TcI, TcII and TcVI Trypanosoma cruzi samples from Chagas disease patients with distinct clinical forms and critical analysis of in vitro and in vivo behavior, response to treatment and infection evolution in murine model

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ARTICLE INFO
Article history:
Received 10 August 2016
Received in revised form 11 November 2016
Accepted 24 November 2016
Available online 28 November 2016

Keywords:
Trypanosoma cruzi
TcI
TcII
TcVI DTUs
Biological characteristics
Drug response
Human disease clinical forms

ABSTRACT
The clonal evolution of Trypanosoma cruzi sustains scientifically the hypothesis of association between parasite’s genetic, biological behavior and possibly the clinical aspects of Chagas disease in patients from whom they were isolated. This study intended to characterize a range of biological properties of TcI, TcII and TcVI T. cruzi samples in order to verify the existence of these associations. Several biological features were evaluated, including in vitro epimastigote-growth, “Vero” cells infectivity and growth, along with in vivo studies of parasitemia, polymorphism of trypomastigotes, cardiac inflammation, fibrosis and response to treatment by nifurtimox during the acute and chronic murine infection. The global results showed that the in vitro essays (acelluar and cellular cultures) TcII parasites showed higher values for all parameters (growth and infectivity) than TcVI, followed by TcI. In vivo TcII parasites were more virulent and originated from patients with severe disease. Two TcII isolates from patients with severe pathology were virulent in mice, while the isolate from a patient with the indeterminate form of the disease caused mild infection. The only TcVI sample, which displayed low values in all parameters evaluated, was also originated of an indeterminate case of Chagas disease. Response to nifurtimox was not associated to parasite genetic and biology, as well as to clinical aspects of human disease. Although few number of T. cruzi samples have been analyzed, a discreet correlation between parasite genetics, biological behavior in vitro and in vivo (murine model) and the clinical form of human disease from whom the samples were isolated was verified.

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1. Introduction

Chagas disease or American trypanosomiasis, caused by the hemoflagellate protozoan Trypanosoma cruzi, was discovered in 1909 by Chagas (1909) and are present in twenty one countries in the Americas (WHO, 2015). New cases of Chagas disease are daily diagnosed in several countries and continents, originally considered non-endemic for Chagas disease (WHO, 2015) where the disease has been transmitted independent of the natural vectors of the parasite such as blood transfusion, congenital, transplantation of organs and tissues and others (Schmunis and Yadon, 2010).

The short-term acute phase of Chagas disease evolves to distinct clinical manifestations during long-lasting chronic phase, ranging from asymptomatic to cardiac, digestive (megalocolon and/or megaesophagus) or cardiac/digestive clinical forms (Rassi et al., 2010).

Currently, six distinct T. cruzi genetic subdivisions or DTU’s (discrete typing units) have been identified (Zingales et al., 2009). The clonal nature of T. cruzi reproduction and propagation is expected to be the basis of genetically stable populations, although the phenomenon of hybridization may rarely occur in this species (Tibayrenc and Ayala, 2015). For these reasons, T. cruzi infection is considered the best model for to study the population genetics and the molecular epidemiology. However, the correlation between the genetic variability and the presence of different clinical forms is still questionable (Lages-Silva et al., 2006; D’Ávila et al., 2009). Several studies have intended to correlate the high rate of T. cruzi genetic variability to its biological characteristics (Toledo et al., 2002) as well as clinical manifestations of Chagas disease, but still unsuccessfully (Macedo et al., 2004).

Different hypotheses have been proposed in an attempt to elucidate the failure to establish this correlation. One of them may be resultant of the mixtures of different T. cruzi subpopulations of distinct genotypes that may lead to different infection evolution and clinical pictures (Macedo and Pena, 2002). Another hypothesis relates the fact that the disease is the result of a multifactorial process that involves aspects of the host, but also of the parasite. The inadequate choice of genetic targets may result in lack of correlation between the clinical forms of Chagas disease with the T. cruzi genetic variability. Moreover, the occurrence of mixed infection, with distinct T. cruzi populations, and the isolation of the predominant T. cruzi genotype in peripheral blood may also appear as a confounding factor in the attempt of the detection of association between parasite genetic and the clinical forms of Chagas disease (Macedo et al., 2004).

Taking into account the current T. cruzi division into distinct genetic groups and that the biological parameters of the parasite strains influence the infection evolution and probably the clinical aspects of Chagas disease (Macedo et al., 2004), we believe that to better understand the current disease of an endemic region firstly is necessary to know the parasite genetic profile affecting the local population. Therefore, the knowledge of parasite genetics would offers notions about patient’s treatment decision, the clinical aspects of the disease, as well as how to establish better epidemiological surveillance and control strategies for Chagas disease (Couro and Junqueira, 2015; Martinez-Perez et al., 2016).

Considering these important statements, the current study aims to characterize a range of biological features of TcI, TcII and TcVI T. cruzi samples from Chagas disease patients with distinct clinical forms of the disease in order to verify possible correlations among the biological aspects of the parasites and the clinical aspects of the patients from whom these samples were isolated.

2. Methods

2.1. T. cruzi samples

The T. cruzi samples were obtained from chronic Chagas disease patients of the endemic areas, Berilo and Januária, Minas Gerais (MG) State, Brazil with different clinical forms. The six T. cruzi samples were properly genotyped by us (Oliveira et al., 2015) as previously proposed by Lewis et al. (2009) and D’Ávila et al. (2009). The criteria of D’Ávila et al. (2009) genotyping was adopted in order to identify safely the distinct DTUs as well as mixed infections in the samples. No evidences of mixed infection were verified by Oliveira et al. (2015). Samples of TcI and TcVI DTUs were chosen because they are present in the endemic region where we have worked in the last years. Although TcI samples were not detected in Berilo, one sample of this DTU was included due its high distribution and importance in Chagas disease epidemiology.

The only sample identified as TcI genotype (sample code: Pr 150) was isolated from a patient presenting the cardiac clinical form (Abolis et al., 2011); three samples were identified as TcI genotype (sample code: 501, 452, 728) and were isolated from patients presenting the indeterminate, cardiac and cardiac/digestive clinical forms, respectively; and two samples were identified as TcVI genotype (sample code: 1337 and 748) and were isolated from patients presenting the indeterminate and cardiac/digestive clinical forms, respectively. The patients were clinically classified according to Brazilian Consensus in Chagas disease (Ministério da Saúde, 2005), based on anamnesis, clinical and physical examination, thoracic and contrasted gastrointestinal tract X-ray along with electrocardiogram after consensus between two independent physicians. The inclusion of patients in the study occurred after signature of the Informed Consent approved by the Ethics Committee on Human Research at the Centro de Pesquisa Renê Rachou, FIOCRUZ, Belo Horizonte, MG, Protocol: 007/2002 and at the Universidade Estadual de Maringá, Paraná state, Brazil. The Table 1 summarizes the main features of T. cruzi samples evaluated.

All T. cruzi samples were isolated by hemoculture (HC) technique as previously described by Chiari et al. (1989). Parasites were growth in LIT medium and stored in liquid nitrogen at first passage in this medium for further use in this study.

2.2. T. cruzi growth in acellular culture in vitro

T. cruzi growth in vitro was evaluated for each sample in LIT medium culture plus 10% of fetal bovine serum starting with a standard inoculum of 1.0 × 10^4 parasites/ml in a final volume 3 ml (Nogueira-Paiva et al., 2015). The cultures were maintained in biological incubator BOD (FANEM® model 347) at 28 °C ± 1 °C. The number of epimastigotes was monitored daily, for 20 consecutive days, in Newbauer chamber examined in optical microscopy (4X40). All experimental batches were carried out in triplicate and data reported as number of epimastigotes/ml.

2.3. T. cruzi growth in “Vero” cells culture in vitro

The T. cruzi amastigote growth in “Vero” cells were evaluated as proposed by Andrade et al. (2010) modified as follows: The cells previously stored in liquid nitrogen stocks were after three passages in D-EMEN seeded into 75 cm^2 polystyrene sterile bottles containing 14 ml of D-MEM medium (Dulbecco’s Modified Eagle Medium, Invitrogen, Carlsbad, CA, USA), supplemented with 5% of heat-inactivated fetal bovine serum (FBS, Nutricell, Campinas, SP, Brazil), added of 2.5% 1 M HEPES pH 7.2, 1.0% of 2 mM glutamine/ml, 0.1% mercaptoethanol 50 mM/ml, 0.2% of gentamicin sulfate 200 mg/ml (Schering-Plough, Kenlworth, NJ, USA). Semi-confluent “Vero” cells monolayer were reached approximately 72 h
after seeding. The “Vero” cells were then trypsinized and approximately $4.0 \times 10^4$ cells were transferred to each well of glass cover slips and incubated at 5% CO$_2$ and at 37°C overnight, prior infection (10:1 ratio) with metacyclic T. cruzi trypanosomates previously differentiated in Grace’s medium as described by Contreras et al. (1985). Twenty four hours after infection, the “Vero” cells monolayers were washed with phosphate buffer-saline (PBS) to remove extracellular parasites and the cultures maintained under the same conditions during 24, 48 and 72 h. Following, the “Vero” cells cultures were fixed in methanol and stained with Fast Panotic dye, and amastigotes growth examined under optical microscopy (100× magnification). The “Vero” cells infectivity was defined for each T. cruzi sample in triplicates and the results expressed as: mean percentage of infected cells per 100 cells; the number of amastigotes/cell and the number of amastigotes per total number of infected cells.

### 2.4. In vivo studies in murine model

For this study female Swiss mouse, 18 and 20 g, 30 days old were used. The parameters parasitemia, bloodstream polymorphism, mortality and response to etiological treatment were evaluated in mice because they are associated to each other (Andrade, 1974) and consequently linked to the parasite genetics (Andrade and Magalhães, 1996; Toledo et al., 2002).

For each sample of T. cruzi four groups of eight mice (total 192) were inoculated intraperitoneally with 10,000 blood trypomastigotes/animal quantified according to Brener (1962), previously maintained up to three passages in Swiss mice. Two groups for each sample of T. cruzi (16 mice) were used for the biological characterization and one was the control untreated group; two others (16 mice) were used for treatment of the acute phase – AP (treatment started in the first day of patent parasitemia) and chronic phase – CP (treatment started on the 90th day of infection) with nitrofuran (nifurtimox, NF) Lampit® produced by Bayer Health Care. Parasitemia was assessed daily (starting at day 4 up to day 50 after infection) by fresh blood examination (FBE) according to Brener (1962). The biological parameters from parasitemia curves were evaluated for each T. cruzi sample, including:

- Pre-patent period (PPP): corresponding to the period (days) between T. cruzi inoculation and the day before the first parasite detection by FBE. The PPP results were expressed as mean ± standard deviation;
- Patent Period (PP): corresponding to the time span between the first and the last day of positive FBE. The PP results were expressed as mean ± standard deviation;
- Maximum Peak of parasitaemia (MPP): corresponding to the maximum number of trypomastigotes/0.1 mL of blood detected by FBE. The MPP results were expressed as mean ± standard deviation;
- Day of Maximum Peak of Parasitemia (DMPP): corresponding to the day of maximum of parasitemia occurrence. The DMPP results were expressed as mean ± standard deviation;
- Morphological characterization of bloodstream trypomastigotes: correspond to the relative of thin, intermedia and large bloodstream trypomastigotes forms observed under optical microscopy (100X) according to Brener and Chiari (1963) methodology.

**Mortality (MOR):** The mortality of the animals was assessed daily until the 90th d.a.i. and weekly until 360th d.a.i. The results were expressed in cumulative percentages.

All in vivo studies in murine models were carried out at Centro de Ciência Animal (CCA-UFO), according to the guidelines established by the Conselho Nacional de Controle em Experimentação Animal (CONCEA) and following the international guidelines. The study was previously approved by the Ethics Committee in Animal Experimentation of the Universidade Federal de Ouro Preto (UFOP) under the protocol number 26/2014.

#### 2.5. Response to nifurtimox treatment during acute and chronic murine infection

##### 2.5.1. Therapeutic schemes

The nifurtimox treatment was carried out by oral route using gavage during the acute phase (starting at the first day of patent parasitemia) and chronic phase of infection (performed at 90 d.a.i.), using daily dose of 100 mg/kg of body weight for 20 consecutive days, according to Brener (1962).

The analysis of the infection evolution and therapeutic efficacy was performed considering several parameters. As cure control was adopted the negativation of three distinct categories of tests: parasitological methods (HC and polymerase chain reaction – PCR) and conventional serological test (IgG anti-T. cruzi enzyme-linked immunosorbent assay – ELISA), as suggested by Ministério da Saúde (2005), and non-conventional serology (flow cytometric anti-live trypomastigotes antibodies – IgG antibody by FC-ALTA), as proposed by Martins-Filho et al. (1995) modified, performed at days 90, 180 and 360 after treatment, as briefly described below:

##### 2.5.2. Hemoculture (HC)

The HC was performed as described by Filardi and Brener (1987).

##### 2.5.3. Polymerase chain reaction (PCR)

The PCR was performed as previously described by Gomes et al. (1998) modified, using the primer #121 (AAATAATGTCGGGTGAGATGCATGA) and primer #122 (GGTTGATTGGTGGTGGTGAATATA). The amplified DNA was visualized by electronic gel electrophoresis (QIAEXCEL Advanced Qiagen). Positive and negative control samples and reagents controls were always examined in parallel.

##### 2.5.4. Conventional serology (ELISA)

The ELISA to detect IgG anti-T. cruzi antibodies was performed according to Voller et al. (1975) modified by Santos da et al. (2012).

##### 2.5.5. Nonconventional serology (FC-ALTA)

The FC-ALTA to detect IgG anti-live trypomastigote antibodies was performed as originally described by Martins-Filho et al. (1995) modified for murine model by Molina et al. (2000).
Based on the cure criteria used, the T. cruzi samples were further classified according to the therapeutic response as resistant (cure rates ≤33%), partially susceptible (cure rates >33% <67%) and susceptible (cure rates ≥67%), according to Toledo et al. (2003).

2.6. Cardiac histopathological analysis

Cardiac histopathological analyses were performed during acute and chronic phases, before and 360 days after nifurtimox treatment. The cardiac inflammatory infiltrate and fibrosis were evaluated in hematoxylin-eosin and Trichrome of Masson respectively, in stained 5 µm tissue sections analyzed in optical microscopy (40X). The inflammation analysis was performed using a Leica QWin software (Leica Microsystems, Wetzlar, Germany). Data were expressed as the mean number ± standard deviation of nucleated cells in a total area of 0.8 × 106 µm², as previously described by Maltos et al. (2004). Collagen deposition was carried out by qualitative comparative analysis in tissues stained with Trichrome Masson.

2.7. Statistical analysis

Overall comparative analysis of epimastigote growth curve profiles from distinct T. cruzi genotypes was performed considering the area under the curve (AUC) via Kruskall-Wallis followed by Dunn’s post-test. Significant differences at p < 0.05 were identified by letters “a”, “b”, “c”, “d”, “e” and “f” as compared to TcI (Pr 150), TcVI (1337), TcII (501), TcII (452), TcII (728) and TcVI (748), respectively. Additionally, comparison amongst distinct T. cruzi genotypes at specific days were carried out by Kruskall-Wallis followed by Dunn’s post-test. AUC values are provided in the figures. Day of maximum parasite growth is identified by arrows.

The in vitro “Vero” cells infectivity obtained for distinct T. cruzi genotypes was evaluated by the two-way analysis of variance ANOVA for comparison of the number of infected cells, number of amastigotes and the ratio of amastigotes/infected cells. Significant differences at p < 0.05 amongst distinct T. cruzi genotypes were identified by letters “a”, “b”, “c”, “d”, “e” and “f” as compared to TcI (Pr 150), TcVI (748), TcII (501), TcII (452), TcII (728) and TcVI (1337), respectively. Moreover, differences amongst the incubation periods (24, 48 and 72 h) were identified by connecting lines.

Comparative analysis of parasitemia curves observed for distinct T. cruzi genotypes was performed considering the area under the curve via ANOVA, followed by Newman-Keuls post-test. AUC values are provided in the figures. Comparison amongst distinct T. cruzi genotypes at specific days were carried out by Kruskall-Wallis followed by Dunn’s post-test. Day of maximum parasite growth is identified by arrows. Analysis of biological parameters derived from the parasitemia curve (PPP, PP, MPP and DMPP) was performed by ANOVA followed by Newman-Keuls post-test. In all cases, significant differences at p ≤ 0.05 were identified by letters “a”, “b”, “c” and “d” as compared to TcII (501), TcII (452), TcII (728) and TcVI (1337), respectively. Observational analysis of relative percentage of bloodstream trypomastigote forms was also performed.

Comparative analysis of parasitemia curves obtained for distinct T. cruzi genotypes, before and after nifurtimox treatment at acute phase, was performed considering the area under the curve via ANOVA followed by Newman-Keuls post-test. AUC values are provided in the figures. Significant differences at p ≤ 0.05 were identified by connecting lines.

The comparison of cardiac inflammatory infiltrate observed for distinct T. cruzi genotypes was carried out by Kruskall-Wallis followed by Dunn’s post-test. Significant differences at p ≤ 0.05 were identified by letters “a”, “b”, “c” and “d” as compared to TcII (501), TcII (452), TcII (728), and TcVI (1337), respectively. Additionally, comparisons of cardiac inflammatory infiltrate before and after nifurtimox treatment during acute and chronic phase was carried out by Man-Whitney and significant differences at p ≤ 0.05 were identified by connecting lines.

All statistical analysis and graphical arts were performed using the GraphPad Prism software (version 5.03, San Diego, CA, USA).

3. Results

3.1. In vitro epimastigote growth curve of TcI, TcII and TcVI T. cruzi samples in acellular culture

The overall epimastigote growth curve profiles for TcI, TcII and TcVI T. cruzi samples in LIT medium are shown in Fig. 1. Data analysis demonstrated that TcII 728 sample presented higher AUC (p ≤ 0.05) as compared with TcI (Pr 150) sample. No differences were observed amongst the other T. cruzi samples. Additional analysis demonstrated that TcI (Pr 150) presented a late maximum peak of parasite growth around 16th day in culture. Both TcVI samples presented maximum peak of parasite growth around 8th day in culture. Interestingly, TcII samples displayed distinct patterns of parasite growth with early maximum peak observed for 501 sample around 6th day, intermediate maximum peak observed for 452 around 8th day and a maximum peak observed for 728 around 10th day in culture (Fig. 1).

3.2. Infectivity of TcI, TcII and TcVI T. cruzi samples in “Vero” cells culture

The infectivity for “Vero” cells evaluated at 24, 48 and 72 h after infection for each T. cruzi sample is presented in Fig. 2. The final infectivity was similar for all samples of the three DTUs although one sample of TcII (501) and one of TcVI (1337) have grown more rapidly. The results showed that the 1337 sample (TcVI) led to an overall higher number of infected cells at 24 and 48 h after infection as compared to other T. cruzi samples. The TcII (501) sample yield higher number of amastigotes per cell at 72 h after infection in relation to all samples. Conversely, the TcII (501) sample also presented the highest ratio of amastigotes per infected cell at 72 h after infection (Fig. 2).

3.3. Parasitemia curves in Swiss mice infected with TcI, TcII and TcVI T. cruzi samples

The overall pattern of parasitemia observed during experimental murine infection with TcI, TcII and TcVI T. cruzi samples is shown in Fig. 3. Data demonstrated that although the TcI (Pr150) and TcVI (748) samples have infected mice, they did not presented patent parasitemia sufficient for inoculation of animals necessary for the comparative analysis with the other samples. The TcVI (1337) sample induced low levels of parasitemia, evident by the low AUC (Fig. 3) and the lowest MPP values (Table 2). The TcII samples yield the highest parasitemia levels as observed by elevated AUC (Fig. 3) and highest MPP values (Table 2). Amongst the TcII sample, the sample 501 led to the most prominent parasitemia level, characterized by outstanding AUC and MPP values (Fig. 3 and Table 2).

The analysis of additional biological features derived from the parasitemia curve demonstrated that TcVI (1337) and TcII (728) samples presented similar and higher PPP as compared to TcII (501) and TcII (452) samples that showed similar PPP. The PP of TcII (501) and TcII (728) sample presented similar and higher PPP values than TcVI (1337) and TcII (452) that showed similar PP. Analysis of DMPP demonstrated that the TcII (728) showed the latest peak of parasitemia observed around 27th day after infection as compared to all other samples evaluated (Table 2).
In vitro epimastigote growth curve in LIT medium

**Fig. 1.** In vitro epimastigote growth curve of TcI, TcVI and TcII *Trypanosoma cruzi* samples in acellular culture. All experimental batches were carried out in triplicate and data reported as number of epimastigotes/mL. Overall comparative analysis of epimastigote growth curve profiles from distinct *T. cruzi* genotypes was performed considering the area under the curve (AUC) via Kruskal-Wallis followed by Dunn’s post-test. Significant differences at p ≤ 0.05 were identified by letters “a”, “b”, “c”, “d”, “e” and “f” as compared to TcI (Pr 150 ), TcVI (1337 = 1337 ), TcVI (748 = ), TcII (501 = ●), TcII (452 = ▲) and TcII (728 = ■), respectively. AUC values are provided in the figures. Day of maximum peak of parasite growth is identified by arrows.

3.4. Morphological analysis of bloodstream trypomastigotes in Swiss mice infected with TcII and TcVI *T. cruzi* samples

The analysis of TcII sample revealed that despite the difference in the length of patent period, all TcII samples displayed a similar kinetic of changes in morphological features of bloodstream trypomastigotes during the evaluation. Indeed, all TcII samples displayed an early predominance of intermediate trypomastigotes that were subsequently changed towards large trypomastigote forms.
Fig. 2. Infectivity of TcI ( ), TcVI ( ) and TcII ( ) Trypanosoma cruzi samples in Vero cells culture in vitro. The Vero cells infectivity was defined at 24, 48 and 72 h after infection for each T. cruzi sample in triplicated and the results expressed as mean percentage of infected cells per 100 cells; the number of amastigotes/cell as well as number of amastigotes per total number of infected cells. Significant differences at p ≤ 0.05 amongst distinct T. cruzi genotypes were identified by letters “a”, “b”, “c”, “d”, “e” and “f” as compared to TcI (Pr 150), TcVI (1337), TcVI (748), TcII (501), TcII (452) and TcII (728), respectively. Moreover, differences amongst the incubation periods (24, 48 and 78 h) were identified by connecting lines.

Table 2

Biological parameters from parasitemia curves of T. cruzi isolates in murine model. Data are expressed as mean ± standard deviation for all parameters (n = 8). PPP: pre-patent period; PP: patent period; MPP: maximum peak of parasitemia; DMPP: Day of maximum peak of parasitemia.

<table>
<thead>
<tr>
<th>T. cruzi Genotype (Isolate code)</th>
<th>Biological parameters</th>
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<tbody>
<tr>
<td>TcI [501]</td>
<td>PPP: 6.8 ± 0.8b,d PP: 21.3 ± 3.4a,c MPP: 200.0 ± 40.5a,c,d DMPP: 17.5 ± 1.4</td>
</tr>
<tr>
<td>TcII [452]</td>
<td>PPP: 6.9 ± 0.3a,b,c,d PP: 13.0 ± 1.9b,d MPP: 102.5 ± 44.9b,c,d DMPP: 13.3 ± 1.3</td>
</tr>
<tr>
<td>TcII [728]</td>
<td>PPP: 12.3 ± 0.5a,b,c,d PP: 28.9 ± 2.1a,b,c,d MPP: 100.0 ± 49.1b,c,d DMPP: 26.9 ± 1.3</td>
</tr>
<tr>
<td>TcVI [1337]</td>
<td>PPP: 10.3 ± 0.9a,b,c,d PP: 16.1 ± 2.4a,b,c,d MPP: 48.5 ± 16.5b,c,d DMPP: 23.5 ± 1.4</td>
</tr>
</tbody>
</table>

3.5. Response to nifurtimox treatment during acute and chronic murine infection with TcI and TcVI T. cruzi samples

Data regarding the parasitemia during the acute infection and acute/chronic cardiac inflammatory infiltrates observed during experimental infection with TcI and TcVI T. cruzi samples are provided in Fig. 5. The results demonstrated that regardless the
parasitemia levels, the nifurtimox treatment during acute infection was able to reduce the parasitemia to almost undetectable levels in animals infected with TcII and TcVI T. cruzi samples (Fig. 5).

3.6. Effectiveness of nifurtimox treatment during acute and chronic murine infection with TcII and TcVI T. cruzi samples

The nifurtimox therapeutic effectiveness was assessed using two parasitological methods (HC and PCR) along with conventional (ELISA) and non-conventional serological (FC-ALTA) performed at day 90, 180 and 360 after treatment (Table 3).

The therapeutic response was evaluated according to the cure criteria assessment adapted from that originally proposed by Ministério da Saúde (2005) and Krettli et al. (1982), using the PCR as an additional parasitological method and the ELISA and FC-ALTA methods as conventional and non-conventional serological approaches, respectively. Adopting the classic criterion of cure proposed by the Ministério da Saúde (2005), only two animals infected with T. cruzi sample 501 (TcII) and treated in the acute phase were considered cured. However, using the criteria proposed by Krettli et al. (1982) modified, it was demonstrated that nifurtimox treatment during acute infection led to higher cure rates in mice infected with TcII (501) and TcII (452) samples (75 and 88% of cure respectively). Lower cure rate was observed in animals infected with TcII (728) sample of this same genotype that displayed 25% of cure. Animals infected with TcVI (1337) were partially resistant (63% of cure) to nifurtimox treatment. An overall therapeutic failure was observed during chronic infection for both criteria of cure (Table 3).

3.7. Cardiac inflammatory infiltrate analysis in Swiss mice infected with TcI, TcII and TcVI T. cruzi samples

Analysis of cardiac inflammatory infiltrate during experimental murine infection with TcI and TcVI T. cruzi samples demonstrated that the TcVI (1337) sample led to significant lower cardiac inflammation at both acute and chronic infection. Interestingly, regardless the similar cardiac inflammation triggered by all TcII samples at acute infection, the TcII (501) induced lower cardiac inflammation at chronic infection as compared to both TcII (452) and TcII (728) samples (Fig. 3).

Moreover, the nifurtimox treatment performed at acute infection was able to significantly reduce the cardiac inflammatory infiltrate in all infected animals, regardless the T. cruzi genotype and sample, the results became similar to the control group INT
(Fig. 5). On the other hand, the nifurtimox treatment applied during chronic infection did not induce significant changes in the cardiac inflammation (Fig. 5).

3.8. Fibrosis analysis in Swiss mice infected with TcI, TcII and TcVI T. cruzi samples

Although apparently, has been verified discreet neoformation of collagen in the control groups (INT) than the treated ones, the quantitative analysis revealed similar collagen deposition in the heart of the animals; regardless the treatment had been done in the acute or chronic phases of infection (data not shown).

3.9. Mortality of Swiss mice infected with TcI, TcII and TcVI T. cruzi samples

No animal death was observed throughout the experimental acute and chronic phases of the infection up to day 360 after inoculation (data not shown).

4. Discussion

The T. cruzi species is divided into six distinct genetic lineages, better known as DTUs I, II, III, IV, V and VI (Zingales et al., 2009), that are distributed differently according to geographic region (Zingales et al., 2012). After established that the T. cruzi presents structure and clonal evolution (Tibayrenc et al., 1986), the correlation between the genetic and biological properties of the parasite (Buscaglia and Di Noia, 2003) is scientifically sustained and corroborated in several studies with strains and clonal stocks of the parasite in vitro (Laurent et al., 1997; Revollo et al., 1998), mice (Carneiro et al., 1991; Andrade and Magalhães, 1996; Toledo et al., 2002) and vectors (Lana et al., 1998). Therefore, the association of these features with the clinical forms of the disease in patients from whom the parasites were isolated is theoretically expected although some studies have failure in corroborate this hypothesis (Lages-Silva et al., 2006; D’Ávila et al., 2009).

Thus, the purpose of the present study was to characterize the biological features of T. cruzi samples (TcI, TcII and TcVI), isolated from chronic Chagas disease patients with distinct clinical forms, considering several parameters, including “in vitro” and “in vivo” biological properties along with the therapeutic response in acute and chronic murine infection, with the perspective of to detect some correlation of these parameters with the clinical characteristics of the disease in the patients from whom the parasites were obtained.

The growth of all six samples studied in LIT medium demonstrated that sample 728 of TcII genetic group presented the highest AUC when compared with TcI (Pr 150) and the other two samples of TcII DTU, which also showed fast or similar growth in LIT.
The response to nifurtimox treatment during acute and chronic murine infection with TcI and TcVI Trypanosoma cruzi samples. Parasitemia data during acute phase are reported as number of trypomastigotes per 0.01 mL of blood of animal infected with TcVI (\(1337^\circ\), □), TcI (501 = ○, ▲), TcII (452 = △, ▼) and TcIII (728 = ■, ▼). Comparative analysis of parasitemia curves observed for distinct T. cruzi genotypes before and after nifurtimox treatment was performed considering the area under the curve via ANOVA followed by Newman-Keuls post-test. AUC values are provided in the figures. Significant differences at \(p \leq 0.05\) were identified by connecting lines. The cardiac inflammatory infiltrate was evaluated in animals infected with TcVI (■, ▼), and TcII (▲, ▼) Trypanosoma cruzi samples. Data are reported as the mean number ± standard deviation of nucleated cells in a total area of \(0.8 \times 10^5\) μm². The comparison of cardiac inflammatory infiltrate observed for distinct T. cruzi genotypes before and after nifurtimox treatment was carried out by Kruskall-Wallis followed by Dunn’s post-test. Significant differences at \(p \leq 0.05\) were identified by letters “a”, “b”, “c” and “d” as compared to TcVI (1337), TcI (501), TcII (452) and TcIII (728), respectively. Additionally, significant differences at \(p \leq 0.05\) for analysis carried out before and after nifurtimox treatment are highlighted by connecting lines. Representative images of changes observed after nifurtimox treatment applied at acute infection and the lack of impact of treatment during chronic infection are provided in the figure.

In “Vero” cells it was verified that the final infectivity of these cells at 72 h after culture was similar for all samples of all DTUs. However, the replication as amastigotes were fast in one sample of TcII (501) and one sample of TcVI (1337) in the previous evaluation (24, 48 h). The TcII (501) sample yield higher number of amastigotes per cell and the highest ratio of amastigotes per infected cell at 72 h after infection. On the other hand, TcI sample displayed the lowest growth in “Vero” cells. These results corroborate the study of Andrade et al. (2010) that also observed higher capacity of TcII genotype strains in infecting “Vero” cells than TcII strains for cardiomyocytes.

Several authors have demonstrated that T. cruzi samples from human cases of Chagas disease of different endemic areas usually have low virulence in mice, sometimes with parasitemia in sub-microscopic levels detected only by hemoculture, as shown by several publications (Abolis et al., 2011; Oliveira-Silva et al., 2015; Teston et al., 2013). Similarly, Sales-Campos et al. (2015) recently demonstrated that strains belonging to genotype TcI have sub-patent parasitemia in mice. Rare exceptions exist such as VL-10 strain (TcII), also isolated from a human case of Virgem da Lapa, MG, (Filardi and Brener, 1987) very close to Berilo municipality, and considered highly virulent leading to intense inflammatory process associated to severe cardiac injury in dog models (Caldas et al., 2013). The TcI (Pr 150) sample here evaluated presented low subpatent parasitemia as previously demonstrated by Teston et al. (2013) and therefore was not further evaluated in mice model. Regarding TcII samples, Andrade et al. (2011) also demonstrated that TcII samples, displayed parasitemia profile, virulence, mortality rate, histopathological lesions and tissue tropism consistent with previous reports for this T. cruzi genotype. In the present study, TcII and TcVI samples showed significant differences in biological parameters derived from the parasitemia curve, including PPP, PP, MPP, DMPP. In the present study, the majority of the samples stud-
ied presented a relative low or subpatent parasitemia (PR-150 of TcI and 748 of TcVI) and when inoculated in mice the animals survived until the end of the experiments (one year of infection) without death consequent of the infection as observed in highly virulent strains. The sample 1337 (TcVI) showed significant differences in all parameters in relation to TcI samples, as observed by Andrade et al. (2011).

The absence of mortality in mice observed with all samples studied are consistent with other previous studies which have also revealed T. cruzi strains of low virulence for samples isolated from human infected of the Central and Southern regions of Brazil (Devera et al., 2002; Rimoldi et al., 2012; Oliveira-Silva et al., 2015). The polymorphism of the blood trypomastigotes of TcI samples is compatible with the expected for parasites of this genetic group (Type II or Zimodeme II) as demonstrated by Andrade (1974) and Andrade and Magalhães (1996) as well as for TcVI sample (hybrid or Zimodeme III).

Data regarding the therapeutic response to nifurtimox during acute infection demonstrated significant reduction in parasitemia of all T. cruzi samples evaluated revealed by the area under the curve of parasitemia (lower AUC and the maximum peak of parasitemia – PMP). The FBE showed limitation to detect parasites in treated and not cured animals due its low sensitivity (Brener 1962; Teston et al., 2013). On the other hand, when hemoculture was performed in animals ITAP with negative FBE the parasite was detected in different percentages of the animals (0–100%) according to the genetic group and sample. When it was analyzed the post-treatment evaluation in animals ITCP, 69% (22/32) of the treated mice were negative in hemoculture. For this reason the use of hemoculture alone is not recommended in post-treatment evaluations due its low sensitivity mainly in certain genetic groups of T. cruzi involved in the infection (Toledo et al., 2002; Toledo et al., 2003).

The therapeutic effectiveness at acute and chronic infections was also assessed by molecular parasitological method. The PCR detected 41% (13/32) of treatment failure in animals ITAP and in 25% (8/32) of the animals ITCP. It is important to highlight the progressive negativation of the parasitological tests used in the successive evaluations (especially at 90 and 360 d.a.t.), differently than the observed with the conventional serology (ELISA).

The positivity of the conventional serology (ELISA) during acute and chronic infection was higher than the observed in parasitological methods (94%). Considering the classic cure criterion (WHO, 2002) these results are indicative of global treatment resistance of the samples to nifurtimox in mice (6.25% of cure). Only 25% of mice infected with the strain TcI (501) treated during acute infection presented negative ELISA at 180 d.a.t. Such discrepancies between parasitological (HC, PCR) and serological methods have also been described in mouse model (Miyamoto et al., 2008; Oliveira-Silva et al., 2015). Therefore, other evaluations of these animals are necessary in order to ascertain the seronegativation, similar to that observed in humans considered cured (Alessio et al., 2014; Lana and Martins-Filho, 2015). However, a decrease of the absorbance was observed throughout the successive evaluations.

In order to better investigate the results of alternative methods applied to post-treatment evaluation we used FC-ALTA in parallel to previous tests (HC, PCR, ELISA) 360 d.a.t., never described in the literature in mice model. We prolonged the time of evaluation because in a recent review about post-treatment evaluation and cure criteria in Chagas disease (Lana and Martins-Filho, 2015) we addressed the need of serological tests able to present early seronegativation. One of the tests applied with this purpose is the FC-ALTA serology that detects the anti-live-tripomastigotes antibodies indicative of active infection in the host. Several publications using this methodology displayed shortly negative results after specific treatment, both in humans (Wendling et al., 2011), and in experimental animals (Martins et al., 2007; Oliveira-Silva et al., 2015). In this work, globally 64% of the animals treated during acute infection evaluated 360 d.a.t. were negative for FC-ALTA. Moreover, it was noteworthy that the parasitological tests (HC and PCR) of the animals were always negative, and in these mice no inflammation and collagen deposit in heart tissue were observed. It was verified higher cure rate by FC-ALTA, with negative results ranging from

### Table 3

<table>
<thead>
<tr>
<th>T. cruzi Genotype (Isolate Code)</th>
<th>Cure Assessment</th>
<th>Parasitological Methods (% of negative results)</th>
<th>Serological Methods (% of negative results)</th>
<th>Conventional</th>
<th>Non-conventional</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>HC (90dat)</td>
<td>PCR (90dat)</td>
<td>ELISA (90dat)</td>
<td>360dat</td>
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<tr>
<td>Acute Phase</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>TcI (501)</td>
<td></td>
<td>63 (5/8)</td>
<td>13 (1/8)</td>
<td>100 (8/8)</td>
<td>100 (8/8)</td>
</tr>
<tr>
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<td>100 (8/8)</td>
<td>0 (0/8)</td>
</tr>
<tr>
<td>TcII (728)</td>
<td></td>
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<td>100 (8/8)</td>
<td>63 (8/8)</td>
<td>0 (0/8)</td>
</tr>
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<td>75 (8/8)</td>
<td>0 (0/8)</td>
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<tr>
<td>Chronic Phase</td>
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<td></td>
<td></td>
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<tr>
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<td>100 (8/8)</td>
<td>100 (8/8)</td>
<td>0 (0/8)</td>
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<tr>
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<td>TcII (728)</td>
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* a Data are presented as percentage of negative results in parasitological e serological methods (number of negative animals/total number of animals); dat: day after treatment HC: hemoculture; PCR: polymerase reaction chain; ELISA: IgG anti-T. cruzi epimastigote enzyme-linked immunosorbent assay; FC-ALTA: Flow cytometric anti-live trypomastigote IgG antibody. Kreteli & Brener cure criteria as proposed by Kreteli & Brener (1982).
25% to 88% when compared with the ELISA (0% of cure). Besides such results of FC-ALTA, the absorbance of ELISA in the animals was decreasing in the successive evaluations, as happened with the PCR positivity, although we have evaluated drug response in only one sample of TcI, it was observed similarity of the rates of cure between TcI (501, 452 and 738) and TcVI (1337) samples.

The apparent lack of correlation between parasite genetic and the response to treatment of mice infected with TcI and TcIV samples here evaluated can be understood since TcVI parasites are resultant from hybridization phenomena between TcI and TcII (Westenberger et al., 2006) and the only sample of this genetic group here studied presented biological behavior more similar to TcI, which is also more genetically closer to TcII than TcI. Similar results were observed regarding treatment for clonal stocks of T. cruzi in hybrid group (Toledo et al., 2003) where parasites resistant, partially resistant and resistant to benznidazole were observed. Unfortunately the other sample of this DTU (TcVI) was not evaluated regarding treatment due its low parasitism insufficient for inoculation of a group of mice.

When FC-ALTA were used in the evaluation of animals ITCP the results were 0% negative as well as ELISA, indicative of therapeutic failure of NF in the chronic phase of infection taking at consideration the classic and novel cure criteria. Thus, in this phase of infection no correlation between response to treatment and parasite genetic was observed. Probably and again, the low number of samples evaluated was not sufficient to verify this correlation, differently from the observed by Toledo et al. (2003), which working with 20 clonal stocks of T. cruzi verified total resistance or partial resistance in parasites of the genetic group I (TcI), partial resistance in the hybrid groups and higher susceptibility of genetic group II (TcII) to benznidazole. However, it is important to note that again, the parasitological tests (HC and PCR) carried using the peripheral blood of the animals showed tendency to negativation (100%), in parallel to decrease of absorbance of the serological tests ELISA (data not shown).

This great discrepancy in the rates of cure using the novel cure criterion for animal ITAP and ITCP may be interpreted as similar as the classically considered for Chagas disease infections in humans (Viotti et al., 2011; Machado-de-Assis et al., 2012) and experimental models (Guedes et al., 2004; Oliveira-Silva et al., 2015). The facility for cure the acute phase of the infection and the difficulty for cure the chronic phase of the infection considering the negative conventional serology as cure criterion. Other possible interpretation of these results is the long duration of the antibodies of the conventional serology (ELISA) following treatment in relation to antibodies of the non-conventional serology (FC-ALTA) (Kretti, 2009), especially in hosts treated with chronic infections or mainly with later chronic infections when their affinity for the T. cruzi antigen is strongly higher.

The histopathological analysis in the acute phase of the infection revealed that cardiac inflammation in mice infected with TcI samples was higher and showed significant difference than the observed for TcVI. The positive impact of treatment was clearly observed in animals infected with TcI and TcVI samples in the reduction or elimination of parasitemia and inflammatory lesions, similarly to the verified by Oliveira-Silva et al. (2015) with others TcI samples of the same locality. No impact of nifurtimox treatment, neither reduction of cardiac lesions (inflammation and fibrosis), were observed during the chronic phase of infection.

In summary, in vitro essays (acellar and cellular cultures) TcI parasites showed always higher values for all parameters (growth and infectivity) than TcVI, followed by TcI. The same tendency was observed in mice infections regarding to parasitemia. An association between parasite genetic and heart lesions in mice model and human disease was verified with the TcI samples (452 and 728). Mice infected with these samples presented severe inflammatory lesions, as well as patients from whom these samples were isolated, which displayed severe clinical forms of the disease, cardiac and cardio-digie- nous, respectively. On the other hand, the TcI sample (501), isolated of a patient with the indeterminate clinical form of Chagas disease, despite had displayed higher parasitemia in mice, the inflammation in the heart was discreet, suggesting the important hole of this lesion on clinical Chagas disease evolution. Interestingly, the only TcVI sample which displayed low values for the same parameters evaluated caused in human the indeterminate form of the disease. The lack of correlation between mice infection evolution and human clinical evolution of the disease may be explained by the fact that we do not know if some patient has parasites of other DTU in the tissues distinct of the sample isolated, taking into account the clonal histotrophic model for T. cruzi (Macedo et al., 2004).

Although the clonal theory accepted for T. cruzi implies in correlation between genetic distance and the biological characteristics of the parasites populations of this specie, the occurrence of genetic exchange (hybridization phenomenon) may not be excluded. Consequently the tentative of to demonstrate these correlations is easier when the parasites are of the DTUs TcI and TcII. In fact, this was also verified in this manuscript and in others already published by our team (Lana et al., 1998; Toledo et al., 2002, 2003) and others (Andrade and Magalhães, 1997). However, when parasites of hybrid DTUs are considerate the correlations are not so clear and may be interpreted as consequence of the genetic background of these parasites which involve genetic components of two DTUs: TcI and TcII for TcV and TcVI or TcIII and TcIV in the case of TcII and TcIII.

Even considering the few number of T. cruzi samples, and the impossibility of to compare in vivo all samples, an apparent or discreet correlation was verified regarding parasite genetic and its general biological characteristics. However, apparently human disease evolution is not exclusively associated to parasite genetics, but also to its ability in develop histopathological lesions in the host, what in turn, is dependent to its genetic and immune response (Tibayrenc and Ayala, 2015). Probably the comparison of more samples of distinct DTUs could to show a stronger association among all aspects as verified by Andrade and Magalhães (1996) and Toledo et al. (2002, 2003) in mice. However this study is still valid and is the first that tried verifying correlation between four aspects in parallel (T. cruzi genetics, its biology in vitro and in vivo and the clinical aspects of the disease in patients from whom the parasites were isolated).

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

Maykon Tavares Oliveira: involved with all laboratorial activities related to the maintenance of parasites in acellar media, cellular media in a murine model; Ana Paula Vieira de Oliveira: undergraduate research student, with all laboratorial activities; Renata Tupinambá Branquinho: assistance in the performance of parasitaemia in mice and help in statistical analysis; Gláucia Diniz Aléssio: responsible for conducting the FC-ALTA; Carlos Geraldo Campos Mello: responsible for carrying out and interpretation of the ELISA data; Nivea Carolina Nogueira de Paiva: responsible for carrying out analysis and histopathology in cardiac tissue of animals; Cláudia Martins Carneiro: responsible for carrying out analysis and histopathology in cardiac tissue of animals; Max Jean de Orelia Toledo: collaborator responsible for the analysis of biological parameters in a murine model; William de Castro Borges:
co-leader of the project; Olindo de Assis Martins-Filho; coordinator flow cytometry platform and collaborator in the analysis of results FC-ALTA; Marta de Lana: coordinator of the study and collabora-

All authors have participated in this study and agree with the final version of the manuscript.

Acknowledgments

We would like to thank FAPESP, CNPq, Universidade Federal de Ouro Preto (UFOP) European Community’s Seventh Framework Program (No. 602773–Project KINDRED), CqPRF/FIOCRUZ for financial support, CMC, MJOT, ABR, WCB, OAMF and ML thank CNPq for fellowships (PQ/CNPq).

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