The Possible Neuroprotective Effect of Astaxanthin on Monosodium Glutamate and Aspartame Induced Hippocampal Changes in Albino Rats: (Histological and Immuno-histochemical Study)

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ABSTRACT

Background: Food additives are substances added to food to improve its safety, freshness, taste, texture, or appearance. They include aroma enhancer eg : monosodium glutamate(MSG) or sweetener eg.: aspartame (ASP).

Aim of the Study: This current study was performed to study the microscopic alterations induced by MSG, ASP individually and in combination and the possible protective effect of astaxanthin (AST) on those induced changes.

Materials and Methods: Forty nine adult male albino rats were used in this study divided into; control group (fourteen rats) two rats for each experimental group and experimental group (thirty five rats) subdivided into seven subgroups:ASP group, MSG group, ASP andMSG group,AST group,ASP andAST group, MSG andAST group, ASPandMSG andAST group. Oral administration was done in the morning daily for 6 weeks for all groups. When the duration of the study ended, blood samples were collected from rat tail and paraffin sections were prepared from the cerebral hemisphere of each animal. They were subjected to hematoxylin and eosin stain and immunohistochemical stain for Caspase3 and Glial Fibrillary Acidic Protein (GFAP). Statistical analysis were done for assessment of body weight, reduced glutathione (GSH) and tumor necrosis factor α (TNF-α) level.

Results: The groups that received ASP (I), MSG (II) individually or in combination (III) exhibited shrunken cells with darkly stained nuclei and surrounded with pericellular haloes and some areas even revealed loss of the cells with increase in immunoreactivity for GFAP and Caspase 3. These groups also showed elevation in the level of TNFα and decrease in the level of GSH. On treatment with AST groups V, VI and VII showed reduced pycknosis, decreased immunoreactivity for GFAP and Caspase 3. It also showed reduction in the level of TNFα and increased level of GSH.

Conclusion: ASP and MSG individually or in combination induce alternation in hippocampus and AST administration ameliorate those changes.

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Key Words: Aspartame, astaxanthin, hippocampus, monosodium glutamate.

INTRODUCTION

Junk foods have proved to be a big problem for human health in general and especially for children, as they contain synthetic flavors and preservatives. Food additives contain two main types, which include aroma enhancers e.g monosodium glutamate and sweeteners e.g aspartame[1,2].

Monosodium glutamate (MSG) (E621) is a salt form of L-glutamic acid and is widely used all over the world to enhance the flavor. It was reported that the mean amount ingested from MSG in the individual within industrialized countries is about 0.3–1.0 g/day. Glutamate, the amino acid in MSG, when taken in extra amounts, it acts as a strong neurotoxin as it increases the excitability and activity of proteolytic enzymes[3].

Aspartame (ASP) is an artificial sweetener that is being used in alteration to sugar in many food products. It is approximately 200 times sweeter than sugar so, used in low-calorie soft drinks and foods[4]. It is rapidly absorbed and metabolized by intestinal esterase and dipeptidase to 40% aspartic, 50% phenyl alanine and10% methanol. Phenylalanine has an important role as a regulator of the process of neurotransmition, moreover, it was reported that aspartic acid has an important role as an excitatory neurotransmitter in the central nervous system[5].

Astaxanthin (AST) is a carotenoid that is separated from shrimp and crab shells. It has strong antioxidant properties. It was revealed that AST has an effective role in scavenging oxygen free radicals, decrease lipid peroxidation, oxidative stress and inhibit reactive oxygen species (ROS)-mediated toxicity[6].

AIM OF THE WORK

The purpose of this study is to shed light on the changes induced by monosodium glutamate and aspartame separately or in combination in hippocampus and to demonstrate possible protective role of astaxanthin on those induced changes.
MATERIALS AND METHODS

a) Materials used

Monosodium Glutamate

Monosodium glutamate (MSG) powder of purity ≥99% was purchased from Beta pharm. It was administrated, orally, via intragastric tube once daily for 6 weeks (120 mg/kg dissolved in distilled water)[7].

Aspartame

Aspartame powder was purchased from beta pharm and administered to the rats by intragastric tube for 6 weeks (250 mg/kg dissolved in distilled water)[8].

Astaxanthin

It was obtained from BIOVE in the form of soft gel container and administered via intragastric tube for 6 weeks (2 mg/kg dissolved in version olive oil)[9].

b) Animals

The study included forty nine adult male albino rats with a weight range of (150-200 gm). They were obtained and handled according to the guidelines after the approval of the ethical committee (CU III F 87 17) from the Animal House-Faculty of Medicine at Cairo University. All the animals were handled for one week prior to experiment to minimize nonspecific stress on the days of the experiment, and were kept at room temperature (24-26°C) with a 12:12h light:dark cycle. They were provided with standard crushed food and water during the experiment. The bodies of the dead animals were disposed of by using the incinerator.

Control Group

Fourteen rats were used, two rats for each of the corresponding experimental groups.

- Six rats received 0.5 ml of distilled water
- Two rats received 0.5 ml of version olive oil
- Six rats received both 0.5 ml of distilled water and 0.5 ml of virgin olive oil.

All animals received the solvent orally once daily via intragastric tube.

Experimental Groups

They were subdivided to seven groups that received the drugs orally via intragastric tube once daily for six weeks.

I. ASP group: five rats received 250 mg/kg ASP dissolved in 0.5ml of distilled water. Dose consumptions between rats and humans were corrected by factor 5 as aspartame metabolizes in rats faster than that in humans.

II. MSG group: five rats received 120 mg/kg MSG dissolved in 0.5ml of distilled water

III. ASP&MSP group: five rats received ASP as in group I and MSG as in group II.

IV. AST group: five rats received AST (2 mg/kg) dissolved in 0.5ml virgin olive oil.

V. ASP& AST group: five rats received ASP as in group I and AST as in group IV.

VI. MSG &AST group: five rats received MSG as in group II and AST as in group IV.

VII. ASP, MSG &AST group: five rats received ASP as in group I, MSG as in group II and AST as in group VI.

The body weight of each rat in all groups was measured at the beginning and at the end of the experiment. Statistical study on the mean weight was done.

At the end of 6th week, blood samples were collected from rat tail. Under general anesthesia (sodium pentobarbital 100mg/kg) administrated intraperitoneally[10], the animals were perfused via the left ventricle with phosphate buffer (PH 7.4) containing 2% paraformaldehyde[11]. Their cerebral hemisphere was dissected and cut sagittally into right and left halves.

I. Light Microscopic Study

The left half of the cerebral hemisphere from each animal was fixed in 10% formalin solution, and then specimens were processed for light microscopic study. Paraffin sections (5 μm thick) were stained with the following:

1. Haematoxylin and eosin stain (Bancroft and Layton, 2013)[12]
2. Immunohistochemical stain for (Jackson and Blythe, 2013)[13].
   - Caspace3 for apoptotic cells
   - Glial Fibrillary Acidic Protein (GFAP) of the astrocytes

2- Statistical Analysis

Body weight, TNF-a level and GSH level of control and experimental animals were subjected to statistical analysis using the PASW software system version 16. The statistical analysis of data was done by using Excel program and statistical package for social science (SPSS) program on windows XP. Comparison between normally distributed (parametric) quantitative data of more than two groups was conducted using ANOVA (analysis of variance) test followed by post-hoc for multiple comparisons

NB: The P value of ≤ 0.05 indicates significant results at confidence interval 95%[14].

3- Morphometric Study

Using Leica Qwin 500 LTD image analysis in medical histology department of Cairo University, the following parameters were measured:

A. Area percent of Caspase 3 +ve cells in Caspase 3 immunostained sections.
The pyramidal layer showed many large pyramidal neurons with vesicular nuclei. The other two layers were devoid of neurons. Scanty number of nuclei of neuroglial cells as astrocytes was detected in these layers beside blood vessels on a pink neuropil background formed of neuronal and glial cell processes (Figure 3A).

In group I (ASP), some pyramidal cells were shrunken with dark stained nuclei and others with pale nuclei. Also some vacuolations appeared in the neuropil (Figure 3B).

In group II (MSG), the pyramidal layer showed marked disorganization with areas of cell loss. Most of large pyramidal cells were shrunken with dark stained nuclei, while few cells showed pale nuclei. There were also vacuolation in neuropil (Figure 3C).

In group III (ASP and MSG) the pyramidal cells layer showed same results as group II (MSG) (Figure 3D). Both molecular and polymorphic layers of CA3 contained dilated blood capillaries, apoptotic dark stained neuroglia (Figures 3B, C and D).

In group IV (AST) the results were similar to the control (Figure 3E). In group V (ASP and AST) and group VI (MSG and AST) the results were comparable to control but still there were few shrunken cells with darkly stained nuclei (Figures 3F and 3G).

In group VII (ASP, MSG and AST) showed scattered pyramidal cells. It had vesicular nuclei relatively similar to control (Figure 3H). In the remaining treated groups, both molecular and polymorphic layers showed relatively normal appearance.

Results of Dentate Gyrus

In all rats of control group, the dentate gyrus was formed of molecular, granular and polymorphic layers. The granular layer is the main layer. It was composed of dense aggregate of granular cells that appeared rounded with vesicular nuclei. The polymorphic layer contains scattered large pyramidal cells with vesicular nuclei among the neuroglia while the molecular layer was formed of neuroglial cells (Figure 4A).

In group I (ASP), the granular layer showed areas of cell loss and some cells were shrunken with marked vacuolation in the neuropil. The molecular layer contained apoptotic neuroglial cells (Figure 4B). Similar results were seen in both group II (MSG) and group III (ASP and MSG) but the polymorphic layer of group III contained dark stained pyramidal cells surrounded by halos (Figures 4C and 4D).

In group IV (AST) the results similar to the control (Figure 4E). In all AST treated groups V, VI, VII the results were comparable to control group with vacuolation in groups VI and VII (Figures 4F and G and H).

2) Immunohistochemical Results

Immunostained sections of hippocampus of all control groups showed few GFAP positive immunoreactive astrocytes dispersed in CA1, CA3, dentate gyrus.
(Figures 5A, B, C). Also, Examined control sections of rats hippocampus showed negative immunostaining with sparse positive immunoreactive cells for Caspase 3 (Figures 6A, B, C).

In group I (ASP), CA1, CA3 and dentate gyrus showed strong GFAP immunoreactive star – shaped astrocytes with intense brownish twisted irregular glial fibers (Figures 7A, B, C). Caspase 3 immunostained section showed strong positive immune staining for hippocampus proper CA1, CA3 and dentate gyrus (Figures 8A, B, C). Similar results were observed in group II (MSG) and group III (ASP and MSG) (Figures 9, 10, 11(A,B,C)). But group III (ASP and MSG) showed moderate reaction for caspase 3 immunostaining in CA1, CA3 and dentate gyrus (Figures 12A, B, C).

In group IV (AST) both GFAP and caspase 3 immunostained sections of hippocampus showed results similar to control (Figures 13 and 14 (A,B,C)).

All treated groups (V, VI, VII) by AST showed moderate positive GFAP immunoreactive astrocytes with faint brownish glial fiber (Figures 15, 17 and 19, (A,B,C)) and moderate reaction for caspase 3 (Figures 16, 18 and 20 (A,B,C))

2- Statistical Study

i- Body Weight

There was no significant difference in the mean of body weight of group III (MSG administrated group), group V (AST treated group), group VI (ASP +AST treated group), group VII (MSG +AST treated group) and group VIII (ASP, MSG and AST) when compared with control rats. On other hand, there was significant decrease in the mean of body weight of rats in both group II (ASP administrated rats) and group IV (ASP +MSG administrated group) with (p < 0.001) when compared with control one (Table 2 and histogram 3).

ii- Assessment of Reduced Glutathione (GSH) and Tumor Necrosis Factor α (TNF-α) Level

The mean GSH level in ASP, MSG administrated group individually or in combination showed a highly significant decrease as compared with that of control rats with (P <0.001). However, the mean GSH level in treated groups with AST treated group showed a significant increase as compared with the of ASP, MSG administrated rats individually and in combination (Table 1 and Histogram 1)

The mean TNF α level in ASP, MSG administrated group individually and in combination showed a highly significant increase as compared with that of control rats with (P <0.001). However, the mean in AST treated groups showed a significant decrease as compared with that diseased rats (group I,II,III) with (P <0.001) (Histogram 2)

iii- Morphometric Results

1) GFAP immunoreactivity (Table 3 and Histogram 4)

The mean area percent of positive GFAP immunoreactivity in CA1, CA3 and dentate gyrus of group IV (AST) showed no significant difference as compared with that of control rats.

Groups (I, II and III) showed a highly significant increase in immunoreactivity compared to the control rats with (P < 0.001). However, in groups (V, VI and VII) the mean area percent of positive GFAP immunoreactivity showed a significant decrease as compared with that of groups (I,II and III) with (P < 0.001) and non-significant increase as compare with control rats with (P > 0.05).

2) Caspase 3 immunoreactivity: (Table 4 and Histogram 5):

The mean area percent of positive Caspase 3 immunoreactivity in CA1, CA3 and dentate gyrus of group IV (AST) showed no significant difference as compared to control group.

Groups (I, II and III) showed a highly significant increase the mean area percent of positive Caspase 3 immunoreactivity compared to control rats with (P <0.001). However, groups (V, VI and VII) the mean area percent of positive caspase 3 immunoreactivity showed a significant decrease compared to groups (I ,II and III) with (P <0.001) and non-significant increase compared to control group with (P > 0.05).
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Fig. 1: A photomicrograph of a section in hippocampal formation of control rat showing the Cornu Ammonis (CA) as CA1, CA2, CA3 & CA4. Dentate gyrus (DG) is seen surrounding CA4 by its upper and lower limbs. (M) denotes molecular layer inside concavity of CA and of DG (H & E x40).

Fig. 2: A photomicrographs showing sections in CA1:(A) control group showed the compact layers of small pyramidal cells of CA1 which have vesicular nuclei (arrows). Both molecular layer (ML) and polymorphic layer (PL) show many glial cells (astircks) among neuropil and blood vessels (arrow head ). (B) group I (ASP) showed decreased thickness of pyramidal layer with many shrunken cells that have darkly stained nuclei and surrounded with preicellular haloes (arrows). Note, absence of pyramidal cells in few areas (curved arrows). Both molecular and polymorphic layers contain congested blood capillaries (astircks). Many apoptotic neuroglial cell (arrow head). In both (C) group II (MSG) and (D) group III (ASP & MSG) some pyramidal cells had pale-stained nuclei, and distorted cellular morphology (arrow head) and other cells were shrunken with darkly stained nuclei and surrounded with preicellular space (arrow). Also, the pyramidal layer in group (C) contain congested blood vessel (black astircks). (E) group IV (AST) the layers were comparable to control group. (F) group V (ASP & AST) showed compact layer of small pyramidal cells of CA1 similar to control most of them have vesicular nuclei (arrows). Note, few shrunken cells with darkly stained nuclei and surrounded with haloes (arrows head). Both molecular layer (ML) and polymorphic layers show many glial cells among neuronal processes (astircks) and blood vessel (curved arrows). Both (G): group VI (MSG & AST) and (H) group VII (ASP, MSG & AST) showed similar result as group V (ASP & AST). (H & E x 200).
Fig. 3: photomicrographs showing sections in CA3: (A) control group showed the three layers of CA3 region of hippocampus proper, molecular layer (ML), pyramidal layer, polymorphis layer (PL). Pyramidal layer contain large pyramidal cells (arrow) with vesicular nuclei. Both molecular and polymorphic layers show neuroglial cells (asterisks) and blood vessel (curved arrows). (B) group I (ASP) showed many shrunken pyramidal cells with dark stained nuclei (arrow) and vacuolated neuropil (asterisks). Both molecular and polymorphic layers contain many apoptotic neuroglial cells (arrow head) surrounded by empty spaces. In both (C) group II (MSG) and (D) group III (ASP & MSG), the pyramidal cell layer showed areas of cell loss (arrows). Most of the large pyramidal cells were shrunken with dark stained nuclei while, few cells had pale nuclei (arrow head) and neuropil showed vacuolation. However, there are clumping of neuronal process (curved arrows). (E) group IV (AST) the layers were comparable to control group. (F) group V (ASP & AST) most of the large pyramidal cells had vesicular nuclei (arrows). In Both (G): group VI (MSG & AST) and (H) group VII (ASP, MSG & AST) showed similar result as group V (ASP & AST). In addition group VI had few cells with pyknotic nuclei and surrounded by halo (arrow head). There are also dilated blood vessel (curved arrow). Group VII showed scattered pyramidal cells. (H&E x 200)

Fig. 4: photomicrographs showing sections in dentate gyrus: (A) control rats showed dentate gyrus forms of molecular layer (ML), granular layer (G) and polymorphic layer (PL). It was formed of dense aggregate of granular cells that appeared rounded with vesicular nuclei (arrow). Dark cells are seen in subgranular zone (arrow head). The polymorphic layer contains scattered large pyramidal cells (p) with vesicular nuclei among the neuroglia while the molecular layer was formed of neuroglial cells (asterisks). (B) group I (ASP) showed many granular cells shrunken (arrow) and some areas with cell loss (curved arrow). Note, marked vacuolation (V) in neuropil and apoptotic dark stained neuroglia (asterisks) in molecular (ML) layer. In both (C) group II (MSG) and (D) group III (ASP & MSG) were showed similar results as group I (ASP). In addition to dark stained apoptotic large pyramidal cells (p) in the polymorphic layer (E) group IV (AST) showed comparable to control group. Note, vacuolation (V) in both groups VI & VII. (H&E x 200)
**Fig. 5:** Photomicrographs of control sections of rat brain showing weak positive immunostaining for GFAP in hippocampus (A) CA1, (B) CA3, (C) dentate gyrus. (Anti GFAP immunostain x200).

**Fig. 6:** Photomicrographs sections of control rats showing negative immune staining for caspase 3 with few positive immune reactive cells (arrows) in hippocampus proper; (A) CA1, (B) CA3 and (C) dentate gyrus. (Anti caspase 3 immunostain x200).

**Fig. 7:** Photomicrograph of ASP administrated sections of hippocampus proper (A) CA1, (B) CA3, (C) dentate gyrus showing strong immunoreactive star-shaped astrocytes (arrows) with intense brownish immunoreaction in glial fibers which appeared twisted and had irregular course (arrow head) (Anti GFAP immunostain x200).

**Fig. 8:** Photomicrographs of ASP administrated rats showing strong reaction for caspase 3 in rat (A) CA1, (B) CA3 and (C) dentate gyrus (arrow). (Anti caspase 3 immunostain x200)
Fig. 9: Photomicrographs of MSG administrated sections in (A) CA1, (B) CA3, (C) dentate gyrus showed strong immunoreactive star-shaped astrocytes (arrows) with intense brownish immunoreaction in glial fibers which appeared twisted and had irregular course (arrow head). (Anti GFAP immunostain x200).

Fig. 10: Photomicrograph in sections of MSG administrated rats showing strong positive immune staining for caspase 3 in (A) CA1, (B) CA3 and (C) dentate gyrus (arrows), some other with moderate reaction (crossed arrow). (Anti caspase 3 immunostain x 200).

Fig. 11: Photomicrographs of ASP and MSG administrated sections of hippocampus proper (A) CA1, (B) CA3, (C) dentate gyrus. Showing strong immunoreactive star-shaped astrocytes (arrows) with intense brownish immunoreaction in glial fibers which appeared twisted and had irregular course (arrow head). (Anti GFAP immunostain x200).

Fig. 12: Photomicrographs section of ASP and MSG administrated rats showing moderate positive immune staining for caspase 3 in hippocampus proper (A) CA1, (B) CA3 and (C) dentate gyrus (crossed arrow) and few cells showing strong reaction (arrow). (Anti caspase 3 immunostain x 200).
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Fig. 13: Photomicrographs of AST treated rat hippocampus showing weak positive immunostaining for GFAP (A) CA1, (B) CA3, (C) dentate gyrus (Anti GFAP immunostain x200).

Fig. 14: Photomicrographs of sections in AST treated rats showing negative immune staining for caspase 3 with few positive immune reactive cells (arrows) in hippocampus proper; (A) CA1, (B) CA3 and (C) dentate gyrus. (Anti caspase 3 immunostain x200)

Fig. 15: Photomicrographs of ASP and AST sections showing moderate positive immunostaining in the astrocytes for GFAP in (A) CA1, (B) CA3, (C) dentate gyrus (Anti GFAP immunostain x200)

Fig. 16: Photomicrographs of ASP and AST treated rats showing few cells with moderate positive immune staining for caspase 3 in (A) CA1, (B) CA3 and (C) dentate gyrus (Anti caspase 3 immunostain x200)
Fig. 17: photomicrographs of MSG and AST administrated sections of hippocampus proper (A) CA1, (B) CA3, (C) dentate gyrus Showing moderate immunoreactive star–shaped astrocytes (arrows) with faint brownish immunoreaction in glial fibers (arrows head) (Anti GFAP immunostain x200).

Fig. 18: photomicrographs of MSG and AST treated rats showing few cells with moderate positive immune staining for caspase 3 in (A) CA1, (B) CA3 and (C) dentate gyrus (Anti caspase 3 immunostain x200).

Fig. 19: photomicrographs of ASP, MSG and AST sections of hippocampus (A) CA1, (B) CA3, (C) dentate gyrus Showing normal distribution as weak positive reaction in the immuno-reactive astrocytes. (Anti GFAP immunostain x200)

Fig. 20: photomicrographs of ASP, MSG and AST administrated rats showing few cells with moderate positive immune staining for caspase 3 in hippocampus proper in (A) CA1, (B) CA3 and (C) dentate gyrus (Anti caspase 3 immunostain x200).
**Table 1:** The mean and SD of GSH and TNFα levels in the control and experimental group

<table>
<thead>
<tr>
<th></th>
<th>Control group</th>
<th>ASP group</th>
<th>MSG group</th>
<th>ASP+MSG group</th>
<th>AST group</th>
<th>ASP+AST group</th>
<th>MSG+AST group</th>
<th>ASP+MSG+AST group</th>
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<tr>
<td><strong>GSH Mean±SD</strong></td>
<td>68.1±4.3</td>
<td>25.7±8.4</td>
<td>36.8±9.1</td>
<td>23.5±7.8</td>
<td>48.8±11.4</td>
<td>55±5.6</td>
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<td><strong>TNFα Mean±SD</strong></td>
<td>27.5±2.4</td>
<td>116.25±25.6</td>
<td>94.25±25.1</td>
<td>115±20.8</td>
<td>60±15.6</td>
<td>46.8±13.9</td>
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**Table 2:** The mean ± SD of body weight of the control and experimental group

<table>
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<th>Control group</th>
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<th>ASP+MSG group</th>
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<tr>
<td><strong>Body Weight</strong></td>
<td>27.5±2.4</td>
<td>116.25±25.6</td>
<td>94.25±25.1</td>
<td>115±20.8</td>
<td>60±15.6</td>
<td>46.8±13.9</td>
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**Table 3:** The mean ± SD area percent of positive GFAP in rats hippocampus of control and experimental groups

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</thead>
<tbody>
<tr>
<td><strong>CA1</strong></td>
<td>1.5±0.1</td>
<td>5.12±0.15</td>
<td>5.59±0.28</td>
<td>7.46±0.13</td>
<td>1.56±0.9</td>
<td>1.98±0.49</td>
<td>1.91±0.4</td>
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<td><strong>CA3</strong></td>
<td>1.46±0.18</td>
<td>8.23±0.7</td>
<td>8.57±1.3</td>
<td>7.84±0.85</td>
<td>1.22±0.17</td>
<td>1.88±0.21</td>
<td>1.5±0.3</td>
<td>1.5±0.2</td>
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<tr>
<td><strong>Dentate gyrus</strong></td>
<td>0.97±0.09</td>
<td>8.17±0.8</td>
<td>7.5±0.52</td>
<td>8.03±0.68</td>
<td>1.7±0.19</td>
<td>1.88±0.06</td>
<td>1.8±0.089</td>
<td>2.18±0.17</td>
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**Histogram 1:** The mean GSH levels in the control and experimental group

**Histogram 2:** The mean TNFα level in control and experimental group

**Table 4:** The mean ± SD of area percent of positive Caspase 3 in rats hippocampus of control and experimental groups

<table>
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<tbody>
<tr>
<td><strong>CA1</strong></td>
<td>0.54±0.09</td>
<td>2.25±0.12</td>
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<td>2.8±0.18</td>
<td>0.75±0.17</td>
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<td>0.65±0.2</td>
<td>0.69±0.22</td>
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<tr>
<td><strong>CA3</strong></td>
<td>0.25±0.1</td>
<td>2.39±0.8</td>
<td>2.86±0.9</td>
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<td>0.35±0.09</td>
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<td>0.81±0.09</td>
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<tr>
<td><strong>Dentate gyrus</strong></td>
<td>0.32±0.17</td>
<td>6.30±0.94</td>
<td>5.62±0.29</td>
<td>7.32±0.92</td>
<td>0.44±0.05</td>
<td>1.07±0.17</td>
<td>0.97±0.2</td>
<td>1.1±0.15</td>
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</table>
Histogram 3: The mean body weight of the control and experimental groups

Histogram 4: The mean area percent of positive GFAP in rats’ hippocampus of the control and experimental groups

Histogram 5: The mean area percent of positive Caspase 3 in rats’ hippocampus of the control and experimental groups
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DISCUSSION

Aspartame (ASP) is one of the most widely used sweeteners in the world. Aspartame is used frequently nowadays to reduce sugar consumption and decrease caloric intake in healthy people as well as in diabetic patients\(^{[15]}\). Several studies reported that chronic use of ASP leads to different neuropsychiatric and neurobehavioral disorders. It also shows marked histopathological changes in different regions of the brain by inducing oxidative stress\(^{[16]}\).

Monosodium glutamate (MSG) is one of the most controversial food additives locally and globally in terms of the safety in its usage\(^{[17]}\). MSG was proved to be toxic for both humans and experimental animals\(^{[18]}\).

In the present study, light microscopic examination of H&E stained sections of hippocampus of group I (ASP) revealed marked histopathological changes. The CA1 and CA3 showed gradual cell loss with a decrease in width of the pyramidal layer and there was no evidence of edema; however, in the dentate gyrus, there was cell death in the granular layer. Similar results were obtained by Onaolapo et al. 2017\(^{[19]}\).

ASP administration alone or in combination was associated by changes in brain morphology which had been occurred secondary to many factors. These were including; generation of large numbers of free-radical species (both nitrogen and oxygen types) that induced disintegration of cellular proteins and DNA, mainly the mitochondrial DNA and excitotoxicity effect of Aspartate\(^{[20]}\).

Histological examination of hippocampus sections in group II (MSG) showed degenerative changes in the pyramidal and granule cells of CA1, CA3 and dentate gyrus however some cells appeared with pale-staining nuclei. Similar findings were reported by Owoeye and Onwuka, 2015\(^{[21]}\).

Halo of empty spaces surrounding neuronal cells in groups (I,II&III) could be explained by the shrinkage, necrosis and apoptosis of cells leaving pericellular spaces. A similar observation was reported in other studies\(^{[22]}\). Pericellular spaces are formed secondary to cytoskeletal affection with subsequent shrinkage of cells and withdrawal of their processes. This finally led to vacuolation in the surrounding neuropil as recorded by Auer and Sutherland, 2002\(^{[23]}\). In the same groups, shrinkage of the cells and acidophilic cytoplasm of the cells could be attributed to ASP and MSG induced apoptosis. The neuronal death might be triggered by excitotoxins. A similar explanation has been reported by other authors as El-Samad, 2010\(^{[24]}\) who reported that excitotoxicity in the neurons is due to the ability of aspartate to open cationic channels leading to uncontrollable calcium (and to a lesser extent sodium) influx, triggering a cascade of intracellular enzyme-controlled reactions culminating in cell death. Most of cells appeared small, darkly stained and surrounded by empty spaces. The failure of the antioxidant mechanisms led to accumulation of free radicals and uncompensated oxidative stress with subsequent accumulation of denatured proteins and dark staining of the degenerated cells. Some authors agreed with this explanation\(^{[25,19]}\).

In the present study, GFAP stained sections of group I (ASP) revealed an apparent increase in the number of the star-shaped astrocytes with intense GFAP immunoreaction. Similar results with ASP treatment were observed by many authors\(^{[26]}\). Increased GFAP-reactivity has also been associated with neuronal injury\(^{[27]}\). Aspartame induces injury in the central nervous system which is compensated by an increase in the astrocytic number (reactive astroglisis). These findings were in accordance with recent studies Cho et al. 2016\(^{[28]}\). Similar results obtained in both group II (MSG) and group III (ASP and MSG). It was reported that an increase in astrocyte immunoreactivity usually occur secondary to toxic insult, because astrocytes share widely in glutamate homeostasis and are important in the reuptake of free glutamate/prevention of glutamate excitotoxicity\(^{[29]}\).

Programmed cell death (apoptosis) might have occurred in response to exposure to toxins\(^{[30]}\). ASP and MSG individually or in combination have been found to be toxic, especially to the nervous system, even in small doses\(^{[31]}\). This was supported by caspase 3 results which revealed strong positive immunoreactions in hippocampus. Similar finding was reported by Omar et al. 2016\(^{[13]}\). Different mechanisms have been proposed for cell death and apoptosis may include the glutathione levels allowing the generation of neurotoxic ROS\(^{[32]}\) and aspartame is also capable of damaging the essential cellular components such as nucleic acid, gene and gene repair activity that can lead to apoptosis\(^{[13]}\).

The reduction in GSH content in groups (I,II and III) might be due to mitochondrial dysfunction and oxidative stress that increase the accumulation of toxicity\(^{[33]}\). While, there was elevation in the level of TNFα in these groups compared to control group which was supported by statistical study. This finding was similar to the results of Riazi et al. 2010\(^{[34]}\). Which explained by Ashok and Sheeladevi, 2014\(^{[13]}\) who observed that aspartame causes increase in the brain TNFα level (a potent proinflammatory cytokine) that is produced by glial cells. Pro-inflammatory cytokines mediate the inflammation and contribute to excitotoxic neuronal damage. High TNF α level in MSG administered group due to accumulation of extracellular glutamate to a level that was high enough to inhibit synaptic activity and kill neurons\(^{[36]}\).

Our results showed that groups treated by AST associated by an increase in the level of GSH. Chan et al. 2009\(^{[37]}\) found that astaxanthin treatment showed neuroprotective effect via the restoration of glutathione level and reduction of ROS formation. Mattei et al. 2011\(^{[38]}\) also demonstrated how AST administration leads to improved antioxidant capacity including increase ratio of GSH: GSSG in the plasma. Results of the present
study indicate that AST decreased TNF-α concentration in all groups (VI, VII and VIII) treated with it. TNF-α inhibition by AST resulting from inhibited NOS activity and decreased NO production. The mechanism of the NO-induced suppression of TNF synthesis is not known this finding supported by Ohgami et al. 2003(40).

AST is potent antioxidant activity, reported to be much more effective than other similar compounds. As it work by different mechanisms as it can reduce radicals by absorption, donation of electrons, and formations of adducts with the reactive species due to presence of hydroxylated ion rings that cap both ends of the carbon backbone distinguish the molecule from other carotenoids in its class, and enables AST energetically favorable spanning the phospholipid bilayer of cell membranes. This orientation and chemical structure effectively protects the membrane against lipid peroxidation(40).

In our present study, we found that astaxanthin treated groups (V, VII and VII) reduce the caspase 3 activity. Lu et al. 2010(41) found that astaxanthin decreases the extent of ischemic infarction in rats by increasing the activities of antioxidants and by maintaining mitochondrial membrane potential (MMP). It is highly possible that these agents via increasing activity of antioxidant enzymes and/or GSH level decreased oxidative stress, which consequently enhanced the stability of mitochondrial membrane, plasma membrane, and transmembrane proteins. Therefore, the application of this agent might provide greater neuroprotection.

In present study, GFAP stained sections of rats treated with AST revealed fewer star-shaped astrocytes with less intense GFAP immunoreaction. This finding coincided with El-Samad (2010)(42) an author who reported that neuroglial cells might support the neuronal recovery through release of some cytokines and growth factors. It seemed that the recovery period was not enough to restore completely the normal structure of the cerebellar cortex. The reduction in the number of astrocytes could be attributed to a gradual return to the normal structure of neurons, which was reflected by the decrease in the number of astrocytes. Astrocytes are a central component of the brain’s antioxidant defense system and activation of astrocytes involves neurotoxic actions induced by oxidative insults(43).

In group II (MSG) showed significant alteration in body weight compared with control group. The effects of MSG on body weight were variable and conflicting according to the previously reported results whether increase(44), decrease(45) or reveal no change(46). This could be explained by change in the body metabolism that occur secondary to increased adipocyte capacity to transport glucose and synthesize lipids, leading to increased insulin sensitivity(47). On the contrary, there was no change in the body weight of rats that were given MSG (in food or water) in the study conducted by Tordoff et al. 2012(48). There was a significant reduction of weight in aspartame administrated group compared with control group. Similar finding was observed by Magalhães et al. 2018(49) because aspartame showed interference with metabolism, liver function and appetite of animals, increasing satiety and thus reducing food intake. However, in the current study, the body weight of the animals that were treated with MSG and ASP were lower than those within the control group. These results came in accordance with other reported studies(50). On the other hand, some studies showed opposite results to these reported in the current study(50). The explanations for these differences could be due to the use of different doses or different duration of exposure and use of animals from other strains.

CONCLUSIONS

From this study, it could be concluded that

1. Aspartame (ASP) and Monosodium glutamate (MSG) individually or in combination induced histopathological and immunohistochemical alterations in hippocampus.
2. Astaxanthin (AST) administration ameliorate those associated changes in rat hippocampus through possible neuroprotective role.

RECOMMENDATION

It is recommended to avoid using ASP and MSG individually or in combination by avoiding the food and products containing it. Astaxanthin should be taken regularly as antioxidant to protect us against neuronal damage.

CONFLICTS OF INTEREST

There are no conflicts of interest.

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الملخص العربي

الوقاية المحتملة من عقار الاستاذنسين على التغيرات المستحدثة بجلوتامات أحادي الصوديوم والاسبرتام على مخ الجرذان البيضاء

(دراسة هستولوجية ومناعية هيستوكيمايئية)

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المقدمه: المحسنات الغذائية هي مواد مضافة إلى الطعام للحفاظ عليه ولتحسين سلامته، أو طعمه، أو ملمسه، أو مظهره. وهي تشمل محسّن الرائحة مثل: جلوتامات أحادي الصوديوم أو مواد تحلية مثل: الأسبارتام.

الهدف من البحث: دراسة التغيرات المجهرية والمرفومتريه التي يسببها كلا من الأسبارتام وجلوتامات أحادي الصوديوم على جهاز مخ انتقال الذرائع الوقائي الممكن من عقار الاستاذنسين على هذه التغيرات المستحدثة.

طرق ومواد البحث: في هذه الدراسة تم استخدام تسعة وأربعون قطعةً من الذرائع البيضاء البالغة تتراوح أوزانها من 150-200 جرام. وقد تم تقسيمهم عشوائيا إلى المجموعة الضابطة التي تضمنت 15 ركبة تم تقديمها بحيث إن كل مجموعة من المجموعات التجريبيه لجاهاز مخ مرتين وثلاثين ثقلاً، 5 مل من الماء المقطر وستة جرذان تلقوا 0.5 مل من زيت الزيتون البكر وستة جرذان اخرون تلقوا كلها 0.5 مل من الماء المقطر و5 مل من مزيج مادة الرطوبة المثبتة.


النتائج: قام الدراسه بقياس مستويات عامل نخز الخلايا ومستويات هرمون الجلوتاثيون المختزل. وتم الحصول على مسافات عامة محددة في كل المجموعات، وتم تطبيقهم لمسافات عامة محددة في كل المجموعات. وتم الحصول على مسافات عامة محددة للخلايا. وتم الحصول على مسافات عامة محددة للخلايا. وتم الحصول على مسافات عامة محددة للخلايا.

الخلاصة: نستنتج من ذلك ان اعطاء عقار الاستاذنسين وجلوتامات أحادي الصوديوم بشكل فردي أو في مركب يمكن أن يؤدي إلى تغييرات في الخصائص الفيزيولوجية للخلايا، ولكن في حالات معينة، قد يؤدي إلى تغيرات في الخصائص الفيزيولوجية للخلايا.

نستنتج من ذلك ان اعطاء عقار الاستاذنسين وجلوتامات أحادي الصوديوم بشكل فردي أو في مركب يحدث تغيرات في الخصائص الفيزيولوجية للخلايا.