Human Mixed Infections of Leishmania spp. and Leishmania-Trypanosoma cruzi in a Sub Andean Bolivian Area: Identification by Polymerase Chain Reaction/hybridization and Isoenzyme


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Parasites belonging to Leishmania braziliensis, Leishmania donovani, Leishmania mexicana complexes and Trypanosoma cruzi (clones 20 and 39) were searched in blood, lesions and strains collected from 28 patients with active cutaneous leishmaniasis and one patient with visceral leishmaniasis. PCR-hybridization with specific probes of Leishmania complexes (L. braziliensis, L. donovani and L. mexicana) and T. cruzi clones was applied to the different DNA samples. Over 29 patients, 8 (27.6%) presented a mixed infection Leishmania complex species, 17 (58.6%) a mixed infection Leishmania-T. cruzi, and 4 (13.8%) a multi Leishmania-T. cruzi infection. Several patients were infected by the two Bolivian major clones 20 and 39 of T. cruzi (44.8%). The L. braziliensis complex was more frequently detected in lesions than in blood and a reverse result was observed for L. mexicana complex. The polymerase chain reaction-hybridization design offers new arguments supporting the idea of an underestimated rate of visceral leishmaniasis in Bolivia. Parasites were isolated by culture from the blood of two patients and lesions of 10 patients. The UPGMA (unweighted pair-group method with arithmetic averages) dendrogram computed from Jaccard’s distances obtained from 11 isoenzyme loci data confirmed the presence of the three Leishmania complexes and undoubtedly identified human infections by L. (V.) braziliensis and L. (L.) chagasi and L. (L.) mexicana species. Additional evidence of parasite mixtures was visualized through mixed isoenzyme profiles. L. (V.) braziliensis-L. (L.) mexicana and Leishmania spp.-T. cruzi. The epidemiological profile in the studied area appeared more complex than currently known. This is the first report of parasitological evidence of Bolivian patients with trypanosomatidae multi infections and consequences on the diseases’ control and patient treatments are discussed.

Key words: leishmaniasis - Chagas disease - mixed infection - isoenzyme characterization - polymerase chain reaction

Among Latin American countries, Bolivia suffers from the highest infection rates of Chagas disease cases with a sero prevalence of 40% (Bryan & Tonn 1990), which represents a significant cause of morbidity and mortality (WHO 1991) for the country. More than 3 million people live in endemic regions and indicators of sero positivity infection reach 75% in several areas (Pless et al. 1992). The National Control Program considers Chagas disease in the Yungas valleys, a Sub-Andean region localized in the La Paz department, as non-endemic. However, vectors (Trypanosoma infestans) have been found in many houses (Le Pont et al. 1992, Brenière et al. 1995) and a recent epidemiological survey detected 12.7% seroprevalence for T. cruzi infection in the North Yungas province (unpublished data).

Leishmaniasis in Bolivia is less frequent than Chagas disease, however this disease affects people in five out of nine Bolivian departments. In the “Yungas” valleys cutaneous and mucocutaneous leishmaniasis constitute a serious public health issue, since consequences for the infected patients are severe mutilations, mostly located on their face (Le Pont et al. 1992). In this area, a prevalence of around 20% was reported based on observation of active lesions and scar without differences according to sex (Le Pont et al. 1992). In this focus cutaneous and mucocutaneous leishmaniasis are due to Leishmania (Viannia) braziliensis (Desjeux et al. 1974). In North and South Yungas provinces (corresponding to the studied area) 353 cases of cutaneous and mucocutaneous leishmaniasis have been registered in 2001 (report of the SNIS, Ministerio de Salud, Bolivia). Whereas parasites belonging to Leishmania (Leishmania) amazonensis were scarcely identified (Desjeux et al. 1987, Le Pont et al. 1992, Dedet 1993), a recent study had revealed a new restricted focus due to L. (L.) amazonensis (Martinez et al. 1998) in a Yungas valley characterized by a severe dry season. The strain of the first case of human visceral leishmaniasis was identified without a doubt by isoenzyme as Leishmania (L.) chagasi (Desjeux et al. 1986). Up to now, six autochthonous case of visceral leishmaniasis have been reported in Bolivia (Dimier-David et al. 1991) but the incidence of this infection and the morbidity are unknown.

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Recently, patient infection caused by Leishmania (V.) lainsoni has been described (Martinez et al. 2001, Bastrenta et al. 2002).

As in many region of South America, in the Yungas valleys there are overlapping endemic areas for Chagas disease and leishmaniasis. The occurrence of leishmaniasis patients carrying a double infection with T. cruzi has been shown by serological specific tests in various works and more scarcely by parasitological methods (Lemesre et al. 1986, Chiller et al. 1990, Chiaramonte et al. 1996). Moreover few cases of simultaneous infections with different Leishmania species have been reported by characterization of isolated stocks in the New World (Silveira et al. 1984, Oliveira Neto et al. 1985, Hernandez-Montes et al. 1998).

The purpose of this work is to identify patients carrying a double infection of Leishmania spp. and Leishmania-T. cruzi by a direct identification of parasites in blood and lesions of patients with active leishmaniasis. The different Leishmania species and the major Bolivian clones of T. cruzi were detected by the PCR amplification of the variable regions of the kinetoplast followed by the hybridization of the PCR products with specific probes of Leishmania complex (L. mexicana, L. donovani and L. braziliensis) and specific probes of T. cruzi clones 20 and 39 (numbering according to Tibayrenc et al. 1986). Mixed infections were also demonstrated by isoenzyme characterization of the parasites isolated from the patients. The epidemiological consequences due to this new data and further implications are discussed.

**PATIENTS AND METHODS**

**Origin of the studied patients** - Patients came from South (16°24’S, 67°31’W) and North (16°15’S, 67°40’W) Yungas in the department of La Paz, Bolivia. The Yungas are steep-sided valleys in the Eastern Andean Cordillera, at an altitude ranging from 1000-1800 m a.s.l. characterized by a subtropical climate. The 29 studied patients were received at the hospital for medical attention concerning typical lesions of leishmaniasis infections (28 cases) and visceral leishmaniasis in one two-year old patient. Moreover, mucosal lesions were suspected in two patients by the presence of nasal blockage and perforation. Leishmaniasis diagnosis was based on clinical observations, epidemiological history and positivity of Giemsa-stained smear of dermal scrapings (11 cases).

**Biological samples** - For the 29 patients, 5 ml of venous blood were collected, 1 ml of which was mixed immediately with an equal volume of 6 M guanidine HCl/0.2 M EDTA, pH 8 for further PCR processing. Some drops of blood were added in two cultures tubes of NNN biphasic medium supplemented with 1.5 ml of Schneider medium and serum was separated from the remaining 4 ml for further serologic tests.

Two fine-needle aspirates were taken from cutaneous ulcers with a 3 ml syringe containing 0.5 ml of sterile normal saline solution. One of the samples was conserved in nitrogen for PCR processing and the other one was distributed in two culture tubes.

A bone-marrow puncture was accomplished on the child with visceral leishmaniasis to process PCR and culture. An oral consent was obtained from the patients and parents for the biological sampling.

**Parasite isolation in culture** - At the laboratory, some drops of heated inactivated foetal calf serum were added to each culture tube and they were incubated at 32°C. Culture were examined daily until their adaptation. Subcultures were realized in Schneider’s medium supplemented with 20% heat-inactivated foetal serum. Promastigotes were grown at 25°C.

**Isoenzyme characterization** - The conditions for electrophoresis on cellulose acetate plates (Helena) were carried out according to Ben Abderrazak et al. (1993). A total of 10 enzyme systems (11 loci) were performed: glucose phosphate isomerase (GPI, EC 5.3.1.9), glucose-6-phosphate dehydrogenase (G6PD, EC 1.1.1.49), isocitrate dehydrogenase (IDH, EC 1.1.1.42), malate dehydrogenase NAD+ (MDH, EC 1.1.1.37), malate dehydrogenase NADP+ or malic enzyme (ME, EC 1.1.1.40), peptidase 1, substrate L-Leucyl-leucine-leucine (Pep-1, EC 3.4.11) phosphoglucomutase (PGM, EC 2.7.5.1), 6-phospho-gluconoate dehydrogenase (6PGD, EC 1.1.1.44), glutamate dehydrogenase NADP+ (GDH-NADP+, EC 1.4.1.4), glutamate dehydrogenase NAD+ (GDH-NAD+, EC 1.4.1.2) and glutamate oxaloacetate transaminase (GOT, EC 2.6.1.1).

The following reference stocks were used: L. (L.) mexicana (MNYC/BZ/M379), L. (L.) chagasi (MHOM/BR/74/PP75), L. (V.) braziliensis (MHOM/BO/84/LPZ595) and T. cruzi (clone 39: MN and clone 20: SO34 c14).

**Phenetic clustering of the isolates** - Phenetic relationships among the isolates and the reference stocks were inferred by computing Jaccard’s distance (Jaccard 1908). The UPGMA method, (unweighted pair-group method with arithmetic averages) was used for the construction of the dendrogram showing the relationship among the stocks (Sokal & Sneath 1963). The dendrogram was obtained using the NTSYS software (Exeter, Setauket, NY, USA).

**Serological techniques** - The detection by immunofluorescence of specific antibodies against L. (V.) braziliensis (MHOM/BR/75/S2903) was carried out according to Camargo (1966) using promastigotes forms fixed by 2.5% glutaraldehyde. Sera were assayed in serial two-fold dilutions from 1/30. The fluorescein isothiocyanate (FITC)-conjugate anti-human IgM (Sigma-Aldrich, St Louis, USA) diluted 1:50 (according to the manufacturer) in PBS containing 0.01% of Evans blue was used as second antibody.

Specific anti-T. cruzi IgG were detected by enzyme-linked immunosorbent assay (ELISA) using a crude antigen obtained from epimastigotes culture forms (Tehuantepec strain), according to previous studies (Breniere et al. 1985). The sera were diluted 1/200. The sheep peroxidase-conjugate anti-human IgG (H-L) (Biosys, Compiène, France) was used at a dilution of 1/2500.

**DNA extraction and PCR conditions** - The method used to treat the samples prior to PCR amplification has been previously described (Britto et al. 1993) and DNA extraction was performed as described by Wincker et al. (1994).
The *Leishmania* kDNA-PCR procedure was performed in duplicate in blood, lesions samples and parasite culture, according to Brenière et al. (1999) with primers L1: 5’-CCT ACC CAG AGG CCT GTC GGG-3’ and L2: 5’-TAA TAT AGT GGG CCG CGC AC-3’, purchased from Eurogentec laboratory (Seraing, Belgium).

PCR detection of *T. cruzi* in blood samples and parasite culture, was performed in duplicate according to Brenière et al. (1998), using the following primers: 121: 5’-AAA TAA TGT ACG G (T/G) GAG A TG CA T GA-3’ and 122: 5’-GGG TTC GAT TGG GGT TGG TGT-3’, purchased from Eurogentec laboratory.

Each run included positive and negative controls using parasite purified DNA and DNA free water as template respectively. PCR products were analysed by electrophoresis on 0.8% agarose gels in TAE X 0.5 and visualised by ethidium bromide staining.

**Probes** - Five kDNA reference probes were used. The three *Leishmania* complex-specific probes, *L. (L.) mexicana* (MNYC/BZ/M379), *L. (L.) chagasi* (MHOM/BR/74/PP75), *L. (V.) braziliensis* (MHOM/BO/90/CJ), were produced by PCR according to Brenière et al. (1999) using L1 and L2 primers, and *T. cruzi* specific probes of clone 39 and clone 20 were produced according to Brenière et al. (1998) using CV1 and CV2 primers (CV1: 5’ CAT TGG GGT TGG ACT AT 3’ and CV2: 5’ TTT AAC GGC CCT CCG AAA AC 3’) and DNA extracts of the reference stocks, TPKI (clone 39) and SO34 cl4 (clone 20). The amount of DNA was quantified by electrophoresis of sequential dilutions.

**Labelling and hybridization conditions** - After nylon membrane transfert of PCR products, labelling probes and hybridization were processed with the enhanced chemiluminescence gene detection system (ECL, Amersham, Buckinghamshire, UK) according to the manufacturer’s instructions. The membranes were washed twice under highly stringent conditions (6 M urea, 0.1 x SSC at 42ºC for 20 min) and then twice in 2 x SSC at room temperature for 10 min. Three exposures were performed (1, 5 and 30 min) on Hyperfilm-MP (Amersham, Buckinghamshire, UK).

**RESULTS**

The Table resumes the results obtained for each analysis: microscopically examination of lesion aspirates (Smear), origin of culture, isoenzyme typing, PCR/hybridization from isolated stains, blood and lesion aspirates and serology. The whole analysis confirmed *Leishmania* spp. infections for the 29 studied patients. Microscopically examination of lesion aspirates showed positivity in 11 cases (25 examined). Isoenzyme typing and PCR hybridization allowed the identification of the different *Leishmania* spp. and the detection of *T. cruzi* concurrent infections in several cases.

Species characterization of the strains - Twelve parasite strains were successfully isolated from the blood of two patients (6.9%), lesions of nine patients (31%) and one strain from the bone-marrow of the child diagnosed with visceral leishmaniasis. Four strains (Chu 02, Chu 04, Lp 06, Lp 07) shown complex patterns for several enzyme systems suggesting mixed infections (Table) and part of these patterns are illustrated in Fig. 1: 6PGD, (Lp 07, *L. (L.) mexicana* and *L. (V.) braziliensis*), PGM, (Lp 06, *L. (V.) braziliensis* and *T. cruzi* clone 39), ME-1, (Chu 02, *L. (L.) mexicana* and *T. cruzi* clone 20), and GDH-1, (Chu 04, *T. cruzi* clones 20 and 39). The dendrogram (Fig. 2) computed from MLEE Jaccard’s distance was established with the other eight strains, which did not show mixture profiles. The dendrogram clustered one strain isolated from a lesion (Lp 02) with *L. (L.) mexicana*, the strain isolated from the bone-marrow (Lp 17) with *L. (L.) chagasi*, and the six other strains isolated from lesions with *L. (V.) braziliensis*. PCR/hybridization applied to the strain cultures provided us confirmation of species and mixtures observed by isoenzymes analysis. Furthermore, PCR/hybridization evidenced the presence of *T. cruzi* in 10 of the 12 strains and one additional mixture of *L. (V.) braziliensis* and *L. (L.) mexicana* (Lp 16, Table).

The current analysis of the strains evidences the occurrence of the three *Leishmania* complexes (*L. donovani*, *L. braziliensis* and *L. mexicana*) in the studied area as well as co-infections with *T. cruzi*.

Detection of *Leishmania* spp. and *T. cruzi* from blood and aspirate samples by PCR-hybridization - PCR-hybridization was applied on blood and aspirate lesion samples of the 29 patients using the five specific probes to identify *Leishmania* complexes and major clones 20 and 39 of *T. cruzi*. A typical result for a series of PCR tests is shown in Fig. 3a. Using L1-L2 primers, multi-sized products ranging from 100 bp to 1300 bp and characteristic of *Leishmania* spp. were amplified and with 121-122 primers
<table>
<thead>
<tr>
<th>Patient code</th>
<th>Smear Origin</th>
<th>Strain with (L. (V.)) braziliensis, (L. (L.)) mexicana, (L. (L.)) chagasi</th>
<th>Lesions</th>
<th>Serological data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lp 26</td>
<td>nd Lesion</td>
<td>(L. (V.)) braziliensis</td>
<td>Lb</td>
<td>IF Leishmannia -</td>
</tr>
<tr>
<td>Lp 10</td>
<td>+ Lesion</td>
<td>(L. (V.)) braziliensis</td>
<td>Lb</td>
<td>ELISA T. cruzi -</td>
</tr>
<tr>
<td>Lp 15</td>
<td>+ Lesion</td>
<td>(L. (V.)) braziliensis + T. cruzi probes</td>
<td>Hyb. (+)</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 45</td>
<td>+ Lesion</td>
<td>(L. (V.)) braziliensis + T. cruzi probes</td>
<td>Hyb. (+)</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 46</td>
<td>+ Lesion</td>
<td>(L. (V.)) braziliensis + T. cruzi probes</td>
<td>Hyb. (+)</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 16</td>
<td>+ Lesion</td>
<td>(L. (V.)) braziliensis + T. cruzi probes</td>
<td>Hyb. (-)</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 17</td>
<td>+ Bone-marrow</td>
<td>(L. (L.)) chagasi</td>
<td>Md, Tc cl 20</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 02</td>
<td>+ Lesion</td>
<td>(L. (L.)) mexicana</td>
<td>Lm, Tc cl 20</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 06</td>
<td>+ Lesion</td>
<td>(L. (V.)) braziliensis + T. cruzi probes</td>
<td>Lm PCR (-)</td>
<td>Lb, Lm nd nd</td>
</tr>
<tr>
<td>Lp 07</td>
<td>+ Lesion</td>
<td>(L. (V.)) braziliensis + T. cruzi probes</td>
<td>Lm PCR (-)</td>
<td>Lb, Lm nd nd</td>
</tr>
<tr>
<td>Chu 04</td>
<td>- Blood</td>
<td>(L. (L.)) chagasi</td>
<td>Lm, Tc cl 20</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 02</td>
<td>- Blood</td>
<td>(L. (L.)) mexicana</td>
<td>Lm, Tc cl 20</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 33</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 16</td>
<td>nd / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 22</td>
<td>nd / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 23</td>
<td>+ / / /</td>
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<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 21</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 24</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 40</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 41</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 27</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 28</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 12</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 40</td>
<td>+ / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 37</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 45</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 08</td>
<td>- / / /</td>
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<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Lp 44</td>
<td>nd / / /</td>
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<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
<tr>
<td>Chu 21</td>
<td>- / / /</td>
<td>(L. (V.)) braziliensis - / / /</td>
<td>Lm other clone</td>
<td>PCR (-) Lb, Lm nd</td>
</tr>
</tbody>
</table>

\(/\): absence of data because the cultures were negative; Tc: \(Trypanosoma cruzi\); cl: clone; Lb, Lm and Ld: \(L. braziliensis\), \(L. donovani\), and \(L. mexicana\) complexes respectively; Hyb: hybridization; \(a\): the PCR was positive but hybridizations with clone 20 and clone 39 probes were negative; nd: not done.
a major 330 bp fragment specific of T. cruzi was obtained. The specificity of the five probes is shown in the right part of the figure. On the left part of the figure examples of hybridization patterns are shown evidencing, in a single patient, mixed infections by Leishmania spp. (lanes 8 and 9), by two Leishmania strains and T. cruzi clone 20 and 39 (lane 1, 2, 3; lane 6, 10, 13) by one Leishmania strain and T. cruzi clone 39 (lane 15, 16) and T. cruzi clone 20 (lane 7, 14) and by one Leishmania strain and T. cruzi clones 20 and 39 (lane 4, 5; lane 11, 12).

Surprisingly, in some cases any probe recognised PCR products amplified by L1-L2 primers from some blood and lesion samples since a multi banding pattern was obtained by PCR (Table). Further study of a large sample (117 samples) of normal blood shown that PCR products should be obtained in 10% of the cases but any hybridization with the three Leishmania complex species was observed (data not shown).

The Table summarizes the results of the PCR-hybridization. Leishmania spp. was identified in blood and/or lesions of the 29 studied patients confirming Leishmania infections. L. mexicana and L. braziliensis complexes species were detected in the majority of the patients (69% and 51.7% respectively). L. braziliensis was more frequently detected in lesion than in blood (41.3% and 10.3% respectively) while inverse result was observed for L. mexicana (48.3% in blood and 31% in lesions). L. donovani complex was detected in the blood and the lesion of the patient with visceral leishmaniasis (Lp 17) and in five others (20.7%), three cases in blood, one case in lesion, and one case in blood and lesion.

Among the 29 patients, eight (27.6%) showed a mixed infection with Leishmania spp., 17 (58.6%) a mixed infections with Leishmania and T. cruzi, and four (13.8%) a multi infections of Leishmania associate with T. cruzi infection (clones 20 and 39). The presence of T. cruzi was detected in 21 patients (72.4%) from blood and in two other patients from the strain isolated from the lesions (Lp 07, Lp 46). The percentage of Leishmania-T. cruzi infections together reaches 79.3% of the patients. The identification of T. cruzi clones confirms earlier work indicating the importance of major clones 20 and 39 in patients, however highest frequency was found for mixed infections by the two clones (52.4%). In two cases (Lp 12 and Chu 33) T. cruzi PCR products amplified from blood samples were not recognised by either probes clone 20 nor clone 39. These cases previously reported represent infections by other clones whose taxonomic status remains to be determined.

SERPOTHERAL ANALYSIS - IgM antibodies against promastigote forms of L. (V.) braziliensis (IF test) and IgG antibodies against total aqueous soluble protein extract of T. cruzi (ELISA) were detected together on 15 patient sera (Table). Seven sera (50%) were positive in both tests, which would confirm mixed infections of Leishmania and T. cruzi. Three sera were negative in both tests, four others were IF positive and ELISA negative and only one sera was IF negative and ELISA positive. PCR/hybridization applied on the same patient group, showed a mixed infection for 11 patients (73.3%). No relationship between antibody patterns and parasite species was observed.

**DISCUSSION**

Isolation and characterization of parasite isolated from patients remain the convincing method to demonstrate mixed infections of Leishmania-T. cruzi. In the present study we have prevailed a low percentage of parasite isolation in culture and the use of Schneider medium most
Fig 3a: ethidium bromide stained 1.5% agarose gel containing kDNA PCR products from *Trypanosoma cruzi* (330 bp) and *Leishmania* spp. (100 to 1300 bp). Lanes - 1-3: patient chu 24, lesion: primer L1-L2, blood: primer L1-L2, blood: primer 121-122; 4-5: patient chu 02, blood culture: primer L1-L2, primer 121-122; 6, 10, 13: patient Lp 16, blood: primer L1-L2, blood: primer 121-122, lesion: primer L1-L2; 7, 14: patient chu 28, blood: primer 121-122, blood: primer L1-L2; 8: patient Lp 40, blood: primer L1-L2; 9: patient chu 37, blood: primer L1-L2; 11-12: patient chu 16, blood: primer L1-L2 and 121-122; 15-16: patient Lp 21, blood primer L1-L2, blood primer 121-122; 18-23: reference strains, respectively: *T. cruzi*: TPK1 clone 39; *T. cruzi*: mixed of TPK1 clone 39 and SO34 cl4 clone 20; *L. (V) braziliensis* (MHOM/BO/84/LPZ595); *L. (L.) mexicana* (MNYC/BZ/M379); *L. (L.) chagasi* (MHOM/BR/74/PP75); *T. cruzi* : SO34 cl 4 clone 20; 17, 24: PCR low Ladder, 100 bp. Hybridizations with *L. (V) braziliensis* probe (MHOM/BO/84/LPZ595); *L. (L.) mexicana* probe (MNYC/BZ/M379); *L. (L.) chagasi* probe (MHOM/BR/74/PP75); *T. cruzi* : SO34 cl 4 clone 20; 17, 24: PCR low Ladder, 100 bp.
likely favoured the isolation of *Leishmania* over *T. cruzi*. Moreover, the percentage of positive cultures from primary skin lesions and mucosal tissues is generally very low due to contamination problems in cultures and the troubles to obtain adequate biopsies. The sampling of isolated strains is probably also biased through the selection that occurs during the culture step (Ibrahim et al. 1994, Bosseno et al. 2000). Even though the isoenzyme analysis remains the basic approach for species characterization, the current results of PCR/hybridization applied to parasite isolates increase the sensitivity, which enables to detect in a few steps multi species infections.

In the framework of the MLEE data, the profiles interrelated to mixed infection could be questioned. The two bands pattern should be equated to an heterozygous state for a monomeric enzyme in a diploid organism (hybrid) or to a double pattern for two different strains with homozygous states. Hybrids between *Leishmania* parasites have already been observed based on phenotypic and genotypic characteristics (Darce et al. 1991, Belli et al. 1994, Dujardin et al. 1995, Noyes et al. 1996, Bañuls et al. 1997) and similarly scarce events of hybridization in *T. cruzi* have been reported (Bogliolo et al. 1996, Carrasco et al. 1996, Brisse et al. 1998). Nevertheless, in both cases the frequency of genetic exchange in natural populations seems to be very low. In the current work, the high percentage of mixed infections of *Leishmania* sp. and *T. cruzi*—*Leishmania* sp. observed by PCR/hybridization support our interpretation. The PCR/hybridization confirmed the MLEE characterization of the strains but detected additional species. As expected, the PCR is more sensitive; experimental *T. cruzi* mixture of different clones showed that MLEE analysis revealed mixtures of at least 10% with only part of the enzyme systems (unpublished data) while Bosseno et al. (2000) reported that to detect mixture in culture, the PCR/hybridization is three times more sensitive than the isoenzyme characterization.

One strain (Lp 02) had a MLEE profile identical to the *L. (L.) mexicana* reference strain. Moreover, two diagnostic loci (*Nhi* and *Pgm*) between *L. (L.) amazonensis* and *L. (L.) mexicana* previously identified by Bañuls (1998) clearly characterized this isolate as belonging to *L. (L.) mexicana* (data not shown). This species had never been identified in Bolivia but it has been characterized in patients, dogs and vectors in an Andean region of Ecuador where cutaneous leishmaniasis is highly endemic (Hashiguchi et al. 1991). Nevertheless, further studies are necessary to confirm the presence of this species in the Andean part of Bolivia.

The direct identification of simultaneous infections using PCR/hybridization applied to blood and lesions surely presented a large advantage over culture typing due to the high sensitivity of the PCR. Nonetheless, processing large samples of blood and lesions with the primers L1 and L2, we questioned the specificity of these primers; PCR products were obtained from 10% of normal blood but their hybridization remain negative with the different *Leishmania* complex probes. Sequences homologies of L1 and L2 primers were then found in a human DNA bank and can explain this cross reactivity. Consequently, the hybridization step is indispensable for the detection and characterization of the *Leishmania* as done for all the studied samples. Several discrepancies of *Leishmania* species identification were noted in blood and lesion of the same patient. Furthermore, parasites belonging to *L. braziliensis* and *L. mexicana* complexes were unequally detected in lesions and bloods. This result could be explained by the higher ability of the current strains of *L. mexicana* complex to reach the circulation, given that the PCR sensibility evaluated by the addition of DNA from *L. (V.) braziliensis*, *L. (L.) mexicana* reference strains, was similar in blood and lesion but more sensitive to *L. braziliensis* complexes than to *L. mexicana* (0.1 pg and 10 pg respectively) (Mita 1999).

*Leishmania* spp. and *T. cruzi* exhibit antigenic cross-reactivity (Manson-Bahr 1987, Chiller et al. 1990) and the serological tests used for these diagnosis (IFA, IHA, ELISA) may lead to confuse epidemiological data and prevent specific diagnosis and treatment. The present data showed that the seven cases with positive serology for leishmaniasis and Chagas disease were actually infected by both parasites. All the same, among the other patients, which were infected with *Leishmania* spp., seven of them had undetectable IgM antibodies against *Leishmania*. Among the 15 patients with *T. cruzi* concurrent infection detected by PCR/hybridization in blood, seven of them showed a negative serology with *T. cruzi* ELISA. No data indicated that an infection with either parasites induced a protective immunity against the other (Chiller et al. 1990, Chiaramonte et al. 1996) but the present serological results demonstrated that a disorder of specific humoral response can be observed in mixed infections.

Owing to this approach the epidemiological facies in the Sub-Andean region appears much more complex than previously reported. In the Yungas valley, *L. (V.) braziliensis* was considered as the agent of cutaneous leishmaniasis (Desjeux et al. 1987). The PCR-hybridization showed the co-existence of three *Leishmania* species complex, and a separated distribution of *L. mexicana* and *L. braziliensis* complexes in Bolivia is no longer accurate. The patients infected by *L. mexicana* complex (69%) had been possibly infected in other endemic areas but a specimen of *Lutzomyia nuneztovari anglesi* (Diptera: Psychodidae) collected in the studied area and naturally infected by *L. (V.) braziliensis* (Torrez et al. 1998), was found positive by PCR-hybridization with both probes *L. (V.) braziliensis* and *L. (L.) mexicana* (data not shown). *L. nuneztovari anglesi* is a candidate vector of the cutaneous and muco cutaneous leishmaniasis in the Yungas valleys and a study in process in our laboratory using PCR/hybridization identification revealed that species of *L. braziliensis* and *L. mexicana* complexes share the same putative reservoir. Herein, visceral cases are not an exception, the declared case of a visceral leishmaniasis concerned a 2-year-old child and the other five infected patients were both children (5, 12 and 15 years old) and adults (29 and 52 years old) who were all healthy people. Obviously, the rate of visceral leishmaniasis is probably underestimated in Bolivia due to the difficulties in reaching correct diagnosis and the scarcity of available prevalence studies. The symptoms of visceral leishmaniasis can be clinically identifiable but some cases are asymptomatic or pociasymptomatic and the illness is
often indistinguishable from other infectious diseases including Chagas disease.

Concurrent infection with two species of *Leishmania* parasites occurs in 41% of the patients and this implies careful attention to the treatment. The clinical manifestations of *Leishmania* infection in humans are largely dependent on the immune responsiveness of the host and the virulence of the infecting parasite strain which explains differences in therapeutical responses (Melby et al. 1992, Grimaldi & Tesh 1993). In some cases of simultaneous natural infection, patients are reported to require distinct treatments (Al-Diwany et al. 1995) or to not respond to the treatment (Rassam et al. 1993).

Chagas disease is widespread in Bolivia and the Yungas valley are not spared. Nevertheless, this area is considered as a low endemic region in Bolivia although *T. cruzi* infected 79.3% of the current *Leishmania* patients. Part of the patients could have been infected in other high endemic areas of the country but this high rate of mixed infection could be also explained by higher susceptibility of chagasic patients to *Leishmania* infection. In such cases of mixed infections, sudden death as a result of chagasic cardiopathy during the pentavalent antimonial therapy and electrocardiograph changes during the course of the leishmaniasis treatment (meglumine antimoniate, sodium stibogluconate) have been observed (Chulay et al. 1985, Sampaio et al. 1988, Antezana et al. 1992). However, antimonial therapy effects in chronic chagasic cardiopathy are not well established. Therefore, chagas infection should be systematically searched for in leishmaniasis patients and antimonial therapy needs to be followed-up by an electrocardiogram.

Among the studied leishmaniasis patients, a high percentage (52.4%) of double infection with clones 20 and 39 was observed. Previous work showed, in Bolivia, similar frequencies of clones 20 and 39 and high rate of mixtures in vectors and during the acute phase of Chagas disease (Brenière et al. 1998) but, in later infection stages, higher prevalence of *T. cruzi* clone 39 over clone 20 in blood was observed with only 11.1% of double infections (clones 20 and 39) (Brenière et al. 1998). These results have been related to biological differences between the clones as virulence and variable susceptibility of the clones to the immune response (control of parasitemia) developed after the acute phase of human infection. By tricking the immune defences of the host, multi-species infections may increase the parasite’s chances of transmission and could play a specific role in the pathogenesis of the disease (Tibayrenc 1999). Obviously, the schedule of *Leishmania* and *T. cruzi* infections on the studied patients is unknown and the follow up of patients with leishmaniasis or Chagas disease living in overlapping endemic areas is necessary in order to reach a better understanding of the interactions between the two parasites when the second infection occurs.

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REFERENCES


