Original Article

Oxidative stress evoked damages leading to attenuated memory and inhibition of NMDAR–CaMKII–ERK/CREB signalling on consumption of aspartame in rat model

Ashok Iyaswamy a,*, Ananth Kumar Kammella d, Citarasu Thavasimuthu c, Wankhar Wankupar a, Wankhar Dapkupar a, Sambantham Shanmugam b, Ravindran Rajan a, Sheeladevi Rathinasamy a

a Department of Physiology, Dr. ALM PG Institute of Basic Medical Sciences, University of Madras, Sekkizhar Campus, Taramani, Chennai 600 113, India
b Molecular Biology Laboratory, Department of Genetics, Dr. ALM PG Institute of Basic Medical Sciences, University of Madras, Sekkizhar Campus, Taramani, Chennai 600 113, India
c Centre for Marine Science and Technology, Manonmaniam Sundaranar University, Rajakkamangalam, Kanyakumari 629 502, India
d Department of Physiology, Michigan State University, East Lansing, MI, USA

Abstract

Many controversial reports are available on the use of aspartame as it releases methanol as one of its metabolite during metabolism. The present study proposes to investigate whether long term (90 days) aspartame (40 mg/kg b.wt) administration could induce oxidative stress and alter the memory in Wistar strain male albino rats. To mimic the human methanol metabolism, methotrexate (MTX)-treated rats were included as a model to study the effects of aspartame. Wistar strain albino rats were administered with aspartame (40 mg/kg b.wt) orally and studied along with controls and MTX-treated controls. Aspartame interfered in the body weight and corticosterone levels in the rats. A marked increase in the mRNA and protein expression of neuronal nitric oxide synthase (nNOS) and induced nitric oxide synthase (iNOS) which resulted in the increased nitric oxide radical’s level indicating that aspartame is a stressor. These reactive nitrogen species could be responsible for the altered cell membrane integrity and even cause death of neurons by necrosis or apoptosis. The animals showed a marked decrease in learning, spatial working and spatial recognition memory deficit in the Morris water maze and Y-maze performance task which could have resulted due to reduced hippocampal acetylcholine esterase (AChE) activity. The animal brain homogenate also revealed the decrease in the phosphorylation of NMDAR1–CaMKII–ERK/CREB signalling pathway, which well documents the inhibition of phosphorylation leads to the excitotoxicity of the neurons and memory decline. This effect may be due to methanol which may also activate the NOS

* Corresponding author.
E-mail address: ashokenviro@gmail.com (A. Iyaswamy).
https://doi.org/10.1016/j.jfda.2017.11.001
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levels, microglia and astrocytes, inducing neurodegeneration in brain. Neuronal shrinkage of hippocampal layer due to degeneration of pyramidal cells revealed the abnormal neuronal morphology of pyramidal cell layers in the aspartame treated animals. These findings demonstrate that aspartame metabolites could be a contributing factor for the development of oxidative stress in the brain.

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1. Introduction

Aspartame, a low-calorie sweetener obtained by the synthesis from two amino acids, L-phenylalanine and L-aspartic acid, is widely used in beverages, soft drinks, food and dietary products [1]. Aspartame’s attractiveness as a sweetener is 180 times sweeter than sugar in typical concentrations without the high energy of sugar. Aspartame on ingestion is metabolised to 40% of aspartic acid, 50% is phenylalanine and 10% is methanol [1]. Being that 10% of the metabolized by product, methanol is oxidized in the liver to formaldehyde which is further oxidized to formic acid in many tissues. In our previous finding it has been reported that aspartame appears to be a potential stressor as aspartame could elevate the corticosterone level and such elevated level may be the reason for the deviations in the free-radical-scavenging system [2–5]. Further, this elevation of corticosterone may be documented by the methanol that has been released from aspartame during its metabolism. The severity of clinical outcomes in methanol intoxication may be interrelated better with formate levels [6]. Also, methanol intoxication is concomitant with mitochondrial damage, augmented microsomal proliferation resulting in increased production of oxygen radicals [7] and altered free-radical-scavenging system in the brain (40 mg/kg b.wt) [8].

All isoforms of nitric oxide synthase (NOS) can be expressed in the brain; endothelial nitric oxide synthase (eNOS) in endothelial cells and neurons, neuronal nitric oxide synthase (nNOS) in neurons (presumably in only 1–2% of the total neuronal population in many brain regions) and induced nitric oxide synthase (iNOS) in activated microglia and astrocytes [9]. When overproduced, much of the newly synthesized nitric oxide (NO) will be converted into peroxynitrite, is an enormously potent free radical [10]. Nitric oxide is also likely to be involved in axonal and neuronal injury in demyelinating conditions [11].

Ionic involvement has been suggested to be involved in memory formation and the Na$^+$/K$^-$-ATPase enzyme is decisive for sustaining ionic gradients in neurons and tissues [12]. Cholinergic transmission is mainly terminated by Acetylcholine (Ach) hydrolysis by the enzyme AChE, this enzyme is widely expressed in tissues that receive cholinergic innervations, in neurons and muscle cells [13]. Hippocampus plays a crucial role in memory and damage to it will cause memory dysfunction and behavioural disturbances. Important molecular feature of learning and memory depends on the NMDAR–CaMKII–ERK/CREB signalling pathway. The synaptic plasticity in the brain is maintained by this signalling pathway. When there is inhibition in this signalling pathway it leads to the memory decline and neurodegeneration. Reduced levels of p-NMDAR1, p-CaMKII, and p-CREB may inhibit the normal function of neuron and it may result in neurodegeneration and memory decline. This may also activate the microglia and astrocytes to clear the degenerated neurons. Heat shock protein70 (Hsp70) is the most important proteins induced by stress in the brain, and its role in neuroprotection has been demonstrated both in vivo and in vitro [14].

Normally formate is metabolized twice as fast in the rat as in the monkey [15]. The rodents do not develop metabolic acidosis during methanol poisoning, due to their high liver folate content and to create similar results in human beings, merely folate deficient rodents are required to accumulate formate in order to develop acidosis [16]. Hence, in this study to mimic the human situation, a folate deficiency status is induced. Over the years researchers have expressed concern over methanol toxicity resulting from aspartame consumption and the chemical has been tested in both nonhuman primates and humans. The central nervous system (CNS) is extremely sensitive to free radical damage due to high polyunsaturated fatty acid content in cell membrane and relatively low total antioxidant capacity which are accountable to neuronal damage. Therefore, in this present study, an attempt has been made to investigate expression of aspartame in NMDAR–CaMKII–ERK/CREB signalling pathway in the central nervous system and the role of NMDAR–CaMKII–ERK/CREB signalling pathway and its inhibition in memory loss.

2. Materials and methods

2.1. Animals

Wistar strain male albino rats (200–220 g) were maintained under standard laboratory conditions with water and food. For the folate-deficient group, folate-deficient diet was provided for 45 days prior to the experiment and methotrexate (MTX) was administered for a week before the experiment. The animals were handled according to the principles of laboratory care framed by the committee for the purpose of Control and Supervision of Experiments on Animals (CPCSEA), Government of India. Prior to the experimentation, proper approval was obtained from the Institutional Animal Ethical Committee (No: 01/032/2010/Aug-11).
2.2. Chemicals

Aspartame and Methotrexate were purchased from Sigma chemical company, St. Louis, MO, USA. Nitric oxide assay kit was purchased from Biovision, USA. Taq-Polymerase, DNTPs from Genet Bio, China, Reverse Transcriptase enzyme kit from Thermo scientific, USA, primary antibody was purchased from Sigma and Cell signalling, USA, secondary antibody was purchased from Merck and other molecular grade chemicals were from Merck Bangalore, India. All other chemicals were of analytical grade obtained from the Sisco research laboratory, Bombay, India.

2.3. Experimental design

The European Food Safety Authority recently confirmed its daily acceptable intake (ADI) for aspartame as 40 mg/kg b.wt./day. In order to confine within the human permitted exposure limit, this dose was selected. Aspartame (Sigma Aldrich, USA) was administered orally; (40 mg/kg b.wt) mixed 1:1 with 0.9% saline and this dosage based on the food and drug administration (FDA) approved ADI limit.

2.4. Groups

The rats were randomly divided into three groups, namely, saline control, folate deficient (MTX treated) control, and MTX-treated (folate deficient) + aspartame administered groups. Each group consisted of six animals. Methotrexate (MTX) in sterile saline was administered (0.2 mg/kg/day) subcutaneously for 7 days to folate-deficient groups [2]. After 7 days of administration with MTX, folate deficiency was confirmed by estimating the urinary excretion of folic acid (FIGLU) [4]. From the eighth day, the MTX-treated (folate deficient) (third) groups were administered orally with aspartame for 90 days, whereas the saline control or vehicle control and (folate deficient) MTX treated control groups received equivalent volumes of saline as an oral dose and all animals were handled similarly.

2.5. Brain collection and dissection

The animals were sacrificed using higher dose of long acting pentothal sodium (100 mg/kg.b.wt). Collection of the blood samples and isolation of brain was performed between 8 and 10 a.m. to avoid circadian rhythm-induced changes. The brain was immediately removed and washed with ice-cold phosphate buffered saline (PBS). Further dissection was made on ice cold glass plate. The discrete regions of the brain (cerebral cortex, cerebellum, midbrain, Pons medulla, hippocampus and hypothalamus) were dissected according the method given in our previous publication [4]. The homogenate (10% w/v) of the individual regions was prepared in a Teflon-glass tissue homogenizer, using ice-cold PBS (100 mM, pH 7.4) buffer and centrifuged separately in refrigerated centrifuge 4 °C at 3000 rpm for 15 min. The supernatant was used for analyzing the biochemical estimations in this study. The body weight was recorded every week to monitor the influence of the aspartame treatment period.

2.6. Estimation of plasma corticosterone and LD50 of aspartame by Karber’s method

Plasma corticosterone was estimated by the method given by Clark [17]. The method is based on the oxidation of corticosteroids with iron (III) in acidic medium and subsequent complex of iron (II) with potassium hexacyanoferrate (III). The method offers the advantage of sensitivity and stability. The absorbance was measured at 780 nm in spectrophotometer (Shimadzu UV-1800). The level of corticosterone in the plasma was expressed as μmolar (μM). The calculation of LD50 by the method of Karber was used for the determination of LD50 of aspartame in Wistar albino rats. LD50 = LD100 - ∑(a x b)n. n = number of rats in a group. a = difference between two successive doses of aspartame administered. b = average number of dead animals in two successive aspartame doses. LD100 = Lethal dose causing the 100% death of all rats.

2.7. Estimation of membrane bound enzymes

The activity of Na+/K+ ATPase (ATP: Phosphohydrolase − EC. 3.6.1.3.) in the tissue was estimated by the method of Bernabe et al. [18]. The activity of Ca2+-ATPase (ATP: Phosphohydrolase − EC. 3.6.1.3.) in the tissues was estimated as described by Hjerten and Pan [19] and the activity of Mg2+-ATPase (ATP: Phosphohydrolase − EC. 3.6.1.3.) in the tissues was estimated by the methods of Ohnishi et al. [20].

2.8. Estimation of acetylcholine esterase (AChE)

The activity of Acetylcholine esterase in the tissues was estimated by the method of Ellman et al. [21]. The Principle of the method being the measurement of the rate of production of thiocholine as acetylthiocholine is hydrolysed. This is accomplished by the continuous reaction of the thiol with 5:5-thiocholine as acetylthiocholine is hydrolysed. This assay relies on a diazotization reaction that was originally described by Griess [22]. The original reaction has been described as a Griess reagent system is based on the chemical reaction, which uses sulfanilamide and N-1-naphthylethenediamine dihydrochloride (NED) under acidic (phosphoric acid) conditions.

2.9. Determination of nitrite using nitric oxide assay kit

To investigate nitric oxide formation it is essential to measure nitrite (NO2) which is one of two primary stable and nonvolatile breakdown products of NO. This assay relies on a diazotization reaction that was originally described by Griess [22]. The original reaction has been described as a Griess reagent system is based on the chemical reaction, which uses sulfanilamide and N-1-naphthylethenediamine dihydrochloride (NED) under acidic (phosphoric acid) conditions.

2.10. Measurement of memory in Y maze and Morris water maze

The Y maze measures spatial working and recognition memory by making use of a rodent’s natural exploratory instincts. In Y-maze the three arms were marked as A, B, C and “C” is marked as the novel arm (blocked arm) to test the spatial working memory. The first session measures spatial working memory in the rat by scoring the number of alternations the
rnat does in Y-maze when the animal visits all three arms without going into the same arm twice in a row. If working memory is intact, the rat should remember the first two arms and be more inquisitive about the third arm (originally blocked arm). In the second session open the blocked arm and the rats were left 5 min of time to evaluate the spatial recognition memory. A rat with a normal memory should spend more time exploring the “new” unblocked arm. The alternatives for the first 2 min were calculated. The overall percentage for 5 min and the percentage for the first 2 min were also calculated for spatial working and recognition memory. The Morris water maze (MWM) test consists of a circular pool made of non-toxic white metal, filled with water and maintained at 21 ± 1 °C. A platform was placed 1 cm below the water surface at a fixed position. Animals were acclimatized and trained. Animals were monitored using an automated tracking system (Ethovision software). Visible platform trials were performed to evaluate sensorimotor and/or motivational deficits that could influence performance during the learning task. Hidden-platform training consisted of 6 days with each day comprised four trials with a 60 min interval. The platform location remained constant in the hidden platform trials, and the entry point was changed randomly in all days. Memory retention was assessed by a probe trial after the 6 days hidden-platform training. The time spent in the target quadrant without the platform was taken to measure level of memory retention in the probe trial.

2.11. Isolation of total RNA & reverse transcription-polymerase chain reaction (RT-PCR)

Total RNA was isolated from cells using Trizol reagent following the method given in our previous publication [4]. The total RNA obtained was free from protein and DNA contamination. The reverse transcription step was performed using the reverse transcriptase (RT) enzyme kit. Each 20 μL reaction mixture contained 5 μL Oligo dT (10 μM), 1 μL dNTP (10 μM), 4 μL First Strand buffer (5×), 0.2 μL super script III reverse transcriptase (200 U/μL) varied quantity of RNA template (dependent on RNA concentration) and RNase free water to make up the volume. Thermal cycling conditions for the first strand reaction consisted of 25 °C for 5 min, 50 °C for 45 min, 70 °C for 15 min and finally maintained at 4 °C for 5 min. PCR amplification was performed using Taq DNA Polymerase. Each 20 μL of sample contained 10 μL Master mix (2 μM), 1 μL forward primer, 1 μL reverse primer (Table 3) for both gene of interest and internal control consecutively, 2 μL RT sample, 4 μL sterile water. The mixture was kept at thermocycler and amplified for 35 cycles. Each thermocycling consisted of 94 °C for 30 s, varied annealing temperature for each gene of interest for 30 s, 72 °C for 30 s. β-actin was co-amplified with c-fos, iNOS, nNOS, Hsp70 genes using the same procedures. The sense and antisense primer for the study is tabulated in table. Ten microliters of each PCR product was analyzed by gel electrophoresis on 2% agarose gel. Agarose gel electrophoresis is an effective method for the identification of purified DNA molecules [2]. Amplified product was analyzed by agarose gel electrophoresis with ethidium bromide staining. Then the gel containing cDNA was visualized with the help of the fluorescent imager (Bio-Rad, USA). The Band intensity was quantified by Quantity One Software. The band intensification for each enzyme mRNA was normalized with that of the internal control β-actin using Quantity One Software.

2.12. Histopathology

Animals were deeply anesthetized with ketamine hydrochloride. Rats were then perfused transcardially with phosphate-buffered saline, followed by buffered 10% formalin. The brain was removed, and preserved in formalin until processed for histology. Then kept on running water to remove formalin pigments and dehydrated with ascending grades of alcohol. After impregnation with paraaffin wax, the paraffin blocks were made. They were processed and sections were cut with 10 μm in thickness using “Spencer Lens, rotary microtome (No 820, New York, USA) and then stained with Cresyl fast violet as follows for the brain.

2.13. Immunohistochemical analysis

Immunohistochemical analysis was carried out using the DAB universal staining kit (Merk Genie, Bengaluru, India). The sections were deparaffinized in xylene and dehydrated in ethanol. After washing with PBS, slides were incubated with 3% H2O2 in at room temperature for 15 min to quench endogenous peroxidase activity. After antigen retrieval (15 min of autoclaving in 10 mM citrate buffer, pH 6.0), the slides were incubated with blocking solution (10% normal goat serum) for 5 min at room temperature. Then, the sections were incubated overnight with primary antibody. Subsequently, the sections were incubated with horseradish peroxidase (HRP) secondary link antibody for 30 min at room temperature, washed with PBS. Then, the sections were treated with 3,3′-diaminobenzidine (DAB) chromogen for 15 min. Finally, the sections were washed with deionised water, counter stained with haematoxylin and mounted. Photographs were taken using Nikon microscope (Nikon 1.25X Japan DS-Fi1).

2.14. Western blot

The brains were washed with ice-cold PBS and powdered in liquid nitrogen, was suspended in the cytobuster buffer (Invitrogen) was used to detect the NMDAR, CaMKII, ERK, CREB, PSD95 and Synaptophysin. Protein concentrations of brain lysates were assayed using BCA reagent (BioRad, USA). Equal amounts of protein were subjected to 8% and 10% SDS polyacrylamide gels at 70 V, and were transferred onto polyvinylidene fluoride (PVDF) membranes (Merck Millipore) at 300 mA for 90 min, and then blocked with 5% fat-free milk or BSA for 1 h at room temperature. The membranes were then incubated overnight with the primary antibodies at 4 °C. Membranes were washed and probed with secondary antibody: goat anti-rat IgG, and goat anti-rabbit IgG (Thermo-scientific). Protein signals were visualized using an ECL chemiluminescent HRP substrate detection system. The band intensity was quantified by ImageJ and Quantity One software (Bio-Rad, USA). The membranes were stripped and reprobed for B-actin (Sigma) (1:10,000) as an internal control.
2.15. Statistical analysis

Statistical analysis was carried out using the SPSS statistical package version 18.0. The results are expressed as Mean ± SD and the data were analyzed by the one-way analysis of variance (ANOVA) followed by Turkey's multiple comparison tests when there is a significant 'F' test ratio. The level of significance was fixed at $p < 0.05$. 

Fig. 1 – Long term aspartame consumption interferes the body weight, plasma corticosterone and brain NO and AcHE levels. A, Effect of aspartame (40 mg/kg b.w) on plasma corticosterone level (mM) in rat. B, Long term consumption of aspartame interferes on body weight of the rats. C, Effect of aspartame (40 mg/kg b.w) on nitric oxide levels (umoles/mg tissue) in rat brain discrete regions. D, Effect of aspartame (40 mg/kg b.w) on acetylcholine esterase levels (umoles/min/mg protein) in rat brain discrete regions. Comparison and analysis were done by the one-way analysis of variance (ANOVA) ($n = 6$). The data from various groups for the individual parameters are presented as bar diagram with mean ± SD. Significance fixed at $F < 0.05$, CC – cerebral cortex, CB – cerebellum, MB – midbrain, PM – pons medulla, HP – hippocampus, HY – hypothalamus. Aspartame treated group when compared to control significance is marked as * and MTX treated groups significance is marked as #.
Table 1 - Effect of Aspartame (40 mg/kg b.wt) on Na⁺/K⁺ ATPase, Ca²⁺ ATPase and Mg²⁺ ATPase (µmoles of phosphorous liberated/min/mg protein) in brain regions.

| Parameter          | Control          | MTX treated     | Asp + MTX treated |
|--------------------|------------------|-----------------|------------------|
| Cerebral cortex    | Na⁺/K⁺ ATPase    | 0.73 ± 0.04     | 0.71 ± 0.04      | 0.29 ± 0.04*#     |
|                    | Ca²⁺ ATPase      | 0.44 ± 0.04     | 0.45 ± 0.04      | 0.28 ± 0.03*#     |
|                    | Mg²⁺             | 0.70 ± 0.03     | 0.69 ± 0.02      | 0.38 ± 0.02*#     |
| Cerebellum         | Na⁺/K⁺ ATPase    | 0.53 ± 0.04     | 0.52 ± 0.03      | 0.35 ± 0.03*#     |
|                    | Ca²⁺ ATPase      | 0.34 ± 0.04     | 0.34 ± 0.03      | 0.20 ± 0.04*#     |
|                    | Mg²⁺             | 0.49 ± 0.03     | 0.51 ± 0.03      | 0.22 ± 0.04*#     |
| Midbrain           | Na⁺/K⁺ ATPase    | 0.70 ± 0.05     | 0.69 ± 0.03      | 0.28 ± 0.03*#     |
|                    | Ca²⁺ ATPase      | 0.45 ± 0.04     | 0.47 ± 0.04      | 0.25 ± 0.01*#     |
|                    | Mg²⁺             | 0.53 ± 0.04     | 0.53 ± 0.04      | 0.26 ± 0.03*#     |
| Hippocampus        | Na⁺/K⁺ ATPase    | 0.63 ± 0.03     | 0.63 ± 0.03      | 0.41 ± 0.03*#     |
|                    | Ca²⁺ ATPase      | 0.39 ± 0.04     | 0.39 ± 0.04      | 0.27 ± 0.02*#     |
|                    | Mg²⁺             | 0.70 ± 0.02     | 0.72 ± 0.03      | 0.38 ± 0.03*#     |
| Hypothalamus       | Na⁺/K⁺ ATPase    | 0.86 ± 0.04     | 0.85 ± 0.05      | 0.49 ± 0.03*#     |
|                    | Ca²⁺ ATPase      | 0.43 ± 0.05     | 0.41 ± 0.03      | 0.27 ± 0.02*#     |
|                    | Mg²⁺             | 0.53 ± 0.03     | 0.51 ± 0.04      | 0.29 ± 0.04*#     |
| The data from various groups for the individual parameters are presented as table with mean ± SD. Significance fixed at P ≤ 0.05. Aspartame treated group when compared to control, MTX treated groups #.

3. Results and discussion

The present study support the fatal effect of aspartame when consumed repeatedly for a prolonged period and it may be probably due to the toxic metabolite methanol. It has been reported that following aspartame consumption, the concentrations of its metabolites are increased in the bloodstream [23]. A small amount of aspartame significantly increases the plasma methanol level [24]. Further, it was reported that not only the metabolites of methanol, but methanol per se as well, is toxic to the brain [4]. In addition, this increase in the blood methanol level was associated with a marked increase in the free radical generation in brain regions of aspartame treated animals [2–5,8,9]. The cells can be injured or killed when the ROS generation overwhelms the cellular antioxidant capacity [25] indicate its vulnerability to free radicals. In this study the LD50 of aspartame was analyzed by karber’s method, the LD50 of aspartame (Table 2) in rats is 10.95 g/kg of body weight. Aspartame interfered in the body weight of the treated animals (Fig. 1B) when compared to the control rats throughout the study duration.

3.1. Aspartame induces the free radical generation

The nitric oxide (•NO), a nitrogen free radical [26], is produced by a number of different cell types. The elevation of nitric oxide (Fig. 1C) observed in this study again support the generation of free radical by aspartame. However, this increase in nitric oxide after aspartame consumption could not be overlooked because according to McCord [27], the prolonged exposure to free radicals, even at a low concentration, may result in the damage of biologically important molecules. In this finding the aspartame treated animals, did showed an elevation of nitric oxide level but not the nitrate and this is supported by the western blot study in which nNOS protein expression was enhanced along with increase mRNA expression of nitric oxide synthesizing enzyme nNOS and iNOS (Figs. 3 and 4). Studies demonstrated that NO produced by iNOS causes damage to brain cells, a process which has been demonstrated in many neurodegenerative diseases and exert numerous toxic effects in a wide variety of mammalian cell targets [28].

3.2. Aspartame induces stress and decreases membrane homeostasis

In the present study, there was an increased plasma corticosterone level (Fig. 1A) in the aspartame treated animals and it is relevant to point out that, in our earlier report also there was a marked increase in the corticosterone level in the plasma on (75 mg/kg) aspartame administered rats [4], indicating that the dietary sweetener aspartame could act as a chemical stressor. Decrease in membrane bound ATPases activity (Table 1) in this study may be due to free radical generation and high NO level, which is supported by few studies. Ashok et al. reported that [2] methanol induced increased production of free radicals and increased oxidative damage to distinct brain regions, retina and optic nerve has also been reported. McIntosh et al. [29] reported a decreased activity of the antioxidant enzymes in the brain of rats treated with glucocorticoids. Indicating that elevated free radical generation overwhelming the antioxidant defence system. High levels of NO have been associated with membrane lipid peroxidation [30] which suggest that the aspartame mediated increase in free radical may be the cause behind the altered membrane integrity observed in this study which eventually lead to decrease in membrane bound ATPases activity.

Na⁺/K⁺-ATPase, the enzyme that maintains Na⁺ and K⁺ gradients across the plasma membrane, was reported to be inhibited by ROS in the brain [31] which strengthen our argument. According to Polizzi et al. [32], the Na⁺/K⁺-ATPase is very sensitive to the plasma membrane structure changes and therefore measuring its activity represents a valuable indicator of the early and late stages of tissue injury. Further, the decrease in the activity of these membrane bound enzymes could not be ignored as Na⁺/K⁺-ATPase is responsible for the generation of the membrane potential through the active transport of sodium and potassium ions in the neurons in CNS necessary and to maintain neuronal excitability. Moreover, the role of Mg²⁺-ATPase is to maintain high brain intracellular Mg²⁺ which can control rates of protein synthesis and cell growth. Na⁺/K⁺, Mg²⁺ and Ca²⁺ ATPase in the plasma
membrane also keeps the intracellular sodium low but intracellular magnesium and potassium high when compared with the levels in extracellular fluids [33]. Hence, the increase in nitric oxide, free radical and corticosteroids in this study after aspartame consumption may be a contributing factor for the alteration observed in the membrane bound enzymes and finally altering the neuronal integrity.

3.3. Aspartame attenuates the learning and memory

The Y-maze task as used in the present study, with one novel arm throughout the experiment, is so-called “reference memory task.” This type of test is considered to be hippocampus dependent. Various reports have shown that bilateral cytotoxic lesions of the hippocampal formation result in severe impairments in the Y-maze reference memory task [34] and NO also contributes to hippocampal dependent learning and memory tasks including spatial memory [35]. AChE is a key enzyme of the muscarinic cholinergic system, involved in learning and memory. However, NO formed as a result of iNOS up regulation during hypoxia has been found to interrupt the memory process by inhibiting acetylcholinergic activity [36]. A marked decrease in the AChE activity (Fig. 1D) in the aspartame treated animals of present study lends a support to alteration in the learning and memory process (Fig. 2A–E) of these aspartame fed animals in both Morris water maze and Y maze task. It is also in agreement with the Irene et al. [37], who reported that aspartame metabolites inhibit the activity of hippocampal AChE. Rico et al. [38], reported that methanol exposure remarkably decreased zebrafish brain AChE activity, suggesting a neurodegenerative event promoted possibly by methanol and suggest that aspartame induced alterations may be mediated by methanol by affecting the hippocampus functioning. Moreover our previous study in 75 mg/kg aspartame consumption also showed alteration in the locomotor, anxiety and emotional behaviour of the treated rats [4]. Thus supporting our study that aspartame attenuates learning, memory (Fig. 2A–E) and declines the memory retention in the Morris water maze task.

3.4. Aspartame inhibits the NMDAR–CaMKII–ERK/CREB signalling pathway

As aspartame attenuated memory, we were curious in finding out the signalling pathway involved in memory decline [38] and neurodegeneration. We further investigated the NMDAR–CaMKII–ERK/CREB signalling pathway. This is an important molecular feature of learning and memory in the neuroplasticity of brain which involves in phosphorylation of receptor and calcium influx [32] leading to activation of several proteins by phosphorylating the proteins. In our study we found that the phosphorylation of NMDAR1, CamKII, ERK and CREB was inhibited (Fig. 4C) which leads to the memory decline and neuronal damage. As this signalling pathway is inhibited the protein levels of presynaptic and postsynaptic markers namely PSD95, Synaptophysin (Fig. 4B) were decreased. The NOS [36] expression levels activated the microglia and astrocytes, there activated protein expression was illustrated by IHC of Iba1 and GFAP (Fig. 4A). These activated proteins illustrates that neuronal degeneration may also have been through gliosis, which clearly confirms that neurodegeneration may be due to inhibition of NMDAR–CaMKII–ERK/CREB signalling pathway activation of glial cells which might have initiated the inhibitory synaptic excitotoxicity in neurons. When methanol from aspartame [2–5] inhibits the phosphorylation of NMDAR1 it leads to the
decrease in the Ca\(^{2+}\) influx, which inhibits the phosphorylation of CaM kinase II further inhibits the phosphorylation of ERK and CREB leading to decrease in memory proteins and other neuronal proteins which damages the neurons and accelerating the generation of free radicals by increasing the expression levels of NOS. Interestingly we also found that NOS also activated the astrocytes and glial cells as neurons were at the verge of cell death. Further the neuroprotective effects of NO may be overcome by the nitrosative stress which leads to prolonged, excessive, and persistent

Fig. 2 – Long term aspartame consumption (40 mg/kg b.wt) attenuates the learning and memory in Wistar albino rats in both Morris water maze and Y maze. A, Effect of aspartame (40 mg/kg b.wt) during the training period of learning and attenuation of memory. B and C, Long term consumption of aspartame (40 mg/kg b.wt) attenuates the memory retention in probe trial. D, Evaluation of aspartame effect (40 mg/kg b.wt) on spatial working memory by Y maze. E, Evaluation of aspartame effect (40 mg/kg b.wt) on spatial recognition memory by Y maze. Comparison and analysis were done by the one-way analysis of variance (ANOVA) (n = 6) control group was compared with MTX control group and aspartame MTX group, MTX control group was compared with Aspartame MTX group. Data are expressed as mean ± SD, n = 6. *P ≤ 0.05. Control, MTX control – methotrexate treated group, Asp + MTX – Aspartame + Methotrexate treated group. Aspartame treated group when compared to control significance is marked as * and MTX treated groups significance is marked as #.
Fig. 3 – Effect of long term aspartame on c-fos, Hsp70, iNOS and nNOS mRNA expression in brain regions of Wistar albino rats. Lanes 1, 4, 7, 10 – Control, Lanes 2, 5, 8, 11 – MTX control, Lanes 3, 6, 9, 12 – MTX + ASP, M-100 bp marker. Lanes 1, 2, 3 – cerebral cortex, Lanes 4, 5, 6 – cerebellum, Lanes 7, 8, 9 – hippocampus, Lanes 10, 11, 12 – hypothalamus. CC – cerebral cortex, CB – cerebellum, HI – hippocampus, HY – hypothalamus. Comparison and analysis were done by the one-way analysis of variance (ANOVA), control group was compared with MTX control group and aspartame MTX group, MTX control group was compared with Aspartame MTX group. Data are expressed as mean ± SD, n = 6. *P ≤ 0.05. Control, MTX control – methotrexate treated group, Asp + MTX – Aspartame + Methotrexate treated group. Aspartame treated group when compared to control significance is marked as * and MTX treated groups significance is marked as #.
Fig. 4 – Effect of long term aspartame on [A] glial cells and astrocytes activation in brain of Wistar albino rats by immunohistochemistry and [B] synaptic markers by western blot [C] mechanism of action on the neuronal memory loss and degeneration. [A] Immunohistomicrograph staining shows the effect of aspartame (40 mg/kg b.w) on iba1 and GFAP in brain of Control, Methotrexate (MTX) control and Aspartame treated animals. Comparison and analysis were done by the one-way analysis of variance (ANOVA), control group was compared with MTX control group and aspartame MTX group, MTX control group was compared with Aspartame MTX group. Data are expressed as mean ± SD, n = 6. *P ≤ 0.05. [B] Effect of Aspartame (40 mg/kg b.w) on synaptic markers namely PSD-95, Synaptophysin, p-ERK and total ERK. [C] Effect of Aspartame (40 mg/kg b.w) on the mechanism of action in memory processing and neuronal degeneration, the proteins involved namely NMDAR1, CaMKII, CREB and its phosphorylated proteins.
Fig. 5 – Effect of long term aspartame (40 mg/kg b.wt) on brain in Wistar albino rat, [A] the histomicrograph of brain stained by cresyl fast violet. [B] Immunohistochemistry of Hsp70, nNOS, c-fos and iNOS protein expression in brain of Wistar albino rats. [A] Histomicrograph Cresyl fast violet (CFV) staining of brain in Control, Methotrexate (MTX) control and Aspartame + Methotrexate treated animals. CA1 – Cornus Ammonis 1, CA2 – Cornus Ammonis 2, CA3 – Cornus Ammonis 3, DG – Dentate Gyrus, DNC – Degenerative neuronal cells (DNC), HL – Hippocampal Layer. [B] Immunohistomicrograph staining of brain in Control, Methotrexate (MTX) control and Aspartame + Methotrexate treated animals. Comparison and analysis were done by the one-way analysis of variance (ANOVA), control group was compared with MTX control group and aspartame MTX group, MTX control group was compared with Aspartame MTX group. Data are expressed as mean ± SD, n = 6. *P ≤ 0.05. Aspartame treated group when compared to control significance is marked as * and MTX treated groups significance is marked as #.
production of NO radical, ultimately leading to neurotoxicity and cell death. Aspartame administration attenuated memory by inhibiting NMDAR–CaMKII–ERK/CREB signalling pathway (Fig. 6), the observed changes may owe to the methanol from aspartame which inhibited the phosphorylation of NMDAR–CaMKII–ERK/CREB signalling pathway.

3.5. Aspartame is a chemical stressor and causes oxidative damage to brain

In the present study a significant increase mRNA expression of c-Fos and increase in the expression of Hsp70 in aspartame treated animals (Figs. 3 and 5B) when compared to controls. Gil et al. [39] reported that Fos protein plays an important role in the cell growth, differentiation and in the recovery from reversible injuries. In various studies c-Fos gene expression has been associated with delayed neuronal cell death, and it has been shown that prolonged c-Fos induction precedes neuronal death [40]. Hsp70 gene encodes a major stress inducible heat shock protein, which plays an important role in protecting cells from deleterious stresses. It was reported that gene expression leading to the synthesis of HSP exerts cytoprotective functions [41]. Moreover, it has been reported that the heat shock system play a vital role in the protective mechanism of brain and the Hsp70 is considered as a critical determinant of brain's stress tolerance [42]. Therefore, the possible cause for the aspartame consumption induced increased expression of these Hsp70 and c-Fos may be to prevent cellular injury, probably caused by the elevated levels of free radicals.

The aspartame treated animals showed a neuronal shrinkage of hippocampal layer (HL) due to degeneration of hippocampal neurons (Fig. 6), the observed changes may owe to the methanol from aspartame. The observed changes may be due to the methanol or its metabolite. Since aspartame is consumed more than the recommended dosage by common people, it is essential to do more work on aspartame and to create awareness regarding the usage of this artificial sweetener.

4. Conclusion

The observed results support the fatal effect of aspartame when consumed repeatedly and aspartame administration attenuated memory by inhibiting NMDAR–CaMKII–ERK/CREB signalling pathway and decreased the membrane bound
infrastructure to conduct the research. We acknowledge University of Madras for providing the financial assistance provided by the Indian Council of Medical Research (ICMR) No. 3/1/2/29/Nut./2012/Dated 29-09-2013 for Senior Research Fellow is gratefully acknowledged.

The authors declare that they have no competing interests.

Conflicts of interest
The authors report no conflicts of interest. The authors alone are responsible for the content and writing of this paper. The authors declare that they have no competing interests.

Acknowledgement
The financial assistance provided by the Indian Council of Medical Research (ICMR) No. 3/1/2/29/Nut./2012/Dated 29-09-2013 for Senior Research Fellow is gratefully acknowledged. We acknowledge University of Madras for providing the infrastructure to conduct the research.

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