**Trypanosoma cruzi** Lineages Detected in Congenitally Infected Infants and *Triatoma infestans* from the Same Disease-Endemic Region under Entomologic Surveillance in Paraguay

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**Abstract.** *Trypanosoma cruzi* II is associated with Chagas disease in the southern part of South America. We analyzed *T. cruzi* variants in field-collected triatomines and congenitally infected infants living in the same disease-endemic region in Paraguay. Results of polymerase chain reactions for *T. cruzi* kinetoplast DNA and satellite DNA were positive in 83 triatomin feces samples and 58 infant blood samples. However, lineages were detected in 33 and 38 samples, respectively. *Trypanosoma cruzi* genotypes were determined in 56 (97%) blood samples after hybridization by using specific probes. The Tc I genotype was not detected. The prevalent sublineage was Tc IId in triatomines (27 of 33) and infant blood (36 of 58) as assessed by amplification of the 24Sα ribosomal RNA and the mini-exon region genes. The Tc Iic genotype was detected in 20 infant blood samples and in 1 triatomine. This study shows *T. cruzi* II is the predominant lineage circulating in triatomines and humans in endemic areas of eastern region of Paraguay.

**INTRODUCTION**

Chagas disease is caused by the protozoan parasite *Trypanosoma cruzi*. In the Southern Cone countries, *Triatoma infestans* is the main domiciliated vector for Chagas disease and over the last 10 years, Uruguay, Chile, Brazil and the Oriental Region of Paraguay (14 of 17 departments) have been certified as being free from disease transmission by *T. infestans*. The epidemiologic and sociocultural aspects encountered during disease management are 1) difficulty of eradication caused by the large number of animal reservoirs that perpetuate the presence of infectious sources; 2) the absence of available drugs that can be used on a large scale for treatment of patients in the chronic phase of the disease; 3) the lack of vaccines for protection of susceptible persons; and 4) the low social demand for medical attention because of chronicity of the infection. It is estimated that approximately 160,000 persons are infected with *T. cruzi* in Paraguay (population of approximately six million); seroprevalence among persons 15–45 years of age is 10–12% from-endemic regions. Extrapolating from this estimation, it is likely that 600 congenitally infected newborns are born per year, yielding a frequency of vertical transmission of 7%. *Trypanosoma cruzi* is genetically classified into two major evolutionary lineages, *T. cruzi* I and *T. cruzi* II, with a significant genetic distance between the two lineages. Linkage between molecular markers, such as the divergent domain of the 24Sα ribosomal RNA (rRNA) and the intergenic region of mini-exon genes, have been extensively analyzed, and clearly demonstrate the division of *T. cruzi* into two distinct phylogenetic lineages. The major lineage, *Tc* II is not homogenous and it can itself be divided into five sublineages on the basis of genotypic and phenotypic properties. According to clinical, epidemiologic, biological, biochemical and immunologic findings, it is assumed by some investigators that Chagas disease is caused by infection with *T. cruzi* II strains. However, isolation and classification of *T. cruzi* I from the myocardial tissue of a chronic chagasic patient with end-stage heart failure has been described. Epidemiologic studies suggest that *T. cruzi* IIB, IId, and IIE are related to anthropoponic environments and chronic Chagas disease patients, *T. cruzi* lineages IIa and Iic to sylvatic environments, and *T. cruzi* lineage I to both environments.

In this study, *T. cruzi* lineages were typed from blood samples obtained from congenitally infected infants identified through a previously described locally sustainable system of prenatal diagnosis and in *T. infestans* collected within the same disease-endemic region under entomologic surveillance. Genetic studies of *T. cruzi* are important to clarify the intraspecies heterogeneity of the parasite. Studies of host-parasite relationships are also crucial to understand the impact of the genetic diversity of the parasite on factors such as pathogenicity and virulence.

**MATERIALS AND METHODS**

**Epidemiology of Chagas disease in study areas.** Congenitally infected infants who were born to chronically infected mothers in an area free of domiciliary vector transmission were identified through a locally sustainable system of prenatal screening of *T. cruzi* infection implemented in 37 rural health care centers of two Chagas disease-endemic departments of the eastern region of Paraguay (Cordillera and Paraguari). The study area is free of domiciliary transmission of *T. infestans* because of integral vector control conducted during 1999–2000, and an established entomologic surveillance system based on community participation implemented since 2001. A description of vertical transmission and laboratory techniques used for analyses have been reported.

The human populations within these regions are stable, of low density, and low migration, and 87% of the population in these two areas were born there and lived there continuously since birth. The seroprevalence of *T. cruzi* infection in pregnant women in these areas range from 10% to 12%. The baseline infestation rates detected in 1999 and 2000 at the departmental level were 1.4% for Cordillera and 2.5% for Paraguari. A total of 1,921 triatomine samples were collected during 2001–2006 during the entomologic surveillance established to impede reinfestation in both areas (37 districts and approximately 95,000 dwellings). We determined that 69% of these triatomines were captured during the first two years of
the surveillance system (2001–2002) because of residual foci if we take into account the colonization index. These triatomi-nes were recorded and analyzed for T. cruzi infection in the Entomology Department of Servicio Nacional de Erradicación del Paludismo SENEPa in Asunción. Most (1,446 [75%] of 1,921) triatomines were T. infestans; the remaining 25% were autochthonous species such as T. sordida, T. guasayana, and T. guasu. With regard to the location of T. infestans species, 688 (48%) were captured from indoor living areas (domiciliary) and 739 (51%) were captured in the outdoor environment immediately surrounding living areas (peridomiciary).

Samples, DNA extraction, and polymerase chain reaction with species-specific primers. Seventy-six blood samples of T. cruzi-infected infants (age range = < 1–5 years) who were born to T. cruzi-infected mothers after 2001 were included in this study. These congenitally infected infants were confirmed as infected in follow-up studies during 2001–2006 by direct microscopic observation and polymerase chain reaction (PCR) or after 8 months of age by serologic techniques (enzyme-linked immunosorbent assay and immunofluorescent antibody assay). All infected children were treated with benznidazole (total daily dose of 5–7.5 mg/kg/day), which was given in two doses at regular intervals over a 60-day period.2

Of 1,921 triatomine specimens captured, 1,020 (53%) were kept at ~20°C at the Molecular Biology Laboratory at the Instituto de Investigaciones en Ciencias de la Salud, Universidad Nacional de Asunción for further analyses. Fecal samples of 320 T. infestans representing 145 infested households detected during 2001–2006 during integrated entomologic surveillance were analyzed; 191 were captured in domiciliary areas and 129 were captured in peridomestic areas (Table 1). Triatoma infestans samples that were subsequently analyzed by PCR were selected to represent 80% all capture sites. DNA extraction from blood samples was performed by using the QIAamp Blood Kit (Qiagen, Valencia, CA), and DNA from triatomines feces was extracted by using the phenol-chloroform method.17

The PCR assay used for detection of T. cruzi involved the amplification of a fragment of 330 basepairs from the variable T. cruzi mini-exon.6 DNA was detected in 58 (76%) of 76 blood samples from infected infants (age range = < 1–5 years) who were born to T. cruzi-infected mothers after 2001 were included in this study. These congenitally infected infants were confirmed as infected in follow-up studies during 2001–2006 by direct microscopic observation and polymerase chain reaction (PCR) or after 8 months of age by serologic techniques (enzyme-linked immunosorbent assay and immunofluorescent antibody assay). All infected children were treated with benznidazole (total daily dose of 5–7.5 mg/kg/day), which was given in two doses at regular intervals over a 60-day period.2

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The PCR assay used for detection of T. cruzi involved the amplification of a fragment of 330 basepairs from the variable regions of the kinetoplast minicircles DNA by using primers TCZ 1–TCZ 2.18–21 DNA extracted from cultured parasites was included as a positive control in PCRs, and a basal product was included as a negative control. All PCRs were performed in a PTC-100 Thermal Cycler (MJ Research Inc., Waltham, MA).

PCR-based identification of T. cruzi phylogenetic lineages. To discriminate between Tc I and Tc IIa–IIe lineages from blood samples positive for T. cruzi by PCR and triatomines feces, two parasite genomic sequences were amplified, the D7 domain of the 24Sα rRNA genes and the mini-exon region.3,22

The combination of both markers distinguishes the lineages and sublineages. The expected PCR products for sublineages are TcIIa: 120, 125, or 130 basepairs for the rRNA genes and no product for mini-exon or a product of 300 basepairs observed with low intensity in combination with 120 basepairs for rRNA; TcIIb: 125 basepairs for rRNA and 300 basepairs for mini-exon; TcIIc: 110 basepairs for rRNA and no product for mini-exon or a product of 300 basepairs observed with low intensity; TcIIId: 110 basepairs for rRNA and 300 basepairs for mini-exon; a low-intensity product of 125 basepairs can be also observed; and TcIe: 125 basepairs for rRNA and 300 basepairs for mini-exon.8

The PCR amplification of the 24Sα rRNA genes was performed under the following cycling conditions: 4 minutes at 94°C; 35 cycles for 1 minute at 94°C, 1 minute at 55°C, and 1 minute at 72°C; and a final extension step for 5 minutes at 72°C. Primers TC-1 (T. cruzi I), TC-2 (T. cruzi II), and TC (common to T. cruzi I and II) were used to amplify the mini-exon gene. The following thermal profile was used: 1 minute at 94°C; 35 cycles for 30 seconds at 94°C, 30 seconds at 65°C, and 30 seconds at 72°C; and a final denaturation step for 10 minutes at 72°C. Amplification products were 300 basepairs (T. cruzi II) and 350 basepairs (T. cruzi I).3 The PCR products were separated by electrophoresis on 2% agarose gels, stained with ethidium bromide, and visualized under ultraviolet light. This system did not differentiate between Tc IIb and Tc Ie sublineages.6

Hybridization. Blood samples that were species-specific positive by PCR, but which had no detectable lineage-specific PCR products by agarose gel electrophoresis, were subjected to hybridization. The PCR amplification products of DNA extracted from these samples, using the 24Sα rRNA gene and the mini-exon gene as targets, were subjected to 2% agarose gel electrophoresis and transferred to a nylon Hybond-N+ membrane (Amersham Biosciences, Little Chalfont, United Kingdom). Probes were prepared with the PCR amplification products of the 24Sα rRNA and the mini-exon genes from reference strains (MN cl2, cl9 for T. cruzi II and X10 cl11, cl20 for T. cruzi I).6 The Enhanced Chemiluminescence Direct Nucleic Acid Labeling and Detection System Kit (Amersham Biosciences) was used for detection.

RESULTS

PCR with species-specific primers. Trypanosoma cruzi DNA was detected in 58 (76%) of 76 blood samples from

### Table 1

| Distribution by habitat of Triatoma infestans positive for Trypanosoma cruzi |
|-------------------------------|-----------------|-----------------|-----------------|-----------------|
| Habitat                      | Total samples   | T. cruzi positive | Total no identified lineages | Cordillera | Paraguari | Total |
| Domicile                     | 191             | 67 (35)          | 7                          | 23            | 30         |
| Peri-domicile                | 129             | 16 (12)          | 0                          | 3             | 3          |
| Total                        | 320             | 83 (26)          | 7                          | 26            | 33         |

* T. cruzi-positive samples identified by using species-specific primers TCZ 1–TCZ 2 and 121–122.

† Lineage-positive samples identified by using lineage-specific primers for mini-exon and 24Sα ribosomal RNA genes.

### Table 2

Distribution by age of 76 infants congenitally infected with Trypanosoma cruzi analyzed by PCR with species-specific primers

<table>
<thead>
<tr>
<th>Age, months</th>
<th>Total</th>
<th>No (%) T. cruzi positive by PCR</th>
<th>Intensity of PCR products by primers TCZ 1–TCZ 2</th>
<th>No PCR positive by primers TCZ 1–TCZ 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>0–12</td>
<td>45</td>
<td>37 (82)</td>
<td>0</td>
<td>17</td>
</tr>
<tr>
<td>13–24</td>
<td>14</td>
<td>10 (71)</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>25–36</td>
<td>6</td>
<td>4 (67)</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>37–48</td>
<td>6</td>
<td>4 (67)</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>49–60</td>
<td>5</td>
<td>3 (60)</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>Total</td>
<td>76</td>
<td>58 (100)</td>
<td>1</td>
<td>28</td>
</tr>
</tbody>
</table>

* T. cruzi-positive samples identified by using species-specific primers TCZ 1–TCZ 2 and 121–122.

† Intensity of PCR products observed by 2% agarose gel electrophoresis and staining with ethidium bromide.

‡ T. cruzi DNA was detected in these samples with primers 121–122; they were not detectable with primers TCZ 1–TCZ 2 primers.
congenitally infected infants by using primers TCZ 1–TCZ 2 and 121–122. The sensitivity of PCR in detecting infected infants was higher in children less than two years of age old (71–82%), and four infants positive by primers TCZ 1–TCZ 2 were also positive by primers 121–122. The intensity of the PCR products observed in a 2% agarose gel, and species-specific primers observed in a 2% agarose gel, and species-specific primers TCZ 1–TCZ 2 was evaluated as weak, average, and strong (Table 2). This variation was used to compare results of lineage and sublineage detection in the same samples. A total of 98% of blood samples had an average and strong intensity of T. cruzi PCR products with the species-specific primers TCZ 1–TCZ 2.

Amplified products of T. cruzi DNA were detected in 83 (26%) of 320 T. infestans fecal specimens sampled from 80% of the capture sites. Detection of T. cruzi DNA was more frequent in the fecal specimens collected from domiciliary T. infestans than from peridomiciliary captures (67 [35%] of 191 versus 16 [12%] of 129, respectively; Table 1). This procedure was conducted to PCR sensitivity and specificity. Twenty-four samples were hybridized with the mini-exon probe and 20 blood samples had no amplification product for the mini-exon gene. Eight of 24 blood samples subjected to hybridization with the 24Sα rRNA this probe were positive (Table 4). The sizes of the 24 Sα rRNA amplification products with the primers D71-D72 are 110 basepairs and 125 basepairs of the 24Sα rRNA gene (Table 4 and Figures 1 and 2).

Hybridization of PCR products to increase sublineage detection in infants blood samples. All samples that were T. cruzi-negative with lineage-specific primers were subjected to hybridization with probes prepared from amplified products of the 24Sα rRNA gene and the mini-exon gene by using reference strains (MN cl2, clon 30 for T. cruzi II and X10 e11, clon 20 for T. cruzi I). This procedure was conducted to PCR sensitivity and specificity. Twenty-four samples were hybridized with the mini-exon probe and 20 blood samples had no amplification product for the mini-exon gene. Eight of 24 blood samples subjected to hybridization with the 24Sα rRNA this probe were positive (Table 4).

The distribution by region and habitat of T. infestans in comparison with T. cruzi is shown in Table 1. The PCR sensitivity was low for T. cruzi lineage determination when DNA extracted from the feces of triatomines was used. Only 33 (40%) of 83 T. cruzi-positive samples could be amplified by the lineage-specific primers, and 30 of 33 lineages detected were from domicile habitats. Hybridization was not performed for triatomine samples.

Trypanosoma cruzi genotypes were determined for 56 (97%) of 58 blood samples from PCR-positive infants (Table 3). Of these 56 blood samples, 38 (68%) were directly detected by agarose gel electrophoresis and 18 were detected after hybridization (all PCR products were 110 basepairs for the 24Sα rRNA gene) (Table 4 and Figures 1 and 2).

Distribution of T. cruzi II subgroups detected in infant blood samples and triatomine feces from Cordillera and Paraguari. Lineage T. cruzi I was not detected in either area, and the PCR product of 350 basepairs was not observed in any of the biological samples when the mini-exon gene was used. Sublineages Tc Iic and Tc IId were found mainly in the 58 infected infants (97%). The proportion of the sublineages detected in blood samples from infants residing in Cordillera and Paraguari areas were 39% (14 of 36) and 27% (6 of 22) for Tc Iic and 58% (21 of 36) and 68% (15 of 22) for Tc IId, respectively. The main sublineage detected among T. infestans captured in both departments was Tc IId; all seven (100%) triatomines from Cordillera and 20 (76%) of 26 from Paraguari (Table 3).

**DISCUSSION**

We describe T. cruzi lineages and sublineages circulating in congenitally infected infants and triatomines collected within the same endemic region in Paraguay under entomologic surveillance during 2001–2006. Amplification of T. cruzi DNA

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**Table 3**

Distribution of Trypanosoma cruzi II subgroups detected in infant blood samples and triatomine feces from Cordillera and Paraguari, Paraguay

<table>
<thead>
<tr>
<th>Department</th>
<th>T. cruzi lineages in infant blood samples</th>
<th>T. cruzi lineages in Triatoma infestans</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T. cruzi lineages</td>
<td>T. cruzi lineages</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>Iic</td>
</tr>
<tr>
<td>Cordillera</td>
<td>36</td>
<td>14</td>
</tr>
<tr>
<td>Paraguari</td>
<td>22</td>
<td>6</td>
</tr>
<tr>
<td>Total</td>
<td>58</td>
<td>20</td>
</tr>
</tbody>
</table>

ND = not described. Atypical pattern not described previously.

* Sample with a PCR amplification product of 300 basepairs with mini-exon primers and no amplification with 24Sα ribosomal RNA primers.

† Sample with PCR amplification products of 300 basepairs and 350 basepairs with mini-exon primers and 24Sα ribosomal RNA primers respectively.

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**Table 4**

Hybridization and PCR results in infected blood

<table>
<thead>
<tr>
<th>PCR Intensity of TCZ amplicons</th>
<th>Total PCR positive</th>
<th>24Sα rRNA Intensity of mini-exon amplicons</th>
<th>Hybridization of Trypanosoma cruzi PCR mini-exon and 24Sα rRNA probes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. (%) positive</td>
<td>Strong</td>
<td>Average</td>
</tr>
<tr>
<td>Weak</td>
<td>1</td>
<td>1 (100)</td>
<td>0</td>
</tr>
<tr>
<td>Average</td>
<td>28</td>
<td>21 (75)</td>
<td>1</td>
</tr>
<tr>
<td>Strong</td>
<td>25</td>
<td>12 (48)</td>
<td>4</td>
</tr>
<tr>
<td>TCZ/121–1228</td>
<td>4</td>
<td>4 (100)</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>58</td>
<td>38 (65)</td>
<td>5</td>
</tr>
</tbody>
</table>

Hybridization by primers TCZ1–TCZ2 was not detectable and hybridization by primers 121–122 was detectable.
directly from biological samples without previous in vitro isolation by culture enabled us to make inferences regarding lineages and sublineages circulating in the human host and in vectors within the same geographic region. The stability of the population in the region studied virtually eliminated migration as a confounding factor in our investigation.

Because typing lineages with genetic markers directly from human blood samples of chronically ill patients is difficult because of the low levels of parasites present in blood, lineages circulating in congenitally infected infants were selected for this study. Trypanosoma cruzi DNA was detected in 58 (76%) of 76 blood samples from infected infants. The PCR sensitivity of primers TCZ 1–TCZ 2 was better than with that of primers 121–122, although T. cruzi DNA detection was not successful in four blood samples with the primers TCZ 1–TCZ 2. Such results might be related to the small amount of blood samples obtained from infants (1–3 mL) and an incomplete dispersion of the kinetoplast minicircle present in the blood samples previously boiled but not treated with guanidine-HCl.

The PCR sensitivity of detection of T. cruzi DNA was higher for blood samples infants less than two years of age. However, when PCR was performed using primers specific for the 24Sα rRNA and mini-exon genes, the sensitivity was substantially lower; only 38 (66%) of the 58 T. cruzi PCR-positive samples were detected by visualization after agarose gel electrophoresis. When hybridization with the lineage-specific primers for 24Sα rRNA and mini-exon genes was used, the sensitivity increased to 97% (56 of 58).

Results of PCR-based assays for detection of T. cruzi DNA using primers 121–122 and TCZ 1–TCZ 2 primers were positive for 83 triatomines feces, and no differences in sensitivity were observed between primer sets. However, further amplification with lineage-specific primers resulted in the successful detection of sublineages in only 33 (40%) of the 83 samples.

Thirty-eight samples were positive by PCR with the 24Sα rRNA primers, and no difference was observed in the sensitivity of these reactions in relation to the amount of parasites present in blood samples (assessed by the intensity of the TCZ 1–TCZ 2 PCR products). Nevertheless, the mini-exon primers seemed to be more dependent on the amount of parasites present; a 72% (18 of 25) detection level was observed in those samples with strong TCZ 1–TCZ 2 PCR-products. The Tc Iic genotypes detected in 20 blood samples were identified after hybridization with the mini-exon probe (when no PCR products were detected).

Sublineages Tc Iic and Tc Iid were the prevalent genotypes among 58 PCR-positive congenitally infected infants, with a distribution of 20 (35%) of 58 for Tc Iic and 36 (62%) of 58 for Tc Iid, respectively. Other genotypes had atypical patterns not described previously and are under study. In addition, Tc IId was the prevalent genotype among 27 of 33 PCR-positive infected triatomines.

On the basis of our findings, it can be inferred that sublineages Tc Iic and Tc Iid are related to cases of chronic disease in the study area. These sublineages were present in 97% of congenitally infected infants, and it can be inferred the same lineages infected their mothers. Five samples, three in triatomines and two in human blood, showed PCR-product patterns that have not been described. These variants are under further study to determine their significance.

In 2008, both areas were declared free of domiciliary vector transmission by international experts and intergovernmental resolutions (official documents of the Pan American Health Organization and the World Health Organization). Our study indicates that we detected T. cruzi in T. infestans during the entomologic surveillance system established in 2001 that were focal/residual triatomines that remained after integral vector control with insecticides performed in 1999–2000. This study identified T. cruzi II strains as the predominant lineages circulating in the main endemic areas of the eastern region of Paraguay, as detected in field-collected triatomines and humans living in the same region. Trypanosoma cruzi I was not found in these regions and does not appear to play a significant role in the epidemiology of Chagas disease in Paraguay.
REFERENCES


