Acute phase proteins: a potential approach for diagnosing chronic infection by *Trypanosoma vivax*

Proteínas de fase aguda: uma possível abordagem para diagnóstico de infecção crônica por *Trypanosoma vivax*

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Abstract

The present study aimed to assess potential changes in acute phase proteins in sheep experimentally infected with *Trypanosoma vivax*. There were studied eight male sheep, four used as controls and four infected with $10^5$ *T. vivax* tripanastigotes. Blood samples were collected at two points times before infection and then at 5, 7, 9, 11, 13, 15, 20, 30, 45, 60, 75, 90, 105 and 120 days post-infection (dpi). Blood samples were centrifuged and allotted, and acute phase proteins were then separated by electrophoresis on acrylamide gel containing sodium dodecyl sulfate. Protein concentrations were determined by computer-assisted densitometry. Total protein was determined by colorimetric biuret method. Trypanosomes were counted daily using a 5 mL aliquot of blood smear on a glass slide under a 22 × 22 mm coverslip. Parasites were counted in 100 microscopic fields (40× magnification), and then multiplied by a correction factor. The results were expressed as parasites per mL of blood. For statistical analyses, we used the Wilcoxon test at 5% significance level. There was found a reduction in several acute phase proteins and increase in antitrypsin and transferrin. This finding can be used for the diagnosis of *T. vivax* infection, especially in chronic infection.

Keywords: Antitrypsin, protein profile, transferrin, sheep.

Resumo

O objetivo do presente estudo foi verificar possíveis alterações nas proteínas de fase aguda em ovinos infectados experimentalmente com *Trypanosoma vivax*. Para tanto, foram utilizados oito ovinos machos, sendo quatro usados como controle e quatro infectados com $10^5$ tripanastigotes de *T. vivax*. Colheram-se amostras de sangue em dois tempos antes da infecção e, posteriormente, aos 5, 7, 9, 11, 13, 15, 20, 30, 45, 60, 75, 90, 105 e 120 dias após a infecção (dpi); após centrifugação e aliquotização das amostras. As proteínas de fase aguda foram separadas por eletroforese em gel de acrilamida, contendo dodecil sulfato de sódio, e suas concentrações foram determinadas através de densitometria computadorizada. A dosagem de proteína total foi realizada pelo método colorimétrico do biureto. A contagem dos tripanossomas foi realizada diariamente, utilizando-se uma aliquota de 5 μL de sangue disperso em lâmina de microscopia, sob lamínula de 22 × 22 mm, contando-se os parasitos em 100 campos microscópicos, com objetiva de 40×, multiplicados pelo fator de correção do microscópio, e o resultado expresso em parasitos por mL de sangue. Para a análise estatística, empregou-se o teste de Wilcoxon a 5% de probabilidade. Foi observada a diminuição de diversas proteínas de fase aguda e aumento de antitripsina e transferrina que podem ser utilizadas para auxiliar no diagnóstico da infecção por *T. vivax*, principalmente na fase crônica da infecção.

Palavras-chave: Antitripsina, proteinograma, transferrina, ovinos.
Trypanosoma vivax infects cattle and other domestic and wild ruminants (DÁVILA; SILVA, 2000). In Brazil this pathogen causes economic losses due to abortions, infertility, reduced performance, decreased production and mortality (SEIDL et al., 1999).

The diagnosis of animal trypanosomiasis may be difficult as there are no pathognomonic clinical signs of infection and standard trypanosome detection methods are not sensitive enough. Despite advances in the diagnosis of animal trypanosomiasis, many acute infections go unnoticed and a chronic form of disease, frequently with no parasitemia, is more prevalent (NANTULYA, 1990).

Animal trypanosomiasis is detected using parasitological, immunological and molecular diagnostic techniques. Micro-hematocrit centrifugation (Woo technique) is the most widely used technique as it can detect parasites six to 10 days earlier than with fresh blood smears. Although it is simple, Woo technique has low sensitivity for chronic infection (MASAKE et al., 1995).

Most immunological methods can detect antibodies to trypanosome antigens. Indirect immunofluorescence and ELISA are the most valuable methods, particularly for T. vivax distribution, because of their sensitivity and specificity. But cross-reactivity between antigens of parasites within the same genus is a major limitation of serological tests, reducing their specificity (NANTULYA, 1990; JONES; DÁVILA, 2001).

Other more sophisticated diagnostic techniques have been employed for T. vivax detection such as polymerase chain reaction (PCR). PCR was evaluated by Desquesnes (1997) and showed excellent results in frozen serum samples. It provided highly specific results with sensitivity similar to parasitological techniques, making it a valuable method for the analysis of field samples when well-equipped laboratories are not available.

The study of acute phase proteins can be a helpful approach for diagnosing T. vivax, especially in chronic infection when parasites are difficult to be detected in the bloodstream. According to Godson et al. (1996), the level of acute phase proteins changes during an infection, depending on the causative agent and the animal species infected. Acute protein profile can be used to facilitate the diagnosis of T. vivax infection.

Cerón et al. (2005) stressed that acute phase proteins increase in animals with different conditions but because of its low diagnostic specificity it is not used for primary diagnosis as a cause of disease cannot be determined. However, it is highly sensitive for detection of inflammation or subclinical infection. Although protein profile does not provide specific information, it can guide the diagnosis and prognosis of diseases.

Protein profiling is a significant approach for diagnosis and monitoring of several diseases in dogs (OHWADA; TAMURA, 1993; CÉRON et al., 2005) and calves infected with Mannheimia haemolytica (FAGLIAI et al., 2003). In addition, it has been used for prognosis postoperatively (FAGLIAI et al., 2008; SAQUETTI et al., 2008).

The present study aimed to assess potential changes in acute phase proteins in sheep experimentally infected with Trypanosoma vivax.

Material and Methods

The T. vivax sample was obtained from cattle in the region of Catolé do Rocha, northeastern Brazil. Two mL of blood with approximately 1.95 x 10^7 T. vivax trypomastigotes were inoculated intravenously in an one-year-old no-breed male ovine for parasite multiplication. A total of eight no-breed male sheep, aged one year were used in the study. They went through a quarantine period for clinical, hematological, biochemical and stool examination, and detection of anti-T. vivax antibodies by indirect immunofluorescence assay (IFA). Healthy animals were kept in single stalls with raised floor and screened panels, fed a corn silage diet and water ad libitum and supplemented with mineral salt for 120 days. They were then randomly assigned to two groups: four were used as controls and the other four were infected intravenously with 1 mL of blood containing 10^8 T. vivax trypomastigotes.

For monitoring parasitemia, from each animal inoculated with T. vivax trypomastigotes, 0.5 mL blood was collected daily from the jugular vein with 10% EDTA up to 120 days post infection (dpi). Trypanosomes were counted daily using a 5 mL aliquot of blood smear on a glass slide under a 22 x 22 mm coverslip. Parasites were counted in 100 microscopic fields (40x magnification), and then multiplied by a correction factor. The results were expressed as parasites per mL of blood.

Three mL of blood were collected from the jugular vein without using an anticoagulant for serum analysis at 14 and 7 days before infection (for baseline, maximum and minimum values for each parameter evaluated) and then at 5, 7, 9, 11, 13, 15, 20, 30, 45, 60, 75, 90, 105 and 120 dpi.

Total protein was determined by the colorimetric biuret method using commercial kits (Labtest® Diagnóstica, Belo Horizonte, Brazil) and then read on a spectrophotometer (LabQuest® Diagnóstica, Belo Horizonte, Brazil).

Acute phase proteins were separated by electrophoresis on acrylamide gel containing sodium dodecyl sulfate (SDS-PAGE), as described by Laemmli (1970). After proteins were separated, the gel was stained with coomassie blue solution for 10 min and then added to a solution of 7% acetic acid to remove excess dye until protein fractions became evident. Protein concentrations were determined by computer-assisted densitometry (Shimadzu CS 9301, Tokyo) using as reference protein molecular weight markers (Sigma, St. Louis, MO, US) (29,000, 45,000, 66,000, 97,400, 116,000 and 205000 daltons), in addition to purified proteins (albumin, IgG, haptoglobin, α1-antitrypsin and transferrin).

All statistical analyses were performed in System for Statistical Analysis (SAEG) and nonparametric Wilcoxon test at 5% significance level was used to assess changes in acute phase proteins.

Results and Discussion

The prepatent period was one day in one animal and two days in the other animals inoculated with parasites. Parasitemia was intermittent and its level varies in all infected animals up to 120 dpi.

Using the SDS-PAGE technique 26 plasma proteins were separated from the eight animals, and their molecular weights ranged from 13,000 to 170,000 daltons. Twelve showed statistically
significant differences (p < 0.05) (molecular weights in daltons): 138,000 (IgA); 130,000 (unknown); 125,000 (ceruloplasmin); 115,000 (unknown); 79,000 (hemopexin); 72,000 (transferrin); 60,000 (antitrypsin); 59,000 (heavy IgG); 55,000 (unknown); 51,000 (unknown); 32,000 (light IgG); 30,000 (unknown).

Of these 12 proteins with significant differences, 10 showed a reduction in the animals infected compared to controls. Four acute phase proteins had a significant reduction which was more markedly seen for ceruloplasmin and hemopexin. And antitrypsin and transferrin had an early reduction (5 dpi) in the animals infected compared to Day zero and controls, and then an increase above baseline in the chronic phase of infection with the highest levels seen for antitrypsin (Table 1).

Acute phase proteins are often measured at 2 hours intervals as they are no longer detectable in the chronic phase. But as parasitemia, though at lower levels, was maintained in sheep infected with T. vivax during the chronic phase of infection, the production of acute phase proteins was stimulated. Thus, the determination of acute phase proteins can help the diagnosis of T. vivax infection.

Ohwada and Tábara (1993) found that parvovirus infection in dogs can be diagnosed two weeks before the onset of clinical signs and/or other diagnostic methods, they may help diagnosing increase above baseline after 30 dpi, and together with clinical signs and/or other diagnostic methods, they may help diagnosing chronic infection (after 30 dpi) given that parasite identification using routine diagnostic methods can be sometimes difficult due to low parasitemia.

Table 1. Means and standard deviations of total protein (g/dL) and acute phase proteins (g/dL) identified in the electrophoretic protein profile (SDS-PAGE) of sheep – controls and those experimentally infected with T. vivax during the experiment period.

| Protein (g/dL) | Days post infection |
|---------------|---------------------|
|               | 0       | 5       | 7       | 9       | 11      | 13      | 15      | 20      | 30      | 45      | 60      | 75      | 90      | 105     | 120     |
| **Total protein** |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |
| Infected      | 5.31a   | 4.77a   | 5.03a   | 5.35a   | 4.81b   | 4.85b   | 4.70b   | 5.03b   | 5.41b   | 5.53b   | 7.17a   | 6.61a   | 6.81a   | 7.28b   |         |
| Controls      | 5.33a   | 6.14*   | 4.99a   | 5.46a   | 5.44a   | 5.85a   | 5.49a   | 6.13a   | 6.02a   | 5.91a   | 5.86a   | 5.38a   | 6.00a   | 6.17a   | 6.30a   |         |
|               | (0.29)  | (0.32)  | (0.27)  | (0.16)  | (0.94)  | (0.55)  | (0.58)  | (0.25)  | (0.42)  | (0.43)  | (0.43)  | (0.39)  | (0.03)  | (0.43)  | (0.52)  | (0.34)  |
| **Ceruloplasmin** |        |         |         |         |         |         |         |         |         |         |         |         |         |         |         |
| Infected      | 0.033a  | 0.018b  | 0.022b  | 0.021b  | 0.020a  | 0.019b  | 0.016b  | 0.014b  | 0.011b  | 0.013b  | 0.016b  | 0.016b  | 0.023b  | 0.019b  |         |
| Controls      | 0.030a  | 0.037b  | 0.030a  | 0.031b  | 0.032b  | 0.038b  | 0.033b  | 0.037b  | 0.036b  | 0.035b  | 0.037b  | 0.034b  | 0.037b  | 0.039b  | 0.036b  |         |
|               | (0.015) | (0.008) | (0.007) | (0.008) | (0.007) | (0.006) | (0.005) | (0.005) | (0.005) | (0.005) | (0.007) | (0.005) | (0.010) | (0.010) | (0.010) |         |
| **Hemopexin** |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |
| Infected      | 0.014a  | 0.009b  | 0.011b  | 0.010b  | 0.011b  | 0.007b  | 0.007b  | 0.005b  | 0.004b  | 0.004b  | 0.005b  | 0.007b  | 0.006b  | 0.005b  | 0.005a  |         |
| Controls      | 0.015a  | 0.017a  | 0.013b  | 0.015b  | 0.017b  | 0.017b  | 0.017b  | 0.018b  | 0.017b  | 0.016b  | 0.016b  | 0.016b  | 0.016b  | 0.017b  | 0.017b  |         |
|               | (0.002) | (0.005) | (0.002) | (0.002) | (0.002) | (0.002) | (0.002) | (0.002) | (0.002) | (0.002) | (0.002) | (0.002) | (0.004) | (0.004) | (0.003) |         |
| **Transferrin** |        |         |         |         |         |         |         |         |         |         |         |         |         |         |         |
| Infected      | 0.360a  | 0.237b  | 0.376b  | 0.405b  | 0.522a  | 0.512a  | 0.485b  | 0.513a  | 0.664a  | 0.744a  | 0.634a  | 0.745a  | 0.620a  | 0.588a  | 0.570a  |         |
| Controls      | 0.361a  | 0.394a  | 0.352a  | 0.389a  | 0.401a  | 0.397a  | 0.407a  | 0.407a  | 0.423a  | 0.424a  | 0.410b  | 0.364b  | 0.422b  | 0.418b  | 0.448a  |         |
|               | (0.064) | (0.120) | (0.050) | (0.073) | (0.096) | (0.078) | (0.082) | (0.073) | (0.083) | (0.064) | (0.056) | (0.072) | (0.081) | (0.074) | (0.062) |         |
| **Antitrypsin** |        |         |         |         |         |         |         |         |         |         |         |         |         |         |         |
| Infected      | 0.183a  | 0.147a  | 0.182a  | 0.198a  | 0.219a  | 0.184a  | 0.206a  | 0.181a  | 0.207b  | 0.291a  | 0.355a  | 0.530a  | 0.548a  | 0.610a  | 0.702a  |         |
| Controls      | 0.171a  | 0.185a  | 0.164a  | 0.169a  | 0.155a  | 0.192a  | 0.181a  | 0.193a  | 0.189a  | 0.183a  | 0.187a  | 0.164a  | 0.169a  | 0.180a  | 0.175a  |         |
|               | (0.015) | (0.028) | (0.048) | (0.025) | (0.030) | (0.025) | (0.030) | (0.025) | (0.040) | (0.031) | (0.031) | (0.047) | (0.046) | (0.027) | (0.045) |         |

Means followed by the same letter are not statistically different in the Wilcoxon test at 5% significant level. ns Non significant in the Wilcoxon test at 5% significant level. Standard deviations in brackets.
The electrophoretic profile of acute phase proteins has the additional advantages of reducing costs and not requiring preparation of glass slides with trypanosomes (as in IFA), and in vitro or in vivo parasite growth, which may be a complex procedure involving inoculation of *T. vivax* in laboratory animals and preparation of antigens for diagnostic use.

**Conclusions**

The electrophoretic protein profile in sheep infected with *T. vivax* showed a different pattern with a reduction of several acute phase proteins and an increase in antitrypsin and transferrin. These proteins can be used to help diagnose infection, especially chronic infection.

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