Growth Response and Ionic Regulation in Common Carp (Cyprinus Carpio L.) After Chronic Dietary Copper Exposure and Recovery

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Abstract

Effects of exposure of common carp juveniles (Cyprinus carpio L.) to dietary copper and its recovery rate were investigated with the aim of determining the growth and physiological impact. The fish were fed elevated copper diets (1000 mgCu/kg and 2000 mgCu/kg as diet 2 and diet 3 respectively) and control diet (5 mgCu/kg, as diet 1) for 42 days and were then fed the control diet for a further 21 days. After 42 days of exposure to elevated copper diets, growth performance examination showed that there was significant increase at (p<0.05) in feed intake, %body weight, weight gain and condition factor by fish fed diet 2 compared to diet 1 and diet 3. There was, however, no difference in specific growth rate, feed conversion ratio, in all treatments (p>0.05). Hepatosomatic index increased significantly in fish fed both elevated diets compared to control diet (p<0.05). Recovery period on normal diet (control) showed no significant effect of copper recovery on feed intake, weight gain, %body weight, specific growth rate and feed conversion ratio in all treatments (p>0.05), but, fish fed diet 2 showed a significant reduction in condition factor compared to other diets (p<0.05). Tissue Na+, Ca++, K+ were disturbed throughout the experiment with sodium increasing from 257.82 ± 2.50 µmol/g to 388.14 ± 1.32µmol/g and calcium increasing from 499.54 ± 6.81 µmol/g to 1025.94 ± 9.16 µmol/g reducing gill copper from 11.63 ± 0.37 mgCu/g to 0.00 ± 0.00mgCu/g. Intestinal copper decreased from 14.93 ± 0.1 mgCu/g to 0.00 ± 0.00 mgCu/g as a result of sodium increasing from 130.30 ± 5.12 µmol/g to 438.72 ± 2.44 µmol/g. The reduction in intestinal calcium was sodium dependent as increasing sodium decreased calcium absorption. Increased gill copper of the 1000 mgCu/kg diet exposed fish during exposure compared to the control was due to copper induced decrease in plasma ion regulatory sodium (Na ATPase activity), which protected fish from direct toxicity effect and could also suggest another pathway other than the common Na/Cu apical channel shared between sodium and copper through which copper binds to fish gill; diet 2 fish showing significant increase at (p<0.05) in haematocrit, red blood cell, white blood cell and neutrophils, and a significant reduction in lymphocyte and mean cell hemoglobin compared to diet 1 and diet 3, (p<0.05). This increase in blood indices is indicative of stress onset to which fish fed diet 2 is subjected. Fish fed diet 3 showed significant reduction in haematocrit, red blood cell, white blood cell and increased lymphocyte (p<0.05) and became anaemic with severe skin discoloration, indicative of a worsening effect of excess dietary copper exposure on the fish. There were not significant differences in moisture content of all tissues during and after copper exposure (p>0.05), although, gills of fish fed diet 2 showed reduction in moisture compared to diet 1 and diet 3-fed fish for both exposure and recovery phases, increasing from 75.3 ± 3.20% to 79.5 ± 6.44% after recovery for 21 days. Gills of fish fed diet 3 also increased post-exposure, indicative of protection of the structural integrity of the gill to prevent hypoxia through oxygen supply from water.

Keywords: Growth; Ionic chronic; Dietary copper Regulation; Exposure; Recovery; Common carp

Introduction

Copper is an essential trace element that plays a vital role in the physiology of animals for fetal growth and early post-natal development, for hemoglobin synthesis, connective tissue maturation especially in the cardiovascular system and in bones for proper nerve function and bone development, and inflammatory process. It is involved in different biochemical process of animal metabolism such as: enzyme – co enzyme catalytic reactions. It is associated with the function of a number of enzymes such as oxygenases including cytochrome C oxidase and copper-zinc super oxide dismutase [1,2] and; ion transport for instance with ceruloplasmin (ferroxidase1), a putative copper transport protein required for the incorporation of iron into transferring for it is transport in plasma [3].

It is an essential micronutrient for vertebrate animals especially fish, and has numerous functions, in addition to the ones stated above, in cellular biochemistry including vital roles in cellular respiration, and a cofactor for over 30 different enzymes [3]. Copper deficiency leads to physiological disturbance. Symptoms include depression of growth, anaemia, bowing of legs, spontaneous fractures, ataxia of new borns, Cardiac and vascular disorders and a depigmentation, decrease in some organs weight, depressed reproductive performance including egg production.Copper, though essential in fish diet, can be harmful when large single or daily intake occurs.

The dietary effect of copper varies from species to species [4-6] and has never been reported for most temperate fish such as Salmon [4] and recently, Nile tilapia [7], but little or no information on the dietary copper exposure and recovery in common carp has been reported. In Salmon, toxic effect of dietary copper includes reduced growth [8] severe lesions in the gut at high concentration (10 gKg-1 food) cell proliferation and metallothionein [1]. Fatty change in the liver of salmon as well as altered haematology has been severally reported that...
copper when in excess, is stored in the liver and tend to increase after feed withdrawal leading to hepatic cell lysis and release of cell content in the liver. This is confirmed by [7] Shaw and Handy. In their report in Nile tilapia, the recovery phase on normal diet without copper was characterized by a reduction in intestinal and branchial copper level after dietary copper exposure was confirmed by elevated copper concentration in the intestine, liver and gills.

Knox reported similar fatty changes in the liver of salmon, but there was altered hematology, contrary to report of [7] Shaw and Handy. Research on copper exposure and recovery in common carp would go a long way to add to sparse literature on dietary copper exposure and recovery in tropical fresh water and to know if difference in climate and region could account for differences in toxicity of dietary copper exposure at same level.

Dietary copper exposure in African catfish has also been reported severally, aqueous copper exposure and recovery on common carp [9]; aqueous cadmium exposure in common carp; aqueous zinc on common carp [10]; dietary copper exposure and recovery in Nile tilapia [7]; as well as threshold for excess dietary copper toxicity on fresh water fish excluding carp [5,8] have all been reported. But to my knowledge and literature review, no study on the dietary copper exposure and recovery in common carp (L) has been reported. This present study will not only examine chronic dietary copper toxicity on common carp, but it will also establish threshold for dietary copper toxicity by investigating growth and ionic response, which has not been reported for the fish.

The common carp, a benthic omnivore, is native to Asia and Eastern Europe [11]. Reputed as a popular food fish and a highly cultivable species with year round breeding under tropical and subtropical conditions, the common carp also plays an important role in polyculture systems in seasonal reservoirs and ponds [12]. It is the only exotic carp species that is known to breed naturally in lake. It has high fecundity and hatchability [13].

It has been introduced into environments worldwide and can grow to a maximum length of 5 feet (1.5m), a maximum weight of over 80lb (37.3kg), and an oldest record age of at least 65 years [14]. This age longevity of common carp makes it good for chronic toxicity test. Similarly, although they are very tolerant of most condition, the common carp prefer large bodies of show or standing water and soft, vegetable sediments. This makes it not unaffected by pollution from heavy metals since they eat anything near bank and bottom, thus ingesting contaminated food and water during feeding [15].

Fish unlike most terrestrial animal, can absorb some minerals, (inorganic elements) not only from their diet, but also from their external aquatic environment [16]. Calcium (ca), sodium (Na), potassium (k), copper (Cu) and other essential minerals are generally derived from the water to satisfy part of the nutritional requirement of fish [17]. Inorganic elements, which are required for the normal life processes of fish, perform the following function: formation of skeletal muscles structures, electron transfer, regulation of acid-base balance/ equilibrium and osmoregulation; they are important component of hormones and enzymes and activate enzymes. Complex biochemical mechanisms control and regulate the uptake, storage, and excretion of various inorganic elements, allowing fish to live in a dynamic equilibrium with their aquatic medium. The electrolytes Na⁺, Mg²⁺, Ca⁴⁺, Cl⁻ and HCO₃⁻ play a major role in the osmotic and ionic regulation of extra –and intra cellular fluids in fish [16].

The exchange of ions from the surrounding water across the gill and skin of fish complicates the measurement of mineral requirement; although most essential elements known for terrestrial animals are also considered important for fish, quantitative requirement have been reported for only nine(9) minerals: calcium, phosphorus, magnesium, iron, copper, manganese, zinc, selenium, and iodine [18-21] for selected fish species.

**Materials and Methods**

**Experimental design**

Common carp were purchased from Oyo state Agricultural Development Programme (ADP) reputable fish farm in Ibadan, Oyo State. They were then placed under laboratory conditions in fish holding tanks with water temperature 27.4 ± 0.42°C and left unfeed in the first 2 days to adapt to a change in environment before feeding them with normal diet.117 fish of average weight 19.43 ± 14.09 g were then placed in 9 plastics of 52 liters each in a water renewal method and were fed a control diet with no added copper to saturation for 14 days in order to acclimatize them to experimental conditions with 13 fish per tank. While fish in the first three containers remained on the control diet, fish in the second and last three containers were fed copper-loaded diets (1000 mg/kg dry weight feed) and (2,000 mg copper/kg dry weight feed) respectively for 42 days. This was then was then followed by a 21 –day recovery period with all containers fed the control diet (no added copper diet). Throughout the experiment fish were fed to satiation twice a day in the morning and evening. Care was taken to ensure no uneaten food remained in the tanks during feeding and copper did not leach from the feed. To achieve these objectives, water was constantly and completely changed daily with fresh well water added and uneaten food removed after satiation was noted. Daily feed intake was calculated by subtracting weight of feed plus container after feeding from feed plus container before feeding. Copper concentrations in the different tank were measured in the analysis of water quality. Growth and nutritional performance in the different treatments were monitored throughout the experiment and the fish randomly sampled from each tank after 42 –day of copper exposure for hematological, tissue ion analysis, and histology. Fish were not fed the day before sampling times in order to empty the gut and to facilitate dissection.

**The Diet**

The control diet was purchased from a commercial animal feed dealer (Adom commercial Feeds, Ibadan, Oyo state, Nigeria) with a proximate composition from manufacturer’s guidelines shown below:

**Dietary Preparations**

The copper-supplemented diet was formulated by starch coating of the commercial feed with copper sulphate. In order to achieve a nominal copper concentration of 1000 mgCu/kg feed 1.1722 g of CuSO₄·5H₂O (Anala R grade, BDH, poole, UK) was dissolved in 35ml of deionised water with 1.2 g of starch to bind the copper to the food sticks. The starch coat dried within minutes, and the copper diet was stored in airtight containers at – 20°C to prevent lipid oxidation. The other copper supplemented diet (2000 mgCu/kg) was also treated, but required higher CuSO₄·5H₂O. In order to achieve a nominal copper concentration of 2000 mgCu/kg feed, 2.3503 g of CuSO₄·5H₂O was also dissolved in 35 ml of deionised water with 1.2 g of starch to bind the copper to the feed. It was then mixed thoroughly; the starch solution was gradually sprayed onto 300 g of the commercial feed with copper sulphate. In order to achieve these objectives, water was constantly and completely changed daily with fresh well water added and uneaten food removed after satiation was noted. Daily feed intake was calculated by subtracting weight of feed plus container after feeding from feed plus container before feeding. Copper concentrations in the different tank were measured in the analysis of water quality. Growth and nutritional performance in the different treatments were monitored throughout the experiment and the fish randomly sampled from each tank after 42 –day of copper exposure for hematological, tissue ion analysis, and histology. Fish were not fed the day before sampling times in order to empty the gut and to facilitate dissection.
by Atomic Absorption Spectrophotometry (model 210 VGP) with the following specification for copper detection and analysis:

**Growth and nutritional performance**

**Growth and nutritional performance was measured and described below:**

Briefly, feed intake was calculated, daily for each tank by weighing feed containers before and after feeding. All fish were individually weighed at the start of the experiment and the end of 42 days of exposure. The individual fish weight was used because the periodic sacrifice of fish during the experiment prevented nutritional parameters being calculated from cumulative tank biomass as follows:

- Specific growth rate SGR (% day\(^{-1}\)) = \(\frac{\log W_2 - \log W_1}{(t_2 - t_1) \times 100}\)
- \(t_1\) = initial time point before exposure (days)
- \(t_2\) = final time point after exposure (days)
- \(W_1\) = fish weights at \(t_1\)
- \(W_2\) = fish weights at \(t_2\)

**Feed Conversion Ratio (FCR) =** \(\frac{\text{feed intake (g)}}{\text{weight gain (g)}}\)

This was calculated from mean gain in body weight for each treatment for:

i) the copper exposure phase (days 0 –42),
ii) recovery phase (days 43 –63)

- Mean weight gain(g)=final weight—initial - weight
- Condition factor(%)= weight(g)/length\(^2\) (cm) \(\times 100\)
- Hepato-somatic index for each fish (%) = liver weight (g)/body weight (g) \(\times 100\).

**Tissue ion and Moisture analysis**

Tissues for trace metal analysis were oven dried to a constant weight, which were subtracted from the initial weight of each tissue before drying to get the moisture content and then expressed as percentage of initial weight; oven dried samples were then digested in nitric acid and then diluted to volume with distilled water. The samples were analyzed by Atomic Absorption Spectrophotometry for copper (at 589.00 nm), K\(^+\) (at 766.5 nm), Na\(^+\) (at 589.00 nm), and Ca\(^2+\) (at 422.7 nm) was using flame photometer (model: corning 410).

**Results and Discussion**

The physiological effects of dietary copper exposure and recovery on normal diet were studies throughout the entire length of the project. Growth performance, histology, hematometry as well as tissue ion and moisture were investigated during the two phase of the experiment (exposure and recovery). Water quality was monitored throughout the exposure phase and result showed that all parameters were within the range required and tolerated by common carp.

This study is a first report of chronic dietary copper exposure/toxicity in common carp; and overall, fish, in this research, accumulated excess copper in the liver and intestine. The results of the experiments are discussions are discussed below with tables showing the effects of dietary copper on each parameter.

### Copper accumulation

Copper accumulations in fish tissue have severally been reported [5,7,22,23]. In this study, copper accumulation in common carp also reflected the route of exposure [24], with large increase in copper content of the liver and intestine [1,23,25] and is consistent with previous studies on temperate species such as rainbow trout [26], Atlantic Salmon [25]. The gills showed increased copper accumulation post exposure in the 1000mgCu/kg (diet 1) and 2000mgCu/kg (diet 2). This cannot be explained by aequous copper uptake, because gill morphology was normal (NVL) during and post exposure period [7]; rather, it was due to increased intestinal absorption of dietary copper.

Copper accumulation in fish does not depend on dietary level [25], but sodium dependent [27-29] as well as being calcium dependent [30]. In this research the time dependent reduction in the control was sodium and calcium dependent [27,30] with sodium increasing from 257.82 ± 2.50\(\mu\)mol/g to 388.14 ± 1.32 \(\mu\)mol/g and calcium increasing from 499.54 ± 6.81 \(\mu\)mol\(^{-1}\) to 1025.94 ± 9.16 \(\mu\)mol\(^{-1}\) reducing gill copper from 11.63 ± 0.37 mgCukg\(^{-1}\) to 0.00 ± 0.00 mgCukg\(^{-1}\). Intestinal copper decreased from 14.93 ± 0.1 mgCukg\(^{-1}\) to 0.00 ± 0.00 mgCukg\(^{-1}\) as a result of sodium increasing from 130.30 ± 5.12 \(\mu\)mol\(^{-1}\) to 438.72 ± 2.44 \(\mu\)mol\(^{-1}\). The reduction in intestinal calcium was sodium dependent as increasing sodium decreased calcium absorption [31]. Increased gill copper of the 1000 mgCukg\(^{-1}\) diet exposed fish during exposure compared to the control was due to copper induced decrease in plasma ion regulatory sodium (Na ATPase activity), which protected fish from direct toxicity effect [32] and could also suggest another pathway other than the common Na/Cu apical channel shared between sodium and copper through which copper binds to fish gill [28]. The later reason could be apt due to the fact that, although, inhibition of copper branchial /basolateral Na+/K+ ATPase cannot extrude intracellular Na+ into the blood (that is influx is inhibited), branchial influx of sodium was stimulated into the blood due to decreased plasma sodium from low intestinal uptake [33]. The copper accumulated in the gill of the 1000 mgCukg\(^{-1}\) exposed fish could, therefore, show that a high affinity mechanism for bronchial copper uptake in the gills was independent of external sodium [34]. In the same vein, reduced gill copper of 2000 mg Cu/kg (diet 3) exposed fish was due to the fact that since sodium and copper shared similarly apical channel, increasing sodium and copper absorption from the intestine elevated plasma concentration of these two mineral elements beyond the needs of the fish; bronchial influx of sodium and copper were inhibited [33].

The increased gill copper of the 1000 mg Cu/kg diet (diet 2) compared to the control (diet 1) and the 2000 mg Cu/kg (diet 3) is an indication of the beginning of stress to which fish is subjected and this response is a result of the accumulation of dietary copper in the gill from direct toxicity effect [32] and could also suggest another pathway other than the common Na/Cu apical channel shared between sodium and copper through which copper binds to fish gill [28]. The later reason could be apt due to the fact that, although, inhibition of copper branchial /basolateral Na+/K+ ATPase cannot extrude intracellular Na+ into the blood (that is influx is inhibited), branchial influx of sodium was stimulated into the blood due to decreased plasma sodium from low intestinal uptake [33].

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in post exposure in all tissues indicated possible delayed toxic effects of dietary copper. This has also been reported for Nile tilapia [7].

**Tissue Sodium, Potassium and Calcium**

The presence of treatment-dependent changes in tissue ions (Na+, K, Ca) suggest that dietary copper caused major osmotic disturbances which stimulated blood production (red blood cell, haematoцит) of fish exposed to diet 2 (1000 mgCu/kg). The time dependent increase in sodium and potassium indicated the ability of common carp to acclimatize to long time dietary copper exposure [24,35] and could also reflect the excretion and detoxification capacity of common carp to copper [35]. The time dependent reduction in gill calcium is sodium dependent [31] which explained the further reduction during elevated dietary exposure to 1000 mg Cu/kg (diet 2) compared to the control as the gill sodium increased. The high level of gill calcium compared to other tissues in all treatments and the low level of intestinal calcium is due to the simple fact that the gill is the major and primary site of calcium absorption even though dietary calcium inclusion exceeded requirement (1-8% vs. 0.34%) and uptake, while the intestinal contribution to calcium uptake comes to 30%. Similarly the presence of hypercalcaemic hormone, cortisol in the gill. Hanssen et al. [36], has been reported to stimulate hypercalcaemia (high calcium), which is calcitropic [37,38] making fish capable of surviving extreme hypercalcaemia (up to 10 mmol-1 total calcium, 4.5 mmol-1 Ca²⁺). Intestinal absorption of Calcium is inhibited by stanniocalcin (calcium reducing hormone) which reduces intestinal calcium absorption. The calcitropic action of cortisol (a stress hormone, [9]), independent of stanniocalcin [39] become noticeable only in the long term [33]. This could explain hypocalcaemia observed during the chronic dietary exposure of common carp to copper, which showed their stress response to elevated dietary copper (1000 mgCu/kg and 2000 mgCu/kg). The post exposure phase was characterized by significant decrease in sodium and potassium in all tissue examined with increasing calcium after all fish were returned or fed normal diet. This reflected the continued calcitropic (hypercalcaemic) stress hormone, cortisol effect on common carp post exposure, which initially protected fish against copper accumulation during the exposure phase, but now induced accumulation of the metal in all tissues post exposure [29]. Increasing cortisol production reflects continued depletion of energy in common after exposure to elevated dietary copper. Similarly, the reduction in sodium accounted for the post exposure increase in all tissue examined of copper concentration post exposure [9]. Table 1 and 2 show ionic response to dietary copper in exposure and recovery phase, respectively.

**Moisture**

Although other parameters, including some regulatory ions, were significantly affected by dietary copper exposure and recovery, moisture remained unaffected and has been consistent with other research on copper toxicity [27]. The effects of dietary exposure and recovery of copper on the moisture content are presented in tables 3 and 4.

**Growth and Nutritional Performance**

- Fish exposed to diet (1000 mgCu/kg-1) showed increased weight gain after 42 and this is reflected in the significant increase in condition factor, % of body weight, increased feed intake, increased specific growth rate and increased hematopoietic index, and reduced feed conversion ratio; although the weight gain, specific growth rate and feed conversion ratio were not statistically significant compared with other treatment. The increased weight gain of the diet 2 fish was an indication of the onset of stress-induced increase in hematological indices (hemoglobin, haematocrit and red blood cells), and could be secondary response of fish to irritants. Thus a physiological mechanism of compensation hemoglobin increased to maintain oxygen supply to the fish which translated to increase in the rate of metabolism induced by Na/K ATPase action on blood [40]. Thus fish fed diet 2 tends to increase their feed intake with consequent increase in weight gain observed after 42 days. However, fish fed diet 3 showed a significant decrease in weight gain which is reflected in decreased condition factor,

| Treatment | Gill | Liver | Intestine |
|-----------|-----|------|-----------|
| Na Initial | 257.82±2.50⁴ | 132.4±1.73⁴ | 130.3±0.51²⁴ |
| Diet 1    | 388.14±1.32⁴ | 40.7±0.29⁴  | 438.72±2²⁴⁴ |
| Diet 2    | 476.65±1.60⁴ | 306.87±1.48⁴ | 356.65±0.32²⁴⁴ |
| Diet 3    | 427.57±0.71⁴ | 356.78±0.47⁴ | 597.74±3.89²⁴⁴ |

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specific growth rate, and an increased feed conversion ratio compared with other treatments. Mortality was recorded in the 2000 mgCu kg⁻¹ diet (diet3) exposed treatment within 3 weeks. Several authors have noted reduction in growth rate during dietary copper exposure in fish [8]. But others have not [5,26]. Reduced growth rate have been observed in Nile tilapia fed 2000 mgCu kg⁻¹ [7] which is consistent with this research on common carp fed 2000 mgCu kg⁻¹ (diet3). The recovery phase was characterized by decrease in feed intake by fish fed 1000 mgCu kg⁻¹ pre-exposure. This was reflected in reduced weight gain, specific growth rate, condition factor and increased feed conversion ratio post exposure. The 2000 mgCu kg⁻¹ diet (diet 3) fed fish also showed reduced weight, specific growth rate and a significantly reduced condition factor post exposure. The effects of dietary exposure and recovery on growth are shown in Tables 3 and 6.

**Table 3**: Tissue moisture during copper exposure for 42 days.

| Parameter | Treatment | Gill | Liver | Intestine |
|-----------|-----------|------|-------|-----------|
| Initial   | 73.2 ± 1.20⁺ | 71.50 ± 1.43⁺ | 75.21 ± 0.44⁺ |
| Diet 1    | 74.2 ± 1.80⁺ | 76.12 ± 1.77⁺ | 72.91 ± 0.43⁺ |
| Diet 2    | 69.5 ± 5.64⁺ | 62.15 ± 5.24⁺ | 76.01 ± 1.43⁺ |
| Diet 3    | 76.1 ± 1.62⁺ | 67.65 ± 4.22⁺ | 72.43 ± 0.33⁺ |

Data are means ± S.E (n=3 per value), expressed as percentage (%). Letters with the same subscript in the same column are not significant different (p>0.05).

**Table 4**: Tissue moisture during recovery phase on normal diet from 21 days.

| Parameter                  | Treatment | Exposure |
|----------------------------|-----------|----------|
| Weight gain (g)            | D₀       | 8.50 ± 5.25⁺ |
|                           | D₁       | 16.2 ± 5.25⁺ |
|                           | D₂       | -1.67 ± 4.37⁺ |
| Feed Conversion Ratio      | D₀       | 0.46 ± 0.18⁺ |
|                           | D₁       | 0.18 ± 0.03⁺ |
|                           | D₂       | 0.51 ± 0.25⁺ |
| Specific Growth Rate (%)   | D₀       | 0.07 ± 0.05⁺ |
|                           | D₁       | 0.14 ± 0.02⁺ |
|                           | D₂       | -0.09 ± 0.10⁺ |
| Condition factor (%)       | D₀       | 1.45 ± 0.02⁺ |
|                           | D₁       | 1.60 ± 0.07⁺ |
|                           | D₂       | 1.41 ± 0.05⁺ |
| Hepatosomatic Index (%)    | D₀       | 0.99 ± 0.07⁺ |
|                           | D₁       | 1.47 ± 0.04⁺ |
|                           | D₂       | 1.74 ± 0.12⁺ |
| Mean Ration Size (%)       | D₀       | 0.84 ± 0.04⁺ |
|                           | D₁       | 1.08 ± 0.04⁺ |
|                           | D₂       | 1.06 ± 0.07⁺ |

Data are means ± S.E (n=3 per value), except for ration size where n= number of daily ration size.

**Table 5**: Growth and Nutritional Performance of common carp fed control (diet 1), 1000mgCu kg⁻¹ (diet 2) and 2000mgCu kg⁻¹ (diet 3) for 42 day (Exposure phase).

| Parameter                  | Treatment | Recovery |
|----------------------------|-----------|----------|
| Weight gain (g)            | D₀       | 5.30 ± 0.60⁺ |
|                           | D₁       | 3.67 ± 2.97⁺ |
|                           | D₂       | 0.83 ± 1.87⁺ |
| Feed Conversion Ratio      | D₀       | 0.26 ± 0.03⁺ |
|                           | D₁       | 1.90 ± 0.86⁺ |
|                           | D₂       | 0.28 ± 0.37⁺ |
| Specific Growth Rate (%)   | D₀       | 0.14 ± 0.03⁺ |
|                           | D₁       | 0.09 ± 0.02⁺ |
|                           | D₂       | 0.02 ± 0.05⁺ |
| Condition factor (%)       | D₀       | 1.54 ± 0.02⁺ |
|                           | D₁       | 1.39 ± 0.01⁺ |
|                           | D₂       | 1.43 ± 0.02⁺ |
| Hepatosomatic Index (%)    | D₀       | 1.24 ± 0.02⁺ |
|                           | D₁       | 1.65 ± 0.02⁺ |
|                           | D₂       | 1.84 ± 0.03⁺ |
| Mean Ration Size (%)       | D₀       | 0.90 ± 0.04⁺ |
|                           | D₁       | 0.90 ± 0.03⁺ |
|                           | D₂       | 0.89 ± 0.06⁺ |

*Data are means ± S.E (n 3 per value), except for ration size where n= number of daily ration size.

Letters with the same subscript in the same column are not significant different (p>0.05).

**Conclusion**

In conclusion, copper, although essential in the diet of fish, involved in many physiological and developmental as well as growth, could be deleterious when dietary inclusions exceed that required for proper function of the body in view of its major effects in ionic misbalances in common carp (*Cyprinus carpio*), which could trigger a whole lot of physiological and enzyme processes in the body, necessary for growth and development. Further research is necessary to determine dietary requirement of tropical fish species, including common carp (*Cyprinus carpio*).

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