Hepatic Histomorphological Changes and Oxidative Stress Profile of
Clarias gariepinus (Burchell, 1822) Juveniles Fed with Sodium
Propanoate-Preserved Diets

Patrick E. ABA1,3*, Ifeanyi E. UZOCHUKWU2, Nelson I. OSSAI3,
Ifeanyi G. EKE1

1University of Nigeria, Department of Veterinary Physiology and Pharmacology, Nsukka, Nigeria;
2University of Nigeria, Department of Animal science, Nsukka, Nigeria
3University of Nigeria, Department of Zoology and Environmental Biology, Nsukka, Nigeria;
Patrick.aba@unn.edu.ng (*corresponding author)

Abstract
Sodium propanoate is in the list of approved feed preservatives. However, there is dearth of information on its biological
effects on the C. gariepinus. The present study investigated the effect of sodium propanoate-preserved feed on the hepatic
histomorphometric changes, oxidative stress and inflammatory parameters of C. gariepinus juveniles. One hundred juveniles of
mixed sexes, assigned into 5 groups of 20 fish per group, with each group consisting of 2 replicates of 10 fish, were used for the
investigation. Group A juveniles were fed basal diet, while groups B-E received basal diet incorporated with sodium propanoate
at the rate of 25, 50, 75 and 100 g/15 kg of feed respectively. Treatments were done two times daily for 8 weeks. Samples (sera
and liver) were collected on the last day for evaluation of a few biochemical parameters (malondialdehyde values, catalase
activity, C-reactive protein levels) and histomorphometric alterations in the liver. Results indicated that fish in groups D and E
had higher catalase activities, lower serum levels of C-reactive proteins and an intact hepatic histomorphometry when
compared with the control group. There was no significant difference in the plasma malondialdehyde values in all the groups.
It was concluded that preservation of fish feed with sodium propanoates improved antioxidant status of C. gariepinus and
protected liver histology.

Keywords: Clarias gariepinus; C-reactive protein; hepatic histomorphometry; oxidative stress markers; sodium propanoate

Introduction
Sodium propanoate or sodium propionate is a white
deliquescent crystalline solid with the chemical formula of
Na (C2H5COO). It can be obtained following a reaction
between propionic acid and sodium carbonate. Sodium
propanoate finds application in food industry as food
additive and as a mold inhibitor in bakery (Brock and
Buckel, 2004). It is an approved food preservative. The use
of non-antibiotic promoters such as sodium propanoate
has been widely advocated for owing to deleterious effects
associated with the use of antibiotics in aquaculture (da
Silva et al., 2014).

Preservation of fish feed is very important not only for
extending the shelf life, but the process may also affect the
health status of the fish. Evaluation of blood biochemistry
parameters is vital in ascertaining the effects of xenobiotics, monitoring of therapy and for assessing the
health status of living organisms (Okorie-kanu and Unkalamba, 2016).

It is also noteworthy that stressful conditions in
aquaculture are a major factor militating against large scale
productions in fish industry (Subramanian et al., 2013). It is
therefore important to monitor oxidative stress indicators
such as malondialdehyde and stabilize them for optimum
operation of aquaculture.

Malondialdehyde is a low molecular weight aldehyde
with three carbon molecules. It is a lipid peroxidation
product which is very reliable in assessing oxidative stress
(Grotto et al., 2009). It indicates the level of lipid
peroxidation by reactive oxygen species (ROS). Consequent
upon metabolic processes, different cells in the body of
living organisms constantly generate ROS, which will
eventually peroxidize lipids (Urso and Clarkson, 2003).
Catalase is a tetrameric antioxidant enzyme which is involved in conversion of harmful hydrogen peroxide to innocuous water and molecular oxygen (Chelikani et al., 2004). Its assessment is crucial as it indicates the antioxidant status of an organism and in turn, the ability of such organism to handle oxidative stress. Catalase is commonly seen in all the cells that are exposed to oxygen (Aba and Okenwani, 2015).

C-reactive protein is an acute phase protein which serves as a reliable marker of inflammation (Young et al., 1999). Its synthesis is in the liver. The serum level of C-reactive protein rapidly increases during inflammation because of its role in opsonization (Ballou and Kushner, 1992). C-reactive protein will bind to a substance known as phosphocholine which is expressed by damaged cells. It also binds to peptosaccharides and polysaccharides expressed by bacteria and fungi (Ballou and Kushner, 1992).

Liver is a vital organ in the body, being involved in a greater percentage of metabolizing foreign materials and drugs. Histomorphometric assessment of the liver always reveals its health status (Aba and Asuzu, 2015). There is dearth of information on the effect of sodium propanoate on some oxidative stress parameters and liver histology of Clarias gariepinus.

The present study therefore was carried out to investigate the effects of preserving fish feed with sodium propanoate on some oxidative stress indicator parameters, antioxidants profile, inflammatory status and hepatic histomorphometry of Clarias gariepinus juveniles.

Materials and Methods

Experimental fish

One hundred Clarias gariepinus juveniles of mixed sexes used in the study were obtained from a cat fish farm, Department of Zoology, University of Nigeria. They were acclimatized for one week during which they were fed normal (basal) fish diet, two times daily and water was changed two times weekly. On the second week, experimental diet (basal diet + sodium propanoate) was introduced.

Experimental diets preparation

The data below shows the composition of the experimental diet (Table 1).

Experimental design

The fish juveniles were randomly assigned into 5 groups (A-E) of twenty (20) per group. The five groups were each, sub-assigned into two (2) replicates of 10 fish per replicate. Each replicate was housed in a plastic bowl containing 15-20 litres of clean tap water. The treatments were as follows:

Group A: Received experimental diet containing 0% sodium propanoate/15 kg of feed (Control).

Group B: Received experimental diet containing 25% sodium propanoate/15 kg of feed.

Group C: Received experimental diet containing 50% sodium propanoate/15 kg of feed.

Group D: Received experimental diet containing 75% sodium propanoate/15 kg of feed.

Group E: Received experimental diet containing 100% sodium propanoate/15 kg of feed.

All treatments were made two (2) times per day and lasted for eight weeks. At the end of the treatments, samples were collected for laboratory analyses.

Sample collections

Blood samples were collected via the mid ventral tail vein into sample bottles devoid of anticoagulant. The blood samples were centrifuged at 10,000 g for 10 mins and sera were harvested for biochemical analyses.

The fish were sacrificed under mild ether anaesthesia and the liver samples were collected for histopathology evaluation.

Evaluation of biochemical parameters

Estimation of lipid peroxidation (Malondialdehyde)

Lipid peroxidation was estimated by measuring spectrophotometrically, the level of the lipid peroxidation product, malondialdehyde (MDA) as described by Wallin et al. (1993).

Lipid degradation occurs forming such products as malondialdehyde (from fatty acids with three or more double bonds), ethane and pentane (from the n-terminal carbons of 3 and 6 fatty acids, respectively). MDA reacts with thiobarbituric acid to form a red or pink coloured complex which in acid solution absorbs maximally at 532 nm.

Exctly 0.1 ml of the serum was mixed with 0.9 ml of H2O in a test tube. A volume of 0.5 ml of 25% TCA (trichloroacetic acid) and 0.5 ml of 1% TBA (thioarbituric acid) in 0.3% NaOH were also added to the mixture. The mixture was boiled for 40 min in a water-bath and then cooled in cold water. Then, 0.1ml of 20% sodium dodecyl sulfate (SDS) was added to the cooled solution and mixed properly. The absorbance was taken at 532 nm and 600 nm against a blank.

\[%\text{TBARS} = (A_{532} - A_{600}/0.527\times0.1) \times 100 (mg/dl)\]

Table 1. Experimental diet formulation with different levels of sodium propanoate (in grams per 15 kg diet)

<table>
<thead>
<tr>
<th>Feed stuff</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fish meal</td>
<td>460</td>
<td>460</td>
<td>460</td>
<td>460</td>
<td>460</td>
</tr>
<tr>
<td>Soybean meal</td>
<td>460</td>
<td>460</td>
<td>460</td>
<td>460</td>
<td>460</td>
</tr>
<tr>
<td>Wheat offal</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
</tr>
<tr>
<td>White corn</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
<td>35</td>
</tr>
<tr>
<td>Vitalyte</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>DCP</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Sodium propanoate</td>
<td>0</td>
<td>25</td>
<td>50</td>
<td>75</td>
<td>100</td>
</tr>
</tbody>
</table>

NB: DCP = Dicalcium phosphate
Estimation of catalase activity

The activity of catalase was assayed by the method according to Sinha (1972). Dichromate in acetic acid was reduced to chromic acetate, when heated in the presence of hydrogen peroxide with the formation of acid as an unstable intermediate. The chromic acetate formed was measured at 590 nm. Catalase was allowed to split H_2O_2 for different periods of time. The reaction was stopped at different time intervals by the addition of dichromate acetic acid mixture and the remaining H_2O_2 was determined by measuring chromic acetate spectrophotometrically after heating the reaction mixture. The chemicals used were as follows:

1. Phosphate buffer, 0.01 M, pH = 7
2. Hydrogen peroxide, 0.2 M
3. Potassium dichromate, 5%
4. Dichromate acetic acid reagent: Potassium dichromate and glacial acetic acid were mixed in ratio of 1:3. From this, 1 ml was diluted again with 4 ml of acetic acid.

Procedure

Exactly 0.1 ml of plasma and 0.4 ml of H_2O_2 were added to 0.9 ml of phosphate. The reaction was stopped after 15, 30, 45 and 60 seconds by adding 2 ml of dichromate acetic acid mixture. The tubes were kept in a boiling water bath for 10 min and cooled. The colour developed was read at 530 nm. Standards in the concentration range of 20-100 µmoles were processed for the test. The activity of catalase was expressed as U/ml for plasma (U = µmoles of H_2O_2 Utilised / second).

Catalase activity = \log A/B \times 0.23/0.00693

CRP assay

The method of Dupuy et al. (2003) was used. Exactly 450 microlitre of R1 (activation buffer) was mixed with 50 microlitre of R2 (latex reagent: uniform suspension of polystyrene latex particles coated with anti-CRP antibody) and incubated at 37 °C for 5 min. Thereafter, 5 ml of serum sample were added to the above mixture and absorbance was read at wavelength 630 nm after 10 sec as A1 and after 2 mins as A2.

Similarly, the CRP standard (calibrator) was processed in the same manner as the samples.

Concentration of CRP in the sample = change in absorbance (A) of the sample \times concentration of standard / change in absorbance (A) of standard.

Where change in A = (A2-A1)

Concentration of the standard = 2mg/dl (A lyophilized preparation of serum equivalent to the stated amount of CRP on a mg/dl basis).

Histopathology examination

The histological examination of the liver tissues of the fish was done using the method of Drury et al. (1967).

A. Fixation and washing

Formalin (10%) was used as fixative and for the purpose of preservation. A thin section of the tissue (about 1 to 2 cm in diameter) was trimmed with a sharp razor blade. The small pieces of the tissue were placed in the 10% formalin, the container was shaken gently several times to make sure that the fluid had reached all surfaces and that pieces were not sticking to the bottom. This was then incubated at 25 °C for 24 h, to allow proper fixing. The fixed tissue pieces were washed with running water for 24 h to free them from excess fixatives.

B. Dehydration

Water was removed from the tissue before embedding the tissue in paraffin. The dehydration was achieved by immersing the thin sections of the tissue in automatic tissue processor containing 12 jars. The first three jars contained 70, 90 and 95% absolute alcohol respectively. This was done to remove the water content in the tissues. The absolute alcohol reduced the shrinking that occurred in the tissue. The time for each step was 30 min. A second change of absolute alcohol was included to ensure complete removal of water. This was achieved in the second three jars of the automatic tissue processor.

C. Clearing

Solutions of xylene were used for clearing the tissue sections. This step was achieved in the third three jars of the automatic tissue processor. Because the alcohol (ethanol) used for dehydration would not dissolve or mix with molten paraffin, the tissue was immersed in xylene solution which was miscible with both alcohol and paraffin before infiltration could take place. Clearing was done to remove opacity from dehydrated tissue. A period of 15 min was allowed to elapse before the tissue was removed from the solution for infiltration with paraffin.

D. Infiltration with paraffin

Paraffin wax at 50 to 52 °C was used to infiltrate the tissue. The tissue was transferred directly from the clearer to a bath containing melted paraffin. After 30-60 min incubation in the first bath, the tissue was then removed to a fresh dish of paraffin contained in the fourth three jars of the automatic tissue processor for a similar length of time.

E. Embedding (blocking) with paraffin

As soon as the tissue was thoroughly infiltrated with paraffin, it was allowed to solidify around and within the tissue.

F. Paraffin sectioning

The embedded blocks were trimmed into squares and fixed in the microtome for sectioning after which the sections were floated on a water bath.

G. Mounting

Glass slides were thoroughly cleaned and a thin smear of albumen fixative was made on the slides. The albumenized slide was used to collect the required section from the rest of the ribbon in the water. The section on the glass slide was kept moist before staining.

H. Staining with haematoxylin

The slides were passed through a series of jars containing alcohols of decreasing strength and various staining solutions.

I. Microscopic observation of slide

The slides prepared were mounted on a photomicroscope and viewed at different magnifications. A photograph of each of the slides was taken.

Statistical analysis

Data on biochemical parameters were analyzed with One-way Analysis of Variance (ANOVA). Variant means were separated with Duncan’s Multiple Range post hoc test. Significance was accepted at probability level less than 0.05.
Results

There were no significant differences (p > 0.05) in plasma malondialdehyde values of all the groups across the treatment period. The serum catalase activities of groups D and E fishes were significantly (p < 0.05) higher than their group A (normal, control) counterpart, while the serum C-reactive protein levels of the control (Group A) and those that received the lowest incorporation of sodium propanoate (Group B) fishes were significantly higher than those of groups C, D and E (Table 2 and Figs 1-5).

Table 2. Effect of sodium propanoate on some oxidative stress parameters of *Clarias gariepinus* (African cat fish)

<table>
<thead>
<tr>
<th>Group/Parameters</th>
<th>Malondialdehyde (mg/ml)</th>
<th>Catalase (IU/L)</th>
<th>C-reactive protein (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>2.04 ± 0.19*</td>
<td>3.58 ± 0.53*</td>
<td>3.76 ± 0.08*</td>
</tr>
<tr>
<td>B</td>
<td>1.76 ± 0.32*</td>
<td>3.77 ± 0.62*</td>
<td>3.73 ± 0.11*</td>
</tr>
<tr>
<td>C</td>
<td>1.74 ± 0.08*</td>
<td>4.83 ± 0.29*</td>
<td>2.95 ± 0.09*</td>
</tr>
<tr>
<td>D</td>
<td>1.41 ± 0.25*</td>
<td>5.54 ± 0.16*</td>
<td>2.65 ± 0.24*</td>
</tr>
<tr>
<td>E</td>
<td>1.33 ± 0.16*</td>
<td>5.29 ± 0.33*</td>
<td>2.53 ± 0.23*</td>
</tr>
</tbody>
</table>

Different superscripts along the same column (across groups) indicate significant differences at p < 0.05

Fig. 1. Group A H&E X400 LIVER showing mild localized periportal infiltration by inflammatory cells (arrows)

Fig. 2. Group B H&E X400 LIVER showing normal hepatocytes arranged in cords with central vein filled with red blood cells

Fig. 3. Group C H&E X200 LIVER showing normal liver architecture

Fig. 4. Group D H&E X400 LIVER normal hepatocytes showing central vein
Discussion

The present study investigated the effects of sodium propanoate-preserved fish feed on a few biochemical parameters and on the hepatic histology.

Upon eight weeks administration of sodium propanoate in feeds, results indicate that the catalase activities of the group that received the highest concentration of sodium propanoate in feed was significantly higher than that of the control group which received basal diet (Table 2). This indicates that inclusion of sodium propanoate enhanced the activity of the antioxidant enzyme. Catalase plays vital role in amelioration of oxidative stress. Catalase causes the decomposition of hydrogen peroxide which is generated in the course of tissue metabolism to water and molecular oxygen (Chelikani et al., 2004). Antioxidant properties of sodium propanoate had earlier been reported in Zebra fish in a study conducted by Safari et al. (2016).

The serum levels of C-reactive protein of the fish group that received the highest incorporation of sodium propanoate were significantly lower than that of the normal control group (Table 1). C-reactive protein is an acute phase protein that indicates the levels of inflammation (Young et al., 1999). The decreases in the serum levels of C-reactive protein in the group that received feed with sodium propanoate indicates that sodium propanoate may have anti-inflammatory properties. Literature search revealed that sodium propanoate is an anti-fungus (Brock and Buckel, 2004). Tedelindet et al. (2007) also submitted the anti-inflammatory properties of short chain fatty acids such as propionate.

The finding of insignificant changes in the plasma malondialdehyde levels of control group when compared with the treated groups (groups that received sodium propanoate -incorporated feeds) indicate that the sodium propanoate may not have interfered with lipid peroxidation of the fish (Table 1). Malondialdehyde is a stable product of lipid peroxidation that is useful in assessing the magnitude of lipid peroxidation in the cell membranes (Urso and Clarkson, 2003; Grotto et al., 2009).

The histopathology results suggest that the dietary sodium propanoate may have protected the liver against any inflammatory or degenerative conditions. The photomicrographs of the liver indicates that the fish groups that received sodium propanoate-incorporated feed showed normal hepatocytes and hepatic architecture (Figs. 2-5) when compared to that of the normal control (Fig. 1) which showed mild degenerative lesion. It can be suggested that the finding of such lesion in the liver of fish fed normal basal diet may be indicative that the feed could have deteriorated since there was no preservative incorporated. Absence of preservative such as sodium propionate predisposes feeds to becoming mouldy (Brock and Buckel, 2004). Mouldy feed could be potentially harmful to the liver, especially if aflatoxin is involved. Aflatoxins particularly have been shown to be involved in oxidative stress and inflammatory processes and could cause hepatocellular disorders (Qin et al., 2016). Literature search also showed that aflatoxin is toxic in channel cat fish and tilapia (Chavez-Sanchez et al., 1994).

Conclusions

It was concluded that preservation of Clarias gariepinus feed with varying levels of sodium propanoate produced desirable effects. Groups that received feeds incorporated with sodium propanoate showed normal liver histology and improved antioxidant status, with no inflammation when compared to the fish group that did not receive the sodium propanoate-incorporated feed. Apart from being used as preservative, sodium propanoate could also positively add to the value of feed.

References


