S-Methyl Cysteine Protective Effects in *Oreochromis Niloticus* Fish Contaminated by Thiobencarb Herbicide

Mostafa Ali Elmadawy*, Walied Abdo², Amira Alaa El-Dein Omar¹, and Nadia B. Mahfouz⁴

¹Department of Forensic Medicine and Toxicology, Faculty of Veterinary Medicine, Kafrelsheikh University, Kafrelsheikh, El Geish Street, 33516, Egypt
²Department of Pathology, Faculty of Veterinary Medicine, Kafrelsheikh University, El-Geish street, 33516 Kafrelsheikh, Egypt.
³Department of Fish Diseases and Management, Faculty of Veterinary Medicine, Kaf El Sheikh University, El-Geish street, 33516 Kafrelsheikh, Egypt.
⁴Department of Pathology, Faculty of Veterinary Medicine, El-Geish street, 33516 Kafrelsheikh, Egypt.

*Corresponding author's Email: drmmostox@yahoo.com. DOI: 0000-0002-7174-1370

ABSTRACT

Thiobencarb which is a carbamate herbicide is used for managing undesirable weeds during rice cultivation in Egypt. This study was designed to investigate the adverse effects of a field dose of thiobencarb on Nile tilapia and ameliorating the role of the low dose of S-methyl cysteine (SMC). Experimental fishes were divided into four groups; first group was reared without any treatments and served as a control group; the second group was exposed to thiobencarb (36µg/L); the third group was fed on a commercial feed containing 200 mg of SMC/Kg in conjunction with thiobencarb added to aquarium (36µg/L) while, the fourth group was fed on a feed containing 200 mg of SMC/Kg only. Fishes were sacrificed at the end of the experimental course (two months) and sampling was carried out. Catalase, Glutathione S Transf erase activities, Glutathione reduced, and brain was also evaluated by several biomarkers. Meanwhile, the comet parameters compared to control values. Moreover, histopathological findings were in line with other results. SMC ameliorated the adverse effects which were effective in the improvement of DNA and oxidative damage in thiobencarb intoxicated fish.

Keywords: Carbamate, Fish, Genotoxic damage, Histopathology

INTRODUCTION

Potential exposure to the toxic chemicals has been progressively increased in the last decades. Scientists have increased their ecological researches in detecting and controlling the hazard chemical agents responsible for unacceptable damages to the ecosystems and human health (Munger et al., 1997; Gorell et al., 1998; Nwani et al., 2011; Nicolopoulou-Stamati et al., 2016). These agents may have adverse physiological, pathological or biochemical effects to living beings, some of them may possess a mutagenic potential which can express their action later in the next generation of the organisms (Anderson, 2005; Recio et al., 2010). Noxious weeds in common are the most remarkable and prevalent biological problem to crop production in rice fields which are commonly controlled by chemical herbicides (Sapari and Ismail, 2012; Hakim et al., 2015). These herbicides can strike their way to the water bodies through surface runoff (Phong et al., 2006; Papadakis et al., 2015) causing impacts on non-target organisms specially those living in the aquatic environment which differs according to degree of dispersion, concentration and toxicity of the herbicide (Van der Werf, 1996). The residues of these herbicides accumulate in the environment due to its widespread use over the last years which negatively damaged the aquatic organisms and living biota (Silva et al., 2019; Valadas et al., 2019).

Thiobencarb is a dithiocarbamate selective herbicide was developed by Kumiai Chem. Ind. Co. Ltd and introduced in the agricultural field to control the unwanted weeds in the rice fields in 1970 (Ishikawa et al., 1977). It has been used extensively by the farmers in Egypt particularly during the rice cultivation season. Previous studies have been conducted to assess the thiobencarb toxicity to the ecosystem (Bailey, 1993; Fernández-Vega et al., 1999; Saka, 2010).

Genotoxic and mutagenic effects of herbicides have been previously evaluated in several model organisms (Sanderson et al., 2001; Hladik et al., 2008; Benfeito et al., 2014) using several biomarkers for genotoxicity assessment, such as DNA adducts, chromosomal aberrations, DNA strands breaks, micronuclei formation and cell death induction (Stein et al., 1993; Sikka et al., 1990; Bombail et al., 2001). *Oreochromis niloticus* fish have been used in several studies as a model for genotoxicity investigation of various environmental pollutants (Lima et al., 2006; Kandiel et al., 2014; Bacolod et al., 2017). Several studies have been conducted implementing relatively simple methods to verify the mutagenic effects of the herbicides. The comet and micronuclei assays can be applied for evaluation of the genetic material damage in the experimental and field studies (Monteith and Vanstone, 1995). S-Methyl Cysteine (SMC) is an...
organosulfur compound naturally discovered in some plants particularly garlic (*Allium sativum*). A promising protective role of SMC had been prophesied as anti-apoptotic (Nasr et al., 2017), antidiabetic (Senthilkumar et al., 2013) and anticancer (Fukushima et al., 2001). However, a high dose of SMC may induce adverse effects (El-Magd et al., 2017).

Present study, aimed to point out the genotoxicity of a field dose of thiobencarb on *Oreochromis niloticus* fish and its modulation by a low dose of SMC.

**MATERIALS AND METHODS**

**Ethical approval**

This study was conducted under the ethical approval from the Experimental Animal Care Committee in accordance with guidelines of Faculty of Veterinary Medicine, Kafrelsheikh University, Egypt.

**Chemicals**

A technical grade of thiobencarb (95 % purity) from Kafrelzyat Chemicals Company, Egypt was used in presenta experimental study. S-methyl cysteine was purchased from Sigma-Aldrich.

**Fish acquisition and maintenance**

One hundred and sixty Nile tilapia (*Oreochromis niloticus*) fish were used for this study. The average weight of fishes was 30±3.5 g. The fish were purchased from a private fish farm in Kafrelsheikh governorate and then carefully transferred to the laboratory in water tanks filled with water from the fish collection source to avoid the stress of another water composition and temperature.

**Aquariums and fish rearing**

Fish were transferred to glass aquariums filled with dechlorinated water, each of them has 200-liter capacity. Fish were acclimatized to the lab conditions for 21 days. Fish were fed on commercial fish diet. Water was replaced every three days during the acclimation period.

**Experimental design**

Fish were divided into four groups; each group contains 40 fish and then each 40 fish divided into two aquariums (20 fish in each). Fish groups were treated as follows; first group (Control group), fish were exposed merely to dechlorinated tap water; second group, fish were exposed to 36 µg/L thiobencarb (this dose was determined as the average concentration of thiobencarb in 10 water samples collected from water drainage bodies near rice filed in Kafrelsheikh, Egypt using gas chromatography); third group, fish in each aquarium were provided every day with 15 g ration containing SMC (200 mg/Kg). Fourth group: fish were exposed to thiobencarb (36 µg/L) and fed on ration containing SMC (200 mg/kg). The experimental course endured 60 days. The water of all aquariums was replaced once a week during the experimental course. The particular herbicide concentration was supplied for the third and fourth groups whenever the water was changed. The laboratory conditions during the experimental period in common were Light period ~12hrs and temperature 25±2°C.

**Sampling**

Fishes were sacrificed at the end of experimental course (two months) and sampling was carried out. Eight fish samples were randomly collected from each group (four samples from each aquarium) at the end of experiment. Fishes were sufficiently dried by paper towels. Heparinized blood samples were collected from the caudal vein puncture for micronuclei analysis. Then fish samples were dissected. Liver and gills samples were picked up and divided into three sufficient parts. First one preserved on PBS buffer solution at -80 °C for carrying out comet assay. Second part processed directly for antioxidants biomarkers assay and the third portion was properly preserved in formalin 10 % for histopathological examination.

**Determining antioxidant biomarkers**

Mincing and homogenization of Liver and gill samples was carried out in phosphate buffer saline and pH 7.4. The homogenate was centrifuged at 4,000 rpm for 20 minutes and supernatants were isolated for measuring antioxidant biomarkers level which include Catalase (CAT), Glutathione S Transferase (GST) activities, total Glutathione reduced (GSH), and Malondialdehyde (MDA) which were assayed by utilizing specified methods described by Beutler et al. (1963), Habig et al. (1974), Ohkawa et al. (1979) and Aebi (1984).

**Micronuclei analysis**

Heparinized blood samples were immediately smeared on clean glass slides and then stained with Giemsa stain. Micronuclei was assayed according to the previously described method (Cavaş and Ergene-Gözükara, 2003). The
number of 1,000 clearly separate cells from the neighboring were scored from each slide by using light microscope (labomed, USA) under 100 × magnifications. Micronuclei were scored, if it was clearly separated from the nucleus (Al-Sabti and Metcalfe, 1995). Other nuclear abnormalities (blebbed nuclei, lobed nuclei, notched nuclei and binuclei) were recorded.

**Comet assay**

Single cell electrophoresis was carried out according to the published method (Singh et al., 1988). Briefly, 10 ul of fish liver or gills homogenate were thoroughly mixed with 90 ul of low melting point agarose (0.7% in PBS) at 37º C. This prepared mixture was then added to a fully frosted microscope slide coated with 110 ul of normal melting point agarose (1% in PBS). A cover slip was directly placed on top of the slide, and the agarose layer was allowed sufficiently to solidify for 10 minutes at 4º C. Afterwards, the cover slip was carefully removed and a second layer of low melting point agarose without cells was added gently, a cover slip was applied, and the slide was held at 4º C for five minutes to allow the agarose layer to solidify. After removal of the cover slip, the slides were placed in lysis buffer (2.5 mol/L NaCl, 100 mmol/L Na2EDTA, 10 mmol/L Tris, (pH 10)) with freshly added 1% Triton X-100 and 10% DMSO for at least one hour at 4º C. Later, slides were placed in the electrophoresis chamber and incubated with electrophoresis alkaline buffer (300 mmol/L NaOH, 1 mmol/L Na2EDTA, pH > 13) for 15 minutes at 4º C to allow DNA unwinding and the expression of alkali-labile DNA damage as strand breaks. Electrophoresis was for 30 minutes at 25 V and 300 mA. The slides were washed three times, with 0.4 mol/L Tris with pH 7.5 as neutralization buffer. Finally, slides were stained with 50 ul of ethidium bromide (2 mg/mL), covered with a cover slip and observed under 400 × magnification with a Optika Axioskop fluorescent microscope.

From each sample, 100 randomly selected cells, respectively, were photographed and scanned. The scanned images were analyzed with the comet score analysis system. For each cell, the length of DNA migration (tail length) was measured in PX from the center of the nucleus to the end of the tail. The percentage of DNA in the tail was determined by measuring the total intensity (fluorescence) in the cells. The tail moment, expressed in arbitrary units which was calculated as tail length multiplied by percentage of migrated DNA/100.

**Histopathological examination**

Samples from the liver, gills and brain were collected from fish of various groups. The samples were fixed in 10% neutral buffered formalin, dehydrated in alcohol, cleared in xylene, embedded in paraffin and sectioned in 5 μm thickness. Then serial sections were subjected to routine hematoxylin and eosin staining (Bancroft and Layton, 2013).

**Statistical analysis**

The data was analyzed by ANOVA procedures of the IBM SPSS software program (Version 16.0; IBM Corp., NY, USA). The differences between different means were estimated by the Duncan Multiple Range Test (DMRT). The differences among means with p < 0.05 were represented statistically as significant differences.

**RESULTS**

**Antioxidant biomarkers**

Catalase, GST activities, GSH and MDA level were demonstrated in homogenate samples from liver and gills of fish of all experimental groups. The results were presented in Figure 1. The CAT activity was significantly (P<0.05) decreased in both hepatic (3.38) and gill (3.32) tissues of thiobencarb treated fish compared to control fish (4.49) and (4.37), respectively. The inhibition of CAT activity was ameliorated by SMC concurrently with thiobencarb which was more prominent in gills than liver. GST activity was significantly elevated in thiobencarb intoxicated fish gills (0.44) and liver (0.36) compared to the control values (0.29) and (0.18), respectively. This elevation was diminished with SMC provided fish. GSH and MDA levels were increased significantly (P<0.05) in the gills and hepatic tissues of intoxicated fish over the control non treated fish. SMC provided in fish ration decreased the level of GSH and MDA compared to the intoxicated fish. Antioxidant biomarkers that measured in the fourth group were evidence to antioxidant system improvement by SMC.

**Micronuclei analysis**

The small non refractive parts that present in cytoplasm and are similar to the nucleus in staining is recorded as micronucleus, it is usually one particle fully separated from the nucleus. The determined percentage of micronuclei was significantly increased (P<0.05) in the second fish group which treated with thiobencarb, compared to the control group. The increased micronuclei level due to thiobencarb was relieved by SMC in the third group compared to the first and second group. Nuclear abnormalities were also recorded in the examined samples (Table 1). Binucleated cells and abnormal nuclei were significantly increased (P<0.05) in fish treated with thiobencarb compared to the control group, fish treated with SMC and thiobencarb revealed improvement in the effect on erythrocyte nuclei. The SMC provided groups revealed non-significant decrease (P>0.05) in nuclear abnormalities compared to the control group.

<table>
<thead>
<tr>
<th>Table 1</th>
</tr>
</thead>
</table>

Table 1. Nuclear abnormalities in erythrocytes of *Oreochromis niloticus* exposed to thiofencarb (36 µg/L) and SMC (200 mg/kg ration) compared with control non treated fish

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Control</th>
<th>Thiofencarb</th>
<th>Thiofencarb + SMC</th>
<th>SMC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Micronucleus (%)</td>
<td>0.032±0.0011&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.040±0.0015&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.033±0.0012&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.030±0.0009&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Binucleated</td>
<td>0.020±0.0012&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.037±0.0005&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.026±0.0009&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.018±0.0011&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Abnormal Nuclei</td>
<td>0.012±0.0008&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.043±0.0012&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0.023±0.0012&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.012±0.0009&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>a</sup> Means with different superscripts in each row are significantly differ at P < 0.05. SMC means S-Methyl Cysteine

**Comet assay**

The DNA damage in liver and gills tissues of *Oreochromis niloticus* exposed to thiofencarb and the potential effect of SMC treatment was investigated by using comet assay. The observed results were presented in Table 2 and Figure 2. Regarding to the Comet parameters in control fish group gills, Comet%, Head Diameter, DNA % Head, Tail Length, DNA % Tail and Tail Moment were 13.99 %, 20.29, 82.3 %, 5.8, 19.32 % and 1.34, respectively. Fish group treated with thiofencarb indicated a significant increase (P<0.05) in the values of Comet % (18.45 %), Head Diameter (25.5), Tail Length (7.59), DNA % Tail (23.5 %), Tail Moment (2.47) and significant decrease in DNA % Head value (76.91 %). The fish group treated with SMC in conjunction with thiofencarb indicated significantly ameliorated DNA damage effects exerted on gill tissue compared to thiofencarb treated fish group. The comet criteria in the fourth fish group which treated with SMC alone represented non significantly difference (P>0.05) with control values of Comet % (13.2 %) and DNA %Head (82.17 %), while significant difference (P<0.05) in Head Diameter (18.28), Tail Length (5.26), DNA %Tail (18.02 %) and Tail Moment (1 %) was observed which indicated that the DNA damage in SMC group was rarely than the control group. Observational data of comet parameters evaluating the DNA damage effects in hepatic cells of thiofencarb and SMC treated fishes clarified a significant increase (P<0.05) in Comet % (13.6%), Head Diameter (24.44), Tail Length (7.07), DNA % Tail (22.7%) and Tail Moment (2.22) compared to the control group values including Comet % (12.4 %), Head Diameter (20.87), Tail Length (4.72), DNA % Tail (19.94 %) and Tail Moment (1.35) and significant decrease (P<0.05) in DNA % Head (75.82 %) compared to control (79.63 %) value. A repair in DNA damage was observed in fish treated with Thiofencarb+SMC which were evidenced by improved comet values in this group compared to the thiofencarb treated group. All comet variables for hepatic cells of fish treated with SMC exhibited non significantly differences (P>0.05) with the same parameters in the control group.

**Histopathological findings**

The histopathological findings were illustrated in Figure 3 and scoring of the lesions was mentioned in Table 3. The liver of the control fish presented normal hepatic and pancreatic tissues. The hepatic tissues consisted of normal hepatocytes with irregular hepatic vacuolation consistent with physiological glycogen storage. The pancreatic tissue was distributed within the hepatic tissues and enclosed venous blood supply. The thiofencarb treated fish presented marked decrease in hepatic vacuolation, marked increase in number of hepatocytes within the examined field, hyperplasia and single cell necrosis within the pancreatic portion. The SMC treated fish had normal hepatic and pancreatic tissues. The Thiofencarb and SMC treated fish demonstrated marked decrease in mitogenic effect which represented with decrease in number of hepatocytes. The gills of control fish revealed normal primary and secondary gill lamellae. Marked loss of secondary gill lamellae associated with necrosis of the lamellae, infiltration of leukocytes and proliferation of goblet cells were noticed within the gills of the intoxicated group. The gills of SMC treated fish were within the normal limits. The intoxicated and SMC treated fish revealed marked decrease in the adhesion of the secondary lamellae and appearance of the secondary gill lamellae. The brain of the normal fish indicated normal nerve cells and nerve fibers. The thiofencarb treated fish revealed multifocal malacia associated with marked gliosis and appearance of gitter-like cells. The brain of SMC treated was normal. The brain of thiofencarb intoxicated fish and SMC treated revealed limited malacic foci with marked decrease in the glia and gitter cells.

Table 2. DNA damage assessed by comet assay in liver and gills of *Oreochromis niloticus* exposed to thiofencarb (36 µg/L) and SMC (200 mg/kg ration) compared with control non treated fish

<table>
<thead>
<tr>
<th>Groups</th>
<th>Comet %</th>
<th>Head diameter</th>
<th>DNA % Head</th>
<th>Tail length</th>
<th>DNA % Tail</th>
<th>Tail moment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gills</td>
<td>Control</td>
<td>13.99±0.135&lt;sup&gt;a&lt;/sup&gt;</td>
<td>20.29±0.631</td>
<td>82.30±0.994&lt;sup&gt;a&lt;/sup&gt;</td>
<td>5.80±0.119&lt;sup&gt;a&lt;/sup&gt;</td>
<td>19.32±0.196&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Thiofencarb</td>
<td>18.45±0.370&lt;sup&gt;b&lt;/sup&gt;</td>
<td>25.50±0.350&lt;sup&gt;b&lt;/sup&gt;</td>
<td>76.91±0.601&lt;sup&gt;b&lt;/sup&gt;</td>
<td>7.59±0.143&lt;sup&gt;b&lt;/sup&gt;</td>
<td>23.50±0.407&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>SMC</td>
<td>16.32±0.448&lt;sup&gt;c&lt;/sup&gt;</td>
<td>22.64±0.495&lt;sup&gt;c&lt;/sup&gt;</td>
<td>79.96±0.275&lt;sup&gt;c&lt;/sup&gt;</td>
<td>6.40±0.039&lt;sup&gt;c&lt;/sup&gt;</td>
<td>21.28±0.292&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Liver</td>
<td>Control</td>
<td>12.04±0.168&lt;sup&gt;a&lt;/sup&gt;</td>
<td>20.87±0.340&lt;sup&gt;a&lt;/sup&gt;</td>
<td>79.63±1.144&lt;sup&gt;a&lt;/sup&gt;</td>
<td>4.72±0.335&lt;sup&gt;a&lt;/sup&gt;</td>
<td>19.94±0.493&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>Thiofencarb</td>
<td>13.60±0.240&lt;sup&gt;b&lt;/sup&gt;</td>
<td>24.44±0.324&lt;sup&gt;b&lt;/sup&gt;</td>
<td>75.82±1.160&lt;sup&gt;b&lt;/sup&gt;</td>
<td>7.07±0.121&lt;sup&gt;b&lt;/sup&gt;</td>
<td>22.07±0.525&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>SMC</td>
<td>12.47±0.428&lt;sup&gt;c&lt;/sup&gt;</td>
<td>20.31±0.781&lt;sup&gt;c&lt;/sup&gt;</td>
<td>78.76±0.622&lt;sup&gt;c&lt;/sup&gt;</td>
<td>5.53±0.455&lt;sup&gt;c&lt;/sup&gt;</td>
<td>20.39±0.398&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
</tbody>
</table>

<sup>*</sup> Means with different superscripts in each column are significantly differ at P < 0.05. SMC means S-Methyl Cysteine
Table 3. Semi quantitative scoring of the lesions within the liver, gills and kidney of *Oreochromis niloticus* exposed to thiobencarb (36µg/L) and SMC (200 mg/kg ration) compared with control non treated fish

<table>
<thead>
<tr>
<th>Groups</th>
<th>Liver</th>
<th>Gills</th>
<th>Brain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Degeneration</td>
<td>Hyperplasia</td>
<td>Adhesion</td>
</tr>
<tr>
<td>Control</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Thiobencarb</td>
<td>++</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>SMC</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>SMC+Thiobencarb</td>
<td>+</td>
<td>++</td>
<td>++</td>
</tr>
</tbody>
</table>

+, ++, +++ and ++++ indicate mild, moderate, severe and focal, severe and diffuse lesions respectively. SMC means S-Methyl Cysteine

Figure 1. Antioxidant biomarkers values in experimental fish (*Oreochromis niloticus*) groups A, B, C and D refers to Catalase, Glutathione S Transferase, Glutathione reduced and Malondialdehyde respectively. SMC refers to S-Methyl Cysteine
Figure 2. Photomicrograph by using comet assay showing the different degrees of DNA damage in liver and gills of *Oreochromis niloticus* exposed to thiobencarb (36µg/L) and SMC (200 mg/kg ration) compared with control non treated fish. (A), (B), (C), (D) refers to control, Thiobencarb, Thiobencarb+ S-Methyl Cysteine and S-Methyl Cysteine treated fish groups respectively. (Ethidium bromide X40)
Figure 3. Histopathological changes in liver, gills and brain of Oreochromis niloticus exposed to thiobencarb (36µg/L) and SMC (200 mg/kg ration) compared with non-treated fish. The liver of control fish (A) showing normal hepatic (arrow) and pancreatic tissues (arrowhead) (the hepatocytes revealing normal vacuolation consistent with glycogen storage), the liver of thiobencarb-treated fish (B) showing marked increase of hepatocytes indicated with increase the number of nuclei (arrow), the liver of normal fish treated with SMC (C) showing normal hepatic (arrow) and pancreatic tissues, and the liver of thiobencarb and SMC-treated fish (D) showing marked decrease the number of the proliferated hepatocytes (arrow), H&E, X200. The gills of control fish (A) showing secondary gill lamellae (arrow), the gill of thiobencarb-treated fish (B) showing marked loss and adhesion of the secondary lamellae (arrow) with increased number of mucous cells, the gill of normal fish treated with SMC (C) showing lamellae tissues (arrow), and the gills of thiobencarb and SMC-treated fish (B) showing marked decrease the adhesion of the secondary lamellae (arrow), H&E, X200. The brain (telencephalon portion) of control fish (A) showing normal neuronal cells and with normal nerve fibers (arrow), the brain of thiobencarb-treated fish (B) showing malacia (arrow) and gliosis (arrowhead), the brain of normal fish treated with SMC (C) showing neuronal tissues with normal glia cells (arrow), and the brain of thiobencarb and SMC-treated fish (B) showing marked decrease the necrotic changes and neuronal vacuolation (arrow), H&E, X200.
DISCUSSION

Aquatic environments contamination level with various chemicals has been increased in Egypt during the last years, which may be attributed to the intensive use of chemicals in the agricultural field (Dahshan et al., 2016). Chemical herbicides are a group of widely spread pesticides used for control of exotic weeds, which may be selective or non-selective. Thiobencarb is one of the intensively used selective herbicides in Egypt, which used primarily for the weeds emerging in the rice fields (Tanetani et al., 2013). Furthermore, the uncontrolled use of these chemicals during rice cultivation season poses a crucial source of aquatic environments pollution (Sapari and Ismail, 2012), it finds their way to the near water bodies through the surface runoff (Phong et al., 2006; Papadakis et al., 2015) causing adverse impacts on aquatic livings ranging from mild effects to life threatening ones (Bailey, 1993).

Fish are one of the most important aquatic organisms affected by water chemical pollution (Sommer, 1983). Moreover, it has been considered as a highly sensitive indicator for aquatic environment contamination (Singh et al., 1988). However, previous studies pointed out the adverse effects of water chemical pollutants particularly herbicides in different fish species (Nwani et al., 2011). Oreochromis niloticus was selected as an indicator for water pollution (Bacolod et al., 2017). Additionally, fish represent different adverse effects from thiobencarb herbicides (Bailey, 1993; Fernández-Vega et al., 1999; Saka, 2010).

This study pointed out the genotoxic effect of thiobencarb exerted on erythrocytes of Oreochromis niloticus which was obvious by significantly increased (P<0.05) micronuclei and nuclear abnormalities in thiobencarb treated fish. Moreover, the genotoxic effect was cleared in hepatic and gill tissues by significant increase (P<0.05) of comet parameters (Comet %, Head Diameter, Tail Length, DNA %Tail and Tail Moment) and significant decrease (P<0.05) of DNA %Head in thiobencarb treated fish compared to control group. Genotoxic effect may be attributed to the oxidative stress of the herbicide (Lima et al., 2006; Bacolod et al., 2017) which was ensured by significantly different antioxidant biomarkers in present results. Furthermore, this adverse toxic effect was reflected on the results of histopathological examination of liver, gills and brain of thiobencarb exposed fish. Hepatic section showed hyperplasia and single cell necrosis, while gills examination revealed marked loss of secondary gill lamellae associated with extensive necrosis of the lamellae, infiltration of leukocytes and proliferation of goblet cells. Also, multifocal malacia associated with marked gliosis and appearance of gitter-like cells was observed in brain tissue of intoxicated fish. The present study results were in line with Elias et al. (2020).

Herbicides induces oxidative stress remains the principal way for production of Reactive Oxygen Species (ROS) which may provoke oxidative damage to DNA, protein and lipids. The ROS detoxification is done by antioxidant system (Fan et al., 2013). Therefore, ROS detoxification failure prompts serious implication on cellular structure, function and life (Bacolod et al., 2017) which was clearly seen in tested parameters of thiobencarb intoxicated fish. The gills are the largest contacting organ of fish with the environment, which play the prime role in fish respiration, osmoregulation, and considered as a crucial part in fish excretion system (Simonato et al., 2008). However, the gills represent the first chief organ critically affected by water contamination, from this stand point the results revealed an elevation in GST activity and GSH level in the gills of intoxicated fish, which means that GST as well as GSH had an important role in thiobencarb detoxification.

SMC is an organosulfur compound naturally found in some plants particularly garlic (Allium sativum). A promising protective role of SMC had been prophesied as anti-apoptotic (Nasr et al., 2017), antidiabetic (Senthilkumar et al., 2013) and anticancer (Fukushima et al., 2001). Sulfur is a crucial component of various cellular proteins impacting the cell health particularly by enzymatic and non-enzymatic antioxidant molecules like glutathione and thioredoxin (Atmaca, 2004). In present study, fish exposed to thiobencarb and fed on ration containing SMC presented a curing effect based on the antioxidant biomarkers. The CAT activity was elevated, GST activity was corrected and MDA level was decreased, while sulfur dependent GSH was elevated after treatment with SMC compared with thiobencarb intoxicated fish. SMC ameliorated the oxidative DNA damage based on the correction of erythrocytes micronucleus and abnormal nuclei frequencies and comet parameters. Moreover, histopathological finding in gills, liver and brain of thiobencarb intoxicated fish were disappeared in SMC treated fish.

CONCLUSION

Oreochromis niloticus were sensitive to thiobencarb contamination which was explained by genotoxic and oxidative damage effects. Moreover, histopathological changes related to herbicide impact were recorded. These adverse toxic effects were ameliorated by S-Methyl Cysteine (SMC). Therefore, controlling of thiobencarb which is used as herbicide in rice fields should be done wisely. In addition, careful monitoring of thiobencarb should be carried out to minimize its adverse impacts on the aquatic ecosystems. S-Methyl Cysteine would be helpful for alleviating the toxic effects exerted by thiobencarb on Oreochromis niloticus. This study should be supported by more field studies to point out the effects of natural factors of aquatic environment on the use of SMC.
DEclarations

Acknowledgments
The authors would like to thank Mahmoud S.El-Tarabany, Department of Animal Wealth Development, Faculty of Veterinary Medicine, Zagazig University, Zagazig, Egypt for his assistance in statistical analysis of this study.

Competing interests
The authors have not declared any conflict of interest.

Author’s contribution
Mostafa Elmadawy, Walied Abdo, Amira Omar, and Nadia Mahfouz designed the study. Mostafa Elmadawy and Amira Omar performed the experiments. Walied Abdo and Nadia Mahfouz supervised the laboratory examinations. Histopathological examination was carried out by Walied Abdo. All authors prepared, read and approved the final manuscript.

Consent to publish
Not applicable

REFERENCES

Bombail V, Dennis AW, Gordon E, and Batty J (2001). Application of the comet and micronucleus assays to butterfish (Pholis gunnellus) erythrocytes from the Firth of Forth, Scotland. Chemosphere, 44: 383–392. DOI: https://doi.org/10.1016/S0045-6535(00)00300-3
Cavas T, and Ergene-Gozukara S (2003). Micronuclei, nuclear lesions and interphase silver-stained nucleolar organizer regions (AgNORs) as cyto-genotoxicity indicators in Oreochromis niloticus exposed to textile mill effluent. Mutation Research/Genetic Toxicology and Environmental Mutagenesis, 538(1-2): 81-91. DOI: https://doi.org/10.1016/S1383-5718(03)00091-3


Monteith DK, and Vanstone J (1995). Comparison of the microgel electrophoresis assay and other assays for genotoxicity in the detection of the DNA damage. Mutation Research, 345(3-4): 97-103. DOI: https://doi.org/10.1016/0165-7992(95)00045-4


Sapari P, and Ismail BS (2012). Pollution levels of thiobencarb, propanil, and pretilachlor in rice fields of the muda irrigation scheme, Kedah, Malaysia: Environmental Monitoring and Assessment, 184(10): 6347-6356. DOI: https://doi.org/10.1016/j.ecoenv.2010.05.008


