Histopathological alterations in gills of freshwater prawn, *Macrobrachium dayanum* (Crustacea - Decapoda) after acute and sub-acute exposure of lead nitrate

Sanjive Shukla*
Department of Zoology, B.S.N.V. P.G. College, Lucknow-226001 (U.P.), India
Kunwer Ji Tiwari
Department of Zoology, University of Lucknow, Lucknow-226007 (U.P.), India
Harnam Singh Lodhi
Department of Zoology, Saket P.G. College, Ayodhya, Faizabad-224123 (U.P.), India
Sandeep Shukla
Department of Zoology, University of Lucknow, Lucknow-226 007 (U.P.), India
Anand Mishra
Department of Zoology, B.S.N.V. P.G. College, Lucknow – 226001 (U.P.), India
U. D. Sharma
Retd. Professor, Department of Zoology, University of Lucknow, Lucknow-226 007 (U.P.), India
*Corresponding author. E-mail: sanjiveshukla@gmail.com

**Abstract**

Lead is a nonessential “grey listed” heavy metal, used in fuels, ceramics, paints and glass wares in industries and vehicles. After taking entry in aquatic ecosystem it becomes toxic and cause serious problem to plants and animals. Haematological, neurological, nephrological and histopathological effects of lead are well known. Fresh water prawn, *Macrobrachium dayanum*, a potential animal for freshwater aquaculture, subjected to acute and sub-acute concentration of lead nitrate (116.46 mg/l; 96h LC₅₀ and 29.12 mg/l; 25% of 96h LC₅₀) showed severe histopathological alterations in gills after 24, 48, 72 and 96h acute exposure. The severity of histopathological alterations was found duration dependant. Present study reveals that histological bio-markers provides complete information regarding heavy metal toxicity particularly lead to these economically important fresh water prawns, which can themselves serve as bio-indicator of worsening status of surface aquatic bodies.

**Keywords:** Fresh water Prawn, Gills, Histopathology, Lead toxicity, *Macrobrachium dayanum*

**INTRODUCTION**

Population explosion, rapid industrialization, intensive chemical use in agriculture and anthropogenic activities led to intensive deterioration of aquatic ecosystem there by adversely affecting the aquatic flora and fauna (Jarup, 2003; Sharma and Agrawal, 2005). Among toxicants, heavy metals are lethal because of their long life half, persistence, bioaccumulation and biomagnification tendency in the food chain there by increasing the problem many folds (Burman and Lal, 1994; Rouzbahani, 2017). Now a day’s heavy metal contamination has become a major environmental issue (Mason, 1996; Chopra et al., 2009; Kumar and Chopra; 2015).

Among heavy metals, lead is a ubiquitous environmental contaminant and belongs to the group of most toxic heavy metal in the biosphere. It is considered as a non-specific poison affecting physiological systems and can cause damage to central nervous system, kidney, gastrointestinal distress and reproductive disorders (Kutlu and Sumer, 1998). Toxic effect of Lead and other heavy metals on gill histopathology have been mostly investigated in fishes (Gill et al., 1988; Khangarot and Tripathi, 1990; Kumari and Kumar, 1995; Parashar and Banerjee, 2002 and Olojo et al., 2005; Mobarak and Sharaf, 2011). Crustaceans, despite being important member of food chain and having high economic and medicinal value are being documented less in reference to gill histopathology against metal toxicity (Costa, 1965; Anderson and Baatrup, 1988; Jadhav, 1993 and Sen et al., 2008; Yamuna et al., 2009; Wu et al., 2009; Mishra et al., 2010).
Smaller fresh water prawns like *M. lamarrei* and *M. dayanum* despite having economic value, potential for good lab model and bioindicator as well as having potential for their aquaculture have not been taken into consideration except few studies (Shukla And Sharma, 2010; Verma et al., 2010; Tripathi et al., 2019). Crustacean gills are important for respiration, excretion, acid-base balance and osmotic and ionic regulation (Pequex, 1995; Soegianto et al., 1999; Lucu and Towle, 2003; Sabu et al., 2017). Gills are the first organs which comes in direct contact with the pollutants present in surrounding medium and are the major sites through which waterborne pollutants take entry in the body as well as affecting gills itself (Soegianto et al., 1999a).

Considering histology as a promising tool which provides complete picture of cell and tissue damage, the present work was undertaken to evaluate the histopathological effects of lead nitrate on gills of fresh water prawn, *Macrobrachium dayanum* (Crustacea–Decapoda), an untapped animal resource for freshwater crustacean aquaculture.

**MATERIALS AND METHODS**

Fresh water prawns, *M. dayanum* (Henderson) – INDIA, with the help of local fisherman and brought to the laboratory (N–26° 49' 55” E–80° 55’ 58”) in large plastic containers. The animals were maintained in glass aquaria of 20 liter capacity containing 10 liter dechlorinated water having physico-chemical characteristics as follows: - pH – 7.66 ± 0.27, Temperature – 27.66 ± 0.66°C, D.O. – 6.6 ± 0.74 mg/l, Total Hardness – 268±2.67 mg/l, Alkalinity – 425±11.36 mg/l (Sharma and Shukla, 1990; APHA et al., 1998).

Stock solution of lead nitrate (Pb(NO\textsubscript{3})\textsubscript{2}), AR Grade molecular weight 331.21 gm/mole, (E-Merck (INDIA) Ltd. Worli Mumbai-400018) was prepared by dissolving weighed amount of salt in double distilled water. Lead nitrate was dissolved in water by adding, 0.3 mg/liter of concentrated Nitric acid (Devi and Fingerman, 1995).

Adult inter-moult staged *M. dayanum* (Average length – 5.64 ± 0.42 cm; weight – 3.262 ± 0.68 gm.) were being utilized in experiments after 5-7 days acclimation to laboratory conditions. Acute exposure was carried out on 96h LC\textsubscript{50} (116.46 mg/liter) for 24, 48, 72 and 96h while sub-acute exposure was carried out on 25% of 96h LC\textsubscript{50} (29.12 mg/liter) for 10, 20 and 30 day respectively. One aquarium containing diluents water and 0.3 ml/liter conc. Nitric acid only served as control for each set. Feeding was suspended 24h prior to acute exposure and through out experiment while change of exposure medium and food supply were maintained on alternate day during sub-acute exposure. Continuous air supply was provided by air diffusers and aerators in both control as well as experimental aquaria in both the experiments. Both acute and sub-acute experiments were carried out according to guideline of APHA et al. (1998). Experiments were replicated thrice. Histopathological observations were taken into account after 24, 48, 72 and 96h in acute, while after 10, 20 and 30 day in sub-acute exposure on live animals only. Gills of control as well as experimental animals were carefully dissected out under stereoscopic dissecting binocular, washed in Crustacean Ringer’s solution (Brown and Creedy, 1970) and fixed in alcoholic Bouin’s fluid for 24h. Tissue were dehydrated and imbedded in paraffin wax (60 – 62°C). Serial sections of 6 - 8µ thickness were cut on rotary microtome, flattened on albuminized slides and stained with Ham’s Haemotoxylin and Eosin by routine histological procedure (Drury and Wallington, 1980). Stained sections were observed and photographed under Olympus trinocular microscope comparing with controls.

**RESULTS AND DISCUSSION**

Freshwater prawn, *M. dayanum* consists of eight pairs of gills on lateral sides which are typically phyllobranch type and almost similar to *Palaemon* (Patwardhan, 1937). Gill consists of gill base on which two rows of gill plates are found. Rhomboidal gill plates are arranged on the gill base like the leaves of a book. Gills are attached to the body through gill roots. The gills of control prawns (Plate-1, Figs. 1-2) appeared normal with spaced gill plates; cuticle appeared smooth and regular along with normal number of circulating haemocytes. Gills are the primary site which directly comes in contact of the surrounding medium and are affected by the toxicants. Gills, the chief respiratory organs of fresh water prawn, *M. dayanum* showed pronounced histopathological changes after acute and sub-acute exposure of Lead nitrate are as follows:

- After 24 hr exposure (Plate 2, Figs. 1, 2), inflammation, vacuolization and distention of gill plates were observed. After 48 hr exposure (Plate 2, Figs. 3, 4), cuticular eruption and bulging of gills plate tip were common. Hyperplasia was seen at most of the places. Mucous coating was observed on gill plates. After 72 hr exposure (Plate 2, Figs. 5, 6), most of the gill plates were found clumped due to hyperplasia reducing interlamellar space further. Leakage of haemocytes and tissue debris in the interlamellar spaces were prominent. Circulating haemocytes showed pyknosis and karyorrhexis. After 96 hr exposure (Plate 2, Figs. 7, 8), due to severe necrotic and degenerative changes general architecture of gills was found almost lost. Tips of gill plates were bulge and almost found stucked together along with granular deposition in it. Haemocytes and tissue debris were found filled in interlamellar spaces.
Plate 1-Figs. 1-2. Photomicrographs of T.S. of Gill (Control), Scale bar= 50 μ [CU=Cuticle, GE=Gill Epithelium, GP=Gill Plate, GPT=Gill Plate Tip, ILS=Inter lamellar space]

Plate 2-Figs. 1-8. Photomicrographs of T.S. of Gills showing the effects of acute exposure of Lead nitrate, Figs. -1 and 2 after 24 hr; Figs.-3 and 4 after 48 hr, Figs.-5 and 6 after 72 hr; Figs.-7 and 8 after 96 hr, Scale bar= 50 μ.
After 10 day exposure (Plate 1, Figs., 4), inflammation and distention in the gill plates was observed. Cuticular eruption and necrosis was common in most of the gill plates while hyperplasia was seen at some places. Very mild leakage of haemocytes in inter tubular spaces was found. After 20 day exposure (Plate 1, Figs. 5, 6), inflammation and hyperplasia was found increasing in gill plates which resulted in highly reduced interlamellar spaces. Increased influx of haemocyte was noticed at this stage. Most of the gill tips were found filled with Granular haemocytes and Cystocytes. After 30 day exposure (Plate 1, Figs. 7 and 7a, 8 and 8a), hyperplasia was pronounced in most of gill plates. Necrotic and degenerative changes were common showing pycnosis and karyolysis. Cuticular eruption was highly common. Gill tips were highly bulged and found filled with granular deposits and haemocytes. Most of the gill tips were found stucked together leaving almost no inter lamellar spaces.

Freshwater prawn, *M. dayanum* consists of eight pairs of gills or branchiae which are typically phyllobranch type having rhomboidal gill plates arranged on triangular gill base (Patwardhan, 1937).

Gill are not only the organ of respiration but are also potent osmoregulatory structure and comes in direct contact of surrounding medium and toxicant present in medium hence are more affected than any other part of the body. Heavy metals are well known to reduce oxygen consumption as well as disruption of osmoregulation in crustaceans (Jadhav, 1993; Sen et al., 2008; Asih et al., 2013; Putranto et al., 2014; Soegianto et al., 2016).

In the present study, histopathological alterations like inflammation, vacuolization, distention of gill plates, hypertrophy and cuticular irruption, cell necrosis, bulging in gill plate tip, hyperplasia, heavy influx of haemocytes, thickening of gill plates and reduction in interlamellar space, collection of granular material below cuticle and leakage of haemocytes in interlamellar spaces, mucous coating after 96 hr acute exposure and clumping of gill plates, necrotic and degenerative changes showing pycnosis and karyolysis, increased influx of haemocytes, fragmentation of nuclear material in haemocyte and aggregate of haemocyte noticed in haemolymph channel in gill base after 30 day sub-acute exposure, were the major alterations after lead nitrate exposure of *M. dayanum*.

Plate 3-Figs.1-6. Photomicrographs of T.S. of Gills showing the effects of sub-acute exposure of Lead nitrate, Figs. -1 and 2 after 10 day; Figs. -3 and 4 after 20 day; Figs.- 5,5a, 6 and 6a after 30 day. Scale bar= 50 μ
Gills showed almost complete disorganization after acute and sub-acute exposure of Lead in *M. dayanum*. Various workers reported almost similar histopathological alterations in gills of fishes and crustaceans after exposure of Lead and other heavy metals (Costa, 1965; Anderson and Bastrup; 1988; Gill et al., 1988; Khangarot and Tripathi, 1990; Kumari and Kumar, 1995; Jadhav, 1993; Dutta et al., 1996; Parashar and Banerjee, 2002; Olojo et al., 2005; Ashi et al., 2013; Putranto et al., 2014; Soegianto et al., 2016). The heavy metal lead was toxic to freshwater fisheries products.

The heavy metal lead was toxic to freshwater fisheries products. Histology of gills can serve as potential and foremost bio-marker in comparison to isolate physiological and biochemical parameters especially in reference to metal toxicity. *M. dayanum* can serve as better bio-indicator in assessment of health of fresh water aquatic bodies.

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