Tandem repeat recombinant proteins as potential antigens for the sero-diagnosis of *Schistosoma mansoni* infection

Yombo Dan Justin Kalenda a,b, Kentaro Kata a, Yasuyuki Goto c, Yoshito Fujii d, Shinjiro Hamano a,e,*

a Department of Parasitology, Institute of Tropical Medicine (NEKKEN), Nagasaki University, 1-12-4 Sakamoto, Nagasaki 852-8523, Japan
b Graduate School of Biomedical Sciences, Nagasaki University, 1-12-4 Sakamoto, Nagasaki 852-8523, Japan
c Laboratory of Molecular Immunology, Department of Animal Resource Sciences, Graduate School of Agricultural and Life Sciences, The University of Tokyo, 1-1-1 Yayoi, Bunkyo-ku, Tokyo 113-8657, Japan
d Department of Eco-epidemiology, Institute of Tropical Medicine, Nagasaki University, 1-12-4 Sakamoto, Nagasaki 852-8523, Japan
e Nagasaki University Nairobi Research Station, NUITM-KEMRI Project, Nairobi, Kenya

**Abstract**

The diagnosis of schistosomiasis infection, followed by effective treatment and/or mass drug administration, is crucial to reduce the disease burden. Suitable diagnostic tests and field-applicable tools are required to sustain schistosomiasis control programs. We therefore assessed the potential of tandem repeat (TR) proteins for sero-diagnosis of *Schistosoma mansoni* infection using an experimental mouse model. TR genes in the genome of *S. mansoni* were searched in silico and 7 candidates, named SmTR1, 3, 8, 9, 10, 11 and 15, were selected. Total RNA was extracted from *S. mansoni* adult worms and eggs. Target TR genes were amplified, cloned, and the proteins were expressed in *Escherichia coli* competent cells. Female BALB/c mice were infected with 100 *S. mansoni* cercariae and sera were collected each week post-infection for 18 weeks. The levels of IgG antibodies to SmTR antigens were compared to those to soluble egg antigen (SEA) and to soluble worm antigen preparation (SWAP). Sera of infected mice reacted to all the antigens whereas those of naïve mice did not. IgG responses to SmTR1, 3, 9, 10 and 15 dramatically decreased 4 weeks after treatment with praziquantel, while those against SEA and SWAP remained elevated. Our study suggests that TR proteins, especially SmTR10, may be suitable antigens for sero-diagnosis of infection by *S. mansoni* and are potential markers for monitoring and surveillance of schistosomiasis, including re-infection after treatment with praziquantel.

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1. Introduction

Schistosomiasis remains a public health problem as one of the major neglected tropical diseases. This helminthic disease is distributed worldwide and mainly affects resource-limited regions, with 74 countries reported to be endemic [1]. Over 230 million people are infected globally, with even more at risk of infection and complications [2]. Schistosomiasis is caused by the trematode, *Schistosoma* spp. In Africa, both *Schistosoma mansoni* (*S. mansoni*) and *Schistosoma haematobium* commonly co-exist, while the less-studied *Schistosoma intercalatum* is found in localized regions of central Africa. The Americas are affected by *S. mansoni* alone, while Asian schistosomiasis is caused by *Schistosoma japonicum* and *Schistosoma mekongi* [3].

The pathogenesis of the disease is related to eggs that are produced in small vessels by adult female worms that have infected the body. In intestinal schistosomiasis, the eggs are trapped in tissues such as liver and intestines, whereas in uro-genital schistosomiasis (caused by *S. haematobium*), the eggs are trapped in the urinary bladder. These eggs are responsible for the formation of granuloma and subsequent associated complications such as hepatic fibrosis and hepatocellular carcinoma, chronic anemia, and stunted growth of chronically infected patients [3–5]. Early detection and treatment of the infection are crucial in preventing severe morbidities [6] and reducing the economic burden of the disease due to consecutive complications among affected populations [7,8]. Current schistosomiasis control and elimination programs rely mainly on mass drug administration (MDA) using praziquantel [9]. This isoquinoline derivative has significantly contributed to reducing both the morbidity and prevalence of schistosomiasis over the past several decades. However, praziquantel is ineffective on parasites in the juvenile stage [10,11]. Furthermore,
reinfection by schistosomes is frequently observed in endemic areas, thus requiring repeated MDA activities in order to interrupt transmission and improve morbidity [12].

The commonly used standard assay for diagnosis of *S. mansoni* infection is the Kato Katz (KK) test [13] and involves the quantitative microscopic observation of parasite eggs in stool samples. It is currently acknowledged that the KK test lacks sensitivity especially in low endemic areas [14]. The search for a reliable tool for monitoring and surveillance of schistosomiasis has led to various antigens being investigated for sero-diagnostic tests [15]: schistosomula tegumental antigens [16,17], soluble egg antigens (SEA) [18], soluble worm antigen preparation (SWAP), and cercarial transformation fluid [19,20]. Tests based on the detection of specific antibodies against these antigens provide variable advantages and drawbacks that limit their applicability [21].

The detection of circulating cathodic antigen (CCA) in urine may be an alternative for monitoring and surveillance [22–24] and to date has provided consistent advantages in both lab and field studies. Although some field studies have reported a decreased sensitivity of the CCA test in low infection settings [25,26], it remains the alternative of choice to the KK test [22]. However, with the increasing widespread use of MDA and ongoing control programs, the diagnosis of schistosomiasis in low-endemic areas requires a test with high sensitivity and appropri-ate specificity. An immunodiagnostic test based on antibody detection has been shown to increase the detection efficiency of schistosomiasis especially in low-endemicity areas [27], and several countries such as China, Venezuela, and Brazil have introduced the antibody detection-based test in their schistosomiasis elimination programs [28]. Although the differentiation of active from past infections has been challenging, the detection of specific antibody against SEA or SWAP has been used in laboratory and field settings to diagnose schistosome infection [29,30]. Immunodiagnostic tests provide increased detection efficiency of schistosome infection in low-endemicity areas and among patients from non-endemic areas [28]. Recombinant antigens have been used as alternatives to SEA and SWAP in order to increase the specificity of the test and overcome the cross-reactions that occur when crude antigens are used; importantly, recombinant antigens are amenable to large-scale production. Thus, it is important to find suitable antigens for a sero-epidemiological approach to diagnose schistosomiasis infection [15,21,31].

Completion of the whole genome sequencing of *S. mansoni* [32] and the availability of increasingly sophisticated bio-informatics tools have provided potential resources in schistosome research for further screening of promising antigen targets useful either for diagnosis [33,34] or vaccination [35].

Tandem repeat (TR) genes have been reported to code polypeptides recognized by B cell receptors and have been studied in different parasites, including *Plasmodium* [36,37], *Leishmania* [38], *Trypanosoma* [39,40] and *S. japonicum* [41]. TR genes in *S. mansoni* have been studied throughout the life cycle of the parasite as modulators of gene transcription [42]. The TR non-coding genes Sm1-7 and Dra-1 have been used as markers of *S. mansoni* and *S. haematobium* infection, respectively, in intermediate host snails [43,44]. Sm1-7 is also used to diagnose *S. mansoni* human infections by detecting this highly repeated gene in biological samples such as stool, blood or urine [25,45,46]. The application of these techniques, however, requires the use of sophisticated tools such Real Time polymerase chain reaction (RT-PCR) or PCR equipment, restricting their field study application in endemic areas often found in resource-limited countries.

TR antigens have been used to diagnose *S. japonicum* infection with interesting outcomes [41]. Here, we investigated the potential of TR antigens in a diagnostic test for *S. mansoni* infection in mice by developing an Enzyme-Linked Immunosorbent Assay (ELISA) to detect reactive immunoglobulin G (IgG) in the sera of infected mice.

2. Materials and methods

2.1. Parasite

A Puerto Rican strain of *S. mansoni* was used. The life cycle of this parasite is maintained in the animal facilities of Nagasaki University by passage through *Biomphalaria glabrata* snails and ICR mice or jirds. ICR mice were percutaneously infected with 250 cercariae, sacrificed 7 weeks after infection, and adult worms were collected from the portal vein by intra-cardiac perfusion with phosphate buffer saline (PBS). Eggs were isolated from the liver of infected mice as previously described [47,48].

2.2. Animals, infection and treatment

Six week old female BALB/c mice were purchased from SLC (Shizuoka, Japan) and maintained in the animal facilities of Nagasaki University. These mice were kept in environmentally controlled, specific pathogen-free conditions, with free access to food and water. Animal housing, handling and feeding were done in compliance with the recommendations of Nagasaki University. All the experiments were approved by the ethical committee of Nagasaki University and were conducted following the animal facility guidelines.

Mice were percutaneously infected with 100 *S. mansoni* cercariae in the inguinal area. At each time point, 6 mice were sacrificed and the liver and intestine were analyzed for the presence of eggs; adult worms were also collected from the portal vein by perfusion, starting 5 weeks post-infection.

A group of 6 mice were treated twice with praziquantel (Sigma, St. Louis, MO) at 300 mg/kg body weight 11 weeks post-infection with a 2 day interval [49]. The drug was dissolved in distilled water and 200 μl of the solution was administrated by gavage; the control group of 6 mice received the equivalent volume of distilled water. One week post-treatment, 2 mice were sacrificed to confirm the absence of adult worms by portal vein perfusion, allowing verification of the efficacy of the treatment.

2.3. Sera

Blood was collected from mice 3 and 5 weeks post-infection, and also weekly from 5 weeks post-infection under anesthesia by per-oral puncture. Mice were sacrificed to confirm infection through the presence of worms and liver lesions. At each time point, 6 mice were sacrificed and submitted to perfusion of the hepatic portal system, as described elsewhere [47]. Sera were collected and aliquots were stored at −30 °C until use. Sera collected from 6 uninfected mice were used as a negative control.

Sera were also collected from the treated group and their matched controls without sacrificing the mice at 11 weeks post-infection, before treatment, and 1, 5 and 7 weeks post-treatment (corresponding to 12, 16 and 18 weeks post-infection, respectively).

2.4. Preparation of SWAP and SEA crude proteins

2.4.1. Soluble worm antigen preparation (SWAP)

Adult worms were harvested from mice infected with *S. mansoni* cercariae 7 weeks post-infection by portal vein perfusion; the worms were washed 3 times with PBS, homogenized using a mechanical grinder, and centrifuged at 10,000 × g for 1 h at 4 °C. The supernatant was collected and filtered through a 0.22 μm mesh. The protein concentration was assessed using the Bradford method (Bio-Rad Laboratory, Hercules, CA) and then the protein was aliquoted and stored at −30 °C.

2.4.2. *S. mansoni* soluble egg antigen (SEA)

Soluble crude egg antigen (SEA) was prepared as previously reported [48]. Briefly, purified eggs from the livers of infected ICR mice were re-
suspended in 5 ml ice-cold PBS at a concentration of 100,000 eggs/ml. The suspended eggs were sonicated for 15 min on ice to prevent heating until more than 95% of the eggs were destroyed. The sonicated eggs were centrifuged at 2000 × g for 20 min at 4 °C. The supernatant was collected in a sterile tube, then ultracentrifuged at 100,000 × g for 1 h at 4 °C. The supernatant was collected and the protein concentration was determined as for SWAP. SEA was aliquoted and stored at −30 °C until use.

2.5. Cloning and expression of recombinant SmTR proteins

S. mansoni total RNA was extracted from adult worms and eggs using the TriZol (Invitrogen, Madison, WI) protocol. Total RNAs were reverse transcribed into a single stranded cDNA using Takara PrimeScript RT reagent kit (Perfect Real Time, Takara Bio Inc., Otsu, Japan) following the manufacturer’s protocol. Sequences encoding the TR domains of the selected genes were PCR-amplified using specific primers containing restriction enzyme ligation sites at both ends: BamHI at the 5′ end and HindIII at the 3′ end of the antigens, except for SmTR10, in which XhoI was inserted instead at the 3′ end. A stop codon was added on the 3′ end of each target gene sequence (Table 1). TR genes were amplified using Hot start Ex Taq DNA polymerase (Takara). Using cDNA from both adult worms and eggs as PCR templates, the amplified TR gene domains were separated on 1.5% agarose gel, the positive band corresponding to one or two periods of the tandem repeat were cut from the gel and purified using a QIAquick gel extraction kit (Qiagen, Hombrechikon, Switzerland). The purified PCR products were cloned into pGEM-T Easy vector (Promega, Madison, WI) and sequenced using an ABI 3730 sequencer (AB Applied Biosystems, Tokyo, USA).

Table 2
Top 20 tandem repeat genes within the whole S. mansoni genome.

<table>
<thead>
<tr>
<th>Score</th>
<th>Gene ID</th>
<th>Gene product</th>
<th>Length (bp)</th>
<th>TR period size</th>
<th>B cell epitope</th>
<th>TR protein size (kDa)</th>
<th>TR copy number</th>
<th>Identity</th>
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<tr>
<td>7990</td>
<td>Smp_187350.1</td>
<td>Hypothetical protein</td>
<td>5634</td>
<td>795</td>
<td>2</td>
<td>33</td>
<td>5</td>
<td>79% S.j</td>
</tr>
<tr>
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<td>Hypothetical protein</td>
<td>8235</td>
<td>1122</td>
<td>13</td>
<td>3.5</td>
<td>39% S.j</td>
<td>SmpTR1</td>
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<td>Serine-rich repeat protein, putative</td>
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<td>429</td>
<td>4</td>
<td>19</td>
<td>6.8</td>
<td>no</td>
</tr>
<tr>
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<td>4392</td>
<td>1023</td>
<td>9</td>
<td>2.7</td>
<td>80% S.j</td>
<td>SmpTR4</td>
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<td>1023</td>
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<td>2.3</td>
<td>SmpTR5</td>
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<td>2</td>
<td>SmpTR6</td>
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<td>Smp_094710.1</td>
<td>Alstrom syndrome protein</td>
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<td>1</td>
<td>21.8</td>
<td>no</td>
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<td>76% S.j, S.h</td>
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<td>Ubiquitin (ribosomal protein 140), putative</td>
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<td>1</td>
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<td>100% S.j</td>
<td></td>
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<td>1873</td>
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<td>Major tegumental antigens Sm15</td>
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<td>13.2</td>
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<td>17</td>
<td>2</td>
<td>80% S.J</td>
</tr>
<tr>
<td>1353</td>
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<td>Translation initiation factor IF-2, putative</td>
<td>792</td>
<td>8</td>
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<td>85.1</td>
<td>no</td>
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<td>SmpTR17</td>
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<td>201</td>
<td>1</td>
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<td>68% S.J</td>
<td></td>
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<td>1</td>
<td>4.3</td>
<td>no</td>
<td>SmpTR20</td>
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Length of the gene in bp, the size of one TR sequence within the gene (bp), number of B cell epitopes within one TR period, Identity: shared homology with S. japonicum; S.h, S. haematobium; and no, no homology with any species. The expressed proteins in this study are marked bold character.
The cloned fragments were excised and sub-cloned into pET28a vector (Novagen, Merck Chemicals, Darmstadt, Germany), then transformed into E. coli BL21 (DE3). Positive colonies of E. coli BL21 (DE3) competent cells were large-scale cultured in 200 ml of 2× YT medium containing 100 μg/ml kanamycin (Sigma) at 37 °C. SmTR protein expression was induced by adding 500 μl 0.1 M Isopropyl β-D-1-thio-galactopyranoside IPTG (Takara) and the culture was kept for 3 h at 37 °C. The pellet was stored at −80 °C until use. Extraction and purification of the target proteins were completed by Nickel-nitrilotriacetic acid Ni-NTA agarose (Qiagen) purification of the pellet under denaturing conditions. SmTR8 and SmTR11 were expressed as single and double period of the TR genes, thus SmTR8 presented 2 bands at 10 and 16 kDa, while SmTR11 showed a faint band at 16 kDa and a strong dimer at 30 kDa.

**Fig. 2.** SDS-PAGE (A) and western blot (B) of recombinant SmTR proteins. SDS-PAGE was conducted under denaturing conditions using MES buffer and a NuPAGE 4–12% Bis-Tris gel. M: molecular weight marker, 1: SmTR1, 3:SmTR3, 8:SmTR8, 9:SmTR9, 11:SmTR10, 11:SmTR11 and 15:SmTR15. SmTR8 and SmTR11 were expressed as single and double period of the TR genes, thus SmTR8 presented 2 bands at 10 and 16 kDa, while SmTR11 showed a faint band at 16 kDa and a strong dimer at 30 kDa.

**Fig. 3.** Comparison of total IgG responses to SmTR antigens between infected and control mice by ELISA. Sera were collected from control (n = 7) and infected (n = 17) mice 8–10 weeks post-infection. Specific IgG responses to TR antigens, SEA and SWAP were measured by ELISA. The IgG levels in the infected group (black dots) were significantly increased as compared to the naive group (black squares). Data are presented as scatter dot plots, lines represent the mean, and dashed lines represent the cut off value (mean + 3SD). Cut off value for each antigen: SmTR1, 0.052; SmTR3, 0.097; SmTR8, 0.200; SmTR9, 0.130; SmTR10, 0.081; SmTR11, 0.045; SmTR15, 0.083; SEA, 0.107 and SWAP, 0.08. p value is indicated for each antigen.
conditions, following the manufacturer’s instructions. The extraction and purification of SmTR8 and 15 followed a slightly different procedure. Instead of lysing of *E. coli* culture pellet with 8 M urea, Bugbuster protein extraction reagent (Novagen) was used following the manufacturer’s protocol as it provided better purification conditions for soluble SmTR8 and 15. Expressed proteins were eluted from the column with 150 mM imidazole. The purified proteins were dialyzed against 20 mM Tris–HCl (pH 8.5), then the buffer was exchanged with PBS using Amicon ultra centrifugal filters 3000 MWCO (Millipore, Volketswil, Switzerland).

The integrity of the proteins was assessed by polyacrylamide gel electrophoresis (SDS-PAGE) using 4–12% NuPAGE® Novex Bis-Tris mini gels (Invitrogen, Carlsbad, CA) under denaturing and reducing conditions. The developed gels were stained with Coomassie blue using Simply Blue Safestain (Invitrogen).

In addition, western blotting was conducted using mouse anti-His tag monoclonal antibody (Novagen). Where the size of the recombinant protein did not correspond to the expected size on SDS-PAGE, the peptide sequence was analyzed by trypsin digestion and liquid chromatography–mass spectrometry (LC–MS/MS) following the detailed protocol described elsewhere [51]. After measuring the concentration of the recombinant proteins as described above for SWAP and SEA, the antigens were aliquoted and stored below −30 °C.

### 2.6. Antibody detection using ELISA

Nunc MaxiSorp microtiter plates (Nalgene Nunc Int., Roskilde, Denmark) were coated with 100 μl of 2 μg/ml (200 ng per well) recombinant SmTR antigen in coating buffer (15 mM Na2CO3, 35 mM NaHCO3, pH 9.6), or 200 ng/ml for SEA and SWAP. Plates were then blocked with 1% bovine serum albumin (BSA) (Sigma) in PBS, pH 7.4 for 2 h at RT. Serum samples were diluted 400 times in PBST containing 0.1% BSA, added to the plates, and incubated at RT for 60 min. Horseradish peroxidase-conjugated goat anti-mouse IgG antibody (R&D, Minneapolis, MN) was used as secondary antibody at a dilution of 1:2000 in PBST containing 0.1% BSA, and tetramethylbenzidine (TMB) (BD Pharmingen, Allschwil, Switzerland) was added as substrate. Absorbance was measured at 450 nm using a Multiskan FC microplate reader (Thermo Scientific). All sera were assayed in duplicate. Sera collected from infected mice 8 and 9 weeks post-infection were pooled and considered as the control standard to assess the ELISA conditions; control negative sera were also included on the ELISA plate.

Specific IgG subtypes and IgE antibodies were detected using 50 times diluted sera of infected mice collected 9 weeks post-infection, and rat anti-mouse IgE-HRP-conjugated antibody (AbD Serotec, Bio-Rad) or rat anti-mouse IgG subtype (1, 2a, 2b or 3)-HRP antibody

![Fig. 4](image_url)
(Invitrogen) were used at 10,000 dilution; the remaining ELISA steps were as described for total IgG detection.

2.7. Statistical analysis

The statistical software Graphpad Prism 5.0 (GraphPad Software Inc., La Jolla, CA) and Excel 2011 (Microsoft Corporation, Seattle, WA) were used for statistical analysis and graph generation. The cutoff point between infected and uninfected group results was estimated as the mean plus 3 standard deviation (mean + 3 SD) of the OD of the control group sera for each respective antigen. ANOVA and t test were used for normally distributed samples, and the Mann–Whitney test or Kruskal–Willis test were used for all other samples. Linear regression was used to evaluate the diagnostic performance between TR antigens and SEA or SWAP. To compare the difference between groups, an r2 value of 0.05 was considered significant.

3. Results

3.1. In silico analysis for the identification of tandem repeat genes

The whole genome of S. mansoni was obtained from GeneDB [52] and screened to detect tandem repeat genes by in silico analysis using "Tandem repeats finder software" [53] following the scoring setting as previously described [38]. Out of 11,809 putative genes of this parasite, the top 20 genes containing TRs with a score equal to or greater than 1000 were selected for further analysis. These genes were named SmTR1 to 20, numbered in descending order of their TR score for easy reference. Among these genes, some homology with the mouse or human proteins were excluded from the study to prevent cross-reaction with the host antigens. The Smp_160680.1 (SmTR12) gene codes for ubiquitin, ribosomal protein L40. This gene shares similarities with genes conserved in various species, including S. japonicum and mouse. Similarly, the Smp_160680.1 (SmTR12) gene contains a sequence coding for a domain of breast cancer-related protein type 2 (BCRA2) and thus was excluded from this study.

3.2. Cloning and expression of SmTR antigens

SmTR1 and 3 were dominantly expressed in S. mansoni at the adult and egg stage (Fig. 1A&B) and were not expressed in the cercaria stage (Fig. 1C). The remaining SmTR genes were expressed at all stages of the parasite life cycle within the host; nevertheless, the expression level of SmTR11 was weak as compared to that of tubulin and other SmTR genes. SmTR8 showed a different expression pattern in adult, egg and cercaria; suggesting that it may have some functional role in the life cycle of the parasite. The recombinant proteins were expressed as a single TR period in the gene; the exceptions were SmTR8 and SmTR11, with TR period sizes of 123 and 171, respectively, which were expressed as both single and double TR periods.

3.3. Antibody response to TR antigens

To evaluate the potential of these antigens to induce antibody responses, sera were collected from naive and infected mice 8 to 10 weeks post-infection and were subjected to ELISA probing for total IgG. The sera from infected mice reacted to each antigen, but not the parasitensis or SEA. The sera from naive mice did not react to any of the antigens tested. The correlation between IgG responses to SmTR8 antigen, SEA and SWAP is given in Fig. 5. The OD values of IgG responses to S. mansoni antigens were assessed using concordance with SEA and SWAP as reference; r²: correlation coefficient value.
sera from naïve mice. These results were consistent with those obtained for SEA and SWAP (Fig. 3).

As the intensity and expression pattern of some SmTR genes is distinct between the egg, cercaria and adult worm stages of the parasite life cycle (Fig. 1), we measured the reaction of IgG antibodies to TR antigens during the course of infection. Significant IgG reaction to crude SWAP was detected in sera 5 weeks post-infection compared to the cut-off value, while IgG responses to SEA increased from 6 weeks post-infection. IgG levels to both SWAP and SEA peaked at 10 weeks post-infection (Fig. 4). IgG to SmTR antigens was detected in sera 5 weeks post-infection, similar to SWAP IgG (Fig. 4). The increase in the antibody to SmTR15 was delayed to 7 weeks post-infection. SmTR1, 8 and 10 induced a higher level of IgG antibodies compared to other recombinant antigens. The IgG response to SmTR10 increased by as early as 3 weeks post-infection when compared with the cut-off value. Nevertheless, a statistically significant increase in IgG antibodies to recombinant antigens could be detected only from either 8 or 9 weeks post-infection when compared to IgG levels in naïve sera. Unlike other SmTR antigens, total IgG levels to SmTR1 and 3 were exceptionally and significantly elevated at 5 weeks post-infection.

3.4. Antibody response after praziquantel treatment

A general characteristic of antibody responses is that their level is maintained in individuals for a certain period of time even after clearance of external antigens. We therefore assessed the levels of IgG response to each SmTR antigen, as well as those to SEA and SWAP, after treatment of infected mice with praziquantel. IgG levels to SEA and SWAP remained high, comparable to those in untreated mice (Fig. 4). Interestingly, antibodies to the recombinant antigens tended to rapidly decrease after treatment with praziquantel; the exception was SmTR8, for which the IgG levels remained high, similar to those for SWAP. The response to SmTR3, 10 and 15 dramatically decreased to baseline level after treatment (Fig. 4).
3.5. Correlation between SmTRs, SEA and SWAP

The diagnostic performance of SEA showed a close correlation with SWAP (Fig. 5), similar to previously reported studies [56]. Among the SmTR antigens, SmTR8 showed the highest correlation with SWAP and SEA (Figs. 5 and S2A & B). However, antibodies to SmTR8 remained increased even after treatment with praziquantel, unlike for SmTR3, 9 and 10 (Fig. 4). SmTR1 and 3 showed the lowest correlation with both SEA and SWAP (Fig. S2A & B).

3.6. Antibody subtype specific reaction

IgGs have different subtypes in human and mouse, with different functions [57]. IgG4 subtypes in human and IgG1 in mouse are involved in resistance and susceptibility to schistosomiasis [58,59] and thus are of great interest for the diagnosis of helminthic diseases [60].

We accordingly attempted to analyze the IgG subtypes raised against these antigens, as well as detecting specific IgE antibody, an important immunoglobulin during helminthic infections, including schistosomiasis [59]. The IgG1 subtype was found to be the dominant IgG subtype and several of these antigens induced an increase in specific IgG4 to SmTR antigens, as well as detecting specific IgE antibodies against these antigens, as well as detecting specific IgE antibody, an important immunoglobulin during helminthic infections, including schistosomiasis [59]. The IgG1 subtype was found to be the dominant IgG subtype and several of these antigens induced an increase in specific IgG4 to SmTR antigens, as well as detecting specific IgE antibodies against these antigens.

Specific IgE responses to SmTR antigens were significantly increased in the infected group, with the exception of IgG responses to SmTR15, which were not induced after infection.

4. Discussion

The main purpose of our study was to evaluate *S. mansoni* TR recombinant proteins as potential candidates for the sero-epidemiology of schistosomiasis. All antigens successfully expressed in this study were previously reported as hypothetical or putative proteins [32]. Our results provided further evidence of their expression at the protein level during the life cycle of the *S. mansoni* parasite. These recombinant TR antigens induced antibody responses that were detected during *S. mansoni* infection. Antibody reaction to SmTR10 was detected from 3 weeks post-infection onwards as compared to the cut off value, much earlier than those to SEA, which are significantly positive 6 weeks post-infection (Fig. 4), corresponding to the release of eggs by the adult female [5]. In addition, the level of specific IgC antibodies to SmTR1, 3, 9, 10 and 15 rapidly decreased after treatment with praziquantel, suggesting that these antigens could be promising candidates not only for the diagnosis of *S. mansoni* infection, but also as potential markers in monitoring re-infection after treatment with praziquantel.

The expression pattern of the SmTR8 gene was distinct in egg, adult worm and cercaria (Fig. 1), suggesting that this gene might have a function within the life cycle of schistosome. Further studies are required to assess the function of this antigen.

Moreover, specific IgG responses to SmTR8 showed a strong correlation with both SEA and SWAP in diagnostic performance. SEA preparation requires a consistent volume of eggs and animals [48] that are time and resources consuming, thus SmTR8 might be an alternative candidate diagnostic test to SEA and SWAP, since these crude antigens are commonly used in non endemic countries and laboratory settings to diagnose schistosomiasis [18,30].

Despite the limitations of antibody detection-based diagnosis in discriminating current from past infection, its high sensitivity provides advantages for detecting schistosomiasis cases in low-endemic areas [16,20].

The size discrepancy of SmTR3 on SDS-PAGE (Fig. 2) might be explained by the serine-rich characteristics of this antigen that induced a slow migration on the gel. A similar condition was described with histidine-rich protein-2 used for the rapid diagnosis of *Plasmodium falciparum* infection [61,62], with the serine-rich repeat invariant surface glycoprotein ISG of *Trypanosoma brucei* [63], and with the glycoproteins p120 and p140 of the bacteria *Ehrlichia chaffeensis* and *Ehrlichia canis* [64].

Most specific antibody responses to TR antigens were of the IgG1 subtype and several of these antigens induced an increase in specific IgE antibodies in sera (Fig. 6). As IgG4 in humans decreases dramatically after treatment with praziquantel, detection of IgG4 antibody responses against helminthic infections can help differentiate active infection from prior exposure [60]. Further study of human schistosomiasis will be required with the aim of detecting specific IgG4 to SmTR antigens in sera and urine, thereby providing more insights into these currently hypothetical proteins. At 5 to 10 weeks after infection with 100 cercariae, the median number of worms collected from BALB/c mice was 30 with interquartile range of 15. We could not find any correlation between the number of worm and the IgG antibody levels in this condition. Although BALB/c mice can tolerate more cercariae than some other mouse strains, 100 is a lot for any mouse. In order to confirm the reproducibility of our data, we infected BALB/c and C57BL/6 mice with 50 cercariae per mouse in a different experiment, and the outcomes obtained were comparable with the present experiments (data not shown).

In silico screening of the TR genes showed shared similarities among *Schistosoma* species, implying potential cross-reaction in co-endemic areas, especially with *S. haematobium* in sub-Saharan regions where many foci of co-infection are reported. It could be considered as an advantage to detect different species with one diagnostic test in field settings [20,65], because the treatment is the same for infection by these schistosome species.

The main purpose of this study was to identify and express TR-containing antigens of *S. mansoni* and focused on a mouse model using a limited number of mice; consequently, this study could not estimate the exact sensitivity and specificity of the proposed diagnostic antigen tests.

5. Conclusions

The availability of suitable markers for monitoring schistosome infections is urgently required for the control and elimination of schistosomiasis. SmTR antigens, and especially SmTR10, are potential markers for the diagnosis of infection, and for monitoring re-infection by *S. mansoni* after treatment with praziquantel.

Since the goal is to use these antigen candidates for identifying human infection cases, our future work will evaluate the validity of the proposed antigen candidates using human samples from a region endemic for schistosomiasis.

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