Parasitological analysis of synovial fluid from dogs naturally infected with *Leishmania* sp.

Análise parasitológica do líquido sinovial de cães naturalmente infectados por *Leishmania* sp.

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ABSTRACT

The aim of this study was to verify the feasibility of synovial fluid (SF) analysis for the parasitological diagnosis of leishmaniasis in naturally infected dogs. Aseptic arthrocentesis of the tarsal and carpal joints of 40 infected dogs (158 *Leishmania* sp samples) was conducted after euthanasia. The SF smears were evaluated under optic microscopy and amastigotes were detected in the SF of 35 dogs (87.5%). Of the 93 positive samples, 53% were obtained from the tarsal joints, and 47% were obtained from the carpal joints. Amastigote forms were confirmed in one (17%), two (26%), three (31%) or all four investigated joints (26%). This research highlights the feasibility and high sensitivity of SF evaluation for the parasitological diagnosis of leishmaniasis in symptomatic dogs.

Key words: leishmaniasis, canine, diagnosis, joint.

Resumo

O objetivo deste estudo foi verificar a viabilidade da análise do líquido sinovial (LS) para o diagnóstico parasitológico de leishmaniose em cães naturalmente infectados. Artrocenteses assépticas das articulações do tarso e do carpo de 40 cães infectados (totalizando 158 amostras articulares) foram realizadas após a eutanásia. Os esfregaços LS foram avaliados sob microscopia óptica e formas amastigotas de *Leishmania* sp. foram detectadas no LS de 35 cães (87,5%). 93 amostras articulares foram positivas, sendo 53% obtidas a partir das articulações tarsais e 47% obtidas de articulações carpais. Formas amastigotas foram confirmadas em uma (17%), duas (26%), três (31%) ou todas as quatro articulações investigadas (26%). Este trabalho destaca a viabilidade e a alta sensibilidade da avaliação LS para o diagnóstico parasitológico de leishmaniose em cães sintomáticos.

Palavras-chave: leishmaniose, canina, diagnóstico, articulação.

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Leishmaniasis is an anthropozoonosis caused by *Leishmania*. A total of 98 countries on 5 continents reported endemic leishmaniasis transmission; official case counts more than 278,000 new cases per year, which are underestimated in one-third of countries which do not have reported data for the last five years (ALV AR et al., 2012). In veterinary medicine, leishmaniasis caused by *L. infantum* is mostly important in dogs, which are considered the main reservoir of this parasite for humans (SOLANO-GALLEGO et al., 2009). This agent is the most important species in the Old World and is synonymous to *L. chagasi* in Central and South America (SOLANO-GALLEGO et al., 2009).

Serological examinations, such as Enzyme-Linked Immunosorbent Assays (ELISAs) and Indirect Fluorescent Antibody Tests (IFATs) are recommended by the World Organization for Animal Health (OIE, 2008) for the diagnosis of leishmaniasis in dogs. However, serological studies, such as IFATs may lead to false-positive results due to cross reactions with other etiological agents like *Trypanosoma cruzi*, *Plasmodium* sp. or *Mycobacterium* sp. Nevertheless, serological tests are more expensive than parasitological diagnosis (GONTIJO & MELO, 2004). Parasitological analysis, on the other hand, has a specificity of 100% and is cheaper but shows variable sensitivity (SILVA, 2009). Samples of bone marrow, lymph node, spleen and skin lesions are currently used for the parasitological diagnosis of leishmaniasis in naturally infected dogs, however...
the use of synovial fluid (SF) for this propose has been rarely reported (SARIDOMICHELAKIS et al., 2005; SILVA, 2009). Although arthritis can be diagnosed in dogs with leishmaniasis, it can occur due to immune complex deposition or to the presence of the *Leishmania* in the joint. Only few case reports have already referred the identification of amastigotes on canine SF (SANTOS et al., 2006; SILVA, 2007). The present study aimed to evaluate the feasibility of SF analysis for the parasitological diagnosis of *Leishmania* sp. infection in naturally infected dogs.

Forty symptomatic dogs, 20 females and 20 males, were included in this study. The dogs were brought to the Veterinary Hospital of the Federal University of Pampa for euthanasia due to leishmaniasis, according to Brazilian Ministry of Health recommendations. The dogs presented signs like cutaneous lesions, loss of weight, enlarged lymphnodes and dehydration. The infection was confirmed by positivity in both serological tests: ELISA and the IFAT, with the cut-off titer of 1:40 as confirmatory, according to brazilian public health authorities (DANTAS-TORRES et al., 2012). The dogs underwent to general anesthesia with Propofol (Propovan®) (8mg kg⁻¹ IV) and euthanasia immediately after the induction, with KCl 19.1% (40mL dog⁻¹, IV).

The samples of SF were obtained from both carpal and tarsal joints of each animal, immediately after death, by aseptic arthrocentesis, as described by CLEMENTS (2006), summing up 160 arthrocentesis. The maximum amount of SF was collected using a fine needle (25x7mm) and a 3ml syringe containing 50μL of ethylenediamine tetraacetic acid (EDTA) in order to prevent clot formation.

The smears were prepared on glass slides with 10μL, without centrifugation, immediately after arthrocentesis and were stained using the quick panoptic method. Two glass slides were prepared from the SF obtained from each evaluated joint (carpal and tarsal joint). The stained smears were examined under a microscope (1000X) and classified as positive or negative for the presence of *Leishmania* sp.

Positive parasitological results of carpal and tarsal SF samples were compared using the Standard Frequentist Hypothesis Testing for two proportions (Minitab Statistical Software) (95% confidence).

The SF-based parasitological examination was positive for *Leishmania* sp. in 35 of the 40 examined dogs (87.5%). Amastigote forms were observed in macrophages and neutrophils (Figure 1) or free in the synovia.

From the 160 joints submitted to arthrocentesis, 158 samples of SF (98.75%) were collected without complications and analyzed. Two tarsal joints from the same dog could not be evaluated due to insufficient material. Of the 158 evaluated samples, 93 were classified as positive, and of these, 49 (53%) were obtained from the tarsal joints and 44 (47%) from the carpal joints. However no significant differences were observed in the frequency distribution between the two different joint categories. Amastigotes were observed in only one joint for 17% of the dogs, in two joints for 26%, in three joints for 31% and in all four evaluated joints for 26%.

Parasitological examination for the diagnosis of leishmaniasis is commonly performed using bone marrow, lymph nodes and spleen aspirates,

![Figure 1 - Parasitological analysis of synovial fluid for the identification of *Leishmania* sp. (quick panoptic method). A macrophage containing several *Leishmania* sp. amastigotes is shown (arrows). Obj.100x (A). A neutrophil containing a few *Leishmania* sp. amastigotes is shown (arrow). Obj.100x (B).](image-url)
the failure to obtain a synovial fluid sample, but it was observed only in two of the 160 joints in the present study. The contamination of samples with blood during arthrocentesis is considered an occasional complication (CLEMENTS, 2006) but it was not observed in the 160 arthrocentesis. When performed properly, complications of arthrocentesis to dogs are rare, but can include hemarthrosis, localized trauma, pain, and iatrogenic infection. In fact, the technique of arthrocentesis is simple to perform for most peripheral appendicular joints in small animals and is associated with minimal morbidity, but remains underused (BERG et al., 2009).

Some complications of arthrocentesis were described on the literature and could interfere on SF analysis. According to CLEMENTS (2006), the most common complication of arthrocentesis is the failure to obtain a synovial fluid sample, but it was observed only in two of the 160 joints in the present study. The contamination of samples with blood during arthrocentesis is considered an occasional complication (CLEMENTS, 2006) but it was not observed in the 160 arthrocentesis. When performed properly, complications of arthrocentesis to dogs are rare, but can include hemarthrosis, localized trauma, pain, and iatrogenic infection. In fact, the technique of arthrocentesis is simple to perform for most peripheral appendicular joints in small animals and is associated with minimal morbidity, but remains underused (BERG et al., 2009).

The sensitivity of parasitological examinations can be affected not only by the origin of the sample but also by the time elapsed between collection, preparation and sample analysis; the time spent on the evaluation; and the number of fields covered (BRUSTOLONI, 2006). This manuscript presents methodological adaptations that probably made the use of SF feasible for the parasitological diagnosis of leishmaniasis, in contrast to the results of other studies (SILVA, 2007; SILVA, 2009). First, EDTA was added to the samples based on the possible presence of joint inflammation and increased levels of fibrinogen, which leads to rapid clot formation in the SF interfering with amastigote visualization (HARRY, 2010). Moreover, the period for slide preparation and analysis was restricted to 24 hours after aspiration, and four joints from each dog were evaluated. This report encourages the use of SF for parasitological leishmaniasis diagnosis in a greater amount of dogs. It should be further tested and compared its sensitivity and complications in dogs with the traditionally aspirates of bone marrow and lymph nodes and for asymptomatic patients.

The parasitological examination of SF for *Leishmania* sp. was feasible and presented positivity of 87.5% for symptomatic naturally infected dogs.

**ETHICS COMMITTEE**

The present study was approved by the Ethics Committee for the Use of Animals (CEUA) of the Federal University of Pampa (protocol no. 029/2012).

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