EFFECTS OF BETAMETHASONE ON THE COURSE OF EXPERIMENTAL INFECTION WITH *TRYPANOSOMA CRUZI*

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In this experiment, the effect of betamethasone administered in the early post-acute infection of mice by *Trypanosoma cruzi* was studied. This drug was administered during 30 days after the 42nd day of infection in a dose of 0.15 mg/day. The betamethasone treatment did not cause fresh outbreaks of parasitemia and the histopathological findings in the chronic phase were not different from those in the control group. The higher cumulative mortality after treatment in the experimental group was due to superimposed bacterial infections. Outbred albino mice infected with low numbers of Y strain *Trypanosoma cruzi* trypomastigotes were not suitable models for Chagas' disease, since after 7 months of observation only mild histological lesions developed in all the animals. Prolonged betamethasone treatment of mice infected with low numbers of *Trypanosoma cruzi* of the Y strain, during the post-acute phase did not aggravate the course of infection.

Key words: Chagas' disease. Trypanosomiasis. Immunosuppression. Betamethasone. Light *Trypanosoma cruzi* infection.

The pathogenesis of chagasic cardiopathy has not yet been completely explained. However, several papers support the hypothesis that autoimmune mechanisms are involved. At present there is no experimental chronic Chagas' disease model of general acceptance, even though the canine models and the rabbit models have been suitable for some purposes. Recently, Laguens et al. proposed a mouse model using the Tulahuen strain of *Trypanosoma cruzi* in a low dose inoculum. The effects of immunosuppressive treatment during the acute phase or the established chronic phase of the disease are known. However immunosuppression in the early post-acute phase has not been extensively investigated.

The purpose of the present paper is to evaluate: 1) the suitability of low dose infection of mice with the Y strain of *Trypanosoma cruzi* as a model of Chagas' disease; 2) if immunosuppression in the early post-acute phase alters the course of infection.

MATERIAL AND METHODS

Animals: Outbred Swiss 3 month old albino male mice were used, weighing 35 ± 4.5 g (mean ± standard deviation).

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Infection: Twenty five blood trypomastigotes of the Y strain of *T. cruzi* (obtained from Dr. M.P. Deane, FIOCRUZ, Rio de Janeiro, in October 1983 and maintained by weekly passages in mice since then) were inoculated intraperitoneally. The inoculum estimation was done using a Neubauer hemocytometer.

Experimental group: Thirty infected and betamethasone treated mice.

Control group: Fifteen infected and non-treated mice.

Betamethasone: This drug was administered from the 42nd to the 75th day after inoculation by esophageal intubation in a dose of 0.15 mg/day.

Parasitemia and mortality: the parasitemia was estimated by Brener's technique at 5, 10, 15, 21, 30, 40, 65, 80 and 120 days after infection. Dead mice were removed daily.

Morphological study: All mice that had died spontaneously and those killed seven months after infection were autopsied. Samples from all the examined organs (heart, striated muscle, liver, spleen, intestines, kidneys and lungs) were fixed in 10% neutral formalin, embedded in paraffin, and sections cut at 5 micrometers and stained by hematoxylin-eosin and Mallory-Russel trichrome.

Statistical analysis: The parasitemias were compared by the "t" test and differences between cumulative mortality rates, were evaluated by normal approximation. The significance level accepted was 0.05.
RESULTS

The levels of parasitemia attained in all T. cruzi infected mice in the acute phase of the infection, are shown in Figure 1. The peak of parasitemia was reached at the 10th day after infection, when mice presented 8 ± 2 (X ± SD) parasites/50 microscopic fields of 400 X magnification. On the 40th day after inoculation only 2/38 of the mice continued with patent parasitemia (1 parasite/50 fields of 400 X) and parasitemia was negative thereafter.

As far as parasitemia was concerned, the betamethasone treated group did not differ from the control group, since no blood parasites were detected after immunosuppression.

The mortality in the acute phase was very low (8.0%). The cumulative mortality between the 42nd and the 200th day after infection is shown in Figure 2. At the end of the experiment the mortality rate observed in the experimental group (69.6%) as compared to that seen in the control (50%), was not significant, but after betamethasone treatment (75 days after inoculation) mortality was significantly higher in the experimental (56.5%) than in the control group (12.5%).

Morphological study: No gross lesions were detected in the hearts examined. In the acute phase of infection, only one mouse out of three that died spontaneously, showed a subacute, diffuse myocarditis with the presence of some pseudocysts in the myocardial fibers (Fig. 3). In the chronic phase of infection, the histopathological findings were mild and not important. Focal mononuclear infiltrates either in the myocardial interstitium or under the epicardium layer were seen as well as in the neighbourhood of parasym pathetic ganglia neurones displaying chromatolysis and other regressive lesions. Microfoci of myocardial fibrosis and myocytolysis were detected.

However these findings were not different from those observed in the control group. In the experimental group, the lesions seen in other organs were most frequent in the lungs and kidneys and resulted from superimposed bacterial infections. Septicemia was the most frequent cause of death, as indirectly evidenced through the frequent development of micro-abscesses in lungs, kidneys, spleen and liver; pulmonary oedema and hemorrhages; intensive generalized congestive changes; acute renal tubular necrosis and development of intensive acute cutaneous and organ infections (splenitis, pneumonitis, nephritis). These histological lesions have been taken as evidence that the administration of betamethasone by esophageal intubation had been effective in inducing immunosuppression in...
the animals. Liver and spleen amyloidosis were observed in seven mice from the experimental group and in none of the control animals.

Variable splenomegaly was observed in mice which died in the acute phase of infection (3 animals), as well as those in the chronic phase showing associated bacterial infections.

**DISCUSSION**

Laguens et al. demonstrated by morphological, immunological and electrocardiographic studies that the infection of three month old outbred mice with a low number of trypomastigotes of a highly pathogenic strain of *T. cruzi* (Tulahuen strain) leads to a chronic illness showing remarkable similarities to human Chagas’ disease. However, in the present experiment the infection of outbred mice with a small inoculum of *T. cruzi* (Y strain) did not seem to be a suitable model for Chagas’ disease. The histopathological findings were very mild, either in acute or chronic phases. Probably, the model proposed by Laguens et al. is dependent on the particular properties of the Tulahuen strain. The parasitemia and mortality rate were low and the parasitemic peak was delayed in our model, in comparison to the behavior of the Y strain of *T. cruzi* when high dose infections are induced.

The betamethasone treatment during 30 days in the period that followed immediately the acute phase, did not seem to reactivate the infection, since blood trypomastigotes could not be detected again. The effects of immunosuppression during acute Chagas’ disease are well known rendering the infection more severe. On the other hand, the administration of cortisone does not seem to alter the course of chronic infection in the rat or mouse. Brener & Chiaré have demonstrated that gamma radiation and cyclophosphamide could induce, in mice inoculated with the CL strain, an acute phase with outbreaks of high parasitemia and high mortality rate, whereas no increase of parasitemia was observed in animals inoculated with the Y, Berenice and PNM strains and similarly treated during the chronic phase. Additionally, some chronic chagasic patients submitted to immunosuppressive treatment did not apparently show any alterations in the course of the disease. However Rassi et al. observed increase of parasitemia in 8 out of 11 chronic chagasic patients treated with corticoid.

The higher cumulative mortality after corticoid treatment as seen in the experimental group in this paper, was due to superimposed bacterial infections.

The occurrence of severe infections complicated by septicemias was very high in this group indicating that the betamethasone dosage was effective. It can therefore be concluded that under the present experimental conditions, betamethasone treatment during the early post-acute phase does not modify the course of chronic Chagas’ disease, since neither the experimental nor the control groups showed characteristic lesions of the chronic illness.

**RESUMO**

Foram estudados os efeitos da betametasona administrada na fase pos-aguda imediata de uma infecção pelo *T. cruzi* em camundongos. O tratamento consistiu de 30 doses diárias de 0,15 mg de betametasona, a partir de 42° dia de infecção, não havendo aparecimento de novos surtos de parasitemia. No tempo de duração do experimento (7 meses) não houve diferença entre as lesões histopatológicas dos animais tratados e dos não tratados. O grupo experimental apresentou uma maior mortalidade acumulada no 75° dia de infecção, o que pode ser atribuído a infecções bacterianas associadas. Por outro lado, camundongos albinos “outbred”, infectados com baixo inoculo, não se apresentaram como bom modelo de doença de Chagas, já que não desenvolveram lesões importantes nem na fase aguda nem após 7 meses de infecção. Em conclusão, o tratamento imunosupressivo prolongado, após a fase aguda de uma infecção mínima com a cepa Y do *T. cruzi* não tem influência sobre o curso da infecção, pelo menos no que tange ao agravo da mesma.


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