Comparison of Trypanosoma cruzi infection in dogs inoculated with blood or metacyclic trypomastigotes of Berenice-62 and Berenice-78 strains via intraperitoneal and conjunctival routes

Maria Terezinha Bahia1, Washington Luiz Tafuri1, Marcelo Vidigal Caliari2, Vanja Maria Veloso1, Cláudia Martins Carneiro3, George Luiz Lins Machado Coelho4 and Marta de Lana3

Abstract  This paper aimed to verify the influence of the inoculum source (blood or metacyclic trypomastigote) and the route of inoculation (intraperitoneal or conjunctival) on the course of T. cruzi infection in dogs, using comparatively the T. cruzi strains Berenice-62 and Berenice-78. All dogs inoculated intraperitoneally became infected independently of the T. cruzi strain and source of trypomastigotes used. High level of infectivity was also observed when metacyclic trypomastigotes of both strains were inoculated by conjunctival route. However, when blood trypomastigotes were inoculated by conjunctival route the percentages of infectivity were significantly lower in dogs inoculated with both strains. Parasitaemia was significantly higher in animals infected with metacyclic trypomastigotes via the conjunctival route independently of the T. cruzi strain used. All animals infected with Berenice-78 strain showed severe acute myocarditis. On the other hand, animals infected with Berenice-62 showed severe acute myocarditis only when infected with metacyclic trypomastigote, via the intraperitoneal route. The results suggest that the source of the inoculum and the route of inoculation remarkably influence the evolution of the infection for the T. cruzi in the vertebrate host even when the same strain of the parasite is used.


Resumo  Neste trabalho foi avaliada a influência da fonte do inóculo (tripomastigota sangüíneo ou metacíclico) e da via de inoculação (intraperitoneal ou conjuntival) na evolução da infecção de cães pelas cepas Berenice-62 e Berenice-78 do Trypanosoma cruzi. O índice de infectividade e os níveis de parasitaemia foram significativamente influenciados pelas condições de inoculação, tendo sido maiores nos animais inoculados pela via conjuntival, com tripomastigotas metacíclicas. Por outro lado, não foi observada relação entre as condições de inóculo e as lesões teciduais nos animais infectados com a cepa Berenice-78, pois todos apresentaram miocardite aguda severa. De forma inversa, nos animais infectados com a cepa Berenice-62, foi observada miocardite aguda intensa apenas nos cães inoculados com tripomastigotas metacíclicas, via intraperitoneal. Estes resultados sugerem que a taxa de infectividade e a evolução da infecção do hospedeiro vertebrado pelo T. cruzi podem ser marcadamente influenciadas pela fonte do inóculo e pela via de inoculação, mesmo quando a mesma cepa do parasita é utilizada.

Trypanosoma cruzi develops a complex biological cycle involving mammal hosts and triatomine bugs. During this cycle, distinct forms with varying morphology and functions can be observed. Amastigotes and epimastigotes are the division stages observed in mammal cells and in the intestinal tract of the triatomine bugs, respectively. Blood (BT) and metacyclic trypomastigotes (MT) are the infective stages encountered in mammals and triatomin excretions. The natural infection of mammals begins when the triatomines, after a blood meal deposit the MT in the host skin or mucous membrane. These trypomastigotes can penetrate through the mucous membrane, especially the ocular one or through the skin lesion caused by the insect bite.

There are many other mechanisms of T. cruzi transmission, including congenital, blood transfusion, laboratorial accidents and oral infection. Regardless of the mechanism of transmission, the host and parasite interaction begins after contact with BT or MT. It seems that only these stages are equipped with important molecules necessary for the interaction and/or host cell invasion. Apparently, MT has a greater ability than BT to penetrate the oral mucosa of mice.

Trypanosoma cruzi strains. T. cruzi strains, Berenice-62 and Berenice-78, both isolated from the patient Berenice, the first human case of Chagas’ disease characterized as T. cruzi II.

Experimental animals and infection. Eight groups of mongrel dogs, two months old were used. Animals were raised in the kennel of the Federal University of Ouro Preto, MG, Brazil. Dogs were fed with commercial chow and water was available ad libitum. Before the experiments, dogs were treated with anthelmintic and immunized against infectious diseases.

The animals were inoculated via intraperitoneal (IP) or conjunctival (CO) routes with 2000 blood (BT) or metacyclic (MT) trypomastigotes per kg bodyweight of the Berenice-62 or Berenice-78 strains made as described in Table 1. BT from mice (peak of parasitaemia) and MT from experimentally infected Triatoma infestans were used as inoculum.

To verify whether components of blood or triatomine feces influence the trypomastigote infectivity, three dogs were inoculated with BT, through CO route, rapidly washed by centrifugation and diluted in PBS, pH 7.2 and three other animals with BT washed in PBS, pH 7.2 and diluted in feces and urine of triatomines.

Parameters evaluated. Parasitaemia curves: parasitaemia was followed from the 10th day of infection up to negativation by fresh blood collected from the marginal ear vein. Parasites were counted according to the technique proposed by Brener, and the parasitaemia curves represent the mean value from all infected dogs at each time point.

Hemoculture: was performed only from sample of dogs without parasites in fresh blood examination. The technique was processed using 10ml of blood and examined monthly up to 120 days for the presence of T. cruzi parasites.

Serological profile: For serological tests the blood was collected 40 days after inoculation. Sera obtained were stored at -20°C and ELISA tests performed according to Voller et al, using antigen obtained from the Y strain of T. cruzi cultivated in LIT medium and peroxidase conjugated goat anti-dog IgG (Sigma Co. Ltd). The cut-off value was determined using the mean absorbance from ten uninfected animals plus two standard deviations.

Histopathological examinations. For histopathological evaluation animals were necropsied during the acute phase (30 to 40 days after inoculation). Fragments of the heart were fixed in 10% buffered formalin, pH 7.2, routinely processed and included in paraffin. Five mm thick sections were stained with Hematoxylin-Eosin. Intensity of parasitism and inflammatory lesions were comparatively analyzed in sections of the right atrium. This tissue was chosen due to previous studies demonstrating that the strains studied here possess special tropism to this region of the heart.

Statistical analysis. To compare the rates of infectivity the X² test was used. The parasitaemia was...
analyzed by a non-parametric test that compares the area under the curve of parasitaemia. The pre-patent
period and patent period were compared using analysis of variance.

RESULTS

Table 1 shows the serological (ELISA test) and parasitological results of dogs inoculated with BT and MT of *T. cruzi* via IP and CO routes. Dogs with positive parasitological and serological tests were considered infected. ELISA results were always in agreement with the parasitological examinations. The rates of infectivity were very similar in animals infected with both *T. cruzi* strains inoculated under the same conditions. All animals inoculated via IP route were infected (100%) independently of the *T. cruzi* strain, source of trypomastigote and route of inoculation used. Dogs inoculated with MT of both strains by CO showed 88.9% of infectivity, however the rates of infectivity were significantly (p<0.05) lower when BT of Berenice-62 (11.1% of infectivity) and Berenice-78 (0% of infectivity) strains were inoculated using the same route. No infection was detected in animals inoculated via CO route with BT washed with PBS, pH 7.2 or diluted in feces and urine of triatomines.

Figure 1 shows the mean of parasitaemia of dogs inoculated with the Berenice-62 strain. Animals inoculated with MT/CO showed significantly higher

Table 1 - Experimental conditions, serological, parasitological and infectivity data of dogs inoculated with blood or metacyclic trypomastigotes of *Trypanosoma cruzi* Berenice-62 and Berenice-78 strains, via intraperitoneal and conjunctival routes.

<table>
<thead>
<tr>
<th><em>T. cruzi</em> strains</th>
<th>Source of inoculum/(+)</th>
<th>Inoculation route</th>
<th>ELISA/(+)</th>
<th>Parasitological test/ dogs examined</th>
<th>Infectivity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Be-62</td>
<td>BT/IP</td>
<td>6/6</td>
<td>6/6</td>
<td>100.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>BT/CO</td>
<td>1/9</td>
<td>1/9</td>
<td>11.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>MT/IP</td>
<td>6/6</td>
<td>6/6</td>
<td>100.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>MT/CO</td>
<td>8/9</td>
<td>8/9</td>
<td>88.9</td>
<td></td>
</tr>
<tr>
<td>Be-78</td>
<td>BT/IP</td>
<td>6/6</td>
<td>6/6</td>
<td>100.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>BT/CO</td>
<td>0/16</td>
<td>0/16</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>MT/IP</td>
<td>6/6</td>
<td>6/6</td>
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<td>8/9</td>
<td>8/9</td>
<td>88.9</td>
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</tbody>
</table>

BT – blood trypomastigotes
MT – metacyclic trypomastigotes
IP – intraperitoneal route
CO – conjunctival route
$X^2 <0.05$

Figure 1 - Mean parasitaemia of dogs inoculated with blood (BT) or metacyclic trypomastigotes (MT) of *Trypanosoma cruzi* Berenice-62 strain via intraperitoneal (IP) or conjunctival (CO) route during the acute phase of the infection.
parasitaemia (p<0.05) than animals inoculated with BT despite the inoculation route.

Figure 2 shows the mean of parasitaemia of dogs inoculated with MT or BT from the Berenice-78 strain. Parasitaemia was significantly higher (p<0.05) in animals inoculated with MT independent of the route of inoculation, but especially when CO route was used. No infection was detected in animals inoculated with BT by CO route.

![Figure 2 - Mean parasitaemia of dogs inoculated with blood (BT) or metacyclic trypomastigotes (MT) of Trypanosoma cruzi Berenice-78 strain via intraperitoneal (IP) or conjunctival (CO) route during the acute phase of the infection.](image)

The parasitaemia of animals infected with MT of the Berenice-78 strain was significantly higher (p<0.05) than the observed in all dogs infected with the Berenice-62. No significant difference of parasitaemia among the strains was observed when dogs were infected with BT/IP. This comparison was not possible when animals were inoculated with BT/CO because only one dog inoculated with Berenice-62 was infected.

The results of pre-patent and patent periods are shown in Table 2. Difference of pre-patent period was observed only among dogs inoculated with the Berenice-62 strain (BT/CO). The patent periods were significantly longer (p<0.05) only in dogs inoculated with MT by CO route of both *T. cruzi* strains. Exactly in these situations, a tendency for more variation in the pre-patent period was observed.

<table>
<thead>
<tr>
<th>Source of inoculum/Route of inoculation</th>
<th>Berenice-62</th>
<th>Berenice-78</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPP (DAI)</td>
<td>13 (13-16)</td>
<td>18&lt;sup&gt;a&lt;/sup&gt; (18)</td>
</tr>
<tr>
<td>PP (DAI)</td>
<td>23 (21-23)</td>
<td>28&lt;sup&gt;a&lt;/sup&gt; (28)</td>
</tr>
</tbody>
</table>

DAI: days after inoculation
PPP – pre-patent period; PP patent period
BT – blood trypomastigotes
MT – metacyclic trypomastigotes
IP – intraperitoneal route
CO – conjunctival route
NI – not infected
<sup>a</sup> - p<0.05
<sup>b</sup> - one animal infected
Histopathological analysis of the right atrium of dogs infected with *T. cruzi* Berenice-62 strain showed mild lesions (Figure 3) when infected with BT, moderate lesions when infected with MT/CO and severe lesions only when infected with MT/IP. All dogs infected with *T. cruzi* Berenice-78 strain always showed severe lesions (Figure 4), independently of the inoculation route used. No infection was detected in animals inoculated with BT/CO.

**DISCUSSION**

The rates of infectivity in dogs inoculated with *T. cruzi* Be-62 and Be-78 strains were very similar when the conditions of inoculation were the same. However, the experiments clearly showed a great difference of infectivity between BT and MT by CO route in relation to IP route regardless of the *T. cruzi* strains used. The demonstration of the inferior capacity of BT in relation to MT to invade the mucosa has been previously described and it was demonstrated that MT were 200 times more efficient in penetrating the oral mucosa of mice than BT. On the other hand, McHardy and Neal, demonstrated that mice immunized with vaccines prepared from freeze-thawed or ultra-sonicated epimastigotes, blood trypomastigotes, or *plasma antigen* of *T. cruzi* strains and challenged by the injection of blood trypomastigotes killed more rapidly than the inoculated with bug-derived trypomastigotes.

Our results showed that these differences are probably not dependent on the components normally present in the blood or feces and triatomine urine. It was demonstrated that BT washed with PBS or mixed with feces and triatomine urine were not able to penetrate ocular mucosa of dogs.

These differences of infectivity are not surprising if the natural mechanisms of Chagas’ disease infection are considered. In natural conditions the infection with BT occurs by blood transfusion, congenital transmission, carnivorism or breast-feeding. Except in carnivorism and breast-feeding, a very unlikely mechanism of transmission, there is no interaction of the BT with the mucosa during the infection. Thus the inoculation of BT by CO route is artificial and was, in our experimental conditions, unable or inefficient to infect dogs (0 and 11.1%). However, when BT are inoculated directly in the peritoneum they infect 100% of the animals.

In contrast, the infection with MT in natural conditions occurs by deposition of the parasites by the triatomines on the skin and penetration via the skin bite or by CO route. Thus, the inoculation of dogs with MT by CO route represents the natural process of Chagas’ disease transmission in rural area by triatomines and our results reinforce the great success of MT to invade mucosa. Finally, the results also confirm the higher infectivity of MT (88.9% and 100%) in relation of BT, independently of the route of inoculation used.

Although the differences between BT and MT are relatively well known, especially in relation to membrane constituents, the real nature of their ability to infect ocular mucosa, reported in this work, as well as oral mucosa has not been determined. MT has biological and biochemical characteristics probably not shared with BT, which could be fundamental in mucosa invasion. The enlargement of the pre-patent period in dogs inoculated with MT of both strains by CO route, larger than that observed with IP inocula, suggests the development of different tissue-parasite interactions dependent on the compartment where the infection begins. The long pre-patent period observed in dogs inoculated with BT/CO may indicate the difficulty of this parasite stage in
penetrating the conjunctival mucosa. Differences in the infectivity between BT/CO and MT/CO could be due to the greater capacity of MT in resisting to the immune defense mechanisms present in the ocular mucosa. When both types of trypomastigotes were inoculated in dogs through IP route the rates of infectivity were the same (100%).

The results showed also the influence of the source of the inoculum and route of inoculation on the parasitaemia of the dogs. Animals inoculated with MT/CO always displayed higher levels of parasitaemia when infected with both *T. cruzi* strains. Although the histopathological data have not been quantified, it was demonstrated that all animals inoculated and infected with Be-78 strain always showed severe acute myocarditis. Similar lesion, with Berenice-78 strain, only occurred in dogs infected with MT/CO. This event, in addition to the histopathological data observed in both strains, evidenced more virulence and pathogenicity in MT than BT forms.

Previous studies, comparing the pathogenicity between MT/CO from Berenice-62 and Berenice-78, showed higher parasitaemia and histopathological lesions in dogs infected with Berenice-78\(^10\). This strain, when applied intraperitoneally, also provoked more severe acute myocarditis\(^3\) and more chronic lesions than the former\(^11\).

In conclusion, the results of this work indicate that the pathogenicity in *T. cruzi* infection in the vertebrate host, specifically in dogs, depends on the strain, source of the inoculum and route of inoculation. The work points to the importance to determine the molecules that take part in host-parasite interface in the site of *T. cruzi* entry into specific organs and cells, and to define what characteristics of each natural and experimental compartment are used to establish the infection.

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**REFERENCES**


