Repertoire, Genealogy and Genomic Organization of Cruzipain and Homologous Genes in Trypanosoma cruzi, T. cruzi-Like and Other Trypanosome Species

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Abstract

Trypanosoma cruzi, the agent of Chagas disease, is a complex of genetically diverse isolates highly phylogenetically related to T. cruzi-like species, Trypanosoma cruzi marinkellei and Trypanosoma dionisii, all sharing morphology of blood and culture forms and development within cells. However, they differ in hosts, vectors and pathogenicity: T. cruzi is a human pathogen infective to virtually all mammals whilst the other two species are non-pathogenic and bat restricted. Previous studies suggest that variations in expression levels and genetic diversity of cruzipain, the major isoform of cathepsin L-like (CATL) enzymes of T. cruzi, correlate with levels of cellular invasion, differentiation, virulence and pathogenicity of distinct strains. In this study, we compared 80 sequences of genes encoding cruzipain from 25 T. cruzi isolates representative of all discrete typing units (DTUs TcI-TcVI) and the new genotype Tcbat and 10 sequences of homologous genes from other species. The catalytic domain repertoires diverged according to DTUs and trypanosome species. Relatively homogeneous sequences are found within and among isolates of the same DTU except TcV and TcVI, which displayed sequences unique or identical to those of TcI and TcII, supporting their origin from the hybridization between these two DTUs. In network genealogies, sequences from T. cruzi clustered tightly together and closer to T. c. marinkellei than to T. dionisii and largely differed from homologues of T. rangeli and T. b. brucei. Here, analysis of isolates representative of the overall biological and genetic diversity of T. cruzi and closest T. cruzi-like species evidenced DTU- and species-specific polymorphisms corroborating phylogenetic relationships inferred with other genes. Comparison of both phylogenetically close and distant trypanosomes is valuable to understand host-parasite interactions, virulence and pathogenicity. Our findings corroborate cruzipain as a valuable target for drugs, vaccine, diagnostic and genotyping approaches.

Introduction

Cathepsin L-like (CATL) are cysteine proteases that play important roles in cell invasion, growth, differentiation, immunity, immune-modulation, virulence, pathogenicity and survival of pathogenic protozoans. Different isoforms of CATL are encoded by a large gene family and perform distinct roles in the interactions of the trypanosomes with vertebrate hosts and vectors, differing in stage, cellular localization and expression level during the life cycle. This functional and structural diversification may have contributed to the adaptation of different trypanosome species to their different life cycles, vertebrate hosts and vectors [1–4].

Trypanosoma cruzi is the type species of the subgenus Schizotrypanum and a complex of genetically heterogeneous isolates distributed in 6 intraspecific subdivisions denominated discrete typing units (DTUs), TcI-TcVI [5,6], and one new genotype (Tcbat) identified in Brazilian bats [7]. Closest relatives of T. cruzi are the bat-restricted T. c. marinkellei followed by T. dionisii, which are referred as T. cruzi-like due to morphologically indistinguishable blood and culture forms [8–11]. Development within mammalian cells in vitro is a feature shared by all species of the subgenus Schizotrypanum, while in vivo only T. cruzi has been proven to infect mammals other than chiropterans [7,9,10,12–14]. Bats infected by T. cruzi-like species show nests of amastigotes in cardiac, skeletal and stomach muscle cells likewise T. cruzi in a range of hosts including man. Recent studies demonstrated that T. dionisii and T. cruzi invade mammalian cells through a common mechanism involving lysosome mobilization to the site of parasite entry [15,16]. Previous studies showed that T. cruzi and T. dionisii share similar molecules with important roles in host-parasite interactions such as phospholipids and cysteine proteases [17,18] as well as epitopes associated to autoimmunity in Chagas disease [19]. Nevertheless, T. dionisii differ from T. cruzi in surface glycoproteins involved in host-cell interactions [16]. Besides
morphism and in vitro and in vivo behavior, T. cruzi-like species share cellular, biochemical and immunological features with T. cruzi and, hence, can be valuable as non-infective to humans models for studies of T. cruzi and as targets for trials of drug, vaccines and diagnosis [10,20].

Similar to T. cruzi, all T. cruzi-like isolates differentiate from epimastigotes to infective metacyclic trypomastigotes in cultures and in the vector gut; bats are infected by lking vector feces contaminated with trypanosomes on their fur and/or by ingesting the infected vectors themselves [8,9]. Differently from T. cruzi that can be transmitted by several genera of triatomine bugs, transmission of T. evansi seems to involve only triatomines of the genus Rhodnius, while cimicids are vectors of T. dimasi [8,9]. In addition to T. cruzi, T. rangeli was reported infecting humans and non-human primates, chiroptera and mammals of other orders and are both transmitted by triatomines [21,22]. The mechanisms underpinning vertebrate and vector specificities of these trypanosomes are unknown.

Previous studies demonstrated that two main CATL enzymes are expressed by T. cruzi, the major isoform (>75 copies) addressed in the present study and designated as cruzipain, is the archetype of a large multigene family organized in tandem repeats expressed in all life cycle stages of T. cruzi [4,23–26]. Analysis of polymorphic cruzipain-encoding genes disclosed the isoform cruzipain 2 (~6 copies), which is expressed preferentially by the mammalian stages and differs markedly from cruzipain with respect to substrate specificity and kinetic properties [27,28].

Cruzipain plays fundamental functions in T. cruzi life cycle with recognized roles in parasite-host interactions, in establishing, maintaining, exacerbating and controlling infections. There are increasing evidence that the immunopathogenesis of experimental Chagas Disease is, at least in part, due to the activity of cruzipain mediating cell invasion, inflammation, tissue damage and immune evasion [29,30]. Cruzipain is an immunodominant antigen, expressed on parasite surface and secreted, which elicits potent humoral [31,32] and cellular immune responses in T. cruzi infected humans and mice [33,34]. Vaccination with recombinant cruzipain trigger strong humoral and cell-mediated immunity controlling parasite load and inflammatory tissue damage [34–37]. Addition of synthetic irreversible inhibitors to cultures of cells infected with T. cruzi blocks parasite replication, intracellular growth and differentiation [38,39]. Treatment of T. cruzi infected mice with inhibitors designed to inactivate cruzipain rescued mice from a lethal infection [40]. T. cruzi ability to invade human cells was modulated by the balance between cruzipain and chagasin, a natural endogenous inhibitor of papain-like cysteine proteases [41,42]. Therefore, inhibitors of cruzipain are among the most promising new drugs for treatment of Chagas disease [1,4,43–45].

Studies have suggested that the variable levels of cruzipain activity correlate to degrees of metacyclogenesis, cellular invasion and virulence of T. cruzi isolates. Comparison of TcI and TcII strains suggested that cruzipain proteolytic profiles could be useful for separating members of these two DTUs and that high expression levels could be linked to enhanced metacyclogenesis and cell infectivity [46]. In a study using flow cytometry and anti-cruzipain antibody both TcI and TcII isolates showed heterogeneous surface cruzipain patterns, however, expression levels were higher in TcI isolates showing higher metacyclogenesis [47]. In T. cruzi Dm28c (TcI), cruzipain is down regulated during metacyclogenesis [48]. The over expression of this enzyme throughout the parasite life-cycle was associated with enhanced metacyclogenesis but not with increased cell infectivity [49]. Differences in cruzipain expression were correlated with differential virulence for mice of T. cruzi isolates of Z3 (TcIII and IV) [50].

A proteomic analysis suggested that significant differences in the expression of cruzipain by isolates of TcIII and TcIV (Z3) could also contribute to their differential infectivity to cells [51]. Interestingly, proteolytic activity was lower in the virulent T. cruzi CL strain compared to the non-virulent CL-14 clone [52]. Overall, these studies pointed toward noteworthy but still controversial association between levels of cruzipain expression/activity and virulence, metacyclogenesis and cell infectivity of T. cruzi strains. Low levels of activity and transcripts of CATL-like in T. rangeli (rangelipain) were related to lack of pathogenicity and intracellular development [53]. However, a broad study demonstrated that isolates of divergent lineages of T. rangeli express high levels of enzymatic activity and transcripts of rangelipain [54].

The major cruzipain isoform has been the subject of extensive biochemical, structural and immunological studies. However, to date, sequences of cruzipain-encoding genes were not yet comparatively examined in T. cruzi isolates of all DTUs displaying different degrees of cellular invasion, virulence and pathogenicity. Despite limited to a few strains and lacking sequence analysis, studies suggested relevant diversity of cruzipain genes [55,56]. Genetic diversity and activity of enzymes from T. cruzi of all DTUs and T. cruzi-like species need to be investigated. An understanding of the expression, repertoires and evolutionary relationships of genes encoding cruzipain in T. cruzi of different DTUs and closest related T. cruzi-like species, and comparison with T. rangeli and T. b. brucei, which are phylogenetically distant from T. cruzi with life cycles (which are phylogenetically distant from T. cruzi and unable to develop within mammalian cells in vitro), whose life cycles differ in vertebrate hosts and vectors and differing in vertebrate hosts and vectors, can assist in clarifying the potential role of these enzymes in the host-parasite interactions, virulence and pathogenicity.

Comparative studies between closely related pathogenic (T. cruzi) and non-pathogenic trypanosomes (T. cruzi-like) can contribute for understanding the evolution of the pathogenicity and virulence of T. cruzi and can be helpful for successful design of control, diagnostic and genotyping strategies. With these purpose, our goals in this study were: to characterize the repertoire of genes encoding cruzipain from isolates of all T. cruzi DTUs and compare them with homologues from T. c. marinkellei, T. dimasi, T. rangeli and T. b. brucei; to analyze the relationships among these genes by network genealogies; to investigate their genomic organization by synteny analysis of loci containing cruzipain genes in trypanosome genomes; to assess the expression of cruzipain homologues by the non-pathogenic T. cruzi-like species by northern-blotting and proteolytic assays.

Results

Comparison of whole Genes Encoding CATL Enzymes in Distinct T. cruzi DTUs, T. dionisii, T. rangeli and T. b. brucei

Like mammalian papain-like enzymes, CATL enzymes of trypanosomatids are synthesized as inactive precursors, consisting of pro, pro, and catalytic domains (cd), and a C-terminal extension. Proteolytic cleavage of the N-terminal pro-domain generates the mature enzyme consisting of a cd domain and a C-terminal extension. Protolytic cleavage of the N-terminal pro-domain generates the mature enzyme consisting of a cd domain and a C-terminal extension of trypanosomatids [24,26]. To compare the entire cruzipain sequences (~450 amino acids), sequences from T. cruzi Sylvio X10.6 and G (TcI), Y and Esmeraldo c13 (TcII), M6241 c6 (TcIII), CL Brener (TcVI, one sequence from each the Esmeraldo-like and non-Esmeraldo-like haplotypes) and from Tcbat 1994 were aligned with homologous sequences from T. c. marinkellei 344 and Tc dimasi 211. Sequences
from non-Schizotrypanum species (T. rangeli and T. b. brucei) were included in the alignments [34]. Overall identities were high in the N-terminal region, either in pre- and pro-domains (~94 and 91%, respectively) and catalytic domains (~90%), and most variable in the C-terminal regions (~85%) (Fig. 1). As typically found in peptidases of Clan CA, which are targeted to intracellular compartments and secreted, all cruzipain and homologous genes have a signal peptide at their N-terminal region, as well as the catalytic triad of cysteine, histidine and asparagine residues (Cys25, His159 and Asn179) and the highly conserved Trp181. Important sites for autocatalytic cleavage, the motifs ERFFIN-like and GNFID-Ii-like of pre-domains are conserved in all trypanosomes. The clan CA is characterized by having substrate specificity defined by the S2 pocket. In cruzipain genes from all T. cruzi DTUs and homologues from T. c. marinkellei, the S2 subsites are conserved whereas in T. dionisii, T. rangeli and T. b. brucei divergent amino acids were found in these regions, suggesting differences in substrate specificities (Fig. 1).

Cruzipain amino acid sequences from T. cruzi isolates were relatively conserved in all domains. The ratio of non-synonymous (dN) to synonymous (dS) substitutions in the catalytic domain was dN/dS < 1 by comparing the distinct DTUs of T. cruzi and T. cruzi-like species, suggesting that the enzymatic domain of cruzipain genes has been subjected to stabilizing selection for the conservation of metabolic function within the subgenus Schizotrypanum. Sequences encoding homologous cruzipain genes of T. c. marinkellei were closely related to those of T. cruzi (~6.5% divergence) but the divergences were larger than those separating the T. cruzi DTUs (maximum of ~2.5%). Sequences from T. cruzi largely diverged from homologues of T. dionisii (~20%), T. rangeli (~33%) and T. b. brucei (~43%) in all domains (Fig. 1). Genealogies based on whole cruzipain genes, or restricted to pre-pro or to catalytic domains, resulted in identical topologies (Fig. 2).

Genealogy of Genes Encoding T. cruzi Cruzipain and Homologues from T. c. marinkellei, T. dionisii and Non-Schizotrypanum Trypanosome Species

To study the relationships of cruzipain genes from all T. cruzi DTUs, T. c. marinkellei and T. dionisii, and homologues from T. rangeli and T. b. brucei (Table 1), we aligned ~500 bp cdCATL sequences obtained in this study with the corresponding genes from T. cruzi CL Brener (Esmeraldo-like and non-Esmeraldo-like haplotypes), Esmeraldo c13, JR e4, M2149 c16 and Sylvio X106. The analyses of either amino acid (Fig. 3) or nucleotide sequences (data not shown) generated networks of cruzipain genes with similar topologies. Sequences of cd-cruzipain from all T. cruzi DTUs always clustered together in a homogeneous assemblage (~1.2% amino acid sequence divergence) separated from T. c. marinkellei (5.5% divergence), and largely divergent from the cluster formed by T. dionisii from Brazil (20%) and Europe (21%). Amino acid sequences of cruzipain genes of all Schizotrypanum species clustered together and largely separated (20%) from the homologous genes of T. rangeli and T. b. brucei (Fig. 3; Table 1).

Polymorphism of Cruzipain Gene Copies within Isolates and DTUs of T. cruzi

Any attempt to associate cruzipain polymorphisms with biological features of T. cruzi requires a good appraisal of the diversity of gene copies within both one strain/isolate and each DTU. We have assessed the polymorphism on cruzipain gene copies by comparing 3 to 8 sequences from each isolate. Larger number of sequences (7–8) was analyzed from the isolates of hybrid DTUs TcV and TcVI (Table 1). Cruzipain gene copies (paralogous) from isolates of TcI, TcIII, TcIV and Tcbat were identical or highly similar in their amino acid sequences (Fig. 3), whereas diverged in 2 to 6 polymorphic sites in their nucleotide sequences (Fig. 4). We identified a total of 23 variant sequences of cruzipain sequences (Fig. 4). Relatively homogeneous but not identical copies were found by comparing sequences from 6 isolates of TcI and two of TcII. No polymorphic sites were found among sequences from 3 isolates of each TcIII and TcV. Sequences from Non-Esmeraldo-like haplotype of CL Brener were identical to those of TcIII (Fig. 4).

Results disclosed high nucleotide polymorphism on cruzipain gene copies of the heterozygous hybrids assigned to TcV (3 isolates, 4 to 9 sequences of each) and TcVI (3 isolates, 24 sequences). Different sequences were found in TcV (at least 6 sequence types) and TcVI (8 types), including sequences identical to those found in TcII or TcIII, sequences found in both TcV and TcVI, and sequences so far detected exclusively in TcV or TcVI (Fig. 3, 4).

Distinguishing DTUs of T. cruzi according to Polymorphisms on Catalytic Domain of Cruzipain Genes

We evaluated the suitability of the polymorphism on nucleotide sequences of catalytic domain (cd) of cruzipain genes in distinguishing among the T. cruzi DTUs by comparing 80 sequences from 25 isolates of all DTUs (66 sequences from 22 isolates were determined in this study). Sequences differed by 17 polymorphic nucleotide sites and were divided in 5 major types in addition to the hybrid sequences from TcV and TcVI (Fig. 4). Despite polymorphisms within DTUs, unique polymorphic sites distinguished all DTUs. Cruzipain polymorphisms also distinguished Tcbat from all established DTUs (Fig. 4).

To evaluate the suitability of cruzipain polymorphisms as markers for genotyping, we examined at least three isolates from each DTU, excepting TcVI for which CL Brener and CL 14 were analyzed. Despite multiple copies and small polymorphisms among repeats within the same DTUs and even the same strain (Fig. 4), cruzipain encoding genes showed high sequence conservation of catalytic domains from isolates of the same DTU and clustering of sequences was according to DTUs. Thus, the cruzipain analysis agreed with genotyping methods based on either multiple copy or single copy gene markers [6]. In contrast, large polymorphism and unique sequences detected in each TcV and TcVI (Fig. 4) suggested that these markers can be valuable to detect hybrids. Nevertheless, the use of cruzipain genes for T. cruzi genotyping demands the sequencing of several PCR-amplified sequences and comparison of homologous sequences through phylogenetic analyses. A method based on PCR-RFLP analysis of cruzipain genes has been currently developed to facilitate the use of cruzipain as marker for T. cruzi genotyping [Lima et al., in preparation].

Genomic Organization and Synteny of Cathepsin-L Genes in Trypanosome Species

Previous studies showed that cruzipain genes are organized in the genomes of trypanosomes as tandem arrays of duplicated and polymorphic genes located in two or more chromosomes [25]. Analysis of the genomic organization of cruzipain genes in T. cruzi CL Brener Esmeraldo-like and non-Esmeraldo-like haplotypes disclosed polymorphisms in number and organization of genes encoding cruzipain (cruzipain, cruzipain 2 and other putative isoforms). There was substantial variation in number, sequence, chromosome and position of the duplicate genes of the two haplotypes; Esmeraldo-like showed 1–4 cruzipain repeats dis-
persed in three loci and non-Esmeraldo-like present 3–5 copies in three loci (data from TriTrypDB).

Figure 1. Alignment of predicted amino acid sequences from entire cruzipain of *T. cruzi* (Tcl, TcII, TclIII, TcVI and TcBat) and homologues from *T. cruzi*-like (*T. c. marinkellei* and *T. dionisii*), *T. rangeli* and *T. b. brucei*. Pre, pro, catalytic domain and C-terminal extension amino acid sequences of cruzipain genes from *T. cruzi* Sylvio X10.6 and G (TcI), TCC1994 (TcII), Y and Esmeraldo cl3 (TcII), M6241 cl6 (TcIII), CL Brener (TcVI) Non-Esmeraldo-like (TcIII) and Esmeraldo-like (TcII) haplotypes and homologues from *T. c. marinkellei* (344), *T. dionisii* (211), *T. rangeli* (LDG and AM80) and *T. b. brucei* (TREU 927). The CATL family signatures of pro-domain motifs ERFININ (ERFN) and GNFD (GTFD) are indicated in bold and underlined, the subsites S1, S2 and S2*9* are in bold, and the conserved Trp181 are indicated by (*). The glutamine [Q] of the oxyanion hole, cysteine [C], histidine [H] and asparagine [N] of catalytic triad in the catalytic domain, and 8 cysteines in the C-terminal extension are indicated by arrow heads. doi:10.1371/journal.pone.0038385.g001

Chromosome segments from the genome of *T. cruzi* CL Brener (TcVI) containing three and four tandem copies of cruzipain in Esmeraldo and non-Esmeraldo haplotypes, respectively, were...
compared with data from the genome drafts of *T. cruzi* G (TcI), M6241 cl6 (TcIII) and *T. dionisii*. Homologous segments of chromosome 6 from *T. cruzi* CL Brener containing cruzipain repeats were found in the genomes of other *T. cruzi* strains: one cruzipain gene copy from *T. cruzi* G (of three copies detected in the genome) and three from M6241 cl6 (6 copies in the genome). In the genomes of these strains, cruzipain and homologous genes were flanked by five orthologous genes thus constituting a syntenic block (Fig. 5). Homologous cruzipain genes arranged in the same order were detected in the genomes of *T. dionisii* (two out of three copies found in the genome draft) and *T. b. brucei* (an array of 11 identical copies of brucipain in the chromosome 1) (Fig. 5). This syntenic block was selected for this study considering the positioning of cruzipain gene copies from distinct *T. cruzi* strains and the synteny shared with *T. dionisii*, *T. marinkellei*, *T. rangeli* and *T. b. brucei* (Fig. 5) and other trypanosome species as showed with *T. vivax* and *T. congolense* genome drafts (data not shown).

Assembly of repetitive sequences in tandem arrays is very problematic, and both the copy number and position are very difficult to be accurately determined. Miss-assembly from collapsed repeat sequence frequently arises during automated genome assembly when small sequence reads originating from distinct repeat copies are incorrectly joined to generate a single unit. This artefact could not be ruled out in the genome drafts analyzed. To avoid mis-assembled genes, we selected contigs containing sequences from both cruzipain and adjacent orthologous genes, which warrant the positioning of the genes encoding cruzipain in this syntenic region (Fig. 5). The degrees of sequence identity between each syntenic gene (Fig. 5) agreed with the phylogenetic relationships among all trypanosome species investigated: *T. cruzi*, *T. c. marinkellei*, *T. dionisii*, *T. rangeli* and *T. b. brucei* [7,10,11,14,21,54].

Comparative Expression Analyses of Cruzipain and Homologues in *T. cruzi*-like and *T. rangeli* by Northern Blot Hybridization and Detection of Proteolytic Activity in Gelatin Gels

To assess the expression of cruzipain and homologous enzymes from virulent and non-virulent and human-infective or bat-restricted trypanosome species, developing or not within cells, total RNA from epimastigotes of *Schizotrypanum* species (*T. cruzi* Y, *T. c. marinkellei* and *T. dionisii*) and *T. rangeli* was compared by northern blot hybridization using *T. cruzi* Y derived catalytic domain probes.
Table 1. Trypanosoma cruzi isolates of all DTUs (TcI-TcVI) and other trypanosome species, and their respective sequences of cruzipain and homologous genes determined in this study or retrieved from data banks.

| TCC\(^a\) code | Trypanosome isolate | Host species | Geographic origin | DTU/ genotype | Accession number of cruzipain and homologous sequences | Origin of sequences \(d\) |
|----------------|---------------------|--------------|------------------|---------------|----------------------------------------------------------|---------------------|
| T. cruzi       |                     |              |                  |               |                                                          |                     |
| 1321           | Dm28                | opossum      | Colombia         | TcI           | JF421288/AF411439\(^{c}\)                                  | PCR                 |
| -              | Sylvio X10.6        | human        | Brazil           | TcI           | U41454\(^d\)                                              | GenBank             |
| -              | JR cl4              | human        | Venezuela        | TcI           | TJR4_1_c6805/712/TJR4_1_c13212\(^{c}\)                    | draft genome        |
| 30             | G                   | opossum      | Brazil           | TcI           | JF421297/AF411439\(^{c}\)                                 | draft genome and PCR|
| 417            | M2542               | bat          | Brazil           | TcI           | JF421292/AF411439\(^{c}\)                                 | PCR                 |
| 507            | MO115               | bat          | Brazil           | TcI           | JF421294/AF411439\(^{c}\)                                 | PCR                 |
| 34             | Y                   | human        | Brazil           | TcII          | AJ319429/AF411439\(^{c}\)                                 | GenBank and PCR     |
| 2120           | Esmeraldo cl3       | human        | Brazil           | TcII          | JF421314/AF411439\(^{c}\)                                 | TriTrypDB and PCR   |
| 844            | MT3869              | human        | Brazil           | TcII          | JF421335/AF411439\(^{c}\)                                 | PCR                 |
| 845            | MT3663              | triatomine    | Brazil           | TcII          | JF421337/AF411439\(^{c}\)                                 | PCR                 |
| 1386           | Unidero             | dog          | Brazil           | TcII          | JF421339/AF411439\(^{c}\)                                 | PCR                 |
| -              | M6241 cl6           | human        | Brazil           | TcII          | cCM62_C86\(^{c}\)                                         | draft genome\(^a\)  |
| 85             | JJ (Jose Julio)     | human        | Brazil           | TcIv          | JF421304/AF411439\(^{c}\)                                | PCR                 |
| 337            | Fusciocolis 15      | monkey       | Brazil           | TcIv          | JF421306/AF411439\(^{c}\)                                | PCR                 |
| 778            | Rb778               | triatomine    | Brazil           | TcIv          | JF421305/AF411439\(^{c}\)                                | PCR                 |
| 187            | Bertha              | human        | Bolivia          | TcV           | JF421316 - JF421319\(^{c}\)                              | PCR                 |
| 186            | Tc186               | triatomine    | Bolivia          | TcV           | JF421320 - JF421327\(^{c}\)                              | PCR                 |
| 967            | NR c13              | human        | Chile            | TcV           | JF421328 - JF421334\(^{c}\)                              | PCR                 |
| 33             | CL                  | triatomine    | Brazil           | TcVI          | JF421312/AF411439\(^{c}\)                                 | PCR                 |
|               | CL14                | triatomine    | Brazil           | TcVI          | JF421306/AF411439\(^{c}\)                                | PCR                 |
|               | CL Brener Esmeraldo and Non-Esmeraldo haplotypes unassigned contigs | triatomine | Brazil           | TcVI          | JF421312/AF411439\(^{c}\)                                 | PCR                 |
|               | -                   | T. infestans  | Brazil           | TcVI          | JF421306/AF411439\(^{c}\)                                | PCR                 |
| 294            | 998                 | bat          | Brazil           | Tcbat         | JF421296/AF411439\(^{c}\)                                | PCR                 |
| 499            | 1336                | bat          | Brazil           | Tcbat         | JF421298/AF411439\(^{c}\)                                | PCR                 |
| 1994           | MD294               | bat          | Brazil           | Tcbat         | JF421300/AF411439\(^{c}\)                                | draft genome and PCR|
| 1122           | 1122                | bat          | Brazil           | Tcbat         | JF421302/AF411439\(^{c}\)                                | PCR                 |
| T. cruzi marinkellei |                     |              |                  |               |                                                          |                     |
| 344            | bat                 | C. perspicillata | Brazil          |               | JF421354\(^{c}\)                                         | PCR                 |
| 501            | bat                 | C. perspicillata | Brazil          |               | JF421343\(^{c}\)                                         | PCR                 |
| 611            | bat                 | A. planirostris | Brazil          |               | JF421344\(^{c}\)                                         | PCR                 |
| TCC code | Trypanosome isolate | Host species | Geographic origin | DTU/genotype | Accession number of cruzipain and homologous sequences | Origin of sequences |
|----------|---------------------|--------------|-------------------|--------------|--------------------------------------------------------|-------------------|
| -        | P3                  | bat          | P. pipistrellus    | England      | JF421345<sup>5</sup>                                    | PCR               |
| 495      | bat                 | C. perspicillata | Brazil            | JF421346<sup>5</sup> |                                          | PCR               |
| 1098     | bat                 | Myotis sp    | Brazil            | JF421347<sup>5</sup> |                                          | PCR               |
| 454      | bat                 | D. rotundus  | Brazil            | JF421348<sup>5</sup> |                                          | PCR               |
| 211      | bat                 | E. brasiliensis | Brazil           | JF421355<sup>5</sup> |                                          | draft genome      |

**T. rangeli**

|            | Lineage<sup>e</sup> | |
|------------|----------------------|---|
| 643        | Tra643               | bat | P. luteus         | Brazil      | E | FJ977568<sup>8</sup> | GenBank          |
| 1719       | Tra1719              | bat | A. planirostris   | Brazil      | A | JF421351<sup>5</sup> | PCR              |
| 031        | SA                   | human | H. sapiens       | Colombia    | A | FJ977565<sup>8</sup> | GenBank          |
| 086        | AM80                 | human | H. sapiens       | Brazil      | B | JF421356<sup>5</sup> | draft genome     |
| 014        | PG                   | human | H. sapiens       | Panama      | C | FJ977564<sup>8</sup> | GenBank          |
| -          | LDG cl1              | human | H. sapiens       | Colombia    | C | L38512<sup>8</sup>  | GenBank          |

**T. b. brucei**

| -          | T. b. brucei TREU927 | tsetse fly | Glossina sp | Kenya | XM_840123<sup>5</sup> | GenBank          |

<sup>a</sup>TCC, Code number of the isolates/strains cryopreserved in the Trypanosomatid Culture Collection (TCC); Sequences from cruzipain and homologous genes:
<sup>b</sup>whole genes,
<sup>c</sup>catalytic domains;
<sup>d</sup>cruzipain sequences obtained by sequencing of PCR-amplified genes or from genome databases: GenBank, TriTrypDB and drafts genomes from our Tree of Life project or from the Washington University School of Medicine project<sup>*</sup>;
<sup>e</sup>lineages of *T. rangeli* defined by Maia da Silva et al. [22,61].

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Figure 3. Network and polymorphism analyses on catalytic domain of cruzipain genes from different trypanosome species. Genes from *Schizotrypanum* species (*T. cruzi*, *T. c. marinkellei* and *T. dionisii*) were compared with homologues from their closest relative species, *T. rangeli*, and the distant related *T. b. brucei*. (A) Network of 33 amino acid predicted sequences constructed using the Neighbour-Net algorithm excluding all conserved sites and with Uncorrected p-distance. The numbers in nodes correspond to bootstrap values from 100 replicates. (B) Polymorphism on cruzipain amino acid sequences from the distinct trypanosome species.

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Figure 4. Polymorphism and network analyses of catalytic domain sequences of cruzipain genes from *T. cruzi* isolates of TcI-VI and Tcbat. (A) Polymorphic nucleotide sites on catalytic domains of cruzipain encoding genes; (B) Network based on polymorphic nucleotides constructed with the Neighbour-Net algorithm excluding all conserved sites and with Uncorrected p-distance. The numbers in nodes correspond to bootstrap values from 100 replicates. CLBrener1-5 are sequences from TriTrypDB: Tc00.1047053509429.320, Tc00.1047053507537.20, Tc00.10470533537.20, Tc00.104705335327.20, Tc00.104705335307.20.
cruzipain probe that strongly cross-hybridized with RNA from other *T. cruzi* strains (G, CL and JJ) and from all other trypanosome species using low stringent conditions (data not shown). However, hybridization signals using more stringent conditions correlated very well with sequence identity displaying high cross-hybridization signal with transcripts of *T. cruzi*, moderate hybridization with *T. c. marinkellei*, weak hybridization signal with transcripts of *T. dionisii* and lack of hybridization with *T. rangeli* isolates. This analysis allowed estimation of transcripts of similar size (~1.9 Kb) for all the *Schizotrypanum* species (Fig. 6). In agreement with the differential cross-hybridization among the transcripts of different species, we previously demonstrated that a probe consisting of *T. rangeli* catalytic domain of CATL (rangelipain) strongly hybridized with *T. rangeli* while cross-hybridization with *Schizotrypanum* species was very weak [54].

Cruzipain, a glycoprotein of ~50–60 kDa, is post-transcriptionally regulated during the *T. cruzi* life cycle [4,24,26]. Here, proteolytic activities related to CATL were detected in epimastigote lysates of all *Schizotrypanum* species examined (Fig. 6B). In gels incubated at pH 5.0 with 5 mM DTT (a condition found to be...
optimal for the detection of cysteine proteases in these trypanosomes), remarkable high activities resolving in the region of ~55 to 35 kDa were detected in lysates of *T. c. marinkellei* and *T. dionisii*. Lower cruzipain activity was detected for *T. cruzi* compared to *T. cruzi* strains such as the virulent Y and CL (TcII) (data not shown). Weaker activity was detected for *T. cruzi* Y cruzipain labeled with 32P *T. cruzi* Y (TcY-CatL probe); agarose gel of RNA used for this analysis was stained with ethidium bromide (EtBr). (B) CATL proteolytic activities detected in epimastigote lysates of *T. cruzi* G, *T. cruzi* marinkellei, *T. dionisii* and *T. rangeli*. Activity banding profiles detected in gelatin gels, pH 5.0 and 5 mM DTT were inhibited in gel incubated with 10 µM E-64.

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**Discussion**

In this study, we compared cruzipain-encoding genes of *T. cruzi* isolates representatives of all known subspecific phylogenetic diversity, including all DTUs (TcI-TcVI) and the new *T. cruzi* genotype Tcbat, as well as homologous genes from *T. cruzi*-like species, *T. c. marinkellei* and *T. dionisii*, the closest relatives of *T. cruzi*. Results demonstrated that cruzipain genes from a large set of *T. cruzi* isolates representative of the overall biological and genetic diversity compared with homologous CATL genes from other species in a phylogenetic framework disclosed divergences that closely parallel their phylogenetic diversity [3–7,10,11,14,57]. Our findings revealed species-specific and DTU-specific variability that may be valuable for elucidation of the roles of cruzipain in host-parasite interactions, virulence and pathogenicity evolution.

Isolates of *T. cruzi* show a range of variation in important biological, immunological, pathological (morbidity and mortality) and clinical characteristics [5,6]. It is likely that much of this variation is probably due to genetic differences among isolates that can be related to specific DTUs. The genomes of the “non virulent” *T. cruzi* Sylvio X10/1 (TcI) and “virulent” *T. cruzi* CL Brener (TcVI) are highly similar in their gene-dense “core” coding regions, which show strongly conserved synteny interspersed with variable repetitive sequences [58]. Large differences within and among DTUs have emerged in several repetitive gene families such as cysteine proteases, mucins, trans-sialidases, surface protease gp63 and amastigote surface glycoproteins (amastins) [58–61].

In this study, a relevant conservation of amino acid sequences of genes encoding cruzipain was found among *T. cruzi* of all DTUs, Tcbat and *T. c. marinkellei* compared to more divergent sequences from *T. dionisii*. Cruzipain sequences from all these species tightly clustered together and were more related to rangelipain from *T. rangeli* than to brucipain of *T. b. brucei*. Notwithstanding hard efforts to find sequences homologous to cruzipain 2, no sequences were found that displayed the signature residues identifying this isoform [28]. Besides sequences homologous to major cruzipain, which comprised the largest part of sequences detected, our searches in the genome data banks and PCR-amplified sequences disclosed a few heterogeneous sequences closely related to cruzipain 2 [Lima et al., in preparation] but lacking the signature residues reported as characteristics for the archetype of this isoform. Variations in residues in the S2 pocket may account for peculiar activity and substrate specificity of cruzipain isoforms, hence, detailed biochemical studies will be necessary to assess expression, activity and functions in order to verify whether these sequences encode new isoforms or variants of cruzipain 2. In any case, our findings confirmed that phylogenetic analyses are valuable to discover new cruzipain isoforms and/or variants as recently shown for congopain variants through combined genetic and functional approaches [62].

In network genealogies, sequences of cruzipain from the same species, as well as from strains/isolates of the same phylogenetic lineages, always clustered together. Even with the high degree of conservation in cruzipain genes from all DTUs, polymorphisms on nucleotide sequences from the catalytic domain of cruzipain generated 5 branches within *T. cruzi*, each one comprising sequences from one DTU (TcI-TcVI) or Tcbat. In general, sequences from cruzipain gene copies within the same DTU showed small or no variation at all, as we verified for isolates of TcI-TcIV. Data from these DTUs were consistent with their epidemiology and evolutionary histories, with distances among cruzipain genes suggesting that they have had more time segregated from each other than the more recently emerged
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TeV-TeVI. Isolates of the same DTUs appear to have had a long association with preferential mammalian hosts and vectors and, consequently, naturally circulate separated by biological, ecological and geographical barriers [5–7,63,64]. TeV and TeVI were confirmed as formed by heterozygous isolates exhibiting polymorphic cruzipain sequences, in addition to sequences identical to those of putative donors TcII or TcIII. In agreement with their hybrid origin, sequences from TeV and TeVI clustered with TcII or TcIII forming a reticulate pattern in the network genealogy of cruzipain genes. Tandem arrays of cruzipain genes in T. cruzi CL Brener genome showed sequences apparently derived from TcII (Esmeraldo-like) and TcIII (Non-Esmeraldo-like), even within a single locus. Unique sequences detected in TeV or TeVI strains could be due to the fact that these strains originated from hybridization between strains of TcII and TcIII not included in this study. In addition, more heterogeneous and unique sequences within hybrid strains could emerge post-hybridization through homologous recombination. Our results corroborate that TeV and TeVI resulted from the hybridization between TcII and TcIII as hypothesized before using other markers. Studies have been performed to better understand the role of hybridization events in shaping the genetic diversity within T. cruzi [65–67]. Our findings confirmed Tcbat as a new T. cruzi genotype not yet assigned to any DTU. Sequences from Tcbat clustered closest to TcI but separated from all DTUs as demonstrated with other markers [6,7,68].

Genealogy of cruzipain and homologous genes inferred in this study confirmed T. c. marinkellei isolates as closest relative and outgroup for T. cruzi isolates, as previously shown by examination of other genes including SSUrRNA, gGAPDH and Cytb [10,14,22]. In a comparative proteomic analysis, T. c. marinkellei was indistinguishable from T. cruzi reinforcing their very close relatedness [69]. To date, Tcbat, T. c. marinkellei and T. dionisi have been found exclusively in bats and all invade and develop in culture cells of a variety of mammals. Tcbat was infective to mice, despite very low virulence. However, these three bat trypanosomes have been reported to be incapable of development in triatominine species commonly infected by T. cruzi [7–10,14,16].

Evolutionary studies of cruzipain and homologous genes from other trypanosome species corroborated phylogenetic relationships based on SSUrRNA and gGAPDH genes of all species investigated: T. carassii from fish [70], T. rangeli [54], T. vivax [71] and T. theileri [72,73], the later three from mammals. Evolutionary relationships of cruzipain and homologues from these trypanosome species indicated that this gene family expanded by successive gene duplications followed by divergences giving rise to genes with greater similarity when originated from the same rather than distinct species/genotypes. This suggests that concerted evolution is a widespread homogenizing force in this trypanosome multigene family. Consequently, CATL-like gene duplicates (paralogous) of all trypanosome species examined clustered by species, i.e. like orthologous genes in different species, as shown for several repeated genes in tandem arrays from African trypanosomes [74]. However, gene conversion and positive selection can generate diversity in sequence, quantity and order among tandemly repeated genes as showed in this and in previous studies on trypanosomatids [59–61,74].

Results from this study disclosed molecular markers able to identify the phylogenetically closely related T. cruzi and T. cruzi-like species. Reliable identification and knowledge of the diversity and epidemiology of these species are crucial to understand the shared evolutionary history of T. cruzi and T. cruzi-like bat trypanosomes [7,10,11,14,57]. We have previously shown that T. cruzi and T. rangeli [54], T. vivax [71] and T. theileri [72] could be diagnosed by specific PCR assays targeting cruzipain sequences. In this study, we demonstrated that cruzipain sequences clustered T. cruzi isolates according to DTUs, similarly to CATL-like gene based genotyping of lineages within T. rangeli [54], T. vivax [71] and T. theileri [72,73]. Altogether, these previous studies and results herein described support the use of these genes as valuable markers for both inter- and intra-species phylogenetic analyses of trypanosomes.

There are clear evidence that immunization with cruzipain catalytic domain, and not with C-terminal domain, confer important cellular protective immunity against T. cruzi, as evidenced by the reduction in parasitemia, tissue parasitism and mortality in mice, indicating the existence of epitopes important for protective immunity in the catalytic domains and, thus, supporting the use of cruzipain for vaccine development [34–37]. The main goal of all vaccination strategies should be a polyclonal vaccine against infection by different strains of T. cruzi of any DTU. Although the polymorphism among cruzipain catalytic domains expressed by distinct strains appears to be reduced, for vaccination purposes it is important to evaluate strain-variant and conserved cross-reactive epitopes in different T. cruzi DTUs, and among strains of the same DTU, and the possible relevance of epitope polymorphism for species- or strain-specific protective immune response [75–77]. Therefore, knowledge on the genetic diversity within the plethora of DTUs and strains of any antigen vaccine candidate for Chagas disease can be decisive for the design of efficient and polyvalent vaccines using, if necessary, a pool of stage- and strain-specific antigens. Here, we showed for the first time that all the six DTUs, and even different strains of DTUs, exhibited specific genetic variants of cruzipain. Any possible implications of the cruzipain polymorphisms to warrant efficient cross-protection of a potential vaccine require investigations to demonstrate the immunogenicity of cruzipain variants from different T. cruzi DTUs.

Drugs currently employed for the treatment of Chagas disease have serious limitations due to their failure in chronic patients and side effects. Inhibitors of cruzipain kill the parasite and cure infected mice, thus validating this enzyme as a very promising target for design of new drugs [1,2,4,3,4,4]. In this study, we demonstrated that substrate-binding subsites of cruzipain are conserved in all T. cruzi DTUs, but diverge in T. dionisi and other trypanosome species. T. c. marinkellei cruzipain share identical subsites and, hence, can be a valuable non-infective to human model to test drugs against trypomastigotes and intracellular amastigotes. The efficacy of the cruzipain inhibitor K777 was evidenced against 6 T. cruzi strains differing in tissue tropisms and drug susceptibility [44]. More strains must be tested to verify the efficacy of inhibitors for the ample diversity of cruzipain variants expressed by T. cruzi of the different DTUs.

Characterization of genes encoding major cruzipain, genealogies of genetic variants and genomic organization were addressed in this study for the first time to compare T. cruzi of all DTUs and closest related T. cruzi-like species. This is the most comprehensive study using sequences from protein-encoding genes to compare isolates of T. cruzi from all DTUs. Of the strains selected for genome sequencing CL Brener and Esmeraldo are virulent whereas Sylvio X10.6 and G strain are considered of low virulence. Most T. cruzi isolates examined here by PCR- amplification of cruzipain genes were also previously analysed regarding virulence for mice: TeV, Tcbat and TeVI isolates were non-virulent whereas TeCIII (TCC1386) and TeV (TCC197) isolates showed to be virulent for mice (data not shown). Interestingly, non-virulent CL14 clone was shown in this study to be a hybrid strain like the virulent CL Brener [32]. All the six
DTUs were supported by cruzipain gene markers. Infections caused by different strains of *T. cruzi* from distinct DTUs extensively diverged in the morbidity and mortality. Attempts to associate *T. cruzi* DTUs with behavioral phenotypes, including virulence and pathogenicity in mice, metacyclogenesis and cell infectivity, suggested an important degree of association [5,6,60]. However, besides some overlap between different DTUs, there is a broad intra-DTU phenotypic diversity (largest differences have been reported within *TcI*) and even among clones of a given strain such as the virulent CL Brener and non-virulent CL14 clones, both of *TcVI*. Therefore, we can associate cruzipain polymorphisms to DTUs. However, available data are insufficient to support strong correlations between behavioural phenotypes and DTUs. This goal will require extensive studies of *T. cruzi* strains from all DTUs through a combination of phylogenetic, biochemical, pathological and immunological approaches.

Our findings indicated conserved major cruzipain in *T. cruzi* of all DTUs while other trypanosome species express diverse homologous enzymes. Results showed, for the first time, the expression of cruzipain transcripts by *T. c. marinkellei* and *T. dionisii* that correlated very well to sequence divergences among all trypanosome species investigated and, in addition, revealed high proteolytic activity in these two non-pathogenic species. An understanding of the cruzipain gene repertoires, expression and functions can help to elucidate the evolutionary history that shaped variability within *T. cruzi* and its divergence from *T. cruzi*-like species and the more distantly related *T. rangeli*. *T. cruzi*-like trypanosomes, despite sharing genomic and proteomic features [10,11,14,69], exhibit many peculiarities that can potentially explain differences in host preference/restiction, virulence and pathogenesis. Results from this comprehensive study on major cruzipain isoform are the initial steps toward understanding the roles played by genetic repertoires of cruzipain enzymes and homologues in the life cycles and infections caused by *T. cruzi* of all DTUs, *T. cruzi*-like species and *T. rangeli*.

**Materials and Methods**

**PCR Amplification, Sequencing and Phylogenetic Analysis of CATL Sequences**

The whole sequences of cruzipain genes (~1.4 Kb) of *T. cruzi* Esmeraldo c3 and CL Brener were retrieved from TriTrypDB. Homologous genes in *T. cruzi* M6241 c6 (Project ID:59941) were obtained from the genome drafts produced by the Genome Institute at Washington University School of Medicine (St. Louis, USA). *T. cruzi* G, Tcbat 1994, *T. dionisii* 211 and *T. rangeli* AM80 were obtained from genome drafts that we are currently performing using standard pyrosequencing shotgun methodology according to Roche 454 protocols as described previously [78]. Resulting reads were submitted to Roche’s Newbler software according to Roche 454 protocols as described previously [78].

**Syntenic genes flanking CATL genes were identified in the genomes of *T. cruzi* CL Brener (Non-Emesraldo-like and Emesraldo-like haplotypes) and *T. b. brucei* 927 genomes (TriTrypDB). Scaffolds comprising cruzipain genes from *T. cruzi* G and M6241 c6 and homologues from *T. dionisii* were aligned with these syntenic segments. Search on the draft genome of Sylvio X10/1 [58], Tcbat 1994 and *T. rangeli* AM80 disclosed cruzipain sequences, most partial, in small reads that could not be positioned in homologous scaffolds. The ratio of non-synonymous to synonymous (dN/dS) amino acid changes was calculated according to Yang and Nielsen [81] using PAML, v.4.2 software to infer relative selection pressures [82].**

**Northern Blot and Proteolytic Activities of Cruzipain**

For Northern blot analysis, 10 pg of total RNA extracted using Trizol (Gibco BRL) from epimastigotes of *T. cruzi* Y, *T. c. marinkellei* 344, *T. dionisii* 211 and *T. rangeli* isolates AM80 and SA was electrophoresed in 1.0% agarose gel and blotted onto nylon membrane. The membrane was hybridized for 14–16 hr at 40°C with a probe consisting of PCR-amplified cd-cruzipain of *T. cruzi* Y labeled with 32P and washed in 1×buffer (0.3 M NaCl, 0.3 mM Na citrate, pH 7.0 containing 0.1% SDS) at 50°C for 1 h as described previously [54].

To assess the proteolytic activities in gelatin gels, lysates of cultured epimastigotes were subjected to electrophoresis in 10% resolving SDS-acrylamide gels containing 500 μg/ml gelatin as described before [54,72]. Briefly, gels were incubated in 2.5% (v/v) Triton X-100 in 0.1 M buffer (acetate; pH 4.0 and 5.0; Tris-HCl; pH 7.5) containing 5 mM DTT (dithiothreitol) and then, for ~18 h in buffer-DTT. Maximum activity was detected in pH 5.0 and was greatly stimulated by DTT. Bands associated to cysteine proteases were identified by incubating gel halves with 10 μM E-
64. Gels were fixed with 10% TCA, and stained with Coomassie blue R-250 [54,72].

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Author Contributions
Conceived and designed the experiments: MMGT LL SCA. Performed the experiments: LL PAO AFC GAB JMBA MG9. Analyzed the data: LL PAO FMS JMBA SCA. Wrote the paper: MMGT LL. Input into design of experiments and advice in manuscript preparation: SCA JMBA GAB.

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