/L2906-MHOM/BR/2002/LPC-RPV), L. braziliensis (IOC/L0566- MHOM/BR/1975/M2903) and L. amazonensis (IOC/L0575-MHOM/BR/1967/PH8) as positive controls and water as the negative control, thus enabling identification of the species of Leishmania. A 100 bp molecular weight marker was used (Invitrogen, Carlsbad, CA 92008, USA).

Results

Indirect ELISA and PCR

All the plasma samples (n = 154) were subjected to indirect ELISA to detect anti-Leishmania antibodies. Among these, 76.6% (n = 118) showed positive results. However, PCR confirmed infection in only 87.3% (n = 103) of the samples that were positive according to indirect ELISA.

PCR-RFLP

The samples positive for ELISA and PCR were subjected to PCR-RFLP and the profile of the restriction fragments indicated that 74.8% (n = 77) of the samples were infected by L. amazonensis, 16.5% (n = 17) by L. infantum and the remaining samples (8.7%; n = 9) exhibited a mixed profile, in which two fragments of less than 100 bp that were identical to L. infantum and two fragments of more than 100 bp that were identical to L. amazonensis were observed. Thus, for these specimens, species identification using this technique was not possible. A representative polyacrylamide gel showing PCR-RFLP is shown in Figure 1.

Discussion

This study was conducted to characterize Leishmania species in dogs in the city of Bauru, SP, Brazil. We observed that the species L. amazonensis predominated (74.8%) in the peripheral blood samples from the dogs evaluated in this city.

The high frequency of infection caused by L. amazonensis (74.8%) seen in the dogs in this study showed that the disease that is treated in many cases as canine VL (caused by L. infantum) is in fact infection by L. amazonensis, which causes the cutaneous form of the disease. It also highlights the affirmation that serological diagnoses cannot differentiate visceral from cutaneous infection (TOLEZANO et al., 2007). The Brazilian leishmaniasis control program recommends that seroreactive dogs should be diagnosed and culled as a control measure for VL, and although there is no effective control measure for cases of CL, the program also suggests that the parasite species may be identifiable in areas of transition or simultaneous occurrence of CL and VL. Parasite species identification needs to be mandatory, so as to avoid unnecessarily putting dogs down. Therefore, leishmaniasis control programs should not be based solely on serological tests, because the technique that has been used does not enable species identification.
Given the high frequency of *L. amazonensis* observed in dogs, these occurrences could be contributing towards increasing numbers of human cases of CL in large cities like Bauru and in adjacent cities (CVE, 2004). Although infection by *L. amazonensis* in humans is not considered common, it has been identified in the Northeast, Southeast, South and Center-West regions of Brazil (ASHFORD, 2000; DORVAL et al., 2006; GRIMALDI et al., 1989; GRISARD et al., 2000; LAIISON, 1985; LAIISON & SHAW, 1987; PASSOS et al., 1999).

Despite wide distribution of leishmaniasis in the state of São Paulo, little is known concerning its clinical and epidemiological features. However, it is known that the distribution of *Leishmania* species is subject to the presence of vectors and/or their reservoirs. Moreover, infection in humans depends on their relationship with the chain of parasite transmission (ASHFORD, 2000; LAIISON, 1985). The latest surveys of phlebotomine fauna conducted in some areas of São Paulo have determined that *Lutzomyia intermedia* is the most abundant species in this state (SHIMABUKURO & GALATI, 2011; SILVA et al., 2012), which coincides with findings of infection by *L. (V) braziliensis* (CAMARGO-NEVES et al., 2002). In contrast, *L. flaviscutellata*, the vector species of *L. (L.) amazonensis*, has not been found. Thus, it seems reasonably likely that *L. amazonensis* has adapted to different vectors, as previously reported for *L. braziliensis* (ODDONE, 2007), where the high number of dogs infected with *Leishmania amazonensis*. The high positivity for *L. amazonensis* seen in the samples from dogs in Bauru may have resulted from interaction between dogs that present the vector and wild animal carriers of *L. amazonensis*. The species *L. amazonensis* has been detected in spider monkeys in Bauru zoo (LIMA et al., 2012).

In the past, São Paulo State had many cases of cutaneous leishmaniasis (VALE & FURTADO, 2005). More recently, there was an outbreak of visceral leishmaniasis, so it is reasonable to assume that some of the dogs involved actually had cutaneous disease, as observed in our results. Little attention has been paid to species determination in the area studied, such that previously, only *Leishmania* spp. was detected (TRONCARELLI et al., 2009).

Similarly to our study, cutaneous infection has also been reported in Araçatuba, SP, Brazil, an area that is endemic for VL in peridomestic areas. The resulting infection presented symptoms similar to VL caused by *L. infantum*. Among the 13 samples that were found to be positive through PCR in the study conducted in Araçatuba, the causative agent in two cases of CL was identified as *L. amazonensis*, even though these presented clinical signs of VL (TOLEZANO et al., 2007).

**Conclusion**

Our data enabled identification of the species *L. amazonensis* and *L. infantum* in dogs in Bauru, SP, Brazil. These findings indicate that the diagnoses of cases of canine leishmaniasis need to be reexamined, since the causative agent identified is not restricted to *L. infantum*, which commonly causes visceral leishmaniasis, but may also include *L. amazonensis*, the *Leishmania* species that can cause the cutaneous form. In endemic areas, more than one etiological agent may be circulating, and therefore there is a need to identify the parasite species, so as to guide leishmaniasis control programs.

**Conflict of Interest Statement**

None of the authors has any financial or personal relationships that could have inappropriately influenced or biased the content of the paper.

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Characterization of *Leishmania* species in the peripheral blood of dogs by PCR-RFLP


