Abstract: Water heavy metals contamination is an international issue due to its detrimental impact on the environment, aquatic organisms and impairment of the ecological balance systems. Cadmium is a heavy metal that is extremely toxic, even in low doses, thus it is a big concern to the ecosystem. Cd can cause several Patho-morphological changes in various organs in fish. This study was evaluated the effect of sublethal concentrations of Cd toxicity and effects on Histopathological of muscle tissues of Planiliza abu Juveniles. Series of Cd concentrations (0.5-2) ppm were used and the 15d LC₅₀ was 0.4 ppm. Several histopathological alterations were observed which increase gradually with increased concentration, included, marked thickening and hypertrophy of muscle bundle were will observed in lowest concentration (0.5 ppm). While in the highest concentration (2 ppm), atrophy, splitting, and separation with intracellular edema were marked in fish muscle bundle.

Key words: Planiliza abu. Cadmium. Sublethal concentrations. Muscles. Histopathology.

Introduction

Human activities play a big role to increase metal pollution in waters (Popek et al., 2006). Because it is threatening aquatic life, available in ecosystem (Abo El Ella et al., 2005) and their persistence in the environment (Biswas et al., 2012). Heavy metals might be entered from different sources such as soil and anthropogenic activities (Abo El Ella et al., 2005) such as, use of fertilizers and pesticides in agriculture and decomposition of organic matter (Bai et al., 2010). Some metals are considered potentially toxic (As, Cd, Pb, Hg, etc.), while others are essential (Cu, Zn, Fe and Mn) (Biswas et al., 2012). Cadmium is an extremely toxic metal, even in low doses. It is used in industry specifically in electroplating, industrial paints and batteries (Taylor, 1997). Cd is a natural element nonessential element, due to its ability to transfer, deposition and dissolved within ecosystems, this leads to concerns in terms of accumulation in the aquatic food chain (Guardiola et al., 2013). Cd element and its concentrations have been detected in the air, water, soil, and plants (Amdur et al., 1992). Cd serves as a big concern to the ecosystem, because it has the ability to accumulate in living organism (Mohamed, 2008). Raissy et al. (2011) found the presence of Cd in the aquatic animals and ecosystem only in the range of trace concentrations. The concentration of Cd range in the ocean seawater from approximately (0.01-42) µg.l⁻¹.
Over time the pollution affecting fish could cause dangerous human health problems by concentrated in an aquatic marine. (Authman et al., 2012). Environmental pollutants can induce oxidative stress in aquatic animals (Ceyhun et al., 2011), for example, Cd can cause hyperactivity and excitability in animals, which can lead to muscular fatigue. Muscle tissue can have direct contact with pollutants (Das & Mukherjee, 2000). Cd also can cause several patho-morphological changes which can be showed in different tissues of aquatic fish. (Beširović et al., 2010).

Humans can be exposed to metals through many routs such as, lungs and skin, drinking contaminated water, and eating contaminated food (Wright & Welbourn, 2002). The impact of Cd toxicity on human health showed on Minamata was Itai-Itai (Ouch-Ouch) Disease in Japanese people (Jintsu River Basin) with many symptoms such as complains of joint, bone, and muscle pains and severe kidney dysfunction (Landis et al., 2003; Bradl, 2005). Cd is toxic to humans when the daily intake is 250 to 300 micrograms (Wright & Welbourn, 2002) and it is found highly persistent in humans, with a biological half-life of 20 to 30 years (Wright & Welbourn, 2002; Landis et al., 2003).

Fish play an important role in human diet, because fishes are rich in omega-3 fatty acids, which are important to maintain balanced levels of blood glucose in the body (Qiu et al., 2011). In addition, it has an important nutrient, due to protein, iodine, different vitamins, minerals and necessary lipid compounds (Vieira et al., 2011). It has a big role in the food chain and is an important marker for many problems in aquatic ecosystems (Aktar et al., 2011). Nevertheless, the contaminated water could have negative effects on the function of the omega-3 fatty acids and protection human health against heart disease. (Ferreira et al., 2010). Among the many aquatic organisms, fish are considered a good indicator of the effects of pollutants absorbed (Shukla et al., 2007). The process of heavy metals storing in animals’ tissue different among animals’ types based on the kind of metal (Patnaik et al., 2011). The sublethal concentrations of Cd and lead can bioaccumulate in different tissues, which lead to damage and impaired physiology (Patnaik et al., 2011). The higher bioaccumulation rate was for Cd in fish tissues and many effects on tissue Histology (Vinodhini & Narayanan, 2008). Ebrahimi & Taherianfard (2011) show that the maximum concentrations of Cd in fish tissues were higher than the permissible levels for human consumption. Therefore, the study of fish tissues is an important mechanism to evaluate the health condition of the fish community. (McHugh et al., 2011). Heavy metals toxicity and muscles Histological changes in fishes have been reported by (Mohammed, 2009; Alak et al., 2013b; Begum et al., 2013; Badr et al., 2014).

Planiliza abu is an economically and ecologically important species as well as a human food. It has been detected through the basin of the Euphrates and Tigris rivers, as well as the Mediterranean basin of Southeastern Turkey (Jawad et al., 2015). Therefore, the objective of this study was to evaluate the effects of cadmium toxicity through the sublethal concentrations exposure and effects on Histo-pathological aspects of muscles structures on Juveniles of P. abu.

Materials & Methods

Collection and Maintenance

P. abu collected from marine science station aquariums. Fishes were acclimated under laboratory conditions at 23 ±1 °C and 4 psu
salinity in 10-gallon aquaria (60 X 30 X 30) cm using a 14-h light: 12-h dark photoperiod. The Juveniles mean weight were (10.4-19.8) g and the mean length were (9.6-18.4) cm. Water quality parameters were recorded every day throughout the acclimation period, including water temperature (thermometer) °C, salinity (American Optics Refractometer-psu), dissolved oxygen (YSI Oxygen Meter Model # Quatro Cable Assay, mg.l⁻¹) and pH (pH meter). Juveniles were daily fed Tetramin® Fish Flakes during lab acclimation but, were not fed during acute toxicity tests.

Preparing of concentration stock (LC₅₀)

The stock solution was made by solving (6.8466) g of aqueous cadmium sulfate (CdSO₄·8H₂O) in one liter of distilled water that is equal to (1000) ppm. A required weight calculated according to the following equation: Molecular weight of a compound/ Atomic weight of the element = weight of the compound required gm of per liter (gm.l⁻¹).

Calculating the median-lethal concentration

Each aquarium (60 x30 x30) cm contained 15 animals with three replicates/ cadmium concentration and controls (n=15 fish/treatment) in each aquarium. Aqueous static renewal tests were conducted to determine 15days LC₅₀ (median lethal concentration) the values concentrations for cadmium under laboratory conditions.

Histo-pathological examination

Parts of skeletal muscles from the control and experimental animals were taken from juveniles exposed to different concentrations of cadmium (0.5, 1 and 2) ppm, fixed in Bouin’s solution for 48 hrs. and dehydrated in a graded series of ethyl alcohol (70%, 80%, 90%, 95% and 100%) then cleared in xylene and embedded in paraffin, and sectioned at 5 mm thick. Tissue stained with hematoxylin-eosin method (Humason, 1979) and examined by light microscopy according to Roberts (2001).

Results

Test conditions measured in the cadmium toxicity test are listed in (table 1). After 15days of exposure there were differences in the toxicity of cadmium at all concentrations. Typically, mortality rate increase with increasing cadmium concentrations. For example, the mortality in 0.5, 1 and 2 ppm were (50%,70% and 92%) respectively (table 2). The 15days LC₅₀ of cadmium was 0.405 ppm, (Fig. 1).

Plate (1) shows numerous Histo-pathological alterations in the muscles of P. abu. juveniles exposed to different concentrations of cadmium. They are increased gradually with increasing Cd concentration. The pathological findings included marked thickening and hypertrophy in lowest concentration (0.5 ppm), degeneration along with coagulative necrosis of muscle bundles have been reported in fish exposed to (1 ppm) of Cd. however, muscle tissue showed atrophy, splitting, and separation with intracellular edema, at the highest concentration (2 ppm).
Table (1): Water quality conditions measured in cadmium toxicity tests.

| Water quality parameter | value measured |
|-------------------------|----------------|
| Temperature             | 23 ±1°C        |
| Salinity                | 4 psu          |
| pH                      | 7.7            |
| Dissolved oxygen        | 7.50 mg. l⁻¹   |

Table (2): Mortality % and LC₅₀ for juveniles *P. abu* exposure to different concentrations of Cadmium (ppm) after 15 days of exposure.

| Duration | Concentrations (ppm) | Mortality % | LC₅₀ (ppm) |
|----------|----------------------|-------------|------------|
| 15-day   | 0.5                  | 50%         | 0.405      |
|          | 1                    | 70%         |            |
|          | 2                    | 92%         |            |

Fig. (1): Mortality percentage of *P. abu* Juveniles exposed to different concentrations after 15 days.
Plate (1): Photomicrograph of the normal structures of the skeletal muscle (Control) of *P. abu* showing; capillaries, bands light and bands dark; (b) & (c) Photomicrographs of muscles (Cadmium treated 0.5) ppm showing; MTMB, marked thickening of muscle bundles, H, hypertrophy; (d), (e), Photomicrographs of muscles (Cadmium treated 1) ppm showing; VD, vacuolar degeneration, NC, coagulative necrosis; (f), (g), (h) Photomicrographs of muscles (Cadmium treated 2) ppm showing; AT, atrophy, SP, splitting, and SE, separation of the muscle bundles with intracellular edema. (H&E). Scale Bar = 40 µm.
Discussion

The mortality percentage results of *P. abu* juveniles exposed to different concentrations of cadmium (0.5, 1 and 2) ppm were 50%, 70% and 92% respectively (table 2). The 15 days LC50 of cadmium was 0.40 ppm (figure1). This agrees with Hassan (2005) when exposed juvenile of *Carassius carassius* for 14 days to sublethal concentrations of Cd (0.05 & 0.5) ppm the mortality percentage were (70% & 40%) respectively, while the LC50 was (0.35) ppm. Sobha et al. (2007) proved that mortality increase over the time, LC50 value of cadmium chloride for the fish *Catla catla* was 4.533 mg.l−1. Hassan (2011) observed the juvenile of common carp *Cyprinus carpio* exposed to sub-lethal concentrations of copper and cadmium (0.1, 0.25, 0.4) ppm led to decreased in survival rate with increasing concentration of copper from 50% to 10% and cadmium from 70% to 20% respectively.

Metals detrimental impact are evident in the environment, aquatic organisms and impairment of the ecological balance systems (Vosyliene & Jankaite, 2006). Moreover, heavy metals could effect on the quality and quantity of fish stocks because of their effects on water quality, physiochemical characteristics and biological components (Mantovi et al., 2005; Singh et al., 2007). Heavy metals have adverse effects on aquatic organism including survival, activity, growth, metabolism and reproduction (Wright & Welbourn, 2002). The aquatic organisms can be exposed to the heavy metals by many routes such as drinking water, eating contaminated sediment and other animals/plants that have been exposed (Wright & Welbourn, 2002). Therefore, fish and other aquatic organism can absorb dissolved metals in water and accumulate in tissues higher than in water, because metals undergo bioconcentration (Wright & Welbourn, 2002). Heavy metals can accumulate in aquatic organisms including fish tissues more than the water concentration with a range up to 20,000-fold (Popek et al., 2006).

This study examined the Histological changes in fish muscles were increase gradually with Cd concentration including, marked thickening and hypertrophy in lowest concentration (0.5) ppm (plate 1). Exposure to the sublethal concentrations of cadmium recorded number of changes in the muscle’s bundles including thickening and intramuscular oedema (Patnaik et al., 2011).

Exposure to the waterborne Cd in aquatic organism causes a significant accumulation of Cd in many organs like muscles (Alak et al., 2013b). Cd can accumulate in the aquatic animals through different routes such as food and water. It then binds to albumins and erythrocytes in blood, which can transfer it into different parts of the body (George et al., 1996; Karadeniz et al., 2009). Damage caused by heavy metals is an important cause of pathological and toxicological processes (Ates et al., 2008). In generally, Histopathological studies can be used as an indicator for determining the effects of heavy metals on aquatic organisms. (Bernet et al., 1999; Mohamed, 2009). The Histopathological alteration was caused by cadmium poisoning (Alak et al., 2013b).

However, in (1) ppm concentration was showed vacuolar degeneration and necrosis of muscle bundles. The results are in accordance with those obtained by (Mohamed, 2009). Different Histological alterations were observed in the muscles of both fish effects of contaminated drainage water from Lake Qarun, Egypt on *Coptodon zillii* (= *Tilapia zillii*) and *Solea vulgaris*, which included
vacuolar degeneration and necrosis of muscle bundles. Histo-pathological examinations on the muscles were found at hypodermis of Nile tilapia Oreochromis niloticus and which included degeneration (myolysis) and arrangement exposed to different metals (Cu, Zn, Cd and Pb) (Ibrahim et al., 2008). Several Histo-pathological alterations in muscle of two types of fish O. niloticus and Lates niloticus, were discovered as a result of the accumulation of metals. One of the Histo-pathological alterations was degeneration in muscle bundles (Mohamed, 2008). Histological examination of O. niloticus muscles showed congestion associated with diffuse infiltration with inflammatory cells, hyperactivity of melanomacrogaphre centers around blood vessels, and focal area of necrosis (Badr et al., 2014).

On the other hand, the highest concentration (2) ppm was showed atrophy, splitting, and separation (plate 2). These results were confirmed with Begum et al. (2013) when exposing stinging catfish Heteropneustes fossilis exposed to (7.0 and 20.0) ppm of arsenic caused a significant increase in Histo-pathology of muscles bundle such as separation atrophy and splitting. Furthermore, when rainbow trout exposed to different sublethal concentrations (0.18 and 0.35) mg. 1\(^{-1}\) of lead caused a significant increase in Histo-pathology in muscles such as myocyte necrosis and atrophy (Alak et al., 2013a). While sublethal concentrations of cadmium caused intramuscular oedema (Patnaik et al., 2011). Brown trout were exposed to 2) ppm of cadmium can cause interstitial mononuclear cell infiltrations in muscle tissues. (Alak et al., 2013b). Some structural changes were observed due to lead acetate exposure, leading to space in muscular bundles and glomerular expansion in Clarias gariepinus (Al-Balawi et al., 2013). Exposure of Cyprinus carpio, to sublethal concentrations of cadmium after 28 days of exposure to (1.6) ppm has led to damage in fish muscles tissue Patnaik et al., 2011). After 60 days of exposure, there were partially damaged muscles, and slightly swollen/shortened villi (Begum et al., 2013).

According to the Authman et al. (2012) exposed Nile tilapia (O. niloticus) to different concentrations of metals (Al, Cd, Pb, Hg and Ni), caused various Histo-pathological in muscles bundles.

The concentrations of cadmium in muscular tissue of O. niloticus and the concentrations in water, from contaminated fish farms, were (4.354 and 1.315) mg.l\(^{-1}\) respectively (Authman et al., 2012). The order of fishes tissues exposed to different concentrations of Cd were as follows: muscle >gill > liver > heart (Mohamed, 2008). Ibrahim et al. (2008) monitors the effects of metals on the biological, Histological and quality aspects of fish. The metals included (Cu, Zn, Cd and Pb) on of fish O. niloticus and C. zillii. Authman et al. (2012) found that heavy metal levels in O. niloticus muscles were not possible range of human consumption.

**Conclusion**

The results were revealed that the mortality increases with increasing cadmium concentrations. Numerous of Histo-pathological changes of Planiliza abu Juveniles muscles have been noted. The Histological changes increased with increasing the concentration of the cadmium included, marked thickening and hypertrophy in lowest concentration (0.5 ppm) while atrophy, splitting, and separation, were observed in the tissues of fish muscles bundles at the highest concentration (2) ppm. Future studies should be including testing of
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cadmium in different types of aquatic organism.

Acknowledgements

Authors are thankful to the Fisheries and Marine Resources department in Agriculture College of Basrah University for providing laboratory facilities during this study.

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تأثير التراكيز تحت المميتة للكادميوم على التغيرات المرضية النسيجية لعضاة اسماك الخشني Planiliza abu (Heckel, 1843)

ال연구ة

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المستخلص:
تلوث المياه بالمعادن قضية عالمية، بسبب تأثيره الضار على البيئة والكائنات المائية وضعف نظام التوازن البيئي. الكادميوم معبد نقل شديد السمية، حتى في الجرعات المنخفضة، وبالتالي فهو مصدر قلق كبير لنظام البيئي. يمكن أن يسبب الكادميوم العديد من التغيرات المرضية في الأعضاء المختلفة في الأسماك. بنيت هذه الدراسة أثر التراكيز تحت المميتة لسمية الكادميوم وأثارها على الأنسجة المرضية للأنسجة العضوية على يافعات الخشني Planiliza abu. كانت تراكيز الكادميوم تتراوح من 0.5-2 جزء في المليون لمدة 15 يومًا من التعرض. بلغ الترتيك المميت LC50 (0.40 جزء في المليون). لوحظ أن العديد من التغيرات النسيجية زادت بشكل تدريجي مع تركيز الكادميوم. بينما في ذلك سماكة ملحوظة وتضخم في أقل تركيز (0.5 جزء في المليون). بينما في أعلى تركيز (2 جزء في المليون)، كانت الأنسجة المرضية ضمورًا، انفصالًا، انصفاً، وانفصالًا، في حزم عضلات. 

الكلمات المفتاحية: الخشني، الكادميوم، تركيزات تحت المميتة، عضلات، أمراض نسيجية.