Intraspecies Variation in *Trypanosoma cruzi* GPI-Mucins: Biological Activities and Differential Expression of α-Galactosyl Residues

Rodrigo P. Soares,* Ana C. Torrecilhas, Rafael R. Assis, Marcele N. Rocha, Felipe A. Moura e Castro, Gustavo F. Freitas, Silvane M. Murta, Sara L. Santos, Alexandre F. Marques, Igor C. Almeida, and Alvaro J. Romanha

Centro de Pesquisas René Rachou, FIOCRUZ, Belo Horizonte, Minas Gerais, Brazil; Departamento de Ciências Biológicas, Universidade Federal de São Paulo, Diadema, São Paulo, Brazil; The Border Biomedical Research Center, Department of Biological Sciences, University of Texas, El Paso, Texas

Abstract. The glycosylphosphatidylinositol (GPI)-anchored mucins of *Trypanosoma cruzi* trypomastigotes play an important immunomodulatory role during the course of Chagas disease. Here, some biological activities of tGPI-mucins from four *T. cruzi* isolates, including benzimidazole-susceptible (BZS-Y), benzimidazole-resistant (BZR-Y), CL, and Colombiana, were evaluated. GPI-mucins were able to differentially trigger the production of interleukin-12 and nitric oxide in BALB/c macrophages and modulate LLC-MK2 cell invasion. The significance of these variations was assessed after analysis of the terminal α-galactosyl residues. Enzymatic treatment with α-galactosidase indicated a differential expression of O-linked α-galactosyl residues among the strains, with higher expression of this sugar in BZS-Y and BZR-Y *T. cruzi* populations followed by Colombiana and CL. Unweighted pair group method analysis of the carbohydrate anchor profile and biological parameters allowed the clustering of two groups. One group includes Y and CL strains (*T. cruzi* II and VI), and the other group is represented by Colombiana strain (*T. cruzi* I).

INTRODUCTION

Chagas disease, a neglected illness caused by *Trypanosoma cruzi*, affects ~15 million people in 18 countries in Latin America. It is transmitted by hematophagous triatomine bugs that are widely distributed from southern regions of United States to northern parts of Argentina. Also, congenital, trans- and oral transmissions occur. Its clinical manifestations initiate with a short acute phase characterized by high parasitemia and various symptoms such as myocarditis. After the development of an acquired immune response during the chronic phase, the parasite numbers decrease in the bloodstream and tissues, remaining very low for most Chagasic patients. During this phase, some symptomatic forms appear in the heart and digestive organs. Chagasis disease chemotherapy is limited to two drugs, nitrofuran nitifurmitox (NFX) and 2-nitrimidazole benzimidazole (BZ). Both drugs present low cure rates in chronic patients and considerable side effects. Differences in the susceptibility to BZ and NFX among *T. cruzi* strains and the genetic background of the host may explain variations in therapeutic success.

During early stages of infection, the pattern of susceptibility/ resistance may be determined before acquired immunity, where innate immune mechanisms are crucial for parasite control. *T. cruzi* employs a highly elaborated array of molecules and strategies to invade a wide range of host cells and escape from host’s immune defense mechanisms. In the site of infection, *T. cruzi* triggers the production of chemokines, proinflammatory cytokines (interleukin-12 [IL-12] and tumor necrosis factor-α [TNF-α]), and reactive oxygen (ROI) and nitrogen (RNI) intermediates by cells from the macrophage lineage. Glycosylphosphatidylinositol (GPI) anchors expressed in the surface of *T. cruzi* such as the GPI-mucins and the glycoinositolphospholipids (GIPLs) are determinant in this process. The GPI-mucins of *T. cruzi* are composed of two subfamilies (TcMUC and TcSMUG), with a total of 863 gene members clustered with other multigene surface protein families. The major subfamily of TcMUC (i.e., TcMUC II [844 gene members]) is mainly expressed in the mammalian trypomastigote stage. It consists of a highly antigenic coat, with variations that account for interstrain features, such as virulence and immunomodulatory properties. Early studies showed that *T. cruzi* glycoconjugates are involved in attachment/invasion of hosts cells, escape from host immunity, and induction of protective lytic antibodies. Most of the studies focused in the host innate immune response used the GPI-mucins from the Y strain of *T. cruzi*. In contrast to epimastigote- or metacyclic-derived GPI-mucins (eGPI- and mGPI-mucins, respectively), GPI-mucins and their GPI-moieties alone have a potent proinflammatory activity in murine and human macrophages. This activity is probably caused by the presence of an unsaturated fatty acid (C18:1 or C18:2) in the phosphatidyl-inositol (PI) moiety (Figure 1). Finally, a heterodimer composed by Toll-like receptor 2 (TLR2) and TLR6 is involved in the recognition of *T. cruzi* tGPI-mucins.

Biochemically, the GPI moiety of *T. cruzi* mucins has the conserved core of Manα1-2-Manα1-2-Manα1-6-Manα1-4GlcNAc1-6-α-galactosyl-1-hydroxyinositol (HPO₄), which can be extended by phosphorylated substituents, including ethanolamine-phosphate (EtNP), 2-aminoethylphosphonate (2-AEP), and extra carbohydrate residues. The EtNP or 2-AEP group serves as the point of attachment for the surface mucin glycoprotein moiety (Figure 1). The hydrophobic tail linked to the myo-inositol phosphate consists of an alkylacylglycerol or ceramide lipid. The GPI-mucins of *T. cruzi* are heavily O-glycosylated with glycan chains containing N-acetylgalcosamine (GlcNAc) and β-galactose (β-Gal) residues, which are acceptors for sialic acid (Neu5Ac) units transferred by a trans-sialidase (TS) reaction. One of the features of tGPI-mucins that distinguishes them from the other parasite stages is the presence of terminal α-galactopyranosyl (α-Gal) residues in O-glycans attached to the polypeptide core. These epitopes are highly immunogenic and elicit anti-α-galactosyl antibodies (anti-α-Gal). These antibodies have protective lytic activity against trypomastigotes and are determinant in both acute and chronic phases of Chagas disease. The O-linked sugar structures in the GPI-mucins have already been fully characterized...
in epimastigotes and metacyclic stages of Colombian, Y, and CL 
strains.22–25 But only partially defined in trypomastigotes of the Y strain.15 Currently, after a consensus, the molecular studies enabled the classification of T. cruzi strains into six (TcI–TcVI) discrete typing units (DTUs).26 The biological activities of GPI-mucins from different DTUs are still mostly unknown. In our study, the strains analyzed (Colombian, Y, and CL) belong to T. cruzi DTUs I, II, and VI, respectively. Here, those features were explored in four T. cruzi strains/isolates (BZ-resistant population [BZR-Y], BZ-susceptible Y strain [BZS-Y], CL, and Colombiana) during interaction with macrophages and LLC-MK2 cells. Moreover, intraspecies variation in the terminal α-galactosyl residues in the O-linked glycans was assessed.

MATERIALS AND METHODS

T. cruzi strains and mammalian cells. The four populations/strains of T. cruzi used in this study are listed in Table 1 and were not cloned. BZR-Y was derived from the BZS-Y after in vivo selection after 25 successive passages in mice treated with a single high dose of BZ (500 mg/kg).6 CL and Colombian strains were susceptible and naturally resistant to BZ, respectively.5 All T. cruzi strains had been classified previously as belonging to T. cruzi DTUs I, II, and VI.26 Epimastigote forms were maintained at 28°C in liver infusion tryptose (LIT) medium supplemented with 10% fetal bovine serum (FBS; Cultilab, Campinas, SP, Brazil).27 Mammalian tissue culture-derived trypomastigotes (TCTs) from the four T. cruzi samples were obtained after infection of green monkey (Rhesus) kidney LLC-MK2 epithelial cells obtained from America Type Culture Collection (Manassas, VA).28 Parasite cultures and cells were tested for Mycoplasma as previously described.29

Extraction and purification of GPI-mucins. T. cruzi GPI-mucins from trypomastigote and epimastigote forms were purified as described (Figure 2A).15 Purified GPI-mucins were analyzed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) and silver staining.30 Protein concentration was determined at 214 nm using bovine serum albumin (BSA; Sigma, St. Louis, MO) as standard.

Purification of murine peritoneal macrophages and cytokine and nitrite measurements. The animals were kept in the Animal Facility of the Centro de Pesquisas René Rachou/FIOCRUZ in strict accordance to the Guide for the Care and Use of Experimental Animals.30 All animals and experiments were approved by the Ethical Committee of Animal Handling (CEUA) from Oswaldo Cruz Foundation (FIOCRUZ protocol P-0289/06). Thioglycollate-elicited peritoneal macrophages were removed from 8- to 11-week-old female BALB/c mouse by peritoneal washing and enriched by plastic adherence. Cells (5 × 10⁶ cells/mL) were kept in RPMI 1640 (Invitrogen, Carlsbad, CA) supplemented with 10% FBS (37°C, 5% CO₂; Cultilab, Campinas, SP, Brazil).31 Peripheral blood mononuclear cells were primed with 50 U/mL recombinant interferon-γ (IFN-γ; 18 hours, 37°C, B&D Biosciences, San Jose, CA) and subsequently incubated with GPI-mucins (1 and 10 μg/mL) from BZS-Y, BZR-Y, CL, and Colombian strains. Supernatants

![Figure 1. Schematic representation of T. cruzi GPI-mucins from different lifecycle stages.](image)

**Table 1. T. cruzi strains/populations analyzed in this study.**

<table>
<thead>
<tr>
<th>T. cruzi</th>
<th>Origin*</th>
<th>Host</th>
<th>Susceptibility†</th>
<th>Group‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>Colombiana</td>
<td>Col</td>
<td>Human (chronic patient)</td>
<td>R</td>
<td>I</td>
</tr>
<tr>
<td>BZS-Y</td>
<td>SP</td>
<td>Human (acute case)</td>
<td>S</td>
<td>II</td>
</tr>
<tr>
<td>BZR-Y</td>
<td>SP</td>
<td>Human (acute case)</td>
<td>R</td>
<td>II</td>
</tr>
<tr>
<td>CL</td>
<td>RS</td>
<td>Triatoma infestans</td>
<td>S</td>
<td>VI</td>
</tr>
</tbody>
</table>

*Brazilian states are Rio Grande do Sul (RS) and São Paulo (SP); Col = Colombia.
†In vivo drug susceptibility to benznidazole as described.5 R = resistant; S = susceptible.
‡Classification as reported.26
were collected after 48 hours, and IL-12 (subunit p40) cytokine measurement was performed by sandwich enzyme-linked immunosorbent assay (ELISA) according to the manufacturer’s instructions using the Opteia kit (B&D Biosciences). Nitric oxide (NO) production was measured indirectly by Griess reaction.32 All experiments were performed in triplicate. Negative controls were incubated with medium with IFN-γ (50 U/mL) (BD Biosciences, San Jose, CA) only.

Infection of LLC-MK2 epithelial Cells. LLC-MK2 cells were grown in a 24-well dish and washed two times with Dulbecco’s Modified Eagles Medium (DMEM; Invitrogen, Carlsbad, CA). Cells were infected with tGPI-mucins (1 ug/mL) from BZS-Y, BZR-Y, CL, and Colombiana strains in DMEM supplemented with 10% FBS (Cultilab, Campinas, Brazil) for 24 hours. Negative controls were incubated with DMEM medium only. After incubation, these cells were infected with TCT from the BZS-Y population with a multiplicity of infection (MOI) of 10 for 1 hour at 37°C, and then, they were washed two times with DMEM without serum. Non-adherent parasites were removed by the addition of Lymphoprep (Axis-Shield, Norton, MA) to the cell layers followed by two washes with DMEM without serum. Cells were incubated with DMEM with 10% FBS (24 hours, 37°C), fixed with 4% methanol and acetone and stained with Hoescht fluorescent dye (Molecular Probes; Invitrogen, Carlsbad, CA).

Chemiluminescence immunoassay and specificity. To ascertain the level of α-galactosylation in the O-linked sugars in the protein motif (Figure 2A), tGPI-mucins were subjected to chemiluminescent CL-ELISA against sera containing α-galactosyl antibodies purified from Chagasic patients.33 Briefly, 96-well ELISA microplates (NUNC; Thermo Scientific, Rochester, NY) were sensitized overnight at 4°C with tGPI-mucins (5 μg/mL) from all strains in 100 mM carbonate-bicarbonate buffer (pH 9.6). To avoid nonspecific binding in
the assay, free-wells sites were blocked for 2 hours at 37°C with 1% BSA in phosphate buffered saline (PBS/BSA). Two-fold serial dilutions of primary purified anti-α-Gal antibody, obtained as described, were made in wash buffer (1X PBS 0.05% Tween 20, pH 7.4), added to antigen-coated wells, and incubated for 1 hour at 37°C. Wells were washed three times, and plates were incubated (1 hour, 37°C) with secondary donkey anti-goat immunoglobulin G (IgG)-biotin conjugate (1:2,000; eBioscience, San Diego, CA) in wash buffer. Wells were washed five times, and streptavidin-peroxidase (1:2,000; eBioscience, San Diego, CA) was added for 1 hour at 37°C. In parallel, for the anti-α-Gal antibody specificity, tGPI-mucins were treated overnight at 28°C with 0.1 U/well coffee beans α-galactosidase (Sigma, St. Louis, MO) before antibody incubations. The reaction was developed using luminol (ECL, Thermo Scientific, Pierce Protein Research Products, Rockford, IL) diluted in 100 mM carbonate-bicarbonate buffer (pH 9.6). The intensity of light emission was expressed in relative luminescence units (RLUs) using a Luminoskan luminometer (Tecan). Negative controls were incubated with secondary antibodies only.

**Statistical analysis.** For the biological studies, the following statistical analysis was performed. The Kolmogorov–Smirnov test was used to test the null hypothesis that the data were sampled from a Gaussian distribution. A non-parametric Kruskal–Wallis test and analysis of variance (ANOVA) were performed to test equality of population medians among groups and independent samples. When samples deviated from Gaussian distribution, a non-parametric Kruskal–Wallis test was performed to compare two independent samples. Data were analyzed by GraphPad Prism 4.0 software, and P < 0.05 was considered significant.

**Purification of GPI-anchors from epimastigotes.** The biological experiments described above used purified GPI-mucins from trypomastigotes (tGPI). The unweighted pair group method analysis (UPGMA) analysis of the carbohydrate profile was based only in the epimastigote anchors (eGPIs) for comparison with reported DTUs.26 eGPI-mucins were subjected to Proteinase K treatment (5 mg/mL; Qiagen Inc., Valencia, CA) in 10 mM Tris-HCl, pH 7.8 (18 hours, 37°C) (Figure 2B). The material was heated (100°C, 5 minutes) to deactivate enzyme and subjected to butanol:water partition (2:1). The remaining GPI-anchors in the butanolic phase were subjected to nitrous acid deamination (300 μL 0.25 M sodium acetate and 300 μL 0.5 M NaNO₂) for 16 hours at 37°C to release the lipid. The suspension was then dried and subjected again to butanol:water partition, and polysaccharides were recovered in the aqueous phase. The oligosaccharides were desalted by passage through a two-layered column of AG50W-X12 over AG1-X8 (acetate; Bio-Rad, Hercules, CA). Anchor oligosaccharides were fluorescently labeled with 0.05 N 8-aminonaphthalene-1,3,6-trisulfate (ANTS; Sigma) and 1 M cyanoborohydride (16 hours, 37°C; Sigma) and subjected to fluorophore-assisted carbohydrate electrophoresis (FACE). Samples were run in polyacrylamide gels and visualized by an ultraviolet (UV) imager.14

**Unweighted pair group method analysis.** All strains/populations used in this work were previously characterized as *T. cruzi* I, II, or VI (Table 1).26 To evaluate if such discrimination would be observed in this work, a taxon/character matrix was constructed based on the presence/absence of bands from FACE analysis and biological techniques performed.15 The relationship among the strains was determined after a phenogram construction using the Dice similarity coefficient16 from the NTSYSpc software package (version 2.02).

**RESULTS**

**Purification of GPI-mucins.** *T. cruzi* GPI-mucins from either epimastigotes or trypomastigotes were extracted and purified from different strains/populations (Table 1). Fractions eluted from Octyl-Sepharose were resolved by SDS-PAGE and silver-stained. All GPI-mucins were eluted between 30% and 50% propan-1-ol gradient in 0.1 M ammonium acetate as previously described15 (data not shown). The GPI-mucins containing fractions were pooled, dried, and resuspended in sterile and lipopolysaccharide-free water before biological and immunological assays. Due to amount limitation, only GPI-mucins from epimastigotes (eGPIs) were used for UPGMA analysis (described later).

**Cytokine and nitrite levels on supernatants of stimulated macrophages.** Previous studies16,31,37 have shown the production of NO and proinflammatory cytokines by macrophages after exposure with either parasites or tGPI-mucins of *T. cruzi* (Y strain). In our work, we extended the number of the strains, including an *in vivo*-induced BZR-Y, its susceptible counterpart BZS-Y, and Colombiana and CL strains.
There were considerable differences in the NO and IL-12 production after exposure to tGPI-mucins from all isolates in the concentration of 1 μg/mL (\(P < 0.001\) and \(P < 0.01\), respectively). A higher production of NO and IL-12 was observed for the BZR-Y population compared with the other \textit{T. cruzi} samples (Figure 3A and B). In the concentration of 10 μg/mL, those differences were abolished (\(P > 0.05\); data not shown).

**LLC-MK2 invasion after tGPI-mucins exposure.** \textit{T. cruzi} trypomastigotes are able to release extracellular vesicles containing Tc85, TS, and tGPI-mucins that increase parasite invasion and cardiac tissue lesions in mice.\(^\text{38}\) To evaluate the ability of purified tGPI-mucins in modulating invasion of LLC-MK2 cells by trypomastigotes, the cells were pre-incubated with the glycoconjugates (1 μg/mL) from each strain/population for 24 hours. Trypomastigotes were able to invade and multiply inside LLC-MK2 cells (Figure 4A). After acquisition of pictures and evaluation of 10 fields per experiment, it was observed that the cells pre-incubated with tGPI-mucins of BZR-Y and CL had a higher infection rate than those cells pre-treated with tGPI-mucins from BZS-Y, Colombiana, and control (\(P < 0.02\), \(t\) test) (Figure 4B). Moreover, a slight increase in the number of intracellular amastigotes (~0.5-fold) was noted for the strains BZS-Y, BZR-Y, and CL compared with control (\(P < 0.05\), \(t\) test) (Figure 4C).

**Reactivity with anti-α-Gal antibodies from chronic Chagasic patients.** As observed above, tGPI-mucins from BZR-Y, BZS-Y, Colombiana, and CL strains had differential biological and immunological activities. Next, we evaluated whether these tGPI-mucins could also be differentially recognized by Chagasic anti-α-Gal antibodies. For this evaluation, tGPI-mucins were immobilized in CL-ELISA microplate and subjected or not to α-galactosidase treatment to remove terminal α-Gal residues and then reacted with Chagasic anti-α-Gal antibodies.

---

**Figure 4.** Cell invasion by \textit{T. cruzi} trypomastigotes (Y strain) previously incubated with GPI-mucins from trypomastigotes (tGPI-mucins). Negative controls (CTRLs) were incubated with DMEM medium only. (A) Intracellular amastigotes are indicated by arrows. (B) Percentage of LLC-MK2-infected cells and (C) number of intracellular amastigotes of \textit{T. cruzi} previously incubated with tGPI-mucins. Values are means for two experiments in triplicate. Bars represent standard errors. Definitions of abbreviations are in Figure 3.

**Figure 5.** Reactivity of tGPI-mucins from different \textit{T. cruzi} strains treated/untreated with α-galactosidase and incubated with purified α-galactosyl antibodies from Chagasic patients. Negative controls (CTRLs) used secondary antibodies only. BZR = benznidazole-resistant population from Y strain; BZS = benznidazole-susceptible population from Y strain; CL = CL strain; Col = Colombiana strain; RLU = relative luminescent unit; − = untreated tGPI-mucins; + = tGPI-mucins treated with α-galactosidase.
antibodies (Figure 5). Higher reactivity was observed for BZS-Y and BZR-Y followed by Colombiana and CL ($P < 0.0001$, ANOVA). A significant decrease (above 70%) in anti-α-Gal reactivity was noticed after enzymatic treatment in all strains. These data are consistent with the presence of not only variable amounts of terminal α-Gal residues but also other changes in the structure of this epitope that may affect antibody binding.

**Analysis of the glycan core moiety.** Epimastigotes and trypomastigotes have a similar linear glycan core of Man$_a$(1-2)Man$_a$(1-6)Man$_a$(1-4)GlcN$_a$(1-6)Ins-1-phosphate. Although they share the same glycan core, the latter may exhibit zero to four galactoses (Figure 1).16 For this reason, the preliminary analysis of carbohydrate anchors was based only in GPI-mucins derived from epimastigotes. To evaluate if carbohydrate variations could reflect previous described DTUs,26 GPI-mucins from epimastigotes were subjected to proteinase K treatment, and anchors were purified (Figure 2B). Those anchors were labeled and subjected to FACE (Figure 6A and B). It was observed that there were longer carbohydrate oligosaccharides for BZS and BZR (lanes 2 and 3) that reached up to six sugars (Figure 6A). CL strain (lane 5) had approximately four to five sugars, and strain Colombiana (lane 6) was very low glucosylated, with up to three sugars in its glycan core (Figure 6B).

**Unweighted pair group method analysis.** The previous results showed variations among the strains with no expected pattern. In some experiments, CL strain, for example, was similar to Colombiana (Figures 3, 4C, and 5) or Y (Figures 3B, 4B, and 6). To better describe the relationships among the strains and evaluate their position compared with molecular designations,26 a UPGMA analysis was performed (Figure 7). Based on the taxon/character similarity matrix constructed on the presence/absence of a given character, the strains fell into two main clusters. BZS and BZR populations ($T. cruzi$ II) were grouped together with CL ($T. cruzi$ VI) with a similarity coefficient index of ~76%. Colombiana strain ($T. cruzi$ I) was completely discriminated from the others with a similarity coefficient index of ~51%.

![Figure 6](image)

**Figure 6.** FACE of GPI-anchor oligosaccharides from different $T. cruzi$ strains. (A) Lane 1 = oligoglucose ladder represented by G$_1$–G$_7$; lane 2 = BZS strain; lane 3 = BZR strain. (B) Lane 4 = oligoglucose ladder represented by G$_1$–G$_7$; lane 5 = CL strain; lane 6 = Colombiana strain.

![Figure 7](image)

**Figure 7.** UPGMA of $T. cruzi$ strains based on presence/absence of characters from all biological and biochemical parameters studied. The similarity coefficient (horizontal scale) was derived from the Dice index.36 On the right, the respective DTUs ($T. cruzi$ I–VI) are indicated as previously reported.26 Definitions of abbreviations are in Figure 3.
DISCUSSION

*Trypanosoma cruzi* belongs to a heterogeneous species consisting of a pool of strains, stocks, or isolates circulating between the wild and domestic environments through different vertebrate and invertebrate hosts. Recently, it has been proposed that *T. cruzi* has six DTUs (*T. cruzi* I–VI). T. cruzi I strains are known to induce low parasitism in human Chagasic patients, whereas *T. cruzi* II strains seem to be responsible for tissue lesions and high parasitemia. T. cruzi VI comprises a group of strains that share common features with hybrid strains. For example, the CL Brener clone, with a genome that was linked to the parasite surface. The CL and BZR-Y, and CL belonging to *T. cruzi* groups I, II, and VI, respectively.

Glycoconjugates are not translated directly from the genome, being secondary genomic products. They are synthesized in the lumen of the Golgi and endoplasmic reticulum because of the action of membrane transporters and glycosyltransferases. In trypanosomatids, some of those components have been reported for *Leishmania*, and *T. rangeli*, and *T. cruzi*. The intra- and interspecies variations in glycoconjugate assembly may originate from innumerable combinatorial sequences dependent on the action and specificity of glycosidases and glycosyltransferases. Consequently, this variation allows carbohydrates to be important sources of biological diversity. Such variability is likely to have implications in antigenicity, pathogenesis, and virulence among different strains. Despite this finding, a number of parasite and host cell molecules are known to be involved in this process. Those molecules include proteins, glycoproteins, and glycoconjugates that are crucial for parasite recognition and invasion.5,9,48 which may include the Glp58/trans-sialidase (Gp85/TS) and the mucin-like superfamilies located on the parasite surface. The GPI-mucins of *T. cruzi*, also known as sialoglycoproteins, are mucin-like molecules that are highly glycosylated and have a conserved GPI-anchor moiety of GPI-mucins exhibiting intraspecies polymorphism from the strains Colombiana, Y (BZS-Y and BZR-Y), and CL belonging to *T. cruzi* groups I, II, and VI, respectively. 

During its life cycle, *T. cruzi* trypomastigotes are able to release extracellular vesicles containing glycoproteins from the parasite surface. Those purified vesicles exhibited a proinflammatory pattern during host–parasite interaction determinant in the immunopathology and tissue lesions. The pre-treatment of BALB/c mice with vesicles, followed by parasite challenge, could significantly exacerbate parasite load and inflammation of the heart, and it could hasten animal mortality. Here, similarly to the vesicles, purified tGPI-mucins were also able to modulate trypomastigote invasion during interaction with LLC-MK2 cells. Higher invasion ability was observed for Y strain after the cells were incubated with tGPI-mucins from BZR-Y population and CL strain. However, pre-incubation with tGPI-mucins slightly increased (~0.5-fold) amastigote proliferation for all strains, except for Colombiana. However, this pattern did not vary among the strains, suggesting a role for tGPI-mucins in invasion with no apparent interference in intracellular parasite development. Because the lipid moiety of GPI-anchors tends to be conserved among strains, one hypothesis that could explain the differential invasion pattern could be related to the polymorphisms in its glycan part. Because of its hydrophobic nature, the GPI-anchors from GPI-mucins could be inserted in the plasma membrane, enabling its O-linked sugars to act as a site for parasite recognition.

In protozoa, inter- and intraspecies polymorphisms were reported in the major *Leishmania* glycoconjugate, the lipophosphoglycan (LPG). In *L. infantum* and *L. major*, a higher NO production was observed in the more complex LPG structures containing glucose and galactose side chains, respectively. In *T. cruzi*, early studies reported intraspecies variations in the GPI-LPG structures containing glucose and galactose side chains, respectively. Extensive studies have reported on the biological activities of GPI-mucins from Y strain trypomastigotes, especially in the induction of NO and proinflammatory cytokines (IL-12 and TNF-α) by macrophages. Furthermore, the molecular mechanisms of this induction were reported to involve members of the nuclear mitogen-activated protein kinase (MAPK), stress-activated protein kinase (SAPK)-β (p38), and inhibitor of κB (IkB). However, an unknown aspect of *T. cruzi* glycoconjugate is how tGPI-mucins from different strains/DTUs would trigger different responses in murine macrophages. Here, a higher production of NO and IL-12 was observed for BZR-Y followed by BZS-Y, CL, and Colombiana. This result is consistent with the previous studies on the ability of tGPI-mucins from Y strain to trigger the production of proinflammatory cytokines and NO. Interestingly, this ability was lower for strains CL and Colombiana, showing that differential stimulation by tGPI-mucins may occur in different degrees in the innate immune system. Because of the fact that mostly *T. cruzi* glycoconjugates are on the external surface of cells or secreted, they are able to modulate and mediate important cell-cell, cell–matrix, and cell–molecule interactions. Infections and immune responses suffer major interferences and depend on N- and O-linked carbohydrates during innate and adaptive responses. During its life cycle, *T. cruzi* trypomastigotes are able to release extracellular vesicles containing glycoproteins from the parasite surface. Those purified vesicles exhibited a proinflammatory pattern during host–parasite interaction determinant in the immunopathology and tissue lesions. The pre-treatment of BALB/c mice with vesicles, followed by parasite challenge, could significantly exacerbate parasite load and inflammation of the heart, and it could hasten animal mortality. Here, similarly to the vesicles, purified tGPI-mucins were also able to modulate trypomastigote invasion during interaction with LLC-MK2 cells. Higher invasion ability was observed for Y strain after the cells were incubated with tGPI-mucins from BZR-Y population and CL strain. However, pre-incubation with tGPI-mucins slightly increased (~0.5-fold) amastigote proliferation for all strains, except for Colombiana. However, this pattern did not vary among the strains, suggesting a role for tGPI-mucins in invasion with no apparent interference in intracellular parasite development. Because the lipid moiety of GPI-anchors tends to be conserved among strains, one hypothesis that could explain the differential invasion pattern could be related to the polymorphisms in its glycan part. Because of its hydrophobic nature, the GPI-anchors from GPI-mucins could be inserted in the plasma membrane, enabling its O-linked sugars to act as a site for parasite recognition.
both acute and chronic Chagas patients, anti-α-Gal antibodies have a higher lytic power against *T. cruzi* trypomastigotes, which is crucial for decreasing parasite levels during infection.\(^{15,21,64}\) Our results showed that the levels of O-linked α-galactosylation may vary among the strains, and therefore, the ability of the host to clear the parasite using these antibodies might depend on the expression of those sugars on tGPI-mucins. Moreover, it is interesting to point out that Colombiana and CL strains expressed the lowest levels of those sugars among the strains. Our data are also consistent with previous biochemical studies showing important variations in O-linked oligosaccharides from strains Y, CL-Brener, and Colombiana. In the Y strain, they were represented by Galpβ1-3GlcNAc-ol, Galpβ1-6(Galpβ1-3)GlcNAc-ol, and Galpβ1-2Galpβ1-6(Galpβ1-3)GlcNAc-ol. Also, other O-linked oligosaccharides had a 1→4 linkage to N-acetylgalactosaminyl with the structures Galpβ1-4GlcNAc-ol, Galpβ1-6(Galpβ1-4)GlcNAc-ol and Galpβ1-2Galpβ1-6(Galpβ1-4)GlcNAc-ol.\(^{23}\) In the CL-Brener strain, the O-linked oligosaccharides are structurally similar to those oligosaccharides found in the Y strain. However, the former exhibited novel sialylated O-glycans represented by Neu5Aca2-3Galp-4(Galpβ1-6)GlcNAc-ol and Galpβ1-6(Galpβ1-4)Neu5Aca2→3Galpβ1-4GlcNAc-ol.\(^{24}\) Differently from Y and CL-Brener strains, Colombiana O-linked assignments were very similar to those assignments previously reported for G strains\(^{22,53}\) represented by O-glycans as belonging to a GPI-mucin family that contain β-galactofuranose (β-Gal/) residue attached to an α-GlcNAc.\(^{24}\) It is interesting to notice that the CL-Brener strain, considered a hybrid, exhibited a carbohydrate profile similar to the Y strain. However, in the present study, it was very similar to Colombiana in triggering NO and cytokine production. For this reason, we decided to compare all parameters in this study using UPGMA analysis (Figure 7). As expected, no substantial differences were observed for the Y populations BZS and BZR (*T. cruzi II*) grouped together with CL strain (*T. cruzi VI*). However, the Colombiana strain (*T. cruzi I*) was completely discriminated from the other, a result consistent with the previous molecular studies.\(^{26}\) The data also confirmed the dual nature of CL strain as a possible consequence of its hybrid nature.\(^{11}\)

The BZR-Y population was selected in *in vivo* from the Y strain, and it is resistant to treatment with BZ.\(^{6} \) Using live trypomastigotes, biological differences regarding phagocytosis and cytokine release have shown that the BZR-Y population is less susceptible to destruction than BZS-Y.\(^{65}\) However, the P-glycoprotein (PGP), a multidrug resistance-associated glycoprotein, presented the same level of protein expression in BZS-Y and BZR-Y populations.\(^{66}\) Many molecular studies have compared those two populations to identify resistance markers.\(^{67–69}\) Although no correlation was detected, other studies have found differences in the *in vitro*-induced BZ-resistant strains (17WTS X 17LER).\(^{67}\) More recently, a proteomic analysis of BZS-Y/BZR-Y and 17WTS/17LER also identified many overexpressed proteins in the BZR-Y population.\(^{70}\) In this work, tGPI-mucins from the BZR-Y population were able to induce higher levels of NO and IL-12 than BZS-Y. Similarly, although in lower levels, tGPI-mucins from the naturally BZ-resistant Colombiana strain were more recognized by anti-α-Gal antibodies than the CL-susceptible strain. Although they were able to trigger different levels of NO and IL-12, the BZR-Y and BZS-Y populations had similar expression of α-Gal epitopes.

In conclusion, we have shown intraspecies variations in the anchor and O-linked sugars in the GPI-mucins in four *T. cruzi* strains. Those variations were important for differential NO and IL-12 production as well as modulation of trypomastigote invasion. Enzymatic treatments confirmed intraspecies polymorphisms in the O-linked α-galactosyl residues. Recently, it was proposed to use structural data for taxonomic purposes for comparing the O-glycans of *T. cruzi* I and II lineages.\(^{5,71}\) Here, we could also discriminate all lineages in *T. cruzi* I, II, and VI using the newly reported consensus.\(^{69}\) We could not correlate those variations to the BZ resistance phenotype, and they are probably strain-specific. However, because tGPI-mucins are important for inducing anti-α-Gal antibodies, variations in their structures are also likely to have implications in diagnosis and immunopathology of Chagas disease depending on the *T. cruzi* strain/DTU.

Received January 6, 2012. Accepted for publication April 7, 2012.

Acknowledgments: The authors thank Dr. Egler Chiari for critical reading and helpful suggestions.

Financial support: R.P.S., S.M.M., and A.J.R. are research fellows supported by the National Council for the Development of Research of Brazil (CNPq) Process 305042/2010-6. R.R.A. is supported by Fundação de Amparo à Pesquisa do Estado de Minas Gerais (FAPEMIG). M.N.R. is supported by CNPq Grant 552072/2009-5. This work was supported by CNPq Grants PAPES-IV-400382006-9 and PDI-150880/2005-7 and FAPEMIG. This work was also funded by National Institutes of Health Grants 1R01AI070655-04, 3R01AI070655-04S1, 2G12RR008124-16A1, and 2G12RR008124-16A1S1. We thank the Biomolecule Analysis Core Facility at the Border Biomedical Research Center/Biology/UTEP University of Texas at El Paso (National Institutes of Health Grants 2G12RR008124-16A1, 2G12RR008124-16A1S1 and G12MD007592).

Authors’ addresses: Rodrigo P. Soares, Rafael R. Assis, Marcele N. Rocha, Felipe A. Moura e Castro, Gustavo F. Freitas, Silvane M. Murta, Sara L. Santos, and Alvaro J. Romanha, Centro de Pesquisas René Rachou, FIOCRUZ, Belo Horizonte, MG, Brazil, E-mails: rsoares@cpqrr.fiocruz.br, raafassis@cpqrr.fiocruz.br, marcele@cpqrr.fiocruz.br, felipeiters@hotmail.com, freitas@cpqrr.fiocruz.br, silvane@cpqrr.fiocruz.br, sara.lopes.santos@gmail.com, and romanhat@cpqrr.fiocruz.br. Ana C. Torrecillas, Departamento de Ciências Biológicas, Universidade Federal de São Paulo (UNIFESP), Diadema, SP, Brazil, E-mail: ana.trocoli@gmail.com. Alexandre F. Marques and Igor C. Almeida, Border Biomedical Research Center, Department of Biological Sciences, University of Texas, El Paso, TX, E-mails: afmarques@utep.edu and icalmeida@utep.edu.

REFERENCES


