Highly diluted medication reduces parasitemia and improves experimental infection evolution by \textit{Trypanosoma cruzi}

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Highly diluted medication reduces parasitemia and improves experimental infection evolution by *Trypanosoma cruzi*

Denise Lessa Aleixo1*, Fabiana Nabarro Ferraz1, Érika Cristina Ferreira1, Marta de Lana2, Mônica Lúcia Gomes1, Benício Alves de Abreu Filho3 and Silvana Marques de Araújo1

**Abstract**

**Background:** There is no published information about the use of different protocols to administer a highly diluted medication. Evaluate the effect of different protocols for treatment with biotherapic *T. cruzi* 17 dH (BIOTc17dH) on clinical/parasitological evolution of mice infected with *T. cruzi*-Y strain.

**Methods:** A blind, randomized controlled trial was performed twice, using 60 28-day-old male Swiss mice infected with *T. cruzi*-Y strain, in five treatment groups: CI - treated with a 7% ethanol-water solution, diluted in water (10 μL/mL) ad libitum; BIOTp1 - treated with BIOTc17dH in water (10 μL/mL) ad libitum during a period that started on the day of infection; BIOT4p1 - treated with BIOTc17dH in water (10 μL/mL) ad libitum beginning on the 4th day of infection; BIOT4-5p6 - treated with BIOTc17dH by gavage (0.2 mL/animal/day) on the 4th, 5th and 6th days after infection; BIOT7-8p9 - treated with BIOTc17dH by gavage (0.2 mL/animal/day) on the 7th, 8th and 9th days after infection. We evaluated: parasitemia; total parasitemia (Ptotal); maximum peak of parasites; prepatent period (PPP) - time from infection to detection of the parasite in blood; patent period (PP) - period when the parasitemia can be detected in blood; clinical aspects; and mortality.

**Results:** Parasitological parameters in the BIOTp1 and mainly in the BIOT4p1 group showed better evolution of the infection compared to the control group (CI), with lower Ptotal, lower maximum peak of parasites, higher PPP, lower PP and longer survival times. These animals showed stable body temperature and higher weight gain and water consumption, with more animals having normal-appearing fur for longer periods. In contrast, groups BIOT4-5p6 and BIOT7-8p9 showed worse evolution of the infection compared to the control group, considering both parasitological and clinical parameters. The correlation analysis combined with the other data from this study indicated that the prepatent period is the best parameter to evaluate the effect of a medication in this model.

**Conclusions:** The BIOT4p1 group showed the best clinical and parasitological evolution, with lower parasitemia and a trend toward lower mortality and a longer survival period. The prepatent period was the best parameter to evaluate the effect of a medication in this model.

**Keywords:** Biotherapy, Mice infection, Parasitological evaluation, Clinical evaluation, Prepatent period

* Correspondence: deniseparasito@gmail.com

1Laboratory of Parasitology, Universidade Estadual de Maringá, Maringá, PR, Brazil

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Background

The use of highly diluted medications in the treatment of parasitic diseases has been investigated, but the available information does not allow us to demonstrate details of their effects. One of the hypotheses regarding the therapeutic effect of these drugs is that they modulate the immune system, and understanding the dynamics of this effect depends on the recognition of the information systems involved [1,2]. Bellavite et al. (2007) [3] suggested that homeopathic medicine can regulate inflammatory and immunopathological processes as well as the neuroendocrine network and peripheral receptors. Specifically in mice infected with *T. cruzi*, one study reported an increase in the apoptosis rate and a decrease in the secretion of TGF-β measured in serum collected from the group treated with biotherapeutic medication, compared with the control group [4].

Although several studies have attempted to explain the clinical results of highly diluted medications, few trials have evaluated the effect of different treatment schemes with these drugs on living organisms [5-7]. Biotherapeutics are highly diluted medications prepared according to homeopathic techniques [8] from biological products, including the etiological agent itself [9]. Although experience in clinical practice using biotherapeutics cannot be ignored, academic research on biotherapeutics has been questioned [10].

Chagas disease, caused by the protozoan *Trypanosoma cruzi*, has been studied for more than a century. However, no effective drug for etiological treatment has been discovered so far [11].

*T. cruzi-Y* is considered a reference strain [12]. Extensively studied in experimental murine models, it shows well-defined characteristics such as high parasitemia, peak of parasites by the 12th day of infection, death of all untreated infected animals, and partial resistance to drugs classically used in the treatment of human Chagas disease. Therefore, the murine model is an excellent experimental model for the evaluation of drug interventions [13].

According to the literature, morbidity is related to the presence of the parasite [14,15], and pathogenesis increase is the result of a disruption of the host-parasite relationship [16]. Therefore, using biotherapeutics as an intervention in murine infection with *T. cruzi* is a possible means to understand the effect of these highly diluted medications and to find new candidates to treat Chagas disease.

The present study evaluated *in vivo* the effect of different treatment schemes using the biotherapeutic *Trypanosoma cruzi* 17DH (BIOTPI7-D) in an experimental infection by this protozoan.

Results and discussion

As far as we are aware, this is the first study to compare different treatment schemes using a biotherapeutic *T. cruzi* in murine infection by the protozoan. Evaluation of parasitological and clinical parameters showed that the different treatment schemes using this medicine produced different profiles in the evolution of the infection of the Y strain of *T. cruzi* in Swiss mice.

Parasitological parameters

The mean and standard deviation of the parasitological parameters for each group are shown in Tables 1 and 2. The evolution of parasitemia in groups of animals treated with biotherapics diluted in water *ad libitum* for a long period (BIOTPI and BIOT4PI) showed better evolution of the infection compared to the control group (CI).

Compared to the control group, animals treated with BIOTPI and mainly BIOT4PI showed lower values of area under the curve of daily parasitemia (BIOTPI p < 0.01 and BIOT4PI p < 0.01), lower total parasitemia (BIOTPI p < 0.01 and BIOT4PI p < 0.01), and lower maximum peaks of parasites (BIOTPI p < 0.01 and BIOT4PI p < 0.01) (Table 1). There was no statistical difference in parasitemia on the 8th day of infection and in the 2nd peak of parasites for the BIOTPI and BIOT4PI groups compared to the control, as well as in the percentage of animals that showed the 2nd maximum peak of parasites. These are characteristics typical of infection by the *T. cruzi-Y* strain: a peak of parasites on the 8th day of infection and a second, lower peak of parasites, later. This second peak occurred earlier in BIOTPI and BIOT4PI than in the control group. Considering the parameters shown in Table 2, compared to the control group, the BIOTPI and mainly BIOT4PI groups showed longer prepatent periods (BIOTPI p < 0.01 and BIOT4PI p = 0.13) and shorter patent periods (BIOTPI p < 0.01 and BIOT4PI p < 0.01), longer survival times, significant for the BIOT4PI group (p < 0.01) and a lower (although not significantly lower) mortality. Together, these results show a better outcome of the infection, since the parasitemia in infection by *T. cruzi* is directly proportional to morbidity [17].

In contrast, the BIOT4.5-6 and BIOT7.8-9 groups showed a worse evolution of the infection compared to the control group. The area under the curve of daily parasitemia in these groups did not differ from the value observed for the control group (p = 0.99), and similarly for total parasitemia (p = 0.99). The BIOT4.5-6 and BIOT7.8-9 groups showed a higher maximum peak of parasites, significantly higher in the BIOT4.5-6 group, as well as a significant advancement of this maximum peak (p < 0.01). There was no significant difference in the parasitemia peak on the 8th day of infection, the parasitemia of the 2nd peak of parasites, nor in the percentage of animals that showed the 2nd peak for the BIOT4.5-6 and BIOT7.8-9 groups compared to the control group. The prepatent period and survival was lower for these
### Table 1 Parasitological parameters of animals subjected to different treatment schemes using biotherapic (BIOT TC17dH)

| GROUP         | AUC (x10^5) | P total (x10^6) | P max (x10^6) | DAY OF P max | P8thDI (x10^6) | 2nd P max (x10^6) | 2nd Peak n/N (% ) | DAY OF 2nd PEAK | 2nd PEAK n/N (% ) |
|---------------|-------------|----------------|---------------|--------------|----------------|------------------|-----------------|----------------|------------------|
| CI            | 10.9±8.20   | 12.5±8.98      | 5.02±7.07     | 10.76±3.15   | 2.82±3.12      | 11.0±10.2        | 14.03±2.63      | 8/12 (66%)      |                  |
| BIOT4-5-6     | 9.23±3.26   | 9.33±3.25      | 5.82±2.14**   | 8.25±0.67**  | 5.11±2.62      | 2.10±17.4        | 12.0±2.0*       | 4/12 (33%)      |                  |
| BIOT7-8-9     | 8.78±2.34   | 9.32±2.48      | 4.05±2.42     | 8.64±0.71**  | 3.96±2.51      | 17.4±0           | 14.25±10.0      | 8/12 (66%)      |                  |
| BIOTPI        | 5.95±4.55** | 6.63±5.02**    | 2.25±1.65**   | 15.46±23.06  | 1.62±1.69      | 11.1±13.6        | 13.67±0.63*     | 6/12 (50%)      |                  |
| BIOT4PI       | 4.28±2.89** | 4.63±3.19**    | 1.45±1.00**   | 10.27±2.03   | 1.01±0.96      | 7.09±10.2        | 12.8±0.68*      | 6/12 (50%)      |                  |

Statistical significance (p < 0.05* and p < 0.01**) compared to control group: CI: control group – infected animals treated with 7% ethanol-water solution diluted in water (10μL/mL) offered ad libitum; BIOTTC infected animals treated with biotherapic BIOTTC17dH diluted in water (10μL/mL) offered ad libitum from the day of the infection until the death of the animals; BIOTTC infected animals treated with biotherapic BIOTTC17dH by gavage on the 4th, 5th and 6th day of the infection; BIOTTC17dH infected animals treated with biotherapic BIOTTC17dH by gavage on the 7th, 8th and 9th day of the infection; AUC = area under the curve (parasites/mL); Ptotal = total parasitemia (parasites/mL); Pmax = maximum peak of parasites (parasites/mL); Day of Pmax = Day of maximum peak of parasites; P8thDI = number of tripomastigotes on 8th day of infection (parasites/mL); 2nd Pmax = number of tripomastigotes in the 2nd peak of parasites (parasites/mL); Day of 2nd Pmax = Day of 2nd peak of parasites; 2nd PEAK n/N = number of animals that showed 2nd peak of parasites (n)/total number of infected animals (N) x 100.
groups compared to the control. Mortality was 100% of the animals, as also observed for the control (Table 2).

Considering the 12 parameters shown in Tables 1 and 2, the BIOT4PI group showed the best overall performance.

In this study, the data obtained for the animals treated with highly diluted medications in the more appropriate treatment schemes (BIOTPI and BIOT4PI) showed better results than those observed in the control group. Notably, in the BIOTPI and BIOT4PI groups (which showed better evolution of the infection) there was a significant individual variation observed among the animals of the same group, as shown by the standard deviations (Tables 1 and 2). This individual variation was also observed in the control group.

These data suggest that some animals naturally evolve with lower parasitemia levels, while other animals evolve with higher parasitemia levels. The more-appropriate treatment schemes using highly diluted medications seemed to favor those animals with a tendency to develop lower parasitemia levels. This phenomenon may have led to the result in which the treatment of the BIOT4PI or BIOTPI groups produced a decrease in the parasitemia levels and a tendency to a lower mortality level with a longer survival period. This contrasts with the study by Sandri et al. (2010) [18], where a different treatment scheme was used in the same experimental conditions.

Clinical parameters
The groups of animals that were given the medication (BIOT<sub>Tc<sub>17Δdh</sub></sub>) in water <i>ad libitum</i> for a long period showed better clinical evolution, with a stable body temperature (<i>p </i>&lt; 0.05), higher weight gain (<i>p </i>&lt; 0.01) and higher water consumption (<i>p </i>&lt; 0.05) compared to the control group. The groups treated for a long period (BIOT<sub>PI</sub> and BIOT<sub>4PI</sub>) contained more animals with ‘normal-looking’ fur for longer periods during the study, and the BIOT<sub>4PI</sub> group showed the best results of all. In the BIOT<sub>4-5</sub> and BIOT<sub>7-8-9</sub> groups, more animals showed bristling hair, with a similar evolution to the control group (CI).

In murine <i>T. cruzi</i> infection, the morbidity is linked to parasitemia [17]. However, the host-parasite relationship is essential for the manifestation of the disease, which appears more lethal when the balance of this relationship is jeopardized.

The functional recovery of complex biological systems occurs in an integrated manner (considering the psycho-neuro-immuno-endocrine-metabolic axis), and sometimes the imbalance of one part is necessary to maintain the balance as a whole, with the appearance of new signs and/or exacerbating old symptoms or manifestations [19].

In this context, the host immune response causes many symptoms of the infection [20,21], and may help to explain the discrepancy between virulence and parasite loads [22,23], suggesting new prospects for the advancement of these studies.

Survival evaluation
Three animals survived in the groups that were given the medication (BIOT<sub>Tc<sub>17Δdh</sub></sub>) in water <i>ad libitum</i> for a long period. Of these three survivors, two were from the BIOTPI group, and only one of them had patent parasitemia. The other had negative PCR (Polymerase Chain Reaction) [24], understood as a protective action of the drug, since the treatment began at the time of the infection. Considering that the <i>T. cruzi-Y</i> strain shows 100% infectivity and mortality in Swiss mice [12,25-27] this result shows a patent benefit of the medication.

The surviving animal in the BIOT<sub>4PI</sub> group, although it showed a positive PCR diagnosis, never showed patent parasitemia. This result is consistent with low levels of parasitemia and provides information about the benefit of the treatment scheme used, since survival in the murine model is directly related to parasitemia levels [27].

The better evolution of the infection, considering both parasitological and clinical parameters in BIOT<sub>PI</sub> and mainly in BIOT<sub>4PI</sub> is expressed by the delayed onset of mortality (16<sup>th</sup> and 13<sup>th</sup> day of infection, respectively) as well as the lower mortality compared to the control group. The survival of 3 animals for more than 360 days, with a mean and standard deviation of 27.1 ± 40.98 (<i>p </i>&lt; 0.01) and 29.0 ± 42.79 (<i>p </i>&lt; 0.05) for the BIOT<sub>4PI</sub> and BIOT<sub>PI</sub> groups (Table 2), respectively, suggests an improved prognosis in a pathogenesis process that shows an irreversible evolution [28].
Correlation analysis

The correlation analysis indicates the relationship between two variables within a group. The relationships between each clinical or parasitological parameter (variable) and survival for the groups CI, BIOTPI, and BIOT4PI were evaluated separately. This relationship is expressed by a number called correlation coefficient (‘r’), which can be significant or not. Their values range from +1 to −1, and its size indicates the force of correlation. Correlations up to 0.299 were considered weak, values from 0.300 – 0.499 were considered intermediate, and strong from 0.500 – 0.999. The signal indicates the direction, if the correlation is positive (direct) or negative (inverse).

Considering that in this model, the evolution of the disease is irreversible, culminating in the death of the animals, an increase in the survival period is an indicator of a good treatment scheme. Some clinical and / or parasitological parameters may be more or less related to this increase in survival. The correlation analysis may indicate which are the more important parameters that could be measured. Therefore, we evaluated the correlations between the parasitological/clinical parameters and survival periods for the control group (CI) and the groups treated with the biotherapeutic (BIOTPI and BIOT4PI) in the treatment schemes that improved the clinical and parasitological evolution (BIOTPI and BIOT4PI) (Table 3).

Table 3 Values of correlation (‘r’)* of parasitological and clinical parameters with survival of animals subjected to different treatment schemes using biotherapeutic (BIOT17dH)

| PARAMETERS | CI | BIOTPI | BIOT4PI |
|------------|----|--------|---------|
| AUC        | -0.2250 | -0.1980 | -0.2349 |
| Ptotal     | -0.3220 | -0.4387 | -0.5227 |
| Pmax       | -0.2989 | -0.3182 | -0.5350 |
| Day Pmax   | NS  | NS     | NS      |
| PPP        | +0.4825 | +0.6704 | +0.9995 |
| PP         | +0.7901 | -0.7000 | -0.8998 |
| T*         | NS  | NS     | NS      |
| WEIGHT     | NS  | NS     | (+0.4797) |
| WATER      | NS  | NS     | NS      |

NS = no significant correlation. ‘r’ values are presented for significant correlations (p < 0.05) between survival and the parameters: AUC = area under the curve (parasites/mL); Pmax = total parasitemia (parasites/mL); Pmax = maximum peak of parasites (parasites/mL); PPP = Prepatent period; PP = patent period; T* = Temperature; Weight and Water. Groups: CI control group – infected animals treated with 7% ethanol-water solution diluted in water (104 μL/mL) offered ad libitum from the day of the infection until the death of the animals; BIOT4PI: infected animals treated with biotherapeutic BIOT17dH diluted in water (104 μL/mL) offered ad libitum from the 4th day of the infection until the death of the animals.

The prepatent period (PPP), patent period (PP) and total parasitemia (Ptotal) showed moderate to strong correlations with the survival period for all groups. For shorter survival periods, the groups showed higher total parasitemia (Ptotal). In groups treated for a long period, the Ptotal was lower and showed increasing ‘r’ values (BIOT4PI > BIOTPI > CI).

The prepatent period (PPP) was best correlated with the survival period. The BIOT4PI group showed a strong and positive correlation (r = 0.9995), indicating that the survival was closely linked to an increase in the prepatent period (PPP). In the different groups, this parameter clearly reflected the differences in performance under the different treatments, showing increasing ‘r’ values for the BIOT4PI, BIOTPI and CI groups. This result, combined with the set of data obtained in this study, indicated that the PPP was the best parameter to evaluate an intervention in this model.

This finding is important considering that in protocols with highly diluted substances, researchers tend to evaluate an excessive number of parameters in an attempt to include the key change that shows a benefit, since benefits may be perceived but cannot always be measured. In the experimental protocol used in this study, the risk of contamination and the workload involved need to be optimized, and it is imperative to select parameters that show the best effects most clearly.

The other parameters evaluated showed weak correlations with survival, although they also showed peculiarities in the comparison among the experimental groups. The maximum peak of parasites (Pmax) was inversely correlated with the survival period; the relationship was moderate for the CI and BIOTPI groups and strong for the BIOT4PI group. These results concord with those for the treatment schemes that began after the fourth day of infection, and again indicate the importance of the treatment scheme in determining the degree of morbidity from the infection.

The correlation between the peak day and the survival period had a particular result: the correlation was strong and direct in the CI group, showing that the later the peak of parasites, the longer is the survival time. In the groups treated with the biotherapeutic, the correlation was strong, but negative, with a higher ‘r’ value for the group that performed best in this study (BIOT4PI > BIOTPI).

This fact is consistent with what we have observed in previous studies: the biotherapeutic medications seem to anticipate several important biological phenomena in the course of experimental infection with T. cruzi [29]. The high standard deviation observed for this parameter in the BIOT4PI, BIOTPI and CI groups indicates the individuality of behavior for this parameter (Table 2). Sandri et al. (2010) [18] has also observed this behavior individually in both the treated and control group and...
found out that the animals that were more resistant to infection had a higher sensitivity to the treatment. In the case of Sandri et al. (2010) [18] the treatment scheme was inadequate and resulted in increased parasitemia.

These results show that some parameters can work as ‘markers’ of the performance of an intervention in the infection model used in this study, clearly reflecting the evolution of infection and these parameters can be prioritized in the experimental design, minimizing the work of researchers.

Although in this study in mice, the clinical parameters of temperature, weight and water consumption, as measured, were not effective to clarify the best therapeutic scheme using BIOT$_{7c17dH}$, the researcher perceive the difference among the groups. Find better methods to evaluate these parameters is an important perspective.

Considering the groups of animals treated for long periods, the BIOT$_{4p1}$ group presented a better performance with an intermediate correlation between weight and survival. These data, besides reflecting more accurately what was observed in the clinical evaluations, also supports a good prognosis for animals treated with this scheme.

Further discussions

The differences observed when comparing the evolution of infection in the groups treated for a long period in water (BIOT$_{p1}$ and BIOT$_{4p9}$), and for a short period of time by gavage (BIOT$_{4p5-6}$ and BIOT$_{7p7-8}$) show a better balance for animals treated for longer periods in water. The discussion of these results can also be based on the fact that the medication administered in water for a long period, besides enabling a higher number of stimuli, also causes less stress to the animal compared to the treatment by gavage, for a short period of time. Also in relation to medication administration, diluted in water, the difference in the intake among the animals, although it may be criticized. According to some authors, the concept of medication dose as the amount of medication that a patient must take to modify his state of illness is not applicable to highly diluted medications, because they do not act based on their mass, but rather by their dynamic effect (qualitative), which continues for a shorter or longer period according to the power of reaction or sensitivity of the organism and the stage of the disease [22,23].

Our data concord with the findings of other investigators that the use of highly diluted medications in the acute phase of infection requires a higher frequency of administered doses [23]. In a randomized controlled experiment, the benefit of the treatment using highly diluted medication continuously and for long periods of time, without interruption of the stimulus, promotes balance in acute infections [23].

Considering that the stimulation provided by the highly diluted medication is related to its ability to affect the psycho-neuro-immune-endocrine-metabolic axis, stress may also be a destabilizing factor that should be considered in evaluating the results [15].

$T. cruzi$ is an obligate intracellular parasite and its interaction with the host cells is complex, and inflammation is a decisive component of the pathogenesis. In developing resistance to $T. cruzi$, the immune system can act through different routes that function more or less effectively, depending on the conditions imposed by the host-parasite relationship [17]. Sandri et al. (2011) [30] demonstrated modulation of the inflammation process in eight-week-old animals infected with $T. cruzi$ and treated with a biotherapic, with a higher rate of apoptosis and reduced production of the cytokine transforming growth factor beta (TGF-β).

In the present study, assuming that the highly diluted medication BIOT$_{7c17dH}$ modulated the host immune system, the finding that the best therapeutic result was achieved with the medication administered beginning after 4 days of infection and for long periods (BIOT$_{4p9}$) is consistent with the following hypothesis: In this scheme, the host’s immune system, already activated by the parasite and affected by the highly diluted medication, leads the infection to evolve less aggressively, as described by Sandri et al. (2011) [30].

Conclusion

The BIOT$_{4d1}$ group showed the best clinical and parasitological evolution, with lower parasitemia and a trend toward lower mortality with longer periods of survival. The PPP (prepatent period = time in days from the infection until the first parasite was detected in the blood) proved to be the best parameter to evaluate the effect of a medication in this model.

Methods

Animals and infection

The study involved 60, 28-day-old male Swiss mice from the Central Animal Facility of the Universidade Estadual de Maringá. The animals were infected intraperitoneally with 1,400 blood trypomastigotes of $T. cruzi$-Y strain [12].

Groups according to treatment

The animals were divided equally into five groups according to the treatment: CI - infected animals treated with 7% ethanol-water solution diluted in water (10 μL/mL) offered ad libitum in an amber bottle; BIOT$_{p1}$ -infected animals treated with BIOT$_{7c17dH}$ diluted in water (10 μL/mL) offered ad libitum; BIOT$_{4d1}$ - infected animals treated with BIOT$_{7c17dH}$ diluted in water (10 μL/mL) offered ad libitum.
in an amber bottle from the 4th day of infection to the
death of the animals; BIOT_{4,5-6} - infected animals treated
with BIOT_{17dH} administered by gavage (0.2 mL / animal / day) on the 4th, 5th and 6th days after
infection; BIOT_{7,8-9} - infected animals treated with
BIOT_{17dH} administered by gavage (0.2 mL / animal / day) on the 7th, 8th and 9th days after infection.

The different treatment schemes were based on the ef-
fect of medication linked to the immunological effect
[3,18] and on the specific evolution of T. cruzi-Y strain
in Swiss mice [12,27]. Thus, BIOT_{PI} and BIOT_{APl} simu-
lated a situation where the medication is offered when the
immunological system is not in contact with the
parasite, and a situation where the medication is
offered when the immunological system is already in
contact with the parasite, respectively. In BIOT_{4,5-6} and
BIOT_{7,8-9}, the medication was offered to animals just
prior to the parasite peak and exactly coincident with
the parasite peak, respectively [12,27].

Experimental design
The experiment was performed twice as a blind random-
ized controlled trial. The animals were divided into
separate cages so that the mean initial weights of mice
in each group were statistically equal. The animals were
kept in a climate-controlled vivarium, under controlled
temperature (22.7 ± 1.2 °C), light/dark cycles of 12 hours,
and water and food offered ad libitum.

Biotherapic T. cruzi 17 dH (BIOT_{17dH})
The highly diluted medication was produced from the
buffy coat of blood collected from the orbital plexus of
three mice, containing blood trypomastigotes on the 7th
day of infection with T. cruzi Y-strain. The BIOT_{17dH}
was prepared by adding 0.9 mL of T. cruzi concentrate
(4.1 x 10^7 trypomastigotes/mL) in 9.1 mL of distilled
water in a 30-mL glass bottle. Subsequently, dilutions
were made in a 70% ethanol-water solution until the
16dH dynamization. The dynamization used, 17dH (cor-
responding to a dilution of 1:10^{17} of the initial suspen-
sion) was prepared with 7% ethanol-water solution. [8]
The microbiological test and the biological risk in vivo
were negative, according to the regulations of the Brazilian
Ministry of Health (RDC No. 67) [31], and the drug was
stored in amber glass for better conservation.

Parameters analyzed
Parasitological and clinical parameters were evaluated.
Parasitemia was evaluated by daily counts of blood tryp-
omastigotes from the day of infection, using the tech-
nique of Brener [1962] [32]. For animals that showed
negative parasitemia, blood was collected for PCR diag-
nosis (Polymerase Chain Reaction) [24].

Using the parasitemia curve, the following data were
obtained: prepatent period (PPP), which is the time from
infection to detection of the parasite in blood; patent
period (PP), the period when the parasitemia can be
detected; and total parasitemia (P_{total}), calculated as the
sum of the mean daily parasitemia levels for each mouse
in each group. The area under the curve (AUC), defined
as the number of parasites/mL of blood observed at a
certain time of infection, was calculated using the soft-
ware Prism Graph 5. The peak of maximum parasitemia
(P_{max}) considered the mean of the highest level of para-
sitemia observed in each animal throughout the ex-
periment, parasitemia on the 8th day of infection, the
2nd peak of parasites, and the percentage of occurrence
of this 2nd peak of parasites. The parasitemia was
expressed in parasites/mL and the occurrence of this
2nd peak was expressed by the percentage of animals
that showed a peak (n) within a group (N). The day
on which these peaks occurred was also considered.
Mortality was computed during the course of the ex-
periment, resulting in the following parameters: days of
survival and percentage of mortality expressed by
the number of animals died (n) within a group (N).
The animals were clinically evaluated on alternate days
from the day of infection until the end of the treatment
for body temperature, water consumption, fur
appearance, and body weight [33].

Statistical analysis
The results were compared statistically using the program
Statistica 7.0, and the Kruskal-Wallis test (ANOVA fol-
lowed by multiple comparisons of mean ranks for all
groups) was used since the analyzed parameters did not
all show a normal distribution. The correlations were ana-
yzed using the R.2.10 program. Correlations with r’ value
up to 0.299 were considered weak, from 0.300 – 0.499 as
intermediate, and from 0.500 – 0.999 as strong. The ac-
ceptable level of significance was 5% for all analyses.

Ethics
The experiment was approved by the Ethics Committee
for Animal Experimentation at the Universidade Estadual
de Maringá (UEM), under protocol number 030/2008.

Abbreviations
BIOT_{Tc17dH}: Biotherapic T. cruzi 17 dH; BIOT_{APl} Group treated with BIOT_{17dH}
diluted in water offered ad libitum during a period that started on the day of
infection and finished when the animals died; BIOT_{APl} Group treated with
BIOT_{17dH} diluted in water offered ad libitum from the 4th day of infection
to the death of animals; BIOT_{APl} Group treated with BIOT_{17dH}
diluted in water offered ad libitum from the 4th day of infection
to the death of animals; BIOT_{4,5-6} Group treated with BIOT_{17dH}
diluted in water offered ad libitum from the 4th day of infection
to the death of animals; BIOT_{7,8-9} Group treated with BIOT_{17dH}
diluted in water offered ad libitum from the 4th day of infection
to the death of animals; CI: Control of infection; AUC: Area under the curve; P_{total}: Total
parasitemia; P_{max}: Maximum peak of parasites; Day of P_{max}: Day of maximum
peak of parasites; P_{8thDI}: Parasitemia of 8th day of infection; 2nd P_{max}: Second
maximum peak of parasites; Day of 2nd P_{max}: Day of 2nd maximum peak of
parasites; PPP: Pre patent period; PP: Patent period.
