Recombinant tegumental protein Schistosoma japonicum very low-density lipoprotein binding protein as a vaccine candidate against Schistosoma japonicum

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A polyhistidine-tagged recombinant tegumental protein Schistosoma japonicum very low-density lipoprotein binding protein (SVLBP) from adult Schistosoma japonicum was expressed in Escherichia coli. The affinity purified rSVLBP was used to vaccinate mice. The worm numbers and egg deposition recovered from the livers and veins of the immunized mice were 33.5% and 47.6% less than that from control mice, respectively (p<0.05). There was also a marked increase in the antibody response in vaccinated mice: the titer of IgG1 and IgG2a, IgG2b in the vaccinated group was significantly higher than that in the controls (>1:6,400 in total IgG). In a comparison of the reactivity of sera from healthy individuals and patients with rSVLBP, recognition patterns against this parasite tegumental antigen varied among different groups of the individuals. Notably, the average titres of anti-rSVLBP antibody in sera from faecal egg-negative individuals was significantly higher than that in sera from the faecal egg-positives, which may be reflect SVLBP-specific protection. These results suggested that the parasite tegumental protein SVLBP was a promising candidate for further investigation as a vaccine antigen for use against Asian schistosomiasis.

Key words: Schistosoma japonicum - Schistosoma japonicum very low-density lipoprotein binding protein - vaccine

Schistosomiasis, also known as bilharzia, is the major health risk in tropical and subtropical countries around the world, which affects more than 200 million individuals in 74 countries and remains among the most debilitating parasitic diseases of humanity (Engels et al. 2002, Ross et al. 2002, Chitsulo et al. 2004). Praziquantel, the current drug of choice for all forms of human schistosomiasis, is safe and effective. However, efficient drug delivery requires a substantial infrastructure to regularly cover all parts of an endemic area and, moreover, in endemic regions people often are re-infected soon after drug cure. In addition, there are continued concerns about the emergence of drug resistance to praziquantel (Cioli et al. 2004). Hence, a new approach is needed for the development of vaccines to control this disease (reviewed in McManus & Bartley 2004). Recent studies aimed at sequencing the genome of S. japonicum have provided new sources for the development of new target antigens (Fan et al. 1998, Hu et al. 2003). In our previous report, an expressed sequence tag (EST) of S. japonicum encoding a S. japonicum very low-density lipoprotein binding protein (SVLBP) arose as interesting candidate for further study its biological functions and role in the host-schistosome relationship (Fan et al. 2003). SVLBP was reported to function in membrane association, and located in the tegument and sub-tegument of adult male schistosomes. SVLBP can play an essential role in lipid acquisition by the parasite, and/or in signal transduction pathways, we considered that it was worthy of further investigation for development as a novel anti-schistosomal intervention. Here we report the expression and purification of the recombinant SVLBP for investigation as a vaccine candidate. We express the SVLBP as an experimental vaccine, vaccination of mice with affinity purified recombinant SVLBP, the IgG subclass antibody responses in the vaccinated mice, and anti-rSVLBP antibody responses in sera of humans naturally infected with S. japonicum.

MATERIALS AND METHODS

Parasites, experimental animals, and patients - A Chinese mainland isolate of S. japonicum from the Anhui
Province was maintained in the laboratory by infecting Oncomelania hupensis hupensis snails and Netherlands rabbits. Active cercariae were freshly shed from the infected snails under light irradiation and used for infection of rabbits or mice. Female mice C57BL/6 were purchased from the Experimental Animal Center, Zhejiang Province, PR China. In addition, 253 sera were collected from human subjects residing in S. japonicum endemic regions of Jiangxi Province, PR China. Stool samples of these volunteers were examined for S. japonicum eggs using the modified Kato-Katz thick smear method (Katz et al. 1972). The faecal egg positive individuals were offered treatment for schistosomiasis with praziquantel. Furthermore, human serum samples were also obtained from 105 other individuals residing in non-schistosome endemic regions (Hangzhou City, Zhejiang Province) in PR China.

Recombinant SVLBP - A cDNA encoding SVLBP was cloned into pQE30 (Qiagen) and the expression of the recombinant protein was induced with a final concentration of 1 mM isopropyl-beta-D-thiogalactopyranoside (IPTG) at 37°C for 4 h with shaking after transformation into E. coli (Fan et al. 2003). The recombinant SVLBP was purified on Ni-NTA Superflow resin by affinity chromatography under native conditions, using a linear imidazole gradient (10-250 mM). Eluted fractions containing the expressed recombinant protein were pooled and concentrated in PBS buffer (20 mM NaH2PO4, 30 mM NaCl, pH 7.5). The presence of the recombinant protein was confirmed by SDS-PAGE analysis. Protein concentration was determined by Lowry’s method (Markwell et al. 1978).

Immunization of mice with the recombinant SVLBP - Female C57BL/6 mice, 18-22 g in weight and 6-8 weeks old were randomly divided into two groups. Thirteen mice in the experimental group were each injected subcutaneously (sc) with 45 µl phosphate buffered saline containing 45 µg purified recombinant SVLBP plus complete Freund’s adjuvant at multiple sites of the abdominal area, at weeks 0, 2, 4, respectively. Twelve mice in the control group received the same volume of PBS plus adjuvant as mice in the immunization group. Serum samples were obtained before vaccination and at the time of necropsy, respectively.

Cercarial challenge, worm recovery and egg count - Mice of both groups were challenged percutaneously with 35 ± 1 cercariae by cover slip method (Ruppel et al. 1990) two weeks after the final vaccine or control injection. Mice were euthanized 45 days after the challenge infection by overdose of heparinized sodium pentobarbitone (Sigma) administered by intraperitoneal injection. Immediately thereafter, the portal system blood vessels perfused in order to recover the worms from the livers and veins. Subsequently, the liver were each weighed and minced and digested in 10 ml of 5% KOH at 37°C for 24 h after which schistosome eggs released from the infected livers were counted in 100 µl aliquots. Five aliquots were counted for per liver samples (Liu et al. 1995) aided by light microscopic examination.

Enzyme-linked immunosorbent assay - Serum anti-SVLBP antibody responses of immunized mice were measured by ELISA. The plates were coated with 2 µg/well of recombinant SVLBP in carbonate/bicarbonate buffer (pH 9.6). After blocking with 3% (w/v) BSA in PBS/0.05% (v/v) Tween-20 (PBS-T20), mouse sera were diluted 1/100 in PBS-T20 and added (100 µl/well), then incubated at 37°C for 1 h. 100 µl/well of horseradish peroxidase (HRP)-labeled anti-mouse IgG, IgG1, IgG2a, IgG2b, IgG3 (Caltag Laboratories, Inc. US) diluted 1/2000 was added and incubated at 37°C for 1 h. One mg/ml TMB (3′, 3′′, 5′, 5′′-tetramethylbenzidine) was used as the substrate, and the absorbance was read at 450 nm in a spectrophotometer 15 min after incubation.

 Statistical analysis - The non-parametric r-test and/or Mann-Whitney U test were used to determine the significance of differences in mean worms and egg counts from the immunized groups and the adjuvant control, and to determine the human antibody responses to rSVLBP. All results including the ELISA optical density (OD) values are presented as mean ± S.D. A p value of ≤ 0.05 was considered statistically significant.

RESULTS

Expression of the recombinant SVLBP in E. coli and its purification by affinity chromatography - Expression of the recombinant SVLBP was achieved by introduction of IPTG to the culture, resulting in that a 22 kDa protein was induced (Fig. 1), which was in accordance with the previous description (Fan et al. 2003). The 22 kilodaltons was approximately the size expected of a full-length recombinant fusion protein, i.e. about 20 kDa for SVLBP and 2 kDa for the N-terminal fusion peptide incorporating the six-histidine tag. A single-step solubilization/purification on Ni-NTA resin under non-denaturing conditions was used to purify the recombinant SVLBP in native form. This chromatographic procedure was able to enrich the SVLBP almost to homogeneity (Fig. 1).

Protective efficacy of rSVLBP against S. japonicum - Two groups of mice were used as experiment and control, respectively. Group 1 was immunized with rSVLBP (n = 13), and group 2 was injected with adjuvant as control (n = 12). Worm numbers and liver egg counts for immunized and control groups are presented in the Table. The mean number (X ± S.D.) of worms recovered from mice vaccinated with rSVLBP (15.5 ± 6.1) (n = 13) was significantly reduced by 33.5% (p < 0.05) compared with the mean worm numbers from control mice (23.3 ± 6.8) (n = 12). There was also a significant 47.6% decrease (p < 0.05) in mean number of eggs in livers from vaccinated mice (9.7 ± 2.7 x103; n = 13) compared with the mean number of eggs in livers from control mice (18.5 ± 4.2 x103; n = 12).

Increase of the specific anti-rSVLBP antibodies after parasite challenge - Anti-rSVLBP IgG antibody levels (mean absorbance values at 450 nm ± S.D.) were measured in sera of vaccinated and control mice (Fig. 2). Antibody responses to mice immunized with rSVLBP (2.167 ± 0.34) were significantly higher than antibody levels in sera of mice injected with adjuvant only (0.039 ± 0.022; p < 0.01). Distinct antibody titres as high as 1:6400 were obtained from immunized group for the total IgG, indicating that rSVLBP may be an ideal immunogen candidate in that it elicits strong antibody responses. The profile of specific IgG subclasses after challenge was dominated by IgG1,
IgG2a, and IgG2b, but not by IgG3. The IgG1/IgG2 ratios in the vaccinated mice were less than 0.62. There was no significant detectable specific antibody response between the control group and before immunization (Fig. 2).

**Recognition of the rSVLBP by human sera from schistosomiasis endemic and non-endemic regions** - Antigenicity of rSVLBP was measured in 253 sera from individuals from endemic regions where schistosomiasis japonica is rampant and in sera from 105 uninfected volunteers from non-endemic regions. Human sera from the individuals in endemic regions with positive – or negative – faecal examinations exhibited higher responses than those from healthy individuals in non-endemic regions. It is noteworthy that antibody responses to rSVLBP were significantly greater in the individuals with faecal examination negative versus the patients with faecal examination positive (Mann-Whitney U test, \( p < 0.006 \)) (Fig. 3). Together, these data confirm that rSVLBP displays strong immune responses during the course of natural infection, and suggests a protective effect in sero-positive, faecal negative individuals.

**TABLE**

<table>
<thead>
<tr>
<th>Group</th>
<th>No. of mice</th>
<th>Worm burden</th>
<th>Eggs per gram of liver tissue</th>
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<tr>
<td></td>
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<td>X ± SD</td>
<td>Reduction</td>
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<tr>
<td>Recombinant SVLBP</td>
<td>13</td>
<td>15.5 ± 6.1</td>
<td>33.5%</td>
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<tr>
<td>Adjuvant control</td>
<td>12</td>
<td>23.3 ± 6.8</td>
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Fig. 1: SDS-PAGE and Coomassie stained gel analysis of the expression of recombinant SVLBP in *E. coli* and affinity purification. Lanes - 1: molecular size standards (kDa); 2: uninduced bacterial cell lysates; 3: bacterial cell lysates after induced 3h by adding IPTG to 1mM; 4: purified recombinant SVLBP (rSVLBP). The arrow indicates the position of rSVLBP.

Fig. 2: levels of serum IgG isotype antibody responses in C57BL/6 mice detected by ELISA to rSVLBP compared for rSVLBP immunization (group 1), adjuvant control (group 2), and before immunization. Bars represent the mean ± SE.

Fig. 3: reactivity of 358 sera from endemic regions and non-endemic (control) with rSVLBP antigens using ELISA test for intestinal schistosomiasis. The box and whisker plot (type: median/quart/range) shows the median and range values of serum anti-rSVLBP levels. The median response is represented by a black filled box within a rectangular box that represents the 25th to 75th percentile range. Whiskers represent the full range of our actual data. Although the % of the positive reaction to rSVLBP in the patients with a negative faecal egg counts was significantly lower compared to the patients with positive faecal examination, there was a remarkable elevated level of anti-rSVLBP antibody in the faecal egg counts-negative group (Mann-Whitney U test, \( p < 0.006 \)).
DISCUSSION

The schistosomule tegument is an enigmatic structure – a double bilayer – that is involved with the transition from free-living to the parasitic life style in the mammalian host, in immunological masking, and in escape from immune defenses. In addition, it plays a vital role in acquisition of host derived nutrients, including carbohydrates, amino acids, and lipids (Abath & Werkhauser 1996, Jones et al. 2004). Along with other key molecules, tegument associated antigens of schistosomes have been investigated as targets of protective immune responses in murine hosts and for defining antibody and cytokine correlates of apparent resistance and susceptibility in human populations, a key approach to establishing the protective potential of a vaccine candidate (Ribeiro de Jesus et al. 2000, Acosta et al. 2002, 2004, Al-Sherbiny et al. 2003, McManus & Bartley 2004, Bergquist et al. 2005). Since SVLBP appears to have VLDL binding activity and to locate prominently on the tegument and sub-tegument of adult male schistosomes, SVLBP may play a key role in lipid acquisition by *S. japonicum* (Fan et al. 2003). Given that anti-SVLBP antibody can block lipid binding capacity of the SVLBP, blockade of SVLBP by antibodies and other immune effectors may result in starving the parasite of essential lipid nutrients from the host and may also disrupt lipid molecule-associated immune evasion processes (Fan et al. 2003). Here we have shown that, when delivered as a vaccine, rSVLBP can induce a significant level of protection evidenced as reduced worm burden and liver egg counts in the vaccinated mice. The level of protection achieved was comparable to that reported with a number of other experimental vaccine preparations, including with many *S. mansoni* antigens tested in experimentally infected mice (reviewed in Pearce 2003, McManus 2005).

Proiling of antibody levels of anti-rSVLBP antibodies revealed that IgG2a and IgG2b antibody subclasses predominated in the serum of the vaccinated mice. Gamma interferon (IFN-gamma) and interleukin 4 (= B cell stimulatory factor-1, BSF-1), which are T cell-derived lymphokines, can stimulate B cells to produce IgG2a and IgG1, respectively. Therefore, the IgG2a and IgG1 can be used as serological markers to indicate the induction of Th1/Th2 immune responses, respectively (Snapper & Paul 1987). In general, IgG1/IgG2 ratio of <1.0 indicates a predominantly Th1 type of response whereas IgG1/IgG2 ratio of >1.0 indicates a Th2 phenotype. The increase in both IgG2a and IgG2b seen in the present study clearly suggests that a predominantly Th1/Th2 type of antibody response contributed to the protective response induced by vaccination with rSVLBP. Previous reports on vaccination against *S. mansoni* challenge in mice have shown that only immunized mice develop higher levels of IgG2a (and lower levels of IgG1) to larval and adult worm antigens (Mountford et al. 1994). In addition, using DNA encoding *S. mansoni* 28 kDa glutathione S-transferase to immunize rats leads to a long-lasting predominance of IgG2a and IgG2b (Dupre et al. 1997), as does the p80 subunit of *S. mansoni* calpain (Siddiqui et al. 2003). DNA vaccination and co-delivery of antigens with IL-12 result in amplified IgG2a synthesis and down-regulation of synthesis of IgG1 (Fonseca et al. 2004, Siddiqui et al. 2003).

Interestingly, our study demonstrated that vaccination with rSVLBP induced a predominantly a Th1-type of immune response in mice. This suggests that the vaccine effect was likely to have been mediated by an antibody-dependent, cell-mediated cytotoxicity (ADCC) response involving parasite killing the parasite by monocytes, eosinophils and/or platelets (Capron et al. 1983, Horta & Ramalho-Pinto 1984, 1987).

This work provides the first evidence that the newly discovered *S. japonicum*-specific molecule, SVLBP, has potential as a vaccine candidate. Further, the immune recognition of rSVLBP in human sera from individuals from schistosomiasis japonica endemic regions confirms that the rSVLBP can elicit an immune response in natural human populations in endemic regions, which is a requirement for a potential vaccine candidate. In future studies it would be instructive to clarify the immuno-modulatory mechanisms responsible for its vaccine action along the lines reported by Capron and coworkers with recombinant 28 kDa glutathione-S-transferase of *Schistosoma haematobium* (rShGST28), which represents the first schistosome vaccine tested in humans. Capron et al. (2005) noted that, in initial trials, rShGST28 produced in yeast and delivered in alum is safe and that it can engender significant antischistosomal effects. Given that *S. japonicum* infection is a true zoonosis, unlike the other species of schistosomes of public health consequence, it has been suggested that if a transmission blocking veterinary vaccine developed for bovines can be put into practice in combination with other control strategies such as human chemotherapy, elimination of *S. japonicum* from China may be achievable (McManus 2005). Furthermore, it has recently become clear that vaccine-linked chemotherapy associated with the artemisinin derivative artemether offers the potential for an integrated – chemotherapy and vaccination – approach to schistosomiasis control (Bergquist et al. 2005). On both these fronts also, SVLBP appears to be worthy of further investigation as a transmission blocking, and chemotherapy-linked, vaccine.

REFERENCES


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