Chronic Effects of Methylmercury on *Astronotus ocellatus*, an Amazonian Fish Species

**Abstract**

**Background:** Bioassays are very useful tools to establish cause-effect relationships as they do not allow other confounding factors to interfere within the results. Methylmercury is known as a highly toxic substance and bioassays have demonstrated several effects of this pollutant on different fish species, especially on reproduction and hematology. The aim of this study was to evaluate the effects of chronic methylmercury exposure on *Astronotus ocellatus* biomarkers (hematology, acetylcholinesterase activity, micronucleus frequency, histopathology of liver and gonads), as consequence of methylmercury exposure through artificially contaminated food (10 μg.g⁻¹).

**Methods and findings:** The experiment was performed during 6 months. Blood samples were collected bimonthly for hematology analysis and total mercury determination. In the end of the experiment, samples of different tissues were collected for total mercury analysis and effect biomarkers assessment. Mercury bioaccumulation was clearly observed in all fish tissues (10² higher on test group). Methylmercury exposure caused hematological damages, especially in corpuscular volume and can activate leukocytes production and possibly influences the acetylcholinesterase activity. The frequency of genotoxic effects (bilobed nuclei) and damage on liver and gonad tissues were also increased.

**Conclusion:** Methylmercury exposure caused, decrease in erythrocytes number, corpuscular volume and in monocytes frequency; increase in hepatocytes volume, degeneration, hypertrophy and hyperplasia of follicular layer, follicular atresia, yolk degeneration and presence of proteic material and inflammatory infiltrate on liver and gonads, genotoxic effects were not clear and acetylcholinesterase activity seemed to be higher after exposure.

**Keywords:** Bioassays; Hematology; Histopathology; Micronucleus; Methylmercury; Tropical fish

**Introduction**

Several effects are described for organisms exposed to pollutants in water, air and soil [1-3]. However, studies performed under controlled conditions, including the absence of any external variable interfering in the occurrence of any kind of effect on organisms are useful for understanding the direct effect of a contaminant. Such studies allow establishing cause effect relationships and avoid masking the action of other toxic substances. Bioassays are very useful tools to establish cause effect relationships as they do not allow other confounding factors, such as other pollutants, physical-chemical changes, to interfere within the results [4]. Besides the lethal concentration (LC₅₀) or dose (LD₅₀) for 50% of the tested population, other endpoints can be assessed, especially to investigate non-lethal effects. In this context, biomarkers could be applied, since they...
are, by definition, indicators of exposure to xenobiotics and of effects related to this exposure [5].

Among the persistent pollutants, mercury (Hg) is one of the most toxic, according to World Health Organization (WHO 1990), especially when it is present in the environment as methylmercury (MeHg), a neurotoxic and teratogenic substance to humans. Methylmercury bioaccumulates and biomagnifies in aquatic food webs, reaching high concentrations in top predators. For this reason, the main pathway of MeHg exposure to humans, aquatic mammals, marine reptiles and sea birds is the consumption of contaminated fish. Because of that, fish are commonly used as bioindicators on studies to evaluate mercury contamination [6].

Berntssen et al. noted significant changes in fish hematology when performing bioassays using Salmo salar fed with MeHg enriched food above 5 µg.g⁻¹ [7]. Concentrations above 5 µg.g⁻¹ are reported for carnivorous fish from artisanal and small-scale gold mining areas in Brazilian Amazon, being 5 times higher than the WHO recommended guideline for human consumption of carnivorous fish species (1 µg.g⁻¹) [8, 9]. Laboratory assays also showed the relationship between MeHg exposure and lower success on fish reproduction, due to the reduction on gonad development, including lower gonadosomatic indexes, increase on apoptosis of follicles and inhibition of testosterone production by males and estradiol by females, decreasing the number of eggs [10-13]. MeHg concentrations in eggs are extremely high, varying with the food items chosen by females [14]. Also, it’s reported on literature that MeHg can cause decrease spawning success, besides the reducing on eggs incubation success and survivor during development stages of embryos [11, 15-18]. All these factors have large effects on reproductive success, showing that methylmercury has multiple important impacts on reproduction.

Acetylcholinesterase is an enzyme that hydrolyzes acetylcholine molecules and it is an important regulatory enzyme that controls the transmission of nerve impulses across cholinergic synapses. Its inhibition has been commonly linked to organophosphorades, carbamates and other pesticides exposure, but some studies observed its inhibition due to mercury exposure [1, 19-27]. Jesus et al. when considering neotropical fish species, evaluated possible effects on acetylcholinesterase activity and internal distribution of mercury in a carnivorous species (Hoplias malabaricus), due to intreperitonial injections of mercury and methylmercury chlorides [28, 29]. Costa et al. observed an inhibition of this enzyme on Hoplias malabaricus, using a methylmercury exposure dose of 75 ng g⁻¹, in laboratory conditions, after 70 days of exposure to this contaminant by contaminated food ingestion (the prey were artificially contaminated with intreperitoneal injections) [30].

Thus, the objective of this study was to evaluate the effects of chronic methylmercury exposure on Astronotus ocellatus bioanalyzers (hematology, acetylcholinesterase activity, micrornucleus frequency, histopathology of liver and gonads), as consequence of methylmercury exposure through artificially contaminated food (10 µg.g⁻¹). This is a well-distributed omnivorous/carnivorous species in Amazon Region and outside this region, due to artificial introductions, it is territorialist, not performing seasonal migration, and it is consumed by humans. We expect to observe effects of experimental chronic methylmercury exposure on micronuclei frequency, liver lesions, anemia and induction of defense system.

Methods

Astronotus ocellatus (Agassiz, 1831)

The oscar (Astronotus ocellatus) is an Amazon Basin teleost fish species from Cichlidae family and it is considered a popular aquarium fish around the world. Also, it is an edible fish and is commonly consumed by Brazilians from North and Middle West Regions. This species reaches 25-30 cm of total length and weighs approximately 1.5 Kg in wildlife and tolerates waters with pH from 6.0 to 8.0 and temperatures from 22 to 25°C. It’s an omnivorous/carnivorous species and includes in its diet zoobenthos (crayfish), zooplankton (insects larvae), fish (finfish) and plants (benthic algae, fruits, flowers and seeds from terrestrial plants). It was chosen for this first experiment because (i) it is easy to maintain in aquarium using a relative small volume of water, (ii) its food habit and (iii) previous studies at Tapajós River, Amazon Basin, noted hematological alterations on tucunarés (Cichla sp.) from artisanal and small-scale gold mining (ASSGM) areas, due to mercury exposure and tucunarés and oscars are from the same taxonomic family [31, 32].

Bioassays

The experiment lasted 180 days (six months), being the test group fed on fish food (commercial pellets; 40% of protein) artificially contaminated with methylmercury chloride solution (10 µg.g⁻¹). This concentration was defined as the first to be tested according to the results described on Hammerschmitd et al. and Berntssen et al., who indicated that concentrations above 5 µg.g⁻¹ were able to induce hematological disturbances on fish [7, 11].

The artificial contamination of the pellets followed the methodology described at Hammerschmitd et al. [11]. The food was stored in sealed plastic bags (Ziploc) and they were locked in sealed flasks and frozen. The total mercury concentrations in pellets were determined before and after the contamination procedure, resulting, respectively, in 0.020 (minimum)-0.037 (maximum) µg.g⁻¹ and 9.0 ± 0.5 µg.g⁻¹ (average ± standard deviation).

For this first study, eight specimens of A. ocellatus for test group and three specimens for control group (fed with the same commercial pellets, but without artificial enrichment on MeHg) were used. The number of specimens was not equal for both groups due to a space restriction at the laboratory, where the isolated room for the experiment did not have sufficient area to include more aquariums. This way, the maximum capacity of the room was for 11 aquariums and we chose to use a higher number of specimens on test group. These specimens were of both sex and were acquired from the same commercial breeder, with 4.0 cm of total length initially. They were maintained in individual aquariums of 50 L, with constant filtration and aeration and fed twice per day. The experiment began after six months of acclimatizing, in order to have sufficient weight and length (above
15 cm) to support healthily the blood collection, bimonthly. The aquariums were submitted to partial changes of water monthly. pH was measured once a week and the temperature was noted daily. Any observation related to behavior changes was recorded in the experiment booklet. Total mercury concentrations in water were monitored each 15 days and the contents were below the detection limit (5 ng.l\textsuperscript{-1}) during all the experiment.

**Sampling**

Four blood sampling were performed during the experiment. The first sampling was on day 0, before any kind of exposure to methylmercury chloride. Once the experiment started, blood samples were collected bimonthly, where sampling 2 was carried out after two months of exposure of the test group, sampling 3, after four months and sampling 4, after six months, being the last day of experiment. For every sampling, biometry parameters were measured and hemogram, micronucleus frequency and total mercury in blood samples were determined. The other biomarkers were measure only on samples from the sampling number 4 (the last one), where different tissues were collected for total mercury determination (erythrocytes, plasma, muscles, liver, gonads), for acetylcholinesterase activity (muscles) and for histopathology (liver and gonads) (Figure 1).

Before every blood sampling, the volume that could be collected without causing any health problems to fish was calculated by using as base the body weight of each specimen, being equivalent to a maximum of 1% of body weight [33]. The blood was obtained by caudal puncture, using 1 ml syringes and needles caliber 20 × 0.55 mm, containing EDTA [34]. The samples were stored in 1.5 ml micro tubes and refrigerated until analysis. In order to minimizing stress during blood sampling, the specimens were calmly captured. Eyes (head) and body were covered with a wet towel. At the sampling days, fish was not fed, with 12 h of fasting. Thus, mercury concentrations in blood could be considered as the basal concentration on this biological matrix.

At the end of the experiment, the specimens were euthanized. Before this procedure, lidocaine was directly sprinkled in the gills of the specimens. After few minutes, if immobility and rigor mortis were observed, the specimens were dissected to separate muscles, liver and gonads samples. Every effort was made to minimize animal suffering, according to ethic code for research with animals (Ethic Committee from Fluminense Federal University approved this experiment, with protocol number 89/2011).

**Blood analysis**

Hemogram followed the methodology described by Almosny & Santos, where all cellular types are present in a Newbauer (Improved) chamber, being proceeded erythrocytes, thrombocytes and leukocytes counting together [33, 35]. Corpuscular volume (CV) was determined using capillary tubes submitted to centrifugation (12,000 rpm/5 min). The mean corpuscular volume (MCV) was calculated as follow: CV × 10/ Erythrocyte counting. Total plasmatic protein (TPP) was measure

![Figure 1](image_url)
using a portable refractometer (BIOBRIX, model 301). The specific leukocytes counting was performed by using blood smears, fixed with methanol and stained in GIEMSA during 30 min. The identification of leukocytes was held in optical microscopy, using objective lens (1,000x) to estimate the frequency of each leukocyte type in 100 leukocytes, resulting in a percentage value.

**Micronucleus frequency**

The frequency of micronucleus was conducted on the same blood smears used for specific leukocytes counting. In optical microscopy, using the same objective lens (1,000x), the occurrence of micronucleus and bilobed nucleus in 1,000 erythrocytes for each specimen was counted. It was classified as micronucleus, every small fragment, non-refractory, with the same color and shape of the cell nucleus, from intact cells and as bilobed nucleus, every erythrocyte with nucleus in shape of beans.

**Total mercury determination**

Wet samples of muscles, liver and gonads were homogenized and the determinations were performed in triplicates. For each replicate, around 0.03 g of tissue was used. The blood samples were centrifuged (3,000 rpm/10 min) for separation of erythrocytes and other cells from plasma. The mercury determination on erythrocytes and on plasma was conducted in duplicates, because there was not sufficient sample mass. Total mercury was measured using a portable atomic absorption (LUMEX), specific for total mercury analysis by spectrophotometer. For this methodology, the samples were only homogenized. None previous treatment was necessary, since this equipment works with pyrolysis, where the samples were submitted to high temperatures (above 800°C) in a pyrolysis chamber, volatilizing mercury (all chemical forms) for its detection. The results are expressed as wet weight.

For accuracy evaluation, certified material (IAEA 407, mercury concentration=222 ng.g⁻¹) analysis were performed every day, with an acceptable error of 10% (maximum). The detection limit was 5 ng.g⁻¹. The mean value found for the certified material was 218.4 ± 14.3 ng.g⁻¹ (n=90), resulting in an error of 1.8%. During international intercalibration of IAEA (International Atomic Energy Agency), the error was below 10%.

**Acetylcholinesterase activities**

The determinations of the AChE activity in fish muscles followed the method described by Oliveira Silva et al. with modifications, whose quantifications were based in the reaction described by Ellman et al. [36]. Small portions of muscle samples were weighted and homogenized in sodium phosphate 0.12 M, pH 7.6 (6:1). The samples were centrifuged (9,000 g, 20 min, 8°C). While the samples were centrifuging, test tubes were prepared containing 2 ml of sodium phosphate and 0.5 ml of DTNB 2 mm. At the time of lecture, 500 μl of acetylcholine and 25 μl of sample were added into those previously prepared tubes. Enzymatic activities were determined in spectrophotometer in kinetic form (λ=412 nm), being obtained after two minutes of reaction, the absorbance per minute. For protein determination, muscles samples were diluted (1:10) in sodium phosphate solution 0.12 M pH 7.6. In test tubes were added 4.3 ml of distilled H₂O, 200 μl of NaOH 25% and 200 μl of diluted sample. The blank contained 4.5 ml of H₂O distilled and 200 μl of NaOH 25%. In the first tube (blank) was added 300 μl of Folin reagent, being homogenized on a vortex during 30 s. The same process of homogenization was made to all tubes. After 5 min, the absorbance was measured in photometric model, in a spectrophotometer Shimadzu UV 1601 (λ=660 nm). The absorbance were converted to protein concentration (mg.ml⁻¹) using an albumin curve. The specific activity of AChE (μmoles.min⁻¹.mg⁻¹ protein) was obtained by ratio of the enzyme activity and the protein concentration [37].

**Histological analysis**

Gonads and liver samples were obtained immediately after the specimen’s death and preserved on a solution of formaldehyde 10% buffered with sodium phosphate. After formaldehyde impregnation, the samples were carefully cut with sharp blades and positioned in the cassette. Next, the cassettes containing fragments of the samples were prepared for histological sections, according to Vazzoler [38]. All the prepared slides were observed on optical microscopy and the presence or the absence of any alteration was recorded for liver (presence of macro vacuoles, degeneration and necrosis, decrease on hepatocytes number, hyperplasia, inflammatory infiltration, hemorrhage, adiposities) and for gonads (atresic follicles, degeneration, hypertrophy, hyperplasia, inflammatory infiltration, primary and secondary follicles number, vacuolization, fibrosis).

**Statistical analysis**

The statistical analysis was performed using the software STATISTICA 8. Correlations between mercury concentrations and growth/effect biomarkers were tested using Spearman Test. Mann-Whitney U Test was applied to verify differences among test and control groups. The significance level accepted was at minimum p<0.05. For dependent samples variance evaluation, it was applied the Wilcoxon matched pairs test and the Friedman ANOVA and Kendall coefficient of concordance for multiple dependent samples. However its interpretation deserves attention, due to the low number of specimens used for this first experiment.

**Results**

The mean values of pH and water temperature of aquariums during the experiment period were, respectively, 6.2 ± 0.2 and 25 ± 2°C. During the experiment, 4 specimens died. At sampling T0 and T1, all specimens were alive. Between T1 and T2, we had the first three losses, one for control group and two for test group. The specimen from control group avoided food during more than three days and no clue of any kind of fish disease was observed during this period. Of the two deaths of test group, one specimen escaped from the aquarium during the night and the other one, there was a mechanical fail on the water pumping during the night and the water in the aquarium had no more circulation, decreasing dissolved oxygen. And finally between T2 and T3 samplings, we have lost one more specimen from test group.
Therefore, the results for each sampling correspond to this number of specimens: T0 and T1=3 specimens on control group and 8 on test group, T2=2 specimens on control group and 6 on test group, T3=2 specimens on control group and 5 on test group.

All the specimens have grown similarly (both control and test groups), indicating, initially, no influence of MeHg exposure on growth rates. Although, as we can see on Figure 2, there was a tendency of 10-20% of difference among control and test groups for weight and length, where control group showed slightly bigger specimens (Figure 2).

**Mercury concentrations**

The test group demonstrated a huge increase on mercury concentrations in blood at samplings T1 (Rank Sum: T0=36/T1=84, U: 0.0, Z: -3.24, p<0.005, valid cases: T0=8/T1=7), T2 (Rank Sum: T0=36/T2=69, U: 0.0, Z: -3.09, p<0.005, valid cases: T0=8/T2=6) and T3 (Rank Sum: T0=36/T1=4; U: 0.0, Z: -2.72, p<0.01, valid cases: T0=8/T1=4), when compared to the first sampling, with values 102 higher than control group (Rank Sum: Test Group=38/Control Group=28, U: 2.0, Z: -2.04, p<0.05, valid cases: Test Group=8/Control Group=3). Control group did not show variation on mercury concentrations in blood during the experiment, being similar to T0. The results found in sampling T3 are shown in Table 1. All the analyzed tissues showed twice higher mercury concentrations than control group. Positive correlations were observed between mercury concentrations in erythrocytes and in muscles (0.83, p<0.05/0.9, p<0.05, respectively) and liver (0.77, p<0.05). The same way, it was observed correlation between mercury concentrations in liver and in muscles (0.96, p<0.001). These results suggest equilibrium on mercury distribution among fish tissues (Figure 3 and Table 1).

**Hematology**

There was a decrease on erythrocytes counting in the first two months of experiment for control group (Rank Sum: T0=15/T1=6, U: 0.0, Z: 1.96, p<0.05, valid cases: T0=3/T1=3). This decrease was also observed for test group, but during all the experiment, in other words, all the erythrocytes counting after the beginning of exposure were significantly different from T0 (Rank Sum: T0=90/T1=30, U: 2.0, Z: 3.01, p<0.005, valid cases: T0=8/T1=7) (Rank Sum: T0=82/T2=23, U: 2.0, Z: 2.84, p<0.005, valid cases:

**Figure 2** Weight and total length (± standard deviation) of *Astronotus ocellatus* during the experiment.

**Figure 3** Total Mercury in blood (± standard deviation, ng.g⁻¹, wet weight) in *Astronotus ocellatus* exposed to methylmercury by food ingestion during the experiment.
The haematocrit slightly decreased over the experiment on control group. Concerning differences between control and test groups, MCV was higher for specimens from test group (Rank Sum: Test Group=33/Control Group=3, U: 0.0, Z: 2.00, p<0.05, valid cases: Test=6/Control=2). This tendency can be corroborated with the found correlations (using data from the four samplings) between mercury concentrations in blood and erythrocytes counting (-0.48, p<0.01, n=29) and MCV (0.56, p<0.001, n=29), indicating that the exposure to MeHg could affect the hematological parameters.

**Table 1** Total mercury concentrations (wet weight, ng.g⁻¹) in erythrocytes, plasma, muscles, liver and gonads of *Astronotus ocellatus* fed with artificially contaminated food with methylmercury (10 µg.g⁻¹) after six months.

<table>
<thead>
<tr>
<th></th>
<th>Control Group</th>
<th>Test Group</th>
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<tbody>
<tr>
<td><strong>Total Mercury (ng/g)</strong></td>
<td></td>
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</tr>
<tr>
<td>Muscles</td>
<td>39.5 ± 31.8 (2)</td>
<td>6,368.7 ± 1,089.3 (4)</td>
</tr>
<tr>
<td>Erythrocytes</td>
<td>9.5 ± 6.4 (2)</td>
<td>1,405.0 ± 121.2 (4)</td>
</tr>
<tr>
<td>Plasma</td>
<td>20.0 (1)</td>
<td>1,150.0 ± 139.0 (4)</td>
</tr>
<tr>
<td>Liver</td>
<td>23.2 ± 3.9 (2)</td>
<td>7,845.8 ± 439.6 (3)</td>
</tr>
<tr>
<td>Gonads</td>
<td>32.0 (1)</td>
<td>4,199.2 ± 1,504.0 (3)</td>
</tr>
</tbody>
</table>

(n)=number of specimens collected at the end of the experiment.

Figure 4  Hemogram and specific leukocytes counting (± standard deviation) of *Astronotus ocellatus* exposed to methylmercury by food ingestion during the experiment.
decrease erythrocytes counting and increase MCV, configuring a non-regenerative anemia scenario (Figure 4).

Furthermore higher frequencies of lymphocytes (Rank Sum: Test=39/Control=82, U: 2.0, Z: -3.01, p<0.005, valid cases: Test=8/Control=7) and monocytes (Rank Sum: Test=47/Control=73, U: 11.0, Z: -1.96, p<0.05, valid cases: Test=8/Control=7) on test group were observed after two months of exposure. Correlations between the frequencies of eosinophil (-0.41, p<0.05, n=29) and monocytes (0.49, p<0.01, n=29) and mercury concentrations were found and could suggest a possible chronic mononcytosis scenario and an increase on immunological reaction (Figure 4).

It’s also interesting to observe that, apparently at the sixth month of experiment, monocytes and eosinophils showed percentages similar to T0, but lymphocytes continued altered. And at T3, neutrophil started to decrease (Rank Sum: T0=52/T3=3, U: 0.0, Z: 2.09, p<0.05, valid cases: T0=8/T3=2).

**Micronucleus frequency**

Micronucleus frequency was basically none for all specimens along the experiment, except for T0, where it was found 1 micronucleus in 8,000 counted cells from test group, and for T2, where it was found 1 micronucleus in 2,000 counted erythrocytes from control group. Bilobed nucleus frequency was higher in test group that in control group, although it was not significantly different (Rank Sum: Test Group=56.5/Control Group=9.5, U: 3.5, Z: 1.74, p=0.083, valid cases: Test group=8/control group=3). No correlation was found between mercury concentrations on any tissues and this effect biomarker. However, the bilobed nucleus frequency seemed to be interesting for chemical stress detection. May be it is necessary a higher exposure period to induce micronucleus formation in this fish species. The smears prepared during the last sampling were not used to investigate genotoxic effects, because they had no clean fields to performing the counting (Figure 5).

**Acetylcholinesterase activity**

The test group showed higher activities than control group (p<0.005), showing a possible activation of this enzyme after MeHg exposure (Figure 6).

**Histopathology**

It was observed for both test and control groups an increase on hepatocytes volume with vesicles formation, due to fat accumulation on liver. All fish specimens showed hepatocytes degeneration. This excessive fat accumulation could cause the observed degeneration on liver (hepatic lipidosis). For gonads analysis (three specimens from test group: Two females and one male), it was observed degeneration of cells in the male gonad, absence of spermatogonia, presence of young spermatooza. On female gonads, it was observed hypertrophy and hyperplasia of the follicular layer and presence of follicular atresia, yolk degeneration and proteic material between cells, besides inflammatory infiltrate (Figure 7and Table 2).

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**Individual analysis**

As the experiment included few specimens, it’s interesting to describe the results found for biometry, mercury in blood, hematology and bilobed nuclei frequency along the six months of exposure. These biomarkers were chosen for this individual assessment, because they were measured throughout the experiment in all samplings (T0, T1, T2 and T3). Although micronuclei were also measured in all samplings, it was not observed an important variation during the experiment for all fish specimens. This analysis was only performed for the specimens that survived all the experiment (two from control group and six from test group) and for one specimen from test group that survived until the third sampling (Table 3).

From control group, the specimen named C01 gained weight linearly throughout the experiment (62.0 g). There was no clear relationship between mercury concentrations in blood and biometry. Also mercury concentrations in blood and in erythrocytes were 9 ng.g⁻¹ and 24.5 ng.g⁻¹, respectively, lower at the end of the experiment. Hematimetry decreased almost four times and CV was twice lower than the first sampling (T0). Consequently, the MCV increased. The total plasmatic protein decreased only 1.0 g.dl⁻¹. There was a decrease on leukocytes number and an oscillation on their specific frequencies, with predominance of lymphocytes at T1, monocytes at T2 and lymphocytes/eosinophil at T3. One nuclear alteration was found
at T1 and T2.

Also from control group, the specimen named C02 gained weight linearly throughout the experiment too (77.7 g). Also no clear relationship between mercury concentrations in blood and biometry and a decreasing on mercury concentrations in blood and in erythrocytes (9 ng.g\(^{-1}\) and 22.0 ng.g\(^{-1}\), respectively) were found. Hematimetry decreased only after six months (T3). Corpuscular volume or hematocrit increased at T1 and stabilized. MCV increased two times at T3. The oscillation on leukocytes frequencies was also observed, but with no monocytes predominance, only an exchange between lymphocytes and eosinophil predominance. And the number of leukocytes decreased, but not so expressive in comparison to C01. Nuclei abnormalities increased at T2.

For test group, the specimen named T03 gained 179.2 g and grew 0.5 cm. Obviously mercury concentrations in blood increased more than 1,600 ng.g\(^{-1}\). An expressive increase on mercury concentrations in erythrocytes was observed, almost four times higher than the increase on plasma. At the first two months, hematimetry had an extreme decrease, but then this counting came back to values very close to T0. So did the MCV, increasing at T1 and then coming back to the blank register (before exposure). Additionally lymphocytes predominated as leukocyte type only after two months of exposure. At T2, appeared one bilobed nucleus in the counting.

The specimen named T04 gained 60.0 g and grew 2.0 cm. No growth was observed between T1 and T2. The relationship between mercury concentrations in blood and length is cleaner. The majority of expressive differences on hematology was observed at T1 (after two months of exposure), where PPT, CV and MCV increased and lymphocytes had higher frequency than eosinophils. Bilobed nuclei were similar, increasing in only one alteration at T2 in comparison to T0.

Fish number T05 gained 55.0 g and grew 0.3 cm. After two months of exposure, hematimetry decreased and CV and MCV increased. Then their values came back to those observed for T0. Between the third and the last sampling, this specimen lost weight; mercury concentrations in erythrocytes and in plasma were quite similar, being the ratio 1.3:1 and TPP increased. In this specimen, no nuclei abnormalities were found at the end of the experiment.

T07 gained 114.8 g and grew 0.5 cm. At the first four months, hematimetry was lower than before exposure. CV had an oscillation, increasing in the first moment (T1) and then decreasing (T2). MCV increased until T2. TPP decreased after four months of exposure (T2). Lymphocytes had higher frequencies than eosinophils until the third sampling (T2). The number of nucleus alterations found at T2 was twice higher than at T0.

T09 gained 50.3 g and grew 2.0 cm. Weight lost was also observed between T2-T3 samplings. Similar to the specimen T05, the ratio between mercury concentrations in erythrocytes and plasma was close to 1 (1.2:1). Oscillation on hematimetry was observed, with an increase at T2 and a decrease at T1 and T3, in
**Table 3** Oscillations (gains and losses) on biomarkers results observed for each specimen after six months of experiment (results before exposure subtracted from those after exposure).

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Weight (g)</th>
<th>Length (cm)</th>
<th>HgB (ng/g)</th>
<th>HgE (ng/g)</th>
<th>HgPl (ng/g)</th>
<th>EC (n/mm³)</th>
<th>GLC (n/mm³)</th>
<th>CV (%)</th>
<th>MCV (fL)</th>
<th>TPP (g/dL)</th>
<th>Eos (%)</th>
<th>Neu (%)</th>
<th>Lym (%)</th>
<th>Mon (%)</th>
<th>BN (n/10⁴)</th>
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<tbody>
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<tr>
<td>C01</td>
<td>62,0</td>
<td>-0,6</td>
<td>-9,0</td>
<td>-24,5</td>
<td>-</td>
<td>-3,6</td>
<td>-335000,0</td>
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<td>-1,0</td>
<td>-10,0</td>
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<td>C02</td>
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<td>81,6</td>
<td>0,0</td>
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<tr>
<td><strong>Test Group</strong></td>
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<tr>
<td>T03</td>
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<td>1648,8</td>
<td>1845,5</td>
<td>582,6</td>
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<td>-70000,0</td>
<td>10,0</td>
<td>54,7</td>
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<td>T10</td>
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<td>-8,0</td>
<td>18,0</td>
<td>-1,0</td>
<td>4,0</td>
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</table>

* In gray, negative values

** In bold and italic font, comparisons calculated using the third sampling (four months of exposure-T2), not the fourth (six months of exposure-T3), because there was no result for that parameter at sampling T3

- = no values after exposure

HgB: Mercury Concentrations in Blood; HgE: Mercury Concentrations in Erythrocytes; HgPl: Mercury Concentrations in Plasma; EC: Erythrocytes Counting; GLC: Global Leukocytes Counting; CV: Corpuscular Volume; MCV: Mean Corpuscular Volume; TPP: Total Plasmatic Protein; Eos: Eosinophil Frequency; Neu: Neutrophil Frequency; Lym: Lymphocyte Frequency; Mon: Monocyte Frequency; BN: Bilobed Nucleus Frequency

Comparison to T0. CV and MCV increased after four months of exposure (T2) and TPP decreased. In this specimen was observed a significant difference on monocytes frequency at T1, where eosinophils decreased, but lymphocytes were not predominant. At sampling T2, a condition near to T0 was observed, where eosinophils predominated after the decreasing on lymphocytes and monocytes frequencies. At the end of the experiment (T3), lymphocytes increased again, but not followed by monocytes, being predominant. This specimen had the highest increase on bilobed nucleus frequency, having 8 abnormalities more than at T0.

The last specimen, T10, gained 55.7 g and grew 1.6 cm, showing the best length: Weight relationship. In this specimen, hematimetry decreased during all the experiment, but CV decreased only at the first two months. After that CV increased until the end of the experiment. MCV increase throughout the experiment and showed, apparently, a positive relationship with mercury concentrations in blood, following mercury increasing. TPP oscillated, showing higher concentrations at T1 and T3 and lower concentration in T2, in comparison to T0. At the first two months, lymphocytes predominated, including an increase on monocytes frequency too. After that, eosinophils were the most frequent leukocyte type found. This specimen had high frequencies of nucleus abnormalities since the beginning, showing four more bilobed nuclei at T2 in comparison to T0.

Overall, test group, besides the massive difference on mercury concentrations after the beginning of exposure (Friedman Anova, n=3, df=3, 8.2, p<0.05), showed a tendency of increase on CV, while for the control group the CV decreased. Also, the variation on specific leukocytes count was clear and stronger on test group, especially for monocytes, which increased for test group, but decreased on control group. Comparing only the beginning and the fourth month of exposure (T0 and T3 samplings, respectively), besides biometry (Wilcoxon, n=5, T=0.0, Z=2.02, p<0.05) and mercury concentrations in blood (Wilcoxon, n=5, T=0.0, Z=2.02, p<0.05), it was observed a significant difference for erythrocytes count, global leukometry and mean corpuscular volume (Wilcoxon, n=5, T=0.0, Z=2.02, p<0.05), corroborating the description mentioned above for each individual of test group. For this individual analysis, statistical tests were not applied to control group due to the low number of specimens that survived until the end of the experiment (n=2) ([Table 3]).

**Discussion**

Some field works use the ratio of MeHg concentrations in muscles and liver to evaluate a possible equilibrium of mercury concentrations in fish [39, 40]. They suggest that a ratio between 1.0-1.5 would indicate equilibrium and stability. In this work, the concentrations on muscles were lower than in the liver (Hg Muscles: Hg Liver=0.81), being at a non-equilibrium state. This was expected since liver is an important organ for detoxication and also it is one of the first organs in contact to xenobiotics such as methylmercury. Mela et al. also found lower mercury concentrations in muscles (4.77 ± 1.99 µg g⁻¹) than in liver (8.26 ± 6.45 µg g⁻¹) for a neotropical fish species (*Hoplias malabaricus*) exposed by contaminated food ingestion during 70 days with a similar dose of the present work (10.5 µg g⁻¹), presenting a ratio of 0.58 [41]. Mercury concentrations in erythrocytes were four times lower than those found in muscles, indicating a high methylmercury accumulation rate. The ratio found for mercury concentrations in erythrocytes and plasma was 1.22. It was expected a higher ratio since that methylmercury has higher affinity for erythrocytes, according to iron and sulfur concentrations in these cells [42]. Rodrigues et al. found for a...
wildlife catfish (Genidens genidens) a ratio of 6:1 [1].

The apparent tendency of decrease on erythrocytes counting and on total plasmatic protein over the time, for both groups, were inconclusive, since they could be linked to the long period of permanence of these species in aquarium or even physiological variations, due to age and spawning period. It was observed that all the specimens in the last months have chosen specific places at the aquarium to clean the bottom and starts incubation of eggs, even without physical or visual contact with other individuals.

Among the observed effects, it was described a non-regenerative anemia scenario for specimens from test group. This same scenario was observed for tucunarés (Cichla sp.) from ASSGM areas, extremely contaminated with mercury in Tapajós River, Amazon, Brazil and for a catfish species (Genidens genidens) from Guanabara Bay, Rio de Janeiro, Brazil [1, 32]. In this respect, Castilhos et al. suggested that mercury could be directly involved in ionic equilibrium, increasing the cell volume [32]. According to Lohner et al., methylmercury could be involved on the erythrocytes turn over, inducing anemia in fish specimens [43]. Also, the negative correlation between mercury concentrations on erythrocytes and erythrocytes counting can suggest a chemical stress inducing lower number of erythrocytes.

For specific leukometry, it is common to have higher lymphocytes frequency and lower proportions of other leucocytes types [44]. However, in this study, for both test and control groups, eosinophils were predominant. After exposure, for test group, lymphocytes frequency oscillated, being predominant at some samplings. Higher frequencies of eosinophil and monocytes were correlated to mercury concentrations, suggesting a possible chronic monocyctosis scenario and an increase on immunological reaction. The eosinophil does not have a well delimited function in fish, but it is believed that it is associated to defense against parasites and it is important on a post-inflammatory scenario, being related to phagocytosis of bacteria [44]. Ribeiro et al. on the other hand, observed an increase on erythrocytes number and on corpuscular volume, for another neotropical fish species (Hoplias malabaricus) fed with a MeHg artificially contaminated food during 70 days with a similar dose of the present work (10.5 µg g⁻¹) [48]. Also the authors described an increase on neutrophil and a decrease on monocytes frequencies.

The microuncelus frequency is related to haematepoiesis, performed by kidneys in fish. If the kidney presents a dysfunction, errors during cell division or new erythrocytes formation are expected. During tests in laboratory conditions, the interval between exposure to a xenobiotic and the initial detection of microuncelus frequency increase is related to the erythrocytes half-life, being substituted by new ones after few days of exposure. These new erythrocytes have higher errors rates during mitosis [46]. Sanchez-Galan et al. demonstrated that both cadmium and mercury could induce microuncelus formation in eels after intraarterional injections (1.7 mg of metal per Kg of specimen weight), observing a frequency of 2.64 and 2.35 microuncelus per 1,000 erythrocytes, respectively [47]. Vicari et al. did not found a clear induction of microuncelus or another abnormalities for Hoplias malabaricus fed with an MeHg artificially contaminated food during 70 days with a similar dose of the present work (10.5 µg g⁻¹) [48].

Porto et al. found significantly higher frequencies of micronucleus in Prochilodus nigricans (detritivorous species), Mylossoma duriventris (omnivorous species) and Hoplias malabaricus (piscivorous species) from Madeira River, a highly contaminated area by ASSGM activities on Amazon Region, compared to frequencies in specimens from Solimões River [49]. Moreover, micronucleus frequency was five times higher in Hoplias malabaricus than in the other two species, which was suggested due to higher mercury levels resulted of its food habit. However, Castilhos et al. did not find differences on micronucleus frequencies comparing a ASSGM site on Tapajós River and a reference area, also from Amazon Region, for Cichla sp. [32].

Souto has found positive correlation between mercury in muscles and acetylcholinesterase activity for Cichla sp. [50]. from ASSGM areas at Amazon Basin. In laboratory assays, its activity in brain, gills and liver was inhibited due to mercury exposure, but no inhibition was found in muscles [51]. Costa et al. registered a reduction of 36% on muscle ChE activity for Hoplias malabaricus fed with a MeHg artificially contaminated food during 70 days (dose=0.075 µg g⁻¹) [30]. Other authors have not found this direct relationship between mercury and acetylcholinesterase in muscles, which may be linked to differences on metabolism, including different isofoms of the cholinesterase group [52, 53].

Magnotti et al. reported the occurrence of important differences between acetylcholinesterase types in several fish species [52]. They suggested the existence of two groups of AChE, one with higher activity and the other with lower activity. Differences on acetylcholinesterase sensibility were reported previously for fish and they are linked to many factors, such as the ability of the enzyme’s active site to join the alkyl chain of a substrate like acetylcholine and butyrylthiocholine iodies [54].

Regarding the observed effects on histology, lipodosis can occur due to long periods of fasting or high fat percentage on food. However, this could be a natural characteristic of this fish species, especially during adult life, during reproduction period. Lipid bodies were sporadically visible in the cytoplasm of hepatocytes on control group of the experiment described by Mela et al. for Hoplias malabaricus fed with a MeHg artificially contaminated food during 70 days [55]. Additionally, the authors reported macrophage aggregates and necrosis areas, the presence of atypical cytoplasmic electron dense granules within hepatocytes and of melanomacrophage centers, and high vacuolization of the endothelial cells in H. malabaricus liver. Despite the reduced number of specimens for histological analysis, fish from test group showed severe damages on gonads, which could prejudice their reproduction inevitably, in case of high doses of methyl mercury exposure on wildlife.

**Context in wild fish populations**

As this experiment was carried out using a high dose with a long time pattern, it is also interesting to discuss the expected exposure concentrations of this and other fish species to methylmercury.
in natural conditions and their health responses. Astronotus ocellatus occurs naturally on the Amazon River basin, including registers in Peru, Colombia and Brazil. However this species can occur as an invasive exotic species at aquatic systems. There is only one published study that includes mercury contamination assessment using A. ocellatus, but it was held at a Amazonian lake (Manacapuru Great Lake), located at the drainage basin of Solimões River, with no contamination or anthropogenic source of mercury [56]. Then the observed mercury concentrations in muscles were low (134.2 ± 113.9 ng.g⁻¹), even for piscivorous species. About the effects in this species in wildlife due to mercury exposure, there is no published data available for comparison.

Although for most natural areas it is not expected such high doses, previous works demonstrated that fish are easily exposed to high mercury doses in old ASSGM areas in Brazilian Amazon, where A. ocellatus species occurs. For example, at Tapajós River and its tributaries show mercury concentrations in sediments and in suspended solids between 10⁻¹ and 10¹µg.g⁻¹, being mercury concentrations in the organic matter content of these suspended solids until 10¹µg.g⁻¹ and in planktonic organisms between 10¹ and 10³µg.g⁻¹ [57, 58]. These are initial concentrations in the trophic chain base, which is able to increase 104-105, or more, considering piscivorous fish species, even in non-contaminated areas [59]. Castilhos et al. estimated a daily dose of 0.8 ng.g⁻¹.day⁻¹ for Cichla sp. from Tapajós River [59].

Overall, a study at an ASSGM area located at one tributary of the Tapajós River, found extremely high mercury concentrations in muscles of a carnivorous species-Hoplias malabaricus (Erythrinidae) (6.11 ± 5.93 µg.g⁻¹, n=20), reaching the maximum of 21.9 µg.g⁻¹, which compared to our experiment, showed quite similar mercury concentrations in muscles [8]. Regarding methylmercury exposure specifically, Huang et al. observed a range of 79.6-100.0% for methylmercury in muscles of 15 Amazonian fish species. Castilhos et al. found about 84.4% of methylmercury in muscles of H. malabaricus. For Cichla sp., from the same family of A. ocellatus, the percentage is even higher about 96.6% [8, 60]. Then, one could suggest that results from this bioassay may be fit with some actual mercury exposure conditions in the field.

Concerning the effect biomarkers, we decided to compare our results with two studies developed in Tapajós River Basin, in some ASSGM sites [32, 61]. The first study was held in 2001, where specimens of Cichla sp. were collected at Tapajós River (near to Itaituba city-an ASSGM site) and at Maicá Lake (Santarém city-a reference area) [33]. The authors described higher MCV and lower erythrocyte count, corpuscular volume and leukocyte count at the contaminated area, all correlated to mercury concentrations, suggesting a non-regenerative anemia and a deficiency on immunological system caused by mercury exposure. Additionally, granulocytes (eosinophil and neutrophil) (55.8 ± 15.1%) were lower at Itaituba City, whereas lymphocytes (31.0 ± 15.3%) and monocytes (13.4 ± 8.6%) were higher comparing to the reference area. Also lymphocytes were not the prevalent leukocyte type [62] (Table 4).

The second study was held in 2003, at two study areas: São Chico and Creporizinho ASSGM sites [60]. The authors evaluated mercury exposure and hematological alterations on fish from tributaries of Tapajós River, using one carnivorous species, Hoplias malabaricus and one omnivorous, Caquetaia spectabilis (Cichlidae). Both species had lower corpuscular volume, erythrocytes count and mean corpuscular volume at São Chico mining site, where the specimens also showed higher mercury concentrations in muscles, suggesting a regenerative anemia. Besides, mercury levels and corpuscular volume showed significant negative correlation for both species (H. malabaricus: -0.82, p<0.005 n=11, C. spectabilis: -0.37, p<0.05 n=29).

<table>
<thead>
<tr>
<th>Area</th>
<th>ASSGM</th>
<th>HgM</th>
<th>Lenght</th>
<th>EC</th>
<th>CV</th>
<th>MCV</th>
<th>GL</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>São Chico</td>
<td>Present</td>
<td>7.71 ± 6.20 (9)</td>
<td>24.1 ± 7.2 (9)</td>
<td>2.2 ± 0.4 (7)</td>
<td>26.9 ± 5.5 (7)</td>
<td>123.4 ± 15.4 (7)</td>
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<tr>
<td>Creporizinho</td>
<td>Present</td>
<td>0.73 ± 0.43 (4)</td>
<td>18.6 ± 9.8 (4)</td>
<td>2.8 ± 0.4 (4)</td>
<td>40.7 ± 5.2 (4)</td>
<td>148.9 ± 26.5 (4)</td>
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<td>Castilhos et al. 2004b</td>
</tr>
</tbody>
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* Caquetaia spectabilis (Cichlidae): Omnivorous

<table>
<thead>
<tr>
<th>Area</th>
<th>ASSGM</th>
<th>HgM</th>
<th>Lenght</th>
<th>EC</th>
<th>CV</th>
<th>MCV</th>
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<tr>
<td>São Chico</td>
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<td>2.16 ± 1.02 (18)</td>
<td>8.2 ± 0.6 (18)</td>
<td>1.9 ± 0.3(17)</td>
<td>24.8 ± 4.0(18)</td>
<td>128.4 ± 19.5(17)</td>
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<td>Castilhos et al. 2004b</td>
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<tr>
<td>Creporizinho</td>
<td>Present</td>
<td>0.42 ± 0.26 (11)</td>
<td>8.7 ± 0.8 (11)</td>
<td>2.1 ± 0.2 (11)</td>
<td>29.3 ± 3.1 (11)</td>
<td>141.7 ± 18.0 (11)</td>
<td>-</td>
<td>Castilhos et al. 2004b</td>
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* Cichla sp. (Cichlidae): Carnivorous

<table>
<thead>
<tr>
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<th>ASSGM</th>
<th>HgM</th>
<th>Lenght</th>
<th>EC</th>
<th>CV</th>
<th>MCV</th>
<th>GL</th>
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<tr>
<td>Santarém</td>
<td>Absent</td>
<td>0.24 ± 0.11 (53)</td>
<td>30.2 ± 3.9 (55)</td>
<td>2.7 ± 0.6 (34)</td>
<td>44.4 ± 8.1 (35)</td>
<td>183.1 ± 36.7 (35)</td>
<td>53.2 ± 11.2 (31)</td>
<td>Castilhos et al. 2004a</td>
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<tr>
<td>Jacareacanga/</td>
<td>Present</td>
<td>0.71 ± 0.22 (55)</td>
<td>30.0 ± 6.7 (56)</td>
<td>2.0 ± 0.4 (53)</td>
<td>40.3 ± 5.6 (51)</td>
<td>211.5 ± 41.4 (49)</td>
<td>36.2 ± 10.0 (49)</td>
<td>Castilhos et al. 2004a</td>
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* Astronotus ocellatus (Cichlidae): Omnivorous

<table>
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<th>Bioassays</th>
<th>Artificial exposure</th>
<th>HgM: Mercury Concentrations in Muscles; EC: Erythrocytes Counting (10⁵ mm⁻³); CV: Corpuscular Volume (%); MCV: Mean Corpuscular Volume (fL); GL: Global Leukometry (10⁵ mm⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present</td>
<td>Study</td>
<td>6.39 ± 1.09 (4)</td>
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The same pattern of hematological response was observed for the test group of this bioassay and it is clear that the observed effects on wildlife specimens were reproduced on a smaller scale, in laboratory conditions, reinforcing the relationship between environmental exposure to mercury and biological effects observed on field work.

**Conclusion**

Concluding, mercury bioaccumulation was clearly observed in all fish tissues ($10^4$) and several effects were observed in *A. ocellatus* due to methylmercury exposure, including: decrease in erythrocytes number, increase in corpuscular volume and in monocytes frequency, increase in hepatocytes volume and their degeneration, hypertrophy and hyperplasia of follicular layer, increase in follicular atresia, yolk degeneration and presence of proteic material and inflammatory infiltrate on liver and gonads. Genotoxic effects were not clear and the acetylcholinesterase activity seemed to be higher after exposure. A study at ASGM found mercury concentrations in muscles quite similar of this present bioassay, which permits to suggest that this bioassay could be fit well with some actual wildlife mercury exposure conditions. Since these are the first data for methylmercury exposure of an Amazon fish species kept in laboratory for a chronic experiment (with 180 days of duration, observing juveniles until adult age), more tests are necessary to understand methylmercury non-lethal effects on *Astronotus ocellatus*. Future steps include rising the number of specimens and doses to observe the reproducibility of these results and to perform a dose response relationship analysis.

**Acknowledgement**

The authors would like to thank Prof. Renato Clapp (Fluminense Federal University, Veterinary School), for offer a part of his laboratory physical space to us, allowing this experiment to be done, to FIOCRUZ (Oswaldo Cruz Foundation), where we made the fish food contamination, to CAPES (Coordination for the Improvement of Higher Level or Education Personnel), for the financial support, and all the veterinary students that helped during the experiment and samplings.
References


5. (1994) Agency for toxic substances and disease registry toxicological profile. US Department of Health and Human Services, TP-93/05, Atlanta, GA.


