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# Genetic variability of *Trypanosoma cruzi* TcI isolates from rural and urban areas of Venezuela

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#### **ABSTRACT**

Background & objectives: Several studies have demonstrated genetic heterogeneity in populations of *Trypanosoma cruzi* that allowed the identification of six different discrete typing units (DTU) classified as TcI, TcII, TcIII, TcIV, TcV and TcVI. Furthermore, some characterization studies have described genetic variability within TcI isolates from endemic regions. The objective of the present study was to analyze Venezuelan *T. cruzi* isolates, obtained from triatomine-vectors, mammal-hosts including infected humans, detected in both rural and urban areas from diverse geographic origins.

*Methods:* Molecular characterization of 44 Venezuelan *T. cruzi* isolates, obtained from triatomine-vectors, mammalian hosts and human patients from both rural and urban areas of different geographic origins, were carried out. Samples were analyzed by PCR amplification of the intergenic region of the mini-exon gene,  $24S\alpha$  rDNA and 18S rDNA, followed by sequencing of the amplification products.

Results: The TcI amplification pattern was found in 42 out of 44 (95.5%) isolates; a TcIII strain and one possible TcIV were also found. The sequence analysis of the TcI Venezuelan isolates showed genetic variability among them. Urban isolates formed a homogeneous group, with differences in their sequences, when compared to rural isolates.

Interpretation & conclusion: The results showed genetic heterogeneity in Venezuelan TcI strains, probably in response to different environmental conditions.

Key words Genetic variability; mini-exon; TcI; Trypanosoma cruzi; Venezuela

# INTRODUCTION

Trypanosoma cruzi, the etiological agent of American Trypanosomiasis or Chagas disease, affects about 10 million people and 25 million are at risk in Latin America<sup>1</sup>. The human pathology includes an acute phase, followed by the chronic phase with an unpredictable clinical course, ranging from no symptom to a severe disease with cardiovascular compromise and/or digestive alterations that could cause death<sup>2</sup>. In Venezuela, number of studies suggest an active transmission and re-emergence of the disease<sup>3-4</sup>.

Several investigations based on biochemical and genetic markers showed that *T. cruzi* strains are highly polymorphic and consist of a variety of parasite subpopulations, with biological, biochemical, immunological and genetic heterogeneity observed in their triatominevectors, reservoir hosts and people living in risk areas where the kinetoplastid is endemic<sup>5</sup>.

*T. cruzi* populations have been classified into six discrete taxonomic units (DTUs), named as TcI, TcII, TcIII, TcIV, TcV and TcVI based on different molecular markers and biological features<sup>6-8</sup>. Although, *T. cruzi* I was considered a homogeneous DTU, genetic variability within *T. cruzi* I has been reported in recent years<sup>9-11</sup>.

Several authors associated the parasite variability with differences in the biological cycle, tissue invasion, virulence, clinical profiles, geographic distribution, *etc*. The molecular epidemiology based on the genetic typing of *T. cruzi* isolates from different sources may be useful to understand the variability of this parasite and its possible relationship to the clinical and epidemiological characteristics of the disease<sup>12-13</sup>.

In Venezuela, some studies revealed TcI, TcIII and TcIV in human beings, triatomine bugs, and other mammals<sup>14-17</sup>. In the present study, 44 Venezuelan *T. cruzi* isolates, obtained from triatomine-vectors, mammalian

hosts including infected humans, detected in both rural and urban areas from diverse geographic origins, were analyzed.

## **MATERIAL & METHODS**

## Parasite isolates

A panel of 44 Venezuelan *T. cruzi* isolates, from rural and urban areas, including both domestic and peri-domes-

tic transmission cycles, were studied (Table 1). Caracas City, the states of Cojedes and Guárico are in the central region of Venezuela, whereas Anzoátegui state is in the northeastern part. Five *T. cruzi* isolates, previously characterized, were also included in the analysis, three TcI and two TcV. Parasites were cultured in liver infusion tryptose (LIT) liquid medium and harvested by centrifugation; parasite pellets were stored at  $-70^{\circ}$ C until use. The kinetoplastids were obtained from mammal blood and vec-

Table 1. Molecular characterization of the Trypanosoma cruzi isolates by PCR

S.No.	Isolates	Host	Locality	Habitat	DTU	
1.	MDID/VE/1984/Dm28c	Didelphis marsupialis	Guárico	Rural	TcIa	
2.	MHOM/VE/2007/EP	Homo sapiens	Guárico	Rural	TcIa	
3.	MHOM/VE/2007/EP6c	Homo sapiens	Guárico	Rural	TcIa	
4.	MHOM/PA/2007/LH31	Homo sapiens	Paraguay	Rural	TcV <sup>a</sup>	
5.	MHOM/PA/2007/LH32	Homo sapiens	Paraguay	Rural	TcV <sup>a</sup>	
6.	TMAC/VE/2007/LH1	Triatoma maculata	Anzoátegui	Rural	TcI	
7.	TMAC/VE/2007/LH4	Triatoma maculata	Anzoátegui	Rural	TcI	
8.	TMAC/VE/2007/LH5	Triatoma maculata	Anzoátegui	Rural	TcI	
9.	TMAC/VE/2007/LH6	Triatoma maculata	Anzoátegui	Rural	TcI	
10.	TMAC/VE/2007/LH12	Triatoma maculata	Anzoátegui	Rural	TcI	
11.	TMAC/VE/2007/LH13	Triatoma maculata	Anzoátegui	Rural	TcI	
12.	TMAC/VE/2007/LH19	Triatoma maculata	Anzoátegui	Rural	TcI	
13.	TMAC/VE/2007/LH20	Triatoma maculata	Anzoátegui	Rural	TcI	
14.	TMAC/VE/2007/LH23	Triatoma maculata	Anzoátegui	Rural	TcI	
15.	TMAC/VE/2007/LH26	Triatoma maculata	Anzoátegui	Rural	TcI	
16.	TPRX/VE/2007/LH2	Rhodnius prolixus	Anzoátegui	Rural	TcI	
17.	TPRX/VE/2007/LH3	Rhodnius prolixus	Anzoátegui	Rural	TcI	
18.	TPRX/VE/2007/LH10	Rhodnius prolixus	Anzoátegui	Rural	TcI	
19.	TPRX/VE/2007/LH18	Rhodnius prolixus	Anzoátegui	Rural	TcI	
20.	TPRX/VE/2007/LH21	Rhodnius prolixus	Anzoátegui	Rural	TcI	
21.	TPRX/VE/2007/LH22	Rhodnius prolixus	Anzoátegui	Rural	TcI	
22.	TPRX/VE/2007/LH25	Rhodnius prolixus	Anzoátegui	Rural	TeI	
23.	TPRX/VE/2007/LH27	Rhodnius prolixus	Anzoátegui	Rural	TcI	
24.	TPRX/VE/2007/LH28	Rhodnius prolixus	Anzoategui	Rural	TcI	
2 <del>5</del> .	MDID/VE/2007/LH7	Didelphis marsupialis	Anzoategui	Rural	TcI	
26.	MDID/VE/2007/LH9	Didelphis marsupialis	Anzoategui	Rural	TcI	
20. 27.	MDID/VE/2007/LH24	Didelphis marsupialis	Anzoategui	Rural	TcI	
27. 28.	MDID/VE/2007/LH24 MDID/VE/2007/LH14	Didelphis marsupialis	Anzoategui	Urban	TcI	
26. 29.	MDES/VE/2007/LH33	Desmodus sp	Anzoategui	Rural	TcI	
29. 30.	MDES/VE/2007/LH43	Desmodus sp	Anzoátegui	Rural	TcI	
30. 31.	MCAN/VE/2007/LH43				TcIV <sup>b</sup>	
31. 32.	TGEN/VE/2007/LH35	Canis familiaris	Anzoátegui	Rural Urban	TcI	
		Panstrongylus geniculatus	Caracas			
33.	TGEN/VE/2007/LH36	Panstrongylus geniculatus	Caracas	Urban	TcI	
34.	MRAT/VE/2007/LH30	Rattus rattus	Caracas	Urban	TcI	
35.	MRAT/VE/2007/LH34	Rattus rattus	Caracas	Urban	TcI	
36.	MHOM/VE/2007/LH37	Homo sapiens	Caracas	Urban	TcI	
37.	MHOM/VE/2007/LH42	Homo sapiens	Caracas	Urban	TcI	
38.	MHOM/VE/2007/LH46	Homo sapiens	Caracas	Urban	TcI	
39.	MHOM/VE/2007/LH47	Homo sapiens	Caracas	Urban	TcI	
40.	MHOM/VE/2007/LH48	Homo sapiens	Caracas	Urban	TcI	
41.	MHOM/VE/2007/LH49	Homo sapiens	Caracas	Urban	TcI	
42.	MHOM/VE/2007/LH51	Homo sapiens	Caracas	Urban	TcI	
43.	MHOM/VE/2007/LH60	Homo sapiens	Caracas	Urban	TcI	
44.	MDID/VE/2007/LH38	Didelphis marsupialis	Cojedes	Rural	TcI	
45.	MDID/VE/2007/LH44	Didelphis marsupialis	Cojedes	Rural	TcI	
46.	MDID/VE/2007/LH45	Didelphis marsupialis	Cojedes	Rural	TcIII	
47.	MCAN/VE/2007/LH11	Canis familiaris	Cojedes	Rural	TcI	
48.	MCAN/VE/2007/LH50	Canis familiaris	Cojedes	Rural	TcI	
49.	MHOM/VE/2007/LH29	Homo sapiens	Guárico	Rural	TcI	

<sup>&</sup>lt;sup>a</sup>Reference strains; <sup>b</sup>Possible DTU identified by two of the three markers employed.

tors feces; subsequently, samples were cultured for at maximum two passages to avoid parasite culture selection.

#### DNA extraction

The DNA extractions from *T. cruzi* cultures was carried out with a mixture of phenol-chloroform-isoamilic alcohol, sodium acetate and ethanol precipitation. DNA concentration and purity were determined by spectrophotometry at 260 and 280 nm (UV/Visible GeneQuant pro RNA/DNA Calculator, Amershan)<sup>18</sup>.

# Molecular characterization of the T. cruzi isolates

The molecular characterization of *T. cruzi* was carried out using the molecular markers previously described<sup>7</sup>: (i) intergenic region of the non-transcribed miniexon gene using the primers: TC 5'-CCCCCTCCCA GGCCACACTG-3', TC1 5'-GTGTCCGCCACCTCCTT CGGGCC-3' and TC2 5'CCTGCAGGCACACGTGTGT GTG-3'; (ii) D7 divergent domain of the 24Sα rDNA employing the primers: D71, 5'-AAGGTGCGTCGACA GTGTGG-3' and D72 5'-TTTTCAGAATGGCCGAA CAGT-3' and (iii) size-variable domain of the 18S rDNA using the primers: V1, 5'-CAAGCGGCTGGGTGGTTA TTCCA-3' and V2, 5'-TTGAGGGAAGGCATGACACA TGT-3'.

For all molecular markers, the amplification reactions included Taq polymerase amplification buffer (100 mM Tris-HCl, pH 8.3), 0.2 mM dNTPs solution, 1.5 mM MgCl<sub>2</sub> solution, 1 U of GoTaq® Flexi DNA Polymerase (Promega, Madison, USA), 0.5  $\mu$ M of each primer, 10  $\mu$ l of DNA template and water till a 25  $\mu$ l final total volume. Amplification cycles were performed according to Brisse *et al*<sup>7</sup>, using a BIORAD Cycler (Bio-Rad Laboratories, Philadelphia, USA). The PCR products for each reaction were analyzed by electrophoresis on a 2% agarose gel and visualized by ethidium bromide staining.

## Purification of PCR mini-exon product

The amplification product of the intergenic region of the *T. cruzi* mini-exon was purified from agarose gels with the commercial Wizard® SV Gel kit and PCR Clean-Up System (Promega, Madison, USA), according to the manufacturer's protocol.

## DNA sequencing

Sequencing of DNA fragments was performed at the Sequencing Department of the National Center for Microbiology, Instituto de Salud Carlos III, Madrid, Spain, using the 373 Asystem, Model 377 (Applied Biosystem); samples to be sequenced were submitted to Big Dye Terminator Cycle Sequencing ready reaction kit protocol

(ABI-PRISM, PE Biosystems, Life Technologies, NY, USA).

## Phylogenetic analysis

The multiple alignments were performed using the ClustalW application and BioEdit Sequence Alignment Editor, version 7.0.5.3<sup>19</sup>. Phylogenetic and molecular evolutionary analyses were conducted by MEGA programme, version 4.1. The evolutionary history was inferred using the Neighbor-Joining method<sup>20–21</sup>.

#### **RESULTS**

## Molecular characterization of T. cruzi isolates

The characterization of *T. cruzi* isolates through the amplification of the intergenic region of mini-exon gene, rDNA 24Sα and 18S rDNA, yielded 42 samples, out of 44 (95.5%), showing a TcI DTU profile, demonstrated by amplification of specific bands of 350, 110 and 175 bp, respectively. This amplification pattern was similar to the ones exhibited by the TcI reference strains (EP, EP6c and Dm28c) included in the experiments. Amplicons of 300, 110 and 165 bp were only observed in the TcV reference isolates, corresponding to the mini-exon, rDNA 24Sα and 18S rDNA genes, respectively. Surprisingly, the mini-exon intergenic region was not amplified in two T. cruzi isolates; Canis familiaris LH8 isolate from a peridomestic rural ecotope of Altos de Guanta village, Anzoátegui state, and Didelphis marsupialis LH45 isolate from a peridomestic rural ecotope of Cojedes state. These two isolates showed rDNA 24Sα PCR amplicons of 125 and 110 bp, corresponding to the TcIV and TcIII lineage respectively. In addition, the LH8 isolate had a 175 bp-18S rDNA amplification product, characteristic of TcI-DTU, instead of the 155 bp of TcIV-; so LH8 isolate was identified as "possible TcIV", considering it showed two out of three markers employed. The LH45 isolate showed a possible TcIII pattern (165 bp band) using this marker (Table 1).

## DNA sequencing

The multiple alignments of sequences showed high similarity among the nucleotide sequences of the isolates characterized as TcI. These TcI isolates were very different from TcV reference strains, showing an important nucleotide variation. Although, almost all isolates from Anzoátegui, Caracas, Cojedes and Guárico were identified as DTU TcI populations, the sequences of the miniexon marker in these isolates show some nucleotide variability. A length of 322 bp was obtained in all isolates; the genetic variability amongst TcI isolates was of 15.5%.

There was nucleotide divergence in 50 positions, corresponding to single nucleotide polymorphisms (SNP) with 15 transitions (4.7%), 17 transversions (5.3%) and 23 insertion-deletions (7.1%). There were 272 constant positions. The TcI isolates obtained from vectors and mammalian hosts in Caracas were a group intrinsically

homogeneous, showing differences with the sequences of the rural isolates from Anzoátegui, Cojedes and Guárico (Table 2).

Phylogenetic analysis

The phylogenetic tree, corresponding to the parasite

Table 2. Variable positions in mini-exon sequences of the TcI isolates studied

Isolate	Locality	Variable position																				
	~ · · ·	39	76	78	81	97	100	122	127	198	289		291	292	293	294	295	296		302	303	
	Guárico	G	A	G	C	A	C	A	A	C	G	T	_	_	_	_	-	_	T	_	G	C
EP	Guárico	-	G	G	G	G	C	G	A	C	A	C	A	C	A	C	A	C	T	G	G	C
EP6c	Guárico	_	A	G	G	G	C	G	A	A	A	C	A	C	Α	C	A	C	T	G	G	C
LH1	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	-	_	_	T	_	_	G
LH4	Anzoátegui	G	A	G	C	G	G	A	A	C	T	_	_	_	_	_	_	-	T	G	G	C
LH5	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	-	G	T	G	_
LH6	Anzoátegui	G	A	G	C	G	G	A	A	C	T	-	_	_	_	_	_	_	G	T	G	G
LH12	Anzoátegui	G	A	G	C	G	G	A	A	C	T	_	_	_	_	_	_	_	G	T	G	G
LH13	Anzoátegui	G	A	G	C	G	G	A	A	C	T	_	_	_	_	_	_	_	G	T	G	G
LH19	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	T	G	_
LH20	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	_	_	G
LH23	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	_	_	G
LH26	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	_	_	G
LH2	Anzoátegui Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	_ T	G	_
LH3 LH10	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	T	G	_
LH18	Anzoátegui	G	A	G	C	G	G	A	A	C	T	_	_	_	_	_	_	_	G	T	G	G
LH21	_	G	A	G	C	G	G	A	A	C	G	т	_	_	_	_	_	_	G	T	G	G
LH22	Anzoátegui Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	_	_	G
LH25	Anzoátegui	G	A	G	C	G	G	A	A	C C	G G	T T	_					_	T	— Т	– G	G
LH27	Anzoátegui	G G	A	G	C	G	G	A	A	C	G								T T			_
LH28	Anzoátegui	G	A A	G G	C C	G G	G G	A A	A A	C	G	T	_	_		_	_	_	G	— Т	– G	G
LH7	Anzoátegui	G	A	G	C	G	G	A	A	C	G	_	_	_	_	_	_	_	T	T	G	G
LH9	Anzoátegui	G	A	G	C	G	G	A	A	C	T	_	_	_	_	_	_	_	G	T	G	G
LH24	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	T	G	G
LH14	Anzoátegui	G	A	G	C	G	G	A	A	C	G	_	_	_	_	_	_	_	G	T	G	_
LH33	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	T	_	_	G
LH43	Anzoátegui	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	G	T	G	U
LH35	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH36	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH30	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH34	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH37	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH42	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH46	Caracas	G	G	A	C	A	C	A	C	A	G	Ť	_	_	_	_	_	_	Ť	_	_	G
LH47	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH48	Caracas	G	G	A	C	A	C	A	C	A	G	Ť	_	_	_	_	_	_	T	_	_	G
LH49	Caracas	G	G	A	C	A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH51	Caracas	G	G	A		A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH60	Caracas	G	G	A		A	C	A	C	A	G	T	_	_	_	_	_	_	T	_	_	G
LH38	Cojedes	G	A	G	C	A	G	A	A	C	T	_	_	_	_	_	_	_	G	T	G	C
LH44	Cojedes	_	G	G	G	G	C	G	A	A	A	C	A	C	A	C	A	C	T	G	T	G
LH11	Cojedes	G	A	G	C	G	G	A	A	C	G	T	_	_	_	_	_	_	G	T	G	G
LH50	Cojedes	_	G	G	G	G	C	G	A	A	A	C	A	C	A	C	A	C	T	G	T	G
LH29	Guárico	_	G	G		G	G	G	A	C	A	C	A	C	A	C	A	C	T	G	G	C
11127	Juaneo		U	U	U	U	U	U	Α	C	А		А	C	А	C	А	C	1	U	U	

<sup>(-)</sup> Denote gaps in these positions. The shading denote specific changes in the sequences of isolates from Caracas.

populations analyzed, showed that the T. cruzi isolates from Caracas had more homogeneous sequences than the ones from the other regions (Fig. 1). To highlight this result, dotted lines box, grouping the T. cruzi isolates from Caracas was included in Fig. 1, as they were the only isolates from urban environments and were clustered with a high bootstrap. In contrast, the other isolates were interspersed throughout the different clades, regardless of their geographic origin, all were from rural environments and bootstrap values were lower compared to the Caracas isolates clade form. Conserved substitutions in several positions (14) were observed among the isolates from both urban and rural areas, highlighting the nucleotide variability between the Caracas isolates (urban isolates) and the other isolates from Anzoátegui, Cojedes and Guárico regions (rural isolates). Indeed, mini-exon sequences com-

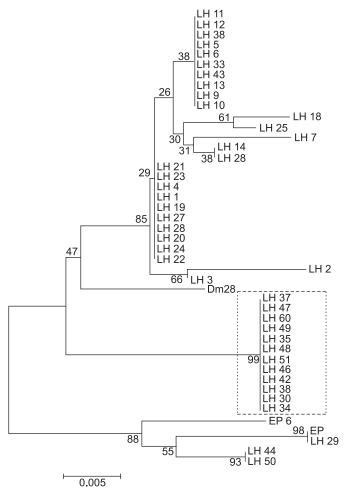


Fig. 1: Evolutionary relationships of Venezuelan Trypanosoma cruzi TcI isolates. The evolutionary history was inferred using the Neighbor-Joining method<sup>20</sup>. The optimal tree is shown. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (500 replicates) is shown next to the branches. Phylogenetic analyses were conducted in MEGA 4<sup>21</sup>. In the dotted lines box, the isolates of T. cruzi from Caracas are grouped.

parisons were made with other *T. cruzi* TcI isolates, obtained in different areas of Venezuela and reported in GenBank. It was observed that most of them were grouped together with the isolates from Cojedes Guárico and Anzoátegui, whereas the Caracas isolates were a separate genetic group (data not shown). Estimates of average evolutionary divergence within groups using the maximum composite likelihood method in MEGA4, showed 0.012 for rural isolates and 0.001 for urban ones, while the divergence between rural isolates and urban ones was 0.039.

#### **DISCUSSION**

T. cruzi is a hemoflagellate of clonal structure with occasional genetic recombination, composed of several subpopulations circulating in vectors, reservoirs and humans from domestic and/or sylvatic cycles. These stable clones are of great significance in terms of adaptive evolution to new environments, as new vectors or reservoirs including humans and its distribution would influence the clinical course of human disease and some epidemiological associations for different regions<sup>22</sup>.

In Venezuela, recent studies suggested the re-emergence of Chagas disease, with important epidemiological changes that favored the presence of the disease not only in endemic regions with rural conditions housing, but also in urban areas with diverse human dwellings<sup>3-4</sup>. Thus, urban areas include housing zinc roof, concrete walls, high class buildings, that permit new habitats for wildlife and synanthropic vectors, such as *Panstrongylus geniculatus or Triatoma maculata* in sympatry with the primary vector, *Rhodnius prolixus*, which represent risk factors for transmission of the disease<sup>3-4, 14, 23-24</sup>. These epidemiological changes could be associated with genetic variability in isolates of *T. cruzi*, which has not been deeply studied in Venezuela.

In the present study, the characterization of the *T. cruzi* isolates from vectors and reservoirs collected in Anzoátegui, Caracas and Cojedes, showed high frequency of TcI-DTU (95.5%), in agreement with other reports of this frequent DTU in at least 17 states of Venezuela, and occasional occurrence of TcIII and TcIV genotypes<sup>15-17, 25</sup>.

TcI has been considered as a homogeneous group; however, some recent studies have reported variations in the intergenic region of the mini-exon sequence in TcI isolates derived from vectors, reservoirs and humans, and collected in different countries as Bolivia, Mexico, Brazil, Colombia, and Argentina, being identified as DTU variants<sup>7, 9–10, 26–27</sup>.

In this study, we found evidence of TcI heterogene-

ity by mini-exon DNA sequencing, as was also suggested in studies of *T. cruzi* isolates from human and vectors from western region or urban Venezuelan capital (Caracas), characterized as TcI but without association with one particular clinical manifestation of Chagas disease<sup>14, 17</sup>. In addition, genetic variability within *T. cruzi* I strains from orally and non-orally transmitted human cases were also reported in Venezuela<sup>25</sup>.

Regarding the present work, the TcI genetic variability observed could be associated with geographical distribution, as it was proposed in other studies<sup>26</sup>. The nucleotide variability found in mini-exon gene could suggest that the SNPs identified have simply accumulated during clonal diversification of TcI in geographically isolated populations, or an adaptive parasite response to different environments<sup>27</sup>. Due to the presence of progressive epidemiological changes in Venezuela, it would be interesting to evaluate more isolates and other markers.

The TcI nucleotide sequences of isolates from vectors and reservoirs, collected in rural habitats from Anzoátegui and Cojedes, showed variations in relation to the sequences of *T. cruzi* isolates from urban habitats of Caracas. This finding could show a possible segregation and selection of subpopulations in function of geographic area.

In Anzoátegui counties, northeastern part of Venezuela, the parasite populations circulate in rural and semirural biotope using different vectors, such as *R. prolixus*, *T. maculata* and *P. geniculatus* and more sporadically *P. rufotuberculatus* and *Eratyrus mucronatus*<sup>24, 28</sup>. These vectors feed on a variety of reservoirs, which may contribute to genetic recombination events, resulting in appearance of new hybrids or genotypes of the parasite, which be infective to domiciled reservoirs, including *C. familiaris*.

The presence of a *C. familiaris* isolate from Anzoátegui, LH8, with a possible TcIV pattern, that did not have all the characteristic markers, could reflect one mixed infection with several DTUs, probably due to the epidemiological role of dogs, as sentinel of *T. cruzi* infections, and the probability of infection in different areas, where owners move with them.

The *T. cruzi* TcI isolates from Caracas showed sequences more homogeneous. Perhaps, in the Caracas region, human induced environmental changes through deforestation and uncontrolled urbanization, which may have altered the original ecological niche, where the parasite circulated in a zoonotic cycle, with a variety of vectors and wild reservoirs, similar to those observed in Anzoátegui and Cojedes regions. In response to the environmental changes, certain vector species could have been adapted to the human habitat, specially attracted by home light and

blood sources in the new urban scenario. In the new areas, the extinction of certain species of vectors and reservoirs favors the dominance of a single vector, *P. geniculatus*, and few reservoirs such as *Rattus rattus*, *D. marsupials* and *Homo sapiens*<sup>29</sup>. These hosts could have acted as biological filters that, over the years, select parasites subpopulations in a closed zoonotic cycle, genetically distinct from the TcI DTU from rural regions. Similar features of genetic variability associated to different epidemiological cycles have been also described in Colombia<sup>9</sup>.

The sequence comparisons between the mini-exon markers of the 42 isolates and other eight sequences from Venezuelans *T. cruzi* TcI isolates deposited in the GenBank showed that they did not have a preferential distribution in relation to geographical area, whereas the Caracas isolates described in the present study remained as a separate group.

Although, the isolates from Caracas characterized in this study were few, its biological characterization in murine models showed higher virulence than the Anzoátegui and Cojedes isolates, and a particular tropism for the central nervous system, ocular tissues, genital organs, bone, cartilage, kidney, lung, liver, and pancreas<sup>30</sup>. In this sense, in the area of Caracas, severe symptomatic cases have been observed recently, many of them associated with oral transmission, showing more pronounced symptoms than those associated to fecal contamination through vector transmission<sup>4, 25</sup>. This increased virulence could be a biological expression in response to genetic recombination events and the result of parasite adaptation to environmental changes, new vectors and reservoirs.

It would be important to have more parasites isolated from humans, vectors and reservoirs of different geographical areas to confirm this trend towards a genetic difference between isolates from rural areas and those from urban areas. However, the difficulties found in *T. cruzi* isolation as well as the classification of some rural, urban and hybrid environments are important limitations to carry out this type of investigations. Therefore, it is advisable to perform characterization studies in order to elucidate the relation of genetic variability within the TcI, with the emergent epidemiological pattern that is currently being observed in Chagas disease in Venezuela.

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#### REFERENCES

- Chagas disease (American trypanosomiasis). 2010. Fact Sheet No. 340. Available from: http://www.who.int/mediacentre/ factsheets/fs340/en/index.html (Accessed on June 24, 2014).
- 2. Prata A. Clinical and epidemiological aspects of Chagas disease. *Lancet Infect Dis* 2001; *1*: 92–100.
- 3. Añez N, Crisante G, Rojas A. Update on Chagas disease in Venezuela A review. *Mem Inst Oswaldo Cruz* 2004; 99: 781–7.
- Alarcón de Noya B, Díaz-Bello Z, Colmenares C, Ruiz-Guevara R, Mauriello L, Zavala-Jaspe R, et al. Large urban outbreak of orally acquired acute Chagas disease at a school in Caracas, Venezuela. J Infect Dis 2010; 201: 1308–15.
- 5. Devera R, Fernandes O, Coura JR. Should *Trypanosoma cruzi* be called "cruzi" complex? A review of the parasite diversity and the potential of selecting population after *in vitro* culturing and mice infection. *Mem Inst Oswaldo Cruz* 2003; 98: 1–12.
- Souto R, Fernandes O, Macedo A, Campbell D, Zingales B. DNA markers define two major phylogenetic lineages of *Trypanosoma* cruzi. Mol Biochem Parasitol 1996; 83: 141–52.
- Brisse S, Verhoef J, Tibayrenc M. Characterisation of large and small subunit rRNA and mini-exon genes further supports the distinction of six *Trypanosoma cruzi* lineages. *Int J Parasitol* 2001; 31: 1218–26.
- 8. Zingales B, Andrade SG, Briones MR, Campbell DA, Chiari E, Fernandes O, *et al.* A new consensus for *Trypanosoma cruzi* intraspecific nomenclature: Second revision meeting recommends TcI to TcVI. *Mem Inst Oswaldo Cruz* 2009; *104:* 1051–4.
- Herrera C, Guhl F, Falla A, Fajardo A, Montilla M, Vallejo GA, et al. Genetic variability and phylogenetic relationships within Trypanosoma cruzi I isolated in Colombia based on mini-exon gene sequences. J Parasitol Res 2009. doi:10.1155/2009/897364.
- Falla A, Herrera C, Fajardo A, Montilla M, Vallejo G A, Guhl F. Haplotype identification within *Trypanosoma cruzi* I in Colombian isolates from several reservoirs, vectors and humans. *Acta Trop* 2009; *110*: 15–21.
- Ramírez JD, Duque MC, Montilla M, Cucunubá ZM, Guhl F. Multilocus PCR-RFLP profiling in *Trypanosoma cruzi* I highlights an intraspecific genetic variation pattern. *Infect Genet Evol* 2012; 12: 1743–50.
- Macedo A, Oliveira R, Pena S. Chagas disease: Role of parasite genetic variation in pathogenesis. *Expert Rev Mol Med* 2002; 4: 1–16.
- Higo H, Miura S, Horio M, Mimori T, Hamano S, Agatsuma T, et al. Genotypic variation among lineages of *Trypanosoma cruzi* and its geographic aspects. *Parasitol Int* 2004; 53: 337–44.
- 14. Añez N, Crisante G, da Silva FM, Rojas A, Carrasco H, Umezawa ES, et al. Predominance of lineage I among Trypanosoma cruzi isolates from Venezuelan patients with different clinical profiles of acute Chagas disease. Trop Med Int Health 2004; 9: 1319–26.
- Llewellyn MS, Miles MA, Carrasco HJ, Lewis MD, Yeo M, Vargas J, et al. Genome-scale multilocus microsatellite typing of Trypanosoma cruzi discrete typing unit I reveals phylogeographic structure and specific genotypes linked to human infection. PLoS Pathog 2009; 5(5): e1000410. doi: 10.1371/

- journal.ppat.1000410.
- MorocoimaA, Carrasco HJ, Boadas J, Chique JD, Herrera L, Urdaneta-Morales S. *Trypanosoma cruzi* III from armadillos (*Dasypus novemcinctus novemcinctus*) from northeastern Venezuela and its biological behavior in murine model. Risk of emergency of Chagas' disease. *Exp Parasitol* 2012; *132*: 341–7.
- Carrasco HJ, Segovia M, Llewellyn MS, Morocoima A, Urdaneta-Morales S, Martínez C, et al. Geographical distribution of *Trypanosoma cruzi* genotypes in Venezuela. *PloS Negl Trop Dis* 2012; 6(6): e1707. doi: 10.1371/journal.pntd.0001707.
- 18. Sambrook J, Russel D. *Molecular cloning: A laboratory manual*. III edn. New York: Cold Spring Harbor 2001; p. 6.4–6.12.
- Hall TA. BioEdit: A user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. *Nucl Acids Symp Ser* 1999; 41: 95–8.
- Saitou N, Nei M. The neighbor-joining method: A new method for reconstructing phylogenetic trees. *Mol Biol Evol* 1987; 4: 406–25.
- Tamura K, Dudley J, Nei M, Kumar S. MEGA4: Molecular evolutionary genetics analysis (MEGA) software version 4.0. *Mol Biol Evol* 2007; 24: 1596–9.
- 22. Gaunt M, Yeo M, Frame I, Stothard J, Carrasco H, Taylor MC, *et al.* Mechanism of genetic exchange in American trypanosomes. *Nature* 2003; *421*: 936–9.
- Morocoima A, Chique J, Zavala-Jaspe R, Díaz-Bello Z, Ferrer E, Urdaneta-Morales S, et al. Commercial coconut palm as an ecotope of Chagas disease vectors in northeastern Venezuela. J Vector Borne Dis 2010; 47: 76–84.
- Morocoima A, Coriano H, Navas C, De Sousa L, Ferrer E, Herrera L. Panstrongylusrufo tuberculatus (Hemiptera, Reduviidae, Triatominae) infected with Trypanosoma cruzi in the state of Anzoátegui (Venezuela). Bol Mal Salud Amb 2012; 52: 135–8.
- Segovia M, Carrasco HJ, Martínez CE, Messenger LA, Nessi A, Londoño JC, et al. Molecular epidemiologic source tracking of orally transmitted Chagas disease, Venezuela. Emerg Infect Dis 2013; 19: 1098–101.
- O' Connor O, Bosseno M, Barnabé C, Douzery E, Breniere S. Genetic clustering of *Trypanosoma cruzi* I lineage evidenced by intergenic mini-exon gene sequencing. *Infect Genet Evol* 2007; 7: 587–93.
- 27. Tomasini N, Lauthier JJ, Monje Rumi MM, Ragone PG, Alberti D'Amato, AA, Pérez Brandan C, *et al.* Interest and limitations of sliced leader intergenic region sequences for analyzing *Trypanosoma cruzi* I phylogenetic diversity in the Argentinean Chaco. *Infect Genet Evol* 2011; *11*: 300–7.
- Morocoima A, Chique J, Herrera L, Urdaneta-Morales S. Eratyrus mucronatus (Stal, 1859) (Hemiptera, Reduviidae, Triatominae): Primer registro para el estado Anzoátegui (Venezuela). Bol Mal Salud Amb 2010; 50: 307–10.
- Herrera L, Urdaneta S. Synanthropic rodent reservoirs of *Trypanosoma (Schizotrypanum) cruzi* in the valley of Caracas, Venezuela. *Rev Inst Med Trop Sao Paulo* 1997; 39: 279–82.
- 30. Morocoima A, Rodríguez M, Herrera L, Urdaneta S. *Trypanosomacruzi*: Experimental parasitism of bone and cartilage. *Parasitol Res* 2006; *99*: 663–8.

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