EFFECT OF MONOSODIUM GLUTAMATE ON THE CEREBELLAR CORTEX OF MALE ALBINO RAT AND PROTECTIVE ROLE OF VITAMIN C

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ABSTRACT
Background: Monosodium Glutamate (MSG) is one of the most widely used food-additives in commercial foods that had effect on various tissues including cerebellum. It acts via creating an oxidative stress. The central nervous system is a target organ for MSG especially cerebellar cortex. The health benefits of vitamin C were derived from its role in the key reactions within immune function, metabolism, and other enzymatic reactions. Aim: To evaluate the effect of MSG on rat’s cerebellar cortex with the possible protective role of Vitamin C. Methods: Twenty one adult male albino rats have been used in this work. The animals were randomly divided into three groups with seven animals in each. Group 1 (Control): Animals were kept without addition of any chemicals. Group 2 (MSG-treated): Animals were subjected to administration of MSG 4g/kg body weight dissolved in 1 ml normal saline. Group 3 (MSG and Vitamin C treated): Animals were subjected to administration of both MSG 4g/kg body weight dissolved in 1 ml normal saline and 500 mg/kg vitamin C, orally. By the end of the experiment which was 10 days. Animals were anaesthetized and sacrificed. Cerebellar hemispheres were obtained and specimens were processed for both light and electron microscopic examination. Results: MSG caused histopathological and morphometric changes in rat’s cerebellar cortex. Vitamin C protected the cerebellar cortex specimens against such changes. Conclusion: MSG could result in hazards to the structure of cerebellar cortex. Fortunately co-administration of vitamin C is suggested to reduce such hazards.
Keywords: Adult rats, cerebellar cortex, monosodium glutamate, vitamin C.

INTRODUCTION
Most food additives act either as preservatives or enhancer of palatability. One of such food additives is monosodium glutamate (MSG) (1). Recently MSG is commonly sold in Egyptian markets and many Egyptians consume it almost daily (2). The central nervous system is an important target organ for MSG (3, 4). A previous study documented that 4 g/kg but not 2 g/kg MSG treatment early in life increased the brain’s ability to propagate cortical spreading depression (CSD) when neonatal animals reached 45 - 60 days of age (5). In addition, various studies reported that 2 g/kg of MSG for 10 consecutive days initiated multiple neurotoxic pathways and affected many neurochemical parameters (6). It has been known that MSG - induced neuronal toxicity that was mediated by oxidative stress (7). Also the increased oxidative stress may mediate the MSG induced cerebellar injury through the depletion of glutathione which resulted in a form of cell injury called oxidative glutamate toxicity (8). Hashem et al. (9) documented that MSG affects Purkinje cells of cerebellum as they appeared with darkly stained cytoplasm and shrunken darkly stained nuclei, while with vitamin C administration these effects were decreased. Ascorbic acid was the antioxidant of choice as dehydroascorbic acid, the main form of oxidized vitamin C in the body, may reduce neurological deficits due to its ability to cross the blood brain barrier (10). This protective role of ascorbic acid was explained by Farombi and

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Onyema (11) who mentioned that, dietary antioxidants as ascorbic acid, vitamin E and quercetin had protection potential against oxidative stress induced by MSG, and also by Pavlovic et al. (12) who found that, the treatment with ascorbic acid may prevent the MSG-induced cytoxicity in rat thymocytes by up-regulating Bcl-2 protein expression (cell lymphoma protein). The aim of this work was to elucidate the possible changes that take place in the structure of cerebellar cortex of adult male albino rats after administration of Monosodium Glutamate (MSG) and the protective effect of vitamin C using different histological procedures.

II. Material and Methods

Chemicals
Monosodium glutamate and vitamin C: They were obtained from Al-kahira Company of pharmaceutical industries as a powder and were dissolved in normal saline solution.

Experimental animals
The study was performed on twenty-one healthy adult male Wistar albino rats (3 months old) weighing 180-220 gm. The animals were obtained from the animal house, Faculty of Medicine, Zagazig University. The animals were housed under controlled laboratory conditions. All experimental procedures were performed in accordance with the guidelines of the Institutional Animal Care and the norms of Ethical Committee of Faculty of Medicine; Zagazig University. The rats were divided into 3 groups as follows:

First (control) group: Animals were kept without any medications.

Second (MSG-treated) group: Rats were subjected to oral administration of MSG 4g/kg body weight dissolved in 1 ml normal saline (12).

Third (MSG and Vitamin C-treated) group: MSG 4g/kg body weight dissolved in 1 ml normal saline and 500 mg/kg vitamin C, orally (12).

Methods
I. Experimental Methodology:
At the beginning of the study all the used rats were weighed and marked in all groups. By the
were obtained. Using the interactive measure, the number of Purkinje cells and thickness of molecular cell layer were measured using magnification X 400 with measure frame 7381.11 µm. Data were analyzed using statistical package of Social Science (SPSS) version 20 (SPSS Inc., Chicago, Illinois, USA), then expressed by using two variants of analytic tests, Kruskal-Wallis test for comparison between more than two groups not normally distributed having quantitative variables or of relatively small sample size and One-way ANOVA (analysis of variance) for comparison of three or more independent quantitative variables normally distributed. The results were considered statistically significant at P-value < 0.05.

RESULTS

I. Rat body weight

By comparing animals’ body weight at the beginning and at the end of the study for each group there was a statistical significant increase in body weight in all groups. By comparing the body weight of the three groups at the end of experiment, there was a statistical significant difference between the three groups (p-value < 0.001), as the rats receiving MSG have the highest body weight (mean body weight of 308 g), compared with control group (mean body weight of 230 g). (Table 1)

II. Light Microscopic Examination

H&E stain

The cerebellar cortex of control group showed that the molecular cell layer and Purkinje cell layer was arranged in one row with densely packed granular cell layer. Meningeal coverings were intact. There was a well-defined core of white matter stuffed with less packed fusiform cells. The cerebellar folia were separated by deep and long sulci (Fig.1a).The cortex showed few scattered cells in molecular cell layer, Purkinje cell layer contained Purkinje cells which were large and flask shaped arranged in one row. Densely packed granular cell layer was with small rounded granular cells (Fig.1b).The cortex of the MSG- treated group showed that there was densely packed granular cell layer. Meningeal coverings were detached. There was a well-defined core of white matter stuffed with less packed fusiform cells (Fig.1c).The cortex of the MSG- treated group showed Purkinje cell layer with necrotic widely spaced cells (Fig.1d). The cortex of MSG + vit. C treated group showed that meningeal coverings were intact and less congested. (Fig.1e).The cortex of MSG +vit.C treated group showed that the Purkinje cell layer contained some cells which were degenerated and necrotic and others were flask shaped with pale nuclei and prominent nucleoli (Fig.1f).

Immunohistochemical results for GFAP

The control sections showed dense positive reaction in astrocytes which appeared large and with long, regular, linear and parallel to each other processes and glial limitans membrane was well developed (Fig. 2a). In MSG- treated group, there was marked positive reaction to GFAP in astrocytes which were large densely packed around Purkinje cells, with long and fragmented, thin and not parallel to each other processes and the cell body is more expressed with the GFAP reaction. The glial limitans membrane was separated from the cortex (Fig. 2b). In MSG +vit. C treated group, there was positive reaction to GFAP in astrocytes which were large densely packed around Purkinje cells, with long and fragmented processes. The glial limitans membrane was preserved and well developed (Fig. 2c). The mean percent of reactivity in control and MSG +Vit C groups were 7.71 ± 2.91 and 8.75 ± 2.31 respectively. One-way ANOVA revealed a statistically significant difference in immunoreactivity between all groups [F (2, 12) =22.94, p<0.001]. Post hoc analyses indicated that the immunoreactivity in MSG treated group was significantly higher than that in control and MSG+ vit. C treated groups respectively. However, the slight differences in immunoreactivity between control and MSG+ vit. C treated groups were statistically non-significant.

Toluidine Blue stain

In control group, Purkinje cells were arranged in one row and appeared large flask-shaped,
near Purkinje cells there were astrocytes (Fig. 3a). In MSG-treated group, the cerebellar cortex was infiltrated with astrocytes which were presented mostly adjacent to Purkinje cells which were shrunken, distorted and deeply stained. Granular layer contained some cells preserving their normal appearance and others were distorted with necrotic nuclei (Fig. 3b). In MSG +vit.C treated group, some Purkinje cells were shrunken, distorted and deeply stained; others were flask-shaped with pale nuclei and prominent nucleoli. Most of granular cells were keeping their rounded or oval nuclei; others were deeply stained and distorted (Fig. 3c).

**III. Electron Microscopic Finding**

The cerebellar cortex of control group showed granule cells which had oval or rounded nuclei with coarse central and peripheral chromatin clumps and scanty cytoplasm containing mitochondria. The surrounding neuropil contained myelinated nerve fibers (Fig.4a). The control groups also showed Purkinje cell which was large flask shaped with euchromatic nucleus with indentations, prominent nucleolus and cytoplasm containing mitochondria and free ribosomes (Fig.5a). The astrocyte also appeared with sharply demarcated nucleus and electron lucent cytoplasm containing mitochondria. The astrocytes were present near the blood capillary which was lined by endothelial cells. The surrounding neuropil contained myelinated nerve fibers. Oligodendrocyte had irregular deeply stained nuclei (Fig.6a).

The cerebellar cortex of MSG-treated group showed granular layer with granular cell containing rounded or oval nuclei with clumped chromatin, vacuoles in cytoplasm and necrotic fragmented cells with intact cell membrane. The surrounding neuropil contained distorted myelinated nerve fibers (Fig.4b). Purkinje cell appeared with euchromatic nucleus with prominent nucleolus; its cytoplasm contained marked dilated rough endoplasmic reticulum and mitochondria. There was also an astrocyte with sharp demarcated nucleus with vacuolated cytoplasmand mitochondria (Fig.5b). The surrounding neuropil contained unmyelinated nerve fibers and granule cells. Purkinje cell was presented with its distinguished damaged appearance (Fig.6b).

The cerebellar cortex of MSG +vit.C treated group showed granule cells which had rounded or oval nuclei with clumped chromatin, vacuoles in cytoplasm which also contained mitochondria (Fig.4c). Purkinje cell contained euchromatic irregular nucleus and prominent nucleolus with indentations all around ill-defined nuclear membrane and its cytoplasm was dark, contained mitochondria, strands of mildly dilated rough endoplasmic reticulum and free ribosomes (Fig.5c). The astrocytes appeared with sharp demarcated nucleus with chromatin and vacuolated cytoplasm containing mitochondria and few vacuoles (Fig.6c).

**IV. Morphometric Results**

There was statistical significant difference between the three groups as regard the number of Purkinje cells (p-value < 0.001), as the rats receiving MSG have the lowest number of Purkinje cells (mean number of 2.1 cells), compared with control group (mean number of 6.1 cells) (Table.2).

There was statistical significant difference between the three groups as regard the thickness of molecular cell layer (p-value = 0.008), as the controlled rats have the thickest molecular cell layer (mean thickness of 145.5 um), compared with MSG-treated group (mean thickness of 114.3 um) (Table.3).
Fig. 1: Photomicrographs of cerebellar cortex of the different groups: 

- **a)** Control group showing: Molecular cell layer (MCL), Purkinje cell layer (PCL) arranged in one row and densely packed granular cell layer (GCL). Meningeal coverings are intact (Ms). There is a well-defined core of white matter (WM) stuffed with less packed fusiform cells. The cerebellar folia (F) are separated by deep and long sulci (S).

- **b)** Molecular cell layer (MCL), Purkinje cell layer (PCL) Purkinje cells (P) which are large flask in shape arranged in one row. Densely packed granular cell layer (GCL) with small rounded granular cells (G) are also observed in granular layer.

- **c)** Molecular cell layer (MCL), Purkinje cell layer (PCL) arranged in one row. Densely packed granular cell layer (GCL). Meningeal coverings (Ms) are detached (zigzag blue arrows). There is a well-defined core of white matter (WM) stuffed with less packed fusiform cells. The cerebellar folia (F) are separated by sulci (S).

- **d)** Molecular cell layer (MCL), Purkinje cell layer (PCL) contains necrotic cells (PN) with spacing between cells (thick arrows), vacuolated granular layer (GCL) appear. Meningeal coverings (Ms) are intact and less congested. There is a well-defined core of white matter (WM) stuffed with less packed fusiform cells. The cerebellar folia are separated by sulci (S).

- **e)** Molecular cell layer (MCL), some Purkinje cell layer (PCL) containing some cells which are degenerated and necrotic (PN) and others are normal (P), vacuolated (V) granular layer (GCL) appear.

- **f)** Molecular cell layer (MCL), some Purkinje cell layer (PCL) containing some cells which are degenerated and necrotic (PN) and others are normal (P), vacuolated (V) granular layer (GCL) appear. 

[a, c & e: H&E X 100; b, d & f: H&E X400]
Fig. 4: a) A photomicrograph of transmission electron microscope of cerebellar cortex of adult control albino rat shows granule cells (G) which have oval or rounded nuclei (Ng) with coarse central and peripheral chromatin clumps (C), scanty cytoplasm (Cg) containing mitochondria (m). The surrounding neuropil contains myelinated nerve fibers (NF). (TEM × 800×17) b) A photomicrograph of transmission electron microscope of cerebellar cortex treated group shows granular layer with granular cells (G) containing rounded or oval nuclei (Ng) with clumped chromatin (c), vacuoles (V) in cytoplasm (Cg), necrotic fragmented cells (K) with intact cell membrane. The surrounding neuropil contains distorted myelinated nerve fibers (NF) (TEM × 800×17) C) A photomicrograph of transmission electron microscope of cerebellar cortex of protected group shows granule cells (G) which have rounded or oval nuclei (Ng) with clumped chromatin (c), vacuoles in cytoplasm (Cg) which also contains mitochondria (m). (TEM × 1500×17)

Fig. 5: a) A photomicrograph of transmission electron microscope of cerebellar cortex of adult control albino rat showing a Purkinje cell (P) which is large flask shaped with euchromatic nucleus (N) with indentations (green arrows), prominent nucleolus (n) and cytoplasm (C) containing mitochondria (m) and free ribosomes (r). (TEM × 800×17) b) A photomicrograph of transmission electron microscope of cerebellar cortex treated adult albino rat shows Purkinje cell (P) with euchromatic nucleus (N) with prominent nucleolus (n); its cytoplasm (C) contains marked dilated rough endoplasmic reticulum (RER) and mitochondria with destructed cristae (m*). An astrocyte (A) with sharp demarcated nucleus (Na) with vacuolated cytoplasm (Ca) and mitochondria (m) (TEM × 800×17). c) A photomicrograph of transmission electron microscope of cerebellar cortex of protected adult albino rat shows Purkinje cell (P) contains euchromatic irregular nucleus (N) and prominent nucleolus (n) with indentations all around ill-defined nuclear membrane (arrows) and its cytoplasm (C) is dark, containing mitochondria (m), strands of mildly dilated rough endoplasmic reticulum (RER) and free ribosomes (r) (TEM × 800×17).
Fig. 6 a) A photomicrograph of transmission electron microscope of a section of cerebellar cortex of adult control albino rat shows an astrocyte (A) with sharply demarcated nucleus (Na) and electron lucent cytoplasm (Ca) containing mitochondria (m). The astrocytes are present near the blood capillary (b,c) which is lined by endothelial cells (e). The surrounding neuropil contains myelinated nerve fibers (NF). Oligodendrocyte (O) has irregular deeply stained nuclei (No). (TEM × 800×17) b) A photomicrograph of transmission electron microscope of a section of cerebellar cortex treated adult albino rat shows astrocytes (A) with sharp demarcated nucleus (Na) with chromatin and vacuolated cytoplasm (Ca) which contains mitochondria (m*). The surrounding neuropil contains unmyelinated nerve Fibers (NF) and granule cell (G). Purkinje cell (P) appears with its distinguished damaged appearance.(TEM × 1000×17) c) A photomicrograph of transmission electron microscope of a section of cerebellar cortex of protected adult albino rat shows astrocytes (A) with sharp demarcated nucleus (Na) with chromatin (blue arrows) and vacuolated cytoplasm (Ca) containing mitochondria (m) and few vacuoles (V).(TEM × 800×17)

Table 1 Comparison between body weights of group 1 (control), group 2 (MSG- treated) and group 3 (MSG +vit C treated).

<table>
<thead>
<tr>
<th>Group</th>
<th>Final body weight at the end of experiment (g)</th>
<th>One-way ANOVA</th>
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<tbody>
<tr>
<td></td>
<td>Range</td>
<td>Mean ± SD</td>
</tr>
<tr>
<td>Control</td>
<td>210 - 270</td>
<td>230.33 ± 15.98</td>
</tr>
<tr>
<td>MSG+ vit C treated</td>
<td>230 - 255</td>
<td>244.00 ± 9.62</td>
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<tr>
<td>MSG treated</td>
<td>295 - 320</td>
<td>308.00 ± 10.37</td>
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</tbody>
</table>

Table 2 Comparison between number of Purkinje cells of group 1 (control), group 2 (MSG- treated) and group 3 (MSG +vit C treated).

<table>
<thead>
<tr>
<th>Group</th>
<th>No. of Purkinje cells (Mean ± 2 SD)</th>
<th>Kruskal-Wallis test</th>
<th>P-value</th>
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</thead>
<tbody>
<tr>
<td>MSG Treated</td>
<td>2.1 ± 1.29</td>
<td>15.661</td>
<td>0.000*</td>
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<tr>
<td>MSG+ Vit. C Treated</td>
<td>4.8 ± 1.87</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>6.1 ± 2.33</td>
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Table 3 Comparison between Molecular cell layer thickness of group 1 (control), group 2 (MSG-treated) and group 3 (MSG + vitC treated).

<table>
<thead>
<tr>
<th>Group</th>
<th>Mean ± SD</th>
<th>One-way ANOVA</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSG treated</td>
<td>114.3 ± 19.9</td>
<td>F 5.737, P-value 0.008*</td>
</tr>
<tr>
<td>MSG+ vit C treated</td>
<td>145.1± 24.5</td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>145.5 ± 25.9</td>
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DISCUSSION
The present study demonstrated that there was a significant increase of the rats’ weight from the beginning to the end of the study which may be caused by the fact that MSG increases appetite, and this is in agreement with El-Helbawy et al. (14) who documented that the effects of MSG evaluated after a period of 14 days’ treatment in young rats showed a statistically highly significant increase (P<0.001) in the mean body weight of rats compared with controls. Another study also was in agreement with current one, demonstrating that low and high doses of MSG treatment increases body weight (15).

The present study clarified that, the molecular cell layer appeared as a zone lying superficial to Purkinje cell layer. The findings of the present study were in agreement with Young and Heath (16) who conveyed that, the mature cerebellar cortex consisted of three layers.

The current study demonstrated that, in control group the Purkinje cells were oval or flask-shaped with big rounded nuclei and prominent nucleoli with the granular cell layer with its distinguished rounded cells. These findings were in agreement with Hashem et al. (9).

With exposure to MSG, the present work elucidated degenerative effects in the form of necrotic Purkinje cells with statistically significant decrease in its number with spacing in between mostly due to edema which was observed on the rat cerebellar cortex. This was in line with Espinar (17) and Eweka and Om’Iniabohs (18).

In addition, Ureña-Guerrero et al. (19) found that, MSG produces neuro-degeneration with severe damaging of the cells in several brain regions when it was administered to neonatal rats, from an early embryonic age to adulthood. These toxic effects have been explained by Pavlovic et al. (20) who mentioned that intake of high concentrations of MSG induced oxidative stress in many organs. All these findings were in agreement with present results. On contrary with Jinap and Hajeb (21) who mentioned that, the Joint Expert Committee on Food Additives of the United Nations Food and Agriculture Organization and World Health Organization sited MSG in the safest class for food additives.

The immune-histochemical studies of control all groups of the current study revealed presence of a positive reaction of glial fibrillary acidic protein (GFAP). The foot processes of astrocytes are thin, short and run in parallel rows in the molecular layer, as Hashish (22) had reported.

In MSG- treated group astrocytes which appeared larger than that in control ones with longer irregular processes across the cerebellar cortex, and this was in agreement with Espinar (17).

As regarding the MSG + vit. C treated group, there was an improvement in all layers of the cortex which appeared in the form, Purkinje cells were mostly keeping their normal appearance with few necrotic cells with less spacing comparing to the MSG- treated group. This was confirmed by a highly statistically significant increase in number of Purkinje cells of the MSG + vit.C treated group in comparison with MSG- treated group with p value < 0.0000 as mentioned by Narayanan et al. (23) and Huang et al. (24).

The GFAP immune-expression of the present study showed increase in group receiving vitamin C, when compared with other groups.
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which was accompanied with preservation of the neurons in the form of long more parallel processes of astrocytes but still are irregular; this was as Hanbury et al. (25) and Hughes et al. (26) documented.

As regarding the ultrastructural results of electron microscopy of the present study, in the control group; the granular layer contained numerous granular cells having rounded or oval nuclei with peripheral or central condensation of chromatin, their thin cytoplasm contains free ribosomes and mitochondria, as documented in previous two studies (13; 27).

The present results revealed that Purkinje cells were fusiform in shape, they contained euchromatic irregular nuclei with prominent nucleolus and their cytoplasm is dark, contains few mitochondria and strands of rough endoplasmic reticulum. This was in agreement with EL Tantawi and El Namshan (28).

The cerebellar cortex of animals treated with MSG showed the nuclei of many Purkinje and granule cells appeared shrunken and densely stained with irregular nuclear envelopes; this was in agreement with other related studies (29; 30; 31).

Ultra-structural observations on vitamin C use showed that most of the granule and Purkinje cells were remarkably healthy while few ones exhibited scarcely degenerative features. The improving effects of vitamin C may be attributed to its antioxidant constituents which increases the circulating level of antioxidants leading to decreasing the oxidative stress induced by MSG. This was in line with Farombi et al. (11).

Conclusion

In conclusion, MSG has toxic effect not only on nerve cells but also on astrocytes, which were reported to protect nerve cells from toxic insults, and hence came the dangerous neurotoxic effect of MSG. Vitamin C supplementation could protect from neurotoxic effect of MSG

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REFERENCES


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21. Jinap S, Hajeb P. Glutamate Food Safety Research (CEFSR), Faculty of Food Science and Technology, Universiti Putra Malaysia, 2010; 43400 UPM, Serdang, Selangor, Malaysia.


25. Hanbury R, Ling ZD, Wuu J, Jeffrey K. GFAP knockout mice have increased levels of GDNF that protect striatal neurons from metabolic and excitotoxic insults. J Comp Neurol., 2003 461:307-316


27. Mahran HA, Arisha SM. The ameliorative effects of the aqueous extract of rosemary against monosodium glutamate neurotoxicity in adult male albino rats: histological, ultrastructural and biochemical studies. Ejpmr, 2018; 5(1), 79-90


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