

Assessment of The Diagnostic Performance of Normal and Irradiated *Schistosoma mansoni* Cercarial Antigens using ELISA

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ABSTRACT

Irradiation of *Schistosoma mansoni* cercariae induces structural and motor changes in its surface molecules converting it from poor immunogenic to highly immunogenic surface. This study aims to comparatively analyze normal and gamma radiation-attenuated *Schistosoma mansoni* cercarial antigens (SCA) by ELISA for serodiagnosis. Total IgG ELISA was assessed to evaluate the potency of two parts of *Schistosoma mansoni* cercarial antigens (2000 each) one normal and other exposed to attenuating dose of 400 Gy gamma radiations. The study was conducted on fifty two individual sera samples divided as follows: Group I included 21 obtained from patients who were positive for schistosomiasis and negative for other parasitic infections; Group II included 18 sera from patients who were negative for schistosomiasis and positive for other parasitic infections; (3) fascioliasis, (5) amoebiasis, (7) toxoplasmosis and (3) hydatidosis and Group III included (13) sera from healthy individuals all were diagnosed parasitologically and serologically. Although both antigens achieved 100 % sensitivity and negative predictive value (NPP), irradiated SCA revealed better specificity, accuracy and positive predictive value (PPV) of 87 %, 84% and 92.3% respectively. While normal SCA revealed 83.9%, 90.4 % and 80.7 % respectively. It was concluded that irradiated SCA achieved better remarkable ODs than normal SCA and better specificity due to low cross-reactivity with other parasites. SO it can be used as a tool for diagnosis of chronic schistosomiasis.

Article Information

Received 01 March 2019

Revised 16 March 2019

Accepted 19 November 2020

Available online 25 February 2021

Authors' Contribution

MME and SAHS designed the research. MME, ENH and MADA performed experiments and contributed in manuscript writing. MME, NIT and BEWE collected the samples and analyzed the data.

Key words

Schistosoma mansoni, Gamma radiation, Cercarial antigen, ELISA, Serodiagnosis

INTRODUCTION

Schistosomiasis is a tropical disease where induction of protective immunity and elimination can be achieved by sustainable surveillance and implantation of a sensitive diagnostic method for control (Amin *et al.*, 2017). Currently used diagnostics as Kato-Katz fecal smear lack sensitivity to accurately determine the prevalence of it in areas with low infection as it is affected by day and day variation in egg excretion (He *et al.*, 2018).

One of advanced strategies to diagnose atypical forms of *Schistosoma* is detection of antibodies which can be used as biomarker of progression of the disease (Ludolf *et al.*, 2014). However, persistence of antibodies in the host makes these tests limited in detection of early infection and in assessing response to treatment so a need for more

sensitive assays is highly desirable (Siqueira *et al.*, 2011; Cavalcanti *et al.*, 2013). The association of antibody and antigen detection assays potentiates the diagnosis. Enzyme linked immunosorbent assay (ELISA), indirect haemagglutination, or immunofluorescence are the most current serological tests to detect antibodies against schistosomal antigens. ELISA was chosen to screen and diagnose patients in residents of a low endemic area (Cristhiany *et al.*, 2012; Turner *et al.*, 2004). It was reported that the use of adult worm antigens (AWA) ELISA for detection of antibodies presents low specificity and cross interaction with other helminthes as hookworms. Also, the use of soluble egg antigen (SEA) ELISA has been shown to cross-react with *Fasciola* and it needs to be harvested from infected laboratory animals (Abdel-Fattah *et al.*, 2011; Chand *et al.*, 2010).

Schistosoma cercarial antigens (SCA) are easier than eggs and adult worm to produce and tests based on it are easier to scale up. Also, cercarial antibodies have been found to be at higher titers in early infection and produce far fewer false-positive results as it lacks epitopes responsible

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for cross reaction with other parasites (Knudsen *et al.*, 2005; Sarhan *et al.*, 2014).

Ionizing radiation was found to have various effects on helminthic parasites. Depending on the dose irradiated, parasites may be attenuated or died. It was reported that attenuation modifies expression of surface antigens and results in greater exposure of them converting the parasite into highly immunogenic however no biochemical or morphological changes can be demonstrated (Wuhrer *et al.*, 1999). Radiation attenuated *Schistosoma mansoni* infective stage can be applied for control measures by immunoprotection or immunodiagnosis. Previous studies reported that vaccination with irradiation stimulates host immune response against challenge infection (Hewitson *et al.*, 2005).

This study aims to comparatively evaluate normal and gamma-irradiated *S. mansoni* cercarial antigen by ELISA for serodiagnoses of chronic *Schistosoma mansoni* infected patients.

MATERIAL AND METHODS

Sample collection and sera preparation

Blood samples were collected from 52 individuals and classified into three groups. Group I included 21 patients who were positive for schistosomiasis from outpatient clinics of Menoufia University Hospitals and Mansoura University Hospitals, Egypt and were negative for other parasitic infections confirmed by examining stool samples by Kato-Katz technique. Group II included 18 patients who were negative for schistosomiasis and positive for other parasitic infections; (3) fascioliasis, (5) amoebiasis, (7) toxoplasmosis and (3) hydatidosis confirmed by parasitological and serological diagnosis. Group III included 13 healthy individuals (negative control). From each individual enrolled in this work, 5 ml blood was taken, centrifuged at 1000 xg for 5 min. After that, the serum was withdrawn, aliquoted in 0.5 ml eppendorf tubes and stored at -80 °C until used.

Ethical approval and/or informed consent

An informed consent was taken from all the patients after explaining the aim of the study to them the study protocol proved by research ethics committee of National Center for radiation Research and Technology (REC-NCRRT).

Preparation of the cercarial antigens

S. mansoni cercariae were purchased from Schistosome Biological Supply Centre (SBSC), Theodor Bilharz Research Institute (TBRI), Cairo, Egypt. Half them were irradiated by exposure to 400 gray gamma irradiation

at a dose rate 2.5 K.Gy/h at intensity of 400 $\mu\text{W}/\text{cm}^2$ for one minute at the National Center for Radiation, Research and Technology (NCRRT) in Cairo, Egypt (Moawad *et al.*, 2016). Antigens preparation were performed and stored at -70 until used according to the method of Feldmeier and Butter (1983).

Enzyme linked immunosorbant assay (ELISA)

Indirect ELISA was done as described by Santiago and Hillyer (1988) with some modifications. The optimal reaction conditions regarding antigens concentration, antibody and conjugate dilutions were chosen after preliminary checker-board titrations. For each cercarial antigen, the ELISA plates (Nunc, Denmark) were coated with 5 μg dissolved in 100 μl coating buffer in each well. The plates were incubated overnight at 4°C. In the next morning, the plates were washed with PBS/0.05 % Tween (T) to get rid of excess unbound antigen. The remaining free binding sites were blocked with 1% BSA (bovine serum albumin) (200 μl / well), and kept for 1 h at room temperature. The plates were then washed 3 times with PBS/ 0.05% T, the human sera were added to the plates in a final volume of 0.1 ml diluted 1/100 in PBS per well and the plates were incubated for 90 minutes at 37°C. After that, the plates were washed with PBS/T, 0.1ml/ well of conjugated horseradish peroxidase anti-human IgG (diluted at 1/1000) was added to all wells and incubated at 37 °C for 1 h. Finally, the plates were washed, 0.1ml ortho-phenylene-diamine substrate was added and the reaction was stopped by adding 100 μl / well of 1 N NaOH. The reaction was read using ELISA reader (ELx800 Absorbance Reader Bio-Tek) at 405 nm wavelength. All OD readings above cut off value were considered positive. Cut off = mean values of negative samples + 2 SD according to Gool *et al.* (2002).

Statistical analysis was performed using one way ANOVA using statically package for social science version 15 (SPSS) computer programs (2002). Descriptive statistics as Mean and standard deviation (SD) were used as they measure central tendency and dispersion of quantitative data. Analytical statistics as comparing groups was done using Student's "t" test for quantitative data of 2 independent samples and ANOVA (Analysis of Variance) for comparison of quantitative data of more than 2 groups. The following were used to calculate the corresponding diagnostic parameters:

Sensitivity = $(a \div (a+c)) * 100$, Specificity = $(d \div (b+d)) * 100$, Positive predictive value (PPV) = $(a \div (a+d)) * 100$, Negative predictive value (NPV) = $(d \div (c+d)) * 100$, Diagnostic accuracy = $(a+d) \div (a+b+c+d) * 100$, Where (a) is true positive, (b) is false positive, (c) is false negative, (d) is true negative.

RESULTS

In the present work, total IgG, ELISA was adopted to compare the potency of the normal and irradiated cercarial antigens using *S. mansoni* and other parasitic infected sera. Reaction of *S. mansoni* infected sera towards normal and irradiated cercarial antigens revealed that there were 14 *S. mansoni* infected sera showed higher antibodies response against irradiated SCA. The difference is highly significant when $P < 0.05$ (Fig. 1).

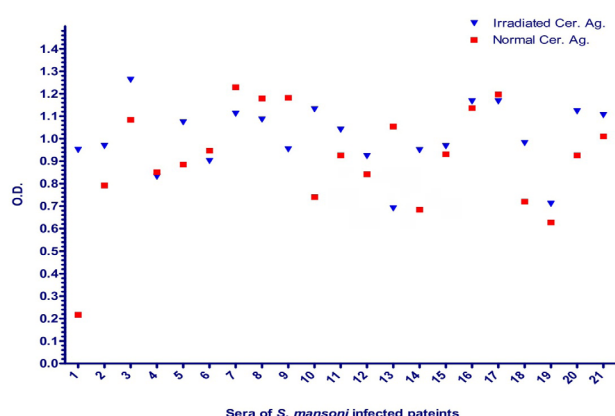


Fig. 1. Reactivity of the individual *S. mansoni* infected sera against irradiated SCA and normal SCA by ELISA.

Concerning cross reaction studies of the selected antigens, it was detected in Table I showing that normal SCA cross-reacted with 5 sera infected by other parasites which were 2 fascioliasis, 2 amoebiasis and 1 toxoplasmosis. While irradiated SCA cross-reacted with 4 sera infected by other parasites which were 1 fascioliasis and 3 amoebiasis.

Table I. Serological reactivity against normal and irradiated cercarial antigens using total IgG ELISA with other parasites infected serum.

Groups	No of sera (+/-)	
	Normal cercaria	Irradiated cercaria
Group I (14)	(21/21)	(21/21)
Group II (18)		
Fascioliasis (3)	(2/3)	(1/3)
Amoebiasis (5)	(2/5)	(3/5)
Toxoplasmosis (7)	(1/7)	(0/7)
Hydatidosis (3)	(0/5)	(0/3)
Group III (13)	(0/13)	(0/13)

The cutoff value for ELISA test using normal and irradiated cercarial antigen with sera were evaluated as 0.18 and 0.168. Optical density values of the normal and irradiated SCA against schistosomiasis sera were ranged from 0.217-1.229 and 0.690-1.262. Optical density values of the positively cross reacted normal SCA with amoebiasis, fascioliasis and toxoplasmosis ranged from 0.55-1.89, 0.19-0.95 and 0.95 respectively. While irradiated SCA cross reacted with amoebiasis OD ranged from 0.22-1.85 and with fascioliasis at 0.96 (Fig. 2).

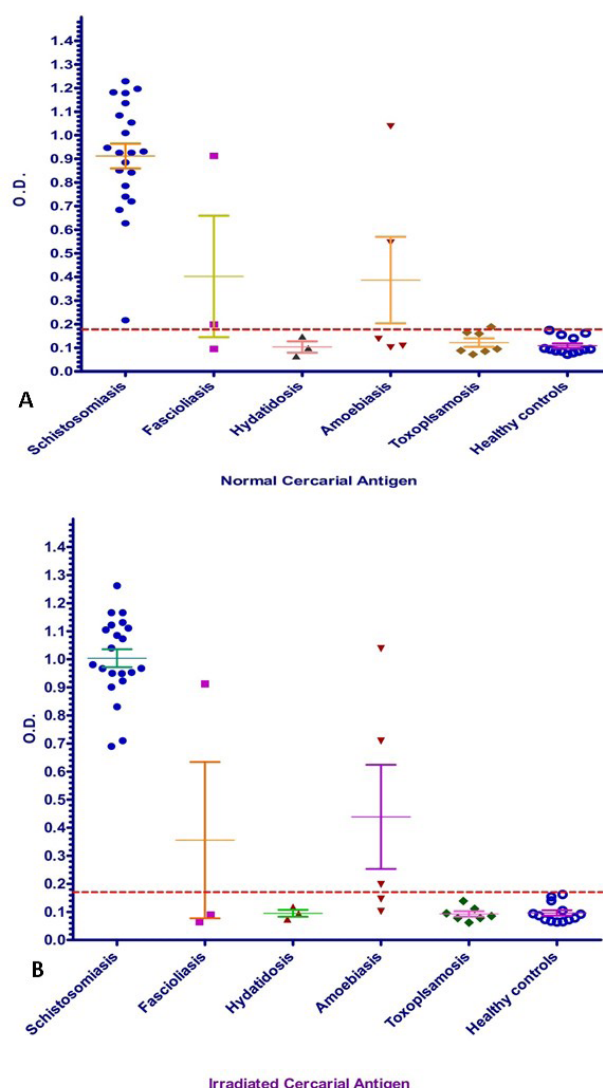


Fig. 2. Serodiagnosis of human schistosomiasis *mansoni* and other parasites using ELISA- normal (A) and irradiated (B) SCA Cut-off of healthy controls.

Both antigens revealed same sensitivity (100 %) and NPV (100 %). Irradiated SCA revealed higher specificity

(87%) compared to normal SCA (83.9). Higher PPV was obtained from Irradiated SCA (84%) while normal SCA achieved 80.7%. The accuracy of Irradiated SCA was 92.3 and normal SCA was 90.4% (Table II).

Table II. Sensitivity, Specificity, PPV, NPV and diagnostic accuracy percentages of both irradiated SCA and normal SCA by ELISA.

Accuracy %	NPV %	PPV %	Specificity %	Sensitivity %	Antigen
92.3%	100%	84%	87%	100%	Irradiated SCA
90.4%	100%	80.7%	83.9 %	100%	Normal SCA.

DISCUSSION

Although great efforts worldwide have been made to control schistosomiasis, it continues to spread to new geographic areas. The problem is linked with diagnosis as we still depend on microscopic examination which proved to be not ideal as light infections are often missed so there is a need to develop practical and scalable techniques (Coulbaly *et al.*, 2016).

Serological examinations as screening for anti-schistosomal antibodies are of limited use for the diagnosis, as large parts of the population may carry antibodies due to past infections so there is an urge for finding the best antigen that can be used in ELISA to diagnose patients from endemic area and follow-up treatment (Kinkel *et al.*, 2012).

In the current study, two types of cercarial antigens were enrolled normal and gamma-irradiated one. All the positive sera as judged by microscopic examination reacted positively with the two antigens. The cutoff value of the examined serum samples of *Schistosoma mansoni* patient against normal cercariae antigen were 0.18 and 0.168. These results nearly coincides with previous study revealed the cutoff value for ELISA test using *Schistosoma* adult worm, cercarial and egg antigens with sera of infected patients as 0.64, 0.82, and 0.892 respectively (El-Aswad *et al.*, 2011).

Studying the antibodies response towards either antigen, it was found that there were 14 *Schistosoma mansoni* infected sera showed higher antibodies response against the irradiated SCA than that of the normal one. Both antigens achieved 100% sensitivity while irradiated SCA showed 87% specificity rather than normal one giving 83.9%. In the same context, previous studies revealed 86.6 % sensitivity and 94.11 % specificity on assessing total IgG ELISA SCA using sera of chronic schistosomiasis patients. Also, ELISA using cercarial transformation fluid

showed sensitivity of 89.7% and an estimated specificity of 100% when used in non-endemic regions (El-Aswad *et al.*, 2011; Eberl *et al.*, 2001). While in endemic areas, the sensitivity has been reported to be between 20 and 65 % which could be explained that in endemic areas people may have subclinical infection with low parasite burden but have high antibodies titers (Doenhoff *et al.*, 2004).

The current study clarified cross reactivity between schistosomiasis, fascioliasis, toxoplasmosis and amoebiasis that was detected using normal SCA while irradiated SCA only cross-reacted with fascioliasis and amoebiasis. This was in accordance to results exhibited by Abdel-Fattah *et al.* (2011) reported serological cross-reactivity of SEA and SCA with sera from *Trichinella*, *Fasciola* and *Hydatid* infections. This may be too high for mass population work causing poor specificity especially in areas where other helminthes are prevalent.

The findings of present study revealed that irradiated SCA achieved better PPV and accuracy (84% and 92.3) than t normal SCA achieving 80.7% and 90.4%. These coincides with studies reported that radiation induces changes in the surface molecule making it highly immunogenic. When cercariae exposed to attenuated dose of gamma irradiation, antigens present on the surface will be altered and certain cryptic antigens having large antigenic loads exposed. Also radiation affects some structural and motor proteins, enzymes in addition to carbohydrates on the parasite surface (Wuhrer *et al.*, 1999; Yang *et al.*, 2009).

It could be concluded that there was a difference between normal and gamma radiation attenuated *Schistosoma mansoni* cercarial antigens in diagnosis of chronic schistosomiasis using ELISA. Irradiated SCA achieved better remarkable ODs, specificity due to low cross-reactivity with other parasites and better diagnostic accuracy so it is an effective tool to make a correct diagnosis.

ACKNOWLEDGEMENTS

The authors wish to acknowledge the assistance given by the National Center of Radiation Research and Technology, which provided all facilities for this study.

Statement of conflict of interest

The authors have declared no conflict of interest.

Funding

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

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