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Immunohistochemical Analysis of P-gp, LC3-II, and Cathepsin-D Associated with Histological Changes in Fish Liver: from the Impacted Environment to Clean Water

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HIGHLIGHTS

- Autophagy and Pg-p in liver from fish can be used as biomarkers of environmental impact.
- In the liver of fish from an impacted environment, autophagy and MXR are noticed.
- Recovery healthy histology liver is possible in animals kept in a good quality water.

Abstract: Herein we evaluated the histopathological alterations and expression patterns of multixenobiotic resistance (MXR) and autophagic proteins in liver samples of fish chronically exposed to anthropogenic contaminants in a highly polluted river, and then again after they had been transferred to good quality water. Two groups were established: euthanized on the day of capture (0 h), and maintained for 30 days in a tank (30 d). The fish of 0 h presented liver with vacuolated and hypertrophic hepatocytes. Also, it was observed strong immunostaining of cathepsin-D, LC3-II and P-gp. Necrosis and apoptosis were also observed throughout the liver. Conversely, the second group (30 d) showed recovery of the liver normal histology and weak immunoreaction of the studied proteins. So, our results indicated that there was a hepatic recovery in the fish kept in good quality water, as showed by the decreased expression of cathepsin-D, LC3-II, and the MXR (P-gp). Therefore, the alterations here observed could be proposed as potential biomarkers to be tested for following the impacts of remediation or mitigation measures to environmental impacts.

Keywords: fish; biomonitoring; ecotoxicology; freshwater.

INTRODUCTION

Urbanization and industrialization have been the main sources of contamination of water bodies contributing to the degradation of the environment, exposing the aquatics organisms to a complex mixture
of chemical pollutants [1,2]. Nowadays, it is emerging the notion that even though the individual contaminants are present at concentrations lower than the No Observable Effect Concentration (NOEC), the toxic effects may be detected, and this phenomenon is known as cocktail effects [3].

The tolerance of aquatic species to toxic agents is highly variable. These agents may generate morphological and biochemical changes in aquatic organisms that are indicative of their environmental conditions [4]. In fish, the liver has become an attractive organ for ecotoxicology studies because it has the potential for biotransformation and excretion of xenobiotics, once it has been in direct contact with environmental pollutants absorbed by an animal [5]. Under stress conditions, the liver of fish exhibits morphological and biochemical modifications, which can be used as biomarkers of environmental impact. Several pathways can be involved in these alterations as phase I and II detoxification enzymes, but some still, not were fully assessed and understood.

Currently, autophagy is not only a route of degradation of cellular components, but it has also been associated with environmental impact conditions in various aquatic organisms [6-9]. Unfavorable conditions trigger autophagy via several cellular stressors, including oxidative stress and DNA damage. The microtubule-associated protein light chain 3 II (LC3-II), a structural component during autophagosome formation, and the Cathepsin D, a lysosomal protease have been used as biomarkers of the autophagic process [6]. In this context, autophagy would act as a mechanism of cellular protection against degraded proteins, xenobiotics, damaged organelles, and other structures that may be toxic to cells, restoring cellular homeostasis [10].

Likewise, other processes related to cell protection and responses to multixenobiotics have been investigated with an emphasis on the expression of P-glycoprotein (P-gp) [11-14]. P-gp is a transmembrane carrier glycoprotein present in several aquatic organisms, also known as ABCB1. It is a representative of the superfamily of ATP-binding cassette transporters. P-gp binds to a variety of substrates and facilitates their efflux, preventing the intracellular accumulation of xenobiotics, and thereby causing resistance to them [15].

The armored catfishes of the genus Hypostomus, belonging to the family Loricariidae, are benthic fish with a wide geographic distribution throughout most of the Neotropical regions of Costa Rica to Argentina [16,17]. The species Hypostomus francisci is widely distributed in the São Francisco basin and frequently captured on the Itapecerica River, a tributary of the Pará River [18]. Because of its constant association with river sediments, this species is in direct contact with several pollutants that may accumulate in this substrate. This characteristic habitat makes this species a good model for environmental impact studies [2]. The aim of the present study was to investigate the histopathology and expression of P-gp and autophagy pathway proteins in the livers of H. francisci captured in the urban river, which has chronic anthropogenic influences. The study also aimed to evaluate the recovery of the healthy structural organization of their livers after fish were maintained in good quality water.

MATERIAL AND METHODS

Experimental design

For the analysis of liver histology and the monitoring of liver recovery after 30 days, 17 adult specimens of H. francisci were captured in the urban area of the Itapecerica River in Divinópolis (20°13′09″ S; 44°54′51″ W), Brazil in December 2015, rainy season in Southeastern, Brazil. A voucher specimens were deposited at the Department of Zoology, access number MHN-UFMG 1456. The water quality of this river was historically monitored in series (1997-2012) where it was evidenced by several thermotolerant coliforms (values between 1.0E+04 and 1.0E+05 mgL⁻¹), oxygen dissolved range 6.0 and 8.0 mgL⁻¹, phosphorus 0.1 and 0.2 mgL⁻¹, and ammonium nitrogen 1.0 and 2.0 mgL⁻¹, respectively, being classified as Water Quality Index (WQI) between medium and poor [19]. This generalized pollution can be linked to various organic and industrial effluents present on this river, as a result of poorly planned municipal sewage systems. In addition, the water of the Itapecerica river has already been evidenced as a cause of inflammatory processes, cellular stress, cell death and several morpho-histological alterations in fish gills [2]. The fish were caught in the early morning using different fishing gear (gill nets, casting nets, and trawls), and then quickly transported to the laboratory in plastic tanks supplied with air. In the lab, the fish were divided into two groups. The first group (N = 9) was euthanized with 250 mgL⁻¹ benzocaine on the day of capture and dissection was performed. The second group (N = 7) was housed for 30 days in a 150 liters tank with tap water dechlorinated, temperature at 25 °C, constant oxygenation, pH 6.8, and photoperiod 14h/10h. The fish were fed daily with commercial feed for 30 days until euthanasia with 250 mgL⁻¹ benzocaine. From both groups of fish, length (L) and weight (W) were obtained and used to calculated the
length-weight relationship (LWR). The parameters a and b were estimated by log-transformed weight and length measurements (logW = log a + log L^b) [20]. All the procedures were conducted in accordance with the recommendations of the Ethics Committee on Animal Use in Research of Federal University of São João Del Rei, nº 49/2010.

Histology and immunohistochemistry

For the histological analysis, liver fragments from a similar region in all fish were collected and fixed in Bouin liquid for 8 h at room temperature. Next, the samples were submitted to paraffin inclusion (one block per fish), sectioned with 5 μm thickness and the histological sections were stained with hematoxylin and eosin (HE) or periodic acid-Schiff reagent (PAS). For immunohistochemistry, histological sections from eight fish (n = 4/group) were submitted to the identification the following primary antibodies: Cathepsin D - CAT-D (C-20) [21] and Microtubule-associated protein light chain 3 II - LC3-II (N-20) [22] from Santa Cruz Biotechnology and P-glycoprotein - P-gp (C219) [12] from Thermo Fisher Scientific. Briefly, the liver sections were deparaffinized and hydrated, and antigen retrieval was performed with 10 mM sodium citrate buffer for 20 min at 100 °C. Next, blocking the endogenous peroxidase with 3% hydrogen peroxide was performed. Blocking of non-specific antigens was done incubating the sections in 2% BSA buffer solution for 30 min. Then, the sections were then incubated with the primary antibody (anti-CAT-D, 1:200; anti-LC3-II, 1:200 or anti-P-gp 1:100) overnight at 4 °C in a humid chamber. For identification of primary antibodies, the LSAB 2 System HRP Kit from Dako Cytomation (K 0675) containing secondary antibody conjugated to biotin (1:200) and streptavidin conjugated with peroxidase was used. Thus, after washing with PBS the sections were incubated with biotinylated secondary antibody for 45 min in a dark chamber. Then a solution of streptavidin conjugated to peroxidase was also applied in slides for 45 min in a dark chamber. The reaction development (primary-secondary antibody-streptavidin-peroxidase) was performed with diaminobenzidine and counterstained with hematoxylin. For negative control, one of the sections do not received primary antibody. The immunoreaction was evaluated by a visual scoring determination in each sample and classified as 0, 1+, 2+, or 3+ [23]. The evaluation was performed by two independent researchers. In the case of a disagreement, discussion was used to reach a consensus.

For morphometry, liver images were captured in a Motic photomicroscope (BA 310) and morphometric analyzes were performed using the Motic Plus 2.0. Thus, five photomicrographs of the liver were consecutively captured on the 40 X objective (N = 3 animals per group). In each micrograph, the area of 20 hepatocytes were measured (N = 300 hepatocyte per experimental group).

The degree of tissue injury (Degree of Liver Damage - DLD) based on the severity of lesions was used to semiquantify the histological alterations in liver sections stained by HE. The method was adapted from Poleksic and Mitrovic-Tundzic [24]. The histopathological findings were classified in progressive tissue damage stages: stage I (SI), which do not alter the normal functioning of the tissue (vacuolated cytoplasm, cellular or nuclear hypertrophy, lateral displacement of the nucleus, glycoegen present); stage II (SII), which are more severe and impair the normal functioning of the tissue (hepatocyte apoptosis, fibrosis, degeneration of cytoplasm or nucleus, rectilneation of vessels and parenchyma); and stage III (SIII) alterations, which are very severe (focal necrosis, tumor). A value of DLD was calculated for each fish by the formula: DLD = (1 X SI) + (10 X SII) + (100 X SIII). DLD values between 0 and 10 indicate normal functioning of the organ; values between 11 and 20 indicate slight damage to the organ; values between 21 and 50 indicate moderate changes in the organ; values between 50 and 100 indicate severe lesions and values above 100 indicate irreversible damage to the organ [24]. Apoptosis was characterized morphologically using the following parameters: nucleus with compacted chromatin at the periphery of the cellular envelope, cellular retraction and formation of apoptotic bodies. Necrosis was hallmark by pyknosis and fragmentation nuclear, cellular membrane rupture and karyolysis.

Statistical analysis

The Shapiro-Wilk test of normality was applied to verify whether the samples present a normal distribution. Once the data failed the Shapiro-Wilk test of normality, the non-parametric Mann-Whitney test was performed, followed by the Dunn’s post hoc test when a significant statistical difference was found. All data were tested using 95% confidence intervals (p < 0.05). All statistical tests were performed using software SigmaPlot (Systat Software Inc.).
RESULTS

Animals

Among the *H. francisci* caught 9 females and 7 males. The biometric data was summarized in the Table 1. The LWR was estimated for *H. francisci* from first group (0 h): parameter b was 1.9227, and parameter a was 0.5445 with a determination coefficient ($r^2$) equal to 0.40, and in the second group (30 d) the parameter b was 2.0303, and parameter a was 0.7589 ($r^2 = 0.98$). The groups were formed by fish with similar size. In addition, the parameter b showed a tendency to increase after 30 days in good water.

Table 1. Biometric data and parameters of the length-weight relationship ($a$ and $b$) for *H. francisci* from first group ($N = 9$) euthanized on the day of capture (0h) and the second group ($N = 7$) euthanized after 30 days in good water (30 d).

| Biometric Data | 0 h ($N = 9$) | 30 d ($N = 7$) |
|----------------|--------------|---------------|
| Lenght (cm)    | 21.0 (max 25.0 - min 17.0) | 22.5 (max 26.0 - min 18.5) |
| Weight (g)     | 100.0 (max 180.0 - min 50.0) | 94.0 (max 132.0 - min 64.0) |
| $a$            | 0.5445       | 0.7589        |
| $b$            | 1.9227       | 2.0303        |

Histology and immunohistochemistry

In both groups of fish, the liver was surrounded by a thin capsule of connective tissue. The hepatic parenchyma was observed with hepatocyte plates bounded by sinusoids. Each plate was formed by polarized hepatocytes with a sinusoidal and biliary face. The hepatocytes were distributed in cord form and arranged in plates, often two hepatocytes thick. In addition, it was possible to observe a biliary system (Figure 1).

In the specimens of group 1 (0 h), the hepatocytes were hypertophics, the nuclei were displaced to the periphery, and abundant cytoplasmic vacuoles were present (Figure 1a and b). Apoptosis in hepatocytes was very frequent (Figure 1b). In addition to apoptosis, necrosis was evident, in several areas of the hepatic parenchyma, being characterized by an area with intense acidophilic properties (Figure 1b). The PAS histochemical in group 1 revealed weak labeling of PAS-positive compounds in the cytoplasm of hepatocytes, indicating few reserves of glycogen (Figure 1c).

In the group 2 specimens (30 days), the liver showed smaller hepatocytes compared with the first group. The vacuolization of the hepatocytes was almost absent. Sinusoid capillaries were less dilated (Figure 1d). Most hepatocytes had central nuclei and eosinophilic cytoplasm (Figure 1d and e). In the hepatic parenchyma, no necrosis areas were registered. The PAS assay showed strong staining in the cytoplasm of hepatocytes, indicating large cytoplasmic glycogen deposits in these cells (Figure 1f).
Figure 1. Histological sections from livers of animals Group 1 0 h (a-c) and Group 2 30 d (d-f). a-b and d-e the sections were stained with Hematoxylin and Eosin (HE); c and f the liver was stained with PAS assay. White (nuclei displaced to the periphery) and black (abundant cytoplasmic vacuolization) arrows. N necrosis area; BV blood vessels; S sinusoid capillaries; asterisk bile duct; white (apoptosis) and black (central nuclei) arrowheads; brackets (hepatocyte plates). Note in c and f the absent and present of glycogen deposits in cytoplasm from hepatocytes, respectively. Bars = 25 µm. Insert shows hepatocytes in apoptosis.

Morphological data showed that in group 1 (0 h), hepatocytes presented a mean area of 88.30 ± 21.31 µm². In contrast, in group 2 (30 days), the hepatocytes were smaller with a mean area of 47.63 ± 11.58 µm² (p < 0.001).

The mean DLD for liver from fish of the first group (0 h) was 54.11, indicating severe damage to the organ. The DTC value was lower in the fish from the second group (30 d) (DTC = 6.42).

In group 1 (0 h), the labeling of P-gp was observed in the plasma membrane of hepatocytes, being more intense in cells close to large blood vessels. The immunoreaction decreased with increasing distance from large vessels (Figure 2a and c). However, in group 2 (30 days), the staining was less frequently restricted to hepatocytes near the large blood vessels (Figure 2b).

In group 1 (0 h), cathepsin-D (Figure 2d and f) and LC3-II (Figure 2g and i) were labeled in hepatocyte cytoplasm with more intensity in cells with cytoplasmic vacuoles. Contrarily, in group 2 (30 days), the immunostaining these proteins was also cytoplasmic, however with less intensity (Figure 2e and h).
Figure 2. Immunohistochemical staining of Pg-P, LC3-II, and Cathepsin-D in the liver of the *H. francisci* from 0 h and 30 d. (a and d) Pg-p reaction in the cytoplasmic membrane, (b and e) LC3-II reaction in cytoplasmic points (see insert in b), (c and f) Cathepsin-D (cat-D) reaction dispersed in the cytoplasm (see insert in c). Positive reactions was detected by intense brown colour. Bars = 25 µm.

Based on the visual evaluation, all proteins evaluated presented an overexpression (score +3) in the fish recently caught in the Itapecerica River, and no or weak expression (score 0, 1+, or 2+) was observed in fish held in clean water (Figure 3).

**DISCUSSION**

In the present study, we observed the recovery of a healthy structural organization in the livers of fish which had been maintained in favorable environmental conditions. In healthy fish, the hepatic parenchyma is organized into hepatocytes arranged in cords, surrounded by sinusoid capillaries, converging toward a vein. The hepatocytes were regulars, with central nucleus, cytoplasm without vacuolization, and often filled with glycogen. The bile ducts have a simple cubic epithelium, which can become columnar in large ducts. The stroma is composed of thin connective tissue, associated with veins, sinusoids and the biliary system.
Increased hepatocyte vacuolization [28], necrosis -y shows the functional activity of this cell type in -n of Cat -r energy demand -es to the ability of cells -y be related by Smital and coauthors [42] who observed that the MXR mechanism -t of -c heir expression. Finally, the recovery of healthy morphology of the liver observed at the 30 days might be the result of a process related to cell proliferation and tissue remodeling.

There is a correlation between the level of pollution identified and the expression of this glycoprotein; therefore, MXR can be considered a generalized biomarker of aquatic contamination [14]. In this study, the results showed that intense labeling in group 1 indicated a high concentration of pollutants in the environment. This result is reinforced by the decreased labeling in group 2, which maintained a basal level of expression. This was also noted by Smital and coauthors [42] who observed that the MXR mechanism can return to its basal level of expression when exposure to contaminants ceases.

In summary, we believe that once a xenobiotic stimulus has been interrupted, the liver of armored catfish showed a return to the healthy morphology. Thus, after a chronic exposition of pollutants, the MXR, herein represented by the P-gp, it was not able to avoid the intense stimulation inside the cell being a trigger of the autophagic pathway (higher expression of Cat-D and LC3-II), that is an attempt to keep cellular viability, and the liver working for the fish survival. Once the stimulus of the generalized pollution was removed, all of the aforementioned pathways stopped to be activated, therefore reducing their expression. Finally, the recovery of healthy morphology of the liver observed at the 30 days might be the result of a process related to cell proliferation and tissue remodeling.

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