Detection and identification of cutaneous leishmaniasis isolates by culture, Polymerase chain reaction and sequence analyses in syrian and central anatolia patients

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ABSTRACT

Objectives: To characterize the cutaneous leishmaniasis (CL) isolates of Syrian and Central Anatolia patients at species levels.

Methods: Skin scrapings of 3 patients (2 Syrian, 1 Turkish) were taken and examined by direct examination, culture in Novy-MacNeal-Nicole (NNN) medium, Internal transcribed spacer Polymerase chain reaction and sequence analysis (PCR).

Results: According to microscopic examination, culture and PCR methods, 3 samples were detected positive. The sequencing results of all isolates in the study were identified as L. tropica. The same genotypes were detected in the 3 isolates and nucleotide sequence submitted into GenBank with the accession number: KP689599.

Conclusion: This finding could give information about the transmission of CL between Turkey and Syria. Because of the Syrian civil war, most of the Syrian citizens circulating in Turkey and different part of Europe, this can be increase the risk of spreading the disease. So, prevention measurements must be taken urgently.

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Leishmaniasis is a vector-borne parasitic disease caused by Leishmania genus protozoa and transmitted to the vertebrate host by female Phlebotominae (Diptera: Psychodidae) sand flies. Leishmaniasis currently ranks second only to malaria in the World Health Organization (WHO) list of protozoan diseases that cause the highest human mortality. It was estimated that approximately 350 million people are under the risk of leishmaniasis infection and 12 million are infected.¹ Leishmania is the main causative agent for cutaneous leishmaniasis in Old World with L. tropica. Whereas L. donovani or L. infantum are reported as visceral leishmaniasis causative agents. However, some cutaneous leishmaniasis (CL) cases have been reported by L. donovani or L. infantum particularly in Tunis and Yemen.² Self-healing zoonotic cutaneous leishmaniasis (ZCL) induced by L. major and anthroponotic cutaneous leishmaniasis (ACL) induced by L. tropica are 2 known types of CL.¹ The CL can be both dry and wet and it is a growing public health problem in Turkey. Southeastern Anatolian and Mediterranean regions are endemic foci for CL caused by L. tropica. Also, in neighboring countries at the southern border of Turkey (Syria, Iraq and Iran), L. major and L. tropica are responsible for CL.³⁻⁴ Most (up to 90%) of the global CL cases are observed in only a few countries: Afghanistan, Algeria, Iran, Iraq, Saudi Arabia, and Syria in the Old World.¹ CL was observed in very early years and is endemic to Syrian Arab Republic. After 1950s, the incidence decreased due to antimalarial sprays. But the number of CL cases sharply increased from the 80s onwards; 4,734 were observed in 1987-1988, 27,739 (new cases) in 2008, 46.148 in 2009 and 58.156 in 2011.³⁻⁵ Syria is the most affected nation among Mediterranean countries with more than 25,000 cases per year.⁶ Due to civil war in Syria, thousands of civilians were forced to leave their country and immigrate to safer countries such as Lebanon, Iraq, Jordan and Turkey. A significant increase in the number of Syrian CL cases was reported in Sanlıurfa Hospitals, Turkey.⁷ Approximately 2 million refugees currently live in different border provinces in Turkey such as Gaziantep, Sanlıurfa and Hatay and cutaneous leishmaniasis cases are expected to increase in these regions.

Clinical evaluation and microscopic demonstration of the parasite are not sufficient for a conclusive leishmaniasis diagnosis. Further molecular techniques are crucial in detection of the pathogen species. Molecular biological tools are widely used to screen Leishmania spp. infections in samples obtained from patients and vector species. Also phylogenetic techniques are used to evaluate genetic distances between strains isolated from humans, canines and vector sand fly species. Thus, we aimed to identify Syrian and Central Anatolian CL patient isolates at the species level using molecular techniques.
Methods. In this descriptive case study, 3 patients with CL suspicion were admitted to the National Reference Laboratories of Parasitology, Turkish Public Health Institution in 2014. Patients were all male and they were 3, 4 and 10 years old (Figure 1). Two patients were from Syria and one from Konya, Turkey. Two Syrian patients mentioned that lesions occurred before leaving Syria. The patients with skin lesions were included in the study.

The suspicious lesions were sterilized with 70% alcohol and allowed to dry. Skin was scraped with a sterile lancet and the smears were taken from the border of the lesion and divided into 2 parts. The first parts of the samples were fixed with methanol and stained with Giemsa. Then slides were examined under the microscope (100x) for the presence of the amastigote form of the parasite. The slides where amastigote was detected were declared positive after 15 minutes of examination.

The second part of the sample (preparation was the same with slide smears, a small amount of the scraped tissue) was inoculated on Biphasic Novy-MacNeal-Nicolle (NNN) medium. The NNN media included fresh, aseptically collected, defibrinated rabbit blood mixed with agar and penicillin G. And the liquid phase was supplemented with 0.85% saline. The cultures were produced at least in duplicates, incubated at 24 °C and observed by microscopy every 72 hours for 1 month. Positive cultures were subcultured in RPMI-1640 media for 3 days, for further propagation of Leishmania promastigotes. The parasites (approximately 50 mL culture) were washed 3 times by centrifuge (500 x g for 5 min) in cold phosphate-buffered saline (PBS) and the final concentration was adjusted to 5x10⁶ promastigotes/ml. The Leishmania promastigote pellets were suspended in 200 µl PBS and Proteinase K was added. Genomic DNA was isolated with QIAamp DNA Mini kit (Qiagen, Hiden, Germany), according to the manufacturer’s protocol.

Internal Transcribed Spacer 1 (ITS-1) gene was amplified to diagnose and identify the Leishmania species, using previously published primers (LITSR/ L5.8S). Amplification reactions were performed with the final 50 µL volume containing 5 µL PCR buffer (0.2 ml, Fermentas), 2 mM MgCl2 (25 mM, Fermentas), 1 ml dNTP (10 mM; Fermentas), 20 pMol of each primer, 5 µL genomic DNA and 2 units Taq polymerase (5U/µL, Fermentas). The cycling was conducted at 95°C for 5 minutes followed by 35 amplification cycles, each consisting of 3 steps: denaturation at 95°C for 20 s, annealing at 53°C for 30 seconds and extension at 72°C for 1 minutes, followed by a final extension at 72°C for 6 minutes. Finally, 8 µL PCR product was loaded in a 1.5% agarose gel electrophoresis and visualized by Ultraviolet (UV) trans-illuminator using a 100 bp DNA marker. Positive PCR products were sequenced as noted elsewhere.

Nucleotide sequences obtained in this study were aligned using ClustalW and the alignment was submitted to the GenBank. Basic Local Alignment Search Tool (BLAST®), used for search for similarities between the sequences, was applied to identify Leishmania species. Sixteen L. tropica sequences previously submitted to the GenBank from different countries were used to generate neighbor-joining (NJ) tree. L. siamensis was used as an out-group in the NJ tree. Maximum Parsimony (MP) and bootstrapping method was conducted with 1000 replicates using Geneious R.

Results. Microscopic examination demonstrated that 3 smears were positive for Leishmania spp. due to the observation of amastigotes in Giemsa stained slides.
(Figure 2). Also all tissue samples scraped from patients were successfully inoculated in the NNN media and promastigote forms were observed in all 3 samples approximately within 3 weeks. The PCR product of all 3 cultured specimens and one international control DNA demonstrated approximately 300-350 bp amplicon of expected size when visualized under UV trans-illuminator.

According to the BLAST® similarity search, the sequencing results for all isolates in the study identified L. tropica, the causative agent of ACL. The same genotypes were detected in 3 isolates and the nucleotide sequence submitted into GenBank with the accession number: KP689599. Neighbor-joining tree results demonstrated a slight diversion in these 3 isolate when compared to other isolates used in the tree except one (KJ002559), which was from Adana/T urkey (Figure 3).

Cutaneous leishmaniasis can be defined as a self-healing dermal disease and typically leaves depressed scars on the infection site. Until today, 3 different pathogens that cause cutaneous leishmaniasis were reported in Turkey. Also L. donovani was identified in a CL patient in France who visited Yemen for 2 weeks. Al-Nahhas and Kaldas analyzed 32 positive samples using ITS1-PCR RFLP method and L. tropica was found as the pathogen in all isolates. Also in the present study, L. tropica was found in 2 isolates that originated in Syria. Latest government report on Syrian refugees in Turkey emphasized the high-risk of new imported cutaneous leishmaniasis cases from Syria. In the first 3 months of 2013, the number of Syrian CL patients rapidly increased in the town of Nizip in the city of Gaziantep. In this period, 67 CL patients were identified and 62 (92%) patients were from Syria. Konya is the seventh populous city in Turkey, located in Central Anatolia. The city is non-endemic for CL, but 72 cases (0.4%) were reported between 1994 and 2000. Toz et al determined L. tropica in Central Anatolia, similar to the findings of the present study. Patient travel history was not available and the infection might be due to a visit to CL endemic region. Thus, it is not possible to conclude that CL agent was L. tropica in Central Anatolia.

In the present study, sequence analysis of Konya and Syrian isolates demonstrated 100% homology with each other and also with one isolate from Adana (KJ002559). There was not any information about the source of the infection in Konya, therefore it could be considered that the patient might have traveled to an endemic region, especially the cities on the border with Syria. Phylogenetic analyzes revealed that the sequence data obtained from patients slightly differed from the other sequences submitted previously to the GenBank.

![Figure 3 - Neighbor-Joining tree for the partial ITS1 rRNA sequences of the isolates.](Image)
The present study demonstrated that utilizing molecular based assays on leishmaniasis provides specific identification of the species.

In this study, existence of *L. tropica* demonstrated the risk of anthroponotic CL in the study areas. Due to the Syrian civil war, several Syrian citizens travel in Turkey and also in different parts of Europe. This may increase the risk of spreading the disease, therefore prevention measures must be taken urgently.

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References


Clinical Practice Guidelines

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