RESPONSES OF THE RODLET CELLS TO METACERCARIAL INFECTIONS IN OREOCHROMIS NILOTICUS

WALAA F.A. EMEISH1, MARWA M. FAWAZ2 and ZEINAB AL-AMGAD3

1 Fish Diseases and Management, Department of Fish Diseases, Faculty of Veterinary Medicine, South Valley University, Qena, Egypt.
2 Department of Parasitology, Faculty of Veterinary Medicine, South Valley University, Qena, Egypt.
3 Veterinary National Service, PhD in Veterinary Pathology and Clinical Pathology, Faculty of Veterinary Medicine, South Valley University, Qena, Egypt.

Received: 24 June 2018; Accepted: 17 January 2019

ABSTRACT

In this report, we describe the responses of Rodlet cells (RCs) of Nile Tilapia, Oreochromis niloticus, with other inflammatory indicators to infections by encysted metacercariae trematodes. RCs are specialized cells of the non-specific immune system of some species of teleosts fish. Sixty eight out of 100 examined fish specimens were found to be naturally infected by the encysted metacercariae in different parts of the body. Digenean metacercariae infection caused significant increase in serum total protein and albumin level concentrations than in uninfected wild caught fish, a possible indication of an infection or inflammatory host response. Significant increase in eosinophils percent revealed parasitic infections. A variable number of RCs were observed in gills and liver tissues of infected fish, with a higher frequency, while couldn’t be found in uninfected tissues. Results propose that RCs represent an inflammatory cell type that is closely linked to other piscine inflammatory cells in respond to the presence of parasites on epithelial surfaces.

Key words: Rodlet cells, Oreochromis niloticus, encysted metacercariae, biochemical parameters.

INTRODUCTION

Recently, RCs of the teleosts have drawn a lot of attention as fish inflammatory cells that are involved in the non-specific immune response against parasitic infections (Reite, 2005). Rodlet cells have been identified as fish blood cells (Weinreb and Bilstad, 1955) and formerly were considered as regulatory elements related to special functions such as transportation of ions (Morrison and Odense, 1978), osmoregulatory mechanisms (Mattey et al., 1979), transport units of genetic material (Viehberger and Bielek, 1982), secretory cells (Leino, 2002), and non-specific immune cells (Dezfuli et al., 1998; Dezfuli et al., 2000; Dezfuli et al., 2002; Dezfuli et al., 2003a; Dezfuli et al., 2003b).

The influence of parasites on fish population is widely recorded (Abdel-Ghaffar et al., 2013). Nile tilapia, Oreochromis niloticus, is a freshwater fish belongs to the Family Cichlidae. It is widely distributed and native to Africa and has been used in aquaculture in different parts of the world because of its good characteristics such as rapid reproduction and growth rates (Nandlal and Pickering, 2004).

Digenean trematodes constitute the largest group of all internal metazoan parasites gathers about more than 2,500 nominal genera, and in general its life cycle includes three hosts, two intermediate and one definitive. Metacercariae are the infective larvae found in the fish that act as the second intermediate hosts. In fish most of detected metacercariae were with unique morphologies so they are morphologically distinguished species by species (Sohn, 2009; 2013).

Blood examination have been applied as physiological indicators to changing external environment (Caruso et al., 2005) and used to evaluate normal health status and to diagnose diseases caused by various factors, like parasitic infections, and others (Fedato et al., 2010). Hence, any alterations associated with Haemato-biochemical parameters due to various parasites establish a data base, which could be used in diagnosis of diseases and in guiding the inquiry of
the treatment or preventive measures (Yaji and Auta, 2007).

The aim of the present study was to describe the response of RCs in gill tissues to infection with a range of parasitic genera including representatives from the Digenea trematodes, metacercariae in wild Nile tilapia, *O. niloticus* collected from the River Nile in Qena Province. Moreover, the potential effects of occurrence of these parasites on selected inflammatory and biochemical indicators and histopathological finding in gills were also studied.

**MATERIALS AND METHODS**

**Fish samples:**
Nile tilapia, *O. niloticus* (n=100) with an average body weight of 50±10 grams were collected alive from two different River Nile Branches at Qena Governorate, South Egypt by the aid of fisherman and then transported alive to the aquatic laboratory at Faculty of Veterinary Medicine, South Valley University, Qena, Egypt, where they were subjected to parasitological, biochemical and histopathological examinations.

**Blood samples:**
Fresh blood samples were collected without anticoagulant from 12 digenetic trematodes metacercariae infected and 12 non-infected wild caught fish from the caudal peduncle according to (Lied et al., 1975). Blood films were prepared, air dried, methanol fixed and stained with diluted Giemsa stain for differential leucocytic count (DLC). Serum was obtained by allowing blood to clot for 30 min – 45 min and then centrifuged at 3000 rpm for 15 minutes and non-hemolysed Serum samples were transferred to separate Eppendorf tubes and preserved at -80°C until use for biochemical analysis.

**Clinical and postmortem examinations:**
The infected fish were subjected to clinical as well as post mortem examinations according to the methods described by Austin and Austin (2016) for detection of any external or internal abnormalities.

**Parasitological examination:**
External and internal examinations were done for detection of metacercaria cysts, by naked eye and/or by using of magnification lenses, smears on glass slides were made from scrapings of the gills, skin and fins for examination of smaller metazoan (Ayanda, 2009). Fish gills were dissected out and each gill filament and arch was examined for the presence of digenean cysts. The fishes were dissected to expose the viscera. The visceral cavities and organs were examined for metacercarial cysts according to (Syme, 1966). Microscopical examination made by put a very small piece of musculature of each sample from different regions and depths, from right and left sides, mixed with few drops of saline solution and compressed between two glass slides and examined microscopically according to the method reported by (Morishita et al., 1965).

**Biochemical analysis:**
Samples were thawed at room temperature and the following analysis were made: levels of serum total protein and albumin were measured using spectrophotometry and “Total Protein” and “Albumin” kits (Spectrum, Egyptian Company for Biotechnology, Obour City, Cairo, Egypt) according to manufacturer’s recommendation. Subtracting the concentration of albumin from that of the total protein was done to obtain blood serum globulin (Coles, 1986).

**Histopathological examination:**
Infected fish with Digenean as well as non-infected fish were subjected to histopathological examination in order to compare and specify the pathological lesions observed if it attributed to Digenean infection or other environmental, nutritional and other infections. Samples from gills and liver were collected, fixed in 10% neutral buffered formalin, washed in running water and dehydrated in different grades of concentrated alcohol, cleared in xylene and embedded in paraffin wax. Paraffin sections of 4.5 µm thickness were obtained and stained by with Harries hematoxylin and eosin for microscopical examinations according to (Culling et al., 1985). Other sections subjected to special stain using Periodic Acid Schiff (PAS) according to (Mc Manus, 1948). Then covered and examined microscopically.

**Statistical analysis:**
Data are presented as mean ± standard error. A two-way analysis of variance (ANOVA) were done to calculate any significant differences between infected and uninfected groups using Graph Pad prism 6, version 6.01, significance was set at 95%.

**RESULTS**

In the present study, digenean metacercariae were detected in 68% fish species, but their all species names could not be determined, while 32% were uninfected. The encysted metacercariae could be recorded mainly in gills tissue, branchiostegal muscle, also in pericardial muscle, chin region, branchiostegel membrane and in internal organs as muscle and kidney tissue (Figure 1).
Figure 1 (A): Encysted metacercaria (Clinostomum sp.) at gills tissue [Arrow] and branchiostegal membrane [Arrow head] (Camera digital).
Figure 1 (B): Unidentified Encysted metacercaria at gills tissue (Microscopically X10).
Figure 1 (C): Clinostomum sp. stained with alum carmine stain (X4).
Figure 1 (D): Encysted metacercaria (Clinostomum sp.) at kidney tissue (Camera digital).

The results of the biochemical indices determined for both infected and uninfected species of O. niloticus are presented in (Table 1). The results indicated vividly that serum total protein and albumin levels concentration were higher in the infected than in the uninfected fishes for studied species. Statistical analysis revealed significant differences between the serum parameters of the infected and uninfected species. However, globulin ratio was lower in the infected than in the uninfected fishes. Statistical analysis revealed insignificant differences.

Results of DLC showed that, the esoinophils and monocytes percent were found to be significantly higher in the infected fishes than in the uninfected ones. While neutrophils percent, were significantly lower in the infected fishes than in the uninfected ones. Results presented in Table 1.

Table 1: Biochemical and inflammatory alterations in Nile Tilapia due to metacercariae infections.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Groups</th>
<th>Non infected (control)</th>
<th>Infected</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=12</td>
<td>n=12</td>
<td></td>
</tr>
<tr>
<td>Total protein g/dl</td>
<td>3.7±0.18b</td>
<td>4.5±0.34a</td>
<td></td>
</tr>
<tr>
<td>Albumin g/dl</td>
<td>1.5±0.1b</td>
<td>1.9±0.16b</td>
<td></td>
</tr>
<tr>
<td>Globulin g/dl</td>
<td>2.2±0.1a</td>
<td>2.6±0.3a</td>
<td></td>
</tr>
<tr>
<td>Lymphocyte %</td>
<td>38.75±1.02a</td>
<td>37.08 ±1.1a</td>
<td></td>
</tr>
<tr>
<td>Monocyte %</td>
<td>32.92±1.5b</td>
<td>36.08±0.8a</td>
<td></td>
</tr>
<tr>
<td>Neutrophil %</td>
<td>16.41±1.04a</td>
<td>13.75±0.7b</td>
<td></td>
</tr>
<tr>
<td>Basophile %</td>
<td>8.8±0.7a</td>
<td>8.6±0.5a</td>
<td></td>
</tr>
<tr>
<td>Eosinophil %</td>
<td>3.08±0.3b</td>
<td>4.4±0.4a</td>
<td></td>
</tr>
</tbody>
</table>
At the normal condition; histological section of gills revealed normal architecture, since RCs couldn’t be distinguished in case of absence encysted metacercarial infection (Fig. 2 a-b). The present study involved Nile Tilapia subjected to parasitic infections which appeared among gills tissues in form of variable sized encysted metacercarial infection surrounded by RCs (Fig. 2 c-d). In response to parasitic infection, there were large population of RCs infiltration among interstitial tissues appeared oval shaped with eosinophilic granular cytoplasm and blue peripheral nuclei when stained by Haematoxylin and Eosin (Fig. 2 e-f). Moreover, primary and secondary lamellae exhibited variable number of RCs related to damage tissues (Fig. 2 g). While when stained PAS, RCs appeared deeply stained blue coloration concentrated at the primary and secondary lamellae in associated with damaged gill tissues (Fig. 2 h).

The histological section of gills of control fish free from parasitic manifestation showed normal architecture composed of gill rackers, gill arch, and filaments. Since, gill arch radiates double rows of paired primary lamellae. Each of these primary lamellae has a series of secondary lamellae located perpendicular to the primary lamellae (Fig. 3 a). While, fish that infected with encysted metacercaria were accompanied with remarkable histopathological changes in the gills tissue characterized by pronounced necrosis of the primary and secondary lamellae (Fig. 3 b). On the hand, gill lamellae showed hyperplasia in the epithelial lining led to focal fusion at the tips of the secondary lamellae (Fig. 3 c). Congestion in the central venous sinus of the primary lamellae was observed, in addition to severe destruction and sloughing of the lamellae (Fig. 3 d). There was curling and curving of some lamellae, besides shortening of others. Gills arch exhibited interstitial mononuclear cells infiltration mainly lymphocytes.

**Figure 2 (a-b):** Histological section in gill tissues showing normal architecture, besides deprived cystic metacercarial infection and free from rodlet cell infiltration (a). High power of Fig. a, was showing normal architecture with absence of rodlet cell infiltration (b). (H&E., bar= 50 & 80 µm).

**Figure 2 (c-d):** Histological sections in gill tissues showing variable sized encysted metacercarial infection (thin arrows) with rodlet cells infiltration (thick arrows). (PAS, bar = 50 & 80 µm).

**Figure 2 (e-g):** Gills associated with encysted metacercarial infection showing extensive rodlet cells infiltration among tissues (e). High power of Fig. e showing oval shaped rodlet cells with eosinophilic granules and blue peripheral nuclei (f). Gills showing rodlet cells infiltration among primary and secondary lamellae with damaged tissues (g). (H&E., bar= 50 & 80 µm).

**Figure 2 (h):** Gills stained with PAS showing deeply blue stained rodlet cells at primary and secondary lamellae (h). (PAS, bar= 50 µm).
Figure 3 (a): Histological section in gills showing normally arranged primary lamella with a series of secondary lamellae located perpendicular to the primary lamellae (a). (H&E., bar= 80 µm)

Figure 3 (b-d): Histopathological changes accompanied by encysted metacercarial infection showing pronounced necrosis of the primary and secondary lamellae with curling of some secondary lamellae (b). Gills were showing focal fusion in the tips of the secondary lamellae (c). Gills were showing congestion in the central venous sinus of the primary lamellae, besides severe destruction and sloughing of the primary and secondary lamellae (d). (H&E., bar= 50 & 80 µm)

The histological section of liver of the infested fish displayed encysted metacercaria embedded among hepatic tissues (Fig. 4 a-b). The liver when stained with PAS showed large encysted metacercarial infection filled with parasitic eggs and surrounded by dense fibrous capsule (Fig. 4 c-d). Also, liver exhibited extensive rodlet cells infiltration with eosinophilic appearance, since there was rodlet cells infiltration related to hepatopancreas (Fig. 4 e-f). The histological section of the control liver was devoid of encysted metacercarial infection or any parasitic cysts, where the hepatic tissue showed normal criteria that composed of branching and anastomosing hepatocytes. Hepatocytes are polygonal with distinctive central nucleus (Fig. 5 a). Liver of the infested fish revealed pronounced histopathological lesions involving hepatic damage and necrosis characterized by small tiny condensed nuclei of the hepatocytes adjacent to central vein (Fig. 5 b-c). Liver showed hyperactivation of kupffer cells among the hepatocytes (Fig. 5 d). Liver showed fatty degenerative changes with fat cells infiltration and mononuclear cells infiltration aggregation mainly lymphocytes (Fig. 5 e). The infested liver revealed proliferation of melanomacrophages. Also, hepatopancreatic area suffered from cytoplasmic vacuolation with inflammatory cells infiltration (Fig. 5 f). There was hemorrhagic inflammation manifested by erythrocytes and lymphocytes cells infiltration (Fig. 5 g). The blood vessels showed thrombotic congestion of the central vein characterized by presence of the fibrinous exudate with lymphocytes and stagnant erythrocytes infiltration attached to the vessels wall (Fig. 5 h).
Figure 4 (a-b): Histological section in liver showing encysted metacercarial infection embedded among hepatic tissues (a). High power of Fig. a showing encysted metacercarial infection embedded among hepatic tissues (b). (H&E., bar= 50 & 80 µm).

Figure 4 (c-d): Liver with especial stain showing encysted metacercarial infection embedded among hepatic tissues (c). High power of Fig. c showing encysted metacercarial infection embedded among hepatic tissues (d). (PAS, bar= 50 & 80 µm).

Figure 4 (e-f): Liver showing extensive rodlet cells infiltration with eosinophilic appearance (e). Liver showing rodlet cells infiltration related to hepatopancreas (f). (H&E., bar= 50 µm).

64
Figure 5 (a): Histological section in control liver showing normal arrangement of the hepatocytes with polygonal shapes (b). (H&E., bar= 50 µm).

**Figure 5 (b-h):** Histopathological changes of the infested liver showing hepatic necrosis adjacent to central vein (b). Liver showing hepatocytes necrosis with tiny condensed nucleus (c). Liver showing hyperactivation of kupffer cells (d). Liver showing fatty infiltration with lymphocytes aggregation (e). Liver showing hepatopancreatic vacuolation with inflammatory cells infiltration (f). Liver showing extravasation of red and white blood cells among hepatocytes (g). Liver showing thrombotic congestion of the central vein (h). (H&E., bar= 50 µm)

**DISCUSSION**

Parasites extensively affect immunity of fish which modulated by pathogen recognizing receptors that can reduce the parasite load (Alvarez-Pellitero, 2008). The cellular involvement in the inflammatory response in teleost fish could be having two phases, beginning with neutrophils inflow and later by monocytes and macrophages (Reite and Evensen, 2006). Furthermore, another inflammatory cellular type is the enigmatic rodlet cells, which located in the epithelial tissues and exclusive to fish species, with a higher variability in their distribution and abundance (Alvarez-Pellitero, 2008). Abd-Elhafeez
and Soliman (2016) concluded that, RCs originated from the stroma of the olfactory organ.

There are arguments over the kind of rodlet cells, but mainly there are three suggestions: one supports that the RCs are glandular components of the connected epithelium (Iger and Abraham, 1997; Imagawa et al., 1998); the second distinguish these cells as protozoan parasites of the Sporozoa (Agullheiro et al., 1986). The main alibi that RCs are parasite is that their numbers differ from fish to fish and often they cannot be found in all individuals of the same species (Manera and Dezfuli, 2004). The third view recognizes RCs as a kind of granulocytes, which is based on common features between RCs and white blood cells (Smith et al., 1995a; Smith et al., 1995b) and aggregation of RCs at the site of infection due to metazoan (Dezfuli et al., 1998; Dezfuli et al., 2000; Dezfuli et al., 2003a) and protozoan parasite (Dezfuli et al., 2004). Several studies have reported that RCs are associated with the immune system of fishes (Manera and Dezfuli, 2004; Bielek, 2005; Reite, 2005). In consequence, although the role of RCs in organisms is not completely understood, RCs was considered as biomarkers in inflammation and parasitic infections, with a similar target to eosinophil, epithelioid and mononuclear cells (Iger and Abraham, 1997; Araujo and Borges, 2015). Previous studies have also reported that exposure to adverse environmental conditions and toxic substances lead to increased numbers of RCs in fish (Hawkins, 1984; Manera and Dezfuli, 2004).

This study has reported increase in the number of RCs in gills and liver tissues of fish exposed to parasitic infections than in uninfected fish, and this was associated with stimulation of antibodies in a response to parasitic infection. There is a correlation between the rising number of RCs and fish parasite infections (Dezfuli et al., 1998; Dezfuli et al., 2000; Dezfuli et al., 2003a). Protozoans, myxozoans and helminths, seem to induce the induction of RCs with variability in their abundance and distribution (Manera and Dezfuli, 2004).

Matisz et al. (2010) studied changes of RCs in the optic lobes of fathead minnows exposed to trematode metacerciae. Smith et al. (1995b) noted higher numbers of RCs around lesions in the head kidney of Angelfish. A variable number of RCs was observed in gills and kidneys tissues, with a higher frequency in gills compared to the kidneys. RCs were observed in healthy fish and in fish parasitized by a myxosporean of the genus Henneguya (Mendonca et al., 2005). Compromise integrity of the RCs in gills tissues occurred in O. niloticus exposed to organophosphate pesticide methyl parathion (Araujo and Borges, 2015).

In addition to the RCs infiltration, there were remarkable histopathological changes encountered in gills of fish infested with encysted metacercaria when compared with non-infested cases in the present study represented by necrosis, focal fusion of gills lamellae, besides congestion of the blood vessels. Aly et al. (2005) displayed that encysted metacercariae of two trematodes (Prohemistomum vivax and Mesostephanus appendiculatus) caused pronounced pathological changes in gills characterized by desquamation of the secondary lamellae, hyperplasia in the epithelium with fusion in their lamellae, besides edema, congestion and mononuclear leukocytic infiltration. Fish parasites are capable of producing proteolytic enzymes responsible for tissue deterioration (Jones et al., 2004; Soror, 2008) and possibly parasite encystations (Martone et al., 1999). The degenerative changes of gills resulted in hypoxia and osmoregulatory failure and consequently death of host. The degree of pathogenicity depends on many factors, such as the species involved, its life cycle and biology, the host species, host age, state of nutrition and host resistance. The degenerative changes could be due to the pressure or toxic products induced by the parasitic cysts (Aly et al., 1995). Yemmen et al. (2011) discussed that the adhesive disc of the parasite lead to decreased respiratory surface and gaseous exchange resulting in hypoxia with lack of oxygen. Encysted metacercariae was associated with destruction of secondary lamellae with loss of the normal gill architecture, similar to (Abdel-Latif, 2007; Reda et al., 2010; Eissa et al., 2011) who recorded congestion of the blood vessels in addition to mononuclear inflammatory cellular infiltration in gills of O. niloticus infected with encysted metacercaria. In addition, (Shoaihi Omrani et al., 2010) noted hyperplasia and hypertrophy with fusion of the gill filament of platy fish infested with metacerceria. (Aly et al., 1995) expressed that inflammatory reactions detected in the gills due to the impact of toxic metabolites produced by the larvae with the prolonged irritation of the parasitic cysts.

Pathological lesions of liver showed hepatic necrosis and damage associated with inflammatory reactions were correlated to a direct effect of the parasites through attachment to the wall of the organ leading to compressing cellular death and damage inflammatory reactions (Hossam et al., 2012). Also hepatic damage and atrophy which may be a result of escape of nutrients from the host tissue to the metacercariae caused shrunken hepatocytes (Harris et al., 2005). Sommerville (1982) recorded that marked inflammatory changes and focal hemorrhages followed penetration and early migration of metacercariae. The inflammatory reaction, predominated by infiltrating macrophages,
is particularly intense around unencysted migrating metacercariae and preceded the eventual enclosure in a fibrous capsule of the encapsulating metacercaria (Yekutiel, 1985). Hyperactivation of kupffer cells among the hepatocytes was detected in response to immune system. Fatty degeneration was noticed with fat infiltration due to the pressure or toxic harmful products released by the parasitic cysts (Aly et al., 1995).

The physiological status of the fish infested by parasites can be diagnosed by blood parameters which act as a reflector of the health of an organism (Joshi et al., 2002) and the normal physiology and nutritional conditions of fish (Chagas and Val, 2003). The total protein and albumin concentrations in fish serum reflect the health of the animal, liver function, metabolic status and stress conditions (Kovyrshina and Rudneva, 2012).

With progress of any infection usually marked changes in the total serum proteins were detected. There may be elevated concentration of the total protein during some stage of the infection. Results of the present study indicated that the blood serum total protein and albumin levels were significantly higher in metacercariae infested fish. In another study on the biochemical parameters of 

Clarias batrachus

infested with trypanosomosis, insignificant higher levels of albumin were reported in infected fish (Kharat and Kothawad, 2012), while Osman et al. (2009) reported low serum albumin levels in C. gariepinus with trypanosomosis.

There are many factors responsible for variation in total protein and albumin concentrations in fish serum other than parasite infection, since it has been noted that diet composition, species, stage of life cycle, age, sexual maturation, environmental and health factors (Patriche et al., 2009; 2011; Chukwuma et al., 2010; Kovyrshina and Rudneva, 2012) also affect total protein and albumin levels. The increase in albumin value consequently leads to decrease in globulin, as globulin was obtained by direct subtracting the values of the albumin from those of the total protein. Globulin and total protein levels indirectly give indication about the condition of specific humeral immunity (Stosik et al., 2001; Maqsood et al., 2009). Albumin is considered as an important serum protein in body and its physiological function is the transportation of steroid hormones (Shahsavani et al., 2010). Also, the importance of albumin has been described in respect to fish pathology, and is widely used as an index of physiological state (Nakagawa, 1978).

During infection white blood cells play a major role by stimulating the immune system and haemopoietic tissues to produce antibodies and chemical mediators which work as defense agent so it considered as a primary line of immunological defense (Tierney et al., 2004). Also, changes in leucocytes occur when fish are stressed and environmental quality is altered (Ponsen et al., 2009). Both innate and adaptive immune response is mounted by fish to control parasite infection (Alvarez-Pellitero, 2008). The innate (non-specific) immune system includes: the phagocytic cells (granulocytes (neutrophil) and monocytes/macrophages) and non-specific cytotoxic cell (Magnadottir, 2006).

In the present study, significantly higher eosinophils count was observed in the infected fishes because of parasitic infestation, where eosinophils were the cells more frequently involved (Alvarez-Pellitero, 2008). An increase in eosinophils count was observed in 

Clarias batrachus
due to helmint infections (Sinha, 2010). Parasitic infection also enhances the phagocytic cells which lead to the clearance of pathogens (Alvarez-Pellitero, 2008), similar reports recorded in this study as there were significant increase in monocyte percent.

The increase or decrease in different types of leucocytes was reported in fish because all leucocytes were calculated as a percentage of the whole leucocytic count which constitutes 100 %. The decrease of the percentage of neutrophils in fish groups may be attributed to the significant increase of other leucocytic cells.

CONCLUSION

Data of this study showed that gill and liver RCs undergo to increase during parasite infection, which could be related to cell activation against parasites. These data also suggest that RCs may play an important role in the non-specific defense mechanisms of teleosts.

REFERENCES

Abdel-Ghaffar, F.; Bashtar, A.R.; Mehlhorn, H.; Abd-Gaber, R. and Saleh, R. (2013): Morphological and molecular characterization of 

Lecithochirium grandiporum

(Digenaea: Hemiuridae) infecting the European eel 

Anguilla anguilla


Epalzeorhynchos frenatum


Diseases and Management). Faculty of Veterinary Medicine, Benha University, Egypt.


Austin, B. and Austin, D.A. (2016): Bacterial Fish Pathogens: Disease in Farmed and Wild Fish, 6th ed. Springer International Publishing Switzerland.


Soror, E.I.M. (2008): Studies on some internal parasitic diseases of Nile tilapia in Kalubia Governorate. M.Sc. Thesis. Fish diseases and management, Faculty of Veterinary Medicine, Benha University, Egypt:


Syme, J.D. (1966): Fish and fish inspection. 2nd Ed. 52-64. University of Toronto.


Weinreb, E.L. and Bilstad, N.M. (1955): Histology of the digestive tract and adjacent structures of the rainbow trout, Salmo gairdneri irideus: Experientia 5: 204.


إستجابة خلايا الرودلت للعواري بيرقات الديدان المفلطحة في أسماك البلطي النيلي

ولاء فتحى على عميش، مروه محمد فواز، زينب الأمجد

Email: walaavet2002@yahoo.com Assiut University web-site: www.aun.edu.eg

في هذا البحث، تم وصف استجابة خلايا الرودلت في أسماك البلطي النيلي إلى العواري بيرقات الديدان المفلطحة مع غيرها من المؤشرات الإلتهابية. خلايا رودلت هي خلايا مناعية غير محددة خاصة ببعض الأسماك. وأوضحت نتائج الفحص أن مجموع الأسماك المصابة بطفيل الميتاسركاريا المتحولى في أماكن مختلفة من جسم السمكة كانت 68 من أصل 100 عينة التي تم فحصها لتكون مصاباً بشكل طبيعي. وأوضحت النتائج أن عدوى الميتاسركاريا المتحولى ثنائية العائل تسببت في زيادة معنوية في تركيز البروتين الكلي والزلال في مصل الدم مقارنة بالأسماك غير مصاباً، وهو مؤشر لأحتمال وجود إصابة أو استجابة الجسم لمؤشرات التهاب. زيادة كبيرة في نسبة الإيزوتبول في الدم كشفت عن الإصابة بالطفيليات. لوحظ وجود عدد متغير من خلايا الرودلت في أنسجة الخياشيم والكبد للأسماك المصابة، في حين لا يمكن العثور عليها في الأنسجة غير المصاباً. تقترح النتائج أن خلايا الرودلت تمثل نوعاً من الخلايا الالتهابية المرتبطة ارتباطًا وثيقًا بالخلايا الالتهابية الأخرى للأسماك والتي تستجيب لوجود طفيليات على الأسطح الطلائية.