INTRODUCTION

Fish are exposed to various contaminants or pollutants in the aquatic environment leading to decreased growth, alterations in physiological processes and ultimately death. These contaminants which are obtained from the aquatic environment and food cover a wide spectrum of gases, chemicals and solids. Most toxicological studies have been limited to the effects of lethal or acute doses of these pollutants (Martinez & Souza 2002, Huertas et al. 2002, Das et al. 2004a), whereas, subtle physiological disorders result with exposure to sub-lethal/low doses of toxicants (Das et al. 2004b, Varo et al. 2007, Kori-Siakpere et al. 2011). Various workers have reported histopathological changes in fish organs (e.g. liver, gills, kidney, lungs etc) exposed to sub-lethal concentrations of pollutants (Stentiford et al. 2003, Rabitto et al. 2005).

Phostoxin which is either in form of tablets or pellets contains 55% aluminum phosphide as its active ingredient and 45 % inert which is used as the carrier of the active ingredient. The use of synthetic pesticides in grain storage is commonly practiced in the West Coast of Africa, particularly in Nigeria. This method of grains preservation by farmers has received strong encouragement from the government for the purpose of achieving food security (Atta et al. 2009). Unfortunately, most of these pesticides have shown to be of health risk to man and the environment (Ofuya et al. 2010).

Aluminum phosphide (phostoxin) commonly called “trebor” by local farmers in Nigeria is used for the eradication of weevils in stored grains especially in maize. In some cases, the tablets are mistakenly ground with the maize used as an ingredient in fish feeds and are fortuitously consumed by fish in their diets (Atta et al. 2009). Although aquatic life can tolerate some measure of stress and occasional adverse effects (Shhuaimi-Othman 2012), the quantity of aluminum phosphide in the aquatic food chain might be sufficient to cause changes in cyto-architecture of the vital organs resulting to alteration in normal body function and growth.

Histopathological effect of sub-lethal concentration of aluminum phosphide (phostoxin) on *Clarias gariepinus* juveniles

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The study evaluated the effect of sub-lethal concentration of phostoxin on *Clarias gariepinus* juveniles. *C. gariepinus* juveniles belonging to the same cohort (40.1±1.2g; 18.1±1.1cm) from a commercial fish farm were randomly placed ten in each of 15 plastic tanks containing 15 liters of water. They were exposed for 96 hrs to three sub-lethal concentrations (treatments) of phostoxin (0.125, 0.250, 0.5mg L⁻¹) and a phostoxin free control. At the end of 96 hrs exposure, they were dissected and the tissues need for histopathology removed and fixed in Bouin’s fluid. The gill filament exhibited fusion at the secondary lamella that was progressive with concentration. At the highest concentration of exposure, the secondary lamellae showed marked pyknotic and necrotic changes characterized by epithelia detachment. The hepatic tissue showed mild inflammatory changes at lower concentrations while at the highest concentration of exposure there was marked inflammation with observed hydropic degeneration. In the kidney, an inflammatory change was only observed in the interstices at the highest dose of exposure with the convoluted tubules showing partial shrinkage. Phostoxin showed to have significantly caused alterations in cyto-architecture of the gills and to a considerable extent liver and kidney of *C. gariepinus*. 

INDEX TERMS: Histopathology, phostoxin, toxicity, organs, *Clarias gariepinus*, juveniles.

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ingredient. When exposed to water reacts to liberate phosphine gas as shown in the equation below (Gordon 1972). Phosphine gas is by far the dominant means of controlling pest insects in stored grain and many other stored commodities (Nath et al. 2011). The phosphine gas emitted is the poison used to kill insect pests in grains without affecting grain viability (Atta et al. 2009). Because phosphine gas is highly toxic to aerobic organisms it could therefore be of considerable health risk to aquatic life. The phosphine gas is colorless and odorless in its pure form but, due to the presence of substituted phosphines and diphosphines, it has a foul odor resembling decaying fish (Chugh 1992). Phosphine is gaseous above -88°C which allows it to disperse readily during fumigation. The concentrated phosphine is potentially explosive in air and can autoignite at near ambient temperatures. However, the commercial aluminium phosphide contains ammonium carbamate which liberates non-flammable and inert agents to reduce fire hazards (Nath et al. 2011):

\[ \text{AlP} + 3\text{H}_2\text{O} \rightarrow \text{Al(OH)}_3 + \text{PH}_3 \]

The mechanism of cytotoxicity of aluminum phosphide on aquatic life is not fully understood. However, in rainbow trout, the reported acute LC50 is 4.1μg/L, indicating very high toxicity (Leuschner 1984). Aluminum phosphide will rapidly react to form phosphine gas which is somewhat soluble in water. The noxious gas which is not known to be absorbed dermally have their main routes of exposure to the body systems through ingestion and inhalation and have shown to be highly toxic via both routes (Degesch 1988). The contact of the gas with the gills readily affects the secondary lamellar, a tender projection from the primary lamellar having delicate epithelial lining for easy gaseous exchange. As reported (Ayoola & Ajani 2007), the effect of the gas on the delicate cellular membrane result to mucus accumulation that cause hypoxia which could lead to death. On the other hand, the ingested phosphine gas is rapidly absorbed throughout the gastrointestinal tract, leading to systemic toxic effects. The action is modulated through an increase in superoxide dismutase activity and decrease in catalase levels which result in increased formation of free radicals and accelerated lipid peroxidation (Wahab et al. 2008). The increase in lipid peroxidation is known to cause membrane damage and then a change in cellular ionic balance leading to cells death (Wahab et al. 2008).

**Clarias gariepinus** is a North Africa catfish that live in a variety of fresh water environment including quiet waters, lakes and pools and prefer rather shallow and swampy areas with a soft muddy substrate and calmer water. They may also occur in fast flowing rivers and in dams (Teugels 1986). They are widely distributed to different parts of African continent being native to streams in the Tolga oasis in Algeria. However, their cultivation in fish farms in Europe, Asia and South America have made them wide spread around the world (Vitule et al. 2006). In Nigeria, with an up surge in demand for fish, there is a huge expanding population of *C. gariepinus* which is the most cultivated catfish in the country aquaculture (De Graaf & Janssen 1996).

**Materials and Methods**

**Experimental fish.** *Clarias gariepinus* juveniles belonging to the same cohort (40.1±1.2g; 18.1±1.1cm) from a commercial fish farm were transported to zoology laboratory in plastic containers to static bath systems well aerated. They were acclimatized in laboratory condition within the temperature range of 15-22°C for two weeks and fed a commercial diet before the commencement of the experiment. Unconsumed feed and faeces were removed daily and water replenished regularly. The investigation was conducted in the month of November 2011.

**Laboratory methods.** Ten fish were randomly placed in each of 15 plastic tanks containing 15 liters of water.

Fish were exposed for 96 hrs using a static bath system to three sub-lethal concentrations (treatments) of phostoxin (0.125, 0.250, 0.5mg L⁻¹) and a phostoxin free control. The doses were determined based on probit analysis which was conducted to determine the LC50 at 96 hrs and the value obtained was 2.75 mg L⁻¹. The 20 % of the LC50 value (2.75 mg L⁻¹) was used as the highest test concentration. The rationale for using 20 % of LC50 value of phostoxin as the highest dose was to keep within the limit of sublethal concentration. The two other doses used were a reduction in concentration of the highest dose in a graded manner. The half concentration of the highest dose (50 % reduction) was used as the second dose while the third dose was 50 % reduction in concentration of the second dose. This progressive reduction of dose in a graded manner was to establish dose dependent effect on the study. Each treatment was triplicated. Crushed tablets of phostoxin were added to the water in the tanks to obtain desired treatment concentrations. Fish were observed at one hour intervals to note behavioral changes in the first 24 hrs, after which observations were made at two hour intervals. The behavior, mortality and other external changes were recorded.

**Fish dissection.** At the end of the exposure period (96 hrs), a fish each was randomly taken from the treatment tanks, sacrificed under chloroform anaesthesia by placing the fish in a closed glass chamber containing soaked chloroform (CHCl₃, 99-99.6%) in a cotton wool. They were later dissected and the gills, hepatic and renal tissues need for histopathology were removed and fixed in Bouin’s fluid for 7 days. The above organs were considered for histopathological examination because in fish, the gill (respiratory organ) is the first contact to toxicant while liver is equally very susceptible to pathological changes because of its unique role in detoxification of ingested toxic materials. To a less extent, the kidney is likeable to inflammatory challenges due to the role it plays in aiding the removal of detoxified waste substances.

**Slide preparation.** The fixed target organs in Bouin’s fluid were removed and dehydrated in increasing concen-
Concentrations of alcohol: 70%, 80%, 90% and absolute alcohol (100%). The organs were treated with acetone and then cleared in xylene for 30 min to enhance the tissue transparency, followed by impregnating and embedding in paraffin wax. Each tissue was then sectioned at 5µm and cleared (dewaxed) for staining with haematoxylin and eosin (Mbaka et al. 2014).

RESULTS

Gills

Gill morphology of the control group (Fig.1a) showed primary and secondary lamellae which was typical of other teleost fish. The lamellae were covered by squamous epithelium composed of undifferentiated pavement cells. The epithelial lining of the primary lamellar was regular showing more than two cell layers with the chloride cells visible which appeared more predominant at the base of the secondary lamellar. Deep to the epithelium were blood sinuses located in the substance of the pillar cells. Traces of vascular channels were spotted within the muscular tissue. At the lamellar core was the rigid mass of cartilaginous tissue and around it were muscle fibres. The secondary lamellar was a tender projection from the primary lamellar having delicate epithelial lining for easy gaseous exchange while the pillar cells assist with support.

Clarias gariepinus was exposed in three different concentrations of phostoxin at sub-lethal levels for 96 hrs. At the end, several pathological changes occurred which increased with concentration. The main changes observed following exposure at 0.125 mg L\(^{-1}\) (Fig. 1b) was non continuous epithelial lining especially at the more delicate respiratory secondary lamellar indicating signs of hyperplasia and epithelial lesion. There was intermittent fusion at the adjacent secondary lamellar as a result of hyperplasia. At 0.250 mg L\(^{-1}\) dose (Fig. 1c), the adjacent secondary lamellar had become almost completely fused forming a continuous lining as a result of more extensive hyperplasia of more delicate respiratory epithelium. At 0.50 mg L\(^{-1}\) (Fig.1d), the secondary lamellae exhibited marked necrotic changes with complete disintegration of the lamellae characterized by accentuated epithelial lifting and detachment. There was intense vasodilatation and many vacuoles. The pillar cells had been altered and blood spaces expanded as a result of the supervening distress.

Liver

A section of hepatic tissue of the control group showed Fig.1. (a-d) Photomicrograph showing transverse section of gill filaments of Clarias gariepinus juveniles. (a) Control showing normal appearance of gill filaments, primary lamellae (PL), secondary lamellae (sc), chloride cell (cc) and polar cell (pc). (b) Gill of exposed at 0.125mg L\(^{-1}\) indicating mild hyperplasia with fused (fc) and un-fused secondary lamellar (vo). (c) Gill of exposed at 0.250mg L\(^{-1}\) indicating severe hyperplasia with complete fusion of secondary lamellae (fc). (d) Gill of exposed at 0. mg L\(^{-1}\) showing complete disintegration of secondary lamellae, epithelial lifting (ef) exposing central venous sinus (vn). Obj.40x.
normal architecture (Fig. 2a). It exhibited typical parenchymal appearance though with indistinct hepatic lobules. The polygonal shaped hepatocytes were arranged as irregular cord-like structure separated by sinusoids. Each cord extended from the peri-lobular margin and showed normal convergence to the central vein. The hepatic tissue after exposure at the dose of 0.125 mg L\(^{-1}\) (Fig. 2b) showed no apparent inflammation within the hepatic parenchyma. The hepatic sinusoids were however ill defined. At the dose of 0.250 mg L\(^{-1}\) (Fig. 2c) there was mild inflammatory changes indicating diffuse presence of lymphocytic cells. There was also an onset of edematous changes and vacuolation. At the dose of 0.5 mg L\(^{-1}\) (Fig. 2d), inflammatory changes had become more apparent indicating more profound edematous changes with the hepatic parenchyma shown to be eroded with infiltrating leukocytes. Also observed in the parenchyma were hydropic degeneration and pyknotic changes of the hepatocytes.

Kidney

The histology of the control group showed (Fig. 3a) normal architecture of the renal cortical tissue. The renal corpuscles appeared as rounded structure surrounded by narrow space, the Bowman’s space. The cortical tubules seen in this section consisted mainly of proximal convoluted tubules with few of the distal convoluted tubules indicated. After exposure at the dose of 0.125 mg L\(^{-1}\) (Fig. 3b) and 0.250 mg L\(^{-1}\) (Fig. 3c) respectively no noticeable distortion was observed. But at 0.5 mg L\(^{-1}\) (Fig. 3d) dose, mild inflammatory change occurred at the interstices. The convoluted tubules exhibited partial shrinkage with a coalesced space around.

DISCUSSION

Fishes are prone to various toxic challenges as these substances are freely disposed into the streams and rivers not considering their deleterious effect on aquatic life. (Jiranngkoorskul et al. 2002). Fish species are most sensitive to aquatic pollutants during their early life stages. It has been reported that when water quality is affected by toxicants, physiological changes occur which are reflected in behavioral changes and swimming activity of fish (Heath 1991, Adeyemo et al. 2004). In this study, severe abnormal behavior was observed such as incessant jumping and gulping of air, restlessness, loss of equilibrium, increase in opercular activity, sudden quick movement and occasional motionlessness at the bottom of the bath. This finding was in agreement with similar behavior observed in *Clarias gariepinus* exposed to certain pesticides (Omoniyi et al. 2002, Rahman et al. 2002). The stressful and erratic behavior of *C. gariepinus* indicated respiratory distress believed to have been caused by the undesirable effect of phostoxin on the gills.

Fig. 2. (a-d) Photomicrograph showing transverse section of hepatic tissue of *Clarias gariepinus* juveniles. (a) Control showing normal parenchyma indicating hepatocytes (h) and central vein (cv). (b) Exposure at 0.125 mg L\(^{-1}\) showing normal hepatocytes (h). (c) Hepatic tissue exposure at 0.250 mg L\(^{-1}\) indicating slight leukocytes infiltration. (d) Exposure at 0.5 mg L\(^{-1}\) indicating severe inflammatory changes with hydropic degeneration (hp). Obj.40x.
Histopathological changes have been widely used as biomarkers in evaluating the health of fish exposed to contaminants, both in the laboratory (Thophon et al. 2003) and in the field studies (Hinton et al. 1992, Teh et al. 1997). Juveniles of C. gariepinus of same cohort showed progressive pathological changes in their vital organs such as the gills and liver when stressed at three different sub-lethal concentrations of phostoxin. At the lowest dose of exposure, the gills showed mild hyperplasia with intermittent fusion at the secondary lamellar indicating inflammatory changes. There was also an indication of increased mucus secretion which has been reported in phosphine gas toxicity to fish (Ayoola & Ajani 2007). When the dose was doubled (0.250 mgL⁻¹), there was severe hyperplasia with nearly complete fusion of secondary lamellar. Lamellar fusion is seen as normal occurrence when fishes are exposed to toxic agents in water (Nowak 1992, Olurin et al. 2006). The reaction could be explained as adaptogenic which may be to minimize surface area contact to toxic agent or body changes in response to irritant substance. The increase in thickness of epithelial layers and fusion of adjacent secondary lamellae would not only decrease the surface area available for oxygen extraction but would also increase the oxygen diffusion distance between water and blood (Skidmore & Tovell 1972). Although this may have served as a protective effect against the noxious agent, it is however an impediment to respiratory process. At the highest dose of sub-lethal concentration of phostoxin exposure, there was complete disintegration of secondary lamellae, increased vacuolation, accentuated lamellae epithelia lifting and epithelia detachment as well as tissue hypoxia and arterial rupture due to the supervening respiratory distress. According to an earlier report, phosphine gas on exposure to fish causes an accelerated lipid peroxidation known to cause membrane damage (Wahab et al. 2008).

In toxic environments, the hepatic tissue is usually a prime target because of its unique role in detoxification process. Fish liver, however, has been reported not to show the diversity of pathological changes seen in higher animals on the account of lack of kupffer cells in the hepatic sinusoids (Ellis et al. 1976). Nevertheless, its susceptibility to a number of toxic and consequential metabolic disturbances has been highlighted (Jiraungkoorskul et al. 2003, Olojo et al. 2005, Mobarak & Sharaf 2011). At the lowest dose of the sub-lethal concentration, there were no traces of inflammatory changes although the sinusoids were ill defined. At increased concentration (0.250 mgL⁻¹) there were signs of leukocytes infiltration and increased vacuolation which were manifestation of inflammatory changes. Inflammatory response is seen as the basic protective response to tissue damage and it may be considered as the product of series of changes that takes place following injury. At the highest concentration (0.5 mgL⁻¹) of exposure, more severe inflammatory changes occurred that include hydropic de-
generation and pyknotic changes of hepatocytes. The hepa-
tocytes appeared disarranged while the hepatic sinusoids
were no longer outlined. It has been established that fish
exposed to toxicants exhibit increase in hepatic vacuol-
zation (Coimbra et al. 2007, Fiuza et al. 2009, Mobarak &
Sharaf, 2011). The increase in vacuolation could have been
as a result of imbalance between the rate of synthesis of
substances and the rate of release to systemic circulation
(Ginerich 1982, Jiraungkoorskul et al. 2002). Pyknosis and
karyolysis have been reported in cases of severe poisoning.

The glomerular complex is vital to renal filtration and
therefore its lesion would lead to impaired renal function.
In this case, no lesion was observed in the kidney after ex-
posure to phostoxin at 0.125mg L⁻¹ and 0.250mg L⁻¹ dosages
respectively while mild inflammatory changes occurred
in the interstices at the highest dose of exposure particularly
with the shrinkage of convoluted tubules. The sub-lethal
concentration of phostoxin used and the duration of expo-
sure may have accounted for less effect of the toxicant on
the kidney. It could also be viewed in the context that the
organ was not the first contact to the toxicant. More often
ingested toxic substances are detoxified in the liver and
before it can reach potentially toxic level to have affected
other internal organs like the kidney, it would have caused
extensive damage to the liver. Damage to kidney has been
observed following fish exposure to toxicants (Jiraungko-
orskul et al. 2003).

CONCLUSIONS

The sub-lethal concentrations of phostoxin exhibited
wide spectrum of histopathological changes on Clarias
gariepinus in which significant alterations occurred in the
tissue morphology and in the cyto-architecture of the gills
and to a considerable extent liver and kidney of the ex-
posed fingerlings.

In the light of the fact that marked deleterious effect oc-
curred at the sub-lethal doses, phostoxin should therefore
be used with serious precaution.

Conflict of interest statement. The authors hereby declare that there is
no conflict of interest in the study.

REFERENCES

Adeyemo O.K., Akintoye O.A. & Oghi M.L. 2004. Acute toxicity of chlorpyri-
fox (Dursban<sup>®</sup>andrel<sup>®</sup>) to <i>Clarias gariepinus</i>/juve-

Atta A.Y., Abubakar M.B. & Adeifa S.S. 2009. Production of alum from

Ayoola S.O & Ajani E.K. 2007. Histopathological Effects of cypermethrin on
Juvenile Nile Tilapia (<i>Oreochromis niloticus</i>). Afr. J. Livestock Extension

Chugh S.N. 1992. Aluminium phospide poisoning: Present status and ma-

tilapia (<i>Oreochromis niloticus</i>), liver morphology, CYP1A activity and
thyroid hormones after endosulfan dietary exposure. Pestic. Biochem. Phys-
iol. 89:230-236.

Degesch America, Inc. 1988. Material Safety Data Sheet: Aluminum Phos-
phide, Phostoxin, Degesch America, Weyers Cave, VA.

of the African catfish Clarias gariepinus in Sub-Saharan Africa: a hand-

nia and its sublethal effects on selected haematological and enzymatic
parameter of mrigal, Cirrhinus mriga (Hamilton). Aquaculture Res.
35:134-143. (Medline)

on selected haematological parameters in fingerling Catla catla (Hamil-
ton). Aquaculture Res. 35:874-880. (Medline)

Rosemead, California, p.123-140.

A study of the phagocytic system and the fate of intraperitoneally inject-
8:67-78. (Medline)

2009. The effect of crude ethanol extract and fractions of Hyptidendron
canum (Pohl ex Benth.) Harley on the hepatic pancreas of Oreochromis

(Ed.), Aquatic Toxicology. Raven Press, New York.

Jiraungkoorskul W., Upatham E.S., Kruatrachue M., Sahaphong S., Vichasri-
-Gramps & Poketheityook P. 2002. Histopathological effects of roundup,
a glyphosate herbicide, on Nile tilapia (<i>Oreochromis niloticus</i>). Science
Asia 28:121-127.

Jiraungkoorskul W., Upatham E.S., Kruatrachue M., Sahaphong S., Vichasri-
effects of glyphosate herbicides on Nile tilapia (<i>Oreochromis niloticus</i>).
Environ. Toxicol. 18(4):260-268. (Medline)

Heath A.G. 1991. Water Pollution and Fish Physiology. 2nd ed. Lewis Publ.,
Boca Raton. 359p.

Hinton D.E., Baumann P.C., Gardner G.R., Hawkins W.E., Hendricks J.D.,
Murchelano R.A. & Okihoro M.S. 1992. Histopathologic biomarkers,
markers Biochemical, Physiological and Histological Markers of Anthro-
pogenic Stress. Lewis Publ., BocaRaton.

E. 2002. Acute exposure of Siberian sturgeon (<i>Acipenser baerii</i>/,
Brandt) yearlings to nitrite: median-lethal concentration (LC<sub>50</sub>)
determination, haematological changes and nitrite accumulation
in selected tissues. Aquat. Toxicol. 57:257-266. (Medline)

Kori-Siakpere O., Ikomi R.B. & Ogbe M.G. 2011. Biochemical and histo-

Leuschner F. 1984. Evaluation of the acute toxicity of Phostoxin (active in-
redient: aluminum phosphide) to rainbow trout. Laboratory for Phar-
macology and Toxicology, Hamburg, German Federal Republic.

Martinez C.B.R. & Souza M.M. 2002. Acute effects of nitrite on ion regula-
tion in two neotropical fish species. Comp. Biochem. Physiol A 133:151-
160.

Mbaka G.O., Ogbonna S.O. & Awoyemo E.O. 2014. Acute and sub-acute to-
xicity studies of ethanol seed extract of Raphia hookerii on Swiss albino

Mobarak Y.M.S. & Sharaf M.M. 2011. Lead acetate-induced histopatho-
logical changes in the gills and digestive system of silver sailfin molly
(<i>Poe-

Nath N.S., Bhattacharya I., Tuck A.G., Schipulais D.J. & Ebert P.R. 2011. Me-
(Medline)

Nowak B. 1992. Histological changes in gills induced by residues of endo-

efficacy of Eugenia aromatic (Bail.) in the control of Gallosoobruchus ma-
culates (Fabricius) (Coleoptera; Bruchidae) Pest. J. Appl. Sci. Environ.
Manage. 14:97-100.


