Acute schistosomiasis diagnosis: a new tool for the diagnosis of schistosomiasis in a group of travelers recently infected in a new focus of Schistosoma mansoni

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ABSTRACT

Introduction: The diagnosis of schistosomiasis mansoni on early stages of infection is important to prevent late morbidity. A simple, cheap, sensitive and specific assay for routine diagnosis of schistosome infection based on the detection of specific IgG for schistosomula tegument antigens (ELISA-Sm-Teg) was developed by our group. Methods: We describe here an acute outbreak involving a travel group of 80 individuals from a non-endemic area of the State of Minas Gerais, Brazil. These individuals were in contact with a freshwater pool where Biomphalaria glabrata was found. Results obtained from our new methodology were compared to IgG antibody titers against soluble worm antigenic preparation (SWAP) by ELISA and, also to parasitological examination, nuclear magnetic resonance and clinical findings. Results: ELISA-SmTeg was capable of detecting 64 positive cases among the 80 individuals participating at the survey with a positivity ratio of 80% and a higher sensitivity than ELISA-SWAP that was only sensitive for 56% of positive cases. Besides, a significant correlation was found for the severity of the infection and the specific IgG titers against SmTeg. Conclusions: Our data showed that ELISA-SmTeg might serve as the initial diagnostic tool for acute stages of the infection in community-based helminth control programs or for the surveillance of individuals from non-endemic areas.

Keywords: Schistosomiasis mansoni. Immunological diagnosis. Schistosomula antigens. Acute outbreak.

INTRODUCTION

Schistosomiasis continues to be a major worldwide public health problem that affects 200 million people and about 779 million people live in endemic areas in the Middle East, South America, Caribbean, Southeast Asia and particularly sub-Saharan Africa. Nonetheless, the disease is seen in growing numbers in recently detected foci of transmission due to increased immigration from endemic areas and tourism. Severe sequelae due to schistosomiasis are rare in travelers from non-endemic areas because the risk of developing significant pathological manifestations after short-term exposure that frequently result in high parasite burden is limited by the life span of the adult worm. The most important factor regarding the difficulty of controlling the spread of the disease is the absence of diagnostic methods capable of detecting the disease, especially in the early patent phase.

Clinical findings make possible the division of the disease in two phases. First is the initial phase, which includes cercarial dermatitis caused by the penetration of cercariae into the skin and the local presence of schistosomula. Together, the acute symptomatic schistosomiasis that may appear 3 to 7 weeks after an exposure and is characterized by cough, fever, anorexia, abdominal pain and headaches. In this phase, the inflammatory reaction is well established and involves predominantly Th1 bias, when immunoglobulin levels are elevated in the serum and hyper eosinophilia. The late acute phase that occurs after 40 to 60 days post-infection is the acute phase, characterized by the egg laying and a predominantly Th2 inflammatory response. Its clinical manifestations vary depending on the parasite load and host immune response. Patients may present gastrointestinal, hepatosplenic, schistosomal myeloradiculopathy and acute forms with the latter being a main indicator of schistosomiasis severity.

Although treatment with oxamniquine and artemisinin on the first days after infection is considered more efficient in obtaining cure than praziquantel, the lack of methods to detect...
the initial phase, before egg laying, precludes the patients from a successful therapy. The diagnosis is first guided by the patient history of water contact in an endemic area. This evidence is further confirmed by the presence of the Schistosoma mansoni eggs in stool samples after egg laying, on oviposition phase\(^6\). Even during early patent phase, the absence of S. mansoni eggs in the faeces does not rule out the diagnosis due to the low sensitivity this methodology may present. There is usually a military distribution of eggs in the organs of the host and laparoscopy frequently reveals whitish nodules, as granulomas, on the surface of liver, intestines and visceral peritoneum. Hence, it is expected that the parasitological methods do not have properly efficiency for the diagnosis, predominantly for patients in acute phase once the female has not yet laid eggs\(^7\).

Serological techniques have proven advantages in diagnosis of schistosomiasis\(^8\) since the humoral response especially related to IgG antibodies in acute patients to egg and worm antigens does not differ from the chronic phase as a significant levels of antibodies are detected, but they are not yet part of the routine of control programs\(^9\).

This work focus on the evaluation of an indirect enzyme linked immunosorbent assay using schistosomula tegument antigen (ELISA-SmTeg). This methodology was first standardized using 7 to 10 days post-infection mice sera when it showed to be a promising diagnostic tool for the acute phase of the disease. In this report, we describe a case involving a group of travelers in Brazil that was exposed to a contaminated freshwater pool. Early detection of the clinical symptoms or signs of schistosomiasis mansoni combined with rapid investigation of the entire group enabled us to observe this group prospectively. The main goal of this work is to offer a new reliable diagnostic tool to determine positive cases before egg laying.

**METHODS**

**Study population**

Eighty individuals were part of a group that was hosted on a country house in Colônia do Teodoro, a recently endemic focus for schistosomiasis mansoni, next to the City of São João Del Rei in the State of Minas Gerais, in southeast Brazil, from December 2009 to February 2010. The individuals were exposed to a freshwater pool where Biomphalaria glabrata snails were found.

**Identifying cases**

To identify symptomatic cases, we initiated the interviews with exposed travelers on March 2010 when samples were collected (1 to 3 months post-infection). Follow-up focused on the detection of symptoms or signs of chronic schistosomiasis previously described in travelers, together with current symptoms and signs, including pulmonary symptoms, gastrointestinal symptoms (e.g. abdominal pain or diarrhea), constitutional symptoms (e.g. fatigue or weight loss), and myeloradiculopathy involvement.

Schistosoma mansoni infection was defined as exposure to the freshwater pool plus one of the following criteria: presence of eggs in faecal samples, IgG antibody titers by ELISA using adult worm antigens, myeloradiculopathy detected by nuclear magnetic resonance, and/or symptoms compatible with acute schistosomiasis. Symptoms defining infection in this group were fever, cough, cerarial dermatitis, and angioedema. Non-specific symptoms (e.g. fatigue, gastrointestinal complaints, and headache) were reported as well. Blood samples were collected by venipuncture of all those patients. Individual serum samples were obtained after centrifugation of blood samples at 3,000g for 5min. These samples were maintained at -20°C. Twenty four out of 80 individuals agreed to submit faeces for examination and these samples were firstly analyzed by Kato-Katz parasitological assay\(^10\). Three glass slides (41.7mg/ smear) of a single fecal sample were examined. Five patients were hospitalized, one with advanced pulmonary stage, one with schistosomal myeloradiculopathy and, three with severe dehydration caused by the hepatointestinal form.

Positive individuals for one of the described criteria were treated with a single oral dose of praziquantel (50mg/kg), according to the recommendation of the Brazilian Ministry of Health.

**ELISA-SWAP**

Enzyme-linked immunosorbent assay with adult soluble worm antigenic preparation (ELISA-SWAP) was performed in microtiter plates MaxiSorp Surface (NUNC, Denmark) sensitized with 100μl/well of 15g/ml of SWAP diluted in buffer 0.05M carbonate-bicarbonate pH 9.6. The incubation was done for 16h at 4°C. The plates were washed three times with 0.15M phosphate buffer saline pH 7.2 with 0.05% of Tween 20 (LCG Biotechnologia, BR) (washing buffer) and, the non-specific sites were blocked with 10% fetal bovine serum in washing buffer at 37°C for 1h. After new washing steps, 100μl of human serum samples diluted 1:50 in PBS were added in triplicate into each well and the plates were incubated at room temperature for 1h. Following, the plates were submitted to washing steps and incubated at room temperature for 1h with conjugated anti-IgG human peroxidase (Southern Biotech, USA) diluted 1:60,000 in washing buffer. The plates were washed again 100μl of substrate 3,3’5,5’-tetramethylbenzidine solution (TMB-H2O2) (Invitrogen, USA) were added to each well. The reaction was stopped after 20min of incubation in the dark by addition of 50μl/well of 2N sulfuric acid. The results were obtained as absorbance values at 450nm in microplate reader (Bio-Rad Laboratories 3550, JA). The cut off value of the ELISA-SWAP was OD=0.188 (standard deviation of 0.08), determined by ROC curve (A=0.70, p=0.0001).

**Groups of study**

The 80 individuals were divided in four groups according to diagnosis results. Group 1 were limited to patients with symptoms and/or signs of acute schistosomiasis but negative for any other diagnostic method used (total of 28 individuals) whereas, group 2, with 26 individuals, involved patients that were positive for clinical examination plus ELISA-SWAP. Patients with positive results for these two last criteria that additionally presented
in stools constituted group 3, with a total of 21 individuals. Finally, all the five hospitalized patients with severe schistosomiasis were represented by group 4.

**Evaluation of the indirect enzyme-linked immunosorbent assay using schistosoma tegument antigen (ELISA-SmTeg)**

Serological studies were conducted at the Brazilian Excellence Center for the Diagnosis of Schistosomiasis mansoni, Oswaldo Cruz Foundation (Brazilian Ministry of Health) in Belo Horizonte. The serologic test performed is based on the detection of specific immunoglobulin G (IgG) against schistosoma tegument antigen by ELISA methodology.

**Preparation of Schistosoma mansoni schistosoma tegument antigen:** cercariae of the LE strain were obtained at the Laboratory of Malacology of the René Rachou Research Center, Oswaldo Cruz Foundation and were mechanically transformed into schistosomula. Briefly, cercariae were placed into conical tubes and left in ice bath for 30 min before centrifugation (Eppendorf Centrifuge 5415R, Hamburg, GR) at 200 G for 3 min at 4°C. The pellet was resuspended in cold Ear’s salts plus lactalbumin hydrolyzate medium. The cercarial tails were broken in vortex (Scientific Industries Genie-2, New York, USA) for 2 min. Later the tails were removed from the medium through repeated washing steps with Ear’s salts media at 37°C, and schistosomula were incubated for 90 min at 37°C. This step was followed by centrifugation at 200 G for 1 min. For **t**eggum removal, 2 ml of 0.3 M calcium chloride were added to the schistosomula that was stirred in vortex for 5 min and centrifuged at 200 G for 1 min. The supernatant was centrifuged at 50,000 G for an hour and the pellet enriched of membrane was resuspended in 0.9% saline and dialyzed against 1.7% saline solution for 24 h. Protein concentration was determined by Bradford method. The final concentration used in standardized tests was 0.52 mg/ml.

**Indirect enzyme-linked immunosorbent assay:** microtiter plates Maxisorp surface were coated with 100 µl per well of SmTeg (1 µg/ml) in buffer 0.05 M carbonate-bicarbonate pH 9.6 for 16 h at 4°C. After plates were washed, the free sites were saturated with 300 µl per well of 2.5% skim milk diluted in washing buffer, incubating at 37°C for 1 h and 100 µl of individual serum sample (diluted 1:100) in 0.15 M phosphate buffer saline pH 7.2 were added to the plates, followed by incubation for 1 h. The plates were submitted to washing steps and incubated for 1 h with anti-IgG conjugated to peroxidase diluted in washing buffer (anti-human IgG Fc specific peroxidase) at the dilution of 1:600. Then, 100 µl of substrate solution were added to each well and the enzymatic reaction was stopped after 10 min by adding 50 µl per well of 2N sulfuric acid. Results were obtained as absorbance values at 450 nm. The cut off value of the ELISA-SmTeg was OD = 0.110 (standard deviation of 0.02), determined by ROC curve (A = 0.92, p < 0.0001).

Negative controls were properly assayed using fifty three volunteers' serum samples as our control group, also wells without antigen and serum samples as control of nonspecific adsorption of conjugate. The standard dilution of sera and conjugated antibodies was determined by a dilution curve, performed with the same reagents and equipment based on six different dilutions.

**Statistical analysis:** data deriving from absorbance values were analyzed with Minitab software (Minitab Inc, College, USA) by Kolmogorov-Smirnov normality test. Normal distributed data were analyzed by Student’s t test and comparisons between methods were done by Chi-square analysis (p < 0.05 as significance level). The sensitivity, specificity, cut off values, receiver operating characteristic (ROC) curve and positive ratios, defined as the capability of detecting real positive and negative individuals, were determined with Prism 5.0 software.

**Ethical considerations**

The Ethical Committee of **Fundação Nacional de Saúde** in Minas Gerais, Brazil granted ethical approval for this study. Written informed consent was obtained from each individual.

**RESULTS**

All the 80 individuals hosted on the country house that had contact with the freshwater pool participated in this survey. Serum samples were submitted to a well established indirect ELISA-SWAP assay that allowed us to identify the levels of specific IgG of each individual patient and a correlation was done with the new assay ELISA-SmTeg, which has been previously standardized in our laboratory for the determination of specific IgG antibody levels against schistosoma tegument antigen.

The cut off value for this new assay had been previously determined (OD = 0.110) using fifty three volunteers' serum samples as our control group by the ROC curve (A = 0.92, p < 0.0001), as shown on Figure 1. It is important to notice that the high sensitivity described was determined with clinical acute phase patients and the same effectiveness was not seen with the chronic form patients (data not shown).

Using the defined cut off value, samples from the 80 individuals were applied on ELISA-SmTeg assay and a first comparison was done with data obtained from another indirect immunoassay usually used as the routine method, the ELISA-SWAP. Table 1 shows the comparison between results obtained from both ELISA assays. Among the 80 positive patients, only 45 individuals were properly diagnosed by ELISA-SWAP. On the other hand, ELISA-SmTeg showed a higher sensitivity by being capable of identifying 64 positive individuals (p = 0.001). The positivity ratio of ELISA-SmTeg assay of 80% was superior to 56% obtained by ELISA-SWAP. The Cohen’s Kappa Index of 0.385 (standard deviation of ± 0.094) indicates fair agreement between both immunological assays.

Fecal samples were obtained in 24/80 (30%) persons, and eggs were detected in 24/24 (100%) fecal samples by Kato-Katz. The patients, additionally, demonstrated high levels of specific IgG antibodies for ELISA-SmTeg with an average of 0.401 (standard deviation of ± 0.016). Three (13%) positive patients
with eggs in stools were not detected by ELISA-SWAP showing an OD average of 0.157 (± 0.026).

Patients with clinical symptoms and/or signs that were properly diagnosed by both methods, Kato-Katz and ELISA-SWAP (21/80) (26%), showed also high levels of IgG against SmTeg (21/21) (100%). The IgG levels for these patients (26/80) were significantly higher (33%) than from patients whom were positive only for ELISA-SWAP, presenting no eggs in stools (p=0.002). Figure 2 shows this relation after patients were divided in four individual groups. We could see that patients that was diagnosed as negative by ELISA-SWAP, but presented all the clinical symptoms compatible with acute schistosomiasis (28/80) (35%), also presented lower IgG titers for SmTeg in comparison to group 2 (p=0.002). The analysis of the patients with severe schistosomiasis (group 4), one with advanced pulmonary stage, one with schistosomal myelodysplasia and, three with severe intestinal form, allows us to reaffirm that ELISA-SmTeg presented once more a high positivity ratio by being capable of diagnosing all the severe cases (5/80) (6%) and, additionally presented a correlation between specific IgG levels and the severity of the infection. Statistical differences were found when comparisons were made for the antibody level of patients on groups 1 and 2 in comparison to group 4 (p=0.003 and 0.001, respectively).

Table 1 - Diagnoses results of enzyme-linked immunosorbent assay with adult worm antigens and with schistosomula filar antigen of serum samples from the 80 travelers positive for schistosomiasis mansoni.

<table>
<thead>
<tr>
<th>TABLE 1</th>
<th>Diagnoses results of enzyme-linked immunosorbent assay with adult worm antigens and with schistosomula filar antigen of serum samples from the 80 travelers positive for schistosomiasis mansoni.</th>
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<tbody>
<tr>
<td></td>
<td>ELISA-SWAP</td>
</tr>
<tr>
<td>Positive</td>
<td>45</td>
</tr>
<tr>
<td>Negative</td>
<td>35</td>
</tr>
<tr>
<td>Total</td>
<td>80</td>
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ELISA-SWAP: ELISA soluble worm antigenic preparation; ELISA-SmTeg: ELISA schistosomula filar antigen. Statistical differences between the number of positive individuals by both assays is represented by (***)(p=0.001) after Chi-square analysis. Artwork created by Prism 5.0 software.

FIGURE 2 - Relation between the number of travelers diagnosed by ELISA-SmTeg and groups divided according to the additional diagnostic results and severity of infection. Group 1: 28 patients positive only for clinical examination; Group 2: 26 patients positive for clinical examination plus ELISA-SWAP; Group 3: 21 patients positive for clinical examination, ELISA-SWAP and Kato-Katz; Group 4: 5 hospitalized patients with severe schistosomiasis. Line represents the cut off of OD=0.110 for ELISA-SmTeg. Results represent the mean of four absorbance values at 450nm. Statistical differences between groups are represented by (***)(p<0.001) after Student t test, for p<0.002 and 0.003, respectively for groups 1 and 2 and, groups 1 and 4; p=0.002 and 0.001, for groups 2 and 3 and, groups 2 and 4. Artwork created by Prism 5.0 software. ELISA-SmTeg: enzyme-linked immunosorbent assay-schistosoma filar antigen; ELISA-SWAP: enzyme-linked immunosorbent assay-schistosoma filar antigenic preparation; OD: optical density.

DISCUSSION

Acute schistosomiasis is one of the clinical manifestations of infection with Schistosoma spp.2,3. The pre-postural phase occurs between cercarial penetration in skin and egg laying by female adult worms after the migration of young parasite to portal system and the maturation of male and female worms (7 to 8 weeks). Oviposition, which defines this postural phase of the disease, typically occurs 45 to 60 days after cercarial penetration. This symptomatic acute phase is mostly severe in non-endemic (primary infection) individuals (i.e. tourists, travelers) exposed to fresh water in endemic areas2,4.
It is considered to be a toxemic and allergic reaction to the migrating and maturing schistosomula. The severity of the clinical presentation varies according to the worm burden and the immune response to the released parasite antigens.

Circulating immune complexes are found in 55-93% of patients with acute schistosomiasis, and their presence and amount are correlated to the intensity and severity of symptoms. These symptoms are typically seen before oviposition, and the appearance of granulomatous reactions around eggs. Oviposition begins at the end of adult maturation and migration to the vesical plexus or mesenteric veins. In the early stages of the acute phase (until 40 days of infection), a search for eggs in stools will be negative. Nonetheless, eggs may still be detected in stools of patients complaining of symptoms compatible with clinical acute form (45-50 days post-infection), although the Kato-Katz technique is rather insensitive in light infections.

Serological testing is an important diagnostic tool to be considered in acute schistosomiasis, especially for the diagnosis before egg laying. Still, it is crucial to select an antigenic portion of the helminth in order to solve the problem of variability on the sensitivity that may occur according to the antigen used. Even so, the primary investigation must be done by a physician in order to determine the medical history of each individual case and then, serological findings ought to be used as a confirmatory methodology. Among the immunological assays available, ELISA is the most commonly used, especially with egg or worm antigens. Its methodology sensitivity can be high for patients with chronic form, reaching values of 89-96% for egg antigens and 90-94% for worm antigens, which is related to the cut off titre defining positivity. It is also possible to use a combination of serological tests and/or antigens, such as SEA and SWAP, to simultaneously establish the diagnosis of schistosomiasis and discriminate the acute from the chronic forms of the disease and to increase sensitivity. Nonetheless, early treatment with praziquantel, although partially active for immature schistosome stages, should be ready for exposed travelers after diagnosis.

Reports of outbreaks in travelers with a single and recent exposure have contributed to a better description of the natural history of schistosomiasis and, particularly in those cases, diagnosis relies mainly on positive serological testing. We report here a recent focus of schistosomiasis mansoni involving 80 individuals that had contact with a contaminated freshwater pool. As a consequence, all the individuals presented clinical findings compatible with acute schistosomiasis and were further investigated for IgG antibody levels by ELISA-SWP. Although the Kato-Katz analysis was not performed for all the 80 patients, the sensitivity of ELISA-SWP was low when it is being used as a tool for the immunodiagnosis of patients with acute form.

Thus, based on the statement that difficulties associated with clinical or parasitological diagnosis in early patent phase might be overcome by adoption of immunological methods, we developed a new indirect ELISA methodology using schistosomula antigen based on the helminth life-cycle. Data from ELISA-SmTeg showed a significant positivity ratio when 64/80 patients presented high levels of specific IgG, superior to ELISA-SWP results (45/80), that is the most used immunological method. Positivity ratio was 56% and 80%, respectively for ELISA-SWP and ELISA-SmTeg. This final analysis corroborates with our first findings showing that ELISA-SmTeg is more sensitive for the differential diagnosis of patients with acute schistosomiasis. There is generally a trade-off between sensitivity and specificity, and the relative performance of a new diagnostic test is therefore calculated as an index in which both these parameters are accounted for. Sensitivity, however, has more relevance in a public health context, especially in situations where the goal is the complete elimination of a new focus.

When comparing the Kato-Katz parasitological result of each individual patient to the specific IgG antibody level for ELISA-SmTeg, once again, the effectiveness of our new immunoassay was confirmed when the assay had properly diagnosed all these 24 positive patients. On the other hand, ELISA-SWP did not achieve the same effectiveness after 3 patients with eggs in stools were not differently diagnosed showing low IgG titers.

When patients were divided into groups based on the results obtained by different diagnostic methodologies and on the severity of infection, it could be seen that the IgG titers against SmTeg antigen significantly increase as the positivity ratio became higher. Individuals that were positive only for the clinical examination (group 1) showed significant lower levels of IgG antibody than those that were also positive for ELISA-SWP assay (group 2). Accordingly, patients that were positive for Kato-Katz other than clinical examination and ELISA-SWP (group 3) demonstrated significantly higher IgG titers in sem than group 1 and group 2. No doubt that including clinical findings as a key to look closely for evidence of infection is important in patient management, as eosinophil count. But correlating immunodiagnostic results and antibody levels with clinical data should allow us improve our practice in rural endemic areas, which in turn would significantly enhance the efficiency of the anti-schistosomiasis control program.

Finally, five hospitalized patients with the severe form of the infection were positive for the presence of eggs in stool and, once more, presented high levels of specific IgG determined by ELISA-SmTeg corroborating with the initial findings that shows a significant sensitivity for this new immunoassay. A positivity ratio of 80% was observed for ELISA-SmTeg and a correlation was confirmed when comparing IgG levels against SmTeg and the severity of the infection. Furthermore, the low sensitivity for ELISA-SWP was again reported when the assay missed the diagnosis of two patients with the severe form of the infection.

While antibody-based methods suffer from low sensitivity, especially for the diagnosis of acute schistosomiasis, the ELISA-SmTeg was noticed here as an important tool to confirm real recently infected individuals during the acute phase. This new tool was superior than the usual ELISA-SWP assay, being also able to be used as a tool for measuring the severity of the infection. This new diagnostic method should be especially considered for groups from non-endemic areas.
such as tourism and social/official activities, including military visits on endemic areas. Hence, an immunoassay capable of diagnosing individuals with acute form, especially before the egg laying will allow the increase on the control programme’s effectiveness. Since ELISA-Sinteg and ELISA-SWAP do not correlate perfectly, these antigens could be used in parallel to further increase sensitivity in a given population.

CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

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