



# Variations of Lipids in Different Tissues (Liver, Intestine and Brain) of Fresh Water Cat Fish [*Heteropneustes fossilis* (Bloch)]

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## Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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## ABSTRACT

In the present study on characterization and identification of lipids by thin layer chromatography in different tissues of *Heteropneustes fossilis* showed a considerable variation. The number of lipid spots was highest in brain compared to other tissues. The Rf values of spots indicate the regions of similarity between tissues. Comparison of Rf values with those of standards indicated that the presence of phospholipids, choline lipids, glycolipids, ninhydrine, amino group containing lipids were present. Intestine showed more amount of glycolipids and low amount of ninhydrine containing lipids, While liver possessed Iodine containing lipids.in addition these, whereas brain showed glycolipids, ninhydrine containing lipids, Iodine containing lipids and choline lipids too. The present study revealed that the Brain contain highest amount of different classes of lipids. The Liver tissue has shown less number of lipids. Glycolipids and general class of lipids are present in all tissues with minor variations, choline lipids present in brain but not in remaining tissues, while Phospho lipids are present in intestine and brain only, whereas amino group containing lipids were more in all tissues when compared with other class of lipids.

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## 1. INTRODUCTION

“Thin-layer chromatography (TLC) is a technique that has been routinely used for the separation and identification of lipids. Here we describe an optimized protocol for the steady state labeling, separation, and quantification of yeast phospholipids using TLC. There is a considerable variation between the tissues of *H. fossilis* was observed during our investigation, the fast moving spots are probably the neutral lipids or cholesterol esters. The staining of gill tissue of *H. fossilis* was deeply stained and moderately stained with sulphuric acid and ninhydrine reagents respectively. The biochemical and pharmacological properties and the lipid composition of epidermal secretions from various catfish of the Arabian Gulf have been investigated” [1]. However, there has been a very little mention about the fatty acid (FA) composition of muscle of catfish lipid, with the report by [2], being the only publication on FA composition in *Arius catfish* n-3 polyunsaturated FA (n-3 PUFA) in the diet has been recognized to have important beneficial properties for the prevention of human coronary artery disease [3].

“Sea and freshwater fish, which constitute the majority of water products, make up an important part of animal food sources for human. Fish are quite different from the other animal food sources, because they provide low energy and have high-level proteins, which contain all essential amino acids. So they are beneficial nutrition sources” [4]) “Fish are not only beneficial protein source but also contain considerable amount of unsaturated fatty acids, and thus the studies on lipid biochemistry have been considered so important recently [5-10]. The lipids are the most important biochemical compounds of fish” [11]. “Fish store the lipids in various organs; particularly in muscles and liver. On the contrary, the mammals store in adipose tissue. A great amount of these lipids are transferred to the different parts of the body to be used for various physiological actions” [12]. “The freshwater fish constitute a great food potential for human. A large part of these fish species are cultivated forms. Since some other new food sources which have the same content are found there is a growing need for information about the chemical composition of the various species and their food contents. For the cultivation of these fish, some important characteristics, such as nutrition properties, biochemical structure and

growth conditions need to be known. It is of great importance to know the seasonal variations of the lipids and the lipid amount of the fish, which is economically important and willingly consumed” [12].

Overall, adequate accuracy was obtained in this study. The proposed method provides a fast and efficient means of identifying fish and feed for quality control purposes.

## 2. METHODOLOGY

The tissues were collected from the fish (*Heteropneustes fossilis*) i.e liver,, intestine, and brain. Tissues were collected by the method of [13] as follows.

The different tissues were collected from various parts of the body and tissues were weighed to nearest milligram and homogenized with chloroform methanol mixture (v/v) (2:1) to a final dilution of 10fold the weight of tissue i.e., the homogenate 1 gm of tissue was diluted to a volume of 10 ml of chloroform methanol mixture solution. The mixture was homogenized and centrifuged at 2000 rpm for 5 mins. The organic layer was collected and evaporated in vacuum at 35°C and the residue was dissolved in 0.5ml of solvent (Chloroform : Methanol. 2:1). The TLC was run on a readymade silica gel plates (E- merck) in a solvent system of Chloroform, methanol, water (65:25: 10) system. The plates were dried at room temperature. The plates were observed under UV lamp and the spots were marked with pencil. For routine analysis of plates were sprayed with 1) Iodine 2) H<sub>2</sub>SO<sub>4</sub> 3) Ninhydrine 4) Dittmer- Lester and 5) Dragendorff's reagent. R<sub>f</sub> values of all the spots were determined immediately.

### 2.1 TLC Plates

Readymade marketed (E. Merck AG, Darmstadt, Germany) silica gel-G coated on aluminium plates with thickness of the layer 0.2mm were used, the solution of total lipids prepared in (Chloroform : Methanol, 2:1) was applied on the plates at a distance of 1 cm from lower edge of the plate. The loaded plates were developed in the solvent system mentioned above.

### 2.2 Reagents Preparation

Following reagent systems were employed in identifying the lipid fractions in accordance with the procedures given by Kates (1986).

### 2.3 Iodine Vapours

Iodine vapours were used as a general purpose detector for all lipids, since both saturated and unsaturated lipids absorb iodine by simple dissolution. Iodine crystals were packed between the blotting papers and wetted with Chloroform. The TLC plates dried at room temperature and covered with Petri dish. Lipid spots were turned into yellow on exposure to Iodine vapours. The spots were scanned immediately and their  $R_f$  values were noted on after exposure to Iodine vapours.

### 2.4 Sulphuric Acid Reagent

Sulphuric acid – di chromate reagent was prepared by dissolving 0.6 gm of  $K_2Cr_2O_7$  in 55%  $H_2SO_4$ . The reagent was sprayed on to the developed plate and then the plates were heated on hot plate from low heat and gradually increasing to high heat for 5-10 minutes. All lipids were charred and appeared as black spots. Glycolipids and sterols appeared as red spots upon low heating.

### 2.5 Nin Hydrin Reagent

This was used to detect Phosphatides or lipids containing a free amino group. Phosphatidyl ethanolamine (PE), Phosphatidyl serine (PS) and Phosphatidyl inositol (PI) were detected by this reagent.

### 2.6 Reagent Preparation

0.25 gm of reagent grade Ninhydrin was dissolved in 100ml of Acetone. After spraying the reagent, reagent plates were heated at  $60^{\circ}C$  for 3-5 minutes the spots appeared purple in colour.

### 2.7 Dittmer – Lester Reagent (Modified by vaskovasky and kostetsky)

This reagent was used to detect phospholipids for which it is specific, and can detect as little as  $10\mu g$  of lipid. The phospholipids appeared as blue spots on a white background with few minutes without heating.

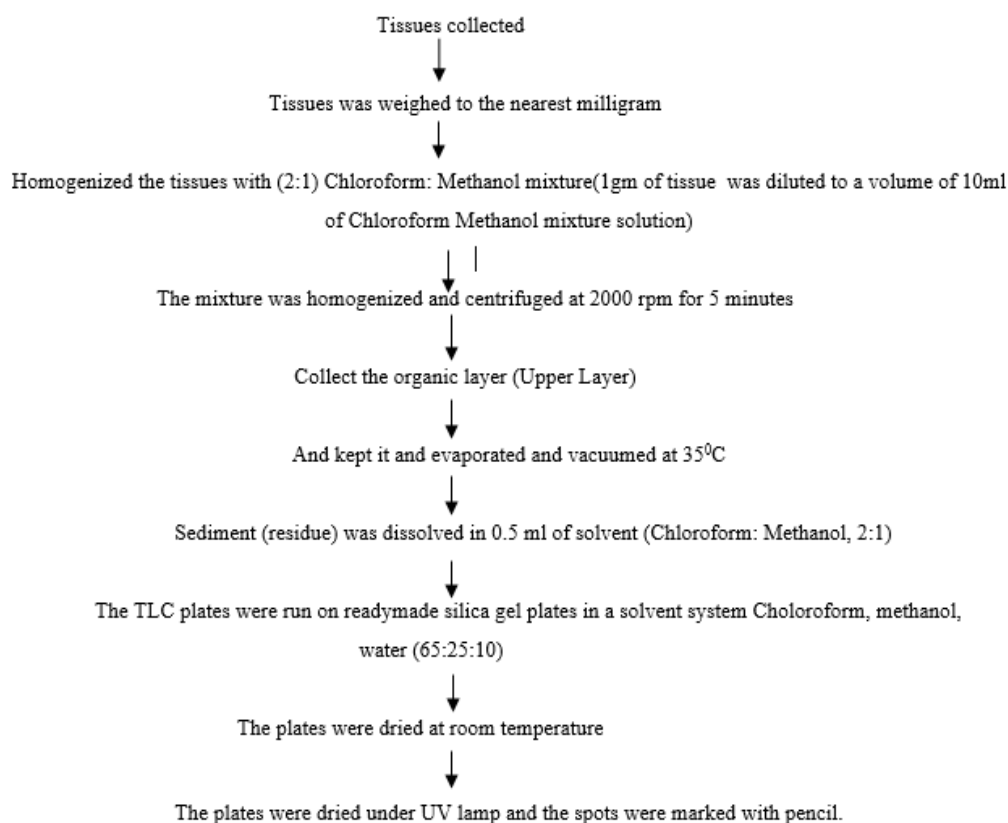


Chart 1. Identification of lipids protocol

## 2.8 Reagent Preparation

### 2.8.1 Solution A

16 gm of Ammonium molybdate in 120 ml distilled water.

### 2.8.2 Solution B

40 ml of Con, HCL, 10ml of mercury and 80 ml of sol. A were added, shaken for 30 minutes and filtered. To remainder of the solution (A) 200 ml of Con. H<sub>2</sub>SO<sub>4</sub> and all the solution (B) were added, cooled and stored. This was bright green coloured solution and was stabled at room temperature. Just before use this solution was diluted with distilled water (1:3 v/v). the final spray reagent was Amber colored.

## 2.9 Dragendorff's Reagent

The stock solution (5ml ) was mixed with 15 ml of distilled water immediately before use.

### 2.9.1 Solution- I

2.7 gm of Bismath nitrate in 100 ml of Acetic acid.

### 2.9.2 Solution- II

10 gm in 25 ml of Potassium Iodide. Before use 20 ml of solution I and 5 ml of solution-II were mixed in and diluted with 70 ml of distilled water.

## 3. RESULTS

The pattern of lipids observed through TLC in the three tissues under examination is presented in Figs. 1,2,3 and R<sub>f</sub> values of individual spots and the staining patterns with different reagents Viz., Dittmer reagent, Dragendorff's, Sulfuric acid, Iodine and Ninhydrin were presented in Table 1, 2,3.

### 3.1 Liver

This tissue showed only one blue color spot of phospholipids when treated with Dittmer reagent with R<sub>f</sub> value 90±0.5, this spot is not clearly visible. Remaining lipid spots did not appeared (lane1). The lipid spots when treated with Dragendorff's, reagent (Lane-2) with R<sub>f</sub> values 20±0.5, 30±0.5, 40±0.5, 80±0.5 and 90±0.5 which were not clearly visible. The lipid spots

with R<sub>f</sub> values 20±0.5, 30±0.5, 40±0.5, 80±0.5 and 90±0.5 were moderately stained with sulphuric acid and exhibited five black, glycol lipid spots, whereas R<sub>f</sub> value 10±0.5, 50±0.5, 60±0.5, 70±0.5 were not appeared (Lane-3). The lipid spots with R<sub>f</sub> values 40±0.5, 50±0.5 and 60±0.5 were moderately stained with iodine vapour and exhibited three yellow spots, with R<sub>f</sub> values 40±0.5, 50±0.5, and 60±0.5 were moderately stained (Lane-4). On staining for lipids with ninhydrine, amino group containing lipids appeared as three purple spots in liver tissue (Lane-5). Whereas R<sub>f</sub> value 70±0.5 and 90±0.5 were deeply stained and R<sub>f</sub> value 60±0.5 was moderately stained (Table.1 and Fig. 1).

### 3.2 Intestine

This tissue showed four blue color spots of phospholipids which stained moderately with Dittmer reagent with R<sub>f</sub> values were 50±0.5, 70±0.5, 80±0.5 and 90±0.5 (lane1). Spots with R<sub>f</sub> value 30±0.5 was moderately stained when treated with Dragendorff's reagent and appeared as an orange lipid spot, remaining spots with R<sub>f</sub> values viz., 20±0.5, 70±0.5 and 90±0.5 were not clearly visible (Lane-2). When this tissue treated with vapors of sulphuric acid, spots with R<sub>f</sub> values 40±0.5, 50±0.5, 60±0.5, 80±0.5 and 90±0.5 were exhibited as five black color spots and all these spots were deeply stained (lane-3). This tissue was treated with Iodine reagent and lipid spots with R<sub>f</sub> values 50±0.5, 60±0.5, 70±0.5, 90±0.5 were faintly stained (lane-4). Spots with R<sub>f</sub> values 20±0.5, 30±0.5, 40±0.5, 60±0.5, 70±0.5 and 80±0.5 were moderately stained when treated with Ninhydrin reagent and exhibited 6 purple color spots of amino acid containing lipid (lane-5) (Table 2 and Fig. 2).

### 3.3 Brain

Lane-1 of brain tissue showed lipid spots with R<sub>f</sub> value 40±0.05, 50±0.05, 60±0.05, 70±0.05, 80±0.05, 90±0.05 treated with Dittmer reagent exhibited blue color spots; in Lane-2 the lipid spots with R<sub>f</sub> value 10±0.05, 20±0.05, 30±0.05, 50±0.05 were not visible clearly. The lipid spots with R<sub>f</sub> value 10±0.05 & 90±0.05 were faintly stained with Dragendorff's reagent and showed nine orange color spots of Choline lipids, in Lane-3 six lipid spots were exhibited and deeply stained, when treated with sulphuric acid, their R<sub>f</sub> values are 20±0.05, 30±0.05, 50±0.05, 60±0.05, 80±0.05, 90±0.05 (Glycolipid spots). Lane-4

when treated with Iodine vapors eight yellow color lipid spots were observed, all these spots were moderately stained. Lane-5 staining with Ninhydrine reagent showed seven purple color

spots for amino group with  $R_f$  values  $10 \pm 0.05$ ,  $20 \pm 0.05$ ,  $30 \pm 0.05$ ,  $40 \pm 0.05$ ,  $70 \pm 0.05$ ,  $80 \pm 0.05$ ,  $90 \pm 0.05$ . All these spots were deeply stained (Table 3 and Fig. 3).

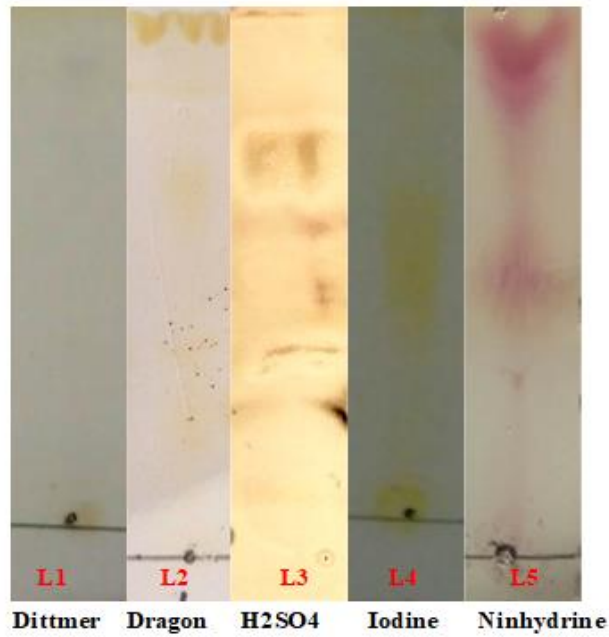


Fig. 1. TLC of lipids in Liver tissue stained with different reagents

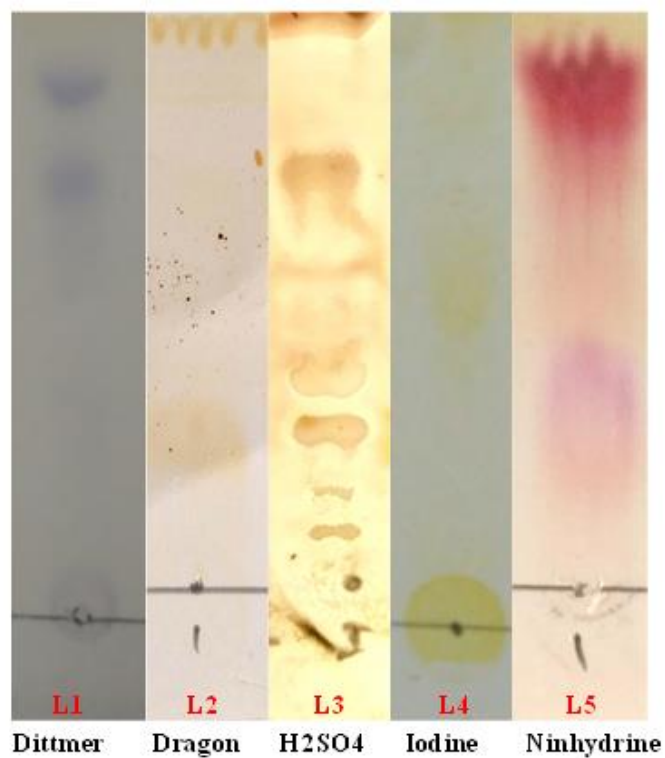


Fig. 2. TLC of lipids in intestine tissue stained with different reagents

**Table 1. TLC of lipids in Liver tissue stained with different reagents**

Liver tissue treated with different reagents for lipid identification R <sub>f</sub> values									
Reagent	10±0.5	20 ± 0.5	30 ±0.5	40 ±0.5	50± 0.5	60± 0.5	70± 0.5	80± 0.5	90± 0.5
Dittmer-Lister	-	-	-	-	-	-	-	-	±
Dragon-Draft	-	±	±	±	-	-	-	±	±
Sulphuric Acid	-	+	+	+	-	-	-	+	+
Iodine	-	-	-	++	++	++	-	-	-
Ninhydrine	-	-	-	-	-	++	+++	-	+++

**Table 2. TLC of lipids in Intestine tissue stained with different reagents**

Intestine tissue treated with different reagents for lipid identification R <sub>f</sub> values									
Reagent	10±0.5	20 ± 0.5	30 ±0.5	40 ±0.5	50± 0.5	60± 0.5	70± 0.5	80± 0.5	90± 0.5
Dittmer-Lister	-	-	-	-	++	-	++	++	++
Dragon-Draft	-	±	++	-	-	-	±	-	±
Sulphuric Acid	-	-	-	+++	+++	+++	-	+++	+++
Iodine	-	-	-	-	+	+	+	-	+
Ninhydrine	-	++	++	++	-	++	++	++	+++

**Table 3. TLC of lipids in Brain tissue stained with different reagents**

Brain tissue treated with different reagents for lipid identification R <sub>f</sub> values									
Reagent	10±0.5	20 ± 0.5	30 ±0.5	40 ±0.5	50± 0.5	60± 0.5	70± 0.5	80± 0.5	90± 0.5
Dittmer-Lister	-	-	-	+	+	-	+	+	+
Dragon-Draft	+	+	+	+	+	+	+	+	+
Sulphuric Acid	-	+++	+++	-	+++	+++	-	+++	+++
Iodine	++	++	-	++	++	++	++	++	++
Ninhydrine	+++	+++	+++	+++	-	-	+++	+++	+++

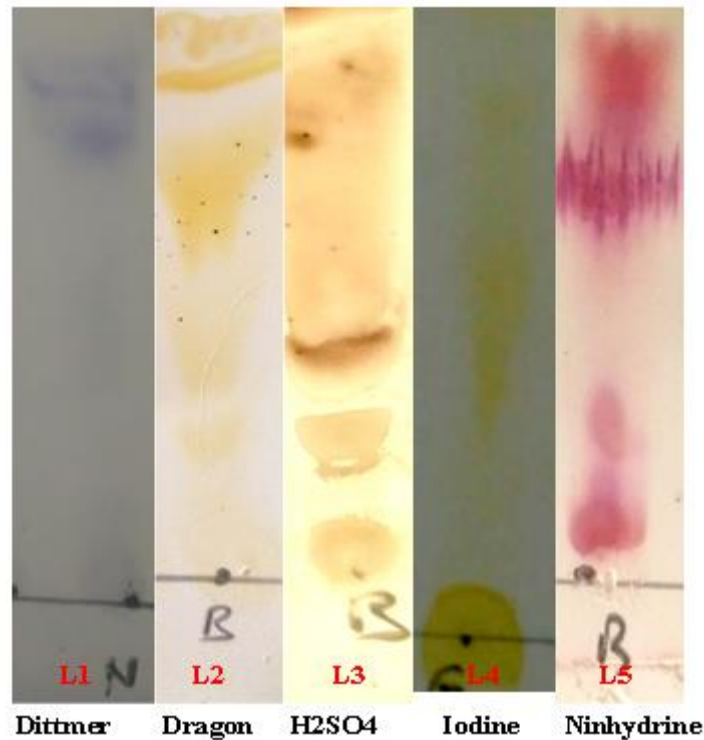


Fig. 3. TLC of lipids in brain tissue stained with different reagents

#### 4. DISCUSSION

There is a considerable variation between the tissues of *H.fossilis* was observed during our investigation, the fast moving spots are probably the neutral lipids or cholesterol esters. The staining of gill tissue of *H.fossilis* was deeply stained and moderately stained with sulphuric acid and ninhydrine reagents respectively. “The biochemical and pharmacological properties and the lipid composition of epidermal secretions from various catfish of the Arabian Gulf have been investigated” [14] “However, there has been a very little mention about the fatty acid (FA) composition of muscle of catfish lipid, with the report by Sinclair *et al.*, 1983, being the only publication on FA composition in *Arius catfish*n-3 polyunsaturated FA (n-3 PUFA) in the diet has been recognized to have important beneficial properties for the prevention of human coronary artery disease” [15].A similar study carried out by [16]) “in muscle and liver of rainbow trout (*Oncorhynchus mykiss*) determined that the variations of the levels of lipids in liver and other organs are the results of irregular seasonal variations which affect the fish diet”. “However, it may be noted that along with seasonal variation, habitat conditions also influence fat synthesis in the fish. In lotic habitat, more amount of lipid

content was observed in liver during winter season but significantly lower during the monsoon but in lentic habitat, apparently low lipid content was observed during winter season” [17-21].The fishes selected in the present study are of high consumer demand having good taste. Present study will provide a detailed understanding of the necessity of the healthy life for the fish consuming people. This information also may be helpful to evaluate the nutritional significance of the fishes. We concluded that fish lipid should not be avoided in human nutrition; its consumption should be minimized by individuals and populations at risk of cardiovascular diseases and its associated sequel [22-24].

#### 5. CONCLUSION

The results obtained from the study should bring about consciousness in preserving this edible fish species not only for its nutritional value but also for maintaining a proper food chain in aquatic zones where they thrive. As we know, the concentration of the lipid and fatty acid content in a fish *Heteropneustes fossilis* depends on its habitat and feeding habit, which in turn depends on the food sources present in the reservoir where the fish completes its entire life cycle, our findings will definitely give an insight

on the type of environment as well as fish feed required for properly maintaining and farming of such fish species and thereby increase its nutritive value. The present study plays an important role in industry effluents risk assessment using a fish model that is a key indicator of environmental toxicity.

## COMPETING INTERESTS

Authors have declared that no competing interests exist.

## REFERENCES

1. Deepasree MI, MS. Rajendran Nair Histological and Protein Profile Alterations of Liver of Freshwater Fish *Channa punctatus* (Bloch), on Exposure to Fungicide, Fytran, International Journal of Science and Research (IJSR), 2017;6(6): 1370-1374.
2. APHA Standard methods for the examination of water and wastewater. 21st Edn., Washington, DC: American Public Health Association, AWWA, WPCP-2005.
3. Barbara Teixeira et al., Changes of Enzymes Activity and Protein Profiles Caused by High-Pressure Processing in Sea Bass (*Dicentrarchus labrax*) Fillets. J. Agric. Food Chem. 2013;61: 2851–2860.
4. Mukesh K N, The effect of pesticides on fish fauna of Bhopal lower lake (M. P.). African Journal of Environmental Science and Technology. 2013;7(7):725-727.
5. Mahvash Jafari; 2012 Effects of paraoxon on serum biochemical parameters and oxidative stress induction in various tissues of Wistar and Norway rats September 2012.
6. Murty et al., The effect of endosulfan and its isomers on tissue protein, glycogen, and lipids in the fish *Channa punctate*. 1982;7(3):280-286.
7. Nagaraju et al. Lethal and sublethal effects of profenofos and carbosulfan on protein pattern of indian major carp, *labeo rohita* (Hamilton) Mai 2016 Biologie 25/2 77-83
8. Laemmli UK Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature. 1970;227: 680-685.
9. Holmes and Master; The developmental multiplicity and isoenzyme status New York, of cavian esterases. Biochimica et Biophysica Acta. 1967;123:379-399.
10. George Jesslin, Arumugam Sabaridasan, Ponnu Edwinthangam, Radhakrishanan Palanikani and Ramaiah Soranam. Electrophoretic studies on the muscle proteins of three species of genus *Puntius* (*Osteichthyes cyprinidae*). International Journal of Research in Fisheries and Aquaculture, 2013;3(2):16-20.
11. Theophilus J, Rao PR. Electrophoretic studies of three species of channa genus. Indian. J. Fish; 1998.
12. Suneetha K et al., Changes in protein subunits induced by endosulfan and fenvalerate in fresh water fish *Labeo rohita* through SDS-PAGE, Journal of Environmental Biology. 2010;31(5):759-763.
13. Folch J, Lees M, Sloane Stanley GH. J Biol Chem. 1957;226:497
14. Al-Hassan JM., Afzal, M., Ali, M., Thomson, M., Fayad, T., Fayad, S., et al. (1986). Lipid composition of the epidermal gel secretion from the Arabian Gulf catfish (*Arius thalassinus* Ruppell). Comp. Biochem. Physiol. B 85 (1), 41–47. DOI: 10.1016/0305-0491(86)90218-X
15. Justin Raj and Baby Joseph, impact of acetamiprid toxicity on electrophoretic patterns in liver, brain and gill tissues of the fish *Oreochromis mossambicus*. International Journal of Zoological Research. 2017;13:120-124.
16. Austin, B. The effects of pollution on fish health. J. Appl. Microbiol. 1999;85:234-242. Elhalwagya EA. et al. Comparative study on pesticide mixture of organophosphorus and pyrethroid in commercial formulation. Environ. Toxicol Pharmacol. 2009;28:219-224.
17. Helan Chandra. Acute toxicity studies on the fresh water fish *Clarias batrachus* exposed to pesticide rogorin J; 2015. Issn: 0974-2115 January 2015.
18. Sridhar et al. Acute toxicity studies on the fresh water fish *Clarias batrachus* exposed to pesticide Rogorin J. Helan Chandra\*1, Sajda 2, S. 3; 2015. Issn: 0974-2115. January-March 2015.
19. Murty et al. The effect of endosulfan and its isomers on tissue protein, glycogen, and lipids in the fish *Channa punctate*. 1982;7(3):280-286.
20. Nagaraju et al., Lethal And Sublethal Effects Of Profenofos And Carbosulfan On Protein Pattern Of Indian Major Carp, *Labeo Rohita* (Hamilton) Mai 2016 Biologie 25/2 77-83.



21. APHA Standard methods for the examination of water and wastewater. 21st Edn. Washington, DC: American Public Health Association, AWWA, WPCP; 2005.
22. Ramadan A, A. genotoxic effects of butataf herbicide on Nile Tilapia. Journal of the Arabian Aquaculture Society. 2007;2(1): 70-89.
23. Sandal S. et al., Genotoxic effects of chlorpyrifos, cypermethrin, endosulfan and 2,4-D on human peripheral lymphocytes cultured from smokers and nonsmokers. Environmental Toxicology. 2011;26:433-442.
24. Ksenia Cheshenko et al. Interference of endocrine disrupting chemicals with aromatase CYP19 expression or activity, and consequences for reproduction of teleost fish. 2008;155(1):31-62. DOI: 10.1016/j.ygcen.2007.03.005

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