The identification of two Trypanosoma cruzi I genotypes from domestic and sylvatic transmission cycles in Colombia based on a single polymerase chain reaction amplification of the spliced-leader intergenic region

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A single polymerase chain reaction (PCR) reaction targeting the spliced-leader intergenic region of Trypanosoma cruzi I was standardised by amplifying a 231 bp fragment in domestic (TcI-dom) strains or clones and 450 and 550 bp fragments in sylvatic strains or clones. This reaction was validated using 44 blind coded samples and 184 non-coded T. cruzi I clones isolated from sylvatic triatomines and the correspondence between the amplified fragments and their domestic or sylvatic origin was determined. Six of the nine strains isolated from acute cases suspected of oral infection had the sylvatic T. cruzi I profile. These results confirmed that the sylvatic T. cruzi I genotype is linked to cases of oral Chagas disease in Colombia. We therefore propose the use of this novel PCR reaction in strains or clones previously characterised as T. cruzi I to distinguish TcI-dom from sylvatic genotypes in studies of transmission dynamics, including the verification of population selection within hosts or detection of the frequency of mixed infections by both T. cruzi I genotypes in Colombia.

Key words: spliced-leader intergenic region - Trypanosoma cruzi I - PCR - Chagas disease - domestic cycle - sylvatic cycle

Trypanosoma cruzi, the causative agent of Chagas disease, has been divided into six discrete taxonomic units (DTUs) named T. cruzi I-VI (Zingales et al. 2009). Recently, many authors have attempted to determine the geographical distribution of T. cruzi DTUs and their relationships with transmission dynamics and clinical outcomes. As summarised by Vallejo et al. (2009a), studies using TCC/TCl/TCl primers designed for the spliced-leader intergenic region (SL-IR) (Souto et al. 1996) have demonstrated that T. cruzi I predominates in Colombian domestic and sylvatic vectors, with 2.1-78.6% having been detected in naturally-infected Rhodnius prolixus, Rhodnius colombiensis and Rhodnius pallescens midgut specimens. Studies of Colombian Chagas disease patients have also revealed the predominance of T. cruzi I over other DTUs in blood and tissue explants (Zafra et al. 2008, 2011, Mantilla et al. 2010, Ramírez et al. 2010).

Although there is a current consensus regarding the predominance of T. cruzi I in Colombian, Venezuelan and Central American countries, T. cruzi I has tremendous genetic variability, as demonstrated by several molecular markers (Guhl & Ramírez 2011). Based on the SL-IR region, four T. cruzi I genotypes associated with Chagas disease transmission cycles have been described in Colombia (Ia, Ib, Ic, Id) and corroborated through the American continent by the detection of a novel genotype, named T. cruzi Ie, associated with Mepraia species in Bolivia and Chile (Herrera et al. 2007, 2009, Cura et al. 2010). Specific primers have also been developed to identify three of these four genotypes (Ia, Ib and Id) (Falla et al. 2009). Other molecular markers have confirmed the intraspecific variability within T. cruzi I, suggesting the emergence of novel genotypes such as a TcI clade, which has been named TcI-dom (formerly TcIa/VEn-dom) and is associated with the domestic transmission cycle throughout the American continent (Ramírez et al. 2012c). Several authors have thus proposed the separation of the two main T. cruzi I groups based on associations with domestic and sylvatic cycles (Llewellyn et al. 2009, Ocaña-Mayorga et al. 2010, Ramírez et al. 2012a, b).

The domestic and sylvatic cycles of T. cruzi are not strictly separated. Several reports have shown that sylvatic triatomines can invade housing units and contaminate food with T. cruzi, causing outbreaks of “oral Chagas disease”. For example, the application of high-resolution molecular markers to biological clones from strains isolated during such outbreaks have incriminated T. cruzi I sylvatic strains and clones as causal agents of oral Chagas disease outbreaks in Colombia (Ramírez et al. 2013). This association between domestic and sylvatic-
ic cycles in *T. cruzi* I highlights the potential for further research in this field. The objective of this present study was to develop a method for single polymerase chain reaction (PCR) amplification of the SL-1R to discriminate between domestic and sylvatic genotypes and to obtain a better understanding of domestic and sylvatic cycles as well as the transmission dynamics of this DTU.

SL-1R sequences (accessions AM259467-AM259477 and EU626722-EU626738) were retrieved from GenBank and subsequently aligned. The 1Am primer (5'-TGTGT-GTGTATGTATGTC-3') was designed and used with primer 1B (5'-CGGACCGGTGTGTGCAG-3') (Falla et al. 2009) to perform PCR reactions in a total volume of 20 µL containing 2 µL of 10X reaction buffer (Invitrogen), 200 µM of a mixture of deoxynucleotide triphosphates, 1.5 mM MgCl₂, 25 µM each primer, 0.5 units of Taq DNA polymerase (Invitrogen). The template (20 ng) included DNA from a strain or clone previously characterised as *T. cruzi* I using different molecular markers, including multilocus enzyme electrophoresis or PCR of large subunit rDNA (Souto et al. 1996), PCR-restriction fragment length polymorphism (RFLP) of heat shock protein 60-ECORV or glycosylphosphatidylinositol-Hhal (Westenberger et al. 2005), PCR-RFLP of COII-Alul (Freitas et al. 2006) and PCR with TCC/TCl/TC2 primers designed in the SL-1R (Souto et al. 1996), as shown in Fig. 1A.

All PCR reactions were conducted for 35 amplification cycles using a thermal minicycler (MJ Research PTC-150-16). Cycling include a denaturing step at 94°C for 30 s (4 min for initial denaturing), annealing at 51°C for 30 s, extension at 72°C for 30 s and a final extension step at 72°C for 10 min.

After standardising the reaction conditions using the previously described primers, the TcI DOM and clones produced the expected 231 bp amplification product; however, the sylvatic clones and strains produced 450-550 bp amplification products when using the same primers (Fig. 1B). Several *T. cruzi* I samples and clones isolated from sylvatic triatomines and reservoirs produced the 450-550 bp profile, as shown in Fig. 1B. The ORT15 strain, isolated by xenoculture from a rectal ampulla from *R. colombiensis* in the department of Tolima, was correctly typed after its isolation in February 2013 (Fig. 1B, Lane 3). *T. cruzi* I clones from the intestinal contents of *R. colombiensis*, *R. pallescens* and sylvatic *R. prolixus*, obtained by direct plating in a sensitive solid medium (Yeo et al. 2007), produced the 450-550 bp profile (Fig. 1B, Lanes 4-8). The D1 and D2 strains, isolated from *Dielphis marsupialis* and maintained in culture in biphasic medium, also produced the sylvatic profile (Fig. 1B, Lanes 9, 10). The existence of mixed profiles in *T. cruzi* I strains is possible, as 175 of 182 (96.2%) clones obtained from sylvatic *R. colombiensis*, *R. pallescens* and *R. prolixus* produced the 450-550 bp profile and seven of these 182 clones (3.8%) produced a 231 bp profile (data not shown). These results support the assertion that the 450-550 bp profile is predominant in sylvatic cycles.

Alternately, MHOM/CO/04/MG, a human strain that was isolated by xenoculture from the intestinal content of domiciliated *R. prolixus* in the Department of Boyaca and kept in culture for several years, displayed the 231 and 450-550 bp profiles (Fig. 1B, Lanes 11, 12). A similar mixed profile was observed in the human EH strain (Lane 14), suggesting a mixed infection of both genotypes, as previously reported by Ramirez et al. (2013).

Domestic or sylvatic reservoirs may also select for subpopulations within *T. cruzi* I. The possibility that domestic and sylvatic vectors may selectively transmit *T. cruzi* I genotypes cannot be excluded, as the selective transmission of both *Trypanosoma rangeli* genotypes has been demonstrated for various *Rhodnius* species (Pulido et al. 2008, Vallejo et al. 2009b, Urrea et al. 2011).

When diagnostic or molecular characterisation techniques are developed, it is necessary to verify that the technique is specific to *T. cruzi* and demonstrate a lack of cross-reaction with *T. rangeli*, as *T. rangeli* frequently occurs in mixed infections with *T. cruzi* in triatomines and in vertebrate reservoirs in Colombia. We emphasise that the purpose of this work was to use a PCR reaction with 1Am/1B primers in strains that had previously been typed as *T. cruzi I sensu lato* and thus we initially used strains or clones that were previously characterised as *T. cruzi I*. Nevertheless, because mixed *T. rangeli* infections in vectors and vertebrates can be found in nearly all *T. cruzi*-endemic areas in Colombia, strains can therefore be isolated together with *T. cruzi I* and *T. rangeli*. *T. rangeli* DNA and 1Am/1B primers, which did not produce any amplification products, were used as controls to rule out cross-reactions (data not shown).

Our results demonstrated that Colombian domestic and sylvatic *T. cruzi* I strains can be discriminated using a single PCR amplification of the SL-1R from TcI DOM (formerly TcIa/VEN DOM) and sylvatic strains. Nine isolates from patients suffering acute Chagas disease involving different transmission routes and different clinically diagnosed symptomatology were used to verify whether this single PCR reaction could be used to discriminate between *T. cruzi* I infections of TcI DOM and sylvatic genotype origin. Six of the nine isolates had the sylvatic profile, corresponding to three cases of acute myocarditis in outbreaks of oral transmission (Santander, Colombia), a congenital case (Santander, Colombia) and two cases of sylvatic vector transmission (Putumayo, Colombia) (Fig. 2). The remaining three cases were identified as *T. cruzi* II when using D71/D72, V1/V2 primers (Brisse et al. 2001). *T. cruzi* II was found in two congenital cases (Boyaca, Colombia), as previously described (Pavia et al. 2009), and in a case of reactivation caused by human immunodeficiency virus/acquired immune deficiency syndrome (Caquetá, Colombia). Human *T. cruzi* II strains were isolated by haemoculture during the acute phase of infection and were typed long after isolation and thus there was no patient follow-up to obtain new isolates for further characterisation. Selection among *T. cruzi* genotype mixtures when maintained in culture or in animal models could not be ruled out, as selection towards TcII in mixed infections with TcI strains (with subtle TcII parasites or clones) has been reported (Pena et al. 2011).
Our single PCR reaction for discriminating *Trypanosoma cruzi* I genotypes could be of great use for determining the genotype frequency as well as follow-up analyses comparing the profiles of initially isolated genotypes with their later stability or selection after maintenance in culture or biological models for varied lengths of time.

The results of this present study corroborated previous reports by Ramírez et al. (2013) highlighting the fact that the sylvatic *Trypanosoma cruzi* I genotype is implicated in oral and sylvatic vector transmission in Colombia. New studies involving a greater number of human isolates should further support these observations.

In summary, after *Trypanosoma cruzi* I has been diagnosed in any isolate, we propose that an additional PCR reaction be used with a single pair of primers for discriminating between TcI_dom and sylvatic *Trypanosoma cruzi* I. This methodology has led to the identification of the genotypes involved in outbreaks of oral Chagas disease as well as supporting studies of the transmission dynamics of *Trypanosoma cruzi* as the causal agent of Chagas disease in Colombia. Our results agree with those of other investigators reporting that *Trypanosoma cruzi* I encompasses two main groups that, to date, are not known to be stable in time and space. Further investigation will be useful for verifying population selection within hosts and detecting the frequency of mixed genotypes of *Trypanosoma cruzi* I in Colombia.

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**REFERENCES**


