Western Blotting Using Strongyloides ratti Antigen for the Detection of IgG Antibodies as Confirmatory Test in Human Strongyloidiasis


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The present study was conducted to evaluate the frequency of antigenic components recognized by serum IgG antibodies in Western blotting (WB) using a Strongyloides ratti larval extract for the diagnosis of human strongyloidiasis. In addition, the WB results were compared to the enzyme-linked immunosorbent assay (ELISA) and the indirect immunofluorescence antibody test (IFAT) results. Serum samples of 180 individuals were analyzed (80 with strongyloidiasis, 60 with other intestinal parasites, and 40 healthy individuals). S. ratti was obtained from fecal culture of experimentally infected Rattus rattus. For IFAT, S. ratti larvae were used as antigen and S. ratti larval antigenic extracts were employed in WB and ELISA. Eleven S. ratti antigenic components were predominantly recognized by IgG antibodies in sera of patients with strongyloidiasis. There was a positive concordance for the three tests in 87.5% of the cases of strongyloidiasis. The negative concordance in the three tests was 94% and 97.5% in patients with other intestinal parasites and healthy individuals, respectively. In cases of positive ELISA and negative IFAT results, diagnosis could be confirmed by WB. ELISA, IFAT, and WB using S. ratti antigens showed a high rate of sensitivity and specificity. In conclusion, WB using S. ratti larval extract was able to recognize 11 immunodominant antigenic components, showing to be a useful tool to define the diagnosis in cases of equivocal serology.

Key words: Strongyloides stercoralis - Strongyloides ratti - strongyloidiasis - immunodiagnosis - Western blotting

Strongyloidiasis, caused by the nematode Strongyloides stercoralis, is one of the major intestinal infections in humans and is distributed worldwide in both tropical and temperate countries (Grove 1996). In Brazil the infection rates vary according to the region. Uberlândia, a city located in the Southeastern region, state of Minas Gerais, was considered as hyperendemic area for S. stercoralis by Machado and Costa-Cruz (1998) through parasitological findings demonstrating that 13% of the children were infected. Considering the severity of this disease in immunocompromised patients, Ferreira et al. (1999) described 25 cases of S. stercoralis and human immunodeficiency virus (HIV) infections in patients followed up in School Hospital, including seven cases of hyperinfection syndrome, which were further confirmed in autopsy through dissemination to various organs.

One of the current problems concerning strongyloidiasis is the difficulty to detect larvae in human feces, because the majority of cases involve chronic, low-level infection, and the larval output is minimal and irregular. Although the agar plate method for stool culture appears to be more sensitive than common parasitological methods, it is more time consuming (Arakaki et al. 1990, Sato et al. 1995, Dreyer et al. 1996, Uparanukraw et al. 1999).

Detection of parasite-specific antibodies by indirect immunofluorescence antibody test (IFAT) and enzyme-linked immunosorbent assay (ELISA) may be a useful complement to the parasitological diagnosis of strongyloidiasis (Conway et al. 1993a, Costa-Cruz et al. 1997, De Paula et al. 2000), but such tests show cross-reactivity with filaria, schistosomes, hookworm, Ascaris lumbricoides, Trichuris trichiura and Echinococcus (Gam et al. 1987, Conway et al. 1993a,b, Lindo et al. 1994). Western blotting (WB) tests have been applied with more specificity in the immunodiagnosis of strongyloidiasis (Genta et al. 1988, Sato et al. 1990, Conway et al. 1994, Atkins et al. 1999, Uparanukraw et al. 1999). It should be emphasized that the presence of chronic strongyloidiasis infections is often very difficult to rule out. Therefore a positive serologic test in someone parasitologically negative may still be a true positive.

Several components of S. stercoralis filarial larvae are recognized by antibodies in sera of infected humans, including surface and excretory-secretory proteins (Brindley et al. 1988, 1995, McKerrow et al. 1990).
Studies on the antigenic composition of extracts from *S. ratti* and *S. venezuelensis* compared to those of *S. stercoralis* suggest that these rodent species might be used as antigen sources for the immunodiagnosis of human strongyloidiasis instead of *S. stercoralis* antigens (Sato et al. 1995). The purpose of this study was to evaluate the frequency of antigenic components recognized by serum IgG antibodies in WB using *S. ratti* larval antigenic extract for the diagnosis of human strongyloidiasis. In addition, the WB results were compared to the ELISA and the IFAT results.

**MATERIALS AND METHODS**

**Patients and serum samples** - Serum samples of 180 individuals were analyzed, from which 80 were from patients who were shedding *S. stercoralis* larvae only in the feces; 60 from patients with other intestinal parasites (hookworm, 20 cases; *Ascaris lumbricoides*, 7 cases; *Enterobius vermicularis*, 7 cases; *Hymenolepis nana*, 9 cases; *Trichuris trichiura*, 6 cases; *Schistosoma mansoni*, 6 cases; *Giardia lamblia*, 5 cases), all from the Clinical Hospital, Federal University of Uberlândia, state of Minas Gerais, Brazil; and 40 serum samples of University students with parasitological analysis negative in three fecal samples (Baermann 1917, Lutz 1919, and Moraes 1948 methods) and with no history of *Strongyloides* infection. Serum from a patient who presented *S. stercoralis* larvae in the feces was used as positive control and three sera from healthy individuals (negative in three fecal samples) were used as negative controls. This study received approval from the Ethical Committee of the Federal University of Uberlândia.

**Parasites** - *S. ratti* larvae were obtained from the feces of experimentally infected rats (*Rattus rattus*). The fecal samples were mixed with an equal part of finely ground wood charcoal, moistened with water, spread equally on Petri dishes and incubated at 25°C for 5 days. Filariform larvae were then harvested according to the Baermann (1917)-Moraes (1948) method, concentrated by centrifugation for 5 min at 1000 g and stored at -20°C until being processed.

**Antigens** - For WB and ELISA, the larval extracts were obtained from 288,000 filariform larvae, resuspended in 1 ml of phosphate buffered saline (PBS) pH 7.2 containing protease inhibitors (1 mM EGTA, 10 mM EDTA, and 0.05 mM PMSF), and sonicated on ice (8 cycles of 20 s). The larval suspension was then centrifuged at 13,000 g for 30 min at 4°C, and the supernatant (larval extract) was submitted to protein estimation by the Lowry et al. (1951) method. For IFAT, *S. ratti* antigens were prepared according to Costa-Cruz et al. (1997).

**Rabbit immune gammaglobulin** - Immune serum from a rabbit experimentally infected with *S. ratti* filariform larvae on successive inoculations was submitted to salt precipitation (3.12 M ammonium sulfate, pH 6.5) to obtain the gammaglobulin fraction. After incubation for 18 h at 4°C the preparation was centrifuged at 3,000 g for 30 min at 4°C. The pellet was washed three times in 1.56 M ammonium sulfate pH 6.5 and dialyzed for 24 h at 4°C against saline solution. Normal rabbit gammaglobulin was used as negative control.

**Electrophoresis and electrophoretic transfer** - Larval extract of the parasite was diluted (v/v) in sample buffer (6.5 μg of final protein content), boiled for 3 min at 100°C, and submitted to sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) using a 12% resolving gel under denaturant and reducing conditions (Biorad System, US), as described by Laemmli (1970). After SDS-PAGE, the gels were either stained by Coomassie brilliant blue or transferred to nitrocellulose membranes (0.45 μm; Sigma, US), as described by Towbin et al. (1979), using a transfer apparatus (Hoefer System, US).

**Serological assays**

**WB-IgG** - Nitrocellulose strips (4 mm wide) were blocked with 5% non-fat milk in PBS containing 0.05% Tween 20 (PBS-T) for 2 h at room temperature and subsequently incubated overnight at 4°C with rabbit or human serum samples diluted at 1:40 in 1% non-fat milk in PBS-T (PBSTM). After washing in PBSTM, the strips were incubated for 2 h at room temperature with the conjugate (anti-human IgG, whole molecule, labeled with peroxidase, Sigma) diluted at 1:200 in PBSTM. The strips were washed in PBS and developed with hydrogen peroxide and 3,3′ diaminobenzidine tetrahydrochloride (DAB-Sigma) for 3 min. The strips were washed with distilled water and the positive reactions were determined by the visualization of clearly defined brown bands. The apparent molecular weight of such bands were estimated in comparison with the linear regression curve obtained from the electrophoretic relative mobility (Rf) versus the molecular masses of the markers (Rainbow™; Amersham, England): myosin (220 kDa), phosphorylase b (97.4 kDa), bovine serum albumin (66 kDa), ovalbumin (46 kDa), carbonic anhydrase (30 kDa), trypsin inhibitor (21.5 kDa) and lysozyme (14.3 kDa).

**ELISA-IgG** - Preliminary experiments were carried out in order to determine the optimal conditions for ELISA-IgG, through block titration of the reagents (antigen, sera, and conjugate). ELISA was carried out in polystyrene microplates (Interlab, São Paulo, Brazil) and immuno-reagents were assayed in 50 μl/well. After each step of the reaction, microplates were washed three times for 5 min with PBS-T. Microplate coating was performed diluting the larval extract at 0.5 μg/well in 0.06 M carbonate-bicarbonate buffer, pH 9.6 and incubating overnight at 4°C. The microplates were washed and incubated with the serum samples diluted at 1:40 in PBS-T for 45 min at 37°C. The conjugate, goat IgG anti-human IgG (Fc chain) labeled with peroxidase (Sigma), was diluted at 1:2000 in PBS-T and incubated for 45 min at 37°C. The enzymatic substrate consisted of H₂O₂ plus o-phenylenediamine (OPD) solution in 0.1 M citrate-Na₂HPO₄ buffer, pH 5.5. The reaction was stopped after 15 min with 25 μl/well of 1 M H₂SO₄ and the absorbance was determined in an ELISA reader (Metrolab, Argentine) at 490 nm. The cut off was established using the mean absorbance of five non-reactive serum samples plus two standard deviations, as described by Bassi et al. (1991). Reactive serum samples were retested in two-fold serial dilutions until the endpoint titer.

**IFAT-IgG** - The immunofluorescence test using *S. ratti* egg antigens • Luciana Pereira Silva et al.
antigen was carried out according to Costa-Cruz et al. (1997). Briefly, *S. ratti* filariform larvae were embedded in Tissue Tek® (O.C.T. compound, Miles) and cut into 4 µ sections with a cryo-microtome (Cryocut 1800 Reichert-Jung). The sections were put onto extra-thin slides (76 mm x 76 mm) previously cleaned in an alcohol-ether solution, then dried at room temperature and preserved at -18°C up to the time of use. Both negative and positive control sera, as well as the test samples, were diluted 1:20 in PBS and added on the slides containing larval antigens. The positive sera were further retested in 2-fold serial dilutions. After incubation for 30 min at 37°C the slides were washed three times in PBS for 5 min and the fluoresceinated anti-human IgG conjugate (Biolab, Brazil) was added at 1:100 in PBS containing 1% Evans blue as contrast stain. After 30 min incubation at 37°C the slides were washed again as described above, mounted in glycerin (pH 8.5) and examined in an Olympus BH 2 RFC immunofluorescence microscope at 200 and 400X. Results were expressed in titers, which were considered positive when ≥ 20. Reactive serum samples were retested in two-fold serial dilutions until the end-point titer.

Statistical analysis - Statistical analysis was performed using the Statistics for Windows software (Stat soft Inc, 1993). The frequencies of antigenic components recognized by IgG antibodies in WB among the different groups were compared using the analysis between two proportions by Z statistics. Differences were considered statistically significant at level of 5%.

RESULTS

Sera were obtained from 85 men and 95 women. In patients with strongyloidiasis, patients' ages ranged from 4 to 80 years (yr) old (36.9 ± 16.5 yr) and in control groups (patients with other intestinal parasitoses and healthy individuals), patients' ages ranged 1 to 70 years old (21.6 ± 12.8 yr).

Thirty-five antigenic components (apparent molecular weight of 8, 10, 14, 17, 20, 23, 26, 28-35, 37, 40, 44, 46, 48, 52, 55, 60, 66, 70, 75, 78, 81, 85, 89, 90, 91, 97, 105, 113, 117, 122, 126, 132, 138, 141 and 148 kDa) were recognized by IgG antibodies in WB using *S. ratti* larval extract in 180 serum samples. Infection-specific immunodominant antigenic components were defined as the ones recognized by at least 25% of strongyloidiasis patients. Thus, 11 immunodominant antigenic components (10, 14, 17, 20, 26, 28-35, 55, 81, 85, 126 and 138 kDa) were significantly more frequent (p < 0.05) in patients with strongyloidiasis as compared to patients with other intestinal parasitoses and healthy individuals (Fig. 1). A representative WB using *S. ratti* larval extract is illustrated in Fig. 2. The criteria for positivity in WB were established as reaction with two or more of the infection-specific immunodominant antigenic components.

The rabbit immune serum recognized 14 antigenic fractions that were coincident with those recognized by serum samples from patients with strongyloidiasis or other parasitoses and healthy individuals. The rabbit normal serum reacted with the bands of 52 kDa and 148 kDa, similarly to those recognized by serum samples of healthy individuals.

**Fig. 1:** frequency of 11 immunodominant antigenic components of *Strongyloides ratti* (kDa)

**Fig. 2:** Western blotting for the detection of IgG antibodies in serum samples (1:40 dilution). *Strongyloides ratti* larval extract was submitted to SDS-PAGE at 12%. Lanes show serum samples from patients with strongyloidiasis (1 to 9), patients with other intestinal parasitoses (10 to 14), and healthy individuals (15 to 17).

Antibody titers in ELISA ranged from 40 to 320 and IFAT titers ranged from 20 to 2 560.

WB, ELISA and IFAT showed sensitivity of 96%, 100% and 90%, respectively, whereas specificity values were 96%, 98%, and 100%, respectively. Sensitivity was found to be significantly different between WB and ELISA, WB and IFAT, and ELISA and IFAT (p < 0.05). In contrast, specificity showed no statistically significant difference among the three tests (p > 0.05). Comparison between sensitivity and specificity values demonstrated statistically significant differences between the tests (p < 0.05).

Fig. 3 shows the comparison of positivity rates among the three tests (WB, ELISA and IFAT) for the detection of IgG antibodies anti-*S. ratti* in serum samples of patients with strongyloidiasis or other parasitoses and healthy individuals. There was 87.5% positive concordance and 12.5% discordance (9% for WB+/ELISA+/IFAT-, 1% for WB-/ELISA+/IFAT- and 2.5% for WB-/ELISA-/IFAT+) in the group of patients with strongyloidiasis. Sera of patients with other intestinal parasites showed 93%
negative concordance and 7% discordance (3.5% for WB+/ELISA+/IFAT- and 3.5% for WB+/ELISA-/IFAT-). Sera of healthy individuals showed 97.5% negative concordance and 2.5% discordance (WB+/ELISA-/IFAT-).

DISCUSSION

*S. ratti* larval extract was studied for the diagnosis of human strongyloidiasis in three tests: WB, ELISA and IFAT. These tests demonstrated specificity and significance values equal or higher than 90%, showing statistically significant differences only with respect to sensitivity, being ELISA the most sensitive test, followed by WB and IFAT.

*S. ratti* larvae provide convenient antigens for the immunodiagnosis of strongyloidiasis, mainly because the logistics of antigen preparation would be greatly simplified, since keeping rats infected with *S. ratti* would be much easier than maintaining immunosuppressed monkeys with *S. stercoralis* with relatively easy mass production of filariform larvae. Therefore, *S. ratti* larvae can be an adequate substitute antigen of *S. stercoralis* using ELISA and IFAT (Grove & Blair 1981, Sato et al. 1995, Costa-Cruz et al. 1997). The WB test was used for the first time in this study and showed immunodominant antigenic components of *S. ratti* larval extract. However, these results should be considered as a first approach for this issue, since further studies are required to compare the immunodominant antigenic components from the studied species (*S. ratti*) to those from *S. stercoralis*. In addition, the use of purified antigens obtained by detergent extraction should be also tested rather than the crude antigenic extract.

In the present work, 35 antigenic components were recognized in WB by serum samples of patients with strongyloidiasis or other intestinal parasitoses and healthy individuals. From these, 11 immunodominant antigenic components (10, 14, 17, 20, 26, 28-35, 55, 81, 85, 126 and 138 kDa) were predominantly recognized by infected human sera.

In this study, immunodominant *S. ratti* antigenic components with apparent molecular weight of 10, 14, 126 and 138 kDa were reported for the first time in the diagnosis of human strongyloidiasis, and these bands were also recognized by rabbit immune sera. Other investigations have employed *S. stercoralis* larval extract in WB (Genta et al. 1988, Sato et al. 1990, Atkins et al. 1999).

*S. stercoralis* immunodominant antigenic components with apparent molecular weight of 28, 31 and 41 kDa were identified as useful antigens for specific immunodiagnosis of strongyloidiasis in serum samples (Conway et al. 1993b, Lindo et al. 1994, Uparanukraw et al. 1999). In the present study, using *S. ratti* antigen, we also verified the importance of these two first components (28 and 31 kDa) by visualization of a smear of 28-35 kDa, as the most frequent band (65%). However the *S. ratti* larval antigen apparently does not have a 41 kDa, thus evidencing a difference between *S. stercoralis* and *S. ratti* antigen. Accordingly, using *S. cebus* and *S. ratti*, Conway et al. (1994) reported recognition of antigenic bands with apparent molecular masses of 28, 31 and 41 kDa in the *S. stercoralis* extract only.

Several proteins on the surface of *S. stercoralis* filariform larvae, including one of 30 kDa, and several excretory-secretory larval products, including those of 40, 30 and 25 kDa, were identified by Brindley et al. (1988) and recognized by antibodies from infected patients. Similar sized bands were seen in this study, in which the range 28-35 kDa showed the higher positivity. The 25 kDa protein may correspond to the 26 kDa band described here.

Cases with strongyloidiasis showed 87.5% positive concordance in the three tests and 93% and 97.5% negative concordance in the three tests, respectively, for patients with other intestinal parasitoses and healthy individuals. These data reinforce the necessity to perform immunological tests that could confirm the diagnosis and avoid false-positive results. In cases of positive ELISA and negative IFAT, WB could define the diagnosis in eight cases (10%) of patients with strongyloidiasis and two cases (3.5%) of patients with other intestinal parasitoses, suggesting that WB may be a useful confirmatory test.

Cross-reactivity occurred with hookworm (1 case) and *T. trichura* (1 case) for both positive WB and ELISA. However, our region is located more than 2 500 km far from the endemic area for filariasis in our country, the latter situated in the Northeastern region. Thus, data about specificity should be restricted to non-endemic areas of filariasis because of the known cross-reactivity (Dreyer et al. 1996).

In conclusion, WB, ELISA and IFAT showed a high rate of sensitivity and specificity in the detection of serum IgG using *S. ratti* antigen in human strongyloidiasis. In addition, WB was able to recognize 11 immunodominant antigenic components of *S. ratti*, showing to be a useful tool to define the diagnosis in cases of equivocal serology.

REFERENCES


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