CONCISE COMMUNICATION

Recombinant *Paragonimus westermani* Yolk Ferritin Is a Useful Serodiagnostic Antigen

Tae Yun Kim,¹ Il-Joong Joo,¹ Shin-Yong Kang,¹ Seung-Yull Cho,² Yoon Kong,² Xiao-Xian Gan,³ Kom Sukomtason,⁴ Kabkaew Sukomtason,⁴ and Sung-Jong Hong¹

¹Department of Parasitology, Chung-Ang University College of Medicine, Seoul, and ²Department of Molecular Parasitology, Sungkyunkwan University School of Medicine, Suwon, South Korea; ³Institute of Parasitic Diseases, Zhejiang Academy of Medical Sciences, Hangzhou, China; ⁴Department of Parasitology, Chiang Mai University Faculty of Medicine, Chiang Mai, Thailand

A recombinant protein of *Paragonimus westermani* yolk ferritin was bacterially produced from a previously cloned complementary DNA and was used as an antigen for an enzyme-linked immunosorbent assay (ELISA) against paragonimiasis- and other helminth-infected sera to evaluate its serodiagnostic potential. The ELISA revealed that paragonimiasis westermani had 88.2% sensitivity and 100% specificity. The positive and negative predictive values of the ELISA were calculated to be 100% and 97.1%, respectively. Sera from cats experimentally infected with *P. westermani* began to produce immunoglobulin G antibodies against the yolk ferritin at 13 weeks after infection, which suggests that the corresponding antigen was derived from the vitellaria in accordance with maturation of *P. westermani*. These results indicate that the recombinant *P. westermani* yolk ferritin is a potent serodiagnostic reagent for paragonimiasis westermani from an early stage of the infection.

Paragonimiasis is a disease caused by lung flukes of genus Paragonimus helminths. Pulmonary paragonimiasis, caused by P. westermani infection, is endemic in the Far East, southeastern Asia, and South America and remains a serious public health problem [1]. The diagnosis of human paragonimiasis westermani is dependent on detection of characteristic eggs in a sputum and/or stool examination. Although there have been some reports about serodiagnosis of the disease by using crude worm extracts as antigens, cross-reactions with other trematode infections, such as schistosomiasis, fascioliasis, or clonorchiasis, decreased the reliability of this diagnostic method [2, 3]. Purified antigens, such as cysteine proteinases of P. westermani, increased sensitivity to paragonimiasis sera and reduced cross-reactivity to fascioliasis sera [4]. Because purification of an antigenic protein from crude preparations to achieve homogeneity is a labor-intensive and time-consuming process, production of specific antigenic molecules into recombinant forms by using appropriate expression systems may be a good alternative way of alleviating the hindrances.

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We cloned a *P. westermani* yolk ferritin cDNA (GenBank accession no. AF188720), produced its recombinant protein in a bacterial system, and characterized its biochemical and molecular biological properties (data not shown). Ferritin is an iron-storage protein identified in a wide range of organisms, from bacteria to mammals [5]. Serodiagnostic potentials of ferritin have been reported to exist in *Echinococcus granulosus* [6] and *Taenia saginata* [7]. We investigate here whether recombinant *P. westermani* yolk ferritin may be useful for serodiagnosis of paragonimiasis westermani.

Materials and Methods

Production of recombinant protein. The coding region of P. westermani yolk ferritin cDNA was cut out by appropriate restriction enzymes and was subcloned into bacterial expression plasmid vector pRSET (Invitrogen). Escherichia coli strain BL21(DE3)pLysS (Novagen) was transformed with the recombined pRSET. Overexpression of the recombinant protein was induced by adding isopropylthio- β -galactoside to the culture medium. Since the recombinant protein was located in insoluble inclusion bodies, it was purified by following the method of van Wuytswinkel et al. [8], with a partial modification. In brief, the bacterial cells were pelleted, resuspended in 20 mM Tris/maleate buffer (pH 8.0), and sonicated on ice. The bacterial lysate was centrifuged at 12,000 g for 10 min at 4°C. The pellet was washed in 20 mM Tris/maleate buffer (pH 8.0) containing 1 mg/mL deoxycholate, 1 mM EDTA, and 0.2 mg/mL lysozyme and was centrifuged, as aforementioned. The pellet was resuspended in 20 mM Tris/maleate (pH 8.0) containing 8 M urea and 0.2% (vol/vol) Triton X-100 and was again

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Reprints or correspondence: Dr. S.-J. Hong, Dept. of Parasitology, Chung-Ang University College of Medicine, Seoul 156-756, South Korea (hongsj@cau.ac.kr).

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centrifuged. The supernatant was saved and was dialyzed against 20 m*M* Tris/maleate (pH 8.0) overnight at room temperature. It was loaded onto Ni-NTA resin (Qiagen) pre-equilibrated with 20 m*M* Tris/maleate (pH 8.0). The resin was washed 3 times with a buffer (50 m*M* NaH₂PO₄, 300 m*M* NaCl, and 20 m*M* imidazole [pH 8.0]). The recombinant protein was eluted with the same buffer, but containing 400 m*M* imidazole, and was dialyzed against 20 m*M* Tris/maleate buffer (pH 8.0) overnight at room temperature. The recombinant protein thus produced was 28-kDa fusion protein with 4-kDa tag protein. Protein concentration was determined by use of Protein Assay Kit II (BioRad).

ELISA. The wells of a 96-well microplate were sensitized with 1 µg/100 µL of the recombinant P. westermani yolk ferritin at 4°C overnight. The wells were incubated with sera from humans infected with paragonimiasis westermani (n = 34), clonorchiasis sinensis (n = 32), opisthorchiasis viverrinii (n = 9), schistosomiasis japonicum (n = 30), fascioliasis hepatica (n = 10), cysticercosis cellulosae (n = 20), and sparganosis (n = 25) and with sera from uninfected humans (n = 10) at a 1:200 dilution at 37°C for 2 h. They were then incubated with peroxidase-conjugated anti-human IgG (Cappel Laboratories) at a 1:2000 dilution for 1 h. Color was developed by using a substrate, o-phenylene diamine, and absorbance was measured at 490 nm (A₄₉₀) with a microplate reader (model 3100; Bio-Rad Laboratories). The data obtained were analyzed for diagnostic sensitivity, specificity, and positive and negative predictive values [9], which were defined and calculated as follows: true negative = no. of control sera (other helminthiasis-infected and healthy control sera) that were negative by the assay; true positive = no. of proven paragonimiasis-infected sera that were positive by the assay; false negative = no. of proven paragonimiasis-infected sera that were negative by the assay; false positive = no. of control sera that were positive by the assay; sensitivity = [no. of true]positives/(no. of true positives + no. of false negatives)] $\times 100$; specificity = [no. of true negatives/(no. of false positives + no. oftrue negatives)] $\times 100$; positive predictive value = [no. of true positives/(no. of true positives + no. of false positives)] $\times 100$; negative predictive value = [no. of true negatives/(no. of true negatives + no. of false negatives)] $\times 100$.

Immunoblot. The recombinant *P. westermani* yolk ferritin was separated by 12.5% SDS-PAGE and was electrotransferred onto a nitrocellulose membrane. Each stripped membrane was incubated, at a 1:200 dilution, with the pooled cat sera, which were collected at 0, 2, 4, 6, 8, 10, 13, and 15 weeks after the experimental infections with *P. westermani* metacercariae. The sera were incubated with peroxidase-conjugated anti–cat IgG (Cappel Laboratories) at a 1:2000 dilution, and color was developed by using 4-chloro-1-naphthol as a substrate.

Results

Absorbances of ELISA, obtained by using the recombinant *P. westermani* yolk ferritin as an antigen, are plotted in figure 1. The cutoff absorbance for a positive reaction was set at $(A_{490}) = 0.214$, which was a mean + 10 SD of the absorbances obtained from uninfected control sera. Of 34 paragonimiasis-infected sera, 30 were determined to be positive, representing

a sensitivity of 88.2% (absorbance range, 0.068-0.660; mean \pm SD, 0.469 ± 0.139). Absorbances of sera of the other helminthiases were below the cutoff absorbance. The mean of the absorbance to paragonimiasis serum was >5 times higher than those of the other helminthiasis sera. Since there were no cross-reactions to other helminthiasis sera, 100% was the specificity of the ELISA using the recombinant *P. westermani* yolk ferritin. The cutoff value employed, 0.214, which was even lower than the conventional cutoff value of 0.25 [10], produced good positive and negative predictive values of 100% and 97.1%, respectively.

IgG antibodies reacting to the recombinant *P. westermani* yolk ferritin were detected by an immunoblot in experimental cat sera after 13 weeks of the metacercarial infection (figure 2).

Discussion

As for molecular biological properties, *P. westermani* yolk ferritin shared a low sequence homology with those of vertebrate and invertebrate animals (data not shown). However, a ferroxidase center was predicted to be formed with a coordination of the highly conserved iron-binding amino acid residues [5], conferring an enzymatic function to the molecule. From a serodiagnostic point of view, the low homology of *P. westermani* yolk ferritin to those of invertebrates, including helminthic parasites, seems to favor its antigenic specificity.

E. granulosus ferritin was reported earlier to show a specific antigenicity with cystic and alveolar echinococcosis sera but not with cysticercosis sera [6], whereas *T. saginata* ferritin cross-reacted to both cysticercosis and hydatidosis sera [7]. Although

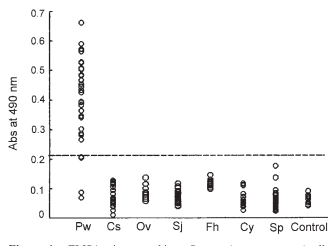


Figure 1. ELISA using recombinant *Paragonimus westermani* yolk ferritin in sera from patients with paragonimiasis westermani or various helminthiases. Each circle (\bigcirc) represents the mean absorbance (Abs) of 3 wells. Pw, paragonimiasis westermani; Cs, clonorchiasis sinensis; Ov, opisthorchiasis viverrinii; Sj, schistosomiasis japonicum; Fh, fascioliasis hepatica; Cy, cysticercosis cellulosae; Sp, sparganosis; control, uninfected human. Cutoff value at absorbance 0.214 is shown (*dashed line*).

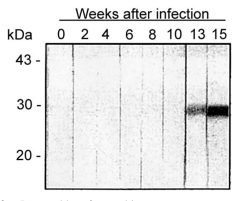


Figure 2. Immunoblot of recombinant *Paragonimus westermani* yolk ferritin probed with pooled sera of cats infected with *P. westermani*. Antibody reactivity was detected after 13 weeks of infection.

cestode ferritins share antigenic epitopes among the systematically related taxa, the recombinant *P. westermani* yolk ferritin do not cross-react to them (figure 1).

The ferritin was localized in vitellaria and eggs of adult *P. westermani* (data not shown). Yolk materials produced in vitellaria are provided to ova and packed in the eggs. In cats, *P. westermani* grows up to be an egg-producing adult between 7 and 8 weeks after the infection [1]. An increase in IgG antibody levels in the cat sera to the egg extract of *P. westermani* was attributed to a 28-kDa molecule after 13 weeks of the infection [11]. The antibody response to the recombinant *P. westermani* yolk ferritin, shown in figure 2 in regard to the 28-kDa molecule, is in accord with the egg production of the worm, with a transition of 4–5 weeks.

Eggs of *P. westermani* remain alive in the pulmonary (their natural habitat) and in extrapulmonary tissues for prolonged periods after the fluke dies. A seropositive result obtained by using egg antigens of various types, including yolk ferritin, may

imply that the patient is in a prolonged chronic stage of the infection. We propose here that the recombinant *P. westermani* yolk ferritin is a potent antigen for serodiagnosis of human paragonimiasis westermani in early and prolonged stages.

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