Immunodetection of *Fasciola gigantica* Circulating Antigen in Sera of Infected Individuals for Laboratory Diagnosis of Human Fascioliasis

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Currently, the laboratory diagnosis of human fascioliasis is based on the parasitological examination of parasite eggs in stool specimens and serological detection of specific antibodies in serum samples, which are often unreliable diagnostic approaches. Ideally, a sensitive and specific diagnostic test for *Fasciola* infection should be based on the detection of circulating *Fasciola* antigen, which implies active infection. Here, a 27-kDa-molecular-mass antigen was identified in a *Fasciola gigantica* adult worm antigen preparation, excretory-secretory products, and sera from *F. gigantica*-infected individuals, and it was not detected in antigenic extracts of other parasites and sera from noninfected individuals. The target antigen was isolated and partially characterized as a protein. Immunoperoxidase staining located the target epitope within teguments and guts of *F. gigantica* adult worms. The performance characteristics of a newly developed enzyme-linked immunosorbent assay (ELISA) based on *F. gigantica* circulating antigen detection in serum (FgCA-27 ELISA) were investigated using sera of 120 parasitologically diagnosed *F. gigantica*-infected individuals and 80 noninfected individuals. The area under the receiving operating characteristic (ROC) curve (AUC) for ELISA was significantly high (AUC = 0.961, P < 0.0001) for discriminating *Fasciola*-infected and noninfected individuals. The developed assay showed high degrees of sensitivity, specificity, and efficiency (>93%), and a significant correlation (r = 0.715, P < 0.0001) between antigen level and parasite egg count was shown. In conclusion, a 27-kDa *Fasciola* antigen was identified in sera of *F. gigantica*-infected individuals. A highly sensitive and specific *Fasciola* antigen detection assay, FgCA-27 ELISA, was developed for laboratory diagnosis of human fascioliasis.

Fascioliasis is a worldwide zoonotic infection caused by liver flukes of the genus *Fasciola*, of which *F. hepatica* and a larger species, *F. gigantica*, are the most common representatives. These two food-borne trematodes usually infect domestic ruminants and cause important economic losses of sheep, goats, and cattle in commercial herds (1). Today, fascioliasis is considered an important human disease, and several areas, e.g., tropical regions of Africa and Asia, have been described as areas of endemicity for the disease in humans, with estimates of 2.4 to 17 million people infected and 91.1 million at risk worldwide (2, 3). Egypt, especially the Nile Delta, is a hot spot for *Fasciola* infection, which has appeared to be endemic, with estimated prevalence varying between 2 and 17% (4–6). Furthermore, fascioliasis has been recognized as an emerging infection in international travelers and migrants, causing significant problems in diagnosis and therapy (7). Owing to the disease’s high-level endemicity in some developing countries and its emergence in nature, the WHO classified it as a neglected parasitic infection and decided to launch a worldwide initiative against this infectious disease (2).

Diagnosis of human fascioliasis is based on clinical findings and laboratory tests. The most reliable means is the finding of parasite eggs in stool of an infected individual (8). However, parasitological diagnosis of human fascioliasis is time-consuming and usually lacks sensitivity and reproducibility, and it is often unreliable because parasite eggs are not found during the prepatent period, which lasts until approximately 3 to 4 months after infection (9). Moreover, once *Fasciola* worms have matured, diagnosis may still remain difficult, since eggs are frequently excreted at irregular intervals, and in most cases, repeated stool examinations are required to accurately diagnose (10). Anti-*Fasciola* antibodies can be detected as early as 2 weeks postinfection, and this can thus facilitate early diagnosis and chemotherapeutic intervention (11). Over the past years, several native and recombinant antigens were identified for the detection of serum-specific anti-*Fasciola* antibodies (12–19), and many antibody-based immunological techniques have been developed for diagnosis of *Fasciola* infection in a trial to replace the classical parasitological techniques (20). However, antibody tests do not distinguish between past, resolving, and current infections, and their results do not correlate with infection intensity (21).

In that sense, the direct detection of parasite antigens in stool (coproantigens) or serum (circulating antigens) of *Fasciola*-infected animals or humans has been reported as a new alternative approach with greater diagnostic accuracy (22–26). In our previous study, we identified a 26- to 28-kDa circulating antigen in sera of cattle infected with *F. gigantica* by using specific rabbit IgG antiserum (22). In the present study, we aimed to identify the target *F. gigantica* circulating antigen in sera of infected individuals and describe a novel enzyme-linked immunosorbent assay (ELISA) based on circulating antigen detection in serum for accurate laboratory diagnosis of human *F. gigantica* infection.

**MATERIALS AND METHODS**

**Stool and blood samples from study subjects.** A total of 200 individuals (101 males and 99 females; age range, 6 to 65 years; mean, 24 years)
admitted at the Tropical Medicine and Parasitology Department, Mansoura University Hospitals, Mansoura, Egypt, were included in the present study. Stool specimens were collected from each individual and examined at the day of collection using simple stool sedimentation by centrifugation. The Kato-Katz thick-smear technique was used for counting *F. gigantica* eggs in 3 to 5 slides, each containing 41.7 mg stool, and the egg count was expressed as eggs per gram of feces (EPG). A total of 120 individuals had *F. gigantica* eggs in their feces, 57 individuals had other parasitic infections, including 38 individuals infected with *Entamoeba histolytica*, 7 infected with *Schistosoma mansoni*, 5 infected with *Ascaris lumbricoides*, 4 infected with *Hymenolepis nana*, and 3 infected with *Giardia lamblia*, and 23 individuals were diagnosed as parasite-free, healthy individuals. According to the Kato-Katz technique results for individuals infected with *F. gigantica*, the intensity of infection was classified as light (<100 EPG) in 76 individuals, moderate (100 to 299 EPG) in 33 individuals, and severe (>300 EPG) in 11 individuals. Blood samples were collected on the day of stool analysis. Sera were separated from blood, aliquoted, and stored at −20°C until used. The Ethical Committee of the Mansoura University Hospitals, Mansoura, Egypt, approved the present study. Informed consents were obtained from all participants, and all individuals were fully informed concerning the diagnostic procedures involved and nature of the disease.

*F. gigantica* and other parasite antigenic preparations. *F. gigantica* soluble worm antigen preparation (FWAP) was prepared as described by Attallah et al. (22). The crude excretory/secretory (E/S) products of adult *F. gigantica* were prepared according to the description of Santiago de Weil and Hillyer (27). Adult *A. lumbricoides* worms, from the stools of infected individuals, were processed in exactly the same way as *F. gigantica* to prepare *A. lumbricoides* soluble worm antigenic preparation (AWAP), while *S. mansoni* soluble worm antigenic preparation (SWAP) was prepared as described by da Silva and Ferri (28). In each run, 250 µl of diluted soluble worm antigenic preparation (SWAP) was applied to prepare gel was stored at −20°C until used. The Ethical Committee of the Mansoura University Hospitals, Mansoura, Egypt, approved the present study.

**Biochemical treatments.** To determine some of the target *Fasciola* antigen biochemical characteristics, sera of the purified 27-kDa antigen were subjected to SDS-PAGE and Western blot. Various samples were subjected to analytical SDS-PAGE, at 50 µg/lane, using vertical slabs of 12% or 16% polyacrylamide (30). Prestained molecular mass standards (Sigma) were run in parallel. Samples separated on SDS-PAGE were electrotransferred onto nitrocellulose (NC) membrane (0.45 µm pore size) in a protein transfer unit (31). The NC membrane was blocked using 5% (wt/vol) nonfat dry milk dissolved in 0.05 M Tris-buffered saline (TBS) containing 200 mM NaCl (pH 7.4), rinsed in TBS, and incubated with anti-27-kDa *Fasciola* antigen IgG antibody, diluted (1:150) in the blocking buffer as described below, with constant shaking. The blots were washed three times (30 min each) in TBS and then incubated for 2 h with goat anti-rabbit IgG–alkaline phosphatase conjugate (Sigma) diluted 1:350 in TBS. After being washed three more times with TBS (15 min each), the blots were soaked in substrate. The color reaction was observed within 10 min, and the reaction was then stopped by dipping the blots in distilled water. To ensure that the 27-kDa protein purified from serum was a parasite molecule, the developed IgG rabbit antiserum (diluted 1:150 in blocking buffer) was absorbed with a proper concentration of the purified protein (200 ng/ml) for 2 h at 37°C. Then, the reactivity of rabbit antiserum was investigated on blots of FWAP and E/S products as described above.

**Gel electroelution and purity of the 27-kDa *Fasciola* antigen.** In preparative slab gel electrophoresis, the SDS-PAGE running condition was used to reduce smear gel electrophoresis, the SDS-PAGE running condition was adapted to reduce smear gel electrophoresis, and to enable a considerably long migration distance between bands in the 27-kDa region according to the prestained molecular mass marker (22). In each run, 250 µl of diluted human serum sample from an infected individual per preparative gel was electrophoresed, and about 35 runs were completed to obtain 1 mg of the 27-kDa *Fasciola* antigen. The protein content of a sample of electroeluted antigen was determined before remainder was stored at −20°C.

**Purity of the electroeluted *Fasciola* antigen.** The purity of the electroeluted 27-kDa *Fasciola* antigen was assessed using gel silver staining (32) and capillary zone electrophoresis (CZE) with a modification of the method described by Gordon et al. (33) using a autosampler (model 1-LIFT; Prince Technologies, Emmen, The Netherlands), a variable UV-visible detector (Lambda 1010; Metrhom, Herisau, Switzerland), and WinPrince software (version 5; Prince Technologies). The signals were analyzed using Dax software (version 5; Prince Technologies).

**Production of specific anti-27-kDa *Fasciola* antigen IgG antibody.** Specific IgG antibodies were produced in 4 New Zealand White rabbits immunized subcutaneously in three different inoculation sites with the purified 27-kDa *Fasciola* antigen according to the method described by Attallah et al. (22). In brief, equal volumes (500 µl) of the antigen (500 µg/ml) and complete Freund’s adjuvant (CFA) or incomplete Freund’s adjuvant (IFA) were homogenized together using two Luer-lock syringes connected to a three-way stainless steel valve. Each rabbit was immunized subcutaneously three times, once with antigen in CFA (on day 0) and twice with antigen in IFA (on days 15 and 28), before being sacrificed on day 32. Blood samples were collected from all rabbits 0, 28 and 32 days after immunization. Sera were separated and its immunoreactivity was tested using ELISA (see below) against *Fasciola* antigenic preparations, and the purified 27-kDa antigen. The specificity of the developed sera was tested using ELISA against adult *S. mansoni* and *A. lumbricoides* antigenic preparations. Sera of nonimmunized rabbits were tested in parallel. The highly reactive rabbit sera were then aliquoted and stored at −20°C until used.

**Immunohistochemical detection of native *Fasciola* antigen by using indirect immunoperoxidase.** A paraffin section (4 µm) of adult worm tissues was deparaffinized and rehydrated through descending grades of alcohols in water. The slides were washed in TBS 3 times for 5 min each. Excess liquid from around the section was wiped, and the slides were laid flat. Four to six drops of 3% hydrogen peroxide (H₂O₂) were applied for each slide, and slides were incubated for 5 min in a dark chamber. Normal goat serum was applied with a dilution of 1:5 with 3% bovine serum albumin (BSA) in TBS for 60 min. Then, the developed rabbit anti-27-kDa *Fasciola* antigen IgG antibody diluted 1:100 in TBS was applied and incubated for 30 min. After the slides were washed, horseradish peroxidase–conjugated goat antibody to rabbit immunoglobulins diluted 1:250 in 1% BSA–TBS was applied and incubated for 60 min. The slides were washed, and then tissue reacted with 3-amino-9-ethyl carbazole–H₂O₂ solution for 30 min in the dark. The reaction was stopped with distilled water. The sections were counterstained with Mayer’s hematoxylin for 2 min and washed in running tap water for 3 min to the desired intensity of blue color. The slides were dried out and applied onto coverslips using glycerol. All washings and incubations were performed at room temperature.

**Development of *F. gigantica* circulating antigen detection assay (FgCA-27 ELISA).** Serial dilutions (1:25 to 1:3200) of selected serum samples from patients with no parasite eggs (n = 4) and low (n = 4, <100 EPG), moderate (n = 4, 100 to 299 EPG), and high (n = 4, >300 EPG) parasite egg counts, in duplicate, were coated with carbonate-bicarbonate buffer pH 9.6, and were investigated to get the proper saturation of the polystyrene solid phase with the target antigen. Checkerboard titrations of the specific anti-27-kDa IgG antibody as well as alkaline phosphatase conjugate were also performed. After optimization of assay conditions, polystyrene flat-bottom microtiter plates (Costar, Acton, MA) were coated with 1:200-diluted human serum samples or serial concentrations (1,024

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to 0 ng/ml) of the purified 27-kDa Fasciola antigen diluted in normal human serum. In parallel and at 2.5 μg/ml, FWAP and E/S products were tested as positive controls and SWAP and EWAP were tested as negative controls for the reactivity of the developed rabbit antibody. After the plate was blocked with 0.3% BSA, 50 μl/well of rabbit anti-27-kDa antigen IgG antibody at a 1:400 dilution, in PBS with 0.05% (vol/vol) Tween 20 (PBS-T20), was added and the plates were incubated at 37°C for 2 h. After the plate was washed, 50 μl/well of anti-rabbit IgG–alkaline phosphatase conjugate (The Binding Site, Birmingham, United Kingdom) diluted 1:600 in 0.2% (wt/vol) BSA in PBS-T20 was added and the plate was incubated at 37°C for 1 h. After the plate was washed, the substrate (p-nitrophenyl phosphate in 0.1 M glycine buffer; pH 10.4) was added and the plates incubated for 30 min at 37°C.

Optical densities (OD) were read at 405 nm using a microplate auto-reader (2860; Metertech Inc., Taipei, Taiwan). The cutoff level of the FgCA-27 ELISA above or below which the tested sample is considered positive or negative was calculated as the mean OD for the FgCA-27 ELISA (range, 0.093 to 0.227) from a control group of 32 serum samples from healthy individuals with no parasitic infection plus 3 standard deviations (SDs) (i.e., cutoff level = 0.131 + [3 × 0.053] = 0.29). The mean OD value of a group of 16 sera from parasitologically diagnosed Fasciola gigantica-infected individuals and investigated in parallel was 0.755 (range, 0.394 to 1.985). The mean OD of another group of 16 serum samples of individuals parasitologically diagnosed as being not infected with Fasciola but having other parasitic infections, including schistosomiasis mansoni (n = 3), ascariasis (n = 3), hymenolepiasis (n = 3), giardiasis (n = 3), and entamoebiasis (n = 4), and investigated in parallel was 0.181 (range, 0.105 to 0.261). To establish the dose-response curve, serial concentrations (1 to 1,024 ng/ml) of the target 27-kDa antigen were mixed with diluted serum samples showing FgCA-27 ELISA-negative results for antigen detection and with no detected antibody levels (n = 3, OD < 0.25; cutoff level of an in-house ELISA based on specific anti-27-kDa IgG antibodies in human serum) and tested in duplicate as well as with selected serum samples with low (n = 3, OD = 0.25 to 0.49), moderate (n = 3, OD = 0.5 to 1.5), and high (n = 3, OD > 1.5) levels of specific anti-27-kDa IgG antibodies.

Statistical analyses. Descriptive results were expressed as means ± SDs and ranges or numbers of patients with a condition. Differences in continuous variables were assessed using Student’s t test or analysis of variance (ANOVA). All tests were two-tailed, and statistical significance was assessed at the 0.05 level. The diagnostic accuracy of the Fasciola antigen detection test was assessed by the area under the receiver operating characteristic (ROC) curve (AUC). An AUC equal to 1.0 is characteristic of an ideal test, whereas 0.5 indicates a test of no diagnostic value. The reproducibility and repeatability of the developed assay, calculated as intra-assay and interassay coefficients of variation (CV%), were assessed using 10 serum samples with different concentrations of Fasciola circulating antigen (17 to 1,200 ng/ml) tested in quadruplicate and undertaken along 4 consecutive weeks. All statistical analyses were done by a statistical software package (SPSS 15.0 for Microsoft Windows; SPSS Inc., Chicago, IL, USA).

RESULTS
Identification of the target Fasciola antigen among F. gigantica antigenic extracts and human sera from infected individuals. FWAP and E/S products of F. gigantica, SWAP of S. mansoni, and AWAP of A. lumbricoides were subjected to SDS-PAGE (Fig. 1A), and Western blotting was carried out to detect the target epitope of Fasciola antigen. It was found that a polypeptide band of 27 kDa in the FWAP and E/S product of Fasciola reacted with anti-27-kDa antigen IgG antibody. This antibody did not recognize any antigens in the Ascaris and Schistosoma antigenic extracts (Fig. 1B). No reaction against E/S products or FWAP using the saturated rabbit antiserum with the 27-kDa protein was shown (Fig. 1C).

The electroeluted 27-kDa antigen gave a single polypeptide band when investigated by SDS-PAGE and silver staining, and the purity of the eluted 27-kDa polypeptide was confirmed using CE; only a single peak (5.9 min) was observed (Fig. 1D). A sharp band was observed at 27 kDa in sera from individuals infected with F. gigantica, but no reaction was observed in sera of uninfected individuals (Fig. 2). Partial biochemical characterization of the reactive epitope confirmed its protein moeity. The reactivity of the anti-27-kDa antibody toward the purified antigen was maintained after periodate oxidation (i.e., a positive result using FgCA-27 ELISA). However, antibody reactivity toward the purified 27-kDa antigen decreased when the incubation time of the antigen with pepsin enzyme was increased, and it was completely lost at 20 min (i.e., a negative result using FgCA-27 ELISA).

Anatomic localization of the native target antigen in an F. gigantica adult worm. The antigenic determinant of the developed IgG antibody was located in the gut and tegumental surface of an adult F. gigantica worm. A strong immunoperoxidase reaction was found within the tegument, the muscularis, and the gut cells (Fig. 3A). No peroxidase reaction was shown within tegument, muscularis, and gut cells of a Fasciola adult worm using normal rabbit serum as a negative control (Fig. 3B).

Detection of Fasciola circulating antigen in human serum samples by using FgCA-27 ELISA. From the established calibration curve (Fig. 4A), the developed FgCA-27 ELISA successfully detected the 27-kDa Fasciola antigen serially diluted in normal human serum up to 1 ng/ml; the point did not overlap the concentration corresponding to 3 SDs above the mean OD of the zero calibrator (n = 4). However, the detection limit of the target circulating antigen in human serum samples at 8 ng/ml corresponds to the cutoff level (OD = 0.29) of the developed assay (n = 32). The presence of different levels of specific anti-27-kDa IgG antibodies in serum decreased the antigen levels but did not reach a significant level of difference (P > 0.05) in comparison with normal human serum, i.e., negative for specific IgG antibodies to Fasciola (data not shown). Serum samples from 120 individuals with fascioliasis and 80 noninfected individuals were tested using FgCA-27 ELISA for the detection of circulating Fasciola antigen. The antigen concentrations (ng/ml) in serum samples are shown in Fig. 4B. The AUC for the Fasciola antigen detection assay was significantly high (AUC = 0.961, P < 0.0001) for discriminating between individuals infected with Fasciola and noninfected individuals (Fig. 5A). The developed assay had a sensitivity of 93.3% and had false-positive results for 4 of 80 noninfected individuals, with a specificity of 95% in detecting human fascioliasis (Table 1).

Cross-reactivity with other parasitic infections was studied based on microscopic stool examinations of 80 noninfected individuals. The FgCA-27 ELISA showed specificities of 100% with S. mansoni (7 cases), 100% with A. lumbricoides (5 cases), 100% with H. nana (4 cases), 100% with G. lamblia (3 cases), 92% with E. histolytica (3 out of 38 cases showing false-positive result), and 96% with parasite-free healthy individuals (1 out of 23 cases showing a false-positive result) (Table 1). All serum samples showing false-negative and false-positive results by FgCA-27 ELISA were investigated using the more-sensitive Western blot analysis. The 27-kDa antigen was identified in all 8 sera showing false-negative ELISA results. Among the four sera showing false-negative ELISA results, only one sample from a parasite-free healthy individual showed a sharp band at 27-kDa, and the remaining 3 sera of E. histolytica-infected individuals showed no bands in their blots. Moreover,
antigen detection rates (percent positivity) increased with increasing severity of infection (expressed as parasite egg count [EPG]) (Table 1), and a highly significant correlation (Spearman $r = 0.715$, $P < 0.0001$) between circulating Fasciola antigen level in serum (ng/ml) and EPG was shown (Fig. 5B). The percent intraassay and interassay coefficients of variation (CV%) in antigen concentrations were 3.3% and 5.8%, respectively.

DISCUSSION

During the last 2 decades, a major focus of research has been directed toward the identification of Fasciola antigens during human infection as a step toward the development of an efficient diagnostic assay (34). Fasciola antigens are mostly released from the gut and the tegument into the blood circulation and excreted in stools of infected animals and humans (35–39). Several Fasciola antigens have been detected as circulating antigens in serum or as coproantigens in feces and were successfully used in immunodiagnosis of human fascioliasis (8, 11, 16, 34, 40). Of these, one antigenic component with an approximate molecular mass of 27 kDa was found to give a consistent reaction with human fascioliasis sera (14, 18, 41–43). Of note, all serologic tests based on the 27-kDa antigen were developed and optimized with an emphasis on the detection of antibodies to F. hepatica (12, 13, 44, 45). However, the direct detection of Fasciola antigens secreted by the living

FIG 1 Identification and purification of the target Fasciola antigen. (A) Silver-stained SDS-PAGE. (B) Western blot for different antigenic sources of F. gigantica, S. mansoni, and A. lumbricoides using rabbit anti-27-kDa Fasciola IgG antibody. (A and B) Lane 1, F. gigantica adult worm antigen preparation (FWAP); lane 2, excretory/secretory (E/S) products from F. gigantica; lane 3, S. mansoni adult worm antigenic preparation (SWAP); lane 4, A. lumbricoides adult worm antigenic preparation (AWAP); lane 5, the purified 27-kDa antigen. The developed anti-Fasciola antibody identified its target epitope at 27 kDa in FWAP, E/S products, and the purified antigen (lanes 1, 2, and 5) but not in SWAP and AWAP (lanes 3 and 4). (C) Inhibition Western blot for different antigenic sources of F. gigantica by using rabbit anti-27-kDa Fasciola IgG antibody saturated with purified 27-kDa antigen. Lane 1, FWAP immunostained with the anti-Fasciola antibody; lane 2, FWAP immunostained with the saturated antibody; lane 3, E/S products immunostained with the anti-Fasciola antibody; lane 4, E/S products immunostained with saturated antibody. The saturated anti-Fasciola antibody did not identify its target epitope at 27 kDa in FWAP and E/S products (lanes 2 and 4). Molecular mass standards (St.) included were phosphorylase B (97.4 kDa), bovine serum albumin (66.2 kDa), glutamate dehydrogenase (55 kDa), ovalbumin (42.7 kDa), aldolase (40 kDa), carbonic anhydrase (31 kDa), and soybean trypsin inhibitor (21.5 kDa). (D) The 27-kDa antigen purified from serum of an F. gigantica-infected individual showing a single peak, at an absorbance of 200 nm, at 5.9 min, using CZE.

FIG 2 Western blot of serum samples from Fasciola-infected and noninfected individuals for the detection of circulating Fasciola antigen. Lane 1, FWAP; lanes 2 to 4, serum samples from 3 noninfected individuals; lanes 5 to 9, serum samples from 5 individuals infected with F. gigantica. Fifty micrograms/lane of each sample was separated on 12% acrylamide gels, transferred to an NC sheet, and reacted with 100 µg/ml rabbit anti-27-kDa IgG antibody. Anti-rabbit IgG-alkaline phosphatase and BCIP (5-bromo-4-chloro-3-indolylphosphate)-nitroblue tetrazolium (NBT) substrate were used to visualize the reaction products. The developed rabbit antibody identified a 27-kDa antigen in all sera of infected individuals (lanes 5 to 9) but not in sera of noninfected individuals (lanes 2 to 4). Molecular mass standards included were phosphorylase B (97.4 kDa), bovine serum albumin (66.2 kDa), glutamate dehydrogenase (55 kDa), ovalbumin (42.7 kDa), aldolase (40 kDa), carbonic anhydrase (31 kDa), and soybean trypsin inhibitor (21.5 kDa).
flukes has apparent advantages over antibody detection tests, in that antigenemia implies active infection, and this approach has the ability to assess efficacy of chemotherapy and determine the effectiveness of future vaccines (24, 46–48).

In our previous study (22), we identified a wide band of 26 to 28 kDa in adult worm and E/S antigenic preparations of *F. gigantica* and also in sera of cattle infected with *F. gigantica* by using specific IgG antisera developed in rabbit to the target 26- to 28-kDa antigen purified from FWAP. This antibody did not recognize any target epitopes in the *Ascaris* and *Schistosoma* antigenic extracts, and its antigenic determinant was located in the guts and teguments of adult *F. gigantica* worms. This enhanced our efforts to identify the target antigen of the specific IgG antibody to the 26- to 28-kDa *Fasciola* antigen in sera of infected cattle infected with *F. gigantica*. We fortunately observed a sharp narrow band at 27 kDa in selected sera from individuals infected with *F. gigantica*, and no specific reaction was observed in sera of uninfected individuals via Western blot analysis (unpublished data).

In the present study, the target 27-kDa circulating antigen was isolated and purified from serum samples from *F. gigantica*-infected humans, and a highly reactive IgG antibody to the purified 27-kDa antigen was developed in rabbit. The newly developed IgG antibody identified a polypeptide band of 27 kDa in adult worm and E/S antigenic preparations of *F. gigantica* but did not recognize any target epitopes in the *Ascaris* and *Schistosoma* antigenic extracts, and its antigenic determinant was located in the guts and teguments of adult *F. gigantica* worms. Moreover, the target circulating antigen was identified at 27 kDa in sera from individuals infected with *F. gigantica*, and no specific reaction was observed in sera of uninfected individuals using Western blotting or ELISA. An inhibition-Western blot assay was performed to confirm that the 27-kDa protein isolated from serum is a parasite molecule. Our antigen is similar in molecular mass to an *F. gigantica* antigen designated FG27 (42, 49). The partial biochemical characterization of our purified antigen confirmed its protein moiety. However, further molecular characterization of our 27-kDa target antigen, including peptide mapping followed by mass spectrometry (MS) amino acid sequence will be performed in collaboration with a well-equipped laboratory to clarify its identity.

The identification of the *Fasciola* antigens in different FWAPs and E/S products and in human serum is of crucial importance for reliable diagnosis of active *Fasciola* infection. Several sensitive and specific methods based on *Fasciola* antigen detection in serum and stool using specific antibodies were applied to *F. hepatica* and *F. gigantica* infections in animals (8, 25, 26, 47). The antigen tests can detect experimental infection a few days after inoculation (23). However, few antigen detection assays have been developed for detection of *F. gigantica* in human fluids, and these have varied ranges of sensitivities and specificities (21). In the present study, we have developed FgCA-27 ELISA, which successfully detected the 27-kDa *Fasciola* antigen with a detection limit of 8 ng/ml in human serum samples. The presence of *Fasciola* antigens in infected sera of infected individuals via the circulation of infected individuals may be attributed to the continuous shedding and release of circulating antigens as a possible escape mechanism by the parasites, to reinfections, and to exposure to maternal antigens that may lead to developing tolerance rather than immunity to free antigens of natural infections (50). Moreover, the presence of high titers of specific IgG antibodies in the sera of infected individuals does not indicate a complete elimination of the pathogen; such high titers of antibodies may potentially neutralize all of the circulating antigens, i.e., via immune complex formation, and also, the specific IgM antibody response will not be stimulated for new infections. The presence of different levels of specific anti-27 kDa IgG antibodies in serum samples did not affect the detection of different concentrations of the target *Fasclola* antigen using the optimized FgCA-27 ELISA. A nonsignificant decrease (P > 0.05) in the antigen concentrations with the increase of anti-27-kDa IgG antibody levels in serum was shown (data not shown). This indicates that the 27-kDa antigen could attach to the solid phase in the presence of specific IgG antibodies or other serum proteins.

The performance characteristics of the developed assay were investigated using sera of 200 parasitologically diagnosed individuals. The developed assay gave high degrees of sensitivity, specificity, and efficiency (>93%). The accuracy of the developed ELISA, calculated by the area under the ROC curve (AUC), yielded a 0.961 value, indicating a high performance of the assay that can significantly (P < 0.0001) discriminate *F. gigantica*-infected and noninfected individuals. Moreover, for the precision of the test, the calculated intra-assay and interassay coefficients of variation were 3.3% and 5.8%, respectively, indicating also high performance in reproducibility and repeatability of the assay. Interestingly, our developed assay shows results that are superior or at least similar to those of the previously reported antigen tests for...
diagnosis of human fascioliasis (37, 51, 52), and the differences may be due to the different natures of antigens used. A recent study evaluated an antigen capture ELISA using a pair of monoclonal antibodies raised against *F. gigantica* E/S antigens in human serum and stools at a lower detection limit of 3 ng/ml (25). The sensitivity (94%) and specificity (94.6%) of the capture ELISA in serum were similar to those shown in our study. Another more recent study showed a similar degree of sensitivity (94.5%) but a lower specificity (84.6%) using sandwich ELISA based on monospecific rabbit IgG antibody to 14.5-kDa protein antigen obtained from *F. gigantica* adult worms (26). Protein-based tests (e.g., protein microarrays) as well as DNA-based molecular tests (including PCR) can be used for clinical detections as well as field screening (53). Further investigations comparing the immunodiagnostic performances of our FgCA-27 ELISA with such tests will be performed in collaboration with a well-equipped laboratory.

FIG 4 Quantification of the 27-kDa *Fasciola gigantica* antigen in human serum by using the developed FgCA-27 ELISA. (A) Calibration curve of the 27-kDa *Fasciola* antigen. The purified 27-kDa antigen (ng/ml) was serially diluted in normal human serum and tested using the developed ELISA. The optical densities (OD) were measured at 405 nm, and the cutoff level was set at an OD of 0.29. (B) Scattergram showing *Fasciola* antigen levels (expressed as ng/ml) in sera of 120 parasitologically diagnosed infected individuals whose infections were classified as light (<100 EPG; *n* = 76), moderate (100 to 299 EPG; *n* = 33), and severe (≥300 EPG; *n* = 11) in comparison with antigen levels in sera of 80 noninfected individuals. The cutoff level of the developed ELISA was set at 8 ng/ml.

FIG 5 Diagnostic accuracy of the developed FgCA-27 ELISA. (A) Receiver operating characteristic (ROC) curve of circulating *Fasciola* antigen detection using ELISA for discriminating *F. gigantica*-infected individuals and noninfected individuals. The true-positive rate (sensitivity) is plotted as a function of the false-positive rate (specificity). Each point on the ROC curve represents a sensitivity/specificity pair corresponding to a particular decision threshold. The area under the curve (AUC) value represents the combined effects of both sensitivity and specificity of circulating *Fasciola* antigen in diagnosing individuals infected with *F. gigantica* (AUC = 0.961, *p* < 0.0001). (B) Correlation between the levels of *Fasciola* circulating antigen measured in human sera by ELISA (expressed in log_{10} ng/ml) and the egg count measured by Kato-Katz technique (expressed in EPG). The results of both assays for 120 serum samples were statistically analyzed and showed a highly significant correlation (Spearman *r* = 0.715, *p* < 0.0001).
In the present study, all serum samples showing false-negative ELISA results, except for one (which was of a moderate parasite burden [i.e., egg count of 100 to 299 EPG]), corresponded to light infection (i.e., egg count < 100 EPG); however, the target 27-kDa antigen was identified in all these samples by using the more sensitive Western blot technique. This result may support the idea of immune complex formation in the samples showing false-negative ELISA results. Therefore, the absence of the target 27-kDa antigen in stool as measured by ELISA, as the egg excretion is probably more erratic in patients with chronic infections. However, the presence of a highly significant correlation between circulating Fasciola antigen levels and parasite egg count which is presumably dependent on the number of flukes in the host is of crucial importance to the use of the developed assay to monitor the efficiency of flukicide treatment in Fasciola-infected individuals and to assess potential new vaccine efficacy.

In conclusion, a 27-kDa Fasciola antigen was identified in sera of F. gigantica-infected individuals. A highly sensitive and specific Fasciola circulating antigen detection assay, FgCA-27 ELISA, was developed for reliable laboratory diagnosis of human fascioliasis. Further characterization of the target antigen will be performed, and further optimization and field studies are necessary to draw a final conclusion regarding the use of FgCA-27 ELISA for diagnosis and follow-up for Fasciola-infected individuals.

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**REFERENCES**


