Bio-Concentration of Some Heavy Metals and Oxidative Stress Enzymes in *Oreochromis niloticus* (Tilapia fish) from the Hadejia- Nguru wetlands, Jigawa State.

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### Abstract

The Hadejia- Nguru wetland is a vast land of intensive agricultural and fishing activities. It receives waste water through discharges from agricultural, sewage and industrial sources. Five points marked as sites A, B, C, D and E were used for the study. The study aimed in evaluating levels of some heavy metals Cadmium(Cd), Chromium(Cr), Aluminum(Al), Lead(Pb) and Mercury(Hg) in *O. niloticus* and their effect in inducing oxidative stress in the fish. The tissue of interest were gills, muscle and liver. Results obtained revealed concentration of heavy metal in the sequence Hg>Cr>Pb>Al>Cd in the fish. The highest level of heavy metals contamination was recorded in fish from sample site B. There was significant difference (P<0.05) in fish from site B compared with sites C, D and E. Mean concentration of Hg recorded in the gills ranged from 249.75 mg/kg in site B to 128.50 mg/kg in site A, 118.75 mg/kg in site D, 77.25 mg/kg in site C and 62.0 mg/kg in site E. Oxidative stress markers such as superoxide dismutase, catalase, lipid peroxidation and reduced glutathione were significantly higher in the fish tissue indicating a stressful condition in the fish. Anthropogenic activities contributed to the high level of metals found in the wetland and this in turn is reflected in the oxidative stress in fish.

**Keywords:** Cichlid fish, Oxidative stress, Pollutants, Toxicity, Wetlands.

### INTRODUCTION

Fish are largely being used for the assessment of the quality of aquatic environment and as such can serve as bio indicators of environmental pollution (Dautremepuits, 2004). In their natural environment they interact with available amount of pollution either from heavy metals or pesticides. These pollutants when accumulated in the tissues of fish may catalyze reactions that brings about reactive oxygen species (ROS) Farombi et al, 2007. This ROS is formed through two mechanisms which includes the formation of reactive oxygen species through redox cycling brought about as a result of the interactions of Redox active metals while metals without redox potential impair antioxidant defences, especially that of thiol-containing antioxidants and enzymes (Sevcikova et al, 2011, Akinwade et al., 2016) When the levels of ROS is raised it leads to oxidative damage including lipid peroxidation, protein and DNA oxidation and enzyme inactivation. Oxidative stress cannot be avoided in aerobic life, it results from imbalance between the production of ROS and antioxidant defensives in living organism (Nishida, 2011). Elevated production of ROS can be readily induced by pesticides,
transitional metal ions and petroleum pollutants (Lushchak, 2011). Elevated production of ROS causes oxidation of proteins and lipids, change/alterations in gene expression and changes in cell redox redox status (Sevcikova et al., 2011). Superoxide dismutase (SOD), Catalase (CAT), Glutathione reductase (GSH) and Glutathione - s - transferase (GST) are the main antioxidant enzymes and important indicators of oxidative stress. When fish is stressed oxidative stress enzymes are readily released becoming inducible and indicating their readiness to adapt to stressful conditions (Nwani et al., 2015). The Hadejia-Nguru wetland (HNW) is a vastland of fisheries and Agriculture known for its immense productivity, it covers both Jigawa and Yobe states of Nigeria and extends to the Chad basin. Activities which includes mining, excavation and chemical applications have made the wetlands prone to pollutants and toxic substances, studies carried out on the soil and water of the wetlands by Sabo et al.,(2016), Egwu et al.,(2018) revealed some level of heavy metal contamination in soil and water but this study seeks to determine levels of heavy metals in O. niloticus. Therefore, this study is aimed at accessing the level of metallic pollution in the HNW and to evaluate the oxidative stress response in the fish.

MATERIALS AND METHODS

Study Area
The Hadejia – Nguru Wetlands (HNW) lies between latitudes 12° 10N and 13° N and longitudes 10° 15E and 11° 30E. The HNW lies within the semi-arid region of Nigeria. The wetland has an area of about 3,500km². The topography of the area is mostly low lying flat surfaces on the north eastern side and limited local relief in the southern and western parts. Rainfall pattern in the Nguru- Hadejia wetland (NHW) has not been stable over the years, but in most cases starts from June and falls through September. Vegetation is mainly Sudan Savanna, with transitional northern Guinea Savanna and Sahel Savanna in the Southern and Northern limits respectively, Abubakar et al (2015)

Site A is Hadejia barrage dam (Kalgwai)
Site B: Kirikasanma
Site C: Maikintari
Site D: Nguru lake
Site E: Dagona
Fish Sampling and Analysis: *Oreochromis niloticus*, were caught from the five sampling sites through the services of a hired fisherman. Samples were collected bi-monthly during the entire period of study. A total of 139 fish was used for the sampling. The Fish samples were gotten very early in the morning by 6a.m. and transported in ice cold containers to the Department of Fisheries and Aquaculture Federal University Dutse, laboratory for dissection and analysis. While the control fish was gotten from Rumbun kifi fish farm located at
Modobbi road, Dutse, Jigawa state. From enquiries and observation it was gathered that the farm is free from pollution coming from any industry or facility that could affect the biochemical responses of the control fish. The fishes were dissected and the gills, liver and muscles were removed then preserved in a refrigerator at a temperature of -4°C for further experimentation.

Analysis of Water sample
Water samples were collected between 7-8:00am. The physicochemical parameters such as water temperature, turbidity, pH, electrical conductivity, TDS, dissolved oxygen were analyzed according to the method of APHA, (1990). The water temperature was measured with a mercury in bulb thermometer, which was dipped into the water and temperature was read off, the turbidity was measured using a secchi disc by lowering into the water and then measuring the depth at which transparency was seen. The pH, Dissolved oxygen, conductivity and total dissolved solid was measured with a compact D.O. meter Hannan model HI9146, Woonsocket, RI USA.

Heavy Metal Analysis
Analysis was carried by the wet method as used by Tyokumbur, (2016). Dissected muscles, gills and liver were removed and oven dried at a temperature of 105°C until a constant weight was achieved, the dried samples were turned in to powdery form using a porcelain mortar prior to digestion. To digest the samples the powdered muscles, gills and liver were homogenized and subjected to concentrated nitric acid and hydrogen peroxide (1:1)v/v of the powdered sample was placed into a 250ml round bottom flask and 10ml each of HNO₃ (65%) and H₂O₂ (30%) were added and the content of the flask was allowed to undergo reactions. The content of the flask was heated on a heating mantle to a temperature of 130°C dissolution inside a fume hood to reduce the volume to 3ml-4ml, the digested sample was allowed to cool and filtered into a conical flask, the filtered sample was transferred to a 50ml volumetric flask. The concentration of Cd, Al, Cr, Pb and Cd was determined using the Atomic Absorption spectrophotometer (Buck scientific model 230) at the soil science Department of Ahmadu Bello University, Zaria. Same was done for water and sediments.

Measurement of Oxidative Stress Markers
The liver samples was assessed for oxidative stress markers- superoxide dismutase (SOD), catalase(CAT), glutathione(GSH)and the lipid peroxidation (Malondialdehyde MDA). These was determined by adopting the methods of analysis as described by Achuba et al., (2014).

Preparation of extract for the determination of lipid peroxidation (MDA)
Of the isolated gills, liver and muscles, 0.5g were separated and homogenized with 10ml of ice-cold 0.05M phosphate butter pH 7.0 containing 1% (w/v) Triton X-100, excess butylated hydroxyl toluene (BHT) and a few crystals of protease inhibitor, phenylmethylsulfonyl fluoride using an MSE blender immersed in ice. Triton X-100 solubilizes membrane-enclosed organelles while BHT prevents in vitro oxidation of lipid during homogenization. The extract was centrifuged at 7000g for 20 min (40°C). The supernatant (S1) was used for the determination of lipid peroxidation by the method of Hunter et al.,(1963) as modified by Gutteridge and Wilkins (1982).

Extraction and Assay of Catalase (CAT)
Catalase activity was determined according to Beers and Sizer (1952) by measuring the decrease in the H₂O₂ concentration, at an absorbance of 240nm. An extinction coefficient for H₂O₂ of 40M-1 cm-1 (Abel, 1974) was used in the calculation.
Extraction and Assay of Superoxide Dismutase (SOD)
The obtained supernatant was used for the assay of superoxide dismutase (SOD) activity, which was based on its ability to inhibit the oxidation of epinephrine by superoxide anion (Aksnes and Njaa, 1981). The enzyme activities were assayed with an SP 1800 UV/VIS Spectrophotometer.

Sample preparation of Glutathione (GSH)
Tissue sample was prepared by washing with PBS twice, 0.1g of the sample was added into homogenizer, 1mL reagent was added (the proportion of tissue and reagents are kept constant) and this was fully grinded on ice (using liquid nitrogen gave a better grinding effect) centrifuge was done at 8000x g for 10minutes at 4°C the supernatant was placed at 4°C. Spectrophotometer was then preheated for 30minutes and adjustment was made to a wave length of 412nm with distilled water.

Statistical Analysis
Data generated were analyzed using statistical package for social sciences (SPSS) version 25. All the results were expressed as means±SD and the data were analyzed using Analysis of variance (ANOVA). Significant difference between the polluted sites and control were determined at 5% (P < 0.05) confidence level using Duncan multiple test range.

Results
Mean Physiochemical parameter result is presented in Table 1; mean range of water temperature was between 25.37°C and 27.23°C with the highest value seen in site E, there was no significant difference (P<0.05) in recorded temperature in all the sampling sites. Dissolved oxygen values ranged from 5.34mg/l in site A to 6.08mg/l in site B. the values for total dissolved solid showed a slight variation in site E, while sites B, C, A and D showed high values of 405mg/l, 352.42 mg/l, 320.10 mg/l and 308.79 mg/l respectively. Electrical conductivity of water samples gave values of 226.89µS/cm in site C to 400.10 µS/cm in site B, there was significant difference (P>0.05) between site B and the other sites in terms of electrical conductivity. Mean values of turbidity ranged between 25.50NTU in site D and 28.76NTU in site C. pH values were between the range of 7.23-7.48 and there was no significant difference (P< 0.05) among the sites.

Heavy metals
Water sample results presented in figure 2 revealed a concentration value of 0.40mg/l for Al and a highest value in site E. The highest concentration of Cd was recorded in site A with a value of 0.008 mg/l followed by 0.007mg/l in site B. Cr in water sample was highest in site E with a value of 1.004mg/l Pb had a highest value of 0.058 mg/l in site E followed by 0.049mg/l in site D respectively.

Results presented in figure 3 showed the heavy metal deposition in the sediments of the HNW, the variation in concentration showed that site A had a high level of Al as 47.1 mg/Kg, 2.0 mg/Kg of Cd, 45.0 mg/Kg of Cr, 116.25 mg/Kg of Pb and 22.8 mg/Kg of Hg. Site B had the following concentration level of the studied metals, Al was 8.19 mg/Kg, Cd had 1.75 mg/Kg in concentration, Cr had 227.3 mg/Kg, Pb 117.3 mg/Kg and Hg 95.8 mg/Kg. Concentration of metals in site C showed that Al had a concentration of 20.71 mg/Kg, Cd 1.7 mg/Kg, Cr 220.0 mg/Kg, Pb 118.75 mg/Kg and Hg 83.0 mg/Kg. Site D showed the following concentration in metal level in the sediments Al was 17.31 mg/Kg, Cd 3.0 mg/Kg, Cr 150.50 mg/Kg, Pb 132.0 mg/Kg and Hg was 115.8 mg/Kg.
Figure 4, illustrates the heavy metals concentration in the gills, liver and muscles of *O. niloticus* in site A. The concentration of heavy metals in gills of *O. niloticus* showed that mercury was the highest with a mean concentration level of 128.5 mg/Kg. Lead had a mean level of 14.0 mg/Kg while Cd and Al had 3.5 mg/Kg and 2.05 mg/Kg. Liver of *O. niloticus* revealed concentration levels of heavy metals which showed Hg>Cr>Pb>Cd>Al in the order of concentration. The muscles of *O. niloticus* showed results in which Hg>Cr>Pb>Al>Cd. Heavy metals accumulation in gills, liver and muscles of the fish in site A revealed that Hg and Cr were the heavy metals that had high concentration level. The concentration of all the metals followed the order Hg>Cr>Pb>Al>Cd.

Concentration of heavy metals in site B is presented in figure 4. Concentration of metals in *O. niloticus* showed that the gills had a high concentration of Cr 210.0 mg/Kg. Liver samples revealed concentration of 249.75 mg/Kg of Hg, 111.25 mg/Kg of Cr, 6.59 mg/Kg of Al, 3.25 mg/Kg of Cd and 3.0 mg/Kg of Pb. There is significantly difference (P<0.05) in the concentration of Hg in comparison with the other heavy metals present in site B.

Metal concentration in *O. niloticus* from site C is presented in figure 4. Concentration of heavy metals in the gills, liver and muscle in *O. niloticus* was 169.0 mg/Kg of Cr, 77.3 mg/Kg of Hg, 3.75 mg/Kg of Cd, 3.5 mg/Kg of Pb and 2.94 mg/Kg of Al. There was significant difference (P<0.05) in the concentration of Cr with other metals as seen in the gills. Concentration of heavy metals in site D is presented in figure 4. Concentration of the studied heavy metals in *O. niloticus* revealed that the gills had a high concentration of Hg with a value of 118.7mg/kg.

Concentration of the heavy metals in *O. niloticus* from site E is presented in figure 4. Hg in gills, liver and muscles showed a concentration level of 62.0 mg/Kg, 120 mg/Kg, and 128.0 mg/Kg, while that of Cr was 177.5 mg/Kg, 28.5 mg/Kg and 33.75 mg/Kg. Concentration of heavy metals in the control fish is presented in figure 4, which showed a relatively lower concentration in all the sampled heavy metals. The result revealed that site B had the highest level of heavy metal contamination followed by site A, there was significant difference (P<0.005) in contamination level in site B with sites C,D and E.

**Oxidative stress Biomarkers**

Oxidative stress biomarkers were studied in the gills, liver and muscles of *O. niloticus* from the HNW. The oxidative stress biomarkers were superoxide dismutase, glutathione, catalase and lipid peroxidation . Table 2 showed the levels SOD in organs of the sampled fish specie. The result showed that in *O. niloticus* the concentration of superoxide dismutase in the gills were highest in site C, while concentration in the liver was 25.37uml⁻¹ also from site C, the SOD concentration of the muscle was 16.40 uml⁻¹.

Table 3 shows the levels of GSH activities in organs of *O. niloticus*. The GSH levels in the various organs of *O. niloticus* was seen to be significantly high (P<0.05) in the liver, gills and muscle of the fish. Fish samples from site B had the highest GSH levels in the gills while, the liver of site E had the value of 97.13.

The levels of CAT formation in *O. niloticus* is shown in Table 4. There are more elevated levels of CAT in the gills of *O. niloticus*, there was markedly increase as seen in the gills (P<0.05). The gills with the highest level of CAT are from site E, followed by site D with values of 10.17 uml⁻¹ and 6.78 uml⁻¹. The CAT level in the liver samples of *O. niloticus* is seen to be highest in site B with a value of 14.92 uml⁻¹ and is significantly different (P<0.05) in the gills of fish.
The levels of MDA formation in *O. niloticus* is shown in Table 5. Site A had a MDA value of 4.72nmol, while fish from site B had a value of 4.73nmol, site C fish had 7.71nmol, while D was 9.03nmol and E was 5.85nmol. The highest value was seen in site D with a value of 9.03 nmol. The MDA formation in the liver was highest in site D with a value of 8.97 nmol and there was no significant difference (P>0.05) between site A, B and C. Levels of MDA seen in the muscles had a level formation of 9.06nmol, this value is of no significant difference (P>0.05) between sites A, B, C, D and E.

**Figure 2**: Heavy metal concentration in water at the HNW

**Figure 3**: Heavy metal concentration in sediments of HNW
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Figure 4: Heavy metals concentration in organs of *O. niloticus* from the Hadejia-Nguru wetlands

C* = Control

Table 1: Mean physicochemical values obtained from the Hadejia-Nguru wetlands

| s  | A   | B   | C   | D   | E   | Standard limits                                                                 |
|----|-----|-----|-----|-----|-----|---------------------------------------------------------------------------------|
|    |     |     |     |     |     | Water temperature(°C)                                                          |
|    |     |     |     |     |     | 25.37±0.31 a                                                                   |
|    |     |     |     |     |     | 26.12±0.53 a                                                                   |
|    |     |     |     |     |     | 26.90±0.54 a                                                                   |
|    |     |     |     |     |     | 27.20±0.05 a                                                                   |
|    |     |     |     |     |     | 27.32±0.02 a                                                                   |
|    |     |     |     |     |     | <40°C                                                                            |
|    |     |     |     |     |     | D.O.(mg/l)                                                                       |
|    |     |     |     |     |     | 5.34±0.57 a                                                                    |
|    |     |     |     |     |     | 6.08±0.51 a                                                                    |
|    |     |     |     |     |     | 6.02±0.08 a                                                                    |
|    |     |     |     |     |     | 5.50±0.01 a                                                                    |
|    |     |     |     |     |     | 5.55±0.05 a                                                                    |
|    |     |     |     |     |     | 5.0-9.0 mg/l                                                                    |
|    |     |     |     |     |     | TDS (mg/l)                                                                       |
|    |     |     |     |     |     | 320.10±0.3 a                                                                   |
|    |     |     |     |     |     | 405.69±0.14 a                                                                   |
|    |     |     |     |     |     | 352.42±0.33 a                                                                   |
|    |     |     |     |     |     | 308.79±0.49 a                                                                   |
|    |     |     |     |     |     | 296.36±0.17 a                                                                   |
|    |     |     |     |     |     | <600 mg/l                                                                       |
|    |     |     |     |     |     | Electrical conductivity(µS/cm)                                                  |
|    |     |     |     |     |     | 267.41±0.8 b                                                                   |
|    |     |     |     |     |     | 400.10±0.82 a                                                                   |
|    |     |     |     |     |     | 352.23±0.59 a                                                                   |
|    |     |     |     |     |     | 226.89±0.99 a                                                                   |
|    |     |     |     |     |     | 270.40±0.41 b                                                                   |
|    |     |     |     |     |     | <1000 µs/cm                                                                     |
|    |     |     |     |     |     | Turbidity(NTU)                                                                   |
|    |     |     |     |     |     | 28.03±0.48 a                                                                   |
|    |     |     |     |     |     | 28.67±0.39 a                                                                   |
|    |     |     |     |     |     | 28.76±0.33 a                                                                   |
|    |     |     |     |     |     | 25.50±0.49 a                                                                   |
|    |     |     |     |     |     | 24.80±0.52 a                                                                   |
|    |     |     |     |     |     | <25NTU                                                                           |
|    |     |     |     |     |     | P*H                                                                             |
|    |     |     |     |     |     | 7.23±0.02 a                                                                     |
|    |     |     |     |     |     | 7.47±0.39 a                                                                     |
|    |     |     |     |     |     | 7.48±0.40 a                                                                     |
|    |     |     |     |     |     | 7.42±0.01 a                                                                     |
|    |     |     |     |     |     | 7.40±0.00 a                                                                     |
|    |     |     |     |     |     | 6.0-9.0*                                                                         |

*Federal Environmental Protection Agency (1991), **Federal Ministry of Environment (2001), ***World Health Organization (1999)

Table 2: Levels of SOD in organs of *Oreochromis niloticus* from the HNW (unit of measurement is U/ml)

| O. niloticus | SITE | Control | A | B | C | D | E |
|-------------|------|---------|---|---|---|---|---|
| Gills       |      | 1.43±0.30 a | 3.41±0.21 b | 15.11±0.11 c | 18.60±3.20 c | 5.54±2.29 b | 2.01±0.01 b |
| Liver       |      | 0.82±0.02 b | 2.50±0.20 b | 12.70±2.14 c | 25.37±5.25 d | 16.41±2.41 c | 11.87±2.17 c |
| Muscle      |      | 0.35±0.03 a | 3.73±2.70 b | 16.40±4.20 c | 5.85±4.58 b | 0.35±0.20 a | 3.60±0.10 b |

Values are means ± SD of determination for a single fish specie from five (5) points in the HNW- means with different superscript letters in the same row are significantly different at P< 0.005. ND = not detected
Table 3: Levels of GSH in organs of *Oreochromis niloticus* from the HNW (unit of measurement is µg/ml)

| *O. niloticus*       | Control | A        | B        | C        | D        | E        |
|----------------------|---------|----------|----------|----------|----------|----------|
| **Organ/Tissue**     |         |          |          |          |          |          |
| Gills                | 7.30±1.05<sup>a</sup> | 151.06±0.99<sup>c</sup> | 388.08±2.07<sup>b</sup> | 66.94±6.12<sup>b</sup> | 104.32±4.21<sup>c</sup> | 121.12±1.04<sup>d</sup> |
| Liver                | 0.81±1.01<sup>a</sup> | ND       | 18.21±8.11<sup>b</sup> | 29.48±0.24<sup>c</sup> | 63.80±3.10<sup>d</sup> | 97.13±7.12<sup>d</sup> |
| Muscle               | 1.61±1.50<sup>b</sup> | 104.32±4.22<sup>b</sup> | 253.15±3.05<sup>d</sup> | 213.44±13<sup>c</sup> | 153.14±4.04<sup>b</sup> | 322.87±21.7<sup>e</sup> |

Values are means ± SD of determination for a single fish specie from five (5) points in the HNW- means with different superscript letters in the same row are significantly different at P< 0.005. ND = not detected

Table 4: Levels of Catalase in organs of *Oreochromis niloticus* from the HNW (unit of measurement is U/ml)

| *O. niloticus*       | Control | A        | B        | C        | D        | E        |
|----------------------|---------|----------|----------|----------|----------|----------|
| **Organ/Tissue**     |         |          |          |          |          |          |
| Gills                | ND      | 0.68±0.00<sup>a</sup> | ND       | 3.39±1.00<sup>b</sup> | 6.78±0.81<sup>c</sup> | 10.17±0.20<sup>d</sup> |
| Liver                | ND      | 0.86±0.38<sup>a</sup> | 14.92±2.81<sup>b</sup> | ND       | 0.65±0.04<sup>a</sup> | 0.68±0.01<sup>a</sup> |
| Muscle               | 0.68±0.01<sup>a</sup> | 4.75±1.23<sup>b</sup> | 0.68±0.01<sup>a</sup> | 2.03±0.05<sup>b</sup> | ND       | 0.68±0.01<sup>a</sup> |

Values are means ± SD of determination for a single fish specie from five (5) points in the HNW- means with different superscript letters in the same row are significantly different at P< 0.005. ND = not detected

Table 5: Levels of Malondialdehyde in organs of *Oreochromis niloticus* from the HNW (unit of measurement is n/mol)

| *O. niloticus*       | Control | A        | B        | C        | D        | E        |
|----------------------|---------|----------|----------|----------|----------|----------|
| **Organ/Tissue**     |         |          |          |          |          |          |
| Gills                | 0.94±0.72<sup>a</sup> | 4.72±1.60<sup>b</sup> | 4.73±0.13<sup>b</sup> | 7.71±0.58<sup>c</sup> | 9.03±0.03<sup>c</sup> | 5.85±0.05<sup>a</sup> |
| Liver                | 0.61±0.37<sup>a</sup> | 6.30±0.30<sup>c</sup> | 6.71±2.10<sup>c</sup> | 6.31±0.19<sup>c</sup> | 8.97±2.12<sup>c</sup> | 3.59±0.01<sup>b</sup> |
| Muscle               | ND      | 1.06±0.03<sup>a</sup> | 1.68±0.43<sup>a</sup> | 2.00±0.50<sup>a</sup> | 2.33±1.00<sup>a</sup> | 2.04±0.00<sup>a</sup> |

Values are means ± SD of determination for a single fish specie from five (5) points in the HNW- means with different superscript letters in the same row are significantly different at P< 0.005. ND = not detected
DISCUSSION

Results revealed that the wetland is contaminated with the studied heavy metals in higher proportion exceeding the maximum recommended limits in aquatic foods, concentration of heavy metals in the body of fish was higher than the concentration in the water body, this is as a result of the ability of fish to bio accumulate metals above the concentration in water. Nsofor et al., (2014). This agrees with the work of Adaka et al., (2017) stating that heavy metals in the body organs of Citharus citharus, T. zilli and Heterotis niloticus from Oguta lake gave a higher concentration than water concentration. Heavy metal contamination in site A resulted to a massive accumulation in fish tissue, The high level of Hg and Cr can be attributed to anthropogenic sources, even though unpolluted water contains trace amounts of Hg which do not exceed 0.1µg/l, the main source of Hg in environment is the fungicides, especially in the organic compounds of mercury. The amount of Hg in the muscles of O. niloticus was higher than the other organs put together, this agrees with the findings of Golovanova, (2006) that visceral distribution of Hg in organs and tissues often shows the following order Muscles>liver > intestine > spleen> brain>gonads this is due to the high content in muscles of functional proteins (-SH, -NH₂, -COOH, _OH) having high affinity to Hg. The presence of Hg even at low concentrations reduces the viability of spermatozoa in fish, reduces production of eggs and affects the survival rate of developing eggs and fry. Concentration of Cr was high in sample site A beyond the recommended acceptable limit in food. The Cr gains entrance in to the aquatic ecosystem through effluents discharged from mining, dyeing, leather tanneries, textiles. The Cr level observed in the study showed a higher concentration and Fish takes in Cr by ingestion or by uptake through the gills, and accumulates morphological alterations. Pb has been classified as one of the most toxic metals, and a high mean concentration was found in the muscles of O. niloticus Pb is naturally available substance with its concentration increased by anthropogenic sources which could be base metal mining, Pb based paints and gasoline. Lead in water may come from industrial and smelter discharges, Pb containing pesticides, street runoff and municipal waste water. The level of Pb observed in the HNW was above the recommended limit as outlined by WHO, (1999), FAO, (2004) which falls between 0.3 mg/Kg and 0.01 mg/Kg in food. When there is exposure to Pb at high concentrations abnormalities such as lead poisoning occurs and manifestations such as hypertension, renal dysfunction, fatigue, sleeplessness, convulsions, abdominal pain and loss of appetite, headache, numbness, arthritis and hallucination occurs. The concentration of Pb observed in this study is similar to the result obtained by Farombi et al., (2007) in C. gariepinus from from the Ogun river where they obtained high values of Pb in the liver and kidney of C. gariepinus. The Cd concentration observed in this study is consistent and of higher value than that obtained by Farombi et al.(2007) in similar study where values of metal concentration was high in organs of C. gariepinus. The mean concentration of metals in all organs was above the permissible levels of food consumption as recommended by the acceptable limit for heavy metals. Cd is a naturally occurring non- essential trace element and its tendency to bio-accumulate in living organisms often in hazadous levels raises environmental concerns, Authman et al., (2015).The use of Cd containing fertilizers, agricultural chemicals, pesticides and sewage sludge in farm land could be the possible source of Cd in the studied water bodies. The concentration level of Cd in the study is similar to the concentration observed in fish as described by Hashim et al., (2014) from studies conducted at the lower reach of the Kelantnan river, Malaysia where the concentrations of Cd was found to be higher above the critical level of the WHO and FAO. The heavy metals concentrations in all the sampling sites showed a higher level in all fish samples even higher than the permissible limit as obtained in the eatable food substances, the level observed showed that they have reached the level of concern. Waakes, (2000) asserted that heavy metals such as Cd
affects the kidneys and causes symptoms of chronic toxicity, such as impairment of kidney function, poor reproductive capacity, hypertension, tumors and hepatic dysfunction.

The activities of SOD, the redox sensitive thiol compound GSH, CAT and MDA were elevated in all the organs (gills, liver and muscle). The significant increase recorded in these organs would be as a result of the presence of heavy metals in the water; these accumulation of heavy metal might have triggered the production of superoxide anions resulting to the induction of SOD to convert the superoxide radical to \( \text{H}_2\text{O}_2 \). SOD catalytically scavenges superoxide radical which appears to be an important agent of oxygen toxicity. GSH showed an elevated level in all the samples and GSH is well understood to be a substrate for the activity of GST. The increase recorded in GSH formation in a high level suggests an adaptation and protective mechanism by this biomolecule against oxidative stress induced by heavy metal and pesticide residues which agrees with the findings of Farombi et al. (2007). The high level of GSH recorded in the gills could result from the gills being more exposed to contaminated water and as such pollutant (heavy metals) can have access through the fish thin epithelia, this can be attributed to the reasons why there was high level of GSH in the fish. Catalase activity was decreased in the fish specie studied although increase in the activity of CAT and SOD is usually observed in the presence of environmental pollutants as documented by Dautremepitis et al. (2004). The reduced rate of CAT in this study can be attributed to the overshadowing ability of superoxide radicals as asserted by Stanic et al., (2005). Significant elevation of lipid peroxidation in all organs indicates the accumulation of heavy metals in the organs as seen in the data obtained in this results, this elevated concentrations can raise high levels of antioxidants and in some cases cause damages in DNA, proteins and lipids, Pandy et al., (2003). The presence of antioxidant in fish establishes oxidative stress conditions in fish as a result of the high concentration of heavy metals analyzed in this study.

Chromium is known to be carcinogenic in humans (WHO, 1988). Ahmad et al., (2006) described the genotoxicity of chromium in the gills and kidneys of Anguilla Anguilla; while Farag et al., (2006) described DNA damage and high elevation of lipid peroxidation in Chinook salmon (Oncorhynchus tshawytscha) during chronic exposure to chromium in water; it is on record that cadmium does not generate reactive oxygen species directly but it can alter GSH levels and influence cell thiol status inducing the expression of metallothioneins in the liver, Sevcikova et al., (2011). Changes in GSH can lead to lipid peroxidation of cell membranes. In this study the increased level of SOD could result from the high level of mercury present, this agrees with the findings of Monteiro et al., (2010) who observed high levels of mercury in liver, gills and heart of fresh water Brycon amazonicus. Other metals such as Aluminium detected is a widespread pollutant capable of inducing oxidative stress in fish. This study showed that fish is a strong bioindicator of metals in the environment and can be used as a measure of the pollution level in an environment.

CONCLUSION
The high level of heavy metals revealed in this study is a reflection of the state of pollution in the HNW. Since aquatic life depends largely on this water for their survival and wellbeing, it is necessary the water bodies in the wetland be monitored by the government in order to regulate and educate people on safety measures when applying some chemicals and to manage waste that makes the presence of heavy metals possible in these water bodies.

REFERENCES

Musa I.M., Imam T.S., DUJOPAS 7 (4b): 168-180, 2021
Bio-Concentration of Some Heavy Metals and Oxidative Stress Enzymes in Oreochromis niloticus (Tilapia fish) from the Hadejia- Nguru wetlands, Jigawa State.

Achuba, F.I., Ebokaiwe, P., Peretiemo-Clarke. B.O. (2014). Effect of environmental pollution on oxidative stress in African Catfish (Clarias hetrobranchus) International Journal of Environmental Monitoring and Analysis. 2(6):297-301.

Aebi, H. (1974). Catalase in method of enzymatic analysis (Brygmeyer, H.U(ED) Academic press, New York Pp. 673-684

Akinwande, A.A., Abdulkadir, J.O. and Adesina, B.T. (2016). Oxidative stress and Antioxidant Response in the Giant African Catfish (Heterobranchus Bidorsalis) under Chronic Paraquat Exposure. Nigerian Journal of Fisheries and Aquaculture, 4(2):30-37.

Alksnes, A., and Njaa, R.L. (1981). Catalase, glutathione peroxidase and superoxide dismutase in different fish species. Comp. Biochem, Physiol. 69B: 893-896.

Arimoro, F.O., Chukwuji, M.A.J and Oghenegholome, O. (2000). Effect of industrial waste water on the physical and chemical characteristics of a tropical coastal river. Res. J. Environ. Sci. 2(3):209-220

Beers, R.F. and Sizer, I.W. (1995). A Spectrophotometric method for measuring the breakdown of hydrogen peroxide by catalas. J boil. Chem. 195; 133-144

Cohen, G, Dembrec, D. and Marcus, J. (1970). Measurement of catalase activity in tissue extracts. Analyt. Biochem: 34:30-38

Crapo, J.D., Mc Cord, J.M. and Fridovich, I. (1978). Preparation and assay of superoxide dismutases. Meth. Enzyme 53:328-393.

Dautremepuits, C., Paris-Palacious, S., Betoule, S., Vernet, G.(2004). Modulation in hepatic and head kidney parameters of crap (Cryptinus carpio L.) Induced by copper and chitosan, Comp. Biochem physical. C. Toxicol pharmacol. 137, 325-333.

Egwu, G.N., Okunola, O.J. and Ugwuike, K.C. (2018). Evaluation of some Heavy Metals in Wetland Soils of Uguru Yobe State, Nigeria. Journal of Applied Science Environ .22(6):987-992

Farag, A.M., May, T., Marty, G.D. Easton, M., Harper, D.D., Little, E.E. and Cleveland, L. (2006). Effect of chronic chromium exposure on the health of chronic salmon Oncorhynchtus tsharytyscha, Aquatic Toxicolgy 76, 246-257.

Farombi, E.O., Adelowo, O.A. and Ajimoko, Y.R (2007). Biomarkers of oxidative stress and heavy metal levels as indicators of environmental pollution in African catfish larias gariepinus from Nigeria, Ogun river. Int.J.Envirnm. Res. Public Health, 4(2):158-165.

Federal Environmental Protection Agency FEPA. (1991). Natural Environmental protection Effluent Limitation Regulations of 1991 Federal Environmental Protection Agency, Lagos, Nigeria. Ref. S. 1-8

Gutteridge, J.M.C and Wilkins, S. (1982). Copper dependant hydroxyl radical charge to ascorbic acid Fed. European society letters. 137:32

Hunter, F.E; Gebrecki, J.M; Hoffstein, P.E; Weinstein, J. and Scott, A. (1963). Swelling and Lysis of rat liver mitochondria induced by Ferrousion J. Biol. Chem. 238, 847-851

Kaur, M and Jindal, R. (2017). Oxidative stress response in liver, kidney and gills of Ctenopharyngodon idellus (Cuvier and Valenciennes) exposed to chlorpyrifos. Biol med. 2017:1 (4): 103-112.

Kefas, M.; Abubakar, K.A and Ali, J. (2015). The assessment of water Quality via physicochemical parameters and macro invertebrates in Lake Geriyo, yola, Adamawa state, Nigeria. The international Journal of Science and Technology, 3(3). 284-290

Misra, H.P. and Fridorch, I., (1972). The role of superoxide ion in the auto-oxidation of epinephrine and a simple assay for superoxide dismutase, J. Bio. Chem. 247:3170-3175.
Bio-Concentration of Some Heavy Metals and Oxidative Stress Enzymes in Oreochromis niloticus (Tilapia fish) from the Hadejia- Nguru wetlands, Jigawa State.

Mwevura, H; Othman, C; Mhehe, L. (2004). Organochlorine Pesticide Residues in Edible Biota from the coastal area of dar es salaam city, *western Indian Ocean J. mar. Sci* 1:91-96.

Nishida, Y. (2011). The chemical process of oxidative stress by copper (II) and iron (III) ions in several neurodegenerative disorders. *Moatshefte for chemie* 142, 375-383.

Nwani, C.D., Ekwueme, H.I., Ejere, V.C., Onyeke, C.C., Chukwuka, C.O., Peace, O.N. and Nwadingwe, A.O. (2015). Physiological effects of paraquat in Juvenile African catfish Clarias grypinus (Burchell, 1822). *Journal of Coastal Life Medicine*, 3(1):35-43.

Ochuwa, O.G., Nnamdi, H.A., Emanuel, B., Adebayo, A.O. (2017). Genitoxic, Histopathological and Oxidative Stress Responses in Catfish, *Clarias grypinus*, Exposed to two Antifouling points. *Journal of health and Pollution, Vol. 7*, No. 16 Dec; 2017.

Pandey, S; Parvez, S; sayeed, I; Haque, R; Bin-hafeez, B; Raisuddin, S.(2003). Biomarkers of Oxidative stress: A comparative study of river Yamuna Fish Wallago attu (BI & schnn), *Sci Total Environ.*, 309 105-15.

Sabo, B.B., Ringim, A.S. and Karaye, A.K. (2016). Evaluation of irrigation water trapped by Typha daimenesis for heavy metals in Hadejia river, Nigeria. International Journal of Public and Environmental Health. 3(5)107-111

Sevcikova, M., Modra, H., Slaninova, A., Svobodova, Z. (2011). Metals as a cause of oxidative stress in fish: a review. *Veterinarni medicina*, 56(11):537-546

Tyokumbur, E.T. (2016). Bioaccumulation of heavy metals in organs and tissues of *Xenopus laevis* and sediment concentrations from Alaro Stream in Ibadan. *New York Science journal* 9(3): 39

Waakis, M.P. (2000). Cadmium carcinogenesis in review. *Journal of inorganic Biochemistry*. 79(1) 241-244.

WHO (1988). Environmental health criteria, Chromium, WHO, Genera.