First Human Cases of *Leishmania (Viannia) lainsoni* Infection and a Search for the Vector Sand Flies in Ecuador

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Abstract

An epidemiological study of leishmaniasis was performed in Amazonian areas of Ecuador since little information on the prevalent *Leishmania* and sand fly species responsible for the transmission is available. Of 33 clinical specimens from patients with cutaneous leishmaniasis (CL), causative parasites were identified in 25 samples based on cytochrome *b* gene analysis. As reported previously, *Leishmania (Viannia) guyanensis* and *L. (V.) braziliensis* were among the causative agents identified. In addition, *L. (V.) lainsoni*, for which infection is reported in Brazil, Bolivia, Peru, Suriname, and French Guiana, was identified in patients with CL from geographically separate areas in the Ecuadorian Amazon, corroborating the notion that *L. (V.) lainsoni* is widely distributed in South America. Sand flies were surveyed around the area where a patient with *L. (V.) lainsoni* was suspected to have been infected. However, natural infection of sand flies by *L. (V.) lainsoni* was not detected. Further extensive vector searches are necessary to define the transmission cycle of *L. (V.) lainsoni* in Ecuador.

Author Summary

In Ecuador, leishmaniasis is endemic in Pacific coast subtropical, Amazonian, and Andean highland areas. Seven *Leishmania* species, *Leishmania (Leishmania) mexicana*, *L. (L.) amazonensis*, *L. (L.) major-like*, *Leishmania (Viannia) guyanensis*, *L. (V.) panamensis*, *L. (V.) braziliensis*, and *L. (V.) naiffi*, are reported to be associated with human cutaneous (CL) and mucocutaneous leishmaniasis (MCL). Causative parasites have been studied extensively in Pacific coast and Andean areas; however, information such as prevalent
Leishmania species and their vector sand fly species is very sparse in Amazonian areas. Giemsa-stained smears taken from patients’ skin ulcers and used for routine diagnosis of CL and Flinders Technology Associates (FTA) card-spotted samples were utilized as DNA sources, and causative parasites were identified on the basis of cytochrome b gene analysis. Causative parasites in 25 samples were successfully identified, and, in addition to previously reported species, L. (V.) guyanensis and L. (V.) braziliensis, L. (V.) lainsoni was identified from two patients living in different areas situated more than 200 km apart. Sand flies were examined in areas where one of the L. (V.) lainsoni infected patient was suspected to have been infected. Although 1,104 female sand flies were dissected and examined for species identification and detection of natural infection with flagellates in the gut, human-infective Leishmania species including L. (V.) lainsoni were not detected. Further extensive investigation of sand fly fauna is necessary to incriminate the vector of this parasite in Ecuador.

Introduction

Leishmaniasis are caused by infection with protozoan parasites of the genus Leishmania transmitted by bites of female sand flies [1, 2]. The genus Leishmania is further divided into two subgenera, Leishmania (Leishmania) and Leishmania (Viannia), originally distinguished by their development in the digestive tract of sand fly vectors and later confirmed by phylogenetic studies. This group of diseases is distributed worldwide, especially in tropical and subtropical areas, affecting at least 12 million people in 98 countries [2]. Approximately 20 Leishmania species are known to be pathogenic to humans, and the infecting species is the major determinant of clinical outcome [2]. Therefore, identification of the parasite species in endemic areas is important for both appropriate treatment and prognosis.

In Ecuador, leishmaniasis is a major public health concern and is reported in 21 of 24 provinces of the country, in Pacific coast subtropical, Amazonian, and Andean highland areas [3]. Currently, seven Leishmania species, Leishmania (Leishmania) mexicana, L. (L.) amazonensis, L. (L.) major-like, Leishmania (Viannia) guyanensis, L. (V.) panamensis, L. (V.) braziliensis, and L. (V.) naiffi, have been identified as causative agents for human cutaneous (CL) and mucocutaneous leishmaniasis (MCL) [4, 5, 6]. In Pacific coast areas and Andean areas, causative parasite species have been studied extensively; L. (V.) guyanensis, L. (V.) panamensis, L. (V.) braziliensis, and L. (L.) amazonensis in Pacific areas, and L. (L.) mexicana as a dominant species and L. (L.) major-like as a minor species in Andean areas [4, 5, 7, 8, 9, 10, 11]. On the other hand, in Amazonian areas, although L. (V.) guyanensis, L. (V.) braziliensis, and recently, L. (V.) naiffi have been identified as causative agents for CL and MCL [4, 5, 6, 12, 13, 14], epidemiological studies on leishmaniasis have been very limited and consequently little information is available regarding prevalent parasite species as well as epidemiological situation.

Molecular biological methods are widely used for identification of Leishmania species using DNA extracted from clinical material of patients’ lesion, and is a powerful tool for epidemiological studies of leishmaniasis [15, 16, 17, 18]. DNA extracted from Giemsa-stained smears obtained from patients’ skin ulcers, which are used routinely for microscopic diagnosis in the laboratory to detect parasites in the lesion, have been used also as templates for detection and identification of Leishmania DNA although the sensitivity is low because of the limitation of the DNA source [19, 20, 21, 22]. Recently, to facilitate sample collection and DNA extraction processes, a Flinders Technology Associates (FTA) card (Whatman), a filter paper that readily lyses the spotted materials and fixes nucleic acids, was used for direct sampling of patients’
material in an epidemiological study of leishmaniasis, and its usability was reported [18, 23]. In
the present study, using smear slides and FTA card-spotted samples as DNA sources, an epidemi-
ological survey of leishmaniasis was performed in Amazonian areas where little information
on the endemic Leishmania species and vector sand flies is available.

Materials and Methods

Sample collection

Clinical samples were collected from patients suspected of having CL who visited health centers
in Cascales, Lago Agrio (Province of Sucumbíos), Coca, La Joya de los Sachs, and Nuevo
Rocafuerte (Province of Orellana) for the diagnosis and treatment of leishmaniasis (S1 Fig).
Tissue samples were taken by scraping the margins of active lesions on patients, spotting them
onto FTA Classic Cards (Whatman, Newton Center, MA) and storing them at room tempera-
ture. Two-mm-diameter disks containing the sample spot were punched out from each filter
paper and washed three times with FTA Purification Reagent (Whatman) and once with Tris-
EDTA buffer. The disks were air-dried and directly subjected to PCR amplification. For the
extraction of DNA from Giemsa-stained smears taken from patients’ skin ulcers that were used
for diagnosis of CL, 30 μl of DNA extraction buffer [150 mM NaCl, 10 mM Tris-HCl (pH 8.0),
10 mM EDTA and 0.1% sodium dodecyl sulfate (SDS)] containing 100 μg/ml of proteinase K
were spotted on each smear, and the tissue sample was collected into 1.5 ml microtube. The
sample was incubated at 37°C overnight, heated at 95°C for 5 min, and then 0.5 μl of each sam-
ple was directly used as a template for PCR amplification.

Sand fly collection

Sand flies were captured with mouth aspirator on protected human bait, CDC light traps, and
the modified Shannon light traps [24] between 18:30 and 21:00 for 14 nights on February 2015
at mountain and forest areas around patient houses in Sucumbíos Province where a patient
was suspected to have acquired an infection. Female sand flies were dissected and identified to
species level based mainly on the morphology of their spermathecae [25]. Sand flies were exam-
ined under light microscopy for natural flagellate infections and positive samples were fixed
individually in absolute ethanol. Ethanol-fixed specimens were individually lysed in 50 μl of
DNA extraction buffer with proteinase K, and 0.5 μl of the extract was directly used as PCR
templates [6, 26, 27, 28].

Identification of Leishmania species

Leishmania species were identified by cytochrome b (cyt b) gene sequence analysis [18, 23].
PCR amplification with a pair of specific primers, L.cyt-AS (5’-GGGGAGAGRRGAGGCGG-3’
and L.cyt-AR (5’-CCACCTCATAATATCTATA-3’), was performed with 30 cycles
of denaturation (95°C, 1 min), annealing (55°C, 1 min) and polymerization (72°C, 1 min)
using Ampdirect Plus reagent (Shimadzu Biotech, Tsukuba, Japan). A portion of the PCR
product was reamplified with L.cyt-S (5’-GGGTAGTTTATAGTYTAGG-3’) and L.cyt-R
(5’-CTACAATAAAAATCATATAATCAATTTAATTTAATTT-3’). For some samples, Leishmania species
were further identified by heat-shock protein 70 (hsp70) gene sequence analysis [29]. PCR
amplification with a pair of specific primers, HSP70sen (5’-GACGGTGCCTGCCTACTTCAA
3’) and HSP70ant (5’-CCGGGCCATGCTCTGTAATTACATTTGAC-3’), was performed with 40 cycles
of denaturation (95°C, 1 min), annealing (55°C, 1 min) and polymerization (72°C, 1 min) using
Ampdirect Plus reagent (Shimadzu Biotech, Tsukuba, Japan). The products were cloned into
the pGEM-T Easy Vector System (Promega, Madison, WI) and sequences were determined by
the dideoxy chain termination method using a BigDye Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, Foster City, CA).

Phylogenetic analysis

The *Leishmania cyt b* gene sequences were aligned with CLUSTAL W software [30] and examined using the program MEGA (Molecular Evolutionary Genetics Analysis) version 6 [31]. Phylogenetic trees were constructed by the neighbor-joining method with the distance algorithms available in the MEGA package [30]. Bootstrap values were determined with 1,000 replicates of the data sets. The database for phylogenetic analyses consisted of *cyt b* gene sequences from *L. (L.) infantum* (GenBank accession number: AB095958), *L. (L.) donovani* (AB095957), *L. (L.) major* (AB095961), *L. (L.) tropica* (AB095960), *L. (L.) amazonensis* (AB095964), *L. (L.) mexicana* (AB095963), *L. (V.) panamensis* (AB095968), *L. (V.) guyanensis* (AB095969), *L. (V.) braziliensis* (AB095966), *L. (V.) lainsoni* (AB433280), *L. (V.) naiffi* (AB433279) and *L. (V.) shawi* (AB433281).

Ethics statement

The collection was performed by local physicians and well-trained laboratory technicians with the approval of the research ethics committee of the Graduate School of Veterinary Medicine, Hokkaido University (license number: vet26-4). Informed consent was obtained from the adult subjects and from the children’s parents or guardians, prior to collection of diagnostic materials at each health center of the Ecuadorian Ministry of Health (Provinces of Sucumbios and Orellana). Signed consent was obtained after the explanation of the process of diagnosis and *Leishmania* species analysis at the time of routine diagnosis carried out at rural health centers, following the guidelines of the Ethics Committee of the Ministry of Health, Ecuador. The subjects studied were volunteers in routine diagnosing/screening and treatment programs promoted by the Ministry. All routine laboratory examinations were carried out free of charge, and treatment with specific drug (Glucantime) was also offered free of charge at each health center of the Ministry.

Results

Identification of *Leishmania* species from patient specimens

Of 33 clinical samples, nine samples from patients in Sucumbios Province and 24 from patients in Orellana Province were obtained (Table 1). Of these, 6 and 14 samples from Sucumbios and Orellana patients, respectively, were Giemsa-stained smears used for routine diagnosis of CL, and others (3 and 10 samples, respectively) which were specimens spotted on FTA cards (Table 1). Patients had one to four skin lesions on their face, arms or/and legs with a diameter of 1 to 3 cm, typical of those observed on patients with CL in the area. The leishmanial *cyt b* gene was successfully amplified from two smears and three FTA card samples from Sucumbios, and seven smear and nine FTA card samples from Orellana (Table 1), and sequences were determined. The *cyt b* gene sequences from two Sucumbios and 13 Orellana patients had a greater degree of homology with those of *L. (V.) guyanensis* (98.1–99.8%), and *cyt b* gene sequences from two Sucumbios and two Orellana patients had a greater degree of homology with those of *L. (V.) braziliensis* (98.6–100%). In addition, *cyt b* sequences of one specimen each from Sucumbios and Orellana patients who have never traveled abroad showed highest homology with those of *L. (V.) lainsoni* (99.0% and 99.1%) (S2 Fig). These results were supported by a phylogenetic analysis showing that parasites from 15 patients were located in the clade of *L. (V.) guyanensis*, four specimens were in the *L. (V.) braziliensis* clade, and two
samples were in the L. (V.) lainsoni clade (Fig 1), indicating that the causative parasite species were L. (V.) guyanensis, L. (V.) braziliensis, and L. (V.) lainsoni, respectively. A sample, 13-8EC7, from Sucumbios, in which causative parasite was identified as L. (V.) lainsoni, was further subjected to hsp70 gene sequence analysis. The hsp70 gene sequence of 13-8EC7 had a greater degree of homology with those of L. (V.) lainsoni (99.3%), and a phylogenetic analysis supported the result (S3 Fig). Nucleotide sequence data reported are available in the DDBJ/EMBL/GenBank databases under the accession numbers LC055616, LC055617, and LC055622-LC055638.

**Sand fly collection and dissection**

A total of 1,104 female sand flies were captured and dissected. Of these, 732 were captured on protected human bait, and 372 flies with light traps (CDC light trap and the modified Shannon
Leishmanial \textit{cyt b} genes were amplified and sequenced from patients with cutaneous leishmaniasis (13-8EC7–13-8EC15, 13AM1–13AM11, 15Ca1–15Ca6, and 15Or1–15Or7), and a phylogenetic analysis of \textit{cyt b} gene sequences was performed.
trap) (Table 2). Four species, *Lu. yuilli yuilli* (71.7%), *Lu. tortura* (12.3%), *Lu. davisi* (7.3%), and *Lu. napoensis* (5.0%) accounted for 96.3% of the sand flies (Table 2). Among these, *Lu. napoensis* was captured only by light traps. In addition to these four species, *Lu. sherlocki*, *Lu. trapidoi*, *Lu. gomezi*, *Lu. marinkelli*, *Lu. dysponeta*, *Lu. camposi*, *Lu. robusta*, *Lu. hirusta hirusta*, *Lu micropyga*, and five unidentified species were captured (Table 2). Natural flagellate infections were observed in the hindguts of 14 *Lu. yuilli yuilli* (1.8%), one *Lu. davisi* (1.2%), and of the only collected specimen of *Lu. camposi* (Table 2). Genomic DNAs were extracted from dissected sand flies infected with flagellates, and parasite *cyt b* genes were amplified. The *cyt b* gene fragments were successfully obtained from nine of the 14 positive *Lu. yuilli yuilli* and from *Lu. davisi*. The unsuccessful amplification of parasite *cyt b* genes from the other five *Lu. yuilli yuilli* and *Lu. camposi* was attributed to the very small number of parasites present in the gut. The *cyt b* gene sequences of parasites from the 10 flagellate-positive sand flies were analyzed and compared to those of related parasite species. The *cyt b* gene sequences from the nine *Lu. yuilli yuilli* showed only 86.6–88.6% homology with those of the *Leishmania* species, and 99.1–99.8% homology with those of *Endotrypanum* species, flagellate parasites of non-human animals transmitted by sand flies [32, 33], indicating that the flagellate infections in *Lu. yuilli yuilli* were *Endotrypanum*. On the other hand, the sequences of the parasite from *Lu. davisi* had only 81.2–82.3% homology with those of *Leishmania* and *Endotrypanum* species, indicating that the flagellate is neither *Leishmania* nor *Endotrypanum*. These results were supported by a phylogenetic analysis showing that nine flagellates from *Lu. yuilli yuilli* were located in the clade of *Endotrypanum* species, while the one from *Lu. davisi* was distant from *Leishmania* and *Endotrypanum* species (Fig 2). We suspect that the flagellates observed in *Lu. davisi* may belong to the genus *Trypanosoma* since some sand fly species are reported to transmit

Table 2. Identification of sand fly species and detection of flagellates within individual sand flies by microscopic examinations.

<table>
<thead>
<tr>
<th></th>
<th>Human bait</th>
<th>Light trap</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Lu. yuilli yuilli</em></td>
<td>540 (10*)</td>
<td>251 (4)</td>
</tr>
<tr>
<td><em>Lu. tortura</em></td>
<td>111</td>
<td>25</td>
</tr>
<tr>
<td><em>Lu. davisi</em></td>
<td>56</td>
<td>25 (1)</td>
</tr>
<tr>
<td><em>Lu. sherlocki</em></td>
<td>12</td>
<td>5</td>
</tr>
<tr>
<td><em>Lu. trapidoi</em></td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td><em>Lu. gomezi</em></td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>Lu. marinkelli</em></td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td><em>Lu. robusta</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>Lu. napoensis</em></td>
<td>0</td>
<td>55</td>
</tr>
<tr>
<td><em>Lu. dysponeta</em></td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td><em>Lu. camposi</em></td>
<td>0</td>
<td>1 (1)</td>
</tr>
<tr>
<td><em>Lu. hirusta hirusta</em></td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td><em>Lu micropyga</em></td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Others</td>
<td>4 [2 spp.]</td>
<td>3 [3 spp.]</td>
</tr>
<tr>
<td>Total</td>
<td>732</td>
<td>372</td>
</tr>
</tbody>
</table>

*flagellate-positive sand flies

Table 2. Identification of sand fly species and detection of flagellates within individual sand flies by microscopic examinations.

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Fig 2. Phylogenetic tree of cyt b gene sequences among species. Leishmanial cyt b genes were amplified and sequenced from flagellates-positive sand flies, *Lutzomyia (Lu.) yuilli* (1, 4, 7, 8, 9, 11, 12, 15, and 16) and *Lu. davisi*, and a phylogenetic analysis of cyt b gene sequences was performed by the neighbor-joining method together with sequences from 12 *Leishmania* and 2 *Endotrypanum* species. The scale bar represents 0.05% divergence. Bootstrap values are shown above or below branches.

doi:10.1371/journal.pntd.0004728.g002
Trypanosoma species [28, 34, 35]. The vector species of L. (V.) lainsoni in Ecuador could not be identified in this study.

Discussion

An epidemiological study of leishmaniasis was conducted in Amazonian areas of Ecuador where very little information is available on prevalent parasite species or sand fly species associated with their transmission. In addition to L. (V.) guyanensis and L. (V.) braziliensis infections, the first cases of CL caused by L. (V.) lainsoni infection in Ecuador were identified. A search for the vector sand fly in the area where one L. (V.) lainsoni infected patient presumably contracted the disease produced no positive flies, thus the vector remains unknown.

Leishmaniasis is widely distributed in Pacific coast subtropical climate areas, Andean highland areas, and Amazonian tropical areas in Ecuador [4, 5]. To date, most clinical and pathological studies of leishmaniasis have been reported from Pacific coast and Andean highland areas [4, 5, 7, 8, 9, 10, 11], and little information is available on endemic parasite and sand fly species in Amazonian areas, mainly because of difficulty in gaining access [4, 5, 6, 12, 13, 14]. In this study, Giemsa-stained smears used for routine diagnosis of CL were utilized as a DNA source in addition to FTA card-collected samples. The detection ratio was lower in DNA samples from smear slides when compared to FTA card collection (2/6 vs. 3/3 in Sucumbíos and 7/14 vs. 9/10 in Orellana), reflecting the amount of DNA recoverable and the condition of the smear slides. The thin smear samples were methanol-fixed, Giemsa-stained, and then examined under oil immersion. Thus, part of specimens and DNA may be lost and damaged when the immersion oil was wiped from the slide. Nevertheless, stored smear slides, which will not be used for any further purpose than diagnosis by microscopic examination, can be useful for identification of Leishmania species in endemic areas where sample collection for epidemiological study is difficult.

In this study, L. (V.) guyanensis and L. (V.) braziliensis were identified as causative agents in Amazonian areas as reported previously [4, 5, 6, 12, 13, 14]. In addition, L. (V.) lainsoni is implicated as a causative agent of CL in Ecuador for the first time. Leishmania (V.) lainsoni was originally identified from patients in Brazilian Amazon [36]. It causes CL with characteristic lesions similar to those caused by other Leishmania (Viannia) species: small ulcers or small self-limiting nodules [37]. The parasite was subsequently identified from patients in Sub-Andean and Amazonian areas of Peru [18, 38, 39, 40], in subtropical climate areas and Sub Andean areas of Bolivia [41, 42], Suriname [43], and French Giana [44]. Two L. (V.) lainsoni-infected patients were found during this study. One was from the northern part of Ecuadorian Amazon (Sucumbíos) near the border with Colombia, and the other was from the eastern frontier with Peru (Orellana), more than 200 km away. These findings suggest that L. (V.) lainsoni is widely distributed in South America.

Natural infections with L. (V.) lainsoni in sand flies have been detected in Lu. ubiquitalis in Brazil [45], Lu. nuneztovari anglesi in Bolivia [42], and Lu. avaraensis in Peru [46]. Microscopic examinations of sand flies collected in this study revealed no natural infection with L. (V.) lainsoni, and none of the above three Lutzomyia species were collected. Of the three dominant human-biting species in our collections, infection with Endotrypanum species, flagellate parasites of sloths, which are non-pathogenic to humans [32, 33], were detected in the most dominant species, Lu. yuilli yuilli, as reported in another Amazonian area of Ecuador [6]. Although natural infection was not detected in this study, the next dominant species, Lu. tortura, was already incriminated as a vector species of L. (V.) naifii in Ecuadorian Amazon [6, 13]. Furthermore, Lu. davisi was reported to transmit L. (V.) naifii in Brazil [47] and L. (V.) braziliensis in Peru [46]. In this study, natural infection of Lu. davisi by flagellates was observed. However,
the parasite was neither *Leishmania* nor *Endotrypanum*. To identify the vector of *L. (V.) lainsoni*, further research is necessary to understand the natural transmission cycle of this parasite in Ecuador. Our finding of the first human cases of CL caused by *L. (V.) lainsoni* infection in two separate areas of Ecuador suggests that this parasite is widely distributed in South America. Extensive countrywide surveillance is necessary to gain a proper understand of the status of *L. (V.) lainsoni*, as well as the sand flies responsible for its transmission in Ecuador.

**Supporting Information**

**S1 Fig.** Map of sample collection sites in Ecuador. 1. Cascales, 2. Lago Agrio (Province of Sucumbios), 3. Coca, 4. La Joya de los Sachas, and 5. Nuevo Rocafuerte (Province of Orellana). (TIF)

**S2 Fig.** Alignment of cytochrome b gene sequences of *Leishmania (Viannia) lainsoni*. Cytochrome b gene sequences obtained from Sucumbios (13-8EC7) and Orellana (13-8EC14) patients were aligned with that of *L. (V.) lainsoni* (AB433280). Black-shaded sequences represent identical nucleotides. (TIF)

**S3 Fig.** Phylogenetic tree of hsp70 gene sequences among species. Leishmanial hsp70 gene was amplified and sequenced from a patient with cutaneous leishmaniasis (13-8EC7), and a phylogenetic analysis of hsp70 gene sequences was performed by the neighbor-joining method together with sequences from 9 *Leishmania* species. The scale bar represents 0.005% divergence. Bootstrap values are shown above or below branches. The database for phylogenetic analyses consisted of hsp70 gene sequences from *L. (L.) tropica* (GenBank accession number: FN395026), *L. (L.) major* (XM_001684512), *L. (L.) donovani* (X52314), *L. (L.) infantum* (XM_001470287), *L. (L.) mexicana* (EU599091), *L. (V.) braziliensis* (XM_001566275), *L. (V.) guyanensis* (EU599093), *L. (V.) naiffi* (FN395056), and *L. (V.) lainsoni* (FN395047 and FN395048). (TIF)

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**Author Contributions**

Conceived and designed the experiments: HK YH. Performed the experiments: HK. Analyzed the data: HK. Contributed reagents/materials/analysis tools: HK AEB TM KH GFS SVG LNV AGG EAG YH. Wrote the paper: HK YH.

**References**


