Impact of *Trypanosoma cruzi* clonal evolution on its biological properties in mice

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Received 7 June 2001; accepted 6 December 2001

Abstract

Twenty *Trypanosoma cruzi* stocks attributed to the 19, 20, 39, and 32 clonal genotypes were comparatively studied in BALB/c mice during the acute and chronic phases of the infection to test the working hypothesis that *T. cruzi* clonal structure has a major impact on its biological properties. Fourteen parameters were assayed: (1) infectivity; (2) prepatent period; (3) patent period; (4) maximum of parasitemia; (5) day of maximum of parasitemia; (6) parasitemia; (7) mortality, (8) percentage of positive hemoculture, (9) tissue parasitism; (10) inflammatory process during the acute phase of the infection; (11) mortality, (12) percentage of positive hemoculture; (13) tissue parasitism; and (14) inflammatory process during the chronic phase of the infection. Statistical comparison showed that the results are overall consistent with the working hypothesis that biological differences are proportional to the evolutionary divergence among the genotypes. Thus, closely related genotypes (19 vs 20 and 32 vs 39) show in general fewer differences than distantly related groups (19 or 20 vs 32 or 39) except for the comparison between 19 and 32. The working hypothesis is even more strongly supported by the result of the nonparametric Mantel test, which showed a highly significant correlation ($P = 2.3 \times 10^{-3}$) between biological differences and genetic distances among all pairs of stocks. These data taken together emphasize that it is crucial to take into account the phylogenetic diversity of *T. cruzi* natural clones in all applied studies dealing with diagnosis, drug and vaccine design, epidemiological surveys, and clinical diversity of Chagas’ disease.

Index Descriptors and Abbreviations: *Trypanosoma cruzi*; phylogenetic distance; biological properties; clonal theory; multilocus enzyme electrophoresis (MLEE); randomly amplified polymorphic DNA (RAPD); acute phase (AP); chronic phase (CP); days after inoculation (d.a.i.); liver infusion tryptose (LIT); gastrointestinal tract (GIT); genitourinary tract (GUT); percentage of infectivity (%INF); percentage of mortality during the acute phase (%MORT AP); percentage of mortality during the chronic phase (%MORT CP); prepatent period (PPP); patent period (PP); maximum of parasitemia (MP); day of maximum of parasitemia (DMP); parasitemia (PAR); percentage of positive hemoculture during the acute phase (% + HC AP); percentage of positive hemoculture during the chronic phase (% + HC CP); tissue parasitism (TP); inflammatory process (IP); tissue parasitism during the acute phase (TP AP); tissue parasitism during chronic phase (TP CP); inflammatory process during acute phase (IP AP); inflammatory process chronic phase (IP CP); Mann–Whitney test (MW); Kruskal–Wallis (KW); Kolmogorov–Smirnov test (KS). © 2002 Elsevier Science (USA). All rights reserved.

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1. Introduction

Trypanosoma cruzi, the aetiological agent of Chagas’ disease, exhibits a great biological diversity for many parameters (Andrade, 1976; Andrade et al., 1985; Dvorak, 1984; Melo and Brener, 1978). Moreover, the clinical manifestations of the disease are also diverse (Brener, 1987; Dias, 1992; Rassi and Luquetti, 1992). Miles et al. (1981b) have suggested that this variability could be in part explained by this parasite’s genetic diversity. Several authors have indeed shown an association between T. cruzi isoenzyme variability and its biological properties (Andrade and Magalhães, 1997; Andrade et al., 1992, 1983; Carneiro et al., 1991; Dvorak et al., 1980; Marques De Araújo and Chiari, 1988). However, these studies lacked either a sufficient sample size or a rigorous population genetic framework or both.

Tibayrenc et al. (1986) and Tibayrenc and Ayala (1988) have proposed that T. cruzi undergoes predominant clonal evolution with only rare events of genetic recombination. This clonal model predicts a parallel evolution between biological differences and genetic divergence among T. cruzi natural clones. Such a statistical association between genetic and biological differences has been found experimentally, in vitro by Laurent et al. (1997) and Revollo et al. (1998), and in the vector by Lana et al. (1998).

The present work explores the same working hypothesis in experimental infection in BALB/c mice.

2. Materials and methods

2.1. Parasites

The same standardized sample of 20 T. cruzi stocks used for previous studies (Lana et al., 1998; Laurent et al., 1997; Revollo et al., 1998) has been used. The stocks were cloned by dilution and micromanipulation, with visual verification under microscope. They were fully characterized by MLEE with 15 “loci” (Tibayrenc and Ayala, 1988), then with both MLEE with 22 different loci (Barnabé et al., 2000) and RAPD with 10 primers (Tibayrenc et al., 1993). They are representative of the “major clonets” (widespread clonal genotypes) 19, 20, 32, and 39 (Tibayrenc and Ayala, 1988). According to the nomenclature recently proposed (Satellite Meeting, Rio de Janeiro, Anon, 1999), major clonets 19 and 20 are included in T. cruzi I (Zymodeme 1 of Miles et al., 1981a; Lineage 1 of Tibayrenc, 1995; and lineage 2 of Souto et al., 1996) whereas both 32 and 39 groups are included in T. cruzi II (Zymodeme 2 of Miles et al., 1981a; Lineage 2 of Tibayrenc, 1995; and lineage 1 of Souto et al., 1996. Clonet 39 is equivalent to lineage 1/2 of Souto et al. (1996) considered as a hybrid clonal genotype (Brisse et al., 1998, 2000; Machado and Ayala, 2001). Information on the laboratory code, host, and geographic origin of these stocks is given in Table 1. The clonal genotypes 19, 20, 39, and 32 illustrate different phylogenetic relationships. Genotypes 32 and 39 are more closely related to each other, whereas genotype 19/
20 is more distantly related to both 32 and 39 (Fig. 1). Lastly, 19 and 20 are very close to each other.

2.2. Experimental conditions

Groups of 20 female BALB/c mice originating from the Instituto de Ciências Biológicas-UFMG, 28–30 days old, were inoculated through intraperitoneal route with 10,000 blood trypomastigotes/animal of each stock studied. Inocula were obtained from breast-fed Swiss mice in which the stocks were previously inoculated with a high number of metacyclic trypomastigotes obtained from late stationary culture in LIT medium after treatment with guinea pig serum according to Deane et al. (1984).

During the AP of the infection, mice were checked every day for PAR and mortality registration until 90 d.a.i. Deaths that occurred between 90 days and 12 months after inoculation were registered as mortality during the CP. Inocula and PAR were counted according to Brener (1962). Sixty and 140 d.a.i. hemoculture in LIT medium (Filardi and Brener, 1987) was made with

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Fig. 1. An unweighted pair-group method with arithmetic averages (UPGMA) dendrogram depicting the phylogenetic relationships among the 20 Trypanosoma cruzi stocks under study, assayed by 22 isoenzyme loci (Tibayrenc et al., 1993). Top cluster corresponds to clonal genotype 19/20 (T. cruzi I), medium cluster corresponds to clonal genotype 39 (T. cruzi II), and bottom cluster corresponds to clonal genotype 32 (T. cruzi II). The scale indicates genetic distances estimated with the index of Jaccard (1908).
approximately 400 μL of blood taken from each mouse (10 for AP and 10 for CP) through the retro-orbital plexus vein. Afterwards, four of these mice, two in the 30th d.a.i. (AP) and two in the 110th d.a.i. (CP), were necropsied for histopathological studies. The following organs and tissues were collected: (1) heart, (2) skeletal muscle, (3) GIT including stomach, small and large intestine, (4) GUT including bladder, kidney and uterine tubes, (5) brain, (6) liver, (7) pancreas, and (8) fat tissue. This material was routine processed, included in paraffin. Sections 5 μm thick of three blocks containing all organs of each mouse were obtained for further histopathological analysis. These preparations were stained with hematoxylin–eosin for microscope examinations.

In some cases (animals infected with the SO34c14, P209c11 and MNC12) the immunohistochemical peroxidase anti-peroxidase preparation (Barbosa, 1985) was processed to increase the detection of amastigotes.

### 2.3. Parameters evaluated

(a) %INF: percentage of mice with positive fresh blood examination and/or hemoculture.

(b) Mortality expressed in cumulative percentage of death registered during the AP (%MORT AP) and during the CP of infection (%MORT CP).

(c) PPP: the first day with positive fresh blood examination.

(d) PP: the period with positive fresh blood examination.

(e) MP expressed in the number of trypomastigotes/0.1 mL of blood, detected daily in fresh blood examination.

(f) DMP.

(g) PAR: area under the curve.

(h) Percentage of mice with positive hemoculture during the acute (%+HC AP-60th d.a.i.) and chronic phases (%+HC CP-140 d.a.i.) of the infection.

(i) TP: number of organs or tissues with parasitism classified as absent (−), mild (+), moderate (++), and severe (+++) during the acute (TP AP) and chronic (TP CP) phases of the infection.

(j) IP: number of organs or tissues with inflammatory process classified as absent (−), mild (+), moderate (++), and severe (+++) during the acute (IP AP) and chronic phases (IP CP) of the infection.

### 2.4. Statistical analysis

The statistical analysis of the parameters of virulence was done as follows: PPP, PP, MP, and DMP were compared with the nonparametric MW (Conares, 1980). Parameters %INF, %MORT, and %+HC were compared by chi-square test ($\chi^2$). The area under the curve of PAR was compared with the nonparametric KS. The histopathologic data (TP and IP) were analyzed with the nonparametric KW (Snedecor and Cochran, 1989).

A global analysis of the data was done with the nonparametric Mantel test (Mantel, 1967), which makes it possible to evaluate a possible correlation between, on one hand, genetic distances measured by either MLEE or RAPD, on the other hand, biological differences among the stocks. Contrary to the classical correlation test, this randomization procedure does not need any assumptions about the number of degrees of freedom. This procedure gives an equal weight to each biological parameter in the overall biological distance.

### 3. Results

#### 3.1. Parameters of virulence

A great standard deviation was observed in PPP, PP, MP, and DMP showing that clones of the same genetic group are not homogeneous in their characteristics (Table 2). The mortality parameter was similar for the different genetic groups during the AP and significantly different only between the genotypes 20 and 39 during the CP of the infection. The rates of mortality during the CP were significantly higher ($p < 0.001$) than those observed during the AP of the infection (Table 3).

#### 3.1.1. Infection with *T. cruzi* I (clonal genotypes 19 and 20)

Significant differences in four out of 14 parameters (Table 3) were observed in mice infected with the genetic groups 19 and 20, phylogenetically closely related, both

### Table 2

Mean values and standard deviations of biological parameters under study in mice infected with *Trypanosoma cruzi* of the genetic groups 19, 20, 39, and 32

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Abbreviation</th>
<th>Genetic groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>19</td>
<td>20</td>
</tr>
<tr>
<td>Prepatent period (days)</td>
<td>PPP</td>
<td>14.4 ± 6.5</td>
</tr>
<tr>
<td>Patent period (days)</td>
<td>PP</td>
<td>28.8 ± 21.8</td>
</tr>
<tr>
<td><em>Maximum of parasitemia</em> ($\times 10^8$)</td>
<td>MP</td>
<td>19 ± 25</td>
</tr>
<tr>
<td>Day of MP (days)</td>
<td>DMP</td>
<td>26.4 ± 13.5</td>
</tr>
</tbody>
</table>

*Number of trypomastigotes/0.1 mL of blood.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Statistic method</th>
<th>19 × 20 ($n_{19}, n_{20}$)</th>
<th>19 × 39 ($n_{19}, n_{39}$)</th>
<th>19 × 32 ($n_{19}, n_{32}$)</th>
<th>20 × 39 ($n_{20}, n_{39}$)</th>
<th>20 × 32 ($n_{20}, n_{32}$)</th>
<th>39 × 32 ($n_{39}, n_{32}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prepatent period</td>
<td>MW</td>
<td>156.3 × 107.6* (110, 146)</td>
<td>133.8 × 88.41* (110, 111)</td>
<td>81.7 × 76.2** (110, 49)</td>
<td>130.4 × 127.1*** (146, 111)</td>
<td>90.5 × 120.4* (146, 49)</td>
<td>72.2 × 99.3* (111, 49)</td>
</tr>
<tr>
<td>Patent period</td>
<td>MW</td>
<td>74.0 × 88.2*** (71, 92)</td>
<td>72.9 × 63.8*** (71, 65)</td>
<td>54.0 × 52.4*** (71, 35)</td>
<td>88.1 × 66.2* (92, 65)</td>
<td>70.6 × 46.6* (92, 35)</td>
<td>49.3 × 52.7*** (65, 35)</td>
</tr>
<tr>
<td>Maximum of parasitemia</td>
<td>MW</td>
<td>48.2 ± 100.0* (70, 81)</td>
<td>61.0 × 74.6* (70, 64)</td>
<td>48.8 × 61.3* (70, 35)</td>
<td>86.7 ± 55.7* (81, 64)</td>
<td>69.9 ± 32.2* (81, 35)</td>
<td>50.0 ± 50.0*** (64, 35)</td>
</tr>
<tr>
<td>Day of maximum of parasitemia</td>
<td>MW</td>
<td>71.9 ± 83.6*** (74, 81)</td>
<td>74.9 ± 63.3*** (74, 64)</td>
<td>54.7 ± 55.6*** (74, 35)</td>
<td>81.9 ± 61.7* (81, 64)</td>
<td>60.3 ± 54.3*** (81, 35)</td>
<td>46.9 ± 55.8*** (64, 35)</td>
</tr>
<tr>
<td>Tissue parasitism acute phase</td>
<td>KW</td>
<td>38.9 ± 42.1*** (40, 40)</td>
<td>42.5 ± 38.5* (40, 40)</td>
<td>41.5 ± 39.5*** (40, 40)</td>
<td>44.0 ± 37.0** (40, 40)</td>
<td>43.1 ± 39.9*** (40, 40)</td>
<td>39.5 ± 41.5*** (40, 40)</td>
</tr>
<tr>
<td>Tissue parasitism chronic phase</td>
<td>KW</td>
<td>40.5 ± 40.5*** (40, 40)</td>
<td>38.5 ± 42.5** (40, 40)</td>
<td>40.5 ± 40.5*** (40, 40)</td>
<td>38.5 ± 42.5** (40, 40)</td>
<td>40.5 ± 40.5*** (40, 40)</td>
<td>42.5 ± 38.5** (40, 40)</td>
</tr>
<tr>
<td>Inflammatory process acute phase</td>
<td>KW</td>
<td>37.6 ± 43.6*** (40, 40)</td>
<td>42.4 ± 38.6*** (40, 40)</td>
<td>39.2 ± 41.9*** (40, 40)</td>
<td>45.5 ± 35.5* (40, 40)</td>
<td>42.5 ± 38.6*** (40, 40)</td>
<td>36.8 ± 44.2*** (40, 40)</td>
</tr>
<tr>
<td>Inflammatory process chronic phase</td>
<td>KW</td>
<td>42.7 ± 38.4*** (40, 40)</td>
<td>46.8 ± 34.2* (40, 40)</td>
<td>42.1 ± 39.0*** (40, 40)</td>
<td>44.3 ± 36.7*** (40, 40)</td>
<td>39.5 ± 41.6*** (40, 40)</td>
<td>34.7 ± 46.3*** (40, 40)</td>
</tr>
<tr>
<td>Infectivity</td>
<td>$\chi^2$</td>
<td>91.4 ± 98.3* (301, 330)</td>
<td>91.3 ± 70.3* (301, 311)</td>
<td>91.3 ± 67.1* (301, 296)</td>
<td>98.3 ± 70.3* (330, 311)</td>
<td>98.3 ± 67.1* (330, 296)</td>
<td>703 ± 67.1*** (311, 296)</td>
</tr>
<tr>
<td>Mortality acute phase</td>
<td>$\chi^2$</td>
<td>1.9 ± 5.4*** (132, 153)</td>
<td>1.9 ± 2.9*** (132, 136)</td>
<td>1.9 ± 2.9*** (132, 136)</td>
<td>5.4 ± 2.9*** (153, 136)</td>
<td>5.4 ± 2.9*** (153, 136)</td>
<td>18.8 ± 18.3*** (136, 136)</td>
</tr>
<tr>
<td>Mortality chronic phase</td>
<td>$\chi^2$</td>
<td>22.4 ± 18.8*** (132, 153)</td>
<td>22.4 ± 13.9*** (132, 136)</td>
<td>22.4 ± 18.3*** (132, 136)</td>
<td>18.3 ± 31.9* (153, 136)</td>
<td>18.3 ± 18.3*** (153, 136)</td>
<td>31.9 ± 18.3*** (136, 136)</td>
</tr>
<tr>
<td>Hemoculture acute phase</td>
<td>$\chi^2$</td>
<td>89.3 ± 95.5*** (49, 44)</td>
<td>89.3 ± 57.6* (49, 48)</td>
<td>89.3 ± 82.8*** (49, 49)</td>
<td>95.5 ± 57.6* (44, 48)</td>
<td>95.5 ± 82.8*** (44, 49)</td>
<td>57.6 ± 82.8* (48, 49)</td>
</tr>
<tr>
<td>Hemoculture chronic phase</td>
<td>$\chi^2$</td>
<td>100.0 ± 98.0*** (49, 58)</td>
<td>100.0 ± 40.9* (49, 56)</td>
<td>100.0 ± 81.0* (49, 51)</td>
<td>98.0 ± 40.9* (58, 46)</td>
<td>98.0 ± 81.0* (58, 51)</td>
<td>40.9 ± 81.0* (46, 51)</td>
</tr>
<tr>
<td>Parasitemia</td>
<td>KS</td>
<td>11.07* (109, 96)</td>
<td>3.39* (109, 107)</td>
<td>4.53* (109, 97)</td>
<td>11.06* (96, 107)</td>
<td>15.37* (96, 97)</td>
<td>5.16* (107, 97)</td>
</tr>
</tbody>
</table>

Mean rank (MW = Mann–Whitney test, KW = Kruskal–Wallis test); Percentual ($\chi^2$ = chi-square test); Z of Kolmogorov–Smirnov (KS = Two-Sample Kolmogorov–Smirnov test).

* $P < 0.005$.

** $P < 0.05$.

*** Not significant.
The genetic group 19 with lower PAR and MP showed a higher value of PPP than genetic group 20 (Tables 3 and 4, Fig. 2). Clones of the genetic group 20 were significantly more infective and they led to a higher PAR (fresh blood examination) during the AP of the infection than clones from the genotype 19 (Table 3).

All five cloned stocks of this genotype induced patent PAR in mice while genotype 19 includes one subpatent stock (SP104 cl1).

### 3.1.2. Infection with T. cruzi II (clonal genotypes 39 and 32)

Mice infected with the stocks pertaining to the clonal genotypes 39 and 32 closely related both *T. cruzi* II and phylogenetically distant from the genotypes 19 and 20, displayed significant differences in 6 out of 14 of the same parameters studied (Table 3). Mice infected with the genetic group 39 with a higher PAR (Fig. 2) showed the lowest values of PPP and percentages of positive hemocultures (in both phases of infection) (Table 2). Mice infected with the five stocks of the genetic group 39 showed the most dispersed results among them in relation to many variables (individual data not shown). This group displayed the second highest MP and PAR (Fig. 2), mostly due to the high values of the stock Bug2149 c110 for these variables.

Only three stocks from the genotype 39 and two from the genotype 32 showed patent PAR in mice.

### 3.2. Histopathological data

The histopathological results revealed in general the presence of few amastigotes and inflammatory lesions. Parasitism was observed only in 5 organs/tissue examined: GIT and GUT were both more involved in this process, followed by skeletal muscles, heart and fat tissue. Inflammatory process was detected in all organs/tissue examined except in the brain, and in general, the organs and tissues more affected were also GUT/GIT, skeletal musculature and heart.

Out of the 20 stocks studied, only nine (45%) displayed a visible TP, generally classified as mild. Parasitism was observed for three stocks of genotype 20 (AP), two stocks of genotype 19 (AP), three stocks of genotype 39 (CP), and one stock of genotype 32 (AP). Parasitism was observed in 8.13% of the organs/tissue in the AP (30th d.a.i.), and in 2.5% of the organs/tissue during the CP (110th d.a.i.) The number of organs/tissue with parasitism in 40 analyzed by genotype during the AP was 4, 7, 0, and 2 for the genotypes 19, 20, 39, and 32, respectively (Table 4). Only mice infected with stocks from genotype 39 showed parasitism (four out of 40 organs/tissue analyzed) during the CP (Table 4). Immuno-histochemical analysis did not change these results.

Inflammatory process was present in mice infected with all 20 stocks. Most cases were classified as mild, whereas severe inflammatory reactions were less. The inflammatory process was observed in 51.30% of the organs/tissue in the AP (30th d.a.i.), and in 45.63% of the organs/tissue during the CP (110th d.a.i.). The numbers of organs/tissue with inflammatory process in 40 analyzed by genotype were: 18, 24, 16, and 24 during the AP and 22, 17, 11, and 23 during the CP for the genotypes 19, 20, 39, and 32, respectively (Table 4).

The genotypes 19, 20, and 32 displayed parasitism only during the AP (Table 4, Figs. 3A, C, and D). The genotype 39 was the only to show parasitism in the CP (three out of the five stocks) (Table 4, Figs. 4A–C).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Phase of infection</th>
<th>Number of organs/tissue</th>
<th>Intensity</th>
<th>Genetic groups</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>19</td>
</tr>
<tr>
<td>Tissue parasitism</td>
<td>Acute</td>
<td>40</td>
<td>–</td>
<td>36</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>+</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>++</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Chronic</td>
<td>40</td>
<td>–</td>
<td>40</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>+</td>
<td>0</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>++</td>
<td>0</td>
</tr>
<tr>
<td>Inflammatory process</td>
<td>Acute</td>
<td>40</td>
<td>–</td>
<td>22</td>
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<td>+</td>
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<td></td>
<td>Chronic</td>
<td>40</td>
<td>–</td>
<td>18</td>
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<tr>
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<td></td>
<td></td>
<td>+</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>++</td>
<td>10</td>
</tr>
</tbody>
</table>

– (Absent); + (mild); ++ (moderate); +++ (severe).
3.2.1. Infection with T. cruzi I (clonal genotypes 19 and 20)

Histopathological results during both phases of the infection were not significantly different between animals infected with genotypes 19 and 20 (Table 3). However, genotype 20 showed more severe parasitism during the AP than the genotype 19 (Table 4 and Figs. 3C and D). During the CP parasites were not observed in mice infected with these genotypes.

During the AP of the infection severe inflammatory reactions were observed only in mice infected with the genotype 20 (Table 4 and Fig. 3B). Moderate inflammatory reaction (Figs. 3A, C, and D) was twice more frequent in mice infected with genotypes 19 and 20 in relation to the other genotypes (Table 4).

3.2.2. Infection with T. cruzi II (clonal genotypes 39 and 32)

Histopathological results were significantly different only during the CP of the infection for both the parasitism and inflammatory process between animals infected with the genotypes 39 and 32 (Table 3). Stocks of T. cruzi II showed during the AP of the infection only mild parasitism and only in mice infected with the genotype 32 (Table 4). Differently from the other three genotypes studied, only mice infected with stocks from the genotype 39 did not show parasitism during the AP of the infection. In contrast, this genotype was the only one that displayed parasitism during the CP of the infection (Table 4) especially in skeletal musculature and TGI. This was observed with three out of five stocks studied (Bug2149 c110, MN c12, and SO3 c15).

Mice infected with genotype 32 showed a higher number of organs/tissue with mild inflammatory reaction (Fig. 4D) in relation to animals of the other genotypes in both phases of the infection (Table 4). The lowest number of organs/tissue with inflammatory reaction among all genotypes was observed in mice infected with genotype 39 during the acute and also during the CP of the infection when only mice infected with this genotype showed TP (Table 4 and Figs. 4A–C).

3.3. Statistical comparisons between T. cruzi I and T. cruzi II genotypes

Significant differences observed between the clonal genotypes 20 vs 39, 19 vs 39, 20 vs 32, and 19 vs 32 were 11, 9, 6, and 4, respectively (Table 3). In the comparison 20 vs 39, genotype 20 showed higher values of PP, MP, DMP, %INF, %+HC (both phases of the infection), PAR, TP AP, and IP AP than genotype 39, except TP CP (Table 3).

When genetic groups 19 vs 39 were compared, genotype 19 showed higher values of PPP, %INF, %+HC in both phases of the infection, TP AP and inflammatory process (CP) than genotype 39. On the other hand genotype 39 showed higher values of MP, PAR, and TP CP than genotype 19 (Table 3).

In the comparison of 20 vs 32, genotype 20 showed higher values of PP, MP, %INF, %+HC (CP), and PAR than genotype 32, except PPP (Table 3).

Finally in the comparison of 19 vs 32, genotype 19 showed higher values of %INF, %+HC (CP), and PAR than genotype 32 (Table 3). The MP was higher in group 32.

Overall, the decreased order of the four genetic groups considering all the parameters here studied for the 20 stocks is 20 > 19 > 32 > 39.

The result of the Mantel test evaluating the correlation between the matrices of 13 biological parameters (except PAR = area under the curve of PAR) and genetic distances for all pairs of stocks was significant ($P = 2.3 \times 10^{-3}$).

4. Discussion

Many authors have tried to find a link between the genetic diversity of T. cruzi and this parasite’s biological diversity. The biological parameters tested were various in experimental conditions in vitro (Dvorak et al., 1980), behavior in vectors and in mice (Garcia and Dvorak, 1982; Andrade et al., 1983; Carneiro et al., 1991), and clinical manifestations of the disease in chagasic patients from which the parasite was isolated (Brenière et al., 1985; Miles et al., 1981b). Andrade and Magalhães (1997) studying 138 T. cruzi stocks isolated from various hosts and vast geographic areas of Latin America classified the stocks in three groups called “biodemes” which were correlated with “zymodemes.” Although the above-cited studies have brought significant contributions, none of them relied on a clear population genetic framework. The present work, as well as previous ones published by other researchers of the same team (Lana et al., 1998; Laurent et al., 1997; Revollo et al., 1998),
aimed to fill up this gap. Based on the working hypothesis of *T. cruzi* clonal evolution (Tibayrenc et al., 1986), a standard set of 20 stocks representative of four “major clonets” (frequently isolated, ubiquitous clonal genotypes of major epidemiological relevance) as well as of the whole phylogenetic diversity of *T. cruzi*, was analyzed for 14 different biological parameters.

The analysis of the 14 biological properties studied (all parameters of virulence and pathogenicity in mice) in relation with the genetic diversity is in agreement with the working hypothesis (Tibayrenc and Brenière, 1988).

Our results are also consistent and corroborate previous observations in vitro and in BALB/c mice (Laurent et al., 1997), in acellular and cellular cultures (Revollo et al., 1998), and in *Triatoma infestans* (Lana et al., 1998) which confirmed the correlation between phylogenetic distance among *T. cruzi* clonal genotypes and their biological properties. Although our data of experimental infection in BALB/c mice show the same general pattern as the results of Laurent et al. (1997), they cannot be compared directly because our animals were not immunosuppressed.
As for these previous analyses, a great standard deviation in the majority of the parameters for a given clonet was observed again. Only the mortality parameter was similar for all genetic groups during AP although significant differences were observed between the genetic groups 20 and 39 during CP of the infection.

The great majority of clonal stocks studied were of low virulence. This fact associated with the low inocula used in the experiments could explain the relatively low TP observed in only 5 out of 8 organs/tissue and a generally discrete or moderate inflammatory reaction in 8 organs/tissue examined. Only the genotype 20, which is the more virulent, displayed severe inflammatory process. Such and other severe degenerative processes were never observed with the other clonal genotypes.

In previous studies (Lana et al., 1998; Laurent et al., 1997; Revollo et al., 1998), clonets 19 and 20 were plotted together, since they were very closely related and did not fall into distinct clusters when they were analyzed with 22 isozyme loci (Barnabé et al., 2000). However, this could be due to the fact that isozyme markers lack resolution for the lowest levels of phylo-

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Fig. 4. Microphotograph of BALB/c mice infected with *Trypanosoma cruzi* II. (A) Skeletal muscle showing parasitism and mild inflammatory process during the chronic phase (110th day of infection) with MN cl2 stock, genotype 39 (660× HE). (B) Fat tissue showing parasitism and mild inflammatory process during the chronic phase with Bug2149 cl10 stock, genotype 39 (660× HE). (C) Mild myocarditis and parasitism during the chronic phase of infection) with SO3 cl5 stock, genotype 39 (330× HE). Insert showing amastigote nest (660× HE). (D) Mild myocarditis and pericarditis with mononuclear infiltrate during the acute phase (30th day of infection) with MAS cl1 stock, genotype 32 (330× HE). Arrows indicate amastigote nest.
genetic divergence. Although their clustering discreteness is not clear, these two clonal genotypes are reliably distinguished by one genetic marker, the isozyme locus 6 phosphogluconate dehydrogenase, which is homozygous for 19 and heterozygous for 20 (Tibayrenc et al., 1986). It is impossible to rule out the hypothesis that their medical and epidemiological relevance is different.

In the present study, we have analyzed them separately. This made it possible to distinguish them for four important variables of virulence during the AP of the infection (PPP, MP, %INF, and PAR), which demonstrated clearly the higher virulence of the genetic group 20 in comparison with genetic group 19. Although not always significant, histopathological analysis also showed that mice infected with clones from this group showed more TP and inflammatory reaction in relation to all genotypes during the AP of infection. Genotypes 19 and 20, as well as "biodeme" III from Andrade and Magalhães (1997), displayed similar behavior and are classified as T. cruzi I (Satellite Meeting, Rio de Janeiro, Anon, 1999).

On the other hand, genetic groups 39 and 32, closely related and phylogenetically distant from the genetics groups 19 and 20, were different in six parameters, four of them related with virulence, specially PAR that was higher due to the Bug2149 c10 stock from the genotype 39. The lowest PPP and DMP of this genotype, not only in relation to genetic group 32 but also in relation to the other genotypes, indicate its high virulence. However the lowest PP and %+HC during both phases of the infection suggest its higher susceptibility to host’s immune response. These results corroborate those observed in mixed experimental infections in mice (Lana et al., 2000). In artificial mixtures involving either 19/20 + 39 or 32 + 39, the rates of successful reisolation were lower than for mixtures involving 19/20 + 32 genotypes.

We have noticed an unexpected result in animals infected with stocks of the genotype 39 which did not show any TP AP of the infection, whereas data from the literature frequently show that detecting TP AP of the infection is easier, but not the contrary.

The other differences between 39 vs 32 were observed during the CP when the genotype 39 was the only one with TP, curiously not associated with inflammatory process, a fact already observed by other authors who have studied virulence and pathogenicity of different T. cruzi stocks in animal experiments (Anselmi et al., 1965; Lana et al., 1992). Maybe this contrasting result observed with stocks of the genotype 39 can be explained by the fact that this genotype is a hybrid clonet, harboring a combination of the genes of the two putative parental genotypes (Brisse et al., 2000).

Some previous results of Diego et al. (1998) concerning histopathological data were confirmed in the present work. These authors studied the AP of infection in Swiss mice infected with the 15 stocks of clonets 19, 20, and 39 that we have also analyzed. Similar results between these authors and us were observed: (i) for these three genotypes, the highest number of parasites and inflammatory lesions was observed in the skeletal muscles; (ii) the highest number of parasite in the skeletal muscles was observed with stocks of the genotype 39 during the CP. Contrary to Diego et al. (1998), we did not observe (i) parasites in lymphoid organs (liver and spleen) from mice infected with stocks of genotype 39; (ii) parasites and inflammatory lesions in the brain of mice infected with any genotype. Maybe the different inocula and lineage of mice used could explain these differences.

Considering the comparison between pairs of phylogenetically distant genotypes the higher number of significant differences was observed between 20 vs 39 (11 differences), and 19 vs 39 (9 differences). These results are overall consistent with the working hypothesis that biological differences are proportional to the evolutionary divergence among the genotypes, since closely related genotypes (19 vs 20—T. cruzi I and 32 vs 39—T. cruzi II) show fewer differences than distantly related groups (19 vs 39 and 20 vs 39).

However only four and six significant differences were recorded between 19 vs 32, and 20 vs 32, respectively, also phylogenetically distant. Differences between genotypes 19 vs 32 were recorded in parameters linked to virulence: higher %INF, PAR, and %+HC during the CP of the infection for the genotype 19. In the comparison of genotypes 20 vs 32 differences were also recorded in parameters of virulence, which show clearly the higher virulence of the genetic 20. This genotype shows higher PP, MP, PAR, %INF, %+HC CP, and the lower PPP.

The working hypothesis of association between biological parameters and evolutionary divergence among natural clones is even more strongly supported by the result of the nonparametric Mantel test, which showed a highly significant correlation ($P = 2.3 \times 10^{-3}$) between biological differences and genetic distances among pairs of stocks. This shows that evolutionary divergence and biological differences do not evolve independently, and can be statistically predicted from each other to a large extent. This is actually another manifestation of the linkage disequilibrium generated by predominant clonal evolution in T. cruzi (Tibayrenc et al., 1986). A similar pattern has been observed in the bacterium Escherichia coli, which has a basically clonal population structure, although horizontal gene transfer is frequent in it (Miller and Hartl, 1986).

Finally, the present results corroborate those of other authors who have analyzed the same set of stocks for other parameters (Lana et al., 1998; Laurent et al., 1997; Revollo et al., 1998), several of which are of medical relevance (virulence in mice, transmissibility through the vector, infectivity of culture cells, in vitro drug sensitivity). Preliminary results showed that the in vivo sus-
ceptibility of the same set of stocks to Benznidazole and Itraconazole (Toledo et al., 2000) is also consistent with this hypothesis. Moreover, experiments dealing with artificial mixture of stocks pertaining to different clonal genotypes in vectors (Pinto et al., 2000) and in mice (Lana et al., 2000) suggested in some cases an interaction between different stocks. All these results taken together emphasize the fact that it is crucial to take into account the phylogenetic diversity of T. cruzi natural clones in all applied studies dealing with diagnosis, drug and vaccine design, epidemiological surveys, and clinical diversity of Chagas’ disease.

Acknowledgments

This work was supported by grants from FAPEMIG (Fundação de Amparo à Pesquisa de Minas Gerais), Brasil.

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