Dot-Blot Immunoassay of *Fasciola gigantica* Infection using 27 kDa and Adult Worm Regurge Antigens in Egyptian Patients

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Abstract: The purpose of the present study was to evaluate the potential role of the 27-Kilodalton (KDa) antigen versus *Fasciola gigantica* adult worm regurge antigens in a DOT-Blot assay and to assess this assay as a practical tool for diagnosis fascioliasis in Egyptian patients. *Fasciola gigantica* antigen of an approximate molecular mass 27- (KDa) was obtained from adult worms by a simple elution SDS-PAGE. A Dot–Blot was developed comparatively to adult worm regurge antigens for the detection of specific antibodies from patients infected with *F. gigantica* in Egypt. Control sera were obtained from patients with other parasitic infections and healthy volunteers to assess the test and compare between the antigens. The sensitivity, specificity, positive and negative predictive values of Dot- Blot using the adult worm regurge were 80%, 90%, 94.1%, and 69.2% respectively, while those using 27-KDa were 100% which confirms the diagnostic potential of this antigen. All patients infected with *Fasciola* were positive, with cross reactivity reported with *Schistosoma mansoni* serum samples. This 27-KDa Dot-Blot assay showed to be a promising test which can be used for serodiagnosis of fascioliasis in Egyptian patients especially, those presenting with hepatic disease. It is specific, sensitive and easy to perform method for the rapid diagnosis particularly when more complex laboratory tests are unavailable.

Key words: *Fasciola gigantica*, 27-KDa, elution SDS-PAGE, dot-blot

### INTRODUCTION

Fascioliasis is a parasitic infection of great concern, not only because of its high prevalence and economic significance to the animal stock [1,2] but also because it appears as an emerging public health problem in many regions of the world [3]. It is estimated that 2.4 million people are infected worldwide and that the number of people at risk is more than 180 million [4]. *Fasciola hepatica* is predominating in temperate zones, while *Fasciola gigantica* is found mainly in tropical and subtropical regions of Asia and Africa [5]. Fascioliasis is an example of an emerging and re-emerging parasitic disease in many countries because of environmental changes as well as man-made modifications [6].

Diagnosis of fascioliasis by finding eggs in stool samples lacks sensitivity, as the eggs are absent in acute and ectopic stages and intermittently released in the chronic stage. Therefore, coprological and immunological methods have to be used to confirm suspected cases [7]. Specific antigens in the somatic extract and excretory-secretory (ES) product of *F. gigantica* adults were identified and characterized by SDS-PAGE and immunoblotting analysis [8]. An antigenic component from the adult ES product with an approximate molecular mass of 27 kDa was found to give a consistent reaction with human fascioliasis sera up to 100% sensitivity and 98% specificity [8].

The purpose of the present study was to isolate the 27 kDa antigen using a simple elution method to evaluate its potential role versus *Fasciola gigantica* adult worm regurge antigens in a dot-blot assay and to assess this assay as a practical tool for diagnosis of fascioliasis in Egyptian patients.

### MATERIALS AND METHODS

#### Worm collection

Adult *F. gigantica* flukes were collected from the bile ducts of naturally infected cattle from Cairo abattoir, Egypt. About 37 flukes were used for the intestinal content preparation [9].
Collection of intestinal content as adult worm regurge

The antigens were obtained as described previously [10]. Briefly, worms were washed 3 times in 0.9 M NaCl at room temperature until the supernatant became clear. The supernatant was placed in a Petri dish with fresh cold 0.9 M NaCl and incubated at 4˚C for 45 min. The supernatant was discarded and the worms were then incubated at 37˚C for 30 min in 0.9 M NaCl. The supernatant with evident black intestinal content was collected. The collected material was centrifuged at 12,000 g for 30 min, and the supernatant was concentrated osmotically using dialysis membrane (Sigma-Aldrich, St. Louis, Missouri, USA) immersed in polyethylene glycol. The concentrated material was aliquoted and stored at -70˚C until use.

SDS-PAGE and elution of the 27 kDa antigen

SDS-PAGE was carried out as described previously [11]. SDS-PAGE in 12% gel was carried using *F. gigantica* adult worm regurge antigens with a concentration of (0.6 mg/ml) and the electrophoretic run took place using the prestained wide range Bio-Rad marker to detect precisely the desired 27 kDa band (Fig. 1). A comb containing 3 lanes was used where the marker was applied to the 1st lane, and the antigen to the 2nd and 3rd lanes. The sample compromised 300 µl of the antigen was added to 300 µl sample buffer from which only 40 µl were added to the 3rd lane, and the rest applied to the 2nd lane. After cutting the stacking gel, the 3rd lane, which was used as a reference to locate the position of the desired protein band on the gel in the 2nd lane, was cut gently and stained in Coomassie blue then destained. It was then compared with the marker. As the destain shrinks the gel, because of its acetic acid massie blue then destained. It was then compared with the marker of the grid through minimizing the area that the solution penetrates (3-4 mm diameter) by applying it slowly. The gel was spotted onto the nitrocellulose membrane at the center of the grid through minimizing the area that the solution penetrates (3-4 mm diameter) by applying it slowly. The membrane was dried. Non-specific sites were blocked by soaking in 5% skimmed milk in PBS-T (5 g milk + 100 ml PBS, pH 7.2 + 30 µl Tween 20) in a 10 cm Petri dish for reaction chamber, then incubated with primary antibody from serum samples diluted to 1:50 and dissolved in 5 % skimmed milk in PBS-T for 1 hr at room temperature, washed 3 times with PBS-T 5 min each. Then, it was incubated with the secondary anti-

derived material was stored at -70˚C until use. before measuring the protein content.

**Determination of the protein content**

Before antigen storage, the protein content was measured [13]. For the *Fasciola* adult worm regurge antigen, the protein concentration was 0.6 mg/ml, and for the 27 kDa band, the protein concentration was 0.06 mg/ml.

**Collection of test and control sera**

Serum samples required for study were separated from blood samples collected from cases referred to the Diagnostic and Research Unit, Parasitology Department, Faculty of Medicine, Ain Shams University, Cairo, Egypt. The study protocol was approved by the Scientific-Ethics Committee of Ain Shams University. Informed consent was obtained from the study subjects using standard guidelines. Whole blood was collected from each subject and centrifuged at 760 g at 4˚C for 10 min and the obtained serum samples were stored at -70˚C until use.

These serum samples were categorized into group I (20 patients with patent fascioliasis passing *Fasciola* eggs in their stool samples and negative for other parasitic infections), group II (10 patients negative for fascioliasis and positive for other parasitic infections by indirect hemagglutination test, IHAT) (3 cases of schistosomiasis mansoni, 3 hydatidosis, and 4 toxoplasmosis), and group III (normal controls that included 10 healthy individuals, free from fascioliasis by IHAT and other parasitic infections by stool examination).

**Dot-blot assay**

Dot-blot assay was applied with some modifications [14] comparatively using *Fasciola* adult worm regurge antigens and the eluted 27 kDa soluble antigen. A grid is drawn by a pencil on the nitrocellulose membrane (Bio-Rad, Hercules, California, USA) to indicate the region to be blotted. Two µl of antigen was spotted onto the nitrocellulose membrane at the center of the grid through minimizing the area that the solution penetrates (3-4 mm diameter) by applying it slowly. The membrane was dried. Non-specific sites were blocked by soaking in 5% skimmed milk in PBS-T (5 g milk + 100 ml PBS, pH 7.2 + 30 µl Tween 20) in a 10 cm Petri dish for reaction chamber, then incubated with primary antibody from serum samples diluted to 1:50 and dissolved in 5 % skimmed milk in PBS-T for 1 hr at room temperature, washed 3 times with PBS-T 5 min each. Then, it was incubated with the secondary anti-
body conjugated with peroxidase-labeled affinity purified anti-human IgG (KPL, Washington DC, Maryland, USA) and diluted to 1 µl: 1,000 µl in PBS-T followed by incubation 1 hr with shaking, washing 2 times with PBS-T 2 min each, and then once with PBS for 5 min. The colour reaction was developed with DAB (5 g from 3,3’-diaminobenzilidine dissolved in 100 ml dH₂O and then aliquoted 500 µl in each tube)/H₂O₂ substrate solution (500 µl of DAB + 50 ml PBS+ 5 µl H₂O₂) for 2 min and stopped by several dH₂O washing. Sera positive for F. gigantica infection showed purple dots on the membrane, while negative sera showed no dots.

Data management and analysis

The chi-square test was used to examine the relationship between 2 qualitative variables. Fisher’s exact test was used to examine the relationship between 2 qualitative variables when the expected count was less than 5 in more than 20% of cells. McNemar test was used to assess the statistical significance of the difference between a qualitative variable measured twice for the same study group. The P-values set for the level of significance were $P>0.05$ (not significant; NS), $P<0.05$ (significant; S), and $P<0.01$ (highly significant; HS).

**RESULTS**

The results showed that, using the adult worm regurgate antigen (antigen I) against serum of diseased patients in the dot-blot assay, 16 out of the 20 patients gave positive results with a sensitivity of 80%. Whereas, using the 27 KDa antigen (antigen II) in the dot-blot assay, all the diseased patients gave positive results with a sensitivity of 100% (Table 1).

The dot-blot assay using antigen I among the negative control cases gave 9 negative results out of the 10 negative controls compared to 10 negative results using antigen II, showing a specificity of antigen I of 90 % compared to 100% specificity using antigen II (Table 1). The results also showed that, using antigen I among positive controls (patients positive for other parasitic infections), 7 cases were negative with a specificity of 70% compared to 9 negative cases among 10 using antigen II with a specificity of 90% (Table 1).

The dot-blot assay done to compare between diseased patients and negative controls for antigen I showed that there was a highly significant difference between them as the test gave positive results in 80% of the diseased cases compared to only 10 % in negative controls (Table 2).

**Table 2.** Comparison of diseased cases to positive and to negative controls using antigen I and antigen II by dot-blot assay

| Disease cases | Negative control | P-value | Significance |
|---------------|-----------------|--------|--------------|
| Antigen I     |                 |        |              |
| Negative      | 4               | 20     | 0.0001       | HS            |
| Positive      | 16              | 80     |               |               |
| Antigen II    |                 |        |              |
| Negative      | 0               | 0      | 0.0001       | HS            |
| Positive      | 20              | 100    |               |               |

The comparison between diseased cases and negative controls for antigen I revealed 94.1% PPV (positive predictive value) and 69.2% NPV (negative predictive value). The comparison between diseased cases and negative controls for antigen II revealed 100% PPV and 100% NPV. The comparison between diseased cases and positive controls for antigen I revealed 84.2% (16/19) PPV and 63.6% (7/11) NPV. The comparison between diseased cases and positive controls for antigen II revealed 95.2% (20/21) PPV and 100% (9/9) NPV.

*Fisher’s exact test.*

**Table 1.** Comparison between antigen I and antigen II as regard to sensitivity for disease diagnosis, specificity among positive controls, and specificity among negative controls with other parasitic diseases using dot-blot assay

| Disease cases | Positive parasitic control | Negative control |
|---------------|----------------------------|------------------|
|               | Positive                   | Negative         |
|               |                           |                  |
| Positive      |                           |                  |
| Antigen I     |                           |                  |
| Negative      | 4                          | 20               |
| Positive      | 16                         | 80               |
| Antigen II    |                           |                  |
| Negative      | 20                         | 100              |
| Positive      | 16                         | 80               |
| $P$-value*    | 0.125                      | 0.5              |
| Significance  | NS                         | NS               |

The sensitivity detected against Fasciola gigantica positive serum samples was 80% for antigen I compared to 100% for antigen II. The specificity detected against negative controls (no parasitic infection) was 90% for antigen I compared to 100% for antigen II. The specificity detected against other parasite positive samples was 70% for antigen I compared to 90% for antigen II.

*McNemar test.*
diseased patients and negative controls for antigen II showed a highly significant difference between both groups as it gave positive results in 100% of diseased group compared to 0% in normal controls (Table 2).

In comparison of diseased cases and positive controls with antigen I, the results showed a significant difference between both groups as it was positive in 80% in diseased cases compared to 30% in positive controls (Table 2). The assay done using antigen II to compare diseased cases and positive controls showed a highly significant difference between the 2 groups as it was positive in 100% of diseased cases compared to 10% in positive controls (Table 2).

DISCUSSION

In humans, fascioliasis is becoming an increasingly important health problem with cases reported from South America, Europe, Africa, Australia, and the Far East [6,15,16]. In endemic areas, clinical examination and imaging techniques are used for the diagnosis of fascioliasis, but these methods cannot always differentiate between fascioliasis and other causes of hepatitis [16]. Therefore, coprological and immunological methods have to be used to confirm suspected cases of fascioliasis. However, diagnosis by coprological examination is complicated by the long prepatent period of \textit{Fasciola} spp., during which no eggs are shed, and by the fact that in humans not all flukes develop to adult egg-laying worms or flukes migrate to non-hepatic areas [7]. Moreover, \textit{Fasciola} eggs may be found in the stools of uninfected persons who have eaten raw infected livers leading to false positive diagnosis. Many factors impose the idea of evaluating and establishing serological methods for the diagnosis of fascioliasis, which allow diagnosis of the disease even in the acute stage, before the parasite eggs can be identified in feces [17].

The immunological techniques offer an interesting option for the diagnosis of fascioliasis [18,19,20]. Among these tests, ELISA has been developed using ES antigens from adult flukes, purified antigens, or recombinant antigens [8,21,22]. An ELISA using the 27 kDa native protein isolated from ES products of \textit{Fasciola} spp. has shown high sensitivity and specificity in the diagnosis of human fascioliasis [23]. Several groups have reported the purification of the immunodominant antigens from the ES products of the liver fluke [24,25]. Of these, only one group described the use of a purified 27 kDa cysteine proteinase from Japanese \textit{Fasciola} spp. and its evaluation as an antigen in the diagnosis of human disease [25]. This proteinase demonstrated a high sensitivity and specificity in ELISA for human fascioliasis, showing only 1 case of cross-reactivity with sera from schistosomiasis japonicum patients. The purified fractions Fas 1 and Fas 2, which contain a cysteine proteinase of \textit{F. hepatica} with an approximate molecular mass of 25-26

![Fig. 1. Characterization of 27-kDa band from Fasciola adult worm regurge separated by SDS-PAGE in 12% gel guided by wide range Bio-Rad marker. Lane 1. Molecular weight standard in kDa. Lane 3. Fractionated adult worm regurge antigens. Lanes are named according to the protocol of materials and methods. Arrow indicates 27 kDa band.](image1)

![Fig. 2. Dot-blot showing reactivity of 20 human samples infected with \textit{F. gigantica} (1st figure), 10 human samples infected with other parasites (positive control samples); samples no.1, 2, 4 (hydatidosis by IHAT), samples no 3,8,10 (S. mansoni infection by IHAT), samples no 5, 6, 7, 9 (toxoplasmosis by IHAT) (2nd figure), and 10 human samples with no parasitic infections (negative controls) (3rd figure) against antigen I (adult worm regurge) and antigen II (27 kDa). Negative control lane shows positive kit guide for colour results, so if a same colour appears it is a negative result. Positive control lane shows positive kit guide for colour results, so if a same colour appears it is a positive result.](image2)
kDa, are specific and sensitive antigens [26].

In the present study, the dot-blot assay using *F. gigantica* adult worm regurgite antigens (antigen I) vs the partially purified fractions of the antigens with an approximate molecular mass of 27 kDa (antigen II) obtained by continuous elution SDS-PAGE, were used in a trial to evaluate their potential role in the diagnosis of fascioliasis. The sensitivity, specificity, positive and negative predictive values using the adult worm regurgite (antigen I) were 80%, 90%, 94.1%, and 69.2%, respectively (Table 2; Fig. 2), while those using 27 kDa (antigen II) were 100% (Table 2; Fig. 2), which confirmed the diagnostic potential of this antigen.

All patients infected with *Fasciola* were positive, with cross-reactivity reported only in 1 schistosomiasis mansoni serum sample. There was significant difference between diseased cases and positive controls in regard to the results of antigen I test, as it was positive in 80% of the diseased group compared to 30% in the positive controls; PPV = 84.2% and NIV = 63.6% (Table 2; Fig. 2). There was a highly significant difference between diseased cases and positive controls in regard to the results of antigen II test, as it was positive in 100% of the diseased group compared to 10% in positive controls; PPV = 95.2% and NIV = 100% (Table 2; Fig. 2).

The cross-reactivity with schistosomiasis was previously reported [27]. This cross-reactivity was explained by the presence of cross-reactive epitopes between *Fasciola* ES products and *S. mansoni*. There was a relatively high cross-reactivity (25.9%) with the 27 kDa protein ELISA in serum samples from cholangiocarcinoma patients in Thailand [8]. Those patients might have been infected with *Opisthorchis viverrini* in association with cholangiocarcinoma development [28]. In another study, 2 out of 3 serum samples from *Clonorchis sinensis* infections gave a false positive result confirming that cross reactions with this parasite should be taken into account [29]. However, in contrast to fascioliasis, clinical manifestations of clonorchiasis and opisthorchiasis occur mainly in the chronic phase of the infection and are usually associated with fecal egg excretion.

There were cross reactions between *Fasciola* crude worm antigen and patients with schistosomiasis mansoni, toxoplasmosis, and ascariasis by western blot technique [30]. Two normal human subjects showed also false reactivity. The presence of cross reactions in sera of normal individuals may be explained by previous subclinical infections with *Fasciola* or other parasites that share *Fasciola* in these cross reactive epitopes or may be due to the presence of a common antigenic band in *Fasciola*

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