Original Article

Epidemiological Study on Cutaneous Leishmaniasis in an Endemic Area, of Qom Province, Central Iran


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Abstract

Background: Cutaneous leishmaniasis (CL) is one of the most important health problems in many areas of Iran. There are two forms of the disease in Iran, anthroponotic and zoonotic CL. This study conducted to assess the epidemiological situation of CL in an endemic area of Qom Province, central Iran from Apr to Nov 2015.

Methods: The sticky paper traps and aspirating tubes were used for collecting adult sand flies. Sherman traps and small insect nets were used to capture rodents and small mammals. Giemsa staining was used for preparing the expanded smear and followed by PCR for identifying the causative agent in human, vectors, and reservoirs. In this study, relative frequency of CL was also calculated.

Results: Fourteen species of Phlebotomine sand flies were collected. Phlebotomus papatasii (61.74%) was the predominant species through the period of activity. Overall, 62 Meriones libycus, 8 Nesokia indica, 4 Mus musculus, 16 Al-lactaga elater and 2 Hemiechinus auritis were caught. PCR technique showed 6 out of 150 P. papatasii (2%), two out of 62 M. libycus (3.23%) and all of suspected human's skin tissue samples (100%) were infected with Leishmania major. The relative frequency of CL was 0.30%.

Conclusion: This is the first detection of L. major within P. papatasii, M. libycus and human in Kahak District in Qom Province of Iran. Zoonotic cycle of CL exists in this area, L. major is the causative agent, P. papatasii is the main vector and M. libycus is the main reservoir of the disease.

Keywords: Zoonotic cutaneous leishmaniasis, Leishmania major, Phlebotomus papatasii, ITS1-PCR, Iran

Introduction

Currently, cutaneous leishmaniasis (CL) is one of the most important vector borne diseases in Iran (1) and still the leading cause of considerable morbidity of a large number of people in the endemic foci characterized by chronic skin lesions followed by scars and deformation of the infected tissue (2).

Phlebotomus papatasii is the main and proven vector and Leishmania (Leishmania) major is the causative agent of Zoonotic CL in Iran. Leishmania major, L. (L.) tropica and L. (L.) aethiopica cause CL in the Old World (3). The parasite has been isolated and identified from naturally infected P. papatasii, P. caucasicus, Rhombomys opimus, Meriones libycus and human in such endemic areas (4-8). The P. papatasii and M. libycus were as proven vector and reservoir in Qomrood, Iran

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district (9) and as probable vector and reservoir in Ghanavat district (10). In addition, a big part of the lowland areas of Qom Province provides good ecological niches for P. papatasi and therefore it has higher transmission potential (11).

In the recent years, molecular methods have been employed for identification of certain species of *Leishmania*, either isolated from cultures or from patients (3) as well as in the detection of the parasite in individual or pooled Phlebotomine specimens (12). Successful establishment of the disease in an endemic area is the outcome of a close association between the *Leishmania* parasite and its natural sand fly vector (13). Thus, vector and parasite identification have great impact on predicting expansions of the disease in endemic area, also help authorities to design new strategic programs to limit spreading vectors and disease (14, 15).

Molecular methods are increasingly employed for diagnostic and epidemiological purposes in order to confirm *Leishmania* infection and to characterize the parasites at the species or genotype level in hosts and vectors (16, 17). The detection of *Leishmania* parasites by PCR methods is highly specific and sensitive, with values reaching up to 100%. Accurate and sensitive diagnostic and identification procedures are required to distinguish *Leishmania* species/strains whose geographic distribution can overlap, which is crucial for adequate treatment and appropriate public health control measures (18).

Based on a rapid increase in incidence of CL reported in an endemic area of Kahak District of Qom Province in central Iran (19), and due to lack of knowledge on the epidemiological situation of CL, this study was conducted to determine the epidemiological features of CL including human infection and the reservoir hosts and their putative vector species in this endemic area during 2015.

### Materials and Methods

#### Study area

The Qom Province is bounded by Tehran Province in the north, Esfahan Province in the south, Semnan Province in the east, and Markazi Province in the west with an area of approximately 11240 square kilometers (0.68% total area of Iran) (Fig. 1). This study was performed from Apr to Nov 2015 in 3 villages (Khor Abad, Sarm, and Ghobadbezan) of Kahak rural district (34°09′–35°11′ N latitude and 50°06′–51°58′ E longitude) of Qom Province with the elevation of almost 1500m above sea level (20). The average annual minimum and maximum temperatures were 16.5 °C and 49 °C in Jan and Jul, respectively. The total annual rainfall was about 150mm and the average Max and Min monthly Relative humidity were 84% and 28% in Dec and Jun, respectively (21).

#### Sand fly collection

Sand flies were collected from indoor (bedroom, bathroom, toilets, hall, and stables) and outdoor (rodent burrows) fixed places from the first half of Apr 2015 to the first half of Nov 2015. To capture the sand flies, 30 Sticky Paper Traps (castor oil coated white paper 20×32cm) was used twice a month from sunset to sunrise. The caught sand flies were transferred to the laboratory in Qom Health Center. To species identification, the head and the last two abdominal segments of the sand flies were mounted in Puris’ medium (24) and identified after 24–72h, using the morphological characters (25). Then, they were counted and segregated by sex. The parous females were distinguished from nulliparous sand flies by observation of the
appearance of the accessory glands (26). In order to determine the natural promastigote infections of female sand flies, some unfed, blood fed, semi-gravid and gravid female sand flies (captured from rodent burrows) dissected in a fresh drop of sterile saline (9/1000) for the presence of promastigotes in alimentary canal during Jun and Sep 2015.

Reservoirs collection and smear preparation

The rodents were captured by Sherman Live Traps (Fig. 2). Collection of dipodids (Rodentia: Dipodidae) was done by using small insect nets. Then the impression smears of ears were fixed with absolute methanol and examined by Giemsa staining method. The Leishman bodies were observed using light microscope and then were subjected to molecular technique for identification. Tissue samples were also taken from the edge of the lesions in 45 suspected CL patients. Lesion smears were fixed with absolute methanol and then stained with Giemsa for CL diagnosis. In this present study, we also calculated the relative frequency of CL.

DNA extraction

DNA extractions were carried out by crushing of the contents on the slides prepared from the digestive system of the 180 dissected female sand flies, the 78 rodent's ear lobes serous fluids and the 45 patient's lesion tissues, using the Bioneer Genomic DNA Extraction Kit. The extracted DNA was stored -20 °C for long storing and at 4 °C for daily working. DNA from L. major (MRHO/IR/75/ER) that provided by Department of Medical Parasitology, School of Public Health, Tehran University of Medical Sciences (TUMS), Iran was used as positive control. Total DNA was extracted from the smears by digestion in 100 μl PBS buffer and the tube was centrifuged at 10000rpm for 10min, then 300μl lysis buffers and 30μl proteinase K added. The tube was incubated for 24h at 37 °C before adding 300μl saccharin phenol. After adding this solution, the tube was centrifuged at 9300 rpm for 5min. After transferring upper phase to new tube, 300μl phenol- chloroform should be added and was centrifuged at 10000rpm for 5min. Again transferred the upper phase to new tube and washed with pure chloroform. Thirty μl MgCl₂ and 1000μl ethanol were added to upper phase and stored at -20 °C for 2h before was centrifuged at 10000rpm for 10min and washed down phase by 70% ethanol with TE and was centrifuged at 1000 rpm for 10min and the TE buffer was added.

Polymerase chain reaction-restriction fragment length polymorphism (PCR-RFLP) for detection of Leishmania infection

The DNA samples were examined for the Leishmania species ITS1 by PCR amplification using the universal primer pair L5 8S (5´-TgA-TaC-CAC-TTA-TCg-CAC-T-<T>-3´) and LITSR (5´-CTg-gAT-CAT-TTT-CCg-AT-<g>-3´) (Table 1).

Molecular study

PCR production was followed by RFLP technique (27). The cycling conditions were 95 °C for 5min., followed by 35 amplification cycles, each consisting of three steps: denaturation at 94 °C for 30sec, annealing 48 °C for 30sec, and extension at 72 °C for one min, followed by a final extension at 72 °C for 7min in thermocycler.

PCR productions were digested with the restriction endonuclease Hae III for 2 h at 37 °C. The restriction fragments were separated by electrophoresis on agarose gel and compared with those of standard reference strain of L. major and negative control (distilled water).

Results

A total of 4164 sand flies (68.90% from outdoors and 31.10% from indoor resting
places) were collected using sticky paper traps. Two genera (Phlebotomus and Sergentomyia) and 14 species were identified, including Phlebotomus (Phlebotomus) papatasi (61.74%), P. (Paraphlebotomus) sergenti (6.87%), P. (Paraphlebotomus) alexandri (1.00%), P. (Paraphlebotomus) caucasicus (2.55%), P. (Paraphlebotomus) caucasicus group (4.13%), P. (Larroussius) kandelakii (5.72%), P. (Larroussius) tobbi (0.10%), P. (Larroussius) major (5.62%), P. (Adleriuss) halepensis (0.10%), P. (Adleriuss) brevis (0.05%), P. adleriuss group (0.31%), Sergentomyia (Ser-gentomyia) sintoni (10.83%), S. (Sergentomyia) theodori (0.91%) and S. (Rondanomyia) pawlowskii (0.07%) (Table 2).

Two species of P. halepensis and the S. pawlowskii were not captured from outdoor and indoor respectively. In the rodent burrows, P. papatasi appeared nearly May and disappeared in the late Oct and in indoors, it appeared in the late Jun and disappeared in the early Oct.

The month wise density of P. papatasi in rodent burrows and indoors are shown in Fig. 3.

The peak of activity was detected for P. papatasi first in the late Jun and the second in the early Aug (Fig. 3).

The sex ratio calculates at 83.71 and 62.25 in outdoors and indoors for P. papatasi respectively as well as 101.5 in outdoors for S. sintoni. In Aug and Sep 2015 a total of 211 sand flies were collected in the vicinity of rodent burrows including P. papatasi (51.66%), P. caucasicus (10.9%), P. sergenti (18%), and S. sintoni (19.44%).

Two percent of P. papatasi species had promastigote infections. The total of 300 P. papatasi specimens was selected and dissected for Leishmania infection, the result showed 67.33%, and 32.67% dissected sand flies were parous and nulliparous, respectively (Table 3).

The analysis of physiological status of dissected P. papatasi revealed (60%) unfed, (20%) blood-fed, (6.67%) semi gravid and (13.33%) gravid (Table 3). The used Leishmania primers of the PCR technique successfully amplified the ITS1 region of the strains L. major and L. tropica revealed that among 360 specimens sand flies examined by PCR, 6 parous specimens (2%) were infected by L. major (Fig. 4, 5).

Among 92 collected rodents (67.39%), (8.70%), (4.35%), (17.39%) and (2.17%) were identified as M. libycus, Nesokia indica, Mus musculus, Allactaga elater and Hemiechinus auritis respectively (Table 4). Leishmania major has been detected in 2 (3.23%) out of 62 M. libycus by PCR (Fig. 6). Both of the infected M. libycus were females.

All of the 45 human cases, which examined through observation passively, confirmed by PCR but only 15 numbers of them were positive by light microscope. In this study, the relative frequency of CL patients was 0.30%.

Twenty-eight (62.22%) of 45 confirmed cases were males and 17 (37.78%) were females. The mean age of patients was 30.78 ± 16.91. The most of the cases (75.60%) occurred in autumn. Eighteen cases (40%) of patients had no positive history of traveling to leishmaniasis endemic areas during the past year. Twenty-two cases (48.89%) of patients had one lesion. The most common location of lesion was on hands (46%). The results of PCR-RFLP indicated that 45 (100%) cases were infected as L. major (Fig. 7).

Table 1. The sequence of primers used in PCR amplification the first internal transcribed spacer (ITS1)

<table>
<thead>
<tr>
<th>Primers</th>
<th>The sequence</th>
<th>Response function</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>LITSR</td>
<td>5'CTGGATCATTTCGGATGG'</td>
<td>Forward</td>
<td>18bp</td>
</tr>
<tr>
<td>L5 8S</td>
<td>5'TGATACACTTATCGCATT3'</td>
<td>Reverse</td>
<td>20bp</td>
</tr>
</tbody>
</table>

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Table 2. The fauna and the number of collected sand flies from the endemic area in Kahak District, Qom Province, 2015

<table>
<thead>
<tr>
<th>Site</th>
<th>Outdoor</th>
<th>Indoor</th>
<th>Total</th>
<th>Each species of the total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Male</td>
<td>Female</td>
<td>Male</td>
<td>Female</td>
</tr>
<tr>
<td></td>
<td>No. (%)</td>
<td>No. (%)</td>
<td>No. (%)</td>
<td>No. (%)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P. papatasi</td>
<td>915 (45.57)</td>
<td>1093 (54.43)</td>
<td>216 (38.37)</td>
<td>347 (61.63)</td>
</tr>
<tr>
<td>P. sergenti</td>
<td>41 (40.20)</td>
<td>61 (59.80)</td>
<td>11 (71.20)</td>
<td>19 (28.80)</td>
</tr>
<tr>
<td>P. alexandri</td>
<td>7 (58.33)</td>
<td>5 (41.67)</td>
<td>11 (36.67)</td>
<td>19 (63.33)</td>
</tr>
<tr>
<td>P. caucasicus</td>
<td>45 (100)</td>
<td>…</td>
<td>61 (100)</td>
<td>…</td>
</tr>
<tr>
<td>P. kandelakii</td>
<td>23 (31.51)</td>
<td>50 (68.49)</td>
<td>15 (21.21)</td>
<td>50 (78.79)</td>
</tr>
<tr>
<td>P. tobbi</td>
<td>1 (100)</td>
<td>0 (0)</td>
<td>3 (100)</td>
<td>0 (0)</td>
</tr>
<tr>
<td>P. major</td>
<td>6 (24.00)</td>
<td>19 (76.00)</td>
<td>60 (28.71)</td>
<td>149 (71.29)</td>
</tr>
<tr>
<td>P. halepensis</td>
<td>0 (0)</td>
<td>…</td>
<td>4 (100)</td>
<td>…</td>
</tr>
<tr>
<td>P. brevis</td>
<td>1 (100)</td>
<td>…</td>
<td>1 (100)</td>
<td>…</td>
</tr>
<tr>
<td>P. adlerius group</td>
<td>…</td>
<td>2 (100)</td>
<td>…</td>
<td>11 (100)</td>
</tr>
<tr>
<td>S. sintoni</td>
<td>207 (50.36)</td>
<td>204 (49.64)</td>
<td>18 (45.00)</td>
<td>22 (55.00)</td>
</tr>
<tr>
<td>S. theodori</td>
<td>15 (51.72)</td>
<td>14 (48.28)</td>
<td>6 (66.67)</td>
<td>3 (33.33)</td>
</tr>
<tr>
<td>S. pawlowski</td>
<td>0 (0)</td>
<td>…</td>
<td>3 (0)</td>
<td>…</td>
</tr>
<tr>
<td>Total</td>
<td>1261 (43.93)</td>
<td>1608 (56.07)</td>
<td>543 (58.08)</td>
<td>752 (41.92)</td>
</tr>
</tbody>
</table>

Fig. 1. Map of Iran and Qom Province, highlighting the location of Kahak District, (2) within Qom Province (22, 23)
Table 3. Natural infection of sand flies with promastigote, Kahak District, Qom Province, 2015

<table>
<thead>
<tr>
<th>Species</th>
<th>Catch place</th>
<th>Physiological status</th>
<th>Age Group</th>
<th>Total</th>
<th>Infected specimens</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>I*</td>
<td>O**</td>
<td>Unfed</td>
<td>Blood fed</td>
<td>Semi gravid</td>
</tr>
<tr>
<td><em>P. papatasi</em></td>
<td>150</td>
<td>50</td>
<td>180</td>
<td>60</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td>20</td>
<td>50</td>
<td>20</td>
<td>50</td>
<td>50</td>
</tr>
<tr>
<td><em>P. sergenti</em></td>
<td>10</td>
<td>50</td>
<td>10</td>
<td>50</td>
<td>50</td>
</tr>
</tbody>
</table>

*: Indoor, **: Outdoor

Fig. 2. Specimens of some collected rodents in Kahak district, Qom Province, 2015

Fig. 3. The month wise density of *Phlebotomus papatasi* in Kahak District, Qom Province, 2015

Fig. 4. Agarose gel electrophoresis of the first internal transcribed spacer (ITS1) - Polymerase chain reaction (PCR) products. M, 50bp ladder: lane1: *Leishmania major* (MRHO/IR/75/ER), 2, 3: *Leishmania major* in human skin samples, lane 4, 5, 6 *Leishmania major* in *P. papatasi*, N: negative control (distilled water), M: Marker

Fig. 5. Agarose gel electrophoresis of Polymerase chain reaction-restriction fragment length polymorphism (HaeIII) products. M, 50bp ladder: lane 1, 2, 3, 4, 5, 6 *Leishmania major* in *P. papatasi*, lane 7 positive *Leishmania major* (MRHO/IR/75/ER), M: Marker

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Fig. 6. Agarose gel electrophoresis of the first internal transcribed spacer (ITS1) - Polymerase chain reaction (PCR) products. M, 50bp ladder: lane 1, negative control (distilled water), lane 2, *Leishmania major* (MRHO/IR/75/ER), lane 3 *Leishmania major* isolated from *Meriones libycus*, M: Marker

Fig. 7. Agarose gel electrophoresis of Polymerase chain reaction-restriction fragment length polymorphism (HaeIII) analysis of ITS region for identification of *Leishmania* species using M, 50bp ladder: lane 1, 2, 3, 4 *Leishmania major* in human skin samples, lane 5, 6 *Leishmania major* isolated from *Meriones libycus*, lane 7, positive *Leishmania major* (MRHO/IR/75/ER), M: Marker

Table 4. Molecular detection of *Leishmania* infection in rodents, Kahak District, Qom Province, 2015

<table>
<thead>
<tr>
<th>Species</th>
<th>No. Rodents</th>
<th>No. examined</th>
<th>No. infected</th>
<th>Infected rate, % (ITS1-PCR)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Meriones libycus</em></td>
<td>62</td>
<td>67.39</td>
<td>62</td>
<td>100</td>
</tr>
<tr>
<td><em>Nesokia indica</em></td>
<td>8</td>
<td>8.70</td>
<td>8</td>
<td>100</td>
</tr>
<tr>
<td><em>Mus musculus</em></td>
<td>4</td>
<td>4.35</td>
<td>4</td>
<td>100</td>
</tr>
<tr>
<td><em>Allactaga elater</em></td>
<td>16</td>
<td>17.39</td>
<td>4</td>
<td>25</td>
</tr>
<tr>
<td><em>Hemiechinus auritis</em></td>
<td>2</td>
<td>2.17</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>92</strong></td>
<td><strong>100</strong></td>
<td><strong>78</strong></td>
<td><strong>84.78</strong></td>
</tr>
</tbody>
</table>

Discussion

We identified *L. major* in *P. papatasi*, *M. libycus* and human by PCR technique. Previous surveys carried out in Qomrood and Ghanavat districts of Qom Province, Iran showed that *P. papatasi* and *M. libycus* were proven (in Qomrood) and probable (in Ghanavat) vector and reservoir of ZCL (9, 10). Awareness of epidemiology of ZCL is an important measure for management and planning of control (28). The entomological and reservoirs (rodents and human) fields and laboratory surveys accompanied by correct identification the common agent in their target part of their bodies is a major component for combatting against disease.

Recently, genetic analyses and genotyping using PCR-RFLP have been applied and performed on vector (s) and reservoir hosts of ZCL (29). This paper reviews recent advances on sand flies vectors, *M. libycus* and human of ZCL, using molecular biological approaches. *P. papatasi* captured in this study is the main vector of *L. major* from animals to man in central Asia and appears to be the vector of *L. major* in Iran and 21 other countries in old world (30).

In this study, *P. alexandri* was also captured. It has been found naturally infected with promastigotes and is suspected vector of VL in Iran (31). *Phlebotomus caucasicus group*, *P. kandelaki* and *P. major* were also collected. *Phlebotomus caucasicus* was naturally infected with promastigotes in a new focus of VL in north-west of Iran (32). *Phlebotomus kandelaki* is naturally infected with *Leishmania* sp. promastigotes in northwest Iran and is sus-
pected as probable vector of VL in the region (33). In Iran, P. major has been found in all area where human cases of VL have been reported and natural promastigote infection of this species has reported in endemic focus VL in Ghar County (Fars Province) South of Iran (34). Moreover, in this area, we captured S. povlovsyki in indoor. This result has been confirmed with the previous study in Esfahan (35). The identified species were consistent with previous study in other parts of Iran (1).

Low percentage of collected sand flies in indoor shows people prevent sand flies to enter their homes. P. papatasi was the predominant species of the genus Phlebotomus in indoors (31.10%) and outdoors (68. 90%). This suggests that P. papatasi is the anthropophilic species in this area. In addition, only P. papatasi is able to develop L. major in its mid gut and transfer the parasite to its proboscis to cause ZCL (36). S. sintoni enters the indoors because of nearby homes to rodent burrows (37). P. major and P. kandelakii are more dominant sand flies were collected from indoor after P. papatasi but due to the low frequency. These species do not involve in leishmaniasis transmission.

In the present study, we distinguished parity of sand flies by observation of the appearance of the accessory glands (26). Accessory glands secretions could not be as an indicator for distinguishing parous from nulliparous of P. papatasi females (38). In this study, the parous rate of P. papatasi was 67.33%, so it is not surprising to see high level of promastigote infection among this sand fly species. The same trend of infection is reported from north Siani in Egypt (39, 40). The results of dissection showed that 2% of P. papatasi species were naturally infected with promastigote infection. The Leishmania infection rate in sand flies is usually very low even in endemic areas (41). The low density of Leishmania parasites in M. libycus in this area may be due to low sensitivity of this rodent compared to R. opimus (4).

In the present study, based on molecular results, L. major was found in M. libycus as a reservoir, P. papatasi as a vector and suspected patients as incidental host. M. libycus as the main reservoir host of ZCL has been distributed in the northeastern, central and southwestern regions in Iran (42, 43). For the first time in 1996, isolation and characterization of L. major from M. libycus in Iran, has been reported (4). Meriones libycus has previously been infected with L. major parasites from Golestan, Esfahan and Yazd Provinces (7, 42, 44-46). Leishmania major was firmly identified in M. libycus that indicates this rodent species can be incriminated as reservoir host of ZCL in this location (47). The results of this study have shown, M. libycus is the main and potential reservoir host of ZCL in Kahak district of Qom Province and it has an important role in stable ZCL in this area, which has pay attention to future rodent control programs with administrative health center.

Conclusion

This study confirms the existence of local transmission of zoonotic cycle of CL in Kahak District of Qom Province, and provides that in this cycle, L. major is the causative agent, P. papatasi is the main vector and M. libycus is the main reservoir of the disease.

Acknowledgments

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References


