Amino acid permease 3 (aap3) coding sequence as a target for *Leishmania* identification and diagnosis of leishmaniases using high resolution melting analysis

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**Abstract**

**Background:** The leishmaniases comprise a spectrum of clinical manifestations caused by different species of *Leishmania*. Identification of species is important for diagnosis, treatment and follow-up management. However, there is no gold standard for species identification. High resolution melting analysis (HRM) offers a possibility to differentiate *Leishmania* species without the need for processing of the PCR-product. The amino acid permease 3 (aap3) gene is an exclusive target for trypanosomatids and is conserved among *Leishmania* spp., thus it can be a valuable target for an HRM assay for diagnosis of the leishmaniases.

**Results:** The HRM dissociation profiles of three amplicons targeting the aap3-coding region allowed the discrimination of *L.* (*Leishmania*) *donovani*, *L.* (*L.* infantum, *L.* major, *L.* tropica, *L.* mexicana, *L.* amazonensis, *L.* (*Viannia*) braziliensis, *L.* (*V.* guyanensis, *L.* (*V.*) laissoni, *L.* (*V.*) naiffi and *L.* (*V.*) shawi using DNA from promastigote cultures. The protocol was validated with DNA samples from clinical infection in humans and a cat, naturally infected sand flies, and experimentally infected mice.

**Conclusions:** HRM analysis using the aap3 coding sequence as target is a relatively cheap, fast and robust strategy to detect and discriminate *Leishmania* species from all the endemic regions worldwide. The target and method proved to be useful in clinical, field and experimental samples, thus it could be used as a tool in diagnosis as well as ecological and epidemiological studies.

**Keywords:** PCR, HRM, Infectious diseases, *Leishmania* discrimination

**Background**

The leishmaniases are a group of diseases caused by *Leishmania* spp. Clinical presentations range from self-healing cutaneous lesions to potentially lethal visceral leishmaniasis [1]. It is defined by the World Health Organization (WHO) as a neglected tropical disease, meaning it is underreported, underestimated, underfunded and underprioritized by the pharmaceutical industry and often by public health authorities alike [2]. Over 350 million people are at risk of being infected, and it is estimated that 20,000 to 40,000 die each year of the leishmaniases [3]. In some regions, it is considered as largely an anthropogetic disease, while in others it is a zoonosis. The reservoir of the parasite can be domestic, sylvatic, and in some areas, both. Over 20 different species of *Leishmania* can cause the disease in humans [1, 4]. The species that cause disease in humans are grouped into two subgenera: *Leishmania* (*Leishmania*) and *Leishmania* (*Viannia*), based on biological features of the parasites [5]. Species identification is important for clinical diagnosis.
prognosis, treatment and follow-up management. For example, some *L. (Viannia)* species, such as *L. (V.) braziliensis*, are known to cause cutaneous lesions, which may later reappear as mucocutaneous lesions, thus requiring systemic treatment, while other lesions infected by strains known to only cause cutaneous manifestations may be treated with local treatment or observation therapy [6].

The leishmaniases can be diagnosed in various ways, with all their strengths and limitations. However, there is no gold standard for diagnosing the diseases. Microscopy is useful in a high-endemic setting, but lacks sensitivity and needs a trained microscopist, not easily found in most non-endemic areas. Diagnostic tests involving serology, such as the direct agglutination test (DAT) and lateral flow immunochromatographic tests can also be useful. Especially lateral flow immunochromatographic tests can give a rapid diagnosis, are simple to use, easy to interpret and relatively cheap - all qualities which are very important in a low-resource setting in many endemic areas. However, serological tests may vary greatly in their sensitivity and specificity between endemic regions [7]. Furthermore, serology is not able to determine the species causing the disease. Diagnosis involving nucleic acid detection is valuable due the high sensitivity and specificity, and for the potential ability to quantify and identify the infecting species. There is a plethora of possible techniques and an equal amount of possible targets [8].

Real-time PCR, followed by high-resolution melting analysis (HRM), generates thermodynamic differences in the dissociation profile of amplicons resulting in specific signatures of polymorphisms due to small differences in nucleotide composition [9]. HRM is rapid, comparatively little laborious and a relatively cheap method where the post-PCR treatment is contained in the tubes with small risk for lab-contamination. HRM has been used for identification of other infectious agents [10–13]. In earlier work, we have already shown that HRM can be a valuable tool for *Leishmania* genotyping, using *hsp70* as a target [14]. AAP3 is an amino acid transporter, which mediates uptake of lysine, histidine, phenylalanine, citrulline and arginine, with the highest affinity for the last one [15–20]. AAP3 is involved in the polyamine pathway, essential for parasite replication [17, 21–23]. The coding sequence for AAP3 is conserved among *Leishmania* spp., indicating its value as the chosen target [17]. We already demonstrated *aap3* as an attractive target for detecting *Leishmania*, by a real-time PCR method, but this approach did not discriminate the species [24]. In this paper, we describe the use of the *aap3* coding sequence as target for differentiation of *Leishmania* spp., using HRM analysis. The *aap3*-HRM method showed to be a specific and sensitive tool to differentiate *Leishmania* spp., using reference strain cultures and validated using clinical samples, naturally infected sand flies and experimentally infected mice samples.

### Methods

#### Organisms

Promastigotes of *Leishmania* spp. (see Table 1) were grown at 25 °C in M199 medium containing Earle’s salts, supplemented with 10% fetal bovine serum, 40 mM HEPES (pH 7.4), 100 μM adenine, 5 mg/l hemin, 0.05 mg/ml streptomycin and 4550 U/ml penicillin. *Trypanosoma cruzi* (Y-strain), *Crithidia fasciculata* (TCC-039) and *Endotrypanum schau diinni* (MCHO/BR/80/M6159) were grown at 28 °C in liver-tryptose medium supplemented with 10% fetal bovine serum and 0.05 mg/ml streptomycin and 4550 U/ml penicillin. *Trypanosoma brucei* (Lister 427) was grown at 28 °C in SDM-79 medium (LGC Biotecnologia, Cotia, São Paulo, Brazil), supplemented with 10% fetal bovine serum, 0.05 mg/ml streptomycin and 4550 U/ml penicillin. Mammalian DNA from BALB/c mouse and Wistar rat were obtained from the DNA repository of the Laboratory of Trypanosomatidae at Physiology - IB-USP and were used as negative controls (see Additional file 1: Figure S1 and Additional file 2: Figure S2).

#### Naturally and experimentally infected samples

To validate the standardized protocols, samples previously identified by other diagnosis tests for the leishmaniases were used as a template in *aap3*-HRM assays [14, 26]. Human paraffin-embedded samples (from patients from Hospital das Clínicas da Universidade de São Paulo or Hospital da Irmandade da Santa Casa de Misericórdia de São Paulo), a sample from an infected cat (from Instituto de Medicina Tropical de São Paulo - USP), naturally infected sand flies (from Superintendência de Controle de Endemias de São Paulo), and samples from experimentally infected mice (from Instituto de Biociências - USP) were included.

#### DNA extraction

The *Leishmania* strains used as assay-standards, *T. cruzi, T. brucei, C. fasciculata, E. schaudinni* and the mammalian DNA was isolated using a modified salting out technique, as previously described [14]. The DNA from the additional strains used for specificity studies was isolated using the QIAamp DNA Blood Mini Kit (Qiagen, Hilden, Germany), according to the manufacturer’s instructions. DNA from mouse was isolated from whole blood and DNA from rat was isolated from liver-tissue, both by using the DNaseasy Blood and Tissue Kit (Qiagen), according to the manufacturer’s instructions. DNA quality and concentration were determined by NanoDrop ND-1000 Spectrophotometer (Thermo Scientific, USA). All DNA was stored at -20 °C until further use.

For the paraffin-embedded tissues, a first step of washing was performed with xylol heated to 95 °C to remove paraffin, followed by repetitive washes of absolute ethanol.
Table 1 *Leishmania* reference strains and additional strains used in this study with information about international code number, clinical form and host according to the World Health Organization classification [25]

| Strain                  | International code number | Clinical form | Isolated from | WHO reference strain |
|-------------------------|----------------------------|---------------|---------------|----------------------|
| Assay standard strains  |                            |               |               |                      |
| *L.* (*L.*) donovani    | MHOM/IN/80/DD8              | Visceral      | *Homo sapiens*| Yes                  |
| *L.* (*L.*) infantum    | MCER/BR/1981/M6445          | Visceral      | Cerdocyon thous | No                  |
| *L.* (*L.*) tropica     | MHOM/SU/60/OD               | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*L.*) major       | MHOM/L/81/Friedlin          | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*L.*) amazonensis| MHOM/BR/1973/M2269          | Cutaneous     | *Homo sapiens*| Yes                  |
| *L.* (*L.*) mexicana    | MNYC/BZ/62/M379             | Cutaneous     | *Nyctomys sumichrasti* | Yes |
| *L.* (*V.*) amazonensis| MHOM/BR/81/M6426            | Cutaneous     | *Homo sapiens*| Yes                  |
| *L.* (*V.*) braziliensis| MHOM/BR/1975/M2903          | Cutaneous     | *Homo sapiens*| Yes                  |
| *L.* (*V.*) guyanensis  | MHOM/BR/1975/M4147          | Cutaneous     | *Homo sapiens*| Yes                  |
| *L.* (*V.*) naiffi      | MDAS/BR/1979/M5533          | na            | *Dasypus novemcinctus* | Yes |
| *L.* (*V.*) shawi       | MCEB/BR/84/M8408            | Cutaneous     | *Cebus apella* | Yes                  |
| Additional strains used for specificity studies |                               |               |               |                      |
| *L.* (*L.*) donovani    | MHOM/CY/2006/CH33           | Visceral      | *Homo sapiens*| No                  |
| *L.* (*L.*) tropica     | MHOM/SU/74/K27              | Cutaneous     | *Homo sapiens*| Yes                  |
| *L.* (*L.*) major       | MHOM/MA/2000/INHW10         | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*L.*) major       | MHOM/AF/2006/LEM5344        | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*L.*) major       | MHOM/MA/2004/LEM4905        | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*L.*) major       | MHOM/TN/2006/LPN296         | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*L.*) major       | MRHO/SU/59/LV39             | na            | *Rhombomyys opimus* | No |
| *L.* (*L.*) mexicana    | MHOM/BZ/82/BEI21            | Cutaneous     | *Homo sapiens*| Yes                  |
| *L.* (*L.*) mexicana    | MHOM/MX/93/CRE47            | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*L.*) mexicana    | MHOM/MX/96/NAN01            | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*V.*) braziliensis| MHOM/BR/75/M2904            | Cutaneous     | *Homo sapiens*| Yes                  |
| *L.* (*V.*) braziliensis| MHOM/BR/87/LTB12MAR87       | Mucocutaneous | *Homo sapiens*| No                  |
| *L.* (*V.*) guyanensis  | MHOM/GF/94/22319            | Cutaneous     | *Homo sapiens*| No                  |
| *L.* (*V.*) naiffi      | MHOM/GF/97/CRE88            | Cutaneous     | *Homo sapiens*| No                  |

Abbreviation: na not applicable

Then DNA was purified by an organic extraction with phenol-chloroform according Uliana et al. [27]. For sand flies, cultured *Leishmania* and mice samples, DNA was purified by silica columns, using DNeasy Blood & Tissue Kit (Qiagen).

### Table 2 Primer sequences used in this study based on the amino acid permease 3 (*aap3*) coding sequence, for amplicon 1 (Amp1), amplicon 2 (Amp2) and amplicon 3 (Amp3)

| Name            | Sequence (5'-3')    | Length (bp) | Orientation | Amplicon size (bp) |
|-----------------|--------------------|-------------|-------------|--------------------|
| AAP3-Amp1-F     | ATCCGCTACGTCTCCGCCATCGG | 23         | Forward     | 123                |
| AAP3-Amp1-R     | CGTGGTGAAATCTCATGTCGC | 23         | Reverse     |                    |
| AAP3-Amp2-F     | GCCGTGCAATAAACACCGAGC | 21         | Forward     | 131                |
| AAP3-Amp2-R     | AAGCGGAAAGATGATTGTGGCACC | 23       | Reverse     |                    |
| AAP3-Amp3-F     | GCGGTGCCTCACATACAGCG | 20         | Forward     | 140                |
| AAP3-Amp3-R     | CGGGCACCATGAACACCGGCATA | 24      | Reverse     |                    |

**Primer design**

The primers (listed in the Table 2) were designed based on the following sequences: *L.* (*L.*) *amazonensis* (HQ912026.1 and HQ912027.1), *L.* (*V.*) *braziliensis* (XM_001567050.2), *L.* (*L.*) *major* (XM_001685021.1), *L.* (*L.*) *donovani* (XM_001685021.1).
DNA from 25 ng to 100 fg was tested, with or without the strains). Overlapped melting profile (see Table 1 for information about the strains).

For sensitivity assays, we tested the performance of the amplicons with standard strains which produced an overlapped melting profile (see Table 1 for information about the strains). For amplicon 1, we tested L. (L.) donovani, L. (L.) mexicana, L. (V.) braziliensis and L. (V.) guyanensis. For amplicon 2, we tested L. (L.) donovani, L. (L.) mexicana, L. (L.) infantum and L. (L.) tropica. For amplicon 3, we tested L. (L.) donovani, L. (L.) mexicana, L. (V.) braziliensis and L. (V.) guyanensis. For all amplicons, a range of parasite DNA from 25 ng to 100 fg was tested, with or without 25 ng/μl of human DNA. The DNA from standard strains was purified from in vitro cultivated parasites, as described above. Efficiency calculations were made for each amplicon from template ranging from 25 ng to 50 pg (see Additional file 4: Figure S3).

Statistics
All samples were tested in duplicate in at least three independent experiments. One-way ANOVA was used to calculate statistical difference between the Tm’s of paired species for each amplicon. The results are presented as mean differences with 95% confidence intervals, and statistical significance set to P < 0.05. The results were analyzed and graphs were produced using GraphPad Prism version 7 (Additional file 5).

Results
Specificity
Three amplicons were designed based on available sequences from GenBank or TriTryp database [28], considering that aap3 is present in two copies situated in tandem around 4 kb apart in most of Leishmania species [16, 47]. As expected, the BLAST search did not reveal any other hits than for Leishmania spp. As shown in Additional file 1: Figure S1, designed primers generated amplicons of the same size for all the species, presenting differences in nucleotide composition. To predict and to delimit potentially informative polymorphic regions, theoretical melting temperatures were calculated in silico using the OligoCalc tool [29] (data not shown), to guide primer design. In addition, the two copies of the gene were aligned concomitantly for the choice of primers that amplify both copies (data not shown). Although some single nucleotide polymorphism (SNP’s) were detected between the two copies of the gene, it did not alter the Tm values. The alignments also allowed the selection of conserved regions for the design of oligonucleotides common to all species, which were used to generate products containing differences in nucleotide composition that were able to distinguish species or groups of species. Furthermore, we analyzed the amplification profile by conventional PCR using the same primers (Additional file 1: Figure S1). We observed specific products of the same size for all species analyzed for each set of primers: 123, 131 and 140 bp for amplicon 1, amplicon 2 and amplicon 3, respectively. No formation of dimers of primers were detected. Control samples were used and no amplification was observed for T. brucei, T. cruzi, E. schaudinni, rat, mouse and human DNA (Additional file 1 and Additional file 2). The analysis of amplicon 3 showed an amplification of C. fasciculata, a non-pathogenic and closely related organism to Leishmania spp., but the Cq and Tm values were able to distinguish C. fasciculata from all Leishmania spp. tested (Additional file 1: Figure S1 and
Individual Tm values and normalized melting profiles of 3 amplicons for reference strains are presented in Fig. 1, indicating that these differential profiles distinguished among *Leishmania* spp. To reliably distinguish different species, we only considered differences in Tm values exceeding ± 0.25 °C, an interval statistically determined as discriminatory (Additional file 5: Figure S3).

According to Tm and melting curves, amplicon 1 was able to distinguish the species into 3 clusters for subgenus *L.* (*Leishmania*): the visceral *L.* (*L.*) *donovani* and *L.* (*L.*) *infantum*, cutaneous strains from Eurasia and Africa [*L.* (*L.*) *major* and *L.* (*L.*) *tropica*], and cutaneous and mucocutaneous strains from the Americas *L.* (*L.*) *amazonensis*, *L.* (*L.*) *mexicana*, and strains of subgenus *L.* (*Viannia*). Furthermore, it was also able to distinguish

![Fig. 1 Melting temperatures and normalized melting profiles obtained with the HRM assays.](image)

**Fig. 1** Melting temperatures and normalized melting profiles obtained with the HRM assays. **a** Melting temperatures (Tm) for all amplicons and standard species. For all amplicons 25 ng of genomic DNA was used as template. The plots show the mean, minimum and maximum of the Tm-values. Each species was tested in duplicate in three independent experiments. **b** Normalized HRM plots for all amplicons: (i), (ii) and (iii) represent the normalized melting profile for amplicon 1, amplicon 2 and amplicon 3, respectively. Each sample was tested in duplicate in three independent experiments. For (i), the bundles circled in the blue line, respectively from the left to the right, represent the profiles for *L.* (*L.*) *tropica*/*L.* (*L.*) *major*, *L.* (*L.*) *donovani*/*L.* (*L.*) *infantum*, *L.* (*L.*) *amazonensis*/*L.* (*L.*) *mexicana*, *L.* (*V.*) *brazilensis*, *L.* (*V.*) *naiffi*, *L.* (*V.*) *guyanensis*/*L.* (*V.*) *shawi* and *L.* (*V.*) *lainsoni*. For (ii), the bundles circled in the blue line, respectively from the left to the right, represent the profiles for *L.* (*L.*) *amazonensis*, *L.* (*L.*) *mexicana*, *L.* (*L.*) *infantum*, *L.* (*L.*) *donovani*, *L.* (*L.*) *major* and *L.* (*L.*) *tropica*. For (iii), bundles circled in the blue line, respectively from the left to the right, represent the profiles for *L.* (*L.*) *infantum*/*L.* (*L.*) *major*, *L.* (*L.*) *tropica*/*L.* (*V.*) *lainsoni*/*L.* (*V.*) *naiffi*, *L.* (*L.*) *donovani*/*L.* (*V.*) *brazilensis*/*L.* (*V.*) *guyanensis*/*L.* (*V.*) *shawi*, *L.* (*L.*) *amazonensis*, *L.* (*L.*) *mexicana*. Abbreviation: RFU: relative fluorescence units.
some species of the subgenus *L.* (*Viannia*) into 3 clusters: *L.* (*V.*) *lainsoni, L.* (*V.*) *brazilensis* from the other species in that subgenus (Fig. 1).

Amplicon 2 was especially designed for strains of the subgenus *L.* (*Leishmania*). No amplification for subgenus *L.* (*Viannia*) was observed (Additional file 1: Figure S1). It was able to further distinguish the visceral strains into two different clusters: *L.* (*L.*) *donovani* and *L.* (*L.*) *infantum*. For the cutaneous species, it was able to distinguish the Eurasian and African cutaneous strains *L.* (*L.*) *major* and *L.* (*L.*) *tropica*. However, no difference in melting temperature was detected for the American cutaneous strains *L.* (*L.*) *amazonensis* and *L.* (*L.*) *mexicana* (Fig. 1). Therefore, amplicon 3 was designed for this purpose.

Amplicon 3 was, as predicted, able to further distinguish *L.* (*L.*) *amazonensis* from *L.* (*L.*) *mexicana*. Furthermore, it was able to distinguish *L.* (*L.*) *donovani* from *L.* (*L.*) *infantum*, thereby distinguishing the two species commonly causing VL. It was also able to distinguish the more common causes of CL in Eurasia and Africa: *L.* (*L.*) *major* and *L.* (*L.*) *tropica*.

Differences in amplification efficiencies could be observed when the amplification curves from real-time PCR were analyzed. Cq (quantification cycle) data were used as a relative parameter of quantification for the 3 targets when normalized amounts of samples were compared (see Additional file 2: Figure S2). Using 25 ng as template, for all *Leishmania* species, the amplification reactions produced curves with similar Cq values for amplicons 1 and 3, indicating a good specificity for all species. For amplicon 2, no amplification of DNA from subgenus *L.* (*Viannia*) was observed. In addition, high Cq-values for *L.* (*L.*) *amazonensis, L.* (*L.*) *major* and *L.* (*L.*) *mexicana* indicate a lower efficiency compared with the other *L.* (*Leishmania*) species, which also could be used as a parameter of discrimination. For amplicon 1, there was an unspecific amplification of human DNA. However, both the Cq and Tm values were distinct.

For all amplicons, additional strains were tested for specificity for all species, except *L.* (*L.*) *infantum, L.* (*L.*) *amazonensis, L.* (*V.*) *lainsoni* and *L.* (*V.*) *shawi*, as we did not have additional strains for these species available. For amplicon 1, we found that most strains, except *L.* (*L.*) *major* and *L.* (*V.*) *naiffi* showed a consistent profile for their melting temperatures. For amplicon 2 a relatively large SD was found for *L.* (*L.*) *donovani* (Table 3).

### Sensitivity

To evaluate the role of DNA-concentration, we tested the performance of the three amplicons in strains that presented close melting temperatures, based on the specificity assays. The limit of detection (LOD) was estimated using ten-times serial dilutions from 25 ng to 100 fg of DNA from standard strains purified from *in vitro* cultured parasites. For amplicon 1 the LOD was 100 fg for all species tested except *L.* (*L.*) *mexicana*, which had a LOD of 250 fg. For amplicon 2 the LOD was 100 fg for all species tested except for *L.* (*L.*) *mexicana*, which had a LOD of 50 pg. For amplicon 3, the LOD was lower for all species, with 500 fg for *L.* (*L.*) *donovani, L.* (*L.*) *mexicana* and *L.* (*L.*) *guyanensis*, while for *L.* (*V.*) *brazilensis* the LOD was 50 pg. The LOD was the same for all amplicons and all species in mixtures of *Leishmania* and human DNA (data not shown). Considering that the *Leishmania* genome sizes range between 29 Mb and 33 Mb, varying from 34 to 36 chromosomes [30], the estimated single-cell DNA is approximately 75 fg. Thus, we can assume that 100 fg is equivalent to just above one parasite in most of the *Leishmania* species.

We further tested if the initial amount of DNA would cause a variation in the melting temperature by serially diluting DNA from 25 ng to 100 fg with or without 25 ng/μl of human DNA. For all three amplicons, the results were the same with or without human DNA. There was a slight variation in melting temperature for all the species, only exceeding 0.2 °C for *L.* (*V.*) *guyanensis* in amplicon 1 with human DNA (Table 4). Furthermore, for amplicon 1 the difference in melting temperature between *L.* (*L.*) *mexicana* and *L.* (*V.*) *brazilensis* could potentially cause a confusion between these two species if melting temperatures were considered for this amplicon, the same can be noticed for *L.* (*L.*) *donovani, L.* (*L.*) *brazilensis* and *L.* (*L.*) *guyanensis* for amplicon 3 (Fig. 2).

### Efficiency

The reaction efficiency for each amplicon was performed using 25 ng, 5 ng and 50 pg of DNA from each reference *Leishmania* species. Efficiency curves, slopes and $R^2$...
values were calculated (see Additional file 4: Figure S3). The template points produced Cq values between 20 and 30, where the PCR reactions produced copies of template at exponential ratios and with 95–105% of efficiency. The efficiency patterns were similar for all species tests for each amplicon, except for L. (L.) mexicana in amplicon 2. The lower efficiency for L. (L.) mexicana for amplicon 2 can be explained by the presence of mismatches comparing in primers (see Additional file 3), as can also be observed in the Cq values showed in Additional file 2: Figure S2.

Validation using naturally and experimentally infected samples
To validate the aap3-HRM protocol with other samples than reference strain cultures, we applied this protocol in seven biological samples from naturally infected humans, naturally infected cat, naturally infected sand flies, and experimentally infected BALB/c mice (Table 5). The results of amplicon 1 showed a correlation with those obtained with other diagnostic targets; small subunit ribosomal DNA (SSU rDNA) [26], glucose-6-phosphate dehydrogenase (g6pd) PCR [31] and heat-shock protein 70 (hsp70) HRM [14]. Although amplicons 2 and 3 were not able to amplify all samples, the positive samples correlated with results obtained with other targets.

Discussion
Diagnosing Leishmania infection at the species level is important, as it may guide treatment options and follow-up protocols. Accurate diagnosis is also important in an ecological and epidemiological sense. Unfortunately, in most endemic regions, the leishmaniases are underreported, and the true burden of the disease remains unknown. Techniques that target genomic or mitochondrial DNA by PCR or related techniques are today the most common for detection and identification of Leishmania spp. However, there is no gold standard in methods and targets [8]. Common targets such as kinetoplast DNA (kDNA) and SSU rDNA gene have been used for parasite detection. On the other hand, the g6pd coding region, internal transcribed spacer 1 (ITS1) rRNA, and hsp70 coding region have been more commonly used to identify the parasite at the species level [14, 32–38]. Conventional PCR and real-time PCR often followed by sequencing is increasingly being used for detection and identification of Leishmania spp. [6, 39]. Both methods are relatively laborious and time-consuming and they require PCR product manipulation, increasing the risk of laboratory contamination. Furthermore, the interpretation of the results requires also considerable skill and experience.

HRM is a relatively new real-time PCR coupled technique, with the first papers appearing in 2003 [40, 41]. The technique identifies nucleotide composition polymorphisms

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**Table 4** Effect of initial amount of DNA on melting temperature. Mean and standard deviation (SD) for serial dilutions of DNA from standard strains. DNA concentration ranged from 25 ng to 100 fg. Human DNA concentration was kept constant at 25 ng/μl.

| Strain             | Without human DNA | With human DNA |
|--------------------|-------------------|----------------|
|                    | Mean Tm ± SD (°C) | Mean Tm ± SD (°C) |
| Amplicon 1         |                   |                 |
| L. (L.) donovani   | 82.52 ± 0.10      | 82.64 ± 0.18    |
| L. (L.) mexicana   | 83.16 ± 0.10      | 83.23 ± 0.13    |
| L. (V.) braziliensis | 83.31 ± 0.09     | 83.43 ± 0.11    |
| L. (V.) guyanensis | 83.83 ± 0.08      | 83.96 ± 0.34    |
| Amplicon 2         |                   |                 |
| L. (L.) donovani   | 85.44 ± 0.14      | 85.44 ± 0.14    |
| L. (L.) mexicana   | 84.40 ± 0.08      | 84.56 ± 0.07    |
| L. (L.) infantum   | 84.92 ± 0.11      | 85.17 ± 0.14    |
| L. (L.) tropica    | 86.33 ± 0.13      | 86.73 ± 0.18    |
| Amplicon 3         |                   |                 |
| L. (L.) donovani   | 86.17 ± 0.12      | 86.15 ± 0.12    |
| L. (L.) mexicana   | 86.77 ± 0.16      | 86.91 ± 0.16    |
| L. (V.) braziliensis | 86.00 ± 0.14     | 86.17 ± 0.15    |
| L. (V.) guyanensis | 85.97 ± 0.13      | 86.36 ± 0.13    |

**Fig. 2** Effect of DNA-concentration on melting temperature (Tm °C). Dispersion graph of Tm values for amplicons 1, 2 and 3 from a range of 25 ng to 100 fg of DNA template. The plot shows mean, minimum and maximum Tm-values.
in real-time PCR products. The HRM methodology presents several attractive features: the whole process is performed in a closed-tube system (avoiding contamination in the laboratory) and it is relatively fast and cheap. The analysis may also be automated. The melting temperature generated depends on a range of factors, where GC-content, sequence, and length of the sequence are central. The melting curves can be objectively differentiated from each other by differences in melting temperature and shape [9].

For Leishmania, there are few studies utilizing HRM for detection and species identification. One of the earliest reports on the usefulness of HRM in differentiation of Leishmania species is by Nicolas et al. [42] who utilized polymorphisms in the coding sequence for minicircle kDNA to differentiate Eurasian and African species, L. (L.) major, L. (L.) donovani and L. (L.) tropica, and L. (L.) infantum. Later, Talmi-Frank et al. [43] utilized the ITS1 rRNA region to identify, distinguish and quantify Eurasian and African species, L. (L.) infantum/L. (L.) donovani, L. (L.) aethiopica, L. (L.) tropica and L. (L.) major. Both Nicolas et al. [42] and Talmi-Frank et al. [43] only targeted Eurasian and African species, making the approaches valuable in these endemic areas, although of limited value in other endemic areas and in a non-endemic settings where species from all endemic regions could be expected. Pita-Pereira et al. [33] also utilized the minicircle kDNA to discriminate among strains from the subgenus L. (Viannia) and L. (L.) infantum and L. (L.) amazonensis, making it an attractive methodology in an American setting to differentiate strains commonly causing cutaneous, mucocutaneous and visceral manifestations. Ceccarelli et al. [44] also demonstrated an ability to differentiate species of the subgenus L. (Leishmania) from L. (Viannia). HRM has also been used for detection of Leishmania in sand flies, where Aghaei et al. [45] used ITS1 to identify L. (L.) tropica in sand flies. Kuang et al. [46] utilized the lacZ gene to differentiate five Eurasian and African species from one American species, L. (V.) braziliensis. Although they showed its usefulness in clinical samples, the relatively few Leishmania species tested made it difficult to conclude if this target could be useful in other endemic areas. With increasing travel to several endemic regions, the correct diagnosis and species identification for the leishmaniases is of paramount interest. Hernandez et al. [36] utilized the ITS1 and hsp70 to differentiate L. (L.) mexicana, L. (L.) infantum, L. (L.) amazonensis, L. (V.) panamensis, L. (V.) guyanensis and L. (V.) braziliensis. Although they found some ambiguities in species identification between the targets, they showed that ITS1 and hsp70 had potential as diagnostic targets utilizing HRM. Zampieri et al. [14] showed, in a recent paper, that HRM targeting several polymorphic sites on the hsp70 coding region could successfully be used to differentiate several Eurasian, African and American species. This makes it the most attractive target reported so far, especially useful in a non-endemic setting where the patient could have traveled to several endemic areas.

In this work, we describe a method for rapid detection and discrimination of most Leishmania species. We used the aap3 coding sequence as target. Leishmania aap3 coding sequences available in the GenBank and TriTryp databases were aligned to search for relatively conserved regions but present polymorphisms that enable the identification of different species. For this, the available coding sequences of some species from the subgenus L. (Leishmania) and L. (Viannia) were analyzed in silico. The coding sequence for aap3 is present in two copies and organized in tandem in most of the Leishmania spp. genomes. For L. (L.) donovani and L. (L.) amazonensis a 98% identity has been described between the copies of coding regions and a 93% identity between these two

### Table 5 Identification of Leishmania in naturally and experimentally infected samples by HRM analysis targeting the aap3 gene

| Sample source | Amplicon 1 Identification | Amplicon 2 Identification | Amplicon 3 Identification | Previous diagnosis |
|---------------|---------------------------|---------------------------|---------------------------|--------------------|
| Human²        | L. (L.) amazonensis       | negative                  | negative                  | SSU rDNA sequencing |
| Human²        | L. (L.) infantum          | L. (L.) infantum          | L. (L.) infantum          | SSU rDNA sequencing |
| Cat³          | L. (L.) infantum          | L. (L.) infantum          | L. (L.) infantum          | hsp70 HRM          |
| Mouse¹        | L. (L.) amazonensis       | L. (L.) amazonensis       | L. (L.) amazonensis       | SSU rDNA sequencing |
| Mouse¹        | L. (V) braziliensis       | negative                  | L. (V) braziliensis       | g6pd PCR           |
| Sand flies⁴   | L. (L.) infantum          | L. (L.) infantum          | L. (L.) infantum          | SSU rDNA sequencing |
| Sand flies⁴   | Subgenus Viannia          | negative                  | Subgenus Viannia          | g6pd PCR           |

Note: The aap3 amplicons 1, 2 and 3 of DNA from each sample was submitted to HRM analysis. The result was compared with previous identification performed by SSU rDNA-sequencing [26], hsp70 HRM [14] or g6pd PCR [31].

²Human paraffin-embedded tissue from Hospital das Clínicas de São Paulo
³Isolated parasites from cat
⁴Experimentally infected BALB/c mice
⁵Naturally infected Lutzomyia (Lutzomyia) longipalpis
⁶Naturally infected Lu. (Nyssomyia) whitmani

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species [16, 47]. In addition, aap3 appeared conserved among other Leishmania species [17]. Although some species present only one aap3 gene copy, for example L. (V.) braziliensis, this could be due to the misannotation in the genome database. Considering these observations, the polymorphisms found between the two copies of the coding region did not affect the Tm analysis.

Amplicon 1 was able to discriminate the two strains causing visceral leishmaniasis from strains causing American cutaneous leishmaniasis. Furthermore, it was able to differentiate the American cutaneous strains and several of the strains of the subgenus L. (Viannia). The inability to differentiate the two visceral strains L. (L.) donovani from L. (L.) infantum was compensated with these strains having distinct Tm's in amplicon 2 and amplicon 3. The same situation is true for the Eurasian cutaneous species: L. (L.) major and L. (L.) tropica were indistinguishable in amplicon 1, but had distinct Tm's in amplicons 2 and 3. Leishmania (L.) amazonensis and L. (L.) mexicana are phylogenetically closely related species [48], and the difficulty in differentiating these species for diagnostic purposes has been described elsewhere [14]. Both amplicon 1 and 2 showed similar profiles for these two species, while amplicon 3 was able to distinguish them. Amplicon 2 was specifically designed for the subgenus L. (Leishmania), and there was no amplification of L. (Viannia) spp. High Cq values, observed for L. (L.) amazonensis, L. (L.) mexicana and L. (L.) major for amplicon 2, could be explained by primer-mismatches. Some caution should be taken when analyzing the result in a diagnostic setting, especially if negative. However, taken together the results from all amplicons can be considered to strengthen the diagnosis.

Diagnosing leishmaniasis relies on patient and travel history, clinical information (symptoms and clinical findings), and results from laboratory tests. Thus, for example, the inability of amplicon 1 to reliably distinguish L. (L.) donovani from L. (L.) infantum, in a patient with suspected visceral leishmaniasis is of little importance in a clinical setting, as the patient would receive the same treatment and follow-up regime regardless. But in an epidemiological setting, it is of importance to distinguish the species from each other to generate reliable data.

In general, multi-copy genes can be expected to yield a higher sensitivity in molecular diagnostics. The assays developed by Talmi-Frank et al. [43], Hernandez et al. [36] and Zampieri et al. [14], all report a limit of detection (LOD) of less than one parasite. Despite that aap3 only comes in two copies we report a relatively good sensitivity with a LOD of 100 fg for amplicon 1 (just above 1 parasite) to 500 fg for amplicon 3 (approximately 5 parasites). This could probably be improved with a pre-amplification step.

Some HRM assays have found little evidence that DNA concentration of the initial template influences the Tm [36, 46]. However, the initial amount of DNA for

![Fig. 3 Proposed strategy for Leishmania species identification. Diagram of a proposed strategy using aap3-HRM for species identification. For VL patients from the Americas, amplicon 1 would suffice, while for VL patients from Eurasia and Africa we suggest the use of either amplicon 2 alone or amplicon 1 and 2. For patients with CL and MCL from the Americas, we suggest the use of both amplicon 1 and 3.](image-url)
this assay influenced the Tm for some of the species. This is in concordance with the findings of Zampieri et al. [14], who also found that initial DNA applied to the assay did affect the Tm for several strains. For the amplifiers investigated in our study, this could lead to a misidentification for *L. (L.) mexicana* and *L. (L.) braziliensis* for amplicon 1. However, it should be noted that we propose the use of the three amplicons in the identification to strengthen the diagnostic validity and avoid species misidentification. A strategy for *Leishmania* species identification in patients with suspected leishmaniases is proposed in Fig. 3.

The validation of the target and the technique with clinical and experimentally infected samples from human, cat, mice and sand flies indicated a good correlation with other diagnostic targets and techniques. Not all amplicons were able to yield a result for all the samples, as could be expected due to differences in specificity and sensitivity of the primers, where amplicon 2 was not produced for any species belonging to the subgenus *L. (Viannia)*. Despite the limited number of samples tested, the data indicate the potential of the *aap3*-HRM method to identify *Leishmania* species.

**Conclusions**

Overall, the *aap3* coding sequence can be a promising target since it is specific and conserved for *Leishmania* spp. The design of the *aap3*-HRM protocol described is a relatively rapid, simple, sensitive and specific method to identify and distinguish several *Leishmania* spp. There is no need for sequencing or gel fractionation to analyze a PCR-product, minimizing the laboratory contamination as all the reactions are performed within a closed tube. The method can be automated, dispensing a trained technician to analyze the results. It also has the potential to quantify parasites present in samples, as it is a real-time PCR technique, able to detect a small number of parasites. In conclusion, the protocol described may offer a relatively low-cost and reliable method for detection and identification of *Leishmania* in biological and clinical samples.
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