Antigenic characterization of *Trypanosoma evansi* using sera from experimentally and naturally infected bovines, equines, dogs, and coatis

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Received May 18, 2009
Accepted June 16, 2009

Abstract

The present research investigated the presence of *T. evansi* antibodies in animals from the subregion of Nhecolandia, in the Pantanal Sul-mato-grossense, by means of an enzyme linked immunosorbent assay (ELISA) and indirect immunofluorescence antibody test (IFAT), and the pattern of polypeptide recognition by sera from experimentally and naturally infected hosts using Western blotting. Serum samples were obtained from bovines (n = 102), horses (n = 98), and dogs (n = 55), and from 32 free-ranging coatis (*Nasua nasua*). None of the bovines were found positive, while sera from 16 dogs (29%) and 23 horses (23.4%) were positive by ELISA. Sera from 8 coatis (25%) were found positive using IFAT. Western blotting revealed major polypeptides of *T. evansi* with molecular weight ranging from 74 to 38 kDa. The polypeptides of 66, 48-46, and 38 kDa were identified by sera from experimentally infected bovines, donkeys, dogs, and coatis. The 48-46 and 38 kDa bands were mainly recognized in chronic phase of infection. The antigen with apparent molecular weight of 66 kDa, revealed by antibodies from all experimental animals, was also recognized in sera of horses and dogs from the Pantanal. The 48-46 kDa polypeptide was identified by antibodies from all naturally infected animals and must be further evaluated for use in specific diagnosis of *T. evansi* infection.

Keywords: *Trypanosoma evansi*, ELISA-test, IFAT, Pantanal mato-grossense, antigenic characterization.
Introduction

Trypanosoma evansi is a widely distributed hemoflagellate parasite that affects a wide range of animals, causing high morbidity and mortality in some species, especially horses and dogs. In Brazil, the disease caused by T. evansi and known as “Mal de Cadeiras” is considered endemic in the Pantanal region, where the parasite is found in horses, dogs, cattle, buffaloes, capybaras, and coatis (STEVENS et al., 1989; NUNES; OSHIRO, 1990; NUNES et al., 1993; FRANKE; GREINER; MEHLITZ, 1994). This trypanosomiasis is the principal protozoal disease of horses in the region, where recent outbreaks with high mortality and mortality have been reported (SILVA et al., 1995a, b), and assumes great importance since horses are indispensable to management in extensive cattle ranching, the main economic activity in the Pantanal (SILVA; BARRÓS; HERRERA, 1995; SEIDL; MORAES; SILVA, 1998).

The diagnosis of “Mal de Cadeiras” remains problematic and commonly used tests have important limitations (DAVISON et al., 1999). Diagnosis of trypanosomes by direct parasitological techniques is not reliable, specially for naturally occurring field cases, where parasitemias are often low and sporadic (RAE; LUCKINS, 1984; BENGAŁY; KANWE; DUVALLET, 1995; NANTULYA, 1994). Serological tests for circulating antibody, ELISA and IFAT, have been intensively employed, showing good sensitivity (PAYNE et al., 1991; MUKANI et al., 1992; MONZON; HOYOS; JARA, 1995). But antibody detection assays are often hampered by the lack of specificity, the result of strong cross-reactions with other pathogenic trypansomones (DESQUENES et al., 2001; UZCANGA et al., 2002). Antigen detection ELISAs have been used in various animal species (DIALL et al., 1992; MONZON; HOYOS; JARA, 1995; KASHIWAZAKI et al., 1998). However, proper evaluation of the diagnostic parameters in these assays showed poor results (DAVISON et al., 1999).

Few studies on antigenic characterization of T. evansi have been undertaken (UCHE; JONES; BOID, 1993; UCHE; JONES, 1994; QUEIROZ et al., 2001) and little is known so far regarding the polypeptide recognition patterns by antibodies from different host species.

The aim of this research was to investigate the presence of T. evansi antibodies in animals from the subregion of Nhecolandia in the Pantanal Sul-mato-grossense, and to study the patterns of polypeptide recognition of a Brazilian T. evansi stock by sera from naturally infected seropositive animals and its correlation with the profile obtained by sera from experimentally infected hosts in acute and chronic stage of the infection.

Material and Methods

1. Animals

A total of 98 horses, 102 bovines, 55 dogs, and 32 coatis from three different properties located in the Pantanal of Nhecolandia - MS, were randomly selected for this study. Blood samples collected from the jugular vein were allowed to clot at room temperature, and then centrifuged. Serum was stored at −20 °C.

Serum samples obtained in previous studies of experimental T. evansi infection in dogs (AQUINO et al., 1999), bovines (POCHINI, 2000), coatis (HERRERA et al., 2001), and donkeys (CADIOLI, 2001) were used. Sera obtained from the experimentally infected animals before inoculation and on different days during infection were used in Western blotting and as reference negative and positive control sera in serology.

2. Antigen preparation

Antigens were prepared from a T. evansi stock originally isolated from a dog naturally infected in the Pantanal of Nhecolandia (AQUINO et al., 1999). Antigenic substrate for IFAT and soluble antigen used in ELISA and Western blotting were prepared according to Aquino et al. (1999). Protein content of the soluble antigen was determined by the bicinecinic acid method (BCA Reagent Kit – Pierce Chemical Company).

3. Serology

Serum samples from horses, bovines, and dogs were evaluated for the presence of anti-trypanosome antibodies by enzyme linked immunosorbent assay (Ab-ELISA). Indirect fluorescent antibody test (IFAT) was used for coati sera. Both tests were carried out essentially as described by Aquino et al. (1999) and briefly described below.

ELISA: Plates were coated with 5, 10, and 15 µg of soluble antigen in horse, dog, and bovine assays, respectively. Bovine, equine, and dog sera were respectively diluted 1:100, 1:200 and 1:400. Alkaline phosphatase conjugated rabbit anti-dog IgG (SIGMA A-6042), anti-horse IgG (SIGMA A-6063), and anti-bovine IgG (SIGMA A-0668) were diluted following manufacturer’s instructions. All test and control sera in serology.

IFAT: Successive coati serum samples at dilutions from 1:40 (1:40, 1:80, 1:160…) were tested. Goat anti-racoon IgG (KPL02-32-06) fluorescent conjugate was used in 1:10 dilution. The IFAT reaction was considered positive at serum dilutions ≥ 1:80.

4. SDS-PAGE

T. evansi soluble proteins were separated by electrophoresis (Mini-Protein II, Bio-Rad) on a 8-18% gradient polyacrylamide gel in the presence of sodium dodecyl sulfate (LAEMMLI, 1970). Samples containing 18.5 µg of soluble proteins were loaded into each well and a mixture of calibration proteins of molecular range between 220 and 14.3 kDa (“Rainbow,” Amersham Life Science RPN 756) was included in one well during each run. All gels were run at 200 volts.
5. Western blotting

Following SDS-PAGE, the polypeptides were electrophoretically transferred at 75 V for 3 hours from the gel to a nitrocellulose membrane by the procedure described by Towbin, Stahelin e Gordon (1979).

Membranes were blocked for 12 hours with 5% nonfat dry milk and 0.05% Tween-20 in Tris buffered saline (TBS: 20 mM Tris, 500 mM NaCl, pH 7.5).

For immuno-detection, strips of nitrocellulose containing individual runs were cut and incubated with test sera. Sera from experimentally infected animals (N donkeys, three donkeys, four dogs, and four coatis) were pooled for each sampling date along the acute and chronic phases of the infection. Sera from seropositive horses, bovines, dogs, and coatis from the Pantanal were evaluated individually against T. evansi antigens. Incubations were performed at room temperature. All sera were diluted 1:100 in blocking buffer (5% nonfat dry milk in TBS-Tween) and added to the membrane strips for 3 hours. Sera were removed and membranes were washed with blocking solution for 10 minutes, followed by two washes (5 minutes each) with TBS-Tween. The strips were then incubated for 90 minutes with appropriate dilution of alkaline phosphatase conjugated antibodies against horse, bovine, dog, and raccoon IgG as already referred to for serology. Anti-raccoon IgG (KPL 05-32-06) was used for detection of antibodies in coati sera, following the instructions of the supplier. Conjugate was removed and three washes (5 minutes each) were performed using TBS-Tween. The polypeptide bands were visualized by the addition of the enzyme substrate 5-bromo-4-chloro-3-indoly phosphate/nitroblue tetrazolium chloride (NBT-BCIP, SIGMA B-5655). Colorimetric reactions were stopped by washing the blots in distilled water.

Sera obtained from experimentally infected animals before and after infection with T. evansi served as the negative and positive controls, respectively. Relative mobility of immunorecognized polypeptides was estimated from a calibration curve based on standards for mobility of proteins.

Results

1. Serology

Antibodies of diagnostic value were detected in sera from 16 dogs (29.1%), 23 horses (23.5%), and 8 coatis (25%). None of the bovines were found positive by serology (Table 1).

2. Antigenic characterization of T. evansi

IgG antibodies to T. evansi polypeptides were first detected in serum of experimentally infected animals from 4 to 7 days after infection. The molecular weight of the T. evansi antigens recognized by IgG in experimentally infected donkeys, bovines, dogs, and coatis ranged from 160 to 15 kDa. The bands corresponding to the polypeptides of 160, 88, 74, 66, 52/50, 48/46, 38, 32/30, 27, 25, 20, and 17 kDa were more consistently and/or more intensely stained. The polypeptides of 74 and 66 kDa were labeled as early as 4 to 5 days after experimental infection by bovine, donkey, and dog antibodies. As the infection progressed increasing numbers of trypanosome antigens were recognized, with the staining of most individual bands becoming more evident. Differences were observed among studied species in the recognition pattern and intensity of labeling of some polypeptides (Table 2 and Figure 1). The polypeptides of 74, 27, and 20 kDa were recognized by dog, donkey, and bovine sera, while those of 25 and 17 kDa were identified by donkeys and dogs; the 88, 52, and 32 kDa polypeptides were common to bovines and donkeys and the 160 kDa band was recognized by dogs and coatis. The polypeptides of 66, 48-46 and 38 kDa were identified by sera from experimentally infected bovines, donkeys, dogs, and coatis. The 48-46 and 38 kDa bands were mainly recognized in chronic phase of infection.

Antibodies of naturally infected animals from the Pantanal recognized the same polypeptides labeled by antibodies of the corresponding experimentally infected species, with variations in the number of antigens recognized occurring between animals (Figure 2). Four lower molecular weight proteins (32, 30, 27, and 25 kDa) that were detected during chronic phase of experimental infection in donkeys were not recognized by sera from 5 of 23 seropositive horses from the Pantanal. Five out of 29 seropositive naturally infected dogs recognized three polypeptides (88, 32, and 30 kDa) not labeled by pooled dog sera collected along experimental infection. Differences in immunorecognition profile were more evident. Differences were observed among studied species in the recognition pattern and intensity of labeling of some polypeptides (Table 2 and Figure 1). The polypeptides of 74, 27, and 20 kDa were recognized by dog, donkey, and bovine sera, while those of 25 and 17 kDa were identified by donkeys and dogs; the 88, 52, and 32 kDa polypeptides were common to bovines and donkeys and the 160 kDa band was recognized by dogs and coatis. The polypeptides of 66, 48-46 and 38 kDa were identified by sera from experimentally infected bovines, donkeys, dogs, and coatis. The 48-46 and 38 kDa bands were mainly recognized in chronic phase of infection.

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Table 1. Detection of anti- T. evansi antibodies (number and percentage of positive sera) in serum samples of horses, dogs, and coatis from Nhecolândia Pantanal, MS.

<table>
<thead>
<tr>
<th>Host species</th>
<th>Test</th>
<th>Positive animals</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>%</td>
</tr>
<tr>
<td>Horses</td>
<td>(Ab - ELISA)</td>
<td>23 (98)</td>
</tr>
<tr>
<td>Dogs</td>
<td>(Ab - ELISA)</td>
<td>16 (55)</td>
</tr>
<tr>
<td>Coatis</td>
<td>(IFAT)</td>
<td>8 (32)</td>
</tr>
</tbody>
</table>

Table 2. T. evansi polypeptides recognized by antibodies of experimentally and naturally infected bovines, donkeys/horses, dogs, and coatis by Western blotting.

<table>
<thead>
<tr>
<th>PM (kDa)</th>
<th>Bovine</th>
<th>Donkey/Horse</th>
<th>Dog</th>
<th>Coati</th>
</tr>
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<tbody>
<tr>
<td>E</td>
<td>N *</td>
<td>E</td>
<td>N</td>
<td>E</td>
</tr>
<tr>
<td>160</td>
<td>x</td>
<td>-</td>
<td>-</td>
<td>x</td>
</tr>
<tr>
<td>88</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>-</td>
</tr>
<tr>
<td>74</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
</tr>
<tr>
<td>66</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
</tr>
<tr>
<td>52/50</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
</tr>
<tr>
<td>48/47/46</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
</tr>
<tr>
<td>38</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
</tr>
<tr>
<td>32/30</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
</tr>
<tr>
<td>27</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
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<tr>
<td>25</td>
<td>-</td>
<td>-</td>
<td>x</td>
<td>o</td>
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<tr>
<td>20</td>
<td>x</td>
<td>-</td>
<td>x</td>
<td>o</td>
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<tr>
<td>17</td>
<td>-</td>
<td>-</td>
<td>x</td>
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E = Experimentally infected; * All cattle were negatives.
N = Naturally infected.
evident between individual free-ranging coatis. Three of them recognized only the 48 kDa polypeptide and a band corresponding to the polypeptides from 110 to 160 kDa. The 66 kDa antigen was labeled by sera from two animals. Serum from one free-ranging coati, however, recognized additionally the polypeptides of 30, 25, and 23 kDa, not labeled by antibodies from experimentally infected coatis.

**Discussion**

In the present study, five major polypeptides with apparent molecular weights 88, 74, 66, 32, and 27 kDa recognized by antibodies of experimental animals, appear to correspond to the 85, 75.5, 67, 32.4, and 28 kDa antigens of *Trypanosoma evansi* revealed by sera of rabbits experimentally infected with an Indonesian stock of the parasite (UCHE; JONES; BOID, 1993). Similarly, the polypeptides of 88, 74, 66, 46, 32, and 17 kDa recognized in our study probably correspond to 86, 73, 68 46, 32, and 17 kDa components labeled by antibodies of horses experimentally infected with a Venezuelan isolate of *T. evansi* (UZCANGA et al., 2002). The increasing numbers of polypeptides recognized by experimental infected bovines, donkeys, dogs, and coatis with the increasing duration of infection in the present study is in agreement with earlier observations made in experimentally infected rabbits (UCHE; JONES; BOID, 1993). The greater number of polypeptides detected in chronic phase of infection is probably due to the release of internal antigens after parasite destruction by variable surface glycoprotein (VSG) specific antibodies. Our results differ however from those of Queiroz et al. (2001), who observed no variation in polypeptide profiles along the course of experimental infection in rats with Brazilian isolates of *T. evansi*.

Differences were observed among studied species in the recognition pattern and intensity of labeling of some polypeptides. The polypeptides of 66, 48-46, and 38 kDa were identified by sera from all experimentally infected hosts in this research.

The analysis of cell membrane preparations of seven different stocks of *T. evansi* isolated from buffaloes, equines, or camels in India revealed polypeptides with molecular weight ranging from 48.4 to 80.2 kDa (SINGH, V.; SINGH, A.; CHHABRA, 1995).

![Figure 1](image-url)  
*Figure 1.* Immunoreactivity of polled sera from experimentally infected bovines (a), horses (b), dogs (c), and coatis (d) obtained before and in different days along infection period. Labels below each blot: “0” indicates sera taken before experimental inoculation, and consecutive numbers indicate days after infection when sera were collected.
Major polypeptides recognized by experimental animals in our study were those with 74 to 48-46 kDa and consequently within the molecular weight range of surface antigens. Recently, Laha and Sasmal, 2008 showed 11 immunogenic proteins using hyperimmune sera from buffalo, horse and cattle, but in naturally *T. evansi* infected immune sera of horse detected 19 immunogenic proteins. Five common immunogenic proteins of relative molecular weight ranged 61-64, 44-47, 33-34, 25-26 and 14-16 kDa using hyperimmune and immune sera. The band corresponding to the 66 kDa polypeptide was firstly recognized as early as five days after infection by all host species in the present study and remained continuously labeled during the experimental period. Besides being revealed by antibodies from all experimental animals, the 66 kDa polypeptide was also recognized by sera of all naturally infected horses and dogs and of most free-ranging coatis. A polypeptide with similar mass (67 kDa) previously identified as a surface antigen of *T. evansi* (UCHE; ROSS; JONES, 1992) was also consistently recognized by rabbits experimentally infected (UCHE; JONES; BOID, 1993) and showed to be associated with protection (UCHE et al., 1994). The 66 kDa polypeptide detected in our experiment may correspond to the 64 kDa polypeptide recognized in a Venezuelan stock of the parasite by sera of experimentally infected horses (UZCANGA et al., 2002).

The same study, however, showed that this polypeptide exhibited cross-reactivity with *T. vivax*. This finding is of concern since *T. vivax* also occurs in the Brazilian Pantanal (SILVA et al.,1996), which limits the possibility of utilizing this polypeptide as a tool for specific diagnosis of *T. evansi* infection in the region.

Additional detailed studies using Western blotting in characterization of different stocks of *T. evansi* by antibodies of various host species may enhance the understanding of the host-parasite relationship and give some insight into the pathogenicity and immunogenicity of this parasite. Specific parasite polypeptides identified could be further evaluated for use in the diagnosis of this trypanosomiasis.
Acknowledgements

To Rosângela Yamazaki de Andrade for technical assistance. This work was supported by Fundação de Amparo a Pesquisa do Estado de São Paulo (FAPESP, nº 2001/ 08097-6).

References


