**Abelmoschus moschatus** extract reverses altered pain and neurohistology of a rat with developmental exposure of fluoride

Kurmeti Sudhakar, Mesram Nageshwar, Karnati Pratap Reddy*

Neuroscience lab, Department of Zoology, Osmania University, Hyderabad-500007, Telangana, India.

**ABSTRACT**

Fluoride causes major human health complications and evidence suggests that exposure of fluoride during developmental periods adversely affect neurodevelopment, behavior, and maturity of the brain as these periods are critical for developing stages for CNS. The present study aimed to assess the mechanically induced pain sensitivity, cell morphology, circuitry in terms of neural connections and networks and maturation of neurons under fluoride-induced toxicity with concurrent treatment of protective effects of *Abelmoschus moschatus* seed extract from day 1 of pregnancy to post-natal day (PND) of a rat with age 30th day. Timed pregnant wistar rats (30) were segregated into six groups, viz. control, sodium fluoride (NaF) (20 ppm), NaF + *A. moschatus* aqueous extract (AMAE), NaF + *A. moschatus* ethanolic extract (AMEE), AMAE, and AMEE and treated for 51 days (21 gestational and 30 post-natal days). On postnatal days 1, 7, 14, 21 and 30, rats were assessed for oxidative stress markers (GSH and GSSG) and neurohistology of the brain. Increased threshold levels to the mechanical stimulus were observed in fluoride-treated rats. Brain stained with Congo red, Cresyl violet and Golgi cox for β-amyloid, Nissl substance and synaptic connections respectively showed cells become amyloidosis with decreased Nissl substance and decreased number of neuronal connections in NaF exposed rats. Reduced content of GSH and increased GSSG levels (P < 0.001) were also recorded in NaF treated rats. These alterations were associated with increased production of free radicals and the effect of fluoride on the brain is inversely proportional with age. These changes were ameliorated by supplementation with AMAE and AMEE with anti-oxidant properties, which reduce the production of free radicals from fluoride. Thus, the seed extract of *A. moschatus* had a protective effect over fluoride induced alteration in neural cell maturation, and the establishment of circuitry, mechanical pain sensitivity, and oxidative stress.

**INTRODUCTION**

Studies have reported various effects of chronic fluorosis on the different organ systems, however, only a few have been on the central nervous system in particular on developmental effects of fluoride. NaF exposed rabbits’ brain histological examinations showed the loss of molecular layer, glial cell layer and Purkinje neurons (Shashi, 2003). Reduced or complete loss of Nissl substance was observed in F intoxicated rabbits (Shashi, 2003) and rats (Shivarajashankara et al., 2002). Developing brain is more prone to fluoride toxicity because of poorly formed protective mechanisms such as blood-brain barrier and anti-oxidant defense. F mediates the generation of superoxide anion (O2−) and the supplementary production of hydrogen peroxide, peroxynitrite, and hydroxyl radicals (Hassan and Yousef, 2009). In addition, F affects glutathione levels. This leads to an excessive mitochondrial production of reactive oxygen species (ROS), triggering mutilation to the cellular constituents. The amplified production of ROS causes membrane damage via lipid peroxidation, membrane depolarization, and apoptosis (Zhang et al., 2007). The generation of ROS and lipid peroxidation has been considered to play an important role in the pathogenesis of chronic fluoride toxicity. Recent reports proved that the possible involvement of reactive oxygen species (ROS) in fluoride-induced toxicity (Cao et al., 2015). Oxidative stress has been implicated for its contribution to fluoride associated tissue injury in the liver, kidney, brain, and other organs. Decreased motor abilities,
thermal and mechanically induced pain responses were noticed in previous research contribution of Mesram et al. (2017) and Sudhakar et al. (2017). Neuropathic pain is a major cause of infirmity, patient’s quality of life and interferes with functions relating to CNS (Soler et al., 2007). Different techniques have been used to quantify and assess the development of pain. Among which Randall-Selitto test (Randall and Selitto, 1957), envisioned to serve as a device to evaluate the effect of analgesic agents on the response thresholds to mechanical pressure stimulation (Anseloni et al., 2003).

Prenatal exposure of fluoride results in significant neurochemical alterations in the rats’ brain such as significantly intensified brain LPO level while reduced glutathione (GSH) content and superoxide dismutase (SOD), glutathione peroxidase (GPx), and glutathione reductase (GR) activities were decreased markedly (Basha and Madhusudan, 2010). Fluoride exposed rats displayed various histological changes in the brain, including cell shape, membrane integrity, decreased Nissl granules, de-myelination, a decrease in the number neural cells, neural connections, circuitry and networks, thickening and disappearance of dendrites (Shivarajashankara et al., 2002; Shashi, 2003; Kamel et al., 2010). We have reported neuroprotective effects of an extract of A. moschatus against F induced motor, nociceptive learning behavior and histological change in the brain of adult rat (Sudhakar et al., 2017).

Recently, researchers have put their attention on the protective effect of natural products against neurodegenerative diseases. The administration of curcumin protect from NaF induced neurodegeneration (Nageshwar et al., 2017) and quercetin diminished the neuronal death and reduced the oxidative stress in aluminum-induced neurodegeneration in the rat hippocampus (Sharma et al., 2016). Resveratrol was used against fluoride induced neurodegeneration through oxidative stress (Nalagoni and Karnati, 2016). Saha et al. (2016) explored that the leaf extract of Acacia catechu is having in anti-neurogenic and antioxidant properties. Neuroprotective effect of Ocimum sanctum leaf extract was studied in the transgenic Drosophila model of Parkinson’s disease (Siddique et al., 2014). All the studies are eventually on adults and no reports are noticed in the literature on protective effects of A. moschatus extract during development. In view of this, we have chosen A. moschatus plant to protect against fluoride induced neurodegeneration during gestational and early post-natal exposure as it has anti-oxidants. The plant consists of quercetin, rutin, catechin, epicatechin, and procyanidin which have good antioxidant properties. Procyanidin oligomers were showed greater antioxidant property than vitamins C and E. According to Shui and Peng (2004) quercetin derivatives and (→)-epigallocatechin as major antioxidant compounds in Okra. Quercetin derivatives (quercetin 3-O-xyllosyl (1’’→2”)) glucoside, quercetin 3-O-glucosyl (1’’→6”) glucoside, quercetin 3-O-glucoside, and quercetin 3-O-(6”-O-malonyl)-glucoside) contributes the major antioxidant property (around 70%) of the plant. Different pectic and hemicellulose (xylan, xyloglucan) structures were recently reported in Okra (Sengkhamparn et al., 2009). By the presence of quercetin derivatives and various carbohydrates, Okra extract is involved in eliminating or reducing in the generation of free radicals and thus prevents neurotoxicity from fluoride. The aim of the present study, therefore, is to obtain a reversal in pain response, anti-oxidant nature and histological alterations induced by fluoride ingestion and neuroprotective effects of Okra seed extract against fluoride during developing stages of the nervous system.

MATERIALS AND METHODS

Adult female Wistar rats (three months old, around 250–300 g body weight) were used. The animals (2 females and 1 male in a separate cage) were kept in standard laboratory conditions with 12 h light/dark periods at a temperature of 22 ± 2°C for breeding and supplied with dry rat food and drinking water ad libitum. The rats were acclimatized for one week prior to the commencement of the experiment. All experimental protocols were approved by the Ethics Committee of our institution as per CPCSEA No: 383/01/a/CPCSEA. After rat’s pregnancy was confirmed, from the first-day pregnancy, the females segregated into six groups and allowed to drink on fluoridated water at the rate of 20 ppm NaF and also Abelmoschus moschatus seed extract was given to the rats at the rate of 300 mg kg1 body weight. The plant was identified and authenticated (No. 282) by Taxonomy lab, Department of Botany, Osmania University, Hyderabad, and voucher specimen also submitted for future reference. The plant material (seeds) was extracted with water and ethanol according to the method described in our earlier report (Sudhakar et al., 2017).

Experimental Design

<table>
<thead>
<tr>
<th>Experimental group</th>
<th>Treatment</th>
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<tbody>
<tr>
<td>Group I</td>
<td>Received normal tap water</td>
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<tr>
<td>Group II</td>
<td>Fed on fluoridated drinking water (20 ppm)</td>
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<tr>
<td>Group III</td>
<td>NaF (20 ppm) + Abelmoschus moschatus seed aqueous extract (AMAE) (300 mg kg1 b. wt.)</td>
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<tr>
<td>Group IV</td>
<td>NaF (20 ppm) + Abelmoschus moschatus seed ethanolic extract (AMAE) (300 mg kg1 b. wt.)</td>
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<tr>
<td>Group V</td>
<td>Abelmoschus moschatus seed aqueous extract (AMAE) (300 mg kg1 b. wt.) only</td>
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<tr>
<td>Group VI</td>
<td>Abelmoschus moschatus seed ethanolic extract (AMAE) (300 mg kg1 b. wt.) only</td>
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Behavioral tests

Mechanical hyperalgesia

The mechanically induced nociceptive withdrawal threshold was assessed by using an apparatus called Randall Selitto electronic algesimeter. The basis for this test is threshold response and stimulus mechanism. The apparatus consists of a pinpoint rod which is used to generate stimulus on the dorsal surface of the hind paw. The maximum force applied was limited to 250 gm (calibrated force) to avoid skin damage. Each experimental rat was exposed to Randall Selitto test and noted down the reads as withdrawal response of paw and expressed in pounds (Randall and Selitto, 1957; Santos-Nogueira et al., 2012).
Oxidative stress

Reduced glutathione

The glutathione content in the brain tissue was estimated by the method of Hissin and Hilf (1976). The method takes advantage of the reaction of GSH with ortho-phthalaldehyde (OPT) at pH 8 and of GSSG with OPT at pH 12; GSH can be complexed to N-ethylmaleimide to prevent interference of GSH with measurement of GSSG. GSH reacts with a fluorescent reagent OPT to form a stable, highly fluorescent tricyclic complex which was measured spectrophotometrically at a wavelength of 350 nm and 420 nm.

Oxidized glutathione

Determination of glutathione disulfide (GSSG) in the brain tissue by the method of Hissin and Hilf (1976). At measurement of GSSG, GSH was complexed to N-ethylmaleimide (NEM) to form GS-NEM. This is followed by the back extraction of excess NEM from the sample with dichloromethane. GSSG concentration is measured by spectrophotometry by GSH recycling method. Fluorescence at emission 420 nm was recorded by excitation at 350 nm.

Histology

Congo red stain

Congo red stain is used for the detection of amyloid in neurons. Amyloid refers to the abnormal, fibrous, extracellular proteinaceous deposits found in organs such as brain, liver, kidneys, spleen etc. Congo red dye forms nonpolar hydrogen bonds with amyloid and appeared in red and the nuclei in blue. This condition of deposition of amyloid in tissues is known as Amyloidosis. The tissue thickness of sections was 10 microns and was stained with Congo red and counterstained with Gill’s hematoxylin. The slides studied under Lawrence digital microscope with 4X (Elghetany et al., 1989).

Cresyl violet stain

Cresyl violet staining allows for analysis of neuronal morphology and the neuronal population. The Nissl-staining method is based on the interaction of basic dye, cresyl violet with the nucleic acid content of cells. It can bind to the DNA content of the cell nuclei, but also to the RNA that is highly concentrated in RER and ribosomes (Nissl substance) in the cytoplasm. The Nissl staining can specifically stain the cytoplasm of neurons as it is an active site for protein synthesis. This is an advantage of Nissl staining. The sections about 10 µ were stained with cresyl violet stain, coverslipped with mounting medium (DPX) and observed under the microscope with 40X magnification (Pilati et al., 2008).

Golgi cox stain

Golgi cox stain help us to study of neural cell growth, connections, circuitry, maturation and neuronal networks. Golgi cox staining solution was prepared by mixing of 5% potassium chromate solution, 5% potassium dichromate solution and 5% mercuric chloride solution in equal proportion, brains were preserved in it for 10 days in the dark room. They were washed and preserved in 30% sucrose solution and made into sections about 10 µ thick on vibratome. These sections were passed through hypo, water, and ammonium solution and dehydrated by graded alcohol baths (30%, 50%, 70%, 90% and 100%). The slides were prepared and observed under a digital microscope with 4X (Gibb and Kolb, 1998).

Statistical analysis

Data for all variables were subjected to analyses of variance (ANOVA) to assess the effect of Na-F administered to rats, duration of administration. When ANOVA was significant, a t-test was conducted in all possible combinations of experimental groups for individual comparison. The probability of p < 0.01 and p < 0.001 was considered significant for all evaluations. All data are expressed as mean ± SEM.

Results

Mechanical hyperalgesia

The treated rats have shown significantly reduced response to mechanical pain stimulus compared to control. This effect is higher in day 21 rats than day 30 of fluoride-treated. The simultaneous treatment with AMAE and AMEE along with fluoride-treated rats showed a reverted response to pain stimulus than fluoride alone treated. AMEE showed better efficacy over AMAE.

Reduced glutathione

Fluoride administered rats showed a significant decrease (P < 0.01) in glutathione (GSH) content as compared to control group. The AMAE and AMEE treated rats along with NaF showed the reverted amount of GSH when compared to NaF alone treated. The percent of decrease in the content was progressively decreased from day 1 to day 14 and its content was reduced in decreasing in day 21 and day 30 of the fluoride-treated group.

Oxidized glutathione

Increased GSSG levels were found in NaF administered rats with increasing age from day 1 to day 14 with compared to the respective age of control group. The increase in the GSSG levels from day 21 to day 30 were not significant. In AMAE and AMEE
treated rats against NaF showed decreased levels of GSSG when compared to NaF received rats.

**Congo red stain**

Cells without any protein aggregations and plaques were seen in histologically examined brain of congo red stained sections in control rat brain. While in NaF treated rats showed aggregations which may be amyloid proteins or lead to the development of plaques along with altered cell structure. These kinds of aggregations were found to be very less in number and cell with normal round shape in AMAE and AMEE treated groups compared to F treated individuals.

![GSH Graph](image1)

**Fig. 2:** Protective effects of *A. moschatus* seed extract to NaF exposure effect on GSH content. *p* < 0.01 probability was considered for all groups. Data expressed as mean ± SEM. GSH content was expressed in terms of µg/mg tissue. Data are presented as the mean of 5 animals per treatment.

![GSSG Graph](image2)

**Fig. 3:** Protective effects of *A. moschatus* seed extract towards NaF toxicity. *p* < 0.01 probability was considered for all groups. GSSG levels were expressed in terms of µg/mg tissue. Data expressed as mean ± SEM. Data are presented as the mean of 5 animals per treatment.

**Cresyl violet stain**

Significant pathological changes were noticed in the neurodegenerative assay of cresyl violet stain as NaF group was perceptibly altered neuronal cell morphology, population and decreased Nissl granules (lightly stained cells). These alterations were obviously progressed in NaF treated group from day 1 to 30. These all changes were reverted to oral administration of AMAE and AMEE towards NaF toxicity.

**Golgi Cox**

The analysis of Golgi cox stained neurons from both cortical and sub cortical regions of the brain showed a decrease in the size of the cell body, in a number of branches (dendrites), in length of branches (dendrites and axon), neural circuitry and networks in NaF fed rats with compared to control group. These changes were reported severely on day 1, gradually progressed from day 7 to day 14 and reduced in the alterations could be found in day 21 and 30. All aforesaid alterations were reverted to normal on the treatment of AMAE and AMEE against NaF.

**DISCUSSION**

In the present study, we demonstrated that the neurotoxic effects of NaF on developing rats’ brain and protective effects of * Abelmoschus moschatus* seed extract here off. Free radicals are usually maintained in balance by the body antioxidant defense system, but with an excessive production of free radicals generate oxidative cell damage. This oxidative damage may be due to immobilization of reactive molecules in the brain cells and increases the rate of lipid peroxidation with the release of free radicals inside the neural cells. NaF mainly acts through the generation of excessive free radicals and which leads to the destruction of membrane lipids and overall anti-oxidant status of the brain. The brain is the rich source of polysaturated fatty acids which makes more prone to oxidative stress. Moreover, the developing brain doesn’t have completely developed anti-oxidant system and blood-brain barrier (BBB) which makes them furthermore susceptible to oxidative stress. Dietary antioxidants cooperate with the body enzymes to protect the brain from free radical damage. *A. moschatus* having antioxidants which helps in hepato-protective (Hu et al., 2014), anti-diabetic (Sabitha et al., 2011), antiulcer (Olorunniwa et al., 2013), anticancer (Gu et al., 2011), anti-inflammatory, laxative, anti-hyperlipidemic, anti-fungal and analgesic activities (Anil and Neeraj, 2017). In this study, we conducted mechanical pain response test, oxidative stress markers such as GSH and GSSG and histopathological studies (Congo red stain, cresyl violet stain, and Golgi cox stain) to evaluate NaF neurotoxic effects during pre- and post-natal exposure and as well neuroprotective effects of seed extract of *A. moschatus*.

In NaF treated rats, nociceptors may get damage and/or weak functioning state take more mechanical pressure to respond pain stimulus. This may be because of inhibitory effects of ROS, inhibition of NF-kB which a key role in the reduced function of nociceptors in NaF received rats (Nageshwar et al., 2017), AMAE and AMEE treated rats against NaF showed decreased in the strength of stimulus to respond to Randall Selitto set up. The possible reason for this is seed extract have anti-oxidant properties thus help in the elimination of free radicals from brain which ultimately improves the function of receptors. These reports are similar to that of curcumin as a protective agent towards NaF intoxication (Nageshwar et al., 2017), and also with Okra extract as reported earlier on adult rats (Sudhakar et al., 2017).
Fig. 4: Cerebral cortex region of rat brain stained with Congo red stain. Protective effect of _A. moschatus_ seed extract on rat brain exposed to NaF. Normal cells with round shape and cell membrane, stained with blue colored were found in control rat brain sections which are indicated by a red color arrow mark. Yellow colored arrow mark showing the cells which were with the destructing membrane, swelling and undergoing necrosis were shown in NaF received group of rats. NaF+AMAE and NaF+AMEE treated rat brain sections were found normal (40X Olympus microscope).
Fig. 5: Cresyl violet stained rat brain (cerebral cortex region). Protective effect of A. moschatus seed extract on rat brain exposed to NaF. Normal cells with round shape, healthy cell membrane and clearly visible Nissl granules in the cells. Black colored arrow mark indicating them. Red colored arrow mark showing the cells with the destructing membrane, the altered shape of the cell and decreased granules of Nissl, undergoing necrosis were shown in NaF received group of rats. NaF+AMAE and NaF+AMEE treated rat brain sections were shown as reverted earlier mentioned alterations (40X magnification).
Fig. 6: Photomicrograph of Golgi cox stained sections of rat brain (cerebral cortex and sub cortex). Protective effect of *A. moschatus* seed extract on rat brain exposed to NaF. In control rats, neural cells with a normal size of the cell body, number of branches (dendrites), length of branches (dendrites and axon), neural circuitry and networks were typical whereas they were decreased in NaF fed rats with compared to control group. Black colored arrow mark is showing the typical circuitry and networks. Red color arrow mark is showing the cells which were altered in terms of earlier named characteristics. NaF+AMAE and NaF+AMEE treated rat brain sections were shown in reverted to earlier mentioned alterations (4X magnification).

The GSH:GSSG ratio used as a marker of oxidative stress, which arises due to various distortions. While GSH levels tended to decrease, GSSG levels were increased in NaF fed rats. The effect of NaF on developing brain is progressively increased with increasing age from day 1 to day 14 as they were not with well-developed anti-oxidant and BBB barrier mechanisms.
Decreased in the content of GSH was observed up to 14 days and further reduced decrement was found on day 21 and 30 as they maturing, develop the earlier said protective mechanisms. In NaF+AMAE and NaF+AMEE treated rats content of GSH and levels of GSSG were reverted. In an earlier report of Flora et al. (2009), decreased GSH:GSSG was found in fluoride treated rats. As fluoride is a chemically active ionized element, it may affect oxygen metabolism and induce oxygen free radicals which play a role in diminishing cognitive ability processes such as learning and memory. Moreover, fluoride binds antioxidants in the body such as N-acetyl cysteine (NAC), glutathione (GSH) and other free-radical destroying enzymes thus triggering oxidative stress which leads to cell damage (Anuradha et al., 2000). In the developing brain absence and/or poorly developed a compensatory antioxidant system with the presence of oxidative stress due to increased free radicals plays a great role in the initiation of damage of nerve cells membrane especially via increased lipid peroxidation (Ghiselli et al., 2000; Gao et al., 2009). The increased LPO levels on NaF intoxication were reverted to treatment with Okra extract in developing brain (Sudhakar et al., 2018).

The main histological alterations were seen in NaF treated group are a change in shape of the cell, size of the cell body, amount of Nissl substance, amyloid-like aggregations, branching patterns such as a number of branches arising from the cell body, length of branches, circuitry in terms of connections and networks. Among some of the more severe on day 1 pups, those include shape and size of cell and networks, others are more profound in day 21 and 30 rats and those are aggregations. The brain areas particularly cerebral cortex, cerebellum white matter and ventricles around choroid plexuses were found with hemorrhage in the brain of rats exposed to high NaF for a long duration. Heba et al. (2010) reported neurodegenerative changes in NaF exposed rats, mainly in their pyramidal cells of Ammon’s horn of hippocampus. The pyramidal cells showed atrophy and necrosis. Nerve cells of cerebral cortex showed neurofilaments accumulation in the cytoplasm and the axons, central chromatolysis, edema, and neuronophagia. Hippocampus and cerebral cortex are two major components which involved in learning and memory process (Vianna et al., 2000). These regions are most affected in NaF intoxication and as a result affect the overall behavior of animal (Varner et al., 1998).

In summary, fluoride produces more amount of free radicals which interact with phospholipid layer of neurons. Due to incompletely developed defense system of developing brain, it is unable to provide protection against NaF. This leads to imbalanced oxidative stress. As result neural cells become paralyzed, thus neurodegeneration follows. PND 1 to 14 rats treated with NaF were displayed more pronounced alterations than PND 21 and 30. Because of poorly developed protective mechanisms in early postnatal pups, the F effect is more. When we supplement with AMAE and AMEE, these may remove free radicals and stabilize the normal chemical milieu of the brain. Thus the structure, shape and overall morphology of brain was protected from NaF intoxication. It is also observed that AMEE has better efficacy over AMAE against F neurotoxicity.

CONCLUSION

There are very few studies on developing brain of rats which are prenatally exposed to NaF and continued until the age of post-natal 30 days as well protecting measures with natural agents. The experimental data of this report clearly showed that the developing brain is much sensitive to NaF toxicity as it has no protective mechanisms. The main protective mechanisms are anti-oxidant status and BBB which were not well established in developing and young rats (PND 1 to 14) make them more prone to NaF neurotoxicity. The products which possess antioxidant nature are useful in reducing the production of free radicals from NaF, thus provide beneficial effects against NaF neurotoxicity. AMAE and AMEE are rich sources for quercetin and its derivatives which reduce the production of free radicals and provide rich nutritional value to the diet. AMEE showed good results over AMAE. Hence, the study concludes that the possible reason for reduced fluoride toxicity is the removal of free radicals with a parallel treatment of AMAE and as well as AMEE.

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CONFICT OF INTERESTS

There are no conflicts of interest.

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