EVALUATION OF HUMERAL IMMUNITY AGAINST THREE HYDATID CYST ANTIGENS OF CAMELS USING SECONDARY CYST DEVELOPMENT IN RABBIT MODEL

By
WALEED M. MOUSA, OLFAT A. MAHDY*, AZZA M. ABDEL-WAHAB AND SOHAILA M. EL-GAMEEL

Department of Parasitology, Faculty of Veterinary Medicine, Cairo University, Giza, P.O. Box 12211, Egypt (*Correspondence: dr.olfat.mahdy@gmail.com)

Abstract

Cystic echinococcosis (CE) is one of the commonest zoonotic parasitosis of worldwide distribution. This study induced a protective immunity against secondary hydatidosis by using conventional vaccination approaches and evaluated the accompanied humeral immune response. For antigen preparation Hydatid cysts (HC) were collected from camels slaughtered at Cairo abattoir, during 2015. Three groups of male rabbits were immunized subcutaneously with crude Hydatid cyst fluid (CHCF), partially purified Hydatid cyst fluid (ppHCF) and protoscolices (PSC) antigens. After two boosters, rabbits were challenged intra peritoneally with 2000 viable protoscolices, and the humeral immune response was analyzed using ELISA. Percent of protection against HC was 93.0 %, 88.4 % and 86.0 % in immunized rabbits with ppHCF, PSC and CHCF antigens respectively. Antibody level in immunized group with ppHCF antigen on day 28 was higher than before immunization and was higher than that in CHCF and PSC antigen groups. The results of this article indicate that ppHCF antigen can be used as a candidate for vaccine production.

Key words: Egypt, Hydatid cyst, Immunization, Antibody, ELISA, Hydatidosis, Histopathology

Introduction

Cystic echinococcosis is one of the most widespread parasitic zoonoses in the world that causes a huge public health problems and economic impacts in several countries (Babazadeh et al, 2015). In the Middle East and Arab North Africa from Morocco to Egypt was considered as one of the major zoonotic parasitosis (El-Madawy et al, 2011), caused by Echinococcus granulosus complex which is endemic parasite in some countries that have a large production of herbivores with usage of dogs as pasturage animals (Thompson et al, 2006). The developing stages of Echinococcus spp is unaffected by the immune response. While the host immune system interacts against the juvenile intestinal stages or oncospheres and may lead to humeral immune response and production of specific antibodies (Eckert and Deplazes, 2004). Hydatidosis in the 1980s years was diagnosed by detection of specific antibody against it in the serum of infected host. These antibodies can be detected by using ELISA (Jenkins and Rickard, 1986). Proteins are probably the preferred choice of most researches on candidates for vaccine antigens preparation. As proteins proved easy to be identified, modified and produced economically in large quantities, for commercial vaccines production. In addition many proteins play key roles in parasite metabolism, so they may be a good target to disturb parasite biology if immune response rose against them (Hein and Harrison, 2005).

This study aimed to evaluate locally prepared three antigens from camel Hydatid cyst (HC); crude hydatid cyst fluid (CHCF), partially purified hydatid cyst fluid (ppHCF) and protoscolices (PSC) antigens. The immunological response against to antigens was evaluated by their effect on secondary HC development in rabbit model and detection of antibody titration using ELISA.

Material and Methods

Animal and parasite: Twenty male New Zealand white rabbits (2 Kg in weight, 4 months old) were reared under good hygienic conditions (clean, well ventilated and warm place) and fed on a balanced diet. All rabbits were examined through coprological
examination daily for 15 day to ensure that they were free from any parasite. Rabbits were randomly grouped into 5 gp (4 rabbits each). The Hydatid cyst fluid (HCF) was aseptically aspirated from freshly collected HC of naturally infected camels. The fluid was centrifuged and PSC were collected, rinsed 5-8 times in a sterile phosphate buffer saline (PBS) (pH, 7.4) containing penicillin (500IU/ml) and streptomycin (100µg/ml) then examined microscopically for viability through flame cell activity. The PSC were re-suspended in PBS containing antibiotics (Hashemi Tabar et al, 2009).

Antigen preparation: Hydatid cysts were collected from the lung and liver of slaughtered camels at Cairo abattoir (El-Basatin), during 2015. The antigens were prepared (Maddison et al, 1989) for partially purified Hydatid cyst fluid antigen (ppHCF), Ramsy et al. (1999) for crude hydatid cyst fluid antigen (CHCF) and Ahmed et al. (2001) for protoscolix antigen (PSC). The protein content of prepared antigens was measured using Lowry’s assay (Lowry et al, 1951).

Immunization and Challenge: Rabbits in G1, G2 & G3 were immunized with CHCF, ppHCF and PSC antigens respectively. The G4 (adjuvant group (Adj)) were injected with PBS and adjuvant only. Second and third immunization were conducted at 14th & 21th day post first immunization using same preparation. The G5 were used as infected non immunized control group (INCG). One week after the third immunization, each rabbit in all groups was challenged with 2000 PSC intraperitoneally (Ito et al, 2001). Blood samples were collected from the ear vein of rabbits at 0, 2, 4, 6, 8, 10, 12, 14 & 16 weeks post immunization (w.p.i) (12 weeks post challenge) for further serological studies (ELISA).

Table 1: Immunization and challenge of different animal groups S.C. subcutaneously in neck.

<table>
<thead>
<tr>
<th>4 Rabbits/group</th>
<th>Immunization and challenge</th>
<th>Challenge/rabbit (at 28th day)</th>
<th>Necropsy (16th w.p.i.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>First immunization (at zero day)</td>
<td>Booster dose (2nd at 14th &amp; 3rd at 21th day)</td>
<td>slaughtered and organs examined for cyst</td>
</tr>
<tr>
<td>Antigen</td>
<td>Dose/rabbit</td>
<td>Dose/rabbit</td>
<td>2000 viable PSC intra peritoneal suspended in 2ml PBS of penicillin (500IU/ml) &amp; streptomycin (100 µg /ml)</td>
</tr>
<tr>
<td>1 CHCF Ag</td>
<td>2 ml (100µg protein in mineral oil adjuvant)</td>
<td>2 ml (50µg protein in mineral oil adjuvant)</td>
<td></td>
</tr>
<tr>
<td>2 ppHCF Ag</td>
<td>2 ml (100µg protein in mineral oil adjuvant)</td>
<td>2 ml (50µg protein in mineral oil adjuvant)</td>
<td>2000 viable PSC intra peritoneal suspended in 2ml PBS of penicillin (500IU/ml) &amp; streptomycin (100 µg /ml)</td>
</tr>
<tr>
<td>3 PSC Ag</td>
<td>2 ml (100µg protein in mineral oil adjuvant)</td>
<td>2 ml (50µg protein in mineral oil adjuvant)</td>
<td>2000 viable PSC intra peritoneal suspended in 2ml PBS of penicillin (500IU/ml) &amp; streptomycin (100 µg /ml)</td>
</tr>
<tr>
<td>4 Adjuvant only</td>
<td>2 ml (PBS Adjuvant)</td>
<td>2 ml (in PBS)</td>
<td></td>
</tr>
<tr>
<td>5 control (INCG)</td>
<td>---</td>
<td>---</td>
<td>slaughtered and organs examined for cyst</td>
</tr>
</tbody>
</table>

Ag: antigen

Necropsy procedure and determination of infection: Rabbits were slaughtered on the 16th W.P.I. The different muscles, all visceral organs and their surrounding membranes particularly lung, liver, kidney, heart and spleen were examined macroscopically according to procedures of FAO (1994). Cyst load was calculated according to Al-Qaoud and Abdel-Hafez (2005) by multiplying the number of cysts with the average size of cysts for each group expressed in millimeters. Percentage of protection = 1- (mean number of cysts in test group/mean number of cysts in the non-immunized control group x 100 (Dempster et al, 1995). Statistical analysis of the obtained results was carried out with ANNOVA test. Collected HC were fixed in neutral-buffered formalin in 10%.

Determination of antibody titer: ELISA was used (De Savgy et al, 1979). The E. granulosus antigens prepared from camel were optimally diluted in coating buffer (carbonate – bicarbonate buffer, pH 9.6) and used 200 µl/ well to coat. Tested sera were added 100 µl / well at different dilutions starting with 1/100, 1/200 up to 1/3200. Protein -A IgG Horseradish peroxidase conjugate (diluted 1/2000) in PBS was added. The optical density (OD) values were read in a micro-ELISA reader system at 450 nm.

Histopathological changes: Collected HC from experimentally infected rabbits were fixed in neutral-buffered formalin 10%, and examined (Carleton et al, 1967).

Statistical analysis: Data were tabulated, computerized and analyzed using analysis of variance (ANNOVA test) and (Scheffe test) where, P-value < 0.01=significant variation.
Results

The immunized groups (Igp) with CHCF and ppHCF and PSC antigens, out of 4 rabbits in each group 2 were found to be infected (50% reduction in established infection). While all rabbits in Adj and INCG were infected with HC (zero reduction in established infection).

Total number and average size of cysts in the three Gs were lower than that of Adj & INCG. Highest number of cysts was six in CHCF followed by five in PSC and the low was three in ppHCF. One was the low number per rabbit was in ppHCF and PSC group and the high was in four/rabbit in CHCF.

<table>
<thead>
<tr>
<th>Animal Group</th>
<th>Cysts No. in individual rabbits</th>
<th>Cysts / group</th>
<th>Mean cysts / animal</th>
<th>Average size of cysts (mm.)</th>
<th>Cyst /group (mm.)</th>
<th>% of protection</th>
</tr>
</thead>
<tbody>
<tr>
<td>R1</td>
<td>0,5</td>
<td>1.5</td>
<td>1.2</td>
<td>12.5</td>
<td>86.0</td>
<td></td>
</tr>
<tr>
<td>R2</td>
<td>1.2</td>
<td>3</td>
<td>0.5-2</td>
<td>7.5</td>
<td>93.0</td>
<td></td>
</tr>
<tr>
<td>R3</td>
<td>1.5</td>
<td>1.25</td>
<td>1.2</td>
<td>7.5</td>
<td>88.4</td>
<td></td>
</tr>
<tr>
<td>R4</td>
<td>1.25</td>
<td>7.75</td>
<td>0.5-4</td>
<td>69.75</td>
<td>27.9</td>
<td></td>
</tr>
<tr>
<td>R5</td>
<td>1.2</td>
<td>10.75</td>
<td>1.8</td>
<td>193.5</td>
<td>0.0</td>
<td></td>
</tr>
</tbody>
</table>

Table 2: parameters of HC in different animal groups.

Antibodies produced in three rabbit Gs immunized with CHCF, ppHCF or PSC antigens was high than in controls. Antibodies appeared at 2 W.P.I and reached a peak at 4 W.P.I (a week after 3rd immunization) antibody level in ppHCF antigen was high (sample to positive (s/p) ratio=1.652) followed by PSC antigen (s/p ratio=1.629), then CHCF antigen (s/p ratio=1.517). Antibody in sera from Adj and INCG appeared on the 6th W.P.I (s/p ratio=0.512) and increased gradually till 12th W.P.I (s/p ratio=0.710) and decreased on the 14th W.P.I (s/p ratio=0.527) till the experimental end.

Histopathological examination of some representative sections of liver showed HC under the surface of liver capsule, embedded in the underlying liver tissue. Cyst was infiltrated by inflammatory cells mostly mono-nuclear cells and encapsulated by fibrous connective tissue. Hepatocytes beneath cyst showed necrobiotic changes from vacuolar degeneration to coagulative hepatic cell necrosis which was characteristically showed in hepatocytes surrounding cyst. Other section was surrounded with laminated layer with abundant degenerative calcific structure of calcareous corpuscles.

Table 3: S/P ratio of ELISA for the sera (1/100 dilution) from three immunized rabbit and control groups

<table>
<thead>
<tr>
<th>(W.P.I.)</th>
<th>CHC-G</th>
<th>ppHCF-G</th>
<th>PSC-G</th>
<th>INC-G</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.099</td>
<td>0.089</td>
<td>0.056</td>
<td>0.045</td>
</tr>
<tr>
<td>2</td>
<td>0.759</td>
<td>0.986</td>
<td>0.804</td>
<td>0.099</td>
</tr>
<tr>
<td>4</td>
<td>1.517</td>
<td>1.652</td>
<td>1.629</td>
<td>0.136</td>
</tr>
<tr>
<td>6</td>
<td>1.219</td>
<td>1.565</td>
<td>1.499</td>
<td>0.512</td>
</tr>
<tr>
<td>8</td>
<td>0.837</td>
<td>1.414</td>
<td>1.365</td>
<td>0.567</td>
</tr>
<tr>
<td>10</td>
<td>0.765</td>
<td>1.023</td>
<td>1.010</td>
<td>0.671</td>
</tr>
<tr>
<td>12</td>
<td>0.723</td>
<td>0.972</td>
<td>0.918</td>
<td>0.710</td>
</tr>
<tr>
<td>14</td>
<td>0.551</td>
<td>0.890</td>
<td>0.870</td>
<td>0.527</td>
</tr>
<tr>
<td>16</td>
<td>0.522</td>
<td>0.832</td>
<td>0.792</td>
<td>0.396</td>
</tr>
</tbody>
</table>

S/P ratio: mean optical density of sample –mean optical density of negative sample /mean optical density of positive sample

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Discussion

A range of different antigens including cystic fluid (Al-Qaoud and Abdel-Hafez, 2005) and protoscolex (Hernandez and Nieto, 1994) were used as vaccines candidates.

In the present study, the 1st assessment (necropsy procedures and cyst evaluation) gave variations between the immunized and the controls. In the group with CHCF and ppHCF and PSC antigens, out of 4 rabbits in each group 2 were found to be infected (50% reduction in established infection). While all rabbits in Adj and INCG were infected with HC. The highest protection rate (93.0% decrease in cyst load) in immunized animals with ppHCF antigen followed by PSC & CHCF antigens (88.4% & 86.0% reduction in cyst load, respectively.) These results agreed with Al-Qaoud and Abdel-Hafez (2005) who found that, mice immunized with antigen B (purified HCF antigen) prepared from sheep cysts showed a significant level of protection (98.3% reduction in cyst load) followed by CHCF and PSC antigens with 79% & 71% protection rate respectively. Hashemi Tabar et al. (2007) who recorded that lamb immunization with HCF antigen showed protection rate (75.75%) higher than that immunized with PSC (54.5%).

In the present study, secondary HC were developed in the omentum, diaphragm and near heart in immunized rabbits. This result agreed with El-Kattan (2017) who found that cysts were mostly distributed in the peritoneal cavity. Presently, in Adj and INCG groups the developed HC were embedded in or adjacent to all internal organs as liver, while chest cavities were free from infection. These results agreed with Theodorides et al. (2001) who recorded the same results.

These variations in cysts’ number and distribution might be related to the route and type of developmental stage used for infection Mousa and El-Massry (1999). Also, Zhang et al. (2001) recorded different degrees of protection against secondary infection when used a primary infection with onchospheres injected by different routes.

In the present study, the 2nd assessment of protection (humeral immune response), showed that antibody level in three immunized groups appeared at 2nd W.P.I and reach to its peak at 4th W.P.I (one week after third immunization) as antibody level was about 11 time higher than before immunization. Antibody level in ppHCF antigen was the highest followed by PSC antigen then CHCF antigen. This result agreed with Youssef et al. (2010) who found that, the mice immunized with purified hydatid fluid antigen produced higher antibody levels than that mice immunized with PSC antigen on the 4th and 8th W.P.I. Also, Al-Qaoud et al. (2008) reported that the highest level of IgG1 was recorded in AgB immunized mice followed by low but protective antibody level in immunized mice with CHF and PSC antigens. Hashemi Tabar and Razmi (2009) recorded that the level of antibody in immunized mice with E. granulosus adult on day 28th & 49th post immunization was higher than in mice immunized with HCF & PSC antigens. On contrary El-Kattan (2017) reported that the level of antibodies produced in response to CHCF antigen was higher than in ppHCF & PSC antigens. Also, Hernandez and Nieto (1994) found that PSC produced high level of antibodies against infection in mice. This variable responses and protection rates were explained by using the different antigen preparations from different antigen sources of variable purities (Al-Qaoud et al, 2008).

Cysts of three groups were collapsed and retarded in its normal growth in comparison with the cysts of the INCG that were more fluidly. Histopathological examination of cysts and surrounding tissues confirmed the results. It showed presence of HC under the surface of liver capsule embedded in the underlying liver tissue. The cyst was infiltrated by inflammatory cells mostly mononuclear cells and encapsulated by fibrous connective tissue. Other sections were surrounded with laminated layer. Also, abun-
dant degenerative calcific structure of calcareous corpuscles was seen. These results agreed with El-Kattan (2017) who found that the collected cysts were collapsed and retarded in its normal growth in apart of immunized rabbit groups and fluid in others.

Generally speaking, in Egypt human hydatidosis was well documented in many governorates (Abdalla et al., 1975; Bebars et al., 1987; Romia et al., 1992; Madwar et al., 1995; Mazyad et al., 1998; 1999; Ramzy et al., 1999; El Shazly et al., 2001; Sadaka et al., 2002; Kandeel et al., 2004; Dyab et al., 2005; Abbas et al., 2006; El-Sebaie et al., 2006; Ibrahim et al., 2007; El Wakil et al., 2007), edible animals (El Kordy, 1946; Hamdy et al., 1980; El-Ridi et al., 1983; Ahmed, 1991; Abdel Rahman et al., 1992; Lotfi et al., 1994; Mohamed et al., 1997; Haridy et al., 1998, 2000; Abdel-Alim et al., 1999, Dyab et al., 2005) and in street dogs (Selim, 1967; Abou-Eisha and Abdel-Aal, 1995; El Shazly et al., 2007; Mazyad et al., 2007). Echinococcosis rate in the stray dogs ranged between 4.6% and 10.2%. Dogs are domestic animals used to tend sheep and goats, and pet dogs enjoy a more intimate contact with man than other animals or birds (Haridy et al., 1998).

Conclusion

No doubt, echinococcosis and hydatidosis are economic and public health problem. Necropsy and ELISA results showed controversial relationship between number of the developed secondary HC and antibody level. As the number of cysts was the lowest in ppHCF antigen (3 cysts) followed by PSC (5 cysts) then CHCF group (6 cysts). Antibody level was the highest in ppHCF antigen (s/p ratio=1.652) followed by PSC antigen (s/p ratio=1.629) then CHCF antigen (s/p ratio=1.517). The ppHCF antigen was the best antigen that induced a protective immunity aganist secondary hydatidosis in rabbit model through reduction in number of the developed cysts and the enhancement of production of the highest antibodies titer. So using ppHCF antigen is a candidate for vaccine production locally.

References


**Explanation of figures**

Fig. 1: Parameters of hydatid cysts in different animal groups.

Fig. 2: S/P ratio of ELISA for the sera (1/100 dilution) obtained from three immunized rabbit and control groups.
Plate 1: HC formed in infected non-immunized control and AGs: A: show HC on spleen (a) and omentum (b) (arrow), B: showed HC on stomach (arrow), C&D: showed HC on liver (arrow), E: showed HC on liver (c) omentum (d) (arrow), F: showed HC on intestine (arrow) G: showed HC on liver (e) omentum (f) (arrow), H: showed HC on omentum (arrow)
Plate 2: HC formed in immunized groups: I: HC in omentum (arrow), J: HC on diaphragm (arrow), K&L: HC on pericardium (arrow)

Plate 3: Liver sections stained with H&E. A: presence of hydatid cyst under surface of liver capsule and embedded in underlying liver tissue (arrow) (H&E x4). B: hydatid cyst infiltrated by inflammatory cells mostly mononuclear cells and encapsulated by fibrous connective tissue. Hepatocytes beneath cyst revealed necrobiosis changes from vacuolar degeneration to coagulative cell necrosis (H&E x20). C: Hydatid cyst surrounded with laminated layer (L) (arrow) (H&E x4). D: Abundant degenerative calcific structure of Calcareous corpuscles (arrow) (H&E x20).