A Study of Genetic Polymorphism of *Crithidia* in Isfahan, Iran

Monir Doudi 1; Marziyeh Karami 2; Gilda Eslami 3; Mahbubeh Setorki 4

1Department of Microbiology, Falavarjan Branch, Islamic Azad University, Isfahan, IR Iran
2Young Researchers Club, Falavarjan Branch, Islamic Azad University, Isfahan, IR Iran
3Department of Parasitology and Mycology, Shahid Sadoughi University of Medical Sciences, Yazd, IR Iran
4Department of Biology, Izeh Branch, Islamic Azad University, Izeh, IR Iran

*Corresponding author: Monir Doudi, Department of Microbiology, Falavarjan Branch, Islamic Azad University, Isfahan, Iran. E-mail: doudi@iaufala.ac.ir*

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**Background:** Genotyping of *Crithidia* species is critical in order to design appropriate programs for control and prevention of cutaneous leishmaniasis (CL).

**Objectives:** The purpose of this project was study and investigation genetic polymorphism of *Crithidia* in Isfahan city.

**Patients and Methods:** In this descriptive-analytical study, 602 samples were isolated from suspicious patients with cutaneous leishmaniasis who referred to Leishmaniasis Research Center or other health centers in the vicinity of Isfahan. After culturing of the samples in NNN (Novy MacNeal Nicole, ©Merck) medium, DNA extraction was performed. Molecular detection and identification was done using PCR-RFLP technique.

**Results:** Results showed that 50 samples (8.31%) were negative. Out of 201 samples (33.39%) were *Crithidia* spp which 27 cases (13.44%) were *Crithidia fasciculata*, 11 cases (5.47%) were *Crithidia luciliae* and the rest (163 cases or 81.9%) were similar to Trypanosomatidae which submitted at gen bank with accession number of GQ331988. BLAST showed that the last ones have 92% and 89% similarity to *C. fasciculata* and *C. luciliae*, respectively. Out of 351 patients (58.31%) identified as *Crithidia* spp.

**Conclusions:** The results of the present study suggested that there is some recombinant species in Isfahan and surrounding area that could be an important agent of cutaneous leishmaniasis.

**Keywords:** Polymorphism; *Crithidia*; Internal Transcribed Spacer 1 (ITS1); Polymerase chain reaction - Restriction Fragment Length Polymorphism (PCR-RFLP)

1. **Background**

Blood and tissue flagellates are a wide range of protozoan parasites, which are, at least in one of the evolutionary stages of flagellum. Blood and tissue flagellates consist of 1-4 forms in a protozoan life cycle, which are the following: 1- Promastigote (*Leptomonas pessowi*), 2- Epimastigote (*Crithidia* form), 3- Tryomastigote (*Trypanosomiasis*), 4- Amastigote (*Leishman body*) [1].

*Crithidia* is considered as insect parasites; however, since in gene sequences of rRNA they are close to *leishmania*, most often they accompany and are mistaken with *leishmania* [2]. *Crithidia* oncopelti is isolated from a sap-feeding insect (presumably the plant *Oncopeltus fasciatus*), *Crithidia deanei* is isolated from a predatory insect called *Zelus leucogrammus* and *Crithidia desouzaii* from a nectar-feeding fly named *Orinidia Obesa* [3-5]. *Crithidia luciliae* and *C. fasciculata* are not only insect parasites, but also are considered as the infecting agents of *leishmania* cultivation [2]. During the past 20 years, a variety of methods have been developed to identify different sub-species of *leishmania* and *Crithidia*, and study the molecular diversity and mutual influences of parasites and host [6-8]. It is worth mentioning that methods with the following features must be used for the distinction and differentiation of species that are very close together: 1- They should be practical, 2- be done quickly, 3- be cost-effective 4- do not require special skills. During recent years, various genetic and molecular techniques to isolate the pathogenic strains of diseases have become important in epidemiological and clinical studies and the study of other infectious diseases. Developments of molecular techniques using molecular markers and different polymerase chain reaction (PCR) methods have been reported for phylogenetic differentiation and identification of parasites [9]. In the recent decade, many of the studied molecular markers for the polymerase chain reaction (PCR) were: Microsatellite DNA, Kinetoplastic DNA (KDNA), Telo-meric sequences, or genes such as gp63, hsp70, mini-ex-
on, tubulin β, and/or ribosomal RNA genes. Among the above mentioned RFLP (Restriction fragment length polymorphism) techniques related to PCR, amplified sequences corresponding to internal transcribed spacer (ITS) gene have suggested promising results [10, 11]. Consequently, it is an appropriate method to investigate phylogenetic polymorphism and relationship between organisms. ITS sequence analysis can show the phylogenetic relationship of many parasites [12, 13]. The structure of ribosomal gene locus has two DNA areas which encode the small subunit of 18s SSU rRNA and the large subunit of 28s LSU rRNA. At the 5’-end region of the 18s, there is a fragment named ETS (External Transcribed Sequences) which continues from 5’-end to the promoter sequence [14, 15].

The PCR-RFLP technique is a sensitive and specific tool for detection and identification. In this technique, the types of endonucleases are used that are able to detect specific nucleotide sequences on DNA organic sequences bases and restriction of the deoxyribonucleic acid (DNA) specifically or non-specifically. Restriction of these enzymes provides DNA fragments in different sizes and comparing these fragments in various isolations can provide a more accurate estimation of sequence difference of DNA in various isolations [16, 17]. This research project aimed to review different genotypic Crithidia through PCR and RFLP with the help of ITS1 gene sequences.

2. Objectives

This infectious agent is genetically polymorphism and it was hypothesized that there could be a probable correlation between the prevalence rate of the infection and genotypic variations of the studied stains and also their geographic origin.

3. Patients and Methods

3.1. Patients and Study Region

This study was descriptive-analytical and carried out in Isfahan and its surrounding areas, located in the central part of Iran. There were 201 study samples. Sampling started from 21 March 2011 to 21 November 2011 (for 8 months). The selected cutaneous leishmaniasis lesions (wounds) in the patients had various forms.

Sample collection and preparation: First, the lesions were sterilized using 70% ethanol and saline serum, and then following a local anesthesia, xylocaine solution was used for them. Thereafter, surface cuts of 2 to 3 mm were made on the edges of the lesions and then the prepared restrictions were impregnated into the Brain Heart Infusion Broth (BHI medium, ©Merck) and sterilized in 4°C or saline solution as the medium and then were transferred into cultured medium NNN (Novy Mac Neal Ni-

cole, ©Merck) close to the flame and finally were placed in incubator in 24 ± 2°C.

Isolation and culture: As mentioned earlier, liquid BHI mediums containing species were also transferred close to flames in sterile conditions in NNN medium in 24 ± 2°C and were examined every three days for 6 weeks before any negative report. Then, the amplified promastigotes were transferred and amplified in culture medium RPMI 1640 with 10% FCS (Sigma). At the late stage of logarithmic growth (5 × 106 parasites/mL), isolated species were taken out and were sterilized with saline solution in phosphate buffer (PBS) and were washed with centrifuge pH = 7.4 and were moved to a freezer in -70°C for further tests and examinations. The reference strains used in this study were: Crithidia luciliae and Crithidia fasciculata.

DNA extraction: Suspension of promastigotes of the 2 washed Leishmania and Crithidia isolates in the previous stage with 5 × 106 parasites/mL were taken out of -70°C freezer. Two hundred microliters of Binding Buffer and 40 μL of protease K were added to its precipitation, and the contents were placed in a water bath (bain-marie) for 10 min in 70°C. The rest of the stages were done according to kit’s instruction (High pure PCR Template Preparation kit, Roche). Finally, DNA was dissolved in 200 μL of elution buffer. The extracted DNA was studied quantitatively and qualitatively by spectrophotometry, and then agarose gel 0.8% in TBE 1 × buffer.

Amplification of ITS1 using PCR: Amplification of ITS1 using PCR was conducted through the LITSr primers L5.8S with the following sequences:

LITSr (5’-CTGGATCATTTTCCGATG-3’)
L5.8S (5’-TGATACATATTGCGACTT-3’)

The best condition for implementing PCR was determined by using the standard strains. The reaction was performed in 25 μL. First, a Master Mix was prepared which consisted of 0.2 μL of Taq polymerase (50 Mm), 2 μL MgCl2, R and F primers with concentration of 10 picomoles 1 μL each, 2 μL PCR10x Buffer, dNTPs (10 Mm) for 1 μL, and finally 1 μL extracted DNA was added to the above mentioned items, and the final volume of the distilled water reached 25 μL. The samples were placed into the thermocycler device (Corbett, Australia) and the following cycles were performed for the PCR implementation:

Thereafter, the PCR products with positive and negative controlling along with molecular marker of 50 bp were placed on 1.5% agarose gel containing 0.7 μg/mL ethidium bromide per ml of gel in TBE 1 × buffer and then were electrophoresed for 25-45 minutes in a voltage of 80 V. The PCR products were observed in transilluminator device using UV lamp [18].

RFLP analysis for amplified ITS1: In order to detection of the genus, the products of ITS1-PCR were digested under the effect of restriction enzyme HaeIII (Fermentas) and the related buffer in 37°C for 2 hours. Analysis of the digestion was done using 2% agarose gel electrophoresis in TBE 1 × buffer under 5 V/cm. Molecular iden-
Identification was performed using RFLP with restriction enzymes of TaqI, ScrFI and DdeI. All of the digestions were compared with the standard strains of *Crithidia* and *Leishmania* [18].

DNA sequencing: Samples with different patterns of RFLP were purified and sequenced. Then, BLAST analysis was done for sequenced samples.

Statistical analysis: Finally, the collected data were analyzed using descriptive statistics and \( \chi^2 \) test in SPSS-16.0 (SPSS Inc., Chicago, IL, USA). P-Values less than 0.05 were considered significant.

4. Results

This 8-month study was carried out on 602 patients with wet CL who referred to the Isfahan Leishmaniasis Research Center or other health centers in the vicinity of Isfahan. Culturing in NNN medium, DNA extraction from promastigotes, implementation of the PCR-RFLP method, amplification of ITS-1, and application of the HaeIII enzyme were carried out. As a result, 201 patients (33.39%) were found with two species of *Crithidia*; 27 cases of *Crithidia fasciculata* (13.44%) and 11 cases of *Crithidia luciliae* (5.47%).

The rest, however, were trypanosomatidae with Accession number of GQ331988; molecular analysis showed that 92% had a tendency of homology with *C. fasciculata* and 8% with *C. luciliae* (163 cases or 81.9%). Out of 351 patients (58.31%), Leishmaniasis parasite was isolated, and out of 50 patients (8.31%), neither Leishmaniasis nor *Crithidia* were isolated. Therefore, their report was negative following cultured samples. The results of the present study showed that of the 201 patients studied in these two areas, 103 of them (51.24%) were related to Isfahan and 98 samples (48.76%) were related to the surrounding areas of Isfahan (P = 0.001).

The results of PCR-RFLP analysis in Isfahan and surrounding areas with HaeIII enzyme restriction suggested that a number of isolated species showed a single-band at 350 bp, which was related to the *leishmania* species. Moreover, a number of them showed no band which was neither *Leishmania* nor *Crithidia*. A number of them also manifested a single-band at 400 bp and many other isolated species showed single-band at 450 bp.

Finally, a large number of isolated species showed two bands at 400 and 450 bp; the last three groups are related to the trypanosomatidae family (Figure 1).

It should be mentioned that the last three groups, which were related to the trypanosomatidae family, had no difference in restriction enzymes compared with standard species after restriction with other enzymes (DdeI, ScrFI, and TaqI) (Figures 2, 3, 4).

The generated band profiles after the effect of DdeI, ScrFI, and TaqI enzymes on isolations with bands at 450 and 400 bp and two bands of 400 and 450 bp simultaneous included.
Following the effect of the DdeI enzyme on all the isolations with band at 400 bp, fragments of 307 and 92 bp were observed. Following the effect of the ScrFI enzyme on all the isolations with band at 400 bp, no fragment was observed. On the other hand, after the effect of the TaqI enzyme on all isolations with band at 400 bp, fragments of 275 and 124 bp were observed.

Following the effect of the DdeI enzyme on all isolations with band at 450 bp, fragments of 331 and 115 bp were observed. Following the effect of the ScrFI enzyme on all isolations with band at 450 bp, fragments of 321 and 125 bp were observed. On the other hand, after the effect of the TaqI enzyme on all isolations with band at 450 bp, fragments of 217 and 229 bp were observed.

Furthermore, after the effect of the DdeI enzyme on all isolations with the two bands at 400 and 450 bp, fragments of 334 and 115 bp were observed. Moreover, after the effect of the ScrFI enzyme on all isolations with the two bands at 400 and 450 bp, fragments of 323 and 126 bp were observed.

On the other hand, following the effect of the TaqI enzyme on all isolations with the two bands at 400 and 450 bp, fragments of 301 and 148 bp were observed.

For genotyping the isolation of trypanosomatidae, the enzyme profiles of which have shown a non-compliance restriction with the standard species, these samples were transferred to Farapajooh Company in Tehran for sequencing, and then their sequences were identified by S. Korea. The results are illustrated in Figures 5 - 7. *Crithidia* phylogenetic analysis based on GenBank (Figure 8).

**Figure 4.** Band Profiles Generated From Non-Consistent DNA Genome with *C. luciliae* and *C. fasciculata* Standards at 400 and 450 bp

![Band Profiles Generated From Non-Consistent DNA Genome](image)

| M, Molecular marker at 50 bp, Line 1, Non-consistent sample with *C. luciliae* and *C. fasciculata* standards, Line 2, Band profiles generated by DdeI enzyme, Line 3, Band profiles generated by ScrFI enzyme, Line 4, Band profiles generated by TaqI enzyme, Line 5, Negative control |

**Figure 5.** Obtained Results From Sequencing the Partial ITS1 Gene Products in Isolations Consistent With the Standard species *C. fasciculata* (Accession Number emb│Y00055.1 │)

TTTCTCTCCTCAACTCTCTCTTTTGTGGGGTTGTTGTGTGGGGGTTTGTGCGCGCGGTGCCGGAACAAGGCCAATCGATGCACGGTGATTGTATTTGTCTTTCTCT

*Crithidia fasciculata* strain in MHOM/IR/90/H.

**Figure 6.** Obtained Results From Sequencing the Partial ITS1 Gene Products in Isolations Consistent With the Standard Species *C. luciliae* (Accession Number emb│AJ627018.1 │)

TTTCTCTCTCCTCCCACTCTCTCTTTGTTGGGGTTGTTGTGTGGGGGTTTGTGCGCGCGGTGCAGGGAACAAAGGCCAATCGATGCACGGTGATTGTATTTGTCTTTCTTT

*Crithidia luciliae* internal transcribed spacer.

**Figure 7.** Obtained Results From Sequencing the Partial ITS1 Gene Products in Isolations From a Species Consistent With Trypanosomatidae With the Accession Number gb│GQ331988.1 │)

TTTCTCTCTCCTACACTCTCTCTCTTTTGTGGGGTTGTTGTGTGGGGGTTTGTGCGCGCGGTGCAGGGAACAAAGGCCAATCGATGCACGGTGATTGTATTTGTCTTTCTTT

Trypanosomatidae SP.66 GE-2009.
Figure 8. Crithidia Phylogenetic Analysis Based on Gen Bank, NCBI
5. Discussion

According to the strategies for controlling cutaneous leishmaniasis in the hyper-endemic regions of Iran and the World Health Organization providing the vaccine to prevent this disease has been proposed [16]. Therefore, one of the basic requirements is the determination of specifications for the species of Crithidia, which unfortunately infects the parasite leishmania cultures and has makes it impossible to prepare a purified antigen from the parasite (genus of leishmania) for preparing the vaccine [17].

Marmur et al. suggested that Crithidia might be a protozoan, which can live within some bacteria such as Bordetella bronchiseptica as endosymbiosis. Furthermore, these researchers found that in addition to the mentioned bacteria, this parasite can also live within B. culicis as a symbiosis. Marmur has proved his results by studying the rRNA ribosomal genes and SSURNA region, i.e. a part of ITS1 gene [4].

Wang et al. proposed that not only can Crithidia be isolated from bacteria such as B. bronchiseptica and B. culicis in an endosymbiotic manner, but also they can be isolated from a mosquito called Aedes vexans; particularly in C. oncophelti from an insect feeding from Oncopeltus fasciatus, Crithidia deanei is isolated from a predatory insect called Zelus leucogrammus, and Crithidia desouzaeii from a nectar-feeding fly called Orinidia obese. These researchers stated that a relationship between this protozoan and a large group of insects from the Depteran and reduviidae hemitran families has caused a great deal of interest about this protozoan [1].

The Crithidiad are able to endosymbiotically live within some parasites related to the trypanosomatid family, and sub-families of Leishmania, and move to different hosts along with them. These researchers believed that the only way to isolate Crithidia from Leishmania is through KDNA and/or rRNA genes. They suggested that by extraction of kinetoplast DNA and/or ribosomal DNA, then amplifying them with PCR, using the PCR-RFLP technique with HaellIII enzyme, and isolation of band at 350 bp for leishmania and at 450 bp for Crithidia these two parasites can be isolated from each other [18].

The results of the present study to some extent are in accordance with the results of the study by Yang et al. [19]. According to data, which illustrates the frequency distribution of isolated trypanosomatid family from culturing in Isfahan and surrounding areas, a large number of samples, i.e. 163 samples (81.09%), showed two bands of 400 and 450 bp after PCR. Using the Hea III enzyme, and after the random sequence of several samples, the results of the molecular analysis via the Software Blast showed 92% homology with C. fasciculata and 89% homology with C. luciae. Moreover, the sequence of the mentioned gene was in accordance with the World Gene Bank with the Accession number of GQ339888 which has been indicated by Doudi et al. [2].

Berzunza-Cruz et al. stated that a great heterogeneity between different strains of Leishmania and its family, i.e. trypanosomatidae, can be found by using PCR-RFLP technique, to which different species show various differences [10]. In this study, by using the molecular PCR-RFLP method, applying Hae III enzyme according to figure 1 and by the generated band profiles formed by the isolated genome we concluded that there was a difference between various species of Crithidia in terms of difference of sequence in the transcribed region of ITS-1. This was in a way that all the isolations following PCR and applying Haell enzyme that had revealed the band at 450 bp were similar to the C. fasciculata in terms of ITS-1 gene sequence, and all the isolations following PCR and applying Haell enzyme that had revealed the band at 400 bp were similar to C. luciae in terms of ITS1 gene sequence.

Yang et al. [19] proposed that similar to many parasites, studying the phylogenetic relationships of different species of Leishmania and Crithidia parasites is extremely complicated. This is because defining and categorizing the subspecies in order to describe them is very hard. This group of researchers collected 17 isolated Leishmania and 2 isolated Crithidia from different areas of China and amplified the ITS1 ribosomal gene using the PCR technique. They then sequenced a number of these isolations in terms of taxonomy, and identified the phylogenetic affinities between species of Leishmania and other parasites [19]. However, it should be mentioned that no difference was observed in analyzing the band profiles using the Ddel, ScrFI and TaqI enzymes on all the isolations, compared with the standard samples. Therefore, it seems that Hae III enzyme played a key role in the isolation of the Crithidia species in this study.

In general, according to the results, it can be concluded that Crithidia, which has been studied for the first time in terms of genetic polymorphism in Iran and Isfahan, is relatively polymorphic. This genetic polymorphism in this parasite showed a direct relationship with PCR-RFLP patterns. However, since Leishmania major is related to the hyper-endemic regions, further studies are required for other geographical regions of Iran and their relationship with these patterns. The results of the present study can be used by physicians, particularly dermatologists, for the appropriate treatment of Cutaneous Leishmaniasis, and can provide a new way for preparing a vaccine for cutaneous leishmaniasis.

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