Complete Immunization against *Trypanosoma cruzi*
Verified in Individual Mice by Complement-Mediated Lysis

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Experimental systems to assay immunity against *Trypanosoma cruzi* usually demonstrate partial resistance without excluding the establishment of sub-patent infections in protected animals. To test whether Swiss mice immunized with attenuated parasites might develop complete resistance against virulent *T. cruzi*, experiments were performed involving challenge with low numbers of parasites, enhancement of local inflammation and the combination of natural and acquired resistance. Absence of infection was established after repeated negative parasitological tests (including xenodiagnosis and hemoculture), and lack of lytic antibody was tested by complement mediated lysis. Immunization with $10^7$ attenuated epimastigotes conferred protection against the development of high levels of parasitemia after challenge with Tulahuen strain, but was unable to reduce the number of infected animals. However, when a strong, delayed-type hypersensitivity reaction was triggered at the site of infection by injecting a mixture of virulent and attenuated *T. cruzi*, a significant proportion of immunized animals remained totally free of virulent infection. The same result was obtained when the immunization experiment was performed in four month old Swiss mice, displaying a relatively high natural resistance and challenged with wild, vector-borne parasites. These experiments demonstrate that complete resistance against *T. cruzi* can be obtained in a significant proportion of animals, under conditions which replicate natural, vector delivered infection by the parasite.

Key words: *Trypanosoma cruzi* - lytic antibodies - immunological resistance

Chagas' disease, caused by *Trypanosoma cruzi*, presents three main clinical forms: acute, inapparent and chronic. The parasite can be detected in blood during the acute stage, but in the remaining stages, parasitemia is usually low and diagnosis is established by detecting specific antibodies using various techniques. Spontaneous cure with negative serology has been observed, but only on rare occasions (Brener & Plessman Camargo 1982, WHO 1991).

Immunologic protection against experimental *T. cruzi* infection has been achieved with several experimental vaccines, (for review see Kierszenbaum 1989) including, more recently, parasite synthetic peptide (Bua et al. 1991). As a result of vaccination, significant reductions have been obtained in parasitemia level (Andrews et al. 1985, McHardy & Elphick 1980), pathology (Kierszenbaum 1989) and transmission to vectors (Basombrío et al. 1993). Stronger effects of protection would be the achievement of “sterile immunity” after immunization with non viable vaccines or, at least, “complete immunity” against a second infection, which we define here as the absence of any positive parasitological test or serological evidence of infection after an infective challenge, in individual mice (not groups) pre-immunized with attenuated vaccines.

The success of experimental vaccination against *T. cruzi* becomes highly questionable when protection is verified with highly sensitive methods (Brener & Plessman Camargo 1982, Kierszenbaum 1989). Mc Hardy and Elphick (1980) demonstrated that infection with small doses, such as might be introduced after a triatomid bite (2 to 100 parasites) infect vaccinated and control mice equally well, a fact confirmed in our laboratory (Basombrío et al. 1989).

Among the humoral responses to *T. cruzi* infection, antibodies that can lyse bloodstream trypomastigotes in the presence of complement are generated (Krettli & Brener 1982). These lytic antibodies can be detected with the complement mediated lysis (CML) technique. Lytic antibodies are important immunologic effectors, associated with active infection and resistance against this parasite (Krettli & Brener 1982). The absence of lytic antibodies is considered to be indicative of cure, in spite of positive findings with other, so called, conventional serological tests (Krettli et al.}

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Lytic antibodies are not equally effective in killing all strains or clones of *T. cruzi* (Krettli & Brenner 1976, Murfin & Kuhn 1988), and CML should not be, therefore, useful to exclude infection by such parasites. For example in our laboratory, CML does not exclude attenuated infections by a low virulence strain (TCC). However, this technique is, in our hands, a good tool to exclude infection with virulent *T. cruzi* of the Tulahuen strain or local wild isolates.

In this work, we studied whether complete immunity against Tulahuen and wild *T. cruzi* parasites, can be induced by prior injection of the TCC strain. To obtain convincing evidence for the lack of residual infection by the virulent challenge, after exhausting the parasitological determinations, we used the CML technique. This study allowed us to identify three possible factors determining complete immunity in a discrete proportion of animals: a high level of natural host resistance, the vectorial origin of infecting parasites, and the development of a strong delayed type hypersensitivity reaction at the site of infection.

**MATERIALS AND METHODS**

**Animals** - All experiments were performed using outbred Swiss mice raised at our breeding facility. Both sexes were used, but were not mixed in any given experiment. Animals were 30 to 60 days old at the beginning of the experiments.

**Parasites** - Culture forms of *T. cruzi* (TCC) were used as sensitizing antigen and as triggering antigen in delayed type hypersensitivity reactions. The TCC subline of Tulahuen *T. cruzi* underwent attenuation in virulence and pathogenicity after more than 17 years of continuous passage in culture (Basombrio et al. 1982). Cultures were grown in Liver Infusion Tryptose (LIT) medium supplemented with 10% fetal bovine serum (GENsa) decomplemented at 68°C for 60 min, 20 µg hemin (Sigma), 100 IU of penicillin and 100 µg streptomycin per ml.

To challenge the mice we used either nonattenuated Tulahuen strain or wild isolates of *T. cruzi*. The former strain was maintained by mouse to mouse passages of infected blood. The wild isolates were obtained from *Triatoma infestans* bugs colonizing a guinea pig yard (Basombrio et al. 1987). The feces of infected bugs were diluted in LIT medium as to obtain the desired concentration of metacyclic trypomastigotes in the inoculum. Contaminating epimastigotes were not counted.

**Delayed type hypersensitivity reactions** - Mice were sensitized by injection of 30 µl of TCC suspension in saline (10^7 TCC epimastigotes), into the left hind footpad. One week later, a triggering inoculum of the same size was given in the right hind footpad. After 20 min and 4, 18, 24, 48, and 72 hr, swelling of the right hind footpad was measured with a caliper. In selected cases, animals were killed 24 hr post-challenge and the right hind footpads were fixed in 10% formalin. Decalcified and paraffin-embedded sections were stained with hematoxylin-eosin and studied histopathologically.

**Fresh blood mounts** - 10 µl of tail vein blood was collected into heparinized capillary tubes and placed between a microscope slide and coverslip, counting the parasites in 100 microscopic fields (480 x).

**Microhematocrit** - Three capillary tubes, each filled with 50 µl of blood were centrifuged at 9500 x g for 2 min and cut at the parasite-enriched leukocyte interphase, which was then examined microscopically.

**Xenodiagnosis** - Ten 2nd or 3rd instar *T. infestans*, fasted for a month were allowed to feed on each pentothal-anesthetized mouse for 15 min in the dark. Thirty and 60 days later, a pool of bug feces diluted in 10 µl of LIT medium was examined microscopically for the presence of parasites.

**Hemoculture** - Blood (0.2 ml) was collected with sterile heparinized Pasteur pipettes from the tail tip of mice, seeded into 75 mm of length and 9 mm of diameter glass tubes containing 1.8 ml of complete LIT medium and incubated at 28°C.

After 15, 30, 45 and 60 days, the presence of motile parasites was established using an inverted microscope.

**CML technique** - Tulahuen bloodstream trypomastigotes were obtained from highly parasitemic Balb/c mice which had been immunosuppressed with a dose of 350 mg/kg cyclophosphamide (LABINCA S.A., Buenos Aires) 24 hr before infection. After isolating the parasites by differential centrifugation at 150 x g for 20 min followed by an incubation at 37°C during 4 hr; they were washed three times at 3000 x g for 6 min, in minimum essential medium Eagle plus 10% inactivated fetal bovine serum (MEM-FBS) and incubated for 30 min in human normal fresh serum. Test sera decomplemented at 56°C for 30 min, were diluted 1:2 in MEM-FBS, mixed with a suspension of 5x10^6/ml washed trypomastigotes and incubated for 30 min at 37°C. The samples were then separated into two 50 µl aliquots (A and B) and kept in an ice bath until 50 µl of human normal fresh serum was added to aliquot A and 50 µl of human heat inactivated serum to aliquot B. After mixing and incubating at 37°C for 60 min, the samples were transferred to an ice bath until motile parasites were counted microscopically using a hemocytometer to calculate the
percent of lysis. More than 80% lysis was considered a positive result, indicating infection by Tulahuen or wild T. cruzi. Less than 20% lysis was taken as negative result. Only 1 of 132 animals fell between 20 and 80% lysis and was considered positive.

**Statistical analysis** - Delayed type hypersensitivity results were evaluated with Mann-Whitney “U” test and rates of infection with Fisher’s exact test.

**RESULTS**

**Correlation between lytic antibodies detection by CML and parasitological tests detecting low T. cruzi infection** - Several experiments were performed, where CML and parasitological tests (fresh blood mount, microhematocrit and xenodiagnosis or hemoculture) were applied to mice infected with virulent T. cruzi. Each of these methods was applied several times to each mouse, during the acute phase of infection. Most of these animals (132/143) had been injected with threshold infective doses 10 to 100 parasites. The load of infection obtained was classified in four grades, according to the sensitivity of the diagnostic method necessary for detection (Table I). The lack of CML closely correlated (p < 10^{-11}) with negative parasitological tests. CML was the most sensitive method to detect low grade infections.

**Failure to induce complete resistance against low doses of T. cruzi** - An immunization-challenge experiment was performed where 45 day-old mice were primed with 10^7 TCC in the left hind footpad and challenged with virulent Tulahuen trypomastigotes in the right hind footpad. This challenge dose was adjusted to a threshold of infectivity (10 Tulahuen trypomastigotes) that would infect about half of the control mice. The average levels of parasitemia in the immunized group were much lower than in the controls (Fig. 1). However, an exhaustive search for parasites in eight fresh blood mounts (day 15 to 49) and one xenodiagnosis (day 57), revealed that such resistance was not complete, since the low challenge doses had infected the same proportion of immunized and control mice.

**Identification of infection-free animals after challenge at the site of delayed type hypersensitivity reaction** - In order to induce an enhanced delayed type hypersensitivity reaction at the site of infection, experiments were performed in which a mixture of 10^7 TCC and 10 virulent Tulahuen blood trypomastigotes was used as the challenge inoculum using mice primed as described above. The sensitized mice developed a strong and typical delayed type hypersensitivity reaction, evidenced by significant footpad swelling, which reached a maximum 18-24 hr post inoculation and disappeared after 3-4 days (Table II). Histologically, the swollen footpads showed edema and infiltration of lymphocytes and macrophages. The presence of systemic T. cruzi infection was studied with progressively sensitive methods in these animals: fresh blood mounts were carried out between day 10 and 30 after challenge. Microhematocrit, xenodiagnosis and hemoculture were performed before day 60. CML was studied after day 90. In most control animals, fresh bloods mounts were enough to detect infection. Search for circulating parasites with microhematocrit and xenodiagnosis in the control groups slightly increased the detection of infected animals, which was close to 100% and was not further increased by hemoculture and CML determinations. In contrast, a significant proportion (59%) of immunized mice was completely resistant to infection in

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**TABLE I**

Comparison of different methods to detect residual infection by **Trypanosoma cruzi**

<table>
<thead>
<tr>
<th>Grade</th>
<th>Result of parasitological test</th>
<th>CML/total</th>
<th>% (+)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FBM</td>
<td>MH</td>
<td>X/H</td>
</tr>
<tr>
<td>I</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>II</td>
<td>-</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>III</td>
<td>-</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>IV</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

* a: grades I, II, III and IV, indicate different degrees of parasitemia. b: no. of mice with lytic antibodies/total mice studied. Abreviations: FBM = fresh blood mount; MH = microhematocrit; X = xenodiagnosis; H = hemoculture; CML = complement mediated lysis.
most experiments, as shown by negative, repeated parasitological results with all methods, including the absence of lytic antibodies 3-6 months after challenge (Fig. 2; Table II).

To further demonstrate the role of locally enhanced delayed type hypersensitivity reaction in the induction of total resistance, mice were challenged with virulent trypomastigotes, mixed or not with delayed type hypersensitivity inducing TCC epimastigotes (Fig. 3). This procedure showed that when a low number of infective parasites was injected alone, the use of progressively sensitive diagnostic methods revealed systemic infection in all control or preimmunized mice. In contrast, when the infective parasites were mixed with $10^7$ attenuated parasites and a strong delayed type hypersensitivity reaction occurred at the site of challenge, a significant proportion (40%) of preimmunized mice, as compared to nonimmunized controls, remained completely free of virulent infection, as demonstrated by all methods, including absence of lytic antibodies.

Selection of naturally resistant mice showing complete immunization against vector-borne \textit{T. cruzi} - Eighteen, 4 month old mice received three immunizing subcutaneous inocula of $2 \times 10^6$ TCC at weekly intervals. A week later they were challenged, together with 20 naive controls with 25 vector-borne, wild \textit{T. cruzi} metacyclic trypomastigotes in the right hind footpad. Two determina-

![Fig. 2: immunization-challenge experiments against \textit{Trypanosoma cruzi}. TCC pre-immunized ( , n=28) or control Swiss female mice ( , n=12) were challenged with 10 Tulahuen strain trypomastigotes admixed to $10^7$ TCC epimastigotes. All immunized mice developed strong delayed type hypersensitivity reactions at the site of infection (not shown). Infection was diagnosed with methods of progressive sensitivity: Fresh Blood Mount (FBM), Microhematocrit (MH), Xenodiagnosis (X) and Complement-Mediated Lysis (CML).](image)

### TABLE II

Selection of mice with complete immunity to a low \textit{Trypanosoma cruzi} challenge by enhancing DTH reactions at the site of infection$^a$

<table>
<thead>
<tr>
<th>Exp. No.</th>
<th>Sex</th>
<th>Group</th>
<th>DTH$^b$</th>
<th>Cumulated number of mice with positive diagnosis/total mice</th>
<th>Infected/Total (final result)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>FBM</td>
<td>MH</td>
</tr>
<tr>
<td>1 (pilot)</td>
<td>F</td>
<td>S</td>
<td>1.44±0.021****/d</td>
<td>2/5</td>
<td>2/5</td>
</tr>
<tr>
<td></td>
<td>C</td>
<td>0.44±0.05</td>
<td>5/5</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>S</td>
<td>1.24±0.07****</td>
<td>17/20</td>
<td>17/20</td>
</tr>
<tr>
<td></td>
<td>C</td>
<td>0.42±0.02</td>
<td>13/15</td>
<td>13/15</td>
<td>13/15</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>S</td>
<td>1.11±0.09****</td>
<td>2/28</td>
<td>2/28</td>
</tr>
<tr>
<td></td>
<td>C</td>
<td>0.27±0.04</td>
<td>7/11</td>
<td>9/11</td>
<td>11/11</td>
</tr>
<tr>
<td>4</td>
<td>M</td>
<td>S</td>
<td>1.20±0.06****</td>
<td>24/51</td>
<td>31/51</td>
</tr>
<tr>
<td></td>
<td>C</td>
<td>0.46±0.04</td>
<td>9/10</td>
<td>10/10</td>
<td>-</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>S</td>
<td>1.40±0.08****</td>
<td>7/19</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>C</td>
<td>0.18±0.02</td>
<td>20/20</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>6</td>
<td>F</td>
<td>S</td>
<td>1.46±0.05****</td>
<td>10/45</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>C</td>
<td>0.30±0.05</td>
<td>5/5</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

$a$: mice were immunized with $10^7$ TCC in the left hind footpad. One week later, immunized and control animals were challenged with 10 blood trypomastigotes of Tulahuen strain (except in experiment No. 5, which was challenged with 150 vector-borne metacyclic trypomastigotes), mixed in 30 µl saline with $10^7$ TCC into the right hind footpad. $b$: DTH = delayed-type hypersensitivity. Footpad swelling (mm) 24 hr after inoculation, mean ± standard error. $c$: mice recorded as positive in a given column include those recorded as positive in previous columns. Abreviations: Exp.= experiment; S = sensitized; C = control; FBM = fresh blood mount; MH = microhematocrit; X = xenodiagnosis; H = hemoculture; CML = complement mediated lysis. $d$: *p<0.09; **p<0.05; ***p<0.01; ****p<0.001.
Fig. 3: immunization-challenge experiments against *Trypanosoma cruzi*. Infection of Swiss female mice by 150 insect-borne, wild *T. cruzi* metacyclic trypomastigotes in control ( , n=5) or TCC-preimmunized ( , n=20) Swiss male mice. In parallel, simultaneous group of mice, the challenge inoculum was admixed with 10^7 epimastigotes and a strong delayed type hypersensitivity reaction (not shown) developed at the site of infection. This modality of challenge infected all control mice ( , n=20) but was totally resisted (p=0.002) by 40% of immunized mice ( , n=20). Systemic infection was studied sequentially by Fresh Blood Mounts (FBM), Xenodiagnosis (X), Hemoculture (H) and Complement Mediated Lysis (CML).

TABLE III

<table>
<thead>
<tr>
<th>Group</th>
<th>Cumulated number of mice with positive diagnosis/total mice</th>
<th>Infected/Total (final result)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fresh blood mount</td>
<td>Microhematocr</td>
</tr>
<tr>
<td>Immunized</td>
<td>3/18</td>
<td>7/18</td>
</tr>
<tr>
<td>Control</td>
<td>17/20</td>
<td>20/20</td>
</tr>
</tbody>
</table>

a: statistic significance for rates of infection was evaluated with Fisher's exact test (p=0.00089)

DISCUSSION

Acquired immunological resistance against protozoan parasites is seldom complete, either in natural or in experimental infections (Andersen 1994). Previous failure to demonstrate sterile or complete immunity to *T. cruzi* challenge (McHardy & Elphick 1980, reviewed in Kierszenbaum 1989, Basombrio et al. 1989) were connected with the capacity of this parasite to establish very low levels of infection and with the low sensitivity of the detection methods. We have attempted to surmount this problem by using the CML technique. The study of lytic antibodies seems to be the most convincing approach to address the demonstration of absence of infection by *T. cruzi* (Krettli & Brener 1982), when the infecting strains or isolates are good inducers of this type of antibody. Such is the case with the Tulahuen and wild isolates used in this work. On the other hand, the TCC organisms we used for immunization do not appear to be good inducers of lytic antibodies, even in parasitologically proven infections (unpublished observations). Therefore, our procedure does not demonstrate sterile immunity in the immunized animals, but it does establish a complete, acquired resistance, at the skin level, against the virulent parasites used for challenge, in a significant number of animals.

When TCC-immunized and control mice were challenged with threshold doses of infecting parasites, the levels of parasitemia were sharply lower in the immunized group, although the proportion of infected mice was comparable in both groups. This indicates that the effects of immunization “are expressed at a stage of infection later than invasion”, confirming the original notion of Mc Hardy and Elphick (1980). This led us to further concentrate efforts in the demonstration of complete resistance using two approaches: enhancing the delayed type hypersensitivity reaction at the subcutaneous level and testing the effects of immunization in animals in which natural resistance to *T. cruzi* was already strong. In the first approach, the 10^7:10 mixture of attenuated and virulent parasites induced a typical delayed type hypersensitivity reaction. This reaction was never seen, at least macroscopically, in mice challenged with virulent parasites alone. The failure of several parasitologic methods to detect systemic
infection plus the absence of lytic antibodies in a significant proportion of immunized-challenged animals suggested that the infecting parasites were in such cases totally eliminated at the site of infection. This modality of challenge does not reflect the normal conditions of vectorial infection by Trypanosoma cruzi and would thus not constitute a model of vaccination applicable to conditions of natural infection. In the second approach, which seems more relevant to natural T. cruzi infection, we tested the combined effects of natural and acquired resistance against vector-borne parasites. Four-month-old Swiss mice are at a time of their life when natural resistance to T. cruzi is highest (Basombrio & Arredes 1987). Using these animals, we could demonstrate that low doses of vector borne, wild T. cruzi injected at the footpad, and which were 100% infective for control animals of the same age and sex, were completely eliminated at the site of infection in a significant proportion of immunized mice.

A survey of 66 published studies on experimental immunization against T. cruzi, involving various animals and immunogens concluded that complete immunological protection against this parasite had never been demonstrated (Kierszenbaum 1989). In most studies, immunized animals were challenged with large infective inocula and the ensuing parasitemias were measured with methods which not always detected very low levels of parasites. Such type of infections contrast with natural parasitism by T. cruzi, which results from low numbers of vector-borne parasites penetrating epithelia and which require more sensitive methods for detection. Only 14 of the 66 studies included methods of higher sensitivity than fresh blood mount (e.g., xenodiagnosis, hemoculture and sub-inoculation) and none of them used serological methods to exclude infection, as practiced by clinicians on patients (Galvão et al. 1993). Two of these studies deserve a special comment in this context, because both used the CML technique and, in one of them, parasitological results suggest total protection. Andrews et al. (1985) immunized A/Snell mice with three doses of 8-methoxypsoralen and U.V. light-treated, culture trypomastigotes and challenged them with blood trypomastigotes. While all control mice developed parasitemia and died, all immunized mice “survived with negative parasitemia (checked daily) and absence of tissue lesions”. However, the immunized group presented lytic antibodies both before and after challenge, a result not consistent with absence of infection. Rodriguez et al. (1983) applied the CML technique to immunized-challenged rats 90 days after challenge, i.e., at a time when the lack of lytic antibodies would indicate complete protection. However, the results were positive for lytic antibodies. Thus, the overall results of this survey indicated that the exclusion of low levels of parasitism by lack of lytic antibodies had never revealed the success of immunoprotection.

In previous field studies using guinea pigs, we observed that a partial resistance, apparent with methods of low sensitivity, is easier to demonstrate than complete protection. Lack of infection was not demonstrated serologically (Basombrio 1990). In another field study performed in dogs (Basombrio et al. 1993) the exclusion of infected animals proceeded a step further by including serological data.

Identification of infected mice by CML showed in this work to be a useful and sensitive diagnostic method, since more advanced procedures, such as Polimerase Chain Reaction, are not yet available in all laboratories.

These results evidence that a significant number of immunized mice can totally reject an otherwise infectious T. cruzi challenge. They can not be attributed to the use of TCC parasites as immunogen, since these most often elicit a partial resistance. Total resistance at the skin level is a part, perhaps minor, among other, more apparent, components of immunity to T. cruzi. Its demonstration in individual animals seemed in this study to be associated with a particular set of experimental conditions, not often combined in previous experimental studies, such as threshold challenge, local delayed type hypersensitivity reaction and high natural host resistance. This set of experimental conditions, however particular, is by no means irrelevant to the usual way of transmission of T. cruzi from its insect vector to its reservoirs, including the human host.

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