Haematological Parameters and Histopathological Alterations in the Gills of Fish, *Catla catla* Exposed to Azo Dye Acid Red -97

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Abstract  Haematological and histopathological parameters have been recognized as valuable tools for monitoring fish health. This study was conducted to evaluate the chronic toxicity of Acid Red 97 textile dyes on haematological and histopathological alterations using *Catla catla* as animal model. Fish were exposed to two Sublethal concentrations (1/100th and 1/50th of LC$_{50}$ = 85mg/l) 0.85 mg/l and 1.7 mg/l of AR 97 for a period of (10, 20 and 30) days. Haematological parameters were observed that with increase of exposure time, total erythrocyte (RBC), haemoglobin (Hb), and packed cell volume (PCV) values decreased but leucocytes (WBC), mean corpuscular hemoglobin concentration (MCHC), mean corpuscular volume (MCV) and mean corpuscular hemoglobin (MCH) values increased. It is believed that observed depression in packed cell volume and haemoglobin values coupled with decreased and deformed erythrocytes are obvious signs of anemia. Resulted changes in erythrocytes and leukocytes after exposing to AR 97 are due to malfunction in hemopoiesis and decrease in non-specific immune system.

Histopathological changes observed in the gills of *Catla catla* were swelling, aneurysm, fusion of lamellae, oedema in primary lamellae, shortened and severe erosions of secondary lamella and lifting epithelial layer and high mucus secretion. Hence, it was concluded that azo dye AR 97 has potential to cause toxicity in fish.

Keywords  Acid Red 97, *Catla catla*, Dye, Haematology, Histopathology

1. Introduction

Most of the azo dyes are released into the environment originate from the textile industry and mainly from dyestuff manufacturing industry [1]. Approximately 10-15% dyes are released into the environment during dyeing process, making the effluent highly coloured and aesthetically unpleasant [2,3]. Textile azo dyes are serious pollutants of the aquatic environment because of their environmental persistence and ability to be accumulated by aquatic organism [4]. Fish can be considered as indicator organisms of environmental pollution [5-7]. This is because they are constantly exposed to toxic elements dissolved in water, through gill breathing and epidermis contact [8,9]. Haematological parameters are patho-physiological reflectors of the whole fish body. Moreover, these blood parameters are too sensitive to any physicochemical, biological and/or environmental alterations. Hence, they are important markers to investigate the structural and functional status of fish exposed to various contaminants [10,11].

Histological changes have been widely used as biomarkers in the evaluation of health of fish exposed to contaminants both in the laboratory [12] and field studies [13,14]. Thus the present study deals with the chronic toxic effect of Acid red 97 on a freshwater fish *Catla catla* with the objective to study the haematological and histological alterations on the treated gill tissues for 10, 20 and 30 days.

2. Materials and Methods

2.1. Experimental Design

AR 97 was purchased from local market and used directly for the experiment. *Catla catla* fingerlings (13±1cm long and 25±2.5g weight) were procured from fish seed farm sivan, surat, Gujarat and acclimatized in the
laboratory condition for 15 days according to APHA [15]. After acclimatization, apparently healthy catla fingerlings were grouped in to 4 (10 in each) and used to carry out toxicity test. Among this one was considered as control and other 3 groups of catla fingerlings were exposed to sublethal concentration 0.85 and 1.7 mg/l (1/100th and 1/50th of LC50 = 85mg/l) of AR 97 for 30 days in triplicates. At fixed interval (10, 20, 30 days) after the exposure blood samples were collected and were analyzed.

2.2. Haematological Studies

Blood was collected by direct puncturing heart and/or caudal vein using sterile syringe (2 ml) pre-rinsed with 2.7 % EDTA solution. Collected blood samples were immediately transferred to vials (2 ml) coated with EDTA. Haemoglobin estimation was done by cyanmethemoglobin method [16]. 20 μl of blood samples were taken in Thoma’s pipettes. They were mixed with diluting fluids Turks’ solution for WBC count and Haem solution for RBC count in the same pipettes. The mixtures were shaken well to suspend cells uniformly in solution. After 10 to 15 minutes, cells were counted using haemocytometer [17].

2.3. Statistical Analysis

Statistical analyses were calculated in triplicate performed using the computer software ‘SPSS’. Data are presented as mean ± SD. The significant means were compared to control and a p<0.05 was considered to be the level of statistically significant.

2.4. Histopathological Studies

The fishes were exposed to both sub lethal concentrations of dyes and at the end of 10, 20 and 30 day exposure period gills were dissected out and fixed in 10% buffered formalin. The tissues were then processed in ascending order of alcohol for dehydration, embedded in paraffin wax and sectioned at 5-6μm on a rotary microtome. Sections were placed on glass slides and stained with haematoxylin and eosin [18]. Approximately 5-6 stained slides were examined with the help of a compound microscope and photographs were taken. Histopathological changes in these tissues were recorded and compared with controls under the guidance of a pathologist.

3. Results

3.1. Haematological Parameters

Fingerlings exposed to the low concentration (0.85 mg/l) of AR 97 azo dye showed decreased haemoglobin level and RBC count where WBC counts were increased. Similarly, the same trends were reported throughout the exposure at 10, 20 and 30 days. When blood sample collected from (1.7 mg/l) exposed fingerlings, also showed similar trend of decreasing in case of RBC and Hb and increasing WBC count throughout the exposure. In short, total RBC count and haemoglobin were decreased but WBC count was continuously increased throughout the exposure in both sublethal concentrations (Bar graph 1-3). All observations and results of AR 97 exposed fingerlings are summarized in the (Table 1). Statistical analysis indicates that all the values were significant.

| Sublethal Concentrations of AR 97 | Exposure (Days) | Blood parameters |
|-----------------------------------|----------------|------------------|
|                                   |                | Hb (g/dl)        | RBC count (10^6 cells/mm³) | WBC count (10^6 cells/mm³) |
| 0.85 mg/l                         | 0              | 6.47±0.03        | 2.42±0.15                  | 1.86±0.05                  |
|                                   | 10             | 6.34±0.18*       | 2.32±0.09*                 | 2.01±0.12*                 |
|                                   | 20             | 6.07±0.07*       | 2.00±0.10*                 | 2.18±0.09*                 |
|                                   | 30             | 5.88±0.12*       | 1.92±0.08*                 | 2.53±0.04*                 |
| 1.7 mg/l                          | 0              | 5.96±0.07*       | 2.11±0.06*                 | 2.31±0.04*                 |
|                                   | 10             | 5.80±0.07*       | 1.89±0.07*                 | 2.46±0.12*                 |
|                                   | 20             | 5.43±0.09*       | 1.59±0.12*                 | 3.06±0.09*                 |

[Each value is mean ± SE of thirty individual observations; Asterisks (*) indicate significant (p<0.05) difference compared to control.]
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**Bar graph 1.** Level of haemoglobin in *Catla catla*

**Bar graph 2.** RBC count in *Catla catla*

**Bar graph 3.** WBC count in *Catla catla*
Haematological indices like PCV, MCV, MCH and MCHC were calculated from obtained results of haemoglobin and RBC value of blood samples collected from control as well as exposed fingerlings. The blood sample collected from the fingerlings exposed to AR 97, showed decreased value of PCV throughout the experiment (Bar graph 4). The MCV and MCH both showed the increased value during the whole exposure period for both sublethal concentrations (Bar graph 5,6). The similar increasing trend was also noticed for MCHC (Bar graph 7). Obtained results are summarized in the Table 2.

Table 2. Effect of AR 97 on haematological indices

| Sublethal Concentrations of AR 97 | Exposure (Days) | Haematological indices |
|-----------------------------------|-----------------|------------------------|
|                                   | PCV %           | MCV (fl)               | MCH(pg)  | MCHC %   |
| 0.85 ppm                          | 0               | 24.84±0.09             | 102.64±0.10 | 26.73±0.12 | 26.06±0.11 |
|                                  | 10              | 23.92±0.10*            | 103.10±0.11* | 27.32±0.20* | 26.50±0.07* |
|                                  | 20              | 20.78±0.06*            | 103.90±0.09* | 30.35±0.13* | 28.93±0.09* |
|                                  | 30              | 19.99±0.09*            | 104.11±0.12* | 30.92±0.09* | 29.42±0.22* |
| 1.7 ppm                          | 10              | 22.04±0.04*            | 104.45±0.07* | 28.24±0.13* | 27.04±0.10* |
|                                  | 20              | 19.94±0.09*            | 105.50±0.05* | 30.68±0.21* | 29.08±0.14* |
|                                  | 30              | 16.91±0.06*            | 106.35±0.13* | 34.15±0.08* | 32.11±0.13* |

[Each value is mean ± SE of thirty individual observations; Asterisks (*) indicate significant (p<0.05) difference compared to control.]
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**Bar graph 5.** Level of MCV in *Catla catla*

**Bar graph 6.** Level of MCH in *Catla catla*

**Bar graph 7.** Level of MCHC in *Catla catla*
3.2. Histopathology of Gills

Primary gill lamellae have rows of secondary gill lamellae ideally arranged and lined up on both sides. The typical histological pattern in secondary lamellae of the normal gill is shown in control. The epithelial cell line covers both primary and secondary gill lamellae. As the gills are the first target organ for any toxicant, they are affected maximally. Many histological changes were observed in gill of exposed fingerlings. These changes were concentration as well as time dependent. After exposure to both (0.85 and 1.7 mg/l) sublethal concentrations of AR 97, gills showed mild to severe haemorrhage at the axis of primary gill lamellae. Extensive hyperplasia of chloride cells at the base of the gill filament and secondary lamellae were common observations in both concentrations in at 10, 20 and 30 days. After 10 days exposure of 0.85mg/l concentration to fingerlings showed histological alteration in gills like oedema in primary lamella, shortened secondary lamella, lamellar fusion, epithelial lifting and mild degeneration of central axis (Fig.1/B). After 20 days erosion of secondary lamella and mild hemorrhage in primary lamella were started also shown oedema in primary lamella, shortened secondary lamella, lamellar fusion (Fig.1/C). After 30 days showed aneurysm, complete erosion of secondary lamella and hemorrhage in primary lamella (Fig.1/D).

At 1.7 mg/l of concentrations, fingerling showed Oedema of the primary gill lamella, disorganization of central axis and epithelial lifting along with erosion of secondary lamella were observed during 10 days of exposure (Fig.2/B). Severe haemorrhage at the axis of the primary gill lamellae, shortened, erosion and curling of secondary lamella along with the minute epithelial lifting from the lamellae and aneurysm were observed after 20 days of exposure (Fig.2/C). In some portions, the gills also showed aneurysm and degeneration of central axis disorganization of the tissue and complete erosion of secondary gill lamellae and mild hemorrhagic tip of the primary gill lamellae at the end of exposures at 30 days (Fig.2/D).

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**Figure 1.** Effect of AR 97 exposed to 0.85mg/l on the structure of gills on *Catla catla* by photomicrograph [A] Control: Normal gill (a) primary lamella (b) secondary lamella (c) central axis [B] 10 days (a) oedema in primary lamella (b) shorten secondary lamella (c) lamellar fusion (d) mild degeneration of central axis (e) epithelial lifting [C] 20days (a) oedema in primary lamella (b) shorten secondary lamella (c) lamellar fusion (d) erosion of secondary lamella (e) hemorrhagic primary lamella [D] 30days (a) aneurysm (b) complete erosion of secondary lamella (c) hemorrhagic primary lamella
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Figure 2. Effect of AR 97 exposed to 1.7mg/l on the structure of gills on *Catla catla* by photomicrograph[A] Control: Normal gill (a) primary lamella (b) secondary lamella (c) central axis [B] for 10days (a) oedema in primary lamella (b) disorganization of central axis(c) epithelial lifting (d) erosion of secondary lamella [C] for 20days (a) oedema in primary lamella (b) shorten secondary lamella (c) lamellar fusion (d) erosion of secondary lamella (e) curling of secondary lamella (f) hemorrhagic primary lamella [D] for 30days (a) aneurysm (b) complete erosion of secondary lamella (c) hemorrhagic primary lamella (d) degeneration of central axis (e) epithelial lifting

4. Discussion

Results obtained from this study show significantly decreased Hb content and RBC count as compared to control. On the other hand, WBC count increased in all AR 97 exposed fingerlings. It clearly indicates the toxic potential of AR 97 even at sublethal concentrations that affect the haematopoiesis of exposed fingerlings. All mentioned alterations indicate that exposed fingerlings suffered from anemia induced by dye. This is an indication of disruptive effects of azo dyes on erythropoietic tissues as well as cells viability [19]. Our previous study also supports the results of the present study [20]. Significant decrease in haemoglobin has been reported in Clarias batrachus by Patnaik and Patrea exposed to sublethal concentration of propoxur and carbaryl [21]. Similar decrease found in Carassius auratus gibelio when they were exposed to the toxic textile dye [22]. Changes in haematological parameters of Heteropneustes fossilis has been supported our results when the exposed to galvanizing industry effluent [23].

When fingerling *Catla catla* were exposed to dye AR 97 total count of RBC’s reduced compared to control. Alterations in RBC count exposed to various toxicants have been reported by many researchers [24,25]. Significant reduction in RBC count was noted in cypermethrin treated *Labeo rohita* [26] and in freshwater *Cyprinus carpio* treated with diazinon [27]. Decrease in the value of Hb, PCV, RBCs may results in hypochromic microcytic anemia which may be due to deficiency of iron and its decreased utility [28]. In the present study, WBC count increased following exposed dye AR 97. Significant increase was also observed in the total WBC count after exposure to malachite green and Pyceze [29]. WBC plays a major role in the defense mechanism of fish. In other words, an immediate activation of the fish immune system is proved by increase in leucocytes [30]. A rise in WBC count following exposure to insecticides has been also reported [31].

Red blood cells indices MCV, MCH, and MCHC are often determined as an index of health status especially in aquatic organisms [32]. In the present study, rise in MCV values seems to be correlated with decline in RBC count. A significant increase in MCV values was also observed in malachite green exposed fish [33,34]. Likewise significantly increased MCHC and MCH level was also noticed in European catfish (*Silurus glanis L.*) exposed to organophosphorous insecticide, diazinon and koprucu et al. in *Salmo salar* exposed to endosulfan [35,36].

The gills are among the most vulnerable structures of the teleost fish because of their external location and intimate contact with the water. So, they are liable to damage by any irritant materials whether dissolved or suspended in the water [18]. Oedema and epithelial lifting are the results of defense mechanism. As the distance of the lamellar epithelium increases from the secondary gill lamellae, it prevents the direct diffusion of the toxicant present in the water body. These kinds of histological
changes in the gills have been observed by researchers due to toxic effects of copper [37].

The gills, which participate in many important functions in the fish, such as respiration, osmoregulation and excretion, remain in close contact with the external environment and particularly sensitive to changes in the quality of the water are considered the primary target of the contaminants [38]. Haemorrhage in gills was also observed in Gambusia affinis and Clarias gariepinus exposed to textile effluents and herbicide [39,40]. Ribeiro et al. observed proliferation of epithelial cells and fusion between the secondary lamellae in Salvelinus alpinus [41].

Most common changes like epithelial lifting, oedema, lamellar fusion, collapsed secondary lamellae and curling of secondary lamellae have been reported by Velmurugan et al. and Jirauungkoorskul et al. in Cirrhinus mrigala and Oreochromis niloticus exposed to pesticide and herbicide [42,43].

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