Alterations in histopathological features and brain acetylcholinesterase activity in stinging catfish *Heteropneustes fossilis* exposed to polluted river water

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**Abstract**

Responses of stinging catfish (*Heteropneustes fossilis*) to pollution were studied in three freshwater rivers, namely Buriganga, Turag, and Shitalakkhya (Dhaka, Bangladesh), which are potentially affected by anthropogenic pollution originating from industrial and sewage dumping. Partial parameters about water quality (temperature, dissolved oxygen, and pH) and seasonal plankton fluctuation were recorded at wet and dry seasons. Histopathology and acetylcholinesterase (AChE) activity were used as biomarkers to assess water toxic effects in 7- and 10-day exposures of *H. fossilis* to three rivers waters, respectively. The lowest level of dissolved oxygen was recorded as $0.7 \pm 0.1$ mg/l, and the lowest count of plankton genera was 21 at lean period. Furthermore, the 7-day exposure of fish to polluted water abruptly altered the normal structure of various organs. Major structural damages were partial and total epidermal loss, dermis and muscle separation, melanin pigment and vacuole in skin muscle; missing of lamellae, clubbing, fungal granuloma, hyperplasia and hemorrhage in gills; hyperplasia, hemorrhage, pyknosis, vacuole, necrosis, nuclear alteration, fatty degeneration, lipid droplets in liver; degenerating glomerular and tubule, hemorrhage, pyknosis and vacuole in kidneys; and scattered spermatozoa and prominent interstitial space in the testis. After subsequent exposure to polluted water, a significant ($P < 0.05$) inhibition of AChE activity in the fish brain was observed with the following order of potency: $102.00 \pm 5.00$ nmol/min/mg protein (Turag) $\geq 104.00 \pm 5.00$ nmol/min/mg protein (Buriganga) $>$ $130.67 \pm 3.51$ nmol/min/mg protein (Shitalakkhya). This study confirmed the utility of biomarkers in biomonitoring studies and reflected the potential hazards of pollution to aquatic biota.

**Keywords:** Acetylcholinesterase activity, *Heteropneustes fossilis*, Histopathology, River pollution

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**Background**

River pollution is one of the recently focused environmental issues where the most attention is drawn to the rivers and canal systems surrounding Dhaka, Bangladesh. Rapid and unplanned urbanization and industrialization centering this area are increasing at an alarming rate. Due to this situation, the major three rivers, Buriganga, Turag, and Shitalakshya, surrounding Dhaka have been steadily experiencing complicated problems of pollution and encroachment that have almost suffocated these valuable lifelines of the city. The presence of pollutants in the environment is partially due to natural processes but mainly as a result of industrial waste. The polluting industries surrounding the capital are mainly concentrated at Hazaribag and Lalbag (Old Dhaka), Tongi (Gazipur), and Demra (Narayanganj) that dispose their huge untreated effluents directly into these three rivers. These rivers specially the Buriganga are also loaded by sewage pollution. However, high concentration of suspended and dissolved solids has also been reported; they previously occurred at low concentration and are now found in high concentrations (Zakir et al. 2006; Mohiuddin et al. 2011).

Occurrence of potential toxicants in aquatic ecosystem causes a reduction in the quality of the aquatic environment that results in impaired level of dissolved oxygen (DO), pH, temperature, biological oxygen demand, and chemical oxygen demand (Roberts 2001). Adverse water quality, moreover, makes the aquatic habitat biologically dead. Availability of a plankton community is an indicator of water quality. Ferdous and Muktadir (2009) described the high potentialities of zooplankton as a bioindicator. However, evaluations on reduced bioavailability of plankton that resulted from pollution have been made earlier (Begum and Khanam 2009; Shah et al. 2008; Begum 2008; Solomon et al. 2009; Sharma et al. 2010).

Though chemical monitoring of water and sediment is a common and reliable measure to describe the degree of contamination, it is not the case for the overall assessment for evaluating the effects of pollution on the environment as toxic or biological effects on organisms cannot be obtained by this method. Recently, different types of biomarkers have been studied and evaluated for their acceptability to detect the biological effects as a biomonitoring tool (Amiard et al. 2006; Magni et al. 2006; Nigro et al. 2006).

Histopathological assessment is a sensitive biomonitoring tool in toxicant impact assessment to indicate the effect of toxicants on fish health in polluted aquatic ecosystems. Histopathological assessment of fish tissue allows for early warning signs of disease and detection of long-term injury in cells, tissues, or organs. Structural changes in various tissues into the polluted ecosystem have also been acknowledged (Peuranen et al. 2000; Marchand et al. 2009). Earlier histopathological assessments of fish exposed to a variety of pollutants reveal the potency of this biomarker against pollution.

Acetylcholinesterase (AChE) is the main cholinesterasic form in all invertebrate and vertebrate tissues such as the brain (Rodrigues et al. 2011), muscles, blood cells, and liver (Valbonesi et al. 2011). This enzyme is found at neuromuscular junctions and cholinergic nervous system where its activity serves to terminate synaptic transmission. It degrades (through its hydrolytic activity) the neurotransmitter acetylcholine, producing choline and an acetate group in both vertebrates and invertebrates (Varo et al. 2008). Cholinesterase inhibitors such as organophosphate and carbamate block the function of AChE and thus provoke excess acetylcholine accumulation in the synaptic cleft that
eventually causes neuromuscular paralysis, leading to death by asphyxiation (Nunes et al. 2003; Purves et al. 2008; Xuereb et al. 2009). Recent studies have shown that AChE is a very useful biomarker of pollution stress under a variety of environmental factors and chemical mixtures in different geographical regions (Baršienė et al. 2006; Kopecka et al. 2006; Schiedek et al. 2006).

In spite of having a lot of deleterious effects of pollution, a limited number of research were conducted in Bangladesh to ascertain their effects on fish at cellular and molecular level. To our knowledge, limited investigations have been made till now to assess the neurotoxic effects (in terms of AChE inhibition) of river pollutants on fish, thus indicating the need for due attention on this matter. Thus, the purpose of the present study is to assess the cellular and neurotoxic effects of river pollution on *Heteropneustes fossilis* through the determination of potency of two biomarkers against pollution.

**Methods**

**Study area**

Buriganga (near Lalbag, Dhaka), Turag (near Tongi Bridge, Gazipur), and Shitalakkhya (near Demra, Narayangang) are supposed to derive massive pollutant loadings from sewage and industrial effluents directly as industries, textiles, pharmaceuticals, and tanneries have clustered here. During the rainy season, the water quality improves moderately, but on the advent of the dry season, pollution concentration increases abruptly because the water level of the rivers reduces a lot at this time, but the rate of pollutants released into the rivers remains identical. Location of the three rivers and sampling sites are shown in Figure 1.

**Water quality parameter measurements**

Some water quality parameters were measured from the three rivers at two different seasons (early September 2010 and early March 2011). A mercury centigrade thermometer was used to measure the water temperature; pH and DO were measured by a pH meter (HI 98127, HANNA Instruments, Beijing, China) and a dissolved oxygen meter (DO-5509, Tyner, Dongguan City Electronic Technology Co., Ltd., Guangzhou, China), respectively. Each parameter was recorded at two different points with three replications in the same river.

**Plankton collection and identification**

With a view to record the seasonal plankton availability, samples were collected in early September 2010 and early March 2011. Planktons were collected randomly from the three rivers by plankton net towing. Collected samples were immediately preserved with 5% buffered formalin in separate tagged plastic bottles. For species identification, the bottle containing plankton samples was gently shaken to resuspend all materials, was poured on water a petri dish, and was allowed to settle for a minute. Two drops of water were placed on a glass slide and covered with a cover slip. The planktons were then identified up to genus level under a compound microscope (OPTIKA B-350, OPTIKA Microscopes, Ponteranica, Bergamo, Italy) according to APHA (1992), Bellinger (1992), and Palmer (1980).
Histopathological study

To assess the effects of polluted water exposure to fish, water samples were separately collected from Buriganga, Turag, and Shitalakkhya rivers in six plastic containers (30 l) and were carried out to the wet laboratory of the Faculty of Fisheries, Bangladesh Agricultural University, Mymensingh. Experiments were conducted in nine aquaria of size $48 \times 26 \times 30$ cm$^3$. Water samples from the three rivers were kept in three different aquaria, each having two replications, whereas one aquarium was kept as control. *H. fossilis* with average size and weight of $12 \pm 1.4$ cm and $10.5 \pm 1.2$ g, respectively, were collected from a local fish market in live condition. Forty-five fishes (five fishes in each aquarium) were exposed to three river water samples for a period of 7 days with continuous aeration. After the 7-day exposure, two fishes were taken from each aquarium and sacrificed. Gills, skin muscle, liver, kidney, and gonads were collected and
preserved in 10% neutral-buffered formalin, while gonads were preserved in Bouin's fluid. The preserved samples were then dehydrated, cleaned and infiltrated in an automatic tissue processor (ThermoFisher Scientific, Waltham, MA, USA), embedded in melted paraffin wax, and sectioned (5 μm) with a microtome machine (Leica Junc 2035, Leica Microsystems Srl, Milan, Italy). Thereafter, the sections were stained with hematoxylin and eosin (H and E) stains. After staining, the sections were mounted with Canada balsam and kept overnight for the permanent slide. Photomicrography of the stained samples was done using a photomicroscope (OPTIKA B-350). The extent of alteration was scored as severe (+++), moderate (++), mild (+), and not found (−). When a pathology occurred in >50% cell or area in maximum investigated slides, it scored severe (+++), followed by >25% for moderate (++), and <25% for mild (+).

**AChE activity measurement**

For the analysis of AChE activity, *H. fossilis* was exposed to river water in glass aquaria for 10 days. Fish exposed to pollutant-free water was kept as control. Following exposure, three fishes were taken from each aquarium (n = 9) for each river. The whole brain was dissected out by sacrificing the fish and was placed in ice-cold 0.1-M sodium phosphate buffer (pH 8.0). In this study, where brain sample was used similarly with teleost, AChE is maximally distributed in the brain (Kopecka et al. 2004; Ferenczy et al. 1997). Tissues were then weighted and homogenized using a glass-Teflon homogenizer in a homogenization buffer (0.1-M sodium phosphate buffer, 0.1% Triton X-100, pH 8.0) to achieve the final concentration of 20 mg tissue/ml phosphate buffer. Tissue homogenate was centrifuged at 10,000 × g for 15 min at 4°C, and the supernatant was removed. An aliquot of supernatant was then removed and measured for protein according to the method of Lowry et al. (1951) using bovine serum albumin in homogenization buffer as a standard.

AChE activity in the fish brain was measured according to the method of Ellman et al. (1961), as optimized by Habig et al. (1988) and Sandahl and Jenkins (2002). Tissue homogenate (50 μl) was added to 900 μl of cold sodium phosphate buffer (0.1 M containing 0.1% Triton X-100, pH 8.0) and 50 μl of 5,5-dithiobis (2-nitrobenzoic acid) (6 mM), then vortexed, and allowed to stand at room temperature for 10 min. Aliquots of 200 μl in triplicate were then placed into microtiter plate wells. The reaction was started with the addition of 50 μl of acetylthiocholine iodide (15 mM) specific for fish (Jash et al. 1982). Changes in absorbance were measured with a microplate reader (SpectraMax 340PC, Molecular Devices LLC, Sunnyvale, CA, USA) at 412 nm for 10 min at 12-s intervals. AChE activity is expressed as nanomole per minute per milligram protein.

**Statistical analysis**

Data were analyzed using one way analysis of variance and expressed as mean ± SD. A post hoc Waller Duncan multiple test range was performed which considered a 5% significant level using SPSS ver. 11.5 computer software program.

**Results and discussion**

**Water quality parameters**

Water quality parameters of the three rivers in two different seasons are presented in Table 1. The temperature in the wet season ranged from 22.5°C to 24.5°C where it
increased to 29°C to 31.5°C in the dry season. Significant reduction in DO level was observed in dry season where Turag River had the lowest DO level (0.7 mg/l). No wide fluctuation in pH level was observed at two different seasons that ranged from 6.9 to 7.2.

Variation in water quality parameters is mainly due to seasonal environmental factors. Increased temperature in the dry season affects the DO level as they are inversely correlated. In the study, very low DO levels (0.7 to 1.9 mg/l) were recorded in the dry season where >5 mg/l is recommended for biological organisms. This anoxic condition especially in Turag reflects the breakdown of untreated organic waste principally received from domestic sewage and chemical residues from various industries surrounding Dhaka (The World Bank 2006). A similar phenomenon was previously reported from these rivers (Begum 2008; Begum and Khanam 2009; The World Bank 2006; Saha et al. 2009).

Plankton composition

The plankton communities of the three rivers were identified at rainy and dry seasons. A total of 33, 26, and 35 genera were identified from Buriganga, Turag, and Shitalakkhya rivers, respectively in the rainy season, whereas, in the dry season, a lesser number of genera were recorded:21, 19, and 26, from Buriganga, Turag, and Shitalakkhya rivers, respectively (Table 2). Among the recorded phytoplankton groups, Bacillariophyceae were found in both seasons in the respective rivers, and Chlorophyceae were recorded more only in the wet season. However, among the recorded zooplankton groups, Cladocera occurred more frequently in the wet season than in the dry season. Frequent abundance of Copepoda represented by two genera was observed from the three rivers in both seasons. Rotifera were recorded low, but the lowest count was made in the dry season; no genera were even found in the dry season from Turag River.

Seasonal variations in plankton are related to a variety of environmental factors in aquatic environments where temperature has been claimed to be the major determining factor in phytoplankton growth and development (Çetin and Şen 2004; Baquero et al. 2006). The less abundance of plankton communities in the dry season mainly due to high temperature and low DO. The occurrence of Chlorophyceae only in the wet season and Bacillariophyceae in both two seasons indicates that Chlorophyceae are more sensitive to pollutant discharge, whereas Bacillariophyceae seem to be very well adapted to polluted zone (Begum and Khanam 2009; Shah et al. 2008). Among the
| Plankton        | Class               | Species          | Wet season | Dry season |
|-----------------|---------------------|------------------|------------|------------|
|                 |                     | Buriganga  | Turag  | Shitalakkhya | Buriganga  | Turag  | Shitalakkhya |
| Phytoplankton   | Bacillariophyceae   | Fragilaria sp.  | ✓        | X          | ✓        | X      | ✓          |
|                 |                     | Cyclotella sp.  | ✓        | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Cymbella sp.    | ✓        | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Gamphonoema sp. | ✓        | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Gyrosigma sp.   | ✓        | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Melosira sp.    | X         | X          | ✓        | ✓      | ✓          |
|                 |                     | Navicula sp.    | ✓        | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Nitzchia sp.    | ✓        | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Synedra sp.     | X         | X          | X        | X      | X          |
| Chlorophyceae   |                     | Ankistrodesmus sp. | ✓        | X          | ✓        | X      | ✓          |
|                 |                     | Chara sp.       | ✓          | ✓          | X        | X      | X          |
|                 |                     | Chlamydomonas sp.| ✓        | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Chlorella sp.   | X         | ✓          | ✓        | ✓      | ✓          |
|                 |                     | Chlorogonium sp.| ✓         | ✓          | X        | X      | X          |
|                 |                     | Cosmarium sp.   | ✓         | X          | X        | X      | X          |
|                 |                     | Eudorina sp.    | X         | X          | ✓        | X      | X          |
|                 |                     | Microspora sp.  | X         | X          | X        | X      | X          |
|                 |                     | Oedagonium sp.  | X         | ✓          | ✓        | X      | X          |
|                 |                     | Pediatrum sp.   | ✓         | X          | X        | X      | X          |
|                 |                     | Pyrobatrix sp.  | X         | X          | X        | X      | X          |
|                 |                     | Scenedesmus sp. | ✓         | X          | ✓        | X      | ✓          |
| Phytoplankton | Buriganga | Turag | Shitalakkhya |
|---------------|-----------|-------|--------------|
| Spirogyra sp. | ✓         | ✓     | ✓            |
| Spirulina sp. | ✓         | ✓     | ✓            |
| Ulothrix sp.  | ✓         | ✓     | ✕            |
| Volvox sp.    | ✓         | ✓     | ✓            |
| Zygnema sp.   | ✕         | ✕     | ✓            |
| Cryptophyceae | ✓         | ✕     | ✕            |
| Cryptomonas sp.| ✕         | ✓     | ✕            |
| Chrysophyceae | ✓         | ✕     | ✕            |
| Mallomonas sp.| ✕         | ✕     | ✓            |
| Cyanophyceae  | ✓         | ✓     | ✓            |
| Anabaena sp.  | ✕         | ✕     | ✓            |
| Aphanizomenon sp.| ✕   | ✕     | ✓            |
| Chlorella sp. | ✓         | ✕     | ✓            |
| Gomphosphaeria sp.| ✕ | ✕     | ✓            |
| Microcystis sp.| ✕         | ✕     | ✕            |
| Oscillatoria sp.| ✕       | ✓     | ✓            |
| Phormidium sp.| ✓         | ✓     | ✓            |
| Euglenophyceae| ✓         | ✓     | ✓            |
| Euglena sp.   | ✕         | ✓     | ✓            |
| Phacus sp.    | ✓         | ✓     | ✓            |
| Trachelomonas sp.| ✕ | ✓     | ✓            |
| Zooplankton   | Bosmina sp.| ✓     | ✓            |
| Cladocera     | Daphnia sp.| ✓     | ✓            |
|               | Diaphanosoma sp.| ✓ | ✓     | ✓            |
|               | Maina sp.  | ✓     | ✓            |
|               | Nauplius   | ✓     | ✓            |
| Copepoda      | Cyclops sp.| ✓     | ✓            |

**Table 2 List of plankton recorded from Buriganga, Turag, and Shitalakkhya rivers at wet and dry seasons** (Continued)
| Rotifera          | Diaptomus sp. | ✓ | ✓ | ✗ | ✓ | ✓ | ✗ |
|------------------|---------------|---|---|---|---|---|---|
|                  | Brachionus sp. | ✓ | ✗ | ✗ | ✓ | ✗ | ✗ |
|                  | Filinia sp.    | ✗ | ✗ | ✗ | ✗ | ✗ | ✗ |
|                  | Keratella sp.  | ✓ | ✗ | ✗ | ✗ | ✗ | ✗ |
| Total            |               | 33| 26| 35| 21| 19| 26|

Table 2 List of plankton recorded from Buriganga, Turag, and Shitalakkhya rivers at wet and dry seasons (Continued)
zooplankton, the members of Cladocera occurred in lower number in the dry season because Cladocera are highly responsive against pollutants; they even react against low concentration of contaminants (Ferdous and Muktadir 2009). Copepoda were represented by two genera and appeared all year round as this group is much more tolerant to O2 deficiency. On the contrary, lower representation by the members Rotifera in the dry season and no genera in Turag is opposed to the studies of Solomon et al. (Solomon et al. 2009) and Sharma et al. (2010) who reported the dominance of Rotifera among all zooplankton. This is because the frequency of that group is lower in polluted water than in unpolluted or lower polluted zones (Eloranta 1980).

**Histopathological observations**

After subsequent exposure of *H. fossilis* to Buriganga, Turag, and Shitalakkhya river water samples, histopathological assessment of skin muscle, gills, liver, kidney, and testis was made by comparing them with the control. Mild to severe alterations in the different organs were assessed (Table 3).

The skin muscle of the control group was in a systemic arrangement of epidermis, dermis, and muscle (Figure 2a). The major pathological signs observed in skin muscle of fish exposed to the water of three different rivers were partial loss of epidermis, totally missing epidermis, separation of dermis from epidermis, separation of muscle from dermis, melanin pigment, and vacuole in muscle and dermis (Figure 2b,c,d). As skin muscle is the primary site of exposure, pollutants affected the epidermis abruptly. Melanin pigment is a prominent feature of chronic inflammatory response.

Gills are the primary site for any histological alteration as it is directly exposed to polluted water. In our present study, the structure of gills in the control group was almost normal. Primary and secondary gill lamellae were found with no pathology (Figure 2e). Moderate to severe structural changes with mentionable pathological signs were observed in the gills of treated fish including missing secondary gill lamellae, hemorrhage, necrosis, hyperplasia and hypertrophy, gill clubbing, and fungal granuloma (Figure 2f,g,h). Disruption in gill structure and function possibly due to various environmental factors, pH, ion concentration, heavy metals, and other pollutants were previously described (Tkatcheva et al. 2004; Peuranen et al. 2000; Playle 1998). Thickening and lifting of the secondary lamellar epithelium due to hypertrophy are the first signs that gills have been exposed to hazardous chemicals, or physical agents may have been a response to increase the diffusion distance between DO and blood, which accordingly was related to hypoxia in fish (Liu et al. 2010). Gill clubbing is due to excess mucus production. In the presence of pollutants, the epithelium of the secondary lamellae has a tendency to increase the number of mucus cell. Excess mucus from mucus cell causes the fusion of secondary gill lamellae resulting in impaired respiration.

Alteration in the liver structure may be used as a biomarker indicating prior exposure to environmental stressors. A constant exposure to toxicants may cause damage to the liver tissue (Nero et al. 2005). In the present study, hepatocytes and other cells were systematically arranged and no structural alteration was assessed in the liver of the control group (Figure 3a) where the treated group revealed a normal liver tissue structure with severe alterations like deposition of body fat, hypertrophy and hyperplasia of hepatocytes, rupture of blood vessel resulting in hemorrhagic area and necrosis, nuclear
alteration, pyknosis, vacuoles, and fatty degeneration (Figure 3b,c,d). A similar assessment was made by Liao et al. (2007) who additionally found basophilic cytoplasm but no lipid droplets in the hepatocytes of medaka (Oryzias latipes) exposed to single mercury and single selenium. Farag et al. (2006) observed lipid droplets in the liver of Chinook salmon (Oncorhynchus tshawytscha) after chronic chromium exposure, whereas livers of the control group had no hepatocellular lipids. Liver necrosis was found in Nile tilapia (Oreochromis niloticus) after exposure to sediment containing a variety of organic chemicals (Perez et al. 2000). Shaw and Handy (2006) found marked hepatic lipidosis (increased intracellular fat stores) in Nile tilapia (O. niloticus). Histopathological alterations in the liver of Clarias gariepinus from polluted aquatic systems were also found by Marchand et al. (2009).

This could be expected as the liver is the main detoxification organ involved in the metabolism and excretion of heavy metals. It is therefore a target organ of various toxic substances. Pollutants cause alterations in the liver, lysis of hepatocytes resulting in

| Histopathological alterations in organ | Extent of alteration |
|--------------------------------------|---------------------|
|                                       | Buriganga | Turag | Shitalakhyha |
| Skin muscle                          | Melanin pigment | +++  | +   | +   |
| Vacuoles                             | –        | ++   | +   |
| Epidermal loss                       | +        | ++   | +++ |
| Muscle separation                    | ++       | +    | +++ |
| Gill                                 | Hemorrhage | +++  | +++ | +   |
| Necrosis                             | ++       | –    | –   |
| Gill clubbing                        | ++       | ++   | +   |
| Fungal granuloma                     | +++      | +++  | +   |
| Missing of secondary gill lamellae   | +++      | ++   | +   |
| Hyperplasia and hypertrophy          | ++       | ++   | +   |
| Liver                                | Hemorrhage | +++  | +++ | +   |
| Pyknosis                             | ++       | ++   | +   |
| Vacuole                              | +        | +    | ++  |
| Necrosis                             | +        | ++   | +   |
| Fatty degeneration                   | ++       | ++   | +   |
| Lipid droplet                        | –        | ++   | –   |
| Nuclear alteration                   | –        | –    | +   |
| Hyperplasia and hypertrophy          | –        | –    | +   |
| Kidney                               | Degenerated kidney tubule | +   | +   | +   |
| Hemorrhage                           | +        | +    | –   |
| Vacuole                              | ++       | ++   | –   |
| Degenerated glomerular tubule        | –        | +    | –   |
| Testis                               | Scattered spermatozoa | ++  | +   | +   |
| Vacuole                              | ++       | +    | ++  |
| Testis interstitial space            | +++      | +++  | +   |

| Extent of alteration: severe (+++), moderate (++), mild (+), not found (−).
Figure 2 Photomicrograph of skin muscle and gills of *H. fossilis* from control and from the three rivers. Photomicrograph of skin muscle from (a) control, (b) Buriganga, (c) Turag, and (d) Shitalakkhya. Photomicrograph of gills from (e) control, (f) Buriganga, (g) Turag, and (h) Shitalakkhya. Ed, epidermis; D, dermis; M, muscle; Mp, melanin pigment; V, vacuole; PEL, partial epidermal loss; TEL, total epidermal loss; Ms, muscle separation; H, hemorrhage; N, necrosis; Gc, gill clubbing; Fg, fungal granuloma; Msgl, missing of secondary gill lamellae; HP, hyperplasia and hypertrophy (H and E × 430).
Figure 3 Photomicrograph of liver and kidney of *H. fossilis* from control and from the three rivers. Photomicrograph of liver from (a) control, (b) Buriganga, (c) Turag, and (d) Shitalakkhya. Photomicrograph of kidney from (e) control, (f) Buriganga, (g) Turag, and (h) Shitalakkhya. H, hemorrhage; Fd, fatty degeneration; Ld, lipid droplet; N, necrosis; P, pyknosis; Na, nuclear alteration; Hp, hyperplasia and hypertrophy; Kt, kidney tubule; Gt, glomerular tubule; Dkt, degenerating kidney tubule; Dgt, degenerating glomerular tubule; V, vacuole (H and E x 430).
necrosis, breakdown of blood vessel resulting in a hemorrhagic area, cirrhosis, and ultimately death. Hepatic lipid accumulation is influenced by toxins and environmental stress factors (Tanaka et al. 2002; Xu et al. 2009).

The kidney is a major organ involved in fluid and ionic balance of fish. In the present study, the normal structures of the kidney tubule, glomerulus, and other hematopoietic cells of kidney were found in a reference group (Figure 3e). Mild structural changes were observed including vacuolation, necrosis, mild pyknosis, and mild tubular degeneration from the kidney of fish exposed to different river water (Figure 3f,g,h). However, necrosis, fibrosis (scarring) and dilation of tubular lumina, and lipid peroxidation were observed in affected kidneys as a result of chromium exposure, but not in reference kidneys (Farag et al. 2006) as the exposure time was longer (105 days). The appearance of vacuoles and tubular degeneration might be a result of a general metabolic disturbance leading to an intensified reabsorption of amino acids and small proteins from the ultrafiltrate (1993). Necrosis that resulted from accumulation of granules is due to karyolysis and vacuolation in the cytoplasm (Rangsayatorn et al. 2004).

The immature testes of the control fish possess a normal arrangement of spermatoocytes in the testicular lumen and a smaller testis interstitial space (Figure 4a). Some structural alterations were observed in the testes of treated fish such as the appearance of some vacuoles, scattered spermatoocytes in the testicular lumen, and larger testis interstitial space (Figure 4b,c,d). Similar findings, except for vacuoles and scattered spermatoocytes, were also made earlier by Schultz et al. (2011) in male fathead minnows exposed to antidepressant pharmaceuticals. Intersex gonads were recorded in several species exposed to estrogen-polluted water source (Barnhoorn et al. 2004, Barnhoorn et al. 2010). However, no intersex testis was observed in the present study. This may be due to less exposure time and sexual immaturity of the fish. A larger testis interstitial space from all treated groups indicated the presence of toxicants in the three reference rivers. These prominent spaces are due to the accumulation of testis interstitial fluid which interrupts the normal functioning of luteinizing hormone.

Figure 4 Photomicrograph of the testis of *H. fossilis* from control and from the three rivers. (a) Control, (b) Buriganga, (c) Turag, and (d) Shitalakkhya. V, vacuole; Sz, spermatoza; Ssz, scattered spermatoza; Tis, testis interstitial space (H and E × 430).
AChE activity of *H. fossilis* exposed to three river water

*H. fossilis* were exposed to polluted water collected from Buriganga, Turag, and Shitalakkhya for 10 days. After 10 days of exposure time, the AChE activities in the fish brain were determined. In the present study, a steady decrease of AChE activity in the fish brain was found in the case of *H. fossilis* exposed to water samples of the three rivers (Figure 5). In the control fish, AChE activities were measured as 202.67 ± 6.51 nmol/min/mg protein. Furthermore, AChE activities in the brain of *H. fossilis* exposed to water samples of Buriganga, Turag, and Shitalakkhya rivers were 104.00 ± 5.00, 102.00 ± 5.00, and 130.67 ± 3.51 nmol/min/mg protein, respectively. Significant inhibition of AChE activity (*P* < 0.05) was observed from all river water samples compared to the control. In this study, it was observed that due to polluted water exposure, AChE activity decreased values were 48.78%, 49.63%, and 35.56% from Buriganga, Turag, and Shitalakkhya river water samples, respectively.

AChE activity in vertebrates and invertebrates is a widely used biomarker of neurotoxicity. Moreover, fish AChE activity has been utilized as a highly sensitive biomarker response of freshwater pollution (de la Torre et al. 2002). A significant decrease in this cholinesterase activity in fish indicates the presence of neurotoxic pollutants in the aquatic ecosystem. In the present study, *H. fossilis* were exposed to polluted water collected from Buriganga, Turag, and Shitalakkhya for 10 days. After 10 days of exposure time, the AChE activities in the fish brain were determined. In the present study, a steady decrease of AChE activity in the fish brain was found in the case of *H. fossilis* exposed to water samples of the three rivers (Figure 5). In the control fish, the AChE activities were measured as 202.67 ± 6.51 nmol/min/mg protein. Furthermore, AChE activities in the brain of *H. fossilis* exposed to water samples of Buriganga, Turag, and Shitalakkhya were 104.00 ± 5.00, 102.00 ± 5.00, and 130.67 ± 3.51 nmol/min/mg protein, respectively. Significant inhibition of AChE activity (*P* < 0.05) was observed from all river water samples compared to the control. In this study, it was observed that due to polluted water exposure, AChE activity decreased values were 48.78%, 49.63%, and 35.56% from Buriganga, Turag, and Shitalakkhya river water samples, respectively. However, the larger decline of cholinesterase activity in the water samples of Turag and Buriganga than that of Shitalakkhya compared to the control suggests that Turag and Buriganga rivers are more highly polluted with neurotoxic compounds. The extent of

![Figure 5 AChE activity measured in *H. fossilis* exposed to three different river water for 10 days.](http://www.intaquares.com/content/5/1/7)
AChE depression required to cause death in aquatic organisms. In fish, most estimates lie in the range of 70% to 85% AChE reduction. Eel (*Anguilla anguilla*), over a 96-h exposure period to pesticide, were able to survive up to 57% reduction in AChE activity (Sancho and Andreu 1998).

Rank et al. (2007) noted a responding change in AChE activity to pollution in native and transplanted mussels in the areas close to coastal chemical dumping sites with some interpretation due to seasonal fluctuation in exposure situation. According to Forget et al. (2003), neurotoxic contaminants brought by rivers caused a 70% to 80% AChE inhibition in copepods from an estuary in comparison with those from upstream. An ascending order of AChE inhibition with the increased concentration of arsenic in *Channa punctatus* was mentioned by Roy et al. (2006). In contrast, the ameliorative potential of selenium on arsenic-mediated inhibition of AChE activity was also described. Combined *in vivo* and *in vitro* effects of cadmium and aluminum like heavy metal on AChE inhibition have also been reported (Carageorgiou et al. 2004; Kohila et al. 2004).

**Conclusions**

This study showed that in dry season, DO of the three rivers existed below the standard level (≥5 mg/l) and were extremely intolerable for plankton community that eventually resulted in the lowest plankton count. Furthermore, structural damage in fish organs and significant inhibition of AChE in the fish brain suggest the potency of the two biomarkers against pollution. The biomarker data from this study can, in the future, be used to evaluate the effects of management actions. With a view to minimize the devastating effects of pollutants, effluent treatment plants are recommended to establish corporately near the major industrial zone to treat industrial effluents before disposal.

**Abbreviations**

AChE: Acetylcholinesterase; H: Hematoxylin; E: Eosin; DO: Dissolved oxygen.

**Competing interest**

The authors have no competing interests.

**Authors’ contributions**

SFR participated in the design of the study, performed the statistical analysis, and drafted the manuscript. AHMMR has made substantial contributions to the concept and design, acquisition, and interpretation of data. MSH helped in the design of the study and in collection of data. ZH supervised the group to coordinate the research, revised the manuscript critically for important intellectual content, and had given the final approval of the version to be submitted. All authors read and approved the final manuscript.

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