

TcI Isolates of *Trypanosoma cruzi* Exploit the Antioxidant Network for Enhanced Intracellular Survival in Macrophages and Virulence in Mice

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Trypanosoma cruzi species is categorized into six discrete typing units (TcI to TcVI) of which TcI is most abundantly noted in the sylvatic transmission cycle and considered the major cause of human disease. In our study, the TcI strains Colombiana (COL), SylvioX10/4 (SYL), and a cultured clone (TCC) exhibited different biological behavior in a murine model, ranging from high parasitemia and symptomatic cardiomyopathy (SYL), mild parasitemia and high tissue tropism (COL), to no pathogenicity (TCC). Proteomic profiling of the insect (epimastigote) and infective (trypomastigote) forms by two-dimensional gel electrophoresis/matrix-assisted laser desorption ionization–time of flight mass spectrometry, followed by functional annotation of the differential proteome data sets (≥ 2 -fold change, $P < 0.05$), showed that several proteins involved in (i) cytoskeletal assembly and remodeling, essential for flagellar wave frequency and amplitude and forward motility of the parasite, and (ii) the parasite-specific antioxidant network were enhanced in COL and SYL (versus TCC) trypomastigotes. Western blotting confirmed the enhanced protein levels of cytosolic and mitochondrial tryparedoxin peroxidases and their substrate (tryparedoxin) and iron superoxide dismutase in COL and SYL (versus TCC) trypomastigotes. Further, COL and SYL (but not TCC) were resistant to exogenous treatment with stable oxidants (H_2O_2 and peroxynitrite [$ONOO^-$]) and dampened the intracellular superoxide and nitric oxide response in macrophages, and thus these isolates escaped from macrophages. Our findings suggest that protein expression conducive to increase in motility and control of macrophage-derived free radicals provides survival and persistence benefits to TcI isolates of *T. cruzi*.

Chagas disease, caused by the unicellular protozoan *Trypanosoma cruzi*, is ranked as the third most important parasitic disease in terms of disability-adjusted life years (1, 2). *T. cruzi* is naturally transmitted by *Triatomine* insects. In recent years, a significant proportion of the infected population has emigrated from rural areas, leading to the urbanization of Chagas disease in countries where the disease is endemic, as well as internationally (1). Chagas disease is therefore an emergent global public health problem associated with congenital transmission (3, 4), blood transfusions (5), and organ transplantations (6). Current estimates suggest that ~70 million people are at risk of infection (7, 8) and that ~6 million individuals are infected with *T. cruzi* in Latin America and Mexico. However, these estimates may not be accurate, since they are not derived from detailed epidemiological studies. For example, in Argentina, the Chagas Disease National Control Program reported 13 provinces located in the Central-Northeast area as endemic for Chagas disease in 2010 (9). Vectorial transmission was reported to be absent in Jujuy, Entre Ríos, La Pampa, Neuquén, and Río Negro but active in provinces such as Formosa, Chaco, Córdoba, Santiago del Estero, and others, with a *Triatoma infestans* infestation index of 18.1% (national average, 10.2%) (9). In the state of Salta that shares borders with Bolivia and Paraguay, national data indicated a 7.5% seroprevalence (5.7% for blood donors and 1.82% among young children); however, several research groups reported up to 27% seropositivity from field surveys

in restricted geographical areas (10, 11). These observations may mean the existence of heterogeneity in the epidemiologic situation related to Chagas disease in regions where it is endemic.

Acute Chagas disease carries a mortality rate of <5% (12), and an asymptomatic or indeterminate stage is most commonly presented in those surviving the acute infection. However, several decades later, infected individuals develop clinically symptomatic cardiac (~30%) or digestive (8 to 10%) forms of Chagas disease (13). The chagasic megacolon is primarily presented in cases in

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Central Brazil and Chile but is largely nonexistent in those in Venezuela and parts of Central America (14). The situation in Argentina is not very clear; however, epidemiological data suggest that genetic variations of both host and the parasite play an important role in determining the development of clinical disease in infected individuals.

Trypanosoma (Schizotrypanum) cruzi belongs to the order Kinetoplastida, which comprises flagellar organisms with a kinetoplast organelle located in the cell mitochondrion. *T. cruzi* is included in the stercorarian section, together with the group of trypanosomes whose infective stage develops in the vectors' digestive tract and contaminates the mammalian hosts through feces (15). The subgenus *Schizotrypanum* has been adopted to identify the intracellular replicative stage. At present, based upon molecular and multilocus enzyme electrophoresis analyses, *T. cruzi* isolates are classified into six discrete typing units (DTU) referred to as TcI to TcVI (16). Of these, TcI has the widest distribution and is the most common cause of disease in Colombia and Venezuela (17), and in the northern region of South America and the United States (4, 18). Although *T. cruzi* isolates belonging to other DTU (e.g., TcII, TcV, and TcVI) have been identified in peripheral blood of seropositive patients in Southern Cone countries of South America (19), TcI isolates were mainly reported in heart biopsy specimens from chronic chagasic patients in the Southern Cone countries (20, 21). These findings suggested to us that patients could be carrying a mixed infection, with TcI associated with the development of clinical cardiac disease. Other studies have, however, failed to find a reliable association between cardiomyopathy and TcI DTU (22).

In this study, we aimed to identify whether diversity at the virulence level exists among the TcI isolates of *T. cruzi*, and we investigated the potential molecular mechanism(s) responsible for diversity in virulence and pathogenicity of the TcI isolates. We examined the infectivity and virulence of three TcI strains chosen based upon their diverse origin, which included the Colombiana strain (COL) recovered from a chronically infected human patient from Colombia (23); the Sylvio-X10 strain (SYL) isolated from an acutely infected human patient in Brazil (24); and the TCC laboratory strain (TCC) cloned from the continuous axenic culture of *T. cruzi* isolated from an infected human patient in Chile (25). We performed global protein expression profiling of the insect (epimastigote) and infective (trypomastigote) stages of the three strains to identify the proteins that might be differentially expressed in a stage- or strain-specific manner and contribute to differences in the disease outcome. We validated the proteome results by Western blotting and performed *in vitro* and *in vivo* studies to look at the protein networks that might be responsible for variations in parasites' survival in macrophages and virulence in mice. Our results suggested that a gene/protein expression profile conducive to enhanced flagellar motility and peroxiredoxin-dependent decomposition of macrophage-derived free radicals provided survival and persistence benefits to TcI DTU isolates. We discuss the significance of our observations in relation to *T. cruzi* modulation of host responses for ensuring its intracellular survival, dissemination, and persistence resulting in chronic disease.

MATERIALS AND METHODS

Parasites, cell culture, and mice. All chemicals were of molecular grade and purchased from Sigma-Aldrich (St. Louis, MO) unless stated otherwise. Epimastigotes of TcI strains of *T. cruzi* (COL, SYL, and TCC) were

cultured in liver infusion tryptose–10% fetal bovine serum (FBS) medium supplemented with 25 µg of hemin/ml, 100 U of penicillin/ml, and 100 mg of streptomycin/ml. *T. cruzi* trypomastigotes were maintained and propagated by continuous *in vitro* passage in C2C12 cells. RAW 264.7 macrophages were cultured in Dulbecco modified Eagle medium high glucose–10% FBS medium (Life Technologies, Grand Island, NY). Macrophages were seeded in six-well (5×10^4 /well), 12-well (2×10^4 /well), or 96-well (0.5×10^4 /well) plates or Nunc Lab-Tek II chamber slides (1×10^4 /well), infected with *T. cruzi* trypomastigotes (cell/parasite ratio, 1:3) for 3 h, washed to remove free parasites, and then incubated according to experimental requirements.

All animal experiments were performed according to the National Institutes of Health *Guide for the Care and Use of Experimental Animals* and approved by the institutional ethics committee at the Universidad Nacional de Salta. Swiss mice were bred at the animal facility at the School of Health Science, National University of Salta, Salta, Argentina. *T. cruzi* epimastigotes were incubated in the presence of 1% triatomine gut homogenate to facilitate differentiation to metacyclic trypomastigotes (26). Complement-resistant metacyclic trypomastigotes were purified by incubation of stationary culture in the presence of normal human serum. Swiss mice (6 to 8 weeks old) were infected by intraperitoneal injection with *T. cruzi* (10^4 metacyclic trypomastigotes per mouse). Survival from infection was monitored daily, and blood parasitemia was monitored at 2-day intervals for up to 56 days postinfection (p.i.). Mice were sacrificed in the chronic disease phase (>120 days p.i.), and blood and tissue samples (heart, skeletal muscle) were harvested and stored at -80°C .

Histology. Tissue sections were fixed in 10% buffered formalin for 24 h, dehydrated in absolute ethanol, cleared in xylene, and embedded in paraffin. Then, 5-µm sections of the heart and skeletal muscle tissue (≥ 3 tissue sections/organ/mouse) were stained with hematoxylin and eosin (H&E) and evaluated by two observers in a blinded fashion by light microscopy using an Olympus BX-15 microscope (Center Valley, PA) equipped with a digital camera and Simple PCI software (v.6.0; Compix, Sewickley, PA). Inflammatory lesions (myocarditis) were scored as follows: 0, absent; 1, focal or mild with ≤ 1 foci; 2, moderate with ≥ 2 inflammatory foci; 3, severe with extensive generalized coalescing of inflammatory foci or disseminated inflammation; or 4, diffused inflammation with severe tissue necrosis, interstitial edema, and loss of integrity (27). Inflammatory infiltrate was characterized as diffused or focal depending upon how closely the inflammatory cells were associated. The parasitic or inflammatory foci in skeletal muscle and heart tissue sections were scored as follows: 0, absent; 1, 0 to 1 foci; 2, 1 to 5 foci; and 3, >5 foci.

T. cruzi homogenates, two-dimensional electrophoresis (2DE), and image analysis. *T. cruzi* epimastigotes or trypomastigotes were washed with cold phosphate-buffered saline and lysed in 7 M urea, 2 M thiourea, 4% CHAPS {3-[(3-cholamidopropyl)-dimethylammonio]-1-propane-sulfonate}, 12 mM EDTA, and 40 mM Tris (pH 7.6) containing protease inhibitor cocktail (1 mM phenylmethylsulfonyl fluoride, 0.001 mM pepstatin, and 0.1 mM leupeptin). Parasite pellets were added to the lysis buffer (1 g [wet weight] per 3 ml of lysis buffer), vortexed for 3 min, incubated on ice for 10 min, and centrifuged at $1,000 \times g$ for 5 min to remove the cell debris. Protein concentrations were determined by using a Bio-Rad protein assay (Bio-Rad, Hercules, CA).

T. cruzi lysates (100 µg of protein) were separated by 2DE, employing an IPGphor multiple sample isoelectric focusing (IEF) device (GE Healthcare, Little Chalfont, United Kingdom) in the first dimension and a Criterion Dodeca cell (Bio-Rad) in the second dimension, as described previously (28). Briefly, samples were loaded onto 11-cm dehydrated, precast, immobilized pH gradient (IPG) strips (pH 3 to 10 and pH 5 to 8; Bio-Rad), and the strips were rehydrated overnight in 1 M thiourea, 8 M urea, 2% CHAPS, 1% dithiothreitol, and 0.2% ampholyte rehydration buffer containing 0.002% bromophenol blue. The IEF was performed at 20°C using the following parameters: 50 V, 11 h; 250 V, 1 h; 500 V, 1 h; 1000 V, 1 h; 8,000 V, 2 h; and then at 8,000 V (for a total of 50,000 V). The IPG strips were transferred to equilibration buffer (50 mM Tris-HCl [pH

6.8], 6 M urea, 20% glycerol) and sequentially incubated for 15 min each in the presence of 2% dithiothreitol–2% sodium dodecyl sulfate (SDS; reducing conditions) and 2.5% iodoacetamide–2% SDS (alkylating conditions). Equilibrated IPG strips were subjected to 2DE by using 8 to 10% linear gradient precast Tris-HCl gels (Bio-Rad) in Tris-glycine-SDS buffer (25 mM Tris-HCl, 192 mM glycine, 0.1% SDS [pH 8.3]). Electrophoresis was performed on a Protean plus Dodeca cell system at 75 V for 1 h and then at 120 V until the dye front reached the bottom of the gel.

Gels were fixed in 10% methanol–7% acetic acid, stained with SYPRO Ruby (Bio-Rad), and destained in 10% ethanol–7% acetic acid to remove the background. Gel images were acquired at an excitation wavelength of 488 nm and an emission wavelength of 560 nm by using a high-resolution ProXPRESS proteomic imaging system (Perkin-Elmer, Waltham, MA). The exposure time was adjusted to achieve a value of ~55,000 to 63,000 pixel intensity (16-bit saturation) from the most intense protein spots on the gel (28).

All 2D gels, representing at least four samples per stage (epimastigote and trypomastigote) from the COL, SYL, and TCC isolates, were scanned and analyzed with the Progenesis SameSpots software 2.0 (NonLinear Dynamics, Durham, NC). After manual and automated pixel-to-pixel alignment, the program performed automatic spot detection on all images. Normalized spot volumes, i.e., the volume of each spot over the volume of all spots in the gel, were used to compare the different groups, and candidates were identified as protein spots that changed in expression level by ≥ 2 -fold versus their specific control. Statistical significance was assessed by a two-tailed Student *t* test and analysis of variance test (analysis of variance [ANOVA]), and *P* values of < 0.05 were considered significant for comparisons between control and experimental data.

Mass spectrometry (MS) and protein identification. Selected spots (< 1 mm) on the 2D gels containing the protein of interest were excised and trypsin digested as described previously by us (29, 30). Briefly, gel spots were incubated at 37°C for 30 min in 50 mM NH_4HCO_3 and dehydrated twice for 5 min each time in 100 μl of acetonitrile, and the proteins were digested in gel at 37°C overnight with 10 μl of trypsin solution (1% trypsin in 25 mM ammonium bicarbonate). Peptide mixtures (1 μl) were directly spotted onto a matrix-assisted laser desorption/ionization–time of flight (MALDI-TOF) tandem MS (MS/MS) target plate with 1 μl of alpha-cyano-4-hydroxycinnamic acid matrix solution (5 mg/ml in 50% acetonitrile) and analyzed by using a MALDI-TOF/TOF ABI 4800 proteomics analyzer (Applied Biosystems, Foster City, CA). The Applied Biosystems software package included 4000 Series Explorer (v3.6 RC1) with Oracle Database Schema Version (v. 3.19.0) and Data Version (3.80.0) to acquire and analyze MS and MS/MS spectral data. The instrument was operated in positive ion reflectron mode with the focus mass set at 1,700 Da (the mass range was 850 to 3,000 Da). For MS data, 2,000 to 4,000 laser shots were acquired and averaged from each protein spot. A peptide mixture with reference masses of 904.468, 1,296.685, 1,570.677, and 2,465.199 kDa was used for automatic external calibration. After MALDI MS analysis, MALDI MS/MS was performed on several (5 to 10) abundant ions from each sample spot. A 1-kV positive-ion MS/MS method was used to acquire data under post-source-decay conditions. The instrument precursor selection window was ± 3 Da. For MS/MS data, 2,000 laser shots were acquired and averaged from each sample spot. Automatic external calibration was performed using reference fragment masses 175.120, 480.257, 684.347, 1,056.475, and 1,441.635 (28).

For protein identification, the MS and MS/MS spectral data were searched against the National Center for Biotechnology Information, UniProtKB/Swiss-Prot, and TryTryp databases (accessed on 21 July 2013) using AB Sciex GPS Explorer (v.3.6) software in conjunction with MASCOT (v2.2.07) as described previously (28). Briefly, protein match probabilities were determined by using expectation values and/or MASCOT protein scores. The MS peak filtering included the indicated parameters—mass range, 800 to 4,000 Da; minimum signal-to-noise ratio (S/N) filter, 10; mass exclusion list tolerance, 0.5 Da—and the mass exclusion list for some trypsin and keratin-containing compounds included masses of

842.51, 870.45, 1,045.56, 1,179.60, 1,277.71, 1,475.79, and 2,211.1 Da. The MS/MS peak filtering included the following parameters: minimum S/N filter, 10; maximum missed cleavages, 1; fixed modification of carbamidomethyl (C); variable modifications due to oxidation (M); precursor tolerance, 0.2 Da; MS/MS fragment tolerance, 0.3 Da; mass, monoisotopic; and peptide charges, +1. The significance of a protein match, based on both the peptide mass fingerprint (PMF) in the MS and the MS/MS data from several precursor ions, is presented as the expectation value. The expectation value is the number of matches with equal or better scores that are expected to occur by chance alone. The default significance threshold is $P < 0.05$; however, we used a stringent threshold of 10^{-3} for protein identification: the lower the expectation value, the more significant the score.

Western blotting. Polyclonal antibodies against *T. cruzi* cytosolic trypanredoxin peroxidase (cTXNPx) and mitochondrial trypanredoxin peroxidase (mTXNPx) were obtained as described previously (31, 32). Polyclonal antibodies against the iron superoxide dismutase (FeSOD) isoforms, FeSODa and FeSODb, and the trypanredoxin isoforms, TXN I and TXN II, were produced as described previously (31, 32).

T. cruzi epimastigotes (5×10^8 cells/ml) and trypomastigotes (1×10^9 cells/ml) were lysed in 10 mM Tris-HCl (pH 7.5), 5 mM EDTA, and 1% Nonidet P-40 containing protease inhibitor cocktail (Sigma-Aldrich). Parasite lysates (30 $\mu\text{g}/10 \mu\text{l}$) were mixed with equal volumes of 2 \times loading buffer (100 mM Tris-Cl [pH 6.8], 4% SDS, 0.2% bromophenol blue, 20% glycerol, and 200 mM β -mercaptoethanol). Samples were resolved by 10% SDS-PAGE, and the proteins were transferred onto polyvinylidene difluoride (PVDF) membranes by using a wet vertical Criterion blotter (Bio-Rad). Membranes were blocked for 1 h with 5% nonfat dry milk (Bio-Rad) in 50 mM Tris-HCl (pH 7.4), 150 mM NaCl, and 0.05% Tween 20. Antibodies were diluted in Tris-buffered saline (TBS; 25 mM Tris-HCl [pH 7.6], 140 mM NaCl, and 3 mM KCl) containing 1% bovine serum albumin and 0.1% Tween 20. Membranes were incubated overnight at 4°C with antibodies against mTXNPx (1:2,000 dilution), cTXNPx (1:5,000 dilution), FeSODa and FeSODb (1:2,000 dilution), and TXN1 and TXN2 (1:1,000 dilution). Membranes were then washed three times in TBS and incubated for 1 h with horseradish peroxidase (HRP)-conjugated anti-rabbit IgG (1:10,000 dilution; Sigma-Aldrich). Signal was developed by using an enhanced chemiluminescence detection system (GE Healthcare). Membranes were stained with Ponceau S to confirm equal loading of samples. Images were visualized and digitized, and densitometry analysis was performed on a Fluorchem HD2 system (Alpha Innotech, Santa Clara, CA).

Parasite susceptibility to oxidants. Activated macrophages exert cytotoxic effects against microorganisms via NADPH oxidase-dependent production of superoxide ($\text{O}_2^{\cdot -}$) that then produces stable and diffusible H_2O_2 oxidant. The reaction of superoxide with nitric oxide produces peroxynitrite (ONOO^-) that is a strong cytotoxic oxidant. To assess the susceptibility of *T. cruzi* isolates to these oxidants, trypomastigotes were distributed to 96-well plates ($10^5/100 \mu\text{l}/\text{well}$), followed by incubation in the presence of 250 μM hydrogen peroxide (H_2O_2) or peroxynitrite (ONOO^-). The selected concentrations of oxidants were chosen based upon the dose response curves using 0 to 500 μg of H_2O_2 and ONOO^- that showed $< 10\%$ of SylvioX10 trypomastigotes died when treated with the highest concentration for 2 h. alamarBlue (10% final concentration; Invitrogen) was added, and its reduction by mitochondrial dehydrogenases of the live parasites, resulting in the formation of resazurin (red, highly fluorescent; excitation, 560 nm/emission, 590 nm), was monitored by using a SpectraMax M5 microplate reader (Molecular Devices, Sunnyvale, CA). The results are expressed as the percentage of viability, which was calculated as follows: (the mean fluorescence of parasites treated with oxidants/the mean fluorescence of parasites untreated) $\times 100$.

Activation of macrophages by *T. cruzi*: oxidative/nitrosative burst and cytokine release. RAW 264.7 macrophages were infected with *T. cruzi* for 3 h (cell/parasite ratio, 1:3), washed with RPMI to remove free parasites, and then incubated for 24, 36, or 48 h. The cells were incubated

with 10 μ M CM-H₂DCF-DA (Molecular Probes, Eugene, OR), and its oxidation by intracellular reactive oxygen species (ROS), resulting in the formation of fluorescent dichlorodihydrofluorescein (DCF; excitation, 498 nm/emission, 598 nm), was monitored on a SpectraMax M5 microplate reader. Fluorescence microscopy for intracellular DCF was performed on an Olympus BX-15 microscope, and images were captured with a mounted digital camera (magnification, $\times 40$).

ROS release in culture supernatants from infected macrophages was measured by an Amplex red assay. Briefly, supernatants (50 μ l) were added in triplicate to 96-well, black flat-bottom plates and mixed with 100 μ M Amplex red (Molecular Probes) and HRP at 0.3 U/ml. The HRP-catalyzed Amplex red oxidation by H₂O₂, resulting in fluorescent resorufin formation, was monitored at an excitation wavelength of 563 nm and an emission wavelength of 587 nm (standard curve, 50 nM to 5 μ M H₂O₂) (33).

The inducible nitric oxide synthase (iNOS) activity was monitored by the Griess reagent assay by using a nitrate/nitrite colorimetric assay kit (Cayman, Ann Arbor, MI). Briefly, culture supernatants (80 μ l) were reduced by the addition of 0.01 U of nitrite reductase, and 100 μ l of 1% sulfanilamide made in 5% phosphoric acid–0.1% *N*-(1-naphthyl) ethylenediamine dihydrochloride (1:1 ratio [vol/vol]) was added. After incubation for 10 min, the formation of diazonium salt was monitored at 545 nm (standard curve, 2 to 100 μ M sodium nitrite). Culture supernatants from infected macrophages were also utilized to measure secreted gamma interferon (IFN- γ), tumor necrosis factor alpha (TNF- α), and interleukin-6 (IL-6) cytokines by using optEIA enzyme-linked immunosorbent assay (ELISA) kits (Pharmingen, San Diego, CA), according to the manufacturer's specifications.

Parasite replication and release in macrophages. RAW 264.7 macrophages were cultured in 12-well plates, infected with *T. cruzi* isolates for 3 h, washed to remove free parasites, and then incubated for 6, 12, or 18 h. Total DNA was isolated from infected and uninfected macrophages by using a QIAamp DNA minikit (Qiagen, Chatsworth, CA). The total DNA (50 ng) was used as a template, and real-time PCR performed on an iCycler thermal cycler with SYBR green Supermix (Bio-Rad) and Tc18S-specific (forward, 5'-TAGTCATATGCTTGTTC-3'; reverse, 5'-GCAA CAGCATTAATATACGC-3') oligonucleotides. Data were normalized to murine GAPDH (glyceraldehyde-3-phosphate dehydrogenase; forward, 5'-TGTGATGGGTGTGAACCACGAGA-3'; reverse, 5'-GAGCCCTCC ACAATGCCAAAGT-3'), and the fold change in *T. cruzi* burden calculated as $2^{-\Delta\Delta CT}$, where ΔCT represents the C_T (sample) – the C_T (control) (27). To assess parasite release, metacyclic trypomastigotes derived from stationary-phase epimastigote cultures of the different *T. cruzi* isolates were added to RAW 264.7 macrophages, and the trypomastigote release in the supernatant was monitored for 2 weeks by light microscopy.

Data analysis. Data (means \pm the standard deviations) were derived from at least three independent experiments (triplicate observations per experiment) performed on different days. All data were analyzed by using Prism 5 (GraphPad Software, La Jolla, CA). Data were tested for normal distribution by histogram and Q-Q test and analyzed by using a Student *t* test (comparison of two groups), and one-way analysis of variance (ANOVA) with Tukey's test (comparison of multiple groups). The level of significance was accepted at $P < 0.05$.

RESULTS

We used a well-established experimental model to assess the virulence of TcI isolates of *T. cruzi*. Swiss mice infected with SYL exhibited patent blood parasitemia that peaked at 21 to 28 days p.i. and was controlled by day 49 p.i. (Fig. 1A). Mice infected with COL exhibited a delayed and up to 2-fold lower level of peak parasitemia compared to that noted in SYL-infected mice ($P < 0.01$ [*t* test]). In comparison, mice infected with TCC exhibited no detectable parasitemia by light microscopy, although blood PCR for Tc18SrDNA (Fig. 1A) and hemoculture (data not shown) con-

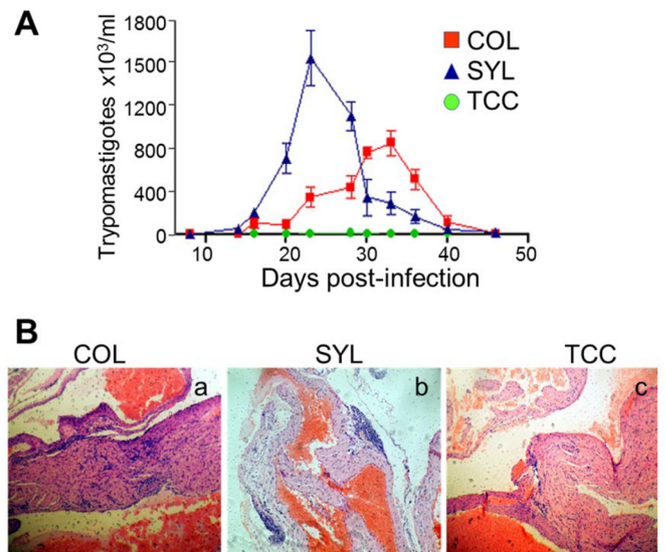


FIG 1 Biological behavior of selected TcI isolates of *T. cruzi* in mice. Swiss mice were infected with Colombiana (COL), Sylvio X10 (SYL), and TCC strains of *T. cruzi* ($n = 6$ per group), as described in Materials and Methods. (A) Blood parasitemia was monitored by light microscopy (10 microscopic fields per sample). (B) Representative H&E staining (blue, nuclear; pink, muscle, cytoplasm, or keratin) of heart tissue sections from COL-, SYL-, and TCC-infected mice, harvested during the chronic disease phase (>120 days postinfection [p.i.]; magnification, $\times 10$). In all figures, data are presented as means \pm the SD, and significance is indicated as follows: *, $P < 0.05$; **, $P < 0.01$; and ***, $P < 0.001$.

firmed that mice were infected. Normal Swiss mice were used as controls in all experiments.

Elicitation of inflammatory response is important for control of acute *T. cruzi* infection, but its chronic persistence can contribute to tissue injury (27). Histological studies were performed in chronic stage (>120 days p.i.) to assess the tissue inflammation and myocarditis. The inflammatory foci were observed in all tissue sections (heart and skeletal muscle) of COL- and SYL-infected mice. Representative images of H&E-stained heart tissue sections are presented in Fig. 1B. Mice infected with COL isolate (versus normal/uninfected mice) exhibited extensive and diffused inflammation consisting mainly of interstitial inflammatory infiltrate throughout the heart (histological score, 2.75 ± 0.4 versus 0.3 ± 0.03 ; $P < 0.05$ [*t* test]; Fig. 1Ba) and skeletal muscle (histological score, 3 ± 0.43 versus 0.32 ± 0.03 ; $P < 0.05$ [*t* test]) tissue sections. The extent of inflammatory infiltrate in the heart tissue (score, 2.5 ± 0.45 ; Fig. 1Bb) and skeletal muscle (score, 3 ± 0.33) of SYL-infected mice was not statistically different than that observed in COL-infected mice. The degeneration of muscle fibers in the heart (Fig. 1B) and skeletal muscle tissue (data not shown) of COL- and SYL-infected mice was also evident. The TCC-infected mice exhibited no overt tissue pathology and minimal inflammatory infiltrate (histological score, 0.25 ± 0.03 and 0.5 ± 0.053 in heart and skeletal muscle, respectively) that was significantly lower than that noted in COL- and SYL-infected mice ($P < 0.05$ [ANOVA-Tukey]; Fig. 1Bc). These data suggested that COL and SYL (but not TCC) strains elicited blood parasitemia during the acute infection period and persistent inflammatory infiltrate and injury in the myocardium and skeletal tissue during the chronic phase. The extent of blood parasitemia was not directly correlated with chronic persistence of the parasite in the tissues.

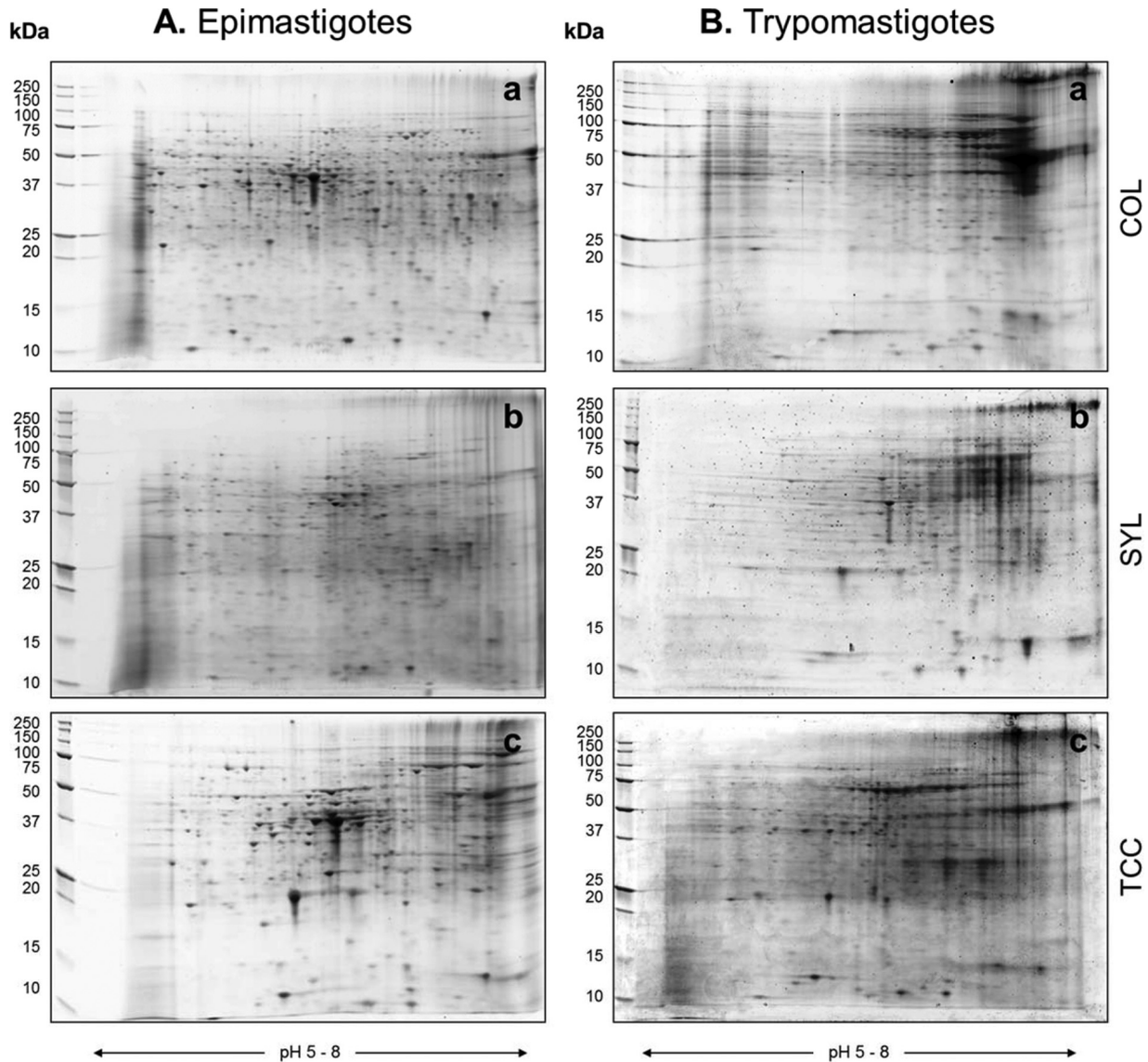


FIG 2 Proteomic profiling of TcI strains of *T. cruzi* by 2DE. Epimastigote (A) and trypomastigote (B) stages of *T. cruzi* COL, SYL, and TCC strains were cultured as described in Materials and Methods. Parasite lysates were separated in the first dimension by IEF on 11-cm linear, pH 5 to 8 immobilized pH gradient strips, and in the second dimension by SDS-PAGE on an 8 to 10% gradient gel. Gel images were obtained by using a high-resolution ProXPRESS proteomic imaging system. Representative gel images of protein lysates from COL (Aa and Ba), SYL (Ab and Bb), and TCC (Ac and Bc) are shown.

To begin to understand why TcI isolates may exhibit a differing virulence in mice, we performed proteomic analysis of the infective trypomastigotes of COL, SYL, and TCC isolates. The proteome profile of the epimastigote (insect, replicative) stage was included to evaluate whether changes in the proteome profile are stage or strain specific. In initial analysis, when using pH 3 to 10 strips, most of the proteins were resolved in the middle or on the acidic side of the second-dimension 8 to 16% gradient gel (data not shown). The narrow-range IPG strips (pH 5 to 8) yielded enhanced resolution of *T. cruzi* proteins and detection of 2-fold-more spots than were detected by using other IPG strips. We therefore used pH 5 to 8 IPG strips for developing the proteome profile of COL, SYL, and TCC strains ($n \geq 4$ per sample). Representative gel images for the epimastigote (Fig. 2Aa to c) and trypomastigote (Fig. 2Ba to c) forms of each *T. cruzi* strain are shown

in Fig. 2. All protein spots were within the relative molecular size of 10 to 250 kDa.

The 2D gel images were assessed for quality control by SameSpots software, and gel images were aligned both manually and automatically against the reference epimastigote stage and trypomastigote stage gels (Fig. 3). Spot boundaries were then detected to identify the protein spots reproducibly found in all samples within a group, as well as those that were differentially abundant between any two groups. On average, 1,153, 1,347, and 1,106 protein spots were observed in epimastigotes, and 1,463, 1,172, and 1,475 protein spots were observed in trypomastigotes of COL, SYL, and TCC isolates, respectively.

For the purpose of selecting protein spots for identification by MS, the data sets were analyzed in a pairwise manner by a *t* test with Welch's correction that accounts for unequal variances. This

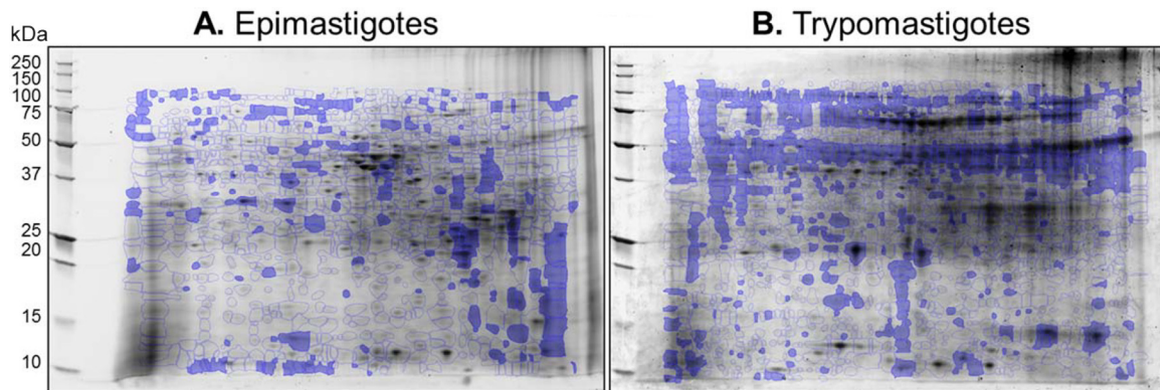


FIG 3 Identification of differentially abundant protein spots in *T. cruzi* TcI strains. Protein spots that exhibited significant changes in abundance in epimastigote (A) and trypomastigote (B) stages of COL and SYL (versus TCC, $P < 0.05$) are marked on representative gel images of SYL lysate. Protein spots were subjected to MALDI-TOF MS analysis for protein identification (see [Tables 1 and 2](#)).

analysis yielded 55, 47, and 30 differentially abundant protein spots (fold change, ≥ 2.0 ; $P < 0.05$ [*t* test]) in COL-versus-SYL, SYL-versus-TCC, and COL-versus-TCC epimastigotes, respectively. Likewise, we identified 50, 33, and 40 differentially abundant protein spots (fold change, ≥ 2.0 ; $P < 0.05$ [*t* test]) in COL-versus-SYL, SYL-versus-TCC, and COL-versus-TCC trypomastigotes, respectively. These protein spots were subjected to MALDI-TOF MS/MS analysis, and homology searches were conducted against the TryTryp database for protein identification. The protein spots that were differentially abundant in epimastigotes and trypomastigotes of COL, SYL, or TCC (fold change, ≥ 2.0 , $P < 0.05$) and identified with high confidence (E value < 0.001) are listed in [Tables 1 and 2](#), respectively.

Bioinformatics analysis using SMART and TARGET software showed that several of the proteins that participate in host cell invasion (e.g., cysteine peptidases, surface glycoprotein gp82, and cruzipain) and cytoskeletal remodeling and motility (e.g., α - and β -isoforms of actin and tubulin, cofilin-1, KMP11, and paraflagellar rod proteins) were significantly increased in abundance in both the epimastigote and trypomastigote stages of the SYL and COL compared to that noted in the TCC strain ([Tables 1 and 2](#)). The antioxidant proteins involved in redox homeostasis (e.g., trypanothione peroxidases, peroxiredoxins, thioredoxin-dependent peroxide reductase [PRDX3], and iron superoxide dismutases [FeSODa/b]) were primarily increased in abundance in the trypomastigote stage of the SYL and COL compared to that noted in TCC trypomastigotes ([Table 2](#)).

We chose to confirm the expression profile of the antioxidant proteins because these exhibited a differential abundance in a stage- (epimastigote versus trypomastigote) as well as strain-specific manner. Parasite lysates were probed by using antibodies against cytosolic (cTXNPx) and mitochondrial (mTXNPx) isoforms of trypanothione peroxidase, trypanothione isoforms TXN I and TXN II, and iron superoxide dismutase isoforms FeSODa and FeSODb ([31, 32](#)). Due to the finding of a differential abundance of α -tubulin and β -actin in *T. cruzi* isolates ([Table 1 and Table 2](#)), we could not use these proteins for normalizing the antioxidant protein content. Instead, equal protein loading was confirmed by Ponceau S staining of the PVDF membranes, as before ([34](#)). Representative Western blot images and densitometry evaluation of the protein abundance in the epimastigote stage of the TcI isolates

are shown in [Fig. 4](#). The cTXNPx expression was increased by 46% in COL epimastigotes (versus TCC, $P < 0.05$ [ANOVA-Tukey]), whereas the abundance of all other members of the trypanothione family and FeSOD isoforms was not changed significantly in the epimastigote stage of the three TcI isolates ([Fig. 4A and B](#)). The Ponceau S-stained gel image ([Fig. 4C](#)) confirmed equal loading of all samples.

The expanded views of the corresponding spot for FeSOD and TXNPx from representative 2D gel images of SYL and TCC trypomastigotes are shown in [Fig. 5A](#). As determined by Western blotting, COL and SYL trypomastigotes exhibited increases in the protein levels of cTXNPx (2.5- and 2-fold, respectively; $P < 0.05$ [ANOVA]), mTXNPx (32 and 28%, respectively; $P < 0.05$ to 0.01 [ANOVA]); TXN I (2.0- and 2.14-fold, respectively; $P < 0.01$), TXN II (39.7 and 31.5%, respectively; $P < 0.05$ [ANOVA]); and FeSODa (33.1 and 49.5%, respectively; $P < 0.05$ [ANOVA]) compared to that noted in TCC trypomastigotes ([Fig. 5B and C](#)). The protein level of FeSODb was not statistically different in trypomastigotes of the three isolates. The Ponceau S-stained gel image ([Fig. 5D](#)) confirmed equal loading of all samples. Together, the results presented in [Fig. 4 and 5](#) validate the 2DE/MS findings and suggest that the antioxidant network was upregulated in the trypomastigote stage of the virulent TcI isolates (COL and SYL) of *T. cruzi*.

To assess the biological significance of the antioxidant status in TcI isolates, we took a three-pronged approach. First, we sought to determine whether the selected TcI strains differed in their ability to resist exogenous oxidative/nitrosative stress ([Fig. 6](#)). Trypomastigotes were challenged for 2 and 4 h with biologically relevant doses of H_2O_2 and $ONOO^-$, the main oxidants present in the phagosomal environment ([35, 36](#)), and parasite viability was determined by an alamarBlue assay. Up to 84.9, 95.7, and 66.8% of COL, SYL, and TCC trypomastigotes, respectively, were viable after H_2O_2 treatment for 2 h, and their viabilities were not further decreased after 4 h ([Fig. 6A](#)). Likewise, 80.6, 93.2, and 49.7%, of COL, SYL, and TCC trypomastigotes, respectively, were viable after $ONOO^-$ treatment for 2 h, and parasite viability was not further reduced at 4 h ([Fig. 6B](#)). In comparison to TCC, the SYL and COL isolates exhibited significantly higher levels of resistance to exogenous H_2O_2 (27 and 46%, respectively; $P < 0.01$ [ANOVA-

TABLE 1 Protein spots that were differentially expressed in epimastigote stage of TcI isolates of *T. cruzi*^a

Protein spot	Protein name (gene)	NCBI accession no.	Mol mass (Da)	Peptide count	Protein score	Protein score		E value
						(CI%)	pI	
Spots conserved in the three isolates								
1	Chain A, tyrosine aminotransferase (TcTAT)	gi 6137418	46,120.3	12	188	100	5.72	3.155E-13
3	Tyrosine aminotransferase (TcTAT)	gi 71407311	46,238.5	13	442	100	5.82	1.256E-38
4	Tyrosine aminotransferase N-term (TcTAT)	gi 71655339	27,698.9	12	400	100	5.71	1.991E-34
5	L-Tyrosine:2-oxoglutarate aminotransferase (TcTAT)	gi 1168606	46,137.4	20	737	100	5.82	3.972E-68
6	Tyrosine aminotransferase (TcTAT)	gi 71413938	46,100.4	16	537	100	5.66	3.972E-48
7	NADPH flavin oxidoreductase (old yellow enzyme, TcOYE)	gi 61741942	42,191.6	19	902	100	6.03	1.256E-84
8	NADPH flavin oxidoreductase (old yellow enzyme, TcOYE)	gi 61741950	42,233.6	18	889	100	5.97	2.506E-83
9	L-Threonine 3-dehydrogenase (TcTDH)	gi 71406160	36,880.7	8	181	100	6.71	1.581E-12
10	NADPH flavin oxidoreductase (old yellow enzyme, TcOYE)	gi 61741952	42,080.5	20	749	100	6.03	2.506E-69
11	Aldo-keto reductase/quinone oxido-reductase (AKR)	gi 189396135	32,485.5	13	499	100	7.18	1.256E-43
12	Aldo-keto reductase/quinone oxido-reductase (AKR)	gi 14279174	31,764.2	13	376	100	8.58	2.506E-31
Protein spots differentially expressed in COL (compared to TCC)								
13	Proline racemase B (TcPRACB)	gi 74812220	38,440.6	12	442	100	5.44	1.256E-38
14	Proline racemase N-term (TcPRACB)	gi 71400794	32,411.5	11	435	100	5.63	6.295E-38
15	Major paraflagellar rod protein 2 (TcPAR2)	gi 162179	69,504.9	19	355	100	5.85	6.295E-30
16	Paraflagellar rod protein N-term (TcPAR)	gi 71411387	42,471.9	16	355	100	6.12	6.295E-30
17	Paraxonemal (paraflagellar) rod protein (TcPAR2)	gi 71398801	31,336.1	10	164	100	5.95	7.924E-11
18	2-Hydroxyacid dehydrogenase N-term, D-isomer (2-Hacid-dh)	gi 71406187	28,516.7	16	567	100	6.17	3.972E-51
20	Short interspersed repetitive element (TcSIRE)	gi 1911172	11,565	10	530	100	5.08	1.991E-47
21	2-Hydroxyacid dehydrogenase, D-isomer (2-Hacid-dh)	gi 71406189	22,228.2	11	296	100	5.87	5E-24
23	HpcH/HpaI aldolase (TcP28)	gi 4558456	29,886.2	7	156	100	6.06	5E-10
24	Pyridoxal kinase (TcPDXX)	gi 71418579	33,273.6	4	125	100	5.93	6.295E-07
25	NADP-dependent malic enzyme N term (TcME1)	gi 71662506	62,705.5	12	307	100	6.55	3.972E-25
26	NADP-dependent malic enzyme N term (TcME1)	gi 71651465	62,634.4	14	300	100	6.37	1.991E-24
27	Cytochrome c oxidase subunit IV (TcCOXIV)	gi 71660419	38,924.4	16	351	100	5.72	1.581E-29
28	Tryparedoxin peroxidase (TcTXNPx)	gi 17224953	22,246.1	11	413	100	5.96	9.976E-36
29	Tryparedoxin peroxidase (TcTXNPx)	gi 71399514	22,375.1	11	411	100	6.32	1.581E-35
30	Tryparedoxin peroxidase (TcTXNPx)	gi 4388655	25,545.3	22	167	100	7.56	2.506E-28
31	Calpain-like cysteine peptidase (TcCAPN1)	gi 71411006	12,798.3	6	179	100	5.07	1.256E-11
32	Beta tubulin of <i>T. cyclops</i> (TcTUBB)	gi 91983197	44,496.8	13	410	100	5.68	1.991E-35
33	Beta tubulin of <i>T. danilewskyi</i> (TdTUBB)	gi 74229926	49,695.8	12	391	100	4.74	1.581E-33
34	Alpha tubulin of <i>T. cyclops</i> (TcTUBA)	gi 91983196	49,777.5	19	677	100	4.93	1.991E-61
35	Alpha tubulin of <i>T. brucei</i> (TbTUBA)	gi 115504283	49,755.5	21	646	100	4.93	2.506E-58
36	Elongation factor 1 α (TcEF1A)	gi 71408910	49,083.4	19	416	100	8.93	5E-36
37	Ribokinase (TcRBKS)	gi 71659743	35,262.6	7	158	100	6.95	3.155E-10
38	Heat shock protein 70 partial (TcHSP70)	gi 10626	74,233.7	31	626	100	5.64	2.506E-56
39	Heat shock protein HSP70 (TcHSP70)	gi 162117	71,210.1	30	623	100	5.42	5E-56
40	Heat shock 70-kDa protein, mitochondrial (TcHSP70-10)	gi 71407515	70,946	24	516	100	5.75	2.506E-45
Spots differentially expressed in SYL (compared to TCC)								
53	Cytochrome c oxidase subunit IV (TcCOXIV)	gi 71667854	38,884.4	15	339	100	5.72	2.506E-28
54	Tryparedoxin peroxidase (TcTXNPx)	gi 71413207	22,376.1	10	357	100	5.69	3.972E-30
55	Tryparedoxin peroxidase (TcTXNPx)	gi 4388655	25,506.1	6	134	100	7.62	7.924E-08
56	Elongation factor alpha G5 (TcEFA-G5)	gi 52424046	49,067.4	18	405	100	8.93	6.295E-35
57	Heat shock protein 70 (TcHSP70)	gi 50659756	71,329.9	30	524	100	5.06	3.972E-46
58	Glucose-regulated protein 78 (TcHSPA5 or TcGRP78)	gi 71415505	71,271.9	29	510	100	5.09	9.976E-45
59	Glucose-regulated protein 78 (TcHSPA5 or TcGRP78)	gi 349838	71,488	25	365	100	5.07	3.155E-30
	75- to 77-kDa antigen (TcHSP70)	gi 1097750	41,716.5	17	273	100	4.99	5E-21
60	Mitochondrial heat shock protein (TcHSP70mt)	gi 205278870	70,969.1	25	527	100	5.87	1.99E-46
61	Heat shock protein 70 (TcHSP70)	gi 205278868	71,193	26	517	100	5.32	2.51E-73

(Continued on following page)

TABLE 1 (Continued)

Protein spot	Protein name (gene)	NCBI accession no.	Mol mass (Da)	Peptide count	Protein score	Protein score		E value
						(CI%)	pI	
62	<u>Alpha tubulin (TcTUBA)</u>	gi 71397525	49,767.5	22	806	100	4.94	2.51E-74
62	<u>Alpha tubulin (TcTUBA)</u>	gi 1314208	46,927.4	20	787	100	5.49	1.99E-72
63	<u>Beta tubulin 1.9 (TcTUBB)</u>	gi 18568139	49,667.8	16	643	100	4.74	5.00E-58
64	<u>Beta tubulin of T. cyclops (TcTUBB)</u>	gi 91983197	44,496.8	14	626	100	5.68	2.51E-56
65	<u>Calpain-like cysteine peptidase (TcCAPN1)</u>	gi 71409962	12,839.3	5	170	100	5.08	1.26E-11
66	Tryparedoxin peroxidase family (TcGPX1)	gi 19171158	16,057.1	5	102	99.9	5.27	9.98E-51
67	Histone H2B of <i>T. brucei</i> (TbHIST1H2B)	gi 71748436	12,588.8	4	84	96.1	11.77	1.26E-54
68	<u>Beta tubulin (TcTUBB)</u>	gi 71656281	49,667.7	15	570	100	4.7	9.51E-56
69	<u>Alpha tubulin of T. damilewskyi (TdTUBA)</u>	gi 74229924	49,765.6	21	722	100	4.94	6.29E-66
70	<u>Cofilin-actin depolymerizing factor (TcCFL1)</u>	gi 71417693	15,735.8	9	300	100	5.69	9.98E-24
71	Prostaglandin F synthase (TcPGFS)	gi 71651365	32,489.5	16	532	100	7.71	6.29E-47
72	Hypothetical protein	gi 71657351	129,881.1	18	77	79.8	6.84	7.92E-41
73	Lipophosphoglycan biosynthetic protein (TcLPG3)	gi 71663313	86,968.5	17	248	100	5.26	1.58E-18
74	3-Hydroxy-3-methylglutaryl coenzyme A synthase-like (TcHMGCS2)	gi 71661816	54,719.2	14	445	100	6.17	3.15E-38
75	Carboxypeptidase (TcCPD)	gi 71667929	57,626.9	19	136	100	5.92	2.51E-07
	Metalloprotease (TcCPD)	gi 47076074	57,686.9	18	127	100	5.85	1.99E-06

^a *T. cruzi* epimastigotes (COL, SYL, and TCC isolates) were cultured in LIT media. Protein homogenates were resolved by two-dimensional gel electrophoresis. Gel images were analyzed on Progenesis SameSpots software, and normalized spot volumes were used to compare different groups. Proteins spots with a >2-fold change were subjected to MALDI-TOF MS/MS analysis. The number of peptides for each protein spot for which peptide mass fingerprint was obtained by MALDI-TOF MS is listed. Protein spots that were identified with high confidence by Mascot search engine (a protein score of >62) and statistically significant at the 95% confidence interval (CI%) are listed. The probability of match of the peptides in database search was accepted as significant at an E value of <0.001. pI, isoelectric pH. Underlined text indicates proteins involved in host cell invasion and cytoskeletal remodeling and motility. Boldface text indicates proteins of the antioxidant family involved in redox homeostasis.

Tukey]) and ONOO⁻ (64.5% and 2.1-fold, respectively; $P < 0.001$ [ANOVA-Tukey]) stress (Fig. 6).

Next, we sought to determine whether TcI isolates exhibited differing capacities to modulate intracellular oxidative/nitrosative stress. For this, macrophages were incubated with *T. cruzi* trypomastigotes for 24, 36, and 48 h, and the intracellular ROS was measured by using a CM-H₂DCF-DA fluorescence probe. Macrophages infected with COL and SYL for 24 h exhibited minimal intracellular DCF fluorescence, while TCC-infected macrophages demonstrated a significant increase in DCF fluorescence compared to that noted in normal controls (Fig. 7A, compare panels a and b to panel c). A similar pattern of increased intracellular DCF fluorescence in TCC-infected macrophages, compared to COL- or SYL-infected macrophages, was observed at 36 h and 48 h p.i. (data not shown).

Subsequently, we measured the release of H₂O₂ (Amplex red assay) and nitrate/nitrite (Griess assay) from the infected macrophages. The H₂O₂ levels in supernatants of macrophages infected with COL and SYL isolates for 24 or 48 h were not statistically different from that noted in normal controls (Fig. 7B). TCC-infected (versus SYL- or COL-infected) macrophages exhibited up to 1.5- and 2.3-fold increases in H₂O₂ levels at 24 and 48 h, respectively (Fig. 7B; all, $P < 0.01$ [ANOVA-Tukey]). The nitrate/nitrite release, an indicator of iNOS activity, was increased by 1.6- to 2.7-fold, 1.7- to 2.9-fold, and 3.7- to 21-fold in COL-, SYL-, and TCC-infected macrophages, respectively (Fig. 7C). The maximal difference in nitrate/nitrite level between COL- and SYL-infected versus TCC-infected macrophages was noted at 24 h p.i. (Fig. 7C, $P < 0.001$ [ANOVA-Tukey]).

To verify that macrophages infected with TCC may simply not be activated, we monitored cytokine (IFN- γ , TNF- α , and IL-6) release in the supernatants of infected cells by an ELISA. These data showed that, irrespective of the parasite strain used, patent increases in TNF- α (33.5- to 35.5-fold) and IL-6 (>100-fold) re-

lease were mounted by macrophages within 24 h p.i., and the cytokine release was maintained at 36 and 48 h p.i. (Fig. 7D and E). No increase in IFN- γ release was observed in any of the infected macrophages. Together, the data presented in Fig. 7 suggested that COL and SYL (versus TCC) trypomastigotes faced compromised oxidative and nitrosative stress in macrophages. The differences in antioxidant capacity of the parasite isolates had no effect on cytokine signaling in macrophages.

Finally, we sought to determine whether the differential abundance of antioxidants conferred a survival advantage to COL and SYL trypomastigotes in macrophages. For this, RAW 264.7 macrophages were infected with *T. cruzi* isolates for 6, 12, and 18 h. Infected cells were submitted to Giemsa staining, and >200 cells/slide (triplicate experiments) were evaluated by light microscopy to determine the number of infected macrophages. We observed no statistically significant difference in the parasite uptake by macrophages incubated with COL, SYL, or TCC isolates (data not shown). A quantitative measure of intracellular parasites by real-time quantitative PCR showed a significant increase in intracellular SYL, followed by COL and TCC, from 0 to 18 h p.i. (Fig. 8A). The rapid replication of SYL in macrophages was evidenced by detection of Tc18SrDNA signal that was 12.6- to 13.5-fold and >27-fold higher than that noted in COL- and TCC-infected macrophages, respectively, from 6 to 12 h p.i. ($P < 0.01$ to 0.001 [ANOVA-Tukey]). By 18 h p.i., intracellular COL level was 7.6-fold higher than that noted for TCC ($P < 0.01$ [t test]) but lagged behind that of SYL by 3.2-fold ($P < 0.01$ [t test]).

We used metacyclic trypomastigotes derived from stationary-phase epimastigote cultures of COL, SYL, and TCC isolates to monitor their ability to escape from macrophages. For this, parasites were added to monolayers of RAW 264.7 macrophages and, after infection for 3 h, the cells were washed and incubated for 2 weeks. The hemacytometer counting of parasites in supernatants

TABLE 2 Protein spots that were differentially expressed in trypomastigote stage of TcI isolates of *T. cruzi*^a

Protein spot	Protein name (gene)	NCBI accession no.	Mol mass (Da)	Peptide count	Protein score	Protein		
						score (CI%)	pI	E value
Spots differentially expressed in COL isolate compared to TCC								
1	Tryparedoxin peroxidase (TcTXNPx)	gi 17224953	22,246.1	10	340	100	5.96	1.99054E-28
2	Tryparedoxin peroxidase (TcTXNPx)	gi 71399514	22,375.1	10	304	100	6.32	3.97164E-24
5	<u>Calpain-like cysteine peptidase (Tc CL)</u>	gi 71411006	12,798.3	5	164	100	5.07	7.92447E-11
6	Tryparedoxin peroxidase family (TcGPX1)	gi 19171158	16,057.1	4	82	98.73	5.27	1.25674E-05
7	<u>Paraflagellar rod protein 3 (TcPFR3)</u>	gi 2226084	68,705.3	24	288	100	5.86	3.15479E-23
8	<u>Paraflagellar rod protein, 69 kDa (TcPFR)</u>	gi 71650963	69,518.9	26	215	100	5.85	6.29463E-16
9	<u>Paraflagellar rod protein 2 (TcPFR2)</u>	gi 162179	69,504.9	24	198	100	5.85	3.15479E-14
10	<u>Paraflagellar rod protein of <i>T. brucei</i> (TbPFR-C/TbPFR-D)</u>	gi 72387193	68,640.3	19	175	100	5.78	6.29463E-12
11	Heat shock protein 70 (TcHSP70/TcHSPA5)	gi 50659756	71,329.9	33	877	100	5.06	3.97164E-82
12	Glucose-regulated protein, 78 kDa (TcGRP78)	gi 349838	71,488	29	603	100	5.07	9.97631E-55
13	<u>Alpha tubulin (TcTUBA)</u>	gi 71397525	49,767.5	19	405	100	4.94	6.29463E-35
14	<u>Alpha tubulin (TcTUBA)</u>	gi 1314208	46,927.4	18	398	100	5.49	3.15479E-34
15	<u>Beta tubulin of <i>T. grayi</i> (TgTUBB)</u>	gi 91983201	42,806.9	13	288	100	5.29	3.15479E-23
16	<u>Beta tubulin of <i>T. danilewskyi</i> (TdTUBB)</u>	gi 74229926	49,695.8	13	281	100	4.74	1.58114E-22
17	Cytochrome <i>c</i> oxidase subunit IV (TcCOXIV)	gi 71667854	38,884.4	15	339	100	5.72	2.50594E-28
18	<u>Kinetoplastid membrane protein 11 (TcKMP11)</u>	gi 71413455	10,876.3	8	349	100	5.73	1.25594E-28
19	<u>Kinetoplastid membrane protein 11 of <i>T. rangeli</i> (TrKMP11)</u>	gi 32892042	11,003.4	8	347	100	6.52	1.99054E-28
20	Phosphoglycerate kinase 1 (TcPGK1)	gi 71411859	44,441.1	16	240	100	6.19	9.97631E-18
21	3-Phosphoglycerate kinase, glycosomal (TcPGKC)	gi 71654900	54,867.6	15	122	100	6.94	6.29463E-06
22	Chain A, tryparedoxin peroxidase (TcTXNPx)	gi 58176947	22,743.3	8	305	100	5.71	3.15479E-24
23	Peroxioredoxin, antioxidant oxidoreductase (TcPRX)	gi 4388655	20,235.2	9	440	100	5.07	9.97631E-38
24	Ubiquitin precursor (TcUBQ)	gi 162337	40,105.6	5	409	100	9.39	1.25594E-34
25	Fe-superoxide dismutase, chain A (TcFeSODA)	gi 158428656	21,800.8	7	441	100	6.65	7.92447E-38
26	Fe-superoxide dismutase (TcFeSOD)	gi 1066332	21,818.9	7	440	100	6.64	9.97631E-38
27	<u>Calpain-like cysteine peptidase (TcCAPN1)</u>	gi 71409962	12,839.3	4	159	100	5.08	1.25594E-09
29	Ascorbate-dependent peroxidase (TcAPX)	gi 24370982	31,855.2	7	269	100	6.69	1.25594E-20
Spots differentially expressed in SYL isolate compared to TCC								
30	<u>Beta tubulin of <i>T. cyclops</i> (TcTUBB)</u>	gi 91983197	44,496.8	14	295	100	5.68	6.29463E-24
31	<u>Beta tubulin 1.9 (TcTUBB)</u>	gi 18568139	49,667.8	14	291	100	4.74	1.58114E-23
32	Prostaglandin F2 alpha synthase (TcPGFS)	gi 71661165	14,648.3	4	119	100	6.05	2.50594E-06
33	Amidinotransferase-like (TcGATM)	gi 71661567	43,051.9	9	184	100	6	7.92447E-13
34	Amidinotransferase-like (TcGATM)	gi 71415202	28,667.4	5	128	100	5.21	3.15479E-07
35	10-kDa heat shock protein (TcHSP1)	gi 71425263	10,692.8	2	66	49.41	9.05	2.50594E-18
36	Tryparedoxin peroxidase, chain A (TcTXNPX)	gi 58176947	22,743.3	9	328	100	5.71	3.15479E-27
37	<u>Alpha tubulin of <i>T. danilewskyi</i> (TdTUBAi)</u>	gi 74229924	49,765.6	18	394	100	4.94	7.92447E-34
38	<u>Tubulin alpha (TcTUBA)</u>	gi 3915082	49,696.5	17	358	100	4.9	3.15479E-30
39	NADPH flavin oxidoreductase (old yellow enzyme, TcOYE)	gi 61741942	42,191.6	12	250	100	6.03	1.99054E-19
40	Glutamate dehydrogenase (TcGLUD1)	gi 71414987	41,680.5	11	239	100	7.1	2.50594E-18
41	Calmodulin (TcCALM1)	gi 71407658	26,740.2	5	89	100	5.1	3.15479E-17
42	Cysteine peptidase inhibitor (TcSERPIN)	gi 71664690	12,114	2	89	98.63	6.58	0.001256238
43	Ubiquitin homolog (TcUBQ)	gi 10673	14,648.8	5	404	100	9.82	3.97164E-34
44	Vacuolar ATP synthase subunit B-like (TcATP5B)	gi 70870795	104,474.5	30	656	100	5.75	2.50594E-59
47	Heat shock protein 85 (TcHSP85)	gi 70876495	82,887.1	18	680	100	5.79	9.97631E-62
48	Ubiquitin-like (TcUBQ)	gi 136677	8,503.6	5	427	100	6.56	1.99054E-36
49	Ubiquitin-like (TcUBQ)	gi 10674	8,762.7	5	425	100	6.56	3.15479E-36
50	<u>Kinetoplastid membrane protein 11 (TcKMP11)</u>	gi 6166388	9,923.9	8	351	100	6.58	7.92447E-29
51	<u>Kinetoplastid membrane protein 11 of <i>T. rangeli</i> (TrKMP11)</u>	gi 24286543	11,048.4	8	349	100	5.96	1.25594E-28
52	Cysteine-protease inhibitor/Chagasin (TcSERPIN)	gi 118137894	14,448	3	185	100	6.66	3.15479E-12

(Continued on following page)

TABLE 2 (Continued)

Protein spot	Protein name (gene)	NCBI accession no.	Mol mass (Da)	Peptide count	Protein score	Protein		
						score (CI%)	pI	E value
53	Cysteine peptidase inhibitor (TcSERPIN)	gi 71664690	12,114	2	127	100	6.58	1.99054E-06
54	Hypothetical protein (Tc CL)	gi 71410630	12,677.5	2	71	23.03	6.58	1.99054E-12
55	Tryparedoxin peroxidase (TcTXNPX)	gi 71413207	22,376.1	8	419	100	5.69	1.25594E-35
56	Thioredoxin-dependent peroxide reductase (PRDX3)	gi 241785831	25,789.7	4	214	100	9.08	3.97164E-15
57	Iron superoxide dismutase (TcFeSOD)	gi 71414374	23,339.6	7	439	100	6.7	1.25594E-37
58	Iron superoxide dismutase B (TcFeSODB)	gi 2149612	21,745.8	6	285	100	6.37	3.15479E-22
59	Heat shock 70-kDa protein (TcHSP70)	gi 71406304	40,831	18	720	100	6.19	9.97631E-66
60	Heat shock protein HSP70 (TcHSP70)	gi 162117	71,210.1	19	685	100	5.42	3.15479E-62
61	Heat shock protein 70 (TcHSP70)	gi 10626	74,233.7	17	609	100	5.64	1.25594E-54
62	Ubiquitin-conjugating enzyme-like (Tc CL)	gi 71414042	15,754.8	9	157	100	5.81	1.99054E-09
63	<u>Beta tubulin 1.9 (TcTUBB)</u>	gi 18568139	49,667.8	12	227	100	4.74	1.99054E-16
64	<u>Beta tubulin (TcTUBB)</u>	gi 3915883	49,540.7	11	262	100	4.69	1.25594E-20
65	<u>Alpha tubulin (TcTUBA)</u>	gi 1314208	46,927.4	18	400	100	5.49	9.97631E-34
66	<u>Alpha tubulin (TcTUBA)</u>	gi 71397525	49,767.5	18	396	100	4.94	2.50594E-33
67	<u>Calpain-like cysteine peptidase (TcCAPN1)</u>	gi 71411006	12,798.3	5	177	100	5.07	1.99054E-11
68	3-Phosphoglycerate kinase (TcPGK)	gi 16945527	54,767.4	15	121	100	6.94	7.92447E-06
69	Phosphoglycerate kinase (TcPGKCL)	gi 71411862	64,528.8	14	114	100	8.04	3.97164E-05
70	Tryparedoxin peroxidase (TcTXNPX)	gi 17224953	22,246.1	9	492	100	5.96	6.29463E-43
71	<u>Cruzipain</u>	gi 1136308	47,852.1	3	348	100	5.35	1.58114E-28
72	<u>69-kDa paraflagellar rod protein (TcPFR)</u>	gi 71411387	42,471.9	18	153	100	6.12	9.97631E-10
73	<u>Surface glycoprotein, 82 kDa (TcGP82)</u>	gi 6049148	59,591.9	20	170	100	6.05	9.97631E-11

^a *T. cruzi* trypomastigotes (COL, SYL, and TCC isolates) were cultured in C2C12 cells. Parasite protein homogenates were resolved by two-dimensional gel electrophoresis. Gel images were analyzed on Progenesis SameSpots software, and normalized spot volumes were used for comparison of different groups. Proteins spots with >2-fold change were subjected to MALDI-TOF MS/MS analysis. The significance of a protein match, based on the peptide mass fingerprint in the MS and the MS/MS data, is presented as expectation values ($P < 0.01$). pI, isoelectric pH. Underlined text indicates proteins involved in host cell invasion and cytoskeletal remodeling and motility. Boldface text indicates proteins of the antioxidant family that are involved in redox homeostasis.

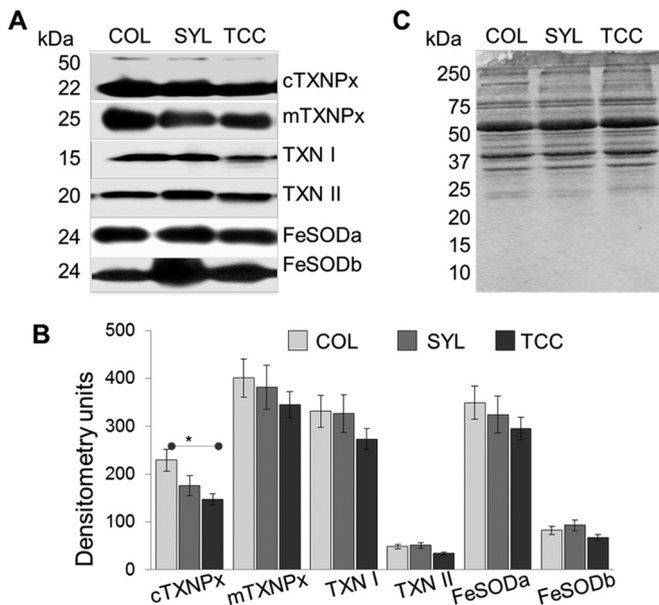


FIG 4 Western blotting for selected proteins of the antioxidant network in TcI epimastigotes. (A) Homogenates of the COL, SYL, and TCC epimastigotes were resolved by SDS-PAGE. Membranes were probed by using specific antibodies against mitochondrial and cytosolic tryparedoxin peroxidases (mTXNPx and cTXNPx, respectively), tryparedoxin substrates (TXN I and TXN II), and Fe²⁺ SODs (FeSODa and FeSODb). (B) Bar graphs show densitometric analysis of the band intensities from Western blotting, normalized to total loaded protein. (C) Ponceau S staining shows equal loading of all samples.

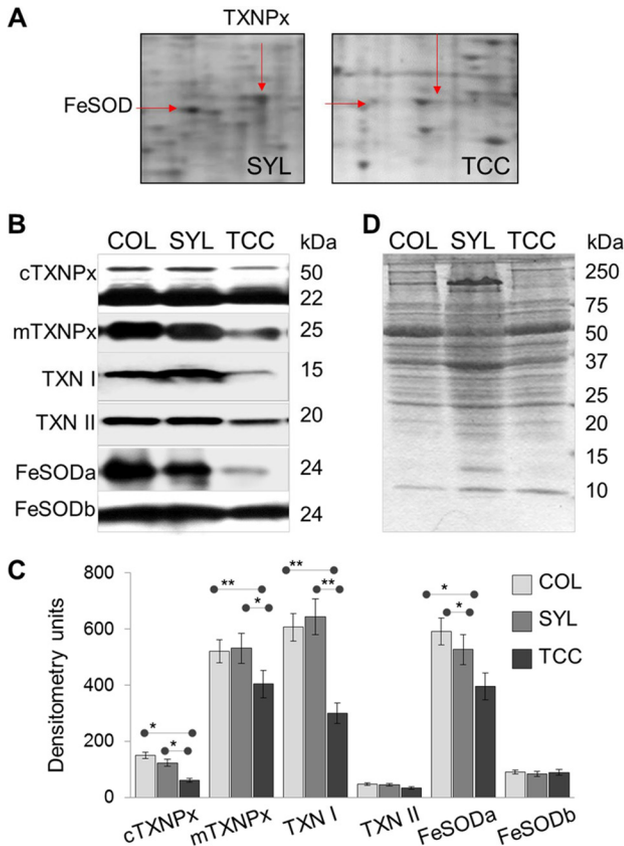
of infected cells at 7 days p.i. showed 36.8- and 24.3-fold-higher levels of parasite release from macrophages infected with COL and SYL isolates, respectively, compared to that observed in supernatants of TCC-infected macrophages (Fig. 8B, $P < 0.01$ to 0.001 [ANOVA-Tukey]). By the end of the second week postinfection, all cells infected with SYL and COL had ruptured, releasing parasites, whereas TCC-infected macrophages continued to exhibit intracellular parasites (data not shown). These data suggested that (i) macrophages were equally competent in their ability to phagocytose different isolates of *T. cruzi*, (ii) COL and SYL (versus TCC) were better equipped to survive and replicate in macrophages, and (iii) COL and SYL trypomastigotes succeeded in efficient release from macrophages.

DISCUSSION

Reported cases of Chagas disease from Latin America have been associated with TcI isolates (10, 11, 37, 38). It has been postulated that TcI isolates exhibit more tissue tropism than do other lineages of *T. cruzi* (39–41) and are therefore responsible for severe pathology of the heart. In this context, we performed biological characterization of three *T. cruzi* isolates of the TcI DTU and showed that COL, SYL, and TCC isolates exhibited differing growth and virulence behavior in mice (Fig. 1). The striking difference in the infectivity and disease severity pattern of COL and SYL versus that of TCC made them an ideal model for global proteomic profiling for the identification of candidates involved with the mechanism of parasite invasion, replication, and survival in the host cells.

The comparative proteome profiling of the three TcI strains that exhibited different virulence levels in mice yielded novel find-

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ings. One, the expression profiles of numerous proteins in COL and SYL epimastigotes and trypomastigotes were consistent with enhanced motility and contractility. For example, upregulation of α -actin, α -tubulin, β -tubulin, cofilin-1, and KMP11 that are central to remodeling of the cytoskeleton and modulation of cell shape for migration (42, 43) was noted in COL and SYL (versus TCC) strains (Tables 1 and 2). Likewise, members of the paraflagellar rod (PFR) and paraxonemal protein families, known to be essential for flagellar wave frequency and amplitude and forward motility of the parasite (reviewed in reference 44), were enhanced in abundance in COL and SYL (versus TCC) strains. Further, SYL and COL trypomastigotes also exhibited an increased abundance of calpain-like cysteine peptidases, surface glycoprotein gp82, and cruzipain that are involved in host cell attachment and invasion (45, 46). We surmise that virulence of the COL, SYL, and possibly other isolates of the TcI DTU is determined by the motility capacity that ensures parasite transmission from triatomines and dissemination to tissues in the mammalian host. Indeed, several studies using RNA interference against *PFR2* in *T. brucei* have shown that ablation of *PFR2* resulted in sluggish parasites that

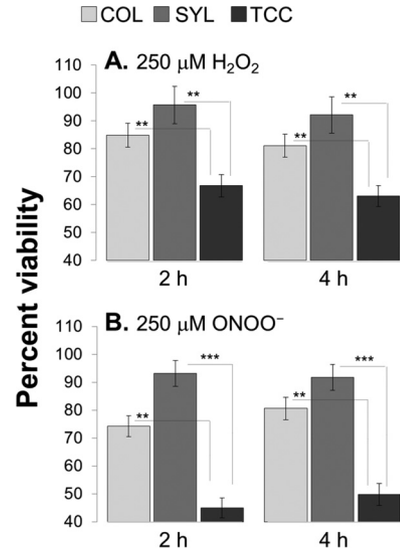


FIG 6 Variable resistance of TcI strains to oxidants. The COL, SYL, and TCC trypomastigotes were *in vitro* incubated with 250 μ M H₂O₂ (A) or 250 μ M ONOO⁻ (B). Parasite viability was determined at 2 and 4 h postincubation by an alamarBlue assay. Data are presented as the percentage of viability with respect to untreated parasites.

lacked cytokinesis capability and were monstrous in size, also showing multiple nuclei, kinetoplasts, and flagella; these PFR2-deficient cells were unable to survive *in vitro* or *in vivo* (47, 48). Further studies will be required to validate the importance of cytoskeletal remodeling and flagellar function as new targets for possible intervention against *T. cruzi* infection.

Comparative proteome analysis of the two parasite forms showed that the differentiation of epimastigotes to trypomastigotes was accompanied by a shift from histidine catabolism to glutamate, a decline in ribosomal proteins, and an increase in the expression level of a battery of antioxidant enzymes (compare Tables 1 and 2). Three major families of peroxidases identified in trypanosomatids included the 2-cysteine peroxiredoxins, as well as the non-selenium glutathione and ascorbate peroxidases (49). Notably, we showed a significant correlation between the abundance of the antioxidant enzymes and the virulent behavior of the COL and SYL trypomastigotes. The increase in trypanredoxin peroxidases occurred in infective trypomastigote and replicative amastigote stages irrespective of the cell types (e.g., muscle cells in this study, and LLC-MK2 and macrophages in other studies [34, 50]) used for culturing the parasites. Regeneration of the reduced TXNPx depends on cytosolic (TXN I)- and membrane (TXN II)-localized thioredoxin substrates (31, 51). Consistent with the increased abundance of TXNPx, the TXN I level was also increased in COL and SYL trypomastigotes (Fig. 5). COL and SYL trypomastigotes also showed an increased abundance of the FeSODa (mitochondrial) enzyme (Fig. 5). Others have also documented the differential expression of peroxiredoxin and FeSOD proteins of the antioxidant network in parasites in correlation with *in vivo* infectivity and pathological behavior (32, 34, 52). The question arises as to how the TXNPx/TXN antioxidant network is associated with the virulence of the TcI isolates of *T. cruzi*. One explanation could be that, upon infection, *T. cruzi* faces the hostile environment of phagocytes, and an increase in antioxidant response would allow adaptation of the infective forms to the oxi-

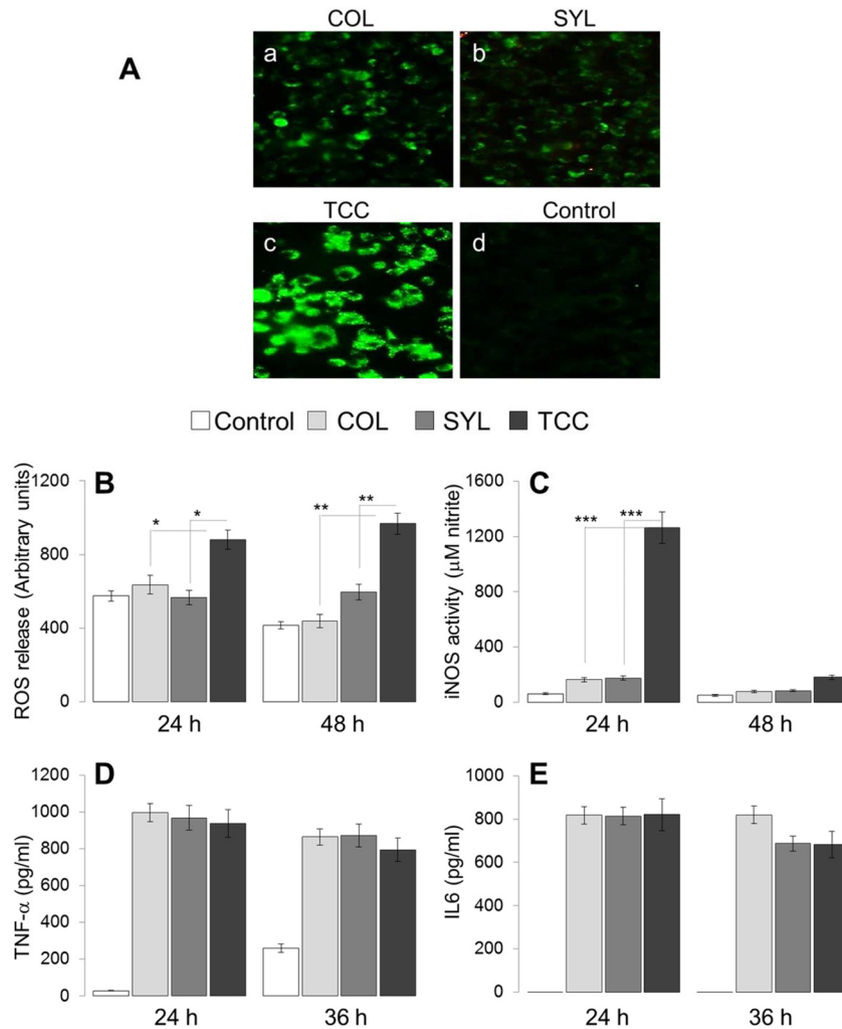


FIG 7 Macrophage activation of oxidative/nitrosative response, but not inflammatory cytokine response, is deregulated by the antioxidant status of the TcI strains of *T. cruzi*. RAW 264.7 macrophages were infected with COL, SYL, and TCC trypomastigotes for 24, 36, or 48 h. Normal (uninfected) cells, incubated for similar time periods, were used as controls. (A) Cells were incubated for 24 h and then loaded with H₂DCFDA. The oxidation of H₂DCFDA by intracellular ROS, resulting in the formation of fluorescent DCF, was monitored by fluorescence microscopy. (B and C) Supernatants were used to measure ROS (B) and nitrite (C) levels by an Amplex red and a Griess reagent assay, respectively. (D and E) TNF-α (D) and IL-6 (E) cytokines in supernatant samples were monitored by an ELISA.

oxidative environment of phagocytic cells in the mammalian host (45, 46). Activated macrophages provide a key defense response to the invading pathogens and exert cytotoxic effects via NADPH oxidase (NOX)-mediated production of superoxide and ROS that result in enhanced oxidative modification of DNA, proteins, and lipids (53, 54). Alternatively, superoxide may combine with nitric oxide to produce peroxynitrite and peroxynitrous acid that can directly kill *T. cruzi* (55). Indeed, COL and SYL (versus TCC) exhibited an enhanced capacity to survive in and escape from macrophages (Fig. 8), and these responses were associated with detection of low levels of oxidative and nitrosative stress in COL-infected and SYL-infected (versus TCC-infected) macrophages (Fig. 7). The *in vitro* (Fig. 7) and *in vivo* (Fig. 1) data presented in this study do not clarify whether the antioxidant network enhanced the capacity of the SYL and COL isolates to efficiently scavenge ROS and nitric oxide (NO) or inhibit the macrophage activation of oxidative and nitrosative response. Yet, our finding of increased resistance of COL and SYL trypomastigotes to exog-

enous H₂O₂ and peroxynitrite-mediated killing (Fig. 6) implies that the parasite's ability to scavenge ROS/NO, at least partially, provides it a survival advantage in immune cells. Others have provided direct biochemical evidence showing that TXNPPx acts as a hub to neutralize free radicals through peroxynitrite reductase activity (56). An enhanced association between the protein level and activity of trypanothione peroxidases (TXNPPx) and virulence of other isolates of DTU TcI-TcIV has also been noted (35, 56). Thus, we surmise that intraphagosomal survival from ROS and NO is the key to the parasite's existence in the host and that, irrespective of the phylogenetic lineage, *T. cruzi* isolates capable of utilizing an antioxidant network to avoid oxidative damage would likely ensure their survival, dissemination, and persistence in the host. The increase in the expression of FeSODA in SYL and COL is in line with our interpretation, since the dismutation of superoxide will lead to a decrease in the concentration of oxidative species.

It is important to note that after escaping from macrophages,

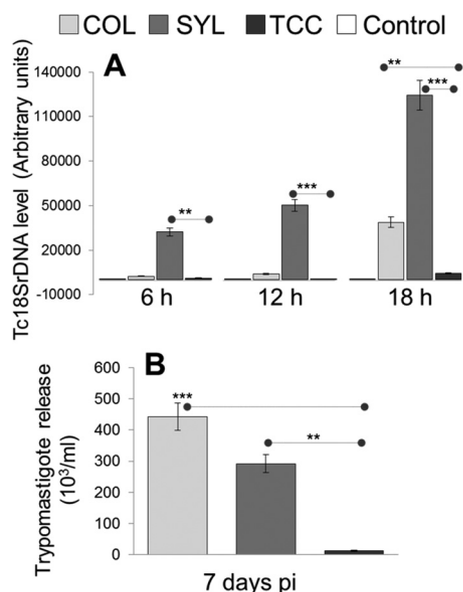


FIG 8 Intracellular replication of TcI isolates and release from infected macrophages. (A) RAW 264.7 macrophages were infected with COL, SYL, and TCC for 6, 12, and 18 h. Real-time quantitative PCR analysis of parasite burden in infected macrophages was performed by using Tc18SrDNA-specific oligonucleotides. The data were normalized to GAPDH expression. Normal/uninfected cells were used as controls. (B) RAW 264.7 macrophages were infected with metacyclic form of COL, SYL, and TCC, and the release of trypomastigotes in supernatants was determined 7 days p.i. by light microscopy.

T. cruzi faces oxidative stress in other tissues as well. For example, we have shown enhanced mitochondrial production of ROS in cardiomyocytes during *T. cruzi* infection (33). It is plausible that *T. cruzi* utilizes antioxidant enzyme network to prevent mitochondrial ROS-mediated direct killing or ROS activation of the apoptosis pathway for removal of the infected (or damaged) cells (57), thereby ensuring its persistence in the host. A consequence of prolonged survival of infected (damaged) cells is the signaling of cell repair and proliferative responses, culminating in cardiac fibrosis (58). Further studies are required to demonstrate the novel role of TXNPs/TXN in establishing a cardiac remodeling environment in Chagas disease; however, if such is the case, inhibiting the parasite antioxidant activity will provide a novel strategy to manage the homeostatic clearance of damaged cells (and parasite) and control the fibrotic processes that culminate in cardiomyopathy in chronic Chagas disease.

In summary, we conclude that *T. cruzi* virulent isolates ensure survival and persistence in the host via peroxiredoxin-dependent decomposition of macrophage-derived free radicals. This is evidenced by our findings in the infective stage of the virulent parasite isolates (COL and SYL), i.e., an upregulation of the protein levels for cTXNPx, mTXNPx, and TXN I that readily detoxified cytotoxic and diffusible reactive oxygen and nitrogen species generated *in vitro* or released by the macrophages. The absence of differences in inflammatory cytokine levels (i.e., TNF- α and IL-6) in macrophages infected with virulent or nonvirulent isolates of *T. cruzi* suggests that the peroxiredoxin-based antioxidant system only helps the parasite to cope with the oxidative challenge without interfering with the cytokine signaling. Further characterization of the parasite's antioxidant network in tissues from infected experimental animals at different stages of infection and disease

development would support its role in the pathogenesis and parasite persistence and provide a strong basis for the development of new chemotherapeutic options against *T. cruzi*.

T. cruzi isolates have been characterized as belonging to the TcI to TcVI lineages. Efforts establishing an association between parasite lineage and pathogenicity have not been successful. We demonstrate that parasite isolates of the same lineage exhibit a variable virulence in mice. An unbiased proteomic approach, followed by targeted studies, showed that the differential abundance of the peroxiredoxin antioxidants and the flagellar proteins allowed intraphagosomal survival from oxidative/nitrosative stress and parasite dissemination and persistence. Thus, gene expression conducive to enhanced antioxidant capacity and motility, irrespective of phylogenetic lineage, provided virulence to *T. cruzi* and should be evaluated for identifying virulent isolates circulating in the Americas.

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