A Real-Time ITS1-PCR Based Method in the Diagnosis and Species Identification of Leishmania Parasite from Human and Dog Clinical Samples in Turkey

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Abstract

Human visceral leishmaniasis (VL) caused by L. infantum and cutaneous leishmaniasis (CL) caused by L. tropica and L. infantum have been reported in Turkey. L. infantum is also responsible for canine leishmaniasis (CanL) and it is widely common in the country. The main aim of the present study was to design a real-time PCR method based on the internal transcribed spacer 1 (ITS1) region in the diagnosis of all clinical forms of leishmaniasis in Mediterranean, and to identify the species directly from clinical samples. Totally, 315 clinical specimens, human/canine visceral (blood, bone marrow, lymph node) and cutaneous (lesion aspiration) samples, and 51 Turkish Leishmania isolates typed by isoenzymatic method were included in the study. For optimization, DNA samples of the 34 strains were amplified by conventional ITS1-PCR and then sequenced for designing the primers and probes, allowing the species identification. Following the validation with the isolates, the test was applied on clinical samples and melting temperatures were used for genotyping. A group of PCR products were further sequenced for confirmation and assigning the inter- and intraspecies heterogeneity. The diagnosis of leishmaniasis is successfully achieved by the new real-time PCR method, and the test identified 80.43% of human and canine VL samples as L.infantum and 65.22% as L.tropica; 52.46% of CL samples as L.infantum and 26.90% as L.tropica. In 13.04% of visceral and 20.62% of cutaneous samples, two peaks were observed. However, the higher peak was found to be concordant with the sequencing results in 96.96%, in terms of species identification. The real-time ITS1 PCR assay clearly identified the leishmanial species in 81.58% of all clinical samples. Genotypic variations of Leishmania parasites in Turkey within species and intraspecies were observed, and L. tropica is also found as causative agent of human and canine VL in Turkey.

Introduction

Leishmaniasis is a group of diseases caused by more than 20 species of the protozoan genus Leishmania in 98 countries and regions with 350 million people living at risk. The main forms of human disease are visceral (VL), cutaneous (CL) and mucocutaneous (MCL) leishmaniasis [1,2].

Turkey is of special epidemiological interest because it lies at the crossroad between Asia and Europe and it comprises seven geographical regions with environmental and ecological differences. Human leishmaniasis, both visceral (>40 cases yearly) and cutaneous (>2000 cases yearly) have been reported for centuries in Turkey. Two species of Leishmania are prevalent in Turkey, causing anthroponotic CL (Leishmania tropica), and zoontic VL (Leishmania infantum) [3,4]. There is also a single report about the occurrence of L. major variants in specific regions of the country [5].

Human VL and CanL are endemic throughout Mediterranean, Ege, Marmara and Black Sea Regions of western Turkey while sporadic in other regions with higher infection rates in dog populations than human cases. There were two zoonodemes of L. infantum (MON1 and MON-98) in dog isolates while all human VL isolates were identified as L. infantum MON-1 by isoenzyme analyses [6-8].

Anthroponotic CL caused by L. tropica is highly endemic in the Southeastern Anatolia, East Mediterranean and Ege Regions of Turkey [3,6-9]. In the South Anatolia, L. infantum in addition to L. tropica has also been reported as a causative agent for human CL [6,10,11]. Leishmania major is known to be endemic in the countries bordering Turkey to the south: Syria, Iraq and Iran [6]; Parasites isolated from CL patients in Sanliurfa province located in Southeastern Region were all identified as L. tropica MON304 while in Aydin province located in western part, Ege region, were identified as L. tropica MON303 (87%) and L. tropica MON304 (13%) [unpublished data].

A universal PCR method targeting the internal transcribed spacer 1 (ITS1) region between the SSU and 5.8S rRNA genes
were described for the direct diagnosis of different clinical manifestations of leishmaniasis and parasite identification. This method is applicable where several more than one parasite species are aetiological relevant. It is highly specific and sensitive detecting approximately 0.2 parasites per sample [12]. Most of the medically important Leishmania species are then readily distinguished by DNA sequencing or restriction enzyme analysis of the PCR product. ITS1 PCR restriction fragment length polymorphism (RFLP) are used for direct species identification in patient tissues, blood or other samples without prior parasite culturing, microscopic analysis or other technique [6,12–14].

In endemic areas, the presence of multiple Leishmania species with overlapping clinical features and geographical distribution requires the development of sensitive laboratory tests with Leishmania species identification in order to evaluate the prognosis of human and canine leishmaniasis and to choose appropriate therapies. Species identification will also contribute to better understanding the epidemiology of leishmaniasis in Turkey [15,16]. Species identification of the agents of leishmaniasis in Turkey is crucial, since the country comprises seven geographical regions with environmental and ecological differences. ITS1 PCR RFLP was successfully performed for clinical samples collected from human leishmaniasis and CanL cases from Turkey [6].

The main aim of the present study was to design a real time PCR method based on internal transcribed spacer 1 (ITS1) region in the diagnosis of all clinical forms of leishmaniasis and identifying of parasite directly from clinical samples (human and canine) or Leishmania isolates. We further sequenced the PCR products for confirmation and assigning the inter- and intraspecies heterogeneity.

**Materials and Methods**

**Clinical samples and Leishmania isolates**

The study was carried out in two steps; (a) designing primer/probes which are specific in genus and species level and optimization of a novel ITS1 real time PCR method using Turkish Leishmania strains previously identified by multilocus enzyme electrophoresis (MLEE) technique in Montpellier Reference Center and four international reference strains; (b) validation of the method using isolates and different types of clinical samples obtained from only confirmed human and dog leishmaniasis cases. Four international reference controls, L. infantum/chania (MHOM/XX/1999/LRC-L774), L. donovani (MHOM/IN/1980/DD8), L. tropica (MHOM/IL/1990/LRC-L590 and MHOM/IL/1996/LRC-L691) and L. major (MHOM/IL/2000/LRC-L779) are included.

A total of 51 Turkish Leishmania strains were isolated from 5 VL, 33 CL, 8 CanL cases between 2000 and 2011, and maintained by subcultures in NNN medium. They were identified by the isoenzymatic method previously. Leishmania promastigotes of all isolates were mass cultivated in RPMI+20%FCS medium and centrifuged in 5th day to obtain pellet in order to use DNA extraction. The concentration of isolates was adjusted to 2.5–3×10^9 promastigotes/mL. A total of 315 clinical samples obtained from human CL (n = 223), human VL (n = 40) and CanL (n = 52) originated from 31 different provinces (mostly from İzmir, Aydın, Hatay and Şanlıurfa provinces) of Turkey were included in the study. The samples were collected between April 2007 and May 2010 and they were all confirmed cases who are found positive/seropositive by different parasitological/serological methods in our laboratory (Table 1, Figure 1).

Blood and bone marrow samples were taken from hospitalized VL patients, and sent to our laboratory in tubes containing EDTA as anticoagulant or on slides. Two hundred microliter sample was used for DNA extraction. Tissue aspirates from CL patients were collected into syringes containing 0.5 mL of saline, a slide was prepared for DNA extraction and part of which was inoculated into NNN medium for isolating the parasite. All the prepared slides were washed with PBS and then this solution was transferred into a 1.5 ml eppendorf tube. The procedures were performed based on the steps mentioned on the DNA extraction kit (Roche Applied Science) for all samples. Quality and quantity of extracted DNA was analysed by spectrophotometry.

**Primer/probe design and optimization**

Conventional ITS1-PCR was applied to 30 Turkish Leishmania strains (obtained from 18 CL, 5 VL, 7 CanL cases) and four international reference strains using the primer set (forward – LITSR; reverse - L5.8S) and conditions published by El Tai et al [17,18]. PCR products were sequenced commercially by RelGen (http://www.relgen.com) and compared by multialignment analysis within each other and with other ITS1 sequences published in BLAST using MultAlin program (CLC Main Workbench v.5.6 genetic program). The forward primer (LITSR) used in the initial experiment was kept but reverse primer (ITS1R-TR1: 5’-GAAGCCAAGTGATCCATCGCCGTTT-3’) and the probes (Probe1: CGGTGTATACAAAAATATACGGGCTTTCGGTTT—FL; Probe 2: GCCTAAGGCTGAGCTCCGGTGAGG—PH) were newly designed according to the variable region for detecting L. donovani complex, L. tropica and L. major, using LightCycler Probe Design Software 2.0 program [19].

The detection of Leishmania DNA in clinical samples and differentiation of species

The real time PCR method targeting ITS1 region between the SSU and 5.8S rRNA genes specific for Leishmania was first applied to 51 Turkish and international isolates to determine the melting temperatures (Tm) for each species. Then, the method was performed using clinical samples and it was repeated twice for each batch of samples. One positive and two negative controls were included for each PCR reaction.
ITS1 real-time PCR method was applied using samples containing 20–50 ng of genomic DNA, 400 nM of each primers, 200 nM of each probes, 2 mM of MgCl₂, 1 µl LightCycler FastStart DNA Master Hybridisation probe (Roche Applied Science), and 1.5 µl PCR grade water (Roche Applied Science) to a reaction total volume of 10 µl. PCR amplification was performed as follows: one cycle of 10 minutes at 95 °C, followed by 45 cycles consisting of denaturation at 95 °C for 10 seconds, annealing at 50 °C for 10 seconds, extension at 72 °C for 20 seconds, and melting at 95 °C for 0 second, 50 °C for 10 second, 40 °C for 10 second, 80 °C for 0 second and cooling at 40 °C for 30 seconds. Melting curves were analysed using channel 2 and 3.

Sequencing and phylogenetic analyses
A group of PCR products from 135 samples (121 human; 14 dogs) including 93 clinical specimens and 42 Turkish Leishmania isolates were sequenced commercially by RefGen (http://www.refgen.com) for the confirmation of the results. The clinical samples group was consisted of DNA samples from 111 lesion aspiration samples of CL patients, 4 blood and 6 bone marrow samples of VL patients; 12 lymph node aspiration and 2 blood samples of dogs. Sequence data were analysed using ClustalW2 (http://www.ebi.ac.uk/Tools/msa/clustalw2/) program and distance of molecular relationship in the group of samples detected as *L. tropica* and *L. infantum* were calculated. Phylogram was generated using CLC Main Workbench (v.5.6) genetic program.

Statistical analysis
Univariant and Tukey analyses were performed using SPSS v.13 program in order to compare the melting temperatures in the group of samples detected as *L. tropica* and *L. infantum*. Statistical significance degree was accepted as <0.05.

Ethical aspects
The study was approved by Local Animal Care and Ethics Committee of the School of Medicine and Ege University Medical School Clinical Research Ethical Committee, Izmir, Turkey.

Results
Optimization using clinical *Leishmania* isolates
The melting temperatures were detected as 68°C Tm for *L. donovani* complex; 62°C Tm for *L. tropica* and 53°C Tm for *L. major*. The sequences of variable region which were used for designing probes in Turkish *Leishmania* isolates were shown in figure S1. The identification results of 51 Turkish *Leishmania* isolates were presented with the results of MLEE comparatively in table S1. Result of the isoenzymatic method was used as gold standard. As indicated in the table S1, species identification of 48 isolates well matched with MLEE results, and sensitivity of the method was found to be 94.11%. Out of three isolates which are not fully in agreement with MLEE, two strains (C010 and C056) are found to be *L. tropica* with PCR while they both were *L. infantum* by MLEE. One strain (C078) gave two peaks with the higher related to *L. tropica* (Figure 2) as concordant with the MLEE analysis (*L. tropica* MON312).

Sequencing results of the 42 strains (10 *L. infantum* and 32 *L. tropica*) were used to construct phylogenetic tree. *L. infantum* isolates

### Table 1. The type and number of the samples included in the study.

| Tissue/Isolates | Type of sample | Clinical feature | Total |
|-----------------|----------------|------------------|-------|
| Blood           | Blood with EDTA| VL 8 - CL - CanL 43 | 51    |
| Bone marrow     | Smear          | - 32 -            | 32    |
| Skin            | Aspiration fluid, smear, biopsy | - 223 - | 223 |
| Lymph node      | Aspiration fluid | - 9 9            |       |
| Isolates        | Promastigotes culture | 5 38 8 | 51 |
| Total           |                | 45 261 60 366      |       |

VL: visceral leishmaniasis; CL: cutaneous leishmaniasis; CanL: canine leishmaniasis.

Figure 1. The Turkey map showing the location and the number of samples in province level.

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constituted a single group while strains determined as *L. tropica* showed several different groups (Figure 3).

The partial sequences of *L. tropica* ITS1 region representing MON200, MON303 and MON304 zymodemes were submitted to GenBank (Accession numbers KC686338, KC679052 and KC609747, respectively).

### Identification of *Leishmania* in clinical samples

Totally, 315 clinical samples were analysed and all of them were diagnosed as positive by ITS1 real time PCR method in genus level. In species level, *L. infantum* and *L. tropica* were detected while no *L. major* was found among clinical samples. Genotyping identified 80.43% (74/92) of human and canine visceral leishmaniasis samples as *L. infantum* and 6.52% (6/92) as *L. tropica*; 52.46% (117/223) of cutaneous samples as *L. infantum* and 26.90% (60/223) as *L. tropica*.

Fifty-eight (18.41%) out of 315 clinical samples (12 of visceral and 46 of cutaneous samples) gave two peaks and two melting temperatures were observed. The standard deviation of melting temperatures was found between 0.4512 and 0.5026 in one peak samples while it was between 0.3718 and 1.0838 in two peak samples.

The sequencing was done for 102 samples (3 blood and 1 bone marrow samples of VL patients; 2 blood samples of dogs; 56 lesion aspiration samples of CL patients; 40 strains from CL, VL and CanL cases) giving one peak and 33 samples (1 blood samples of VL patient; 1 dog blood sample; 29 lesion aspiration samples of CL patients and 2 strains from CL cases) giving two peaks. The sequencing results were concordant with the real time ITS1 PCR results in 98.05% (100/102) samples. For the two peaks samples, a 96.96% (32/33) concordance were detected between the higher peak in the real time ITS1 PCR and sequencing results indicating that the higher peak can be decisive for the species identification.

The ITS1 real time PCR results of clinical samples were classified into three groups which were clearly distinguishable as (a) 191 samples diagnosed as *L. infantum* including 29 visceral, 45 dog and 117 cutaneous samples; (b) 66 samples of *L. tropica* including 60 cutaneous and four visceral and two dog samples and (c) 58 samples of two peaks including 46 cutaneous, seven visceral and five dog samples. The working process and the results were summarized in Figure 4. *L. infantum* and *L. tropica* were found to be causative agents of both clinical forms of human leishmaniasis as well as canine leishmaniasis in Turkey.

### Discussion

The various leishmaniases are caused by different species of *Leishmania*, some of which co-exist in the same region like several endemic areas in Turkey; therefore it is crucial to distinguish the species for diagnosis, treatment and epidemiological purposes. Microscopical examination of stained tissue preparations and culture of tissue aspirates for diagnosis, and multilocus enzyme electrophoresis (MLEE) for species identification are accepted as gold standards. However, all the conventional methods applied in the diagnosis of leishmaniasis have medium to low sensitivity and the amplification of DNA by PCR, using different genomic and kDNA targets which is shown to be more sensitive is gradually replacing the traditional methods for the diagnosis of leishmaniasis. Besides, molecular phylogeny studies of the parasite have increasingly suggested new approaches regarding treatment, prognosis of the disease, the distribution of *Leishmania* species in human and animal hosts, as well as in insect vectors for designing appropriate control measures [12, 20–22].

Different PCR/PCR-RFLP based methods targeting kinetoplastid DNA, telomeric sequences, gp63, minieeks, β-tubulin, or ribosomal RNA encoding genes (particularly the internal transcribed spacers, ITS) and recently microsatellites, have been proposed for species identification in *Leishmania* parasites using isolates and clinical samples [23, 24]. The ITS1 of the ribosomal DNA repeat unit (rDNA-ITS1) has previously been exploited for Old World *Leishmania* species discrimination using RFLP [12–14], reverse hybridization assays [25], and sequencing [26]. There are an estimated 20 to 200 identical copies in the *Leishmania* genome, making it a good target for analyzing low parasite quantities [13, 14, 27]. A new technique applying PCR-SSCP (single-strand conformational polymorphism) was published recently by Chargui et al. The authors notified that SSCP provides more resolution especially when PCR product has weak band on agarose gel and is less expensive than RFLP method [24].

In our study, we present a real-time ITS1-PCR method that can diagnose three Old World *Leishmania* species *L. donovani* complex, *L. tropica* and *L. major*, using newly designed probes to diagnose and simultaneously differentiate between Turkish species in clinical samples. The real time ITS1 PCR assay clearly identified the leishmanial species in 81.58% of all clinical samples. *L. infantum* was identified in 80.43% of human and dog visceral samples while *L. tropica* was detected in 6.52%. In five isolates (C41, C56, C087, C39, C40) from 3 VL patient and 2 dogs, the agent was diagnosed as *L. tropica*. This finding was also supported by isoenzyme typing of the strain C87, resulting as *L. tropica* MON-315 in Montpellier leishmaniasis reference center. This is the first report notifying *L. tropica* as a causative agent of human and canine VL in Turkey although, *L. tropica* has been reported to be isolated from human and canine VL cases in different countries [28–34]. Several studies and case reports published from neighbouring country, Iran, showed that *L. tropica* can rarely cause visceral leishmaniasis. Alborzi et al [35] identified only one *L. tropica* among 64 bone marrow/spleen samples while another group also found only one *L. tropica* out of 11 isolates obtained from dogs in Iran [36].

*L. infantum* is reported as a rare causative agent of CL most notably in the Mediterranean Basin countries such as Tunisia, Algeria, Morocco, Spain, Italy, Portugal, Greece and France [2, 37–40]. Although *L. tropica* is the main causative agent of CL in Turkey, *L. infantum* has dominance in the South Anatolia of Turkey, mainly in Hatay and Adana provinces according to

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**Figure 2. The representative figure of two peaks obtained in some samples.**

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identification of *Leishmania* strains by molecular techniques using miniexon gene PCR-RFLP [10,11], kDNA-PCR [41] and ITS1 PCR-RFLP [6] assays. In our study, it is confirmed that both *Leishmania* species can be found in clinical samples from CL patients in this area. *L. infantum* is also isolated from proven vector sand fly species, *P. tobbi* by Svobodova et al. in the region [11].

Figure 3. Phylogenetic tree based on the alignment of the amplified section of the ITS1 region of 42 Turkish *Leishmania* isolates and international reference strains (for *L. tropica* MHOM/IL/1990/LRC-L590 and MHOM/IL/1996/LRC-L691; for *L. major* MHOM/IL/2000/LRC-L779; for *L. infantum/chagasi* MHOM/XX/1999/LRC-L774). REF*: ITS1 sequences for *L. infantum* MHOM/TN/1980/IPT1; for *L. donovani* MHOM/IN/1980/DD8; for *L. major* MHOM/TM/1973/5ASKH were taken from Talmi-Frank et al. [47] and included the tree. (VL: visceral isolates; CL: cutaneous isolates; CanL: dog isolates). doi:10.1371/journal.pntd.0002205.g003
So far only little is known about the population structures of the two *Leishmania* species in Turkey and the correlation with geographical origin, biogeographical parameters, clinical outcome, involved animal reservoir (dogs) and the transmitting sand fly species. Several pioneer studies were performed that used PCR-RFLP and/or sequence analysis of the ITS rDNA, minicircle kinetoplast DNA, miniexon, NAGT [5,6,10,42] mainly for species identification. In Turkey, nine zymodemes of *L. tropica* and four of *L. infantum* were described so far [43, unpublished data]. The high degree of heterogeneity in *L. tropica* species has been reported [13,14,44] and shown by MLEE [30]. This variation was reported in different level in Old World *Leishmania* species by ITS PCR-RFLP analysis from highest to lowest in order of *L. tropica>*L. aethiopica>*L. major>*L. donovani [44]. A study using PCR-RFLP/sequencing based on ITS1 region was performed in Iran using clinical samples from CL patients and they showed six different genotype groups of *L. tropica* [45]. In our study, the heterogeneity of *L. tropica* was also observed in phylogenetic analysis. *L. infantum* isolates constituted a single group while strains determined as *L. tropica* showed several different groups.

The main difficulty in the present study was to observe two peaks in some samples. It is probably due to genetic variety in ITS1 region. Genetic polymorphism in ITS region of different strains of same *Leishmania* species and possibility of heterogeneity in individual copies was described by El-Tai et al [17]. In a recent study, after digestion of the amplification product with the HaeIII, the ITS1 PCR assay clearly identified the leishmanial species in CL samples in only 72.3% and *L. tropica* was found to be the most dominant [20]. In our study, the failing of species identification in 18.42% of the samples could be either due to the minuteness of the DNA, possibility of a mix infection, hybridization of different species or intraspecies variations in Turkish *Leishmania* parasites as...
also commented by Kifaya et al. [20]. We also found that there are 6 copies of our probe region containing variable part in ITS1 sequence (data not shown). Therefore, we evaluated that if all copies are identical in the DNA sample one peak was obtained; if not, several copies have one or two bases difference, two peaks were obtained but always one peak is higher. In the case of detection of two peaks, we decided to take higher one for species identification with the support of sequence analysis and it can be acceptable that the identification in species level was done in all clinical samples. Hereby, we can speculate that our assay could also propose intragenic heterogeneity of particular isolates and/or *Leishmania* DNA samples obtained from clinical materials. The statistical analyses of melting temperatures in the samples having one and two peaks was also performed and standard deviation was found very low in one peak samples than two peak samples. After we compared these results with sequencing results, we also found highest variability in *L. tropica* group. Gelanew et al. performed PCR-RFLP/direct sequencing assay using *L. donovani* strains and the direct sequencing of both strands of ITS1 DNA showed the presence of multiple peaks in the chromatograms, which could possibly have resulted from: (i) the presence of multiple strains or clones of *Leishmania*; (ii) the presence of a hybrid genotype; (iii) intragenomic variation in the multicye ITS1; or a combination of these [46]. This should be studied furtherly by multiple gene targets and innovative methods like Multilocus Sequence Typing (MLST) of the genes encoding the proteins used for MLEE and Multilocus Microsatellite Typing (MLMT) as well as experimental animal infection studies using Turkish isolates.

Two aims were achieved through the analysis of real time ITS1 PCR products by sequencing of the reference and local strains as designing new probes and proving the sensitivity of the technique at the genus and species identification level. The results of isolates in the optimization step also showed that real time ITS1 PCR results are highly concordant with MLEE analysis (96.07%). We can also propose that the method can identify inter- and intraspecies variability based on ITS1 region but cannot differentiate *L. donovani* complex species within each other like other molecular assays using ITS1 region [26]. In this point, we would like to emphasize that the studies addressed to *L. donovani* complex species (*L. donovani* and *L. infantum*) identification should be planned.

In conclusion, the proposed method presents a sufficient sensitivity for fast and correct diagnosis of leishmaniasis in all type of clinical samples but due to the samples giving two peaks the ability of the method for species identification is limited and needs further analyses. However, the higher peak was always very well matched with the results of sequence analysis. Genetic variations based on ITS1 region of *Leishmania* parasites in Turkey within species and intraspecies were determined. The findings in this study were showed that *L. tropica* is one of the causative agents of human and canine visceral leishmaniasis in Turkey.

**Supporting Information**

**Figure S1** The ITS1 region including the variable region used for designing probes in Turkish Leishmania isolates. [REF]: the sequences were taken from published paper, Talhi-Frank et al. (47); VL: visceral isolates; CL: cutaneous isolates; CanL: dog isolates. (TIF)

**Figure S2** STARD Flowchart of the present study. (TIF)

**Table S1** Identification of Turkish *Leishmania* isolates by ITS1 real time PCR method and isoenzymatic method. (DOCX)

**Table S2** STARD checklist for reporting of the study. (DOCX)

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**Author Contributions**

Conceived and designed the experiments: SOT YO MZA. Performed the experiments: ATV SOT GG. Analyzed the data: CG MZA. Contributed reagents/materials/analysis tools: FYZ HE GC. Wrote the paper: SOT YO MZA CG. We would like to thank the Balamuthia Committee on the Control of Leishmaniases. World Health Organ Tech Rep Ser 949, Geneva.

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