TRYPANOSOMA CRUZI: AVIRULENCE OF THE PF STRAIN TO CALLITHRIX MARMOSETS*

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Callithrix jacchus geoffroy marmosets (Humboldt 1812) were injected once subcutaneously with 10,000 parasites/g body weight and followed for a period of six months. The PF strain of *Trypanosoma cruzi* was used. Follow-up was done through blood cultures, xenodiagnosis, serological tests, and ECG. A small number of normal animals served as control.

INTRODUCTION

Blanchard¹, Laveran⁶, Brumpt², Mayer and Rocha Lima⁹, and recently, other investigators have reported that animals which survived a virulent infection with *Trypanosoma cruzi* became resistant to a subsequent challenge with this parasite.

Different procedures have been utilized over the years by several investigators searching for protection against further virulent *T. cruzi* infections: injection of low-virulence parasites (Norman and Kagan²²; Kagan and Norman⁵; Marr and Pike⁹; Seah and Marsden²⁶); injection of killed parasites (Soltys²⁸; Neal and Johnson²¹); of flagellate fractions (Segura, Cura, Paulone, Vasquez and Cerisola²⁷); of molecular fractions of cells from immunized hosts (Lemos and Menezes⁷); and of subcellular fractions of related parasites (Grymberg, Neri-Guimarães, Castro and Oliveira Lima⁸).

Injection of low-virulence parasites (Pizzi and Prager²⁴) or of a virulent strain, immediately followed by treatment (Pizzi²⁵) seems to be the most effective technique so far employed to obtain efficient protection. These observations have been confirmed by immunological techniques, which demonstrate that the infected animals develop antibodies and immunologically competent cells against the flagellates. A problem posed by this technique is the danger of infection with the low-virulence protozoa, which, although low virulent, could induce patent parasitemia in the injected animals. An additional problem is the mandatory treatment of all animals injected with a virulent strain.

Injection of parasitic fractions or other molecular derivatives of the infected animals opens a promising new field in trypanosomiasis research, but will not be discussed here. My discussion will be limited to the avirulence of a strain of *T. cruzi* i.e. a trypanosoma unable to induce disease or parasitemia, or to produce positive serological tests when injected in a suitable dose and by an adequate route. In 1968, I described such a parasite (Menezes¹⁰) as a probable mixture of mutants of the y strain, reported by Pereira da Silva and Nussenzweig²³ (as being highly virulent to mice. In a

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paper co-authored by Albuquerque (Menezes and Albuquerque13) this strain was named PF, and it has been my objective ever since to demonstrate its avirulence, emphasizing at the same time that, due to its live vaccine nature, it should not be used indiscriminately.

In 1970 (Menezes12), I described the first examples of animals injected with a very high dose of parasites which, although they presented positive blood cultures, gave negative results when inoculated into baby mice in culture form. The study presented here was carried out on Callithrix jacchus geoffroy marmosets, and concerns the tests performed before vaccination and throughout the six month period following injection of the avirulent strain.

MATERIALS AND METHODS

Animals

Sixteen Callithrix jacchus geoffroy marmosets (Humboldt 1812) — 9 males and 7 females — were vaccinated once subcutaneously (Fig. 1). Mean weight at the beginning of the experiment was 316.3 ± 79.5 g for the males, and 331.7 ± 37.7 g for the females. The controls were 8 similar marmosets, 5 males with a mean weight of 281.8 ± 72.2 g, and 3 females with a mean weight of 384.0 ± 72.2 g. All animals were trapped in their natural habitat (all at the same location) and were housed in cages containing one couple each whenever possible. All animals were kept under the same environmental and nutritional conditions. The diet consisted mainly of fruit (banana, papaya and oranges), with protein provided by non-fat milk and dehydrated soups (meat or chicken with vegetables) mixed with bread. All animals received a vitamin supplement once a week by oral route (10.000 IU vitamin A, 2000 IU vitamin D, and 100 mg vitamin C), and had free access to water.

Vaccine

The vaccine used was a suspension of the PF strain of T. cruzi harvested according to a personal modification of the Nakamura20 technique. Uncoagulated and unfiltered Warren medium (Warren30) was put into cellophane bags, which were placed into Erlenmeyer's flasks containing twice the amount of PBS solution, pH 7.2. The material was sterilized in an autoclave for 15 minutes at 115°C. The parasite cultures, 14 days old, were rinsed twice in saline solution and resuspended in sterile saline after centrifugation at 1500 rpm. The flagellate suspension contained 2 x 10^7 parasites/ml, with about 80% mobile forms, and 0.26% trypomastigote forms (over 5000 parasites were counted on five different slides stained with Giemsa).

Fig. 1 — Callithrix jacchus geoffroyi (Humboldt, 1812)

Each animal was injected subcutaneously with 10,000 parasites/g body weight shortly after preparation of the vaccine. An ECG each marmoset was taken once before, and several times after, vaccination. The test was performed in a FUNBEC apparatus, using the 12 standard leads. Prior to the test, each animal was injected subcutaneously with 0.1 ml Inoval (Johnson & Johnson Laboratories) (Fentanyl, 0.5 mg; Droperidol, 25 mg; sterile vehicle, qsp 10 ml).
**Blood Cultures**

Hemocultures were carried out in tubes (2 per animal) containing 5 ml original Warren medium (Warren\(^3\)). The cultures were examined 30 and 45 days later. This test was also performed before and after vaccination.

**Xenodiagnosis**

This test was carried out before and after vaccination of the animals. Six nymphs, 5th instar, of *Rhodnius neglectus* were used for each test. The triatominae were examined 30 and 45 days after the insects fed on marmoset blood. The latest examination was always carried out on the sediment of a pool of the intestinal contents of the remaining insects, after suspension in saline solution and centrifugation at 1500 rpm.

**Indirect Hemagglutination Test (IHT)**

This test was carried out using a filter paper blood smear, following the technique described by Souza and Camargo\(^2\) for the immunofluorescence test. The smears were kept in a deep freezer until the time of the test. The eluates were employed according to the method of Camargo et al.\(^3\) after inactivation at 56°C for 30 minutes, and centrifugation at 1500 rpm for 15 minutes. The sensitized red cells were kindly supplied by the laboratory of Dr. M.E. Camargo (Instituto de Medicina Tropical, USP, São Paulo).

**RESULTS**

**Body Weight**

Table I shows that both vaccinated and control animals lost weight during the 6 months duration of the experiment.

<table>
<thead>
<tr>
<th>Group</th>
<th>Total Number</th>
<th>Sex</th>
<th>Initial Weight</th>
<th>Final Weight</th>
<th>Weight Change Total (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>M</td>
<td>F</td>
<td>M</td>
<td>F</td>
</tr>
<tr>
<td>Vaccinated</td>
<td>16</td>
<td>9</td>
<td>7</td>
<td>316±79 g</td>
<td>332±38 g</td>
</tr>
<tr>
<td>Control</td>
<td>8</td>
<td>5</td>
<td>3</td>
<td>282±72 g</td>
<td>384±71 g</td>
</tr>
</tbody>
</table>

**ECG**

This test revealed abnormalities for only a few animals. The reader is referred to Table II where the results have been summarized. The cardiologist who collaborated in this investigation concluded that no essential abnormalities were found in the experimental animals after vaccination. No positive serological tests were obtained for any animal showing ECG abnormalities.

**Xenodiagnosis**

Table III summarizes the xenodiagnosis results obtained for experimental and control animals. On the average, two tests were performed before, and 5 after, vaccination of the experimental group. An average of 4 xenodiagnoses were performed for the control during the 6 months of observation. The results were always negative for both groups.

**Blood Cultures**

About 2 hemocultures were performed per each experimental animal before vaccination, and about 3 after vaccination. About 4 blood
cultures per animal were performed for the controls throughout the 6 months of observation. As in the preceding test, all results were negative (Table III).

### TABLE II

**ECG Results for the Experimental and Control Groups**

<table>
<thead>
<tr>
<th>Marmosets</th>
<th>Experimental</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Identification</td>
<td>Sex</td>
<td>Before Vaccination</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>LAVB 1/B</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>12</td>
<td>M</td>
<td>LAVBB 1st</td>
</tr>
<tr>
<td>13</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>14</td>
<td>F</td>
<td>N</td>
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<td>37</td>
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<td>N</td>
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<tr>
<td>22</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>25</td>
<td>F</td>
<td>N</td>
</tr>
<tr>
<td>30</td>
<td>F</td>
<td>N</td>
</tr>
<tr>
<td>31</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>32</td>
<td>F</td>
<td>RAVB 1/2B</td>
</tr>
<tr>
<td>33</td>
<td>F</td>
<td>N</td>
</tr>
<tr>
<td>34</td>
<td>F</td>
<td>N</td>
</tr>
<tr>
<td>35</td>
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<td>N</td>
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<td>44</td>
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<td>99</td>
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<td>N</td>
</tr>
<tr>
<td>17</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>19</td>
<td>F</td>
<td>Sinus rhytm</td>
</tr>
<tr>
<td>26</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>47</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>RVBB</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>RVBB</td>
</tr>
<tr>
<td>11</td>
<td>M</td>
<td>N</td>
</tr>
<tr>
<td>8</td>
<td>F</td>
<td></td>
</tr>
</tbody>
</table>

LAVB 1/2B = Left anterior ventricular branch hemi-block  
LAVBB 1st = Left anterior ventricular branch block — first degree. ( ) = Number of tests  
RAVB 1/2B = Right anterior ventricular branch hemi-block.  
N = normal

**Indirect Hemagglutination Test (IHT)**

Before vaccination the IHT was negative in 100% of the marmosets. After vaccination, 93.8% of the tests remained negative while only of 16 animals (6.2%) became serologically positive, but with all the parasitological tests negative (Table III), i.e. the two positive tests of the experimental group belonged to the same animal.
DISCUSSION

The main objective of this study was to demonstrate the avirulence of the PF strain of T. cruzi to marmosets, when administered in appropriate doses and by an advantageous route, as is the case with any other type of vaccine, live or not.

The only reliable means of detection of presumptive trypanosomiasis available to us at present are those utilized in this study. Until the time when an infallible diagnostic test for trypanosoma infections is developed we must be satisfied with what parasitologists the world over consider available evidence of trypanosomiasis, i.e. positive xenodiagnosis, blood culture, serological tests and, whenever possible, histopathological findings. Each of these tests has its limitations when performed separately, but the chances of diagnosis increase when they are employed together and repeated more than once.

Considering these arguments and on the basis of the present experiment, I can conclude once more that, when utilized adequately, the PF live vaccine of T. cruzi is avirulent.

I have already demonstrated that no infection could be detected in different animal species, including man, after the proper use of the PF strain of T. cruzi (Menezes ; Menezes and Ribeiro ; Menezes ).

RESUMO

Calitricideos da espécie C. jacchus geoffroyi (Humboldt, 1812) vacinados uma única vez, subcutaneamente, com 10.000 parasitas/g de peso corporal, foram seguidos durante um período de seis meses. Usou-se como vacina a cepa PF de T. cruzi.

O seguimento foi feito através de hemoculturas, xenodiagnósticos, testes serológicos e ECG. Um pequeno número de animais serviu como controle normal.

ACKNOWLEDGMENTS

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