Ameliorative Effect of Vitamin C against Monosodium Glutamate Induced Toxicity on Testes and Cauda Epididymal Sperm Reserves in Adult Male Albino Rats

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Abstract

Background: Monosodium glutamate (MSG) is a commonly used food enhancer which causes a wide range of endocrinial disorders including reduced fertility rate in both sexes. Earlier studies have shown that chronic administration of MSG induces oxidative stress in some organs of experimental animals. Accordingly, this work was carried out to study the possible ameliorative effect of vitamin C on monosodium glutamate induced testicular toxicity in adult male albino rats.

Study design: Sixty adult male albino rats were included in this study and divided into four main groups. Group I used as control group (10 rats). Group II (10 rats) received 100mg/kg/day vitamin C. Group III (20 rats) was subdivided into two subgroups (10 rats each): 100mg/kg/day and 4gm/kg/day MSG treated rats. Group IV (20 rats) was also subdivided into two subgroups and received both MSG and vitamin C in their previous doses. All animals were treated orally by gastric tube for two months. Body weight, serum testosterone level, cauda epididymal sperm reserves, histopathological examination using H&E stain and survivin immunostaining were studied.

Results: There was a significant increase in the body weight (p<0.05) and in the percentage of sperm abnormal forms (p<0.05), while a significant reduction in the sperm count and motility (p<0.05) was noticed. These results were dose dependent. Animals treated with 4gm/kg MSG showed severely damaged seminiferous tubules with few primary spermatocytes in H&E stained sections and marked reduction in survivin expression in histochemical study compared to other groups. Moreover, there was significant improvement in all studied parameters on administration of vitamin C concomitant with MSG which was more observed with low MSG dose (100mg/kg). Thus, vitamin C holds a promise as an agent that can reduce MSG induced toxic effects in testes.

Introduction

A variety of environmental chemicals, industrial pollutants and food additives have been implicated as causing harmful effects (Moore, 2003). Most food additives act either as preservatives, or enhancer of palatability. One such food additive is monosodium glutamate (MSG) which is one of the most flavor enhancers commonly used all over the world by millions of people in their food (Farombi and Onyema, 2006 and Eweka and Om’Iniabohs, 2011). Unfortunately, many Egyptians consume MSG in both ready and home-made food (Swelim, 2004).

Monosodium glutamate (MSG), a white crystalline powder, is the sodium salt of a naturally occurring non-essential amino acid, glutamic acid (Furst and Stehle, 2004). It triggers the taste buds sensitive to ‘UMAMI’ taste which is referred as the fifth primary taste (Kurihara and Kashiwayanagi, 2000). Umami can be translated “pleasant savory taste”. People taste Umami via receptors for glutamate, commonly found in its salt form. The Umami taste often acts to improve or enhance the acceptance response to a variety of food (Bellisle, 2008). MSG is present in: flavored chips and snacks, soups or sauces (canned, packed), prepared meals, frozen foods and...
meals, fresh sausages, marinated meats, stuffed or seasoned chicken, bottled soy or oriental sauces, manufactured meats, luncheon chicken and turkey, flavored tuna, vegetarian burgers and sausages (Bojanic et al., 2009). The Food and Drug Administration (FDA) of the United States reported that monosodium glutamate is harmless and that it should be kept on the “Generally Recognized as Safe” (GRAS)-list of foods (Eweka and Om’Iniabohs, 2011). As a consequence, it requires no specified daily intake, or an upper limit intake requirement (Moore, 2003). However, inadvertent abuse of this food additive may occur because of its abundance, mostly without labeling, in many food ingredients (Egbruonu et al., 2009).

Some studies indicated that MSG could exhibit harmful effects on human body including: obesity with affected locomotor activity (Egbruonu et al., 2010), altered lipid metabolism (Egbruonu and Osakwe, 2011), asthma (Scopp, 1991 and Ehlers et al., 1998), epilepsy, stroke and depression (Samuels,1999 and Mozes and Sefcikova, 2004).

More recent reports have confirmed the adverse effects of MSG on the hypothalamic-pituitary axis of the brain, including neuro-excitatory/neuroendocrine effects (Feldman and Weidenfeld, 2005 and Seo et al., 2010).

Several studies were done to observe the effects of MSG on the organs concerned with the reproduction as testis, ovary, uterus etc. in the neonatal animals (rats & mice), fertility rate has been reported to be reduced in both sexes (Mohamed, 2012).

More recent studies have examined the induction of oxidative stress in different tissues of experimental animals after administration of chronic doses of MSG (Singh et al., 2003; Diniz et al., 2004; Farombi and Onyema, 2006 and Onyema et al., 2006).

Vitamin C (ascorbic acid) which is a water-soluble micronutrient, well absorbed by the gastrointestinal tract has a considerable antioxidant activity: it scavenges reactive oxygen species and may, thereby, prevent oxidative damage to the important biological macromolecules, such as DNA, proteins, and lipids (Konopacka, 2004).

Thus, this study aimed to investigate the possible toxic effect of MSG in different doses, as a natural constituent of many food items, on the testes and cauda epididymal sperm reserves of adult male albino rats and to evaluate the possible efficiency of vitamin C, as a natural available cheap antioxidant, in ameliorating such toxicity.

Material and methods

Chemicals

Monosodium glutamate was obtained in the form of L-Glutamic acid Sodium Salt 500 gm (Eastrin Fine Chemicals. LTD).

Vitamin C was obtained in the form of L-ascorbic acid 100 gm (Universal Fine Chemicals PVT. LTD India A.R. grade).

Animals

The present study was carried out on 60 adult male albino rats, weighing (150-200 gm). They were housed in clean properly ventilated cages under the same environmental conditions with free access to food and water throughout the whole period of the experiment. The room temperature was about 22-24 °C and the animals were exposed to 12:12 hours light dark cycles.

Experimental design

This current work was carried out in Forensic Medicine & Clinical Toxicology and Pathology Departments in accordance to the guidelines of the Ethical Committee of Medical Research, Faculty of Medicine, Tanta University, Egypt.

After one week of acclimatization, the animals were divided into four main groups:

- **Group I: (control group):** included 10 animals that further subdivided into two subgroups (five rats each)
  - Subgroup (i): received no treatment.
  - Subgroup (ii): received 0.2 ml of distilled water (the diluting vehicle for MSG and vitamin C).

- **Group II: (Vitamin C treated animals):** included 10 animals that received vitamin C in a dose of 100 mg/kg/day (Obochi et al., 2009), prepared as 20mg/0.2ml solution in distilled water.

- **Group III: (MSG treated animals):** included 20 rats that further subdivided into two subgroups (ten rats each)
  - Subgroup (i): received MSG in a dose of 100 mg/kg/day (Obochi et al., 2009), prepared as 20mg/0.2ml solution in distilled water. The average human daily intake of MSG is estimated to be 0.3-1.0 gm in industrialized countries (Geha et al.,
2000). MSG in a dose of 100mg/kg/day used in this group was equal to the average intake in human. It was calculated by the use of the formula of dose translation based on body surface area (Reagan-Shaw et al., 2008).

- **Subgroup (ii):** received MSG in a dose of 4 gm/kg/day (Monno et al., 1995), prepared as 800mg/4ml solution in distilled water. This MSG dose has been widely used in studying MSG-induced toxic reactions in experimental animals (Diniz et al., 2004 and Farombi and Onyema, 2006). Moreover, this dose is ¼ LD50 in rats and mice as LD50 in rats and mice is 15,000-18,000 mg/kg body weight (Walker and Lupien, 2000).

- **Group IV:** (Protective group): included 20 rats that were further subdivided into two subgroups (ten rats each):
  - Each subgroup received vitamin C concomitantly with MSG (at the same dose and duration of their correspondents).
  - MSG, vitamin C and their vehicle were given orally by gastric tube once daily for two months.

### Methods

**1- Body weight**

Body weight was recorded at the beginning of the experiment of all studied groups and after two months.

**2- Hormonal assay**

Prior to sacrificing each rat, 2ml of blood was collected from the tail vein into a test tube and allowed to clot. The serum was obtained by centrifugation and kept frozen at -4°C for performing testosterone assay. This assay was carried out using ELISA technique.

The animals were anaesthetized with diethyl ether inhalation and afterward sacrificed by cervical dislocation.

**3- Tissue Processing**

- **For histological examination:** Orchidectomy was performed by open castration method. A midline or pre-scrotal incision was made and the testicles were milked out of the incision site. The testicles were exposed by incising the tunica vaginalis. The spermatic cord was exposed, ligated and incised. The testes were fixed by immersion in Bouin’s fluid for 48 hours and processed for paraffin sections of 5 µ thickness. Sections were stained with hematoxylin and eosin for light microscopy (Bancroft and Gamble, 2002). Caudae epididymes from each animal were also dissected for the study of sperm abnormalities.

- **For immunohistochemistry**, Four micrometer thick sections of formalin-fixed, paraffin-embedded tissue were immuno-stained for survivin (antiapoptotic). By using streptavidin-biotin peroxidase complex method, immunohistochemical staining was performed using the primary antibody for survivin (Cat. #RB-9245-R7, rabbit polyclonal antibody, Lab Vision/NeoMarkers, Fremont, CA, USA). The paraffin-embedded sections were dewaxed and rehydrated in ethanol. Endogenous peroxidase activity was blocked by incubation with H2O2/phosphate buffer solution, whereas the other steps of the procedure were the same. The cellular labeling of survivin was visualized in the nucleus and the cytoplasm as brown coloration (Lechler et al., 2007).

**4- Semen examination**

Semen examination of all groups was performed according to Bearden and Fuquay, (1997) as following:

The epididymis was dissected and its content was obtained by cutting the epididymis and squeezing its content into one centimeter saline in a clean Petri dish. The aliquot was subjected to the following:
A. **Progressive sperm motility:** A small droplet of semen was placed on a clean slide and covered with a cover slip and then several fields were examined under microscope and the number of progressively motile sperms per 100 sperms was estimated and recorded.

B. **Sperm cell concentration:** The sperm cell concentration was estimated by direct cell count using a hemocytometer slide after diluting semen to the desired concentration. The cover slip was placed over the counting chamber and a small amount of diluted semen is placed at the edge of the cover slip then the slide is placed under the microscope to count the number of the sperm in the four corners and the central square, then, the number of the sperm per epididymis was counted using the following formula:

\[
\text{No. of sperms/epididymis (million) = number of sperm x dilution rate x 50,000}
\]

C. **Epididymal sperm abnormalities:** A drop of epididymal content of each rat was mixed with equal drops of Eosine 1% and nigrosine 5% stain and carefully mixed, two dry clean slides were used to make the smear and the stained slide was allowed to dry and examined under high power of microscope (x40). The percentage of abnormal sperms was calculated as mean from counting 100 sperm in each of the two slides. Photomicrographs of the desired sections were made for further observations.

### 5. Morphometric study

The morphometric measurements were carried out using the Leica Qwin 500 (Leica, London, UK) image analyzer computer system (Central Laboratory, Faculty of Medicine, Tanta University). Ten high-power fields, from each slide of all animals of each group, were used to evaluate the immunohistochemistry results by counting survivin positively stained cells. Immunoreactivity was assigned one of the following four categories: 0 (<20% of cells show staining); +1, (20-50%); +2, (50-75%); and +3, (>75%). Immunoreactivity in more than 20% of cells was considered positive i.e. score equal to or above +1(Kennedy et al., 2003).

### Statistics

Data were expressed as means ± standard deviation. ANOVA test was used for comparison among different groups in quantitative data. Scheffe test was used to compare each group versus another. P<0.05 was considered significant.

### Results

A. **Initial and final body weights**

Table (1) and figure (1) show mean values of the initial and final body weights of adult male rats of all studied groups. There was no significant difference of the mean values of the initial body weight between all the studied groups. At the end of the experiment, the mean values of the final body weight in MSG- treated rats (group III) were increased significantly (P<0.05) compared with other studied groups (I, II, IV), while treatment with vitamin C along with MSG in group IV significantly reduced this increase.

The mean values of the final body weight in 4gm/kg MSG- treated rats (326.4 ± 13.81) were significantly higher than those of 100mg/kg MSG-treated rats (287.6 ± 13.86). There was also significant decrease in the final body weight in 100mg/kg MSG-vitamin C treated animals (252.5±10.61) compared to 4gm/kg MSG-vitamin C treated animals (265.2±14.10).

B. **Serum testosterone levels**

Table (2) and figure (2) show mean values of serum testosterone levels (ng/ml) in adult male rats of all studied groups. There was significant decrease (P<0.05) in mean values of serum testosterone levels in MSG- treated animals (group III) compared with other studied groups (I, II, IV), while treatment with vitamin C along with MSG in group IV significantly improved this reduction. The mean values of serum testosterone levels in 4gm/kg MSG- treated rats (0.15± 0.031) were significantly decreased compared to those in 100mg/kg MSG- treated rats (0.32±0.047).

There was significant improvement in 100mg/kg MSG-vitamin C treated animals (0.94±0.036) compared to 4gm/kg MSG-vitamin C treated animals (0.83±0.020).

C. **Assessment of caudaepididymal sperm reserves**

Table (3) and figure (3) Show mean values of cauda epididymal sperm reserves as regard sperm count (X10⁶), sperm motility (%) and incidence of abnormal forms (%).
The sperm count (X10⁶) showed high significant decrease (P<0.05) in rats on MSG alone (group III) compared with other studied groups (I, II, IV), while treatment with vitamin C along with MSG in group IV significantly improve this decrease. The mean values of sperm count of 4gm/kg MSG-treated rats (9.3±1.15) were significantly decreased compared to 100mg/kg MSG-treated rats (21.5±4.1). Moreover, sperm count was significantly improved in 100mg/kg MSG-vitamin C treated animals (34.5±2.91) compared with 4gm/kg MSG-vitamin C treated group (26.2±3.33).

The sperm motility (%) showed significant decrease (P<0.05) in rats treated with MSG alone (group III) compared with other studied groups (I, II, IV), while treatment with vitamin C along with MSG in group IV significantly improve this reduction. The mean values of sperm motility in 4gm/kg MSG-treated rats (29.9±2.33) were significantly decreased compared to 100mg/kg MSG-treated rats (60.4±3.59).

Moreover, there was significant decrease in sperm motility (%) in 100mg/kg MSG-vitamin C treated rats (81.6±2.71) compared to 4gm/kg MSG-vitamin C treated group (77.2±3.25).

The incidence of abnormal forms (%) showed high significant increase (P<0.05) in MSG treated animals (group III) compared with other studied groups (I, II, IV), while treatment with vitamin C along with MSG in group IV significantly reduced this increase. The mean values of abnormal forms of sperms in 4gm/kg MSG-treated group (38±2.16) were significantly increased compared to 100mg/kg MSG-treated group (24.9±3.84).

Moreover, sperm count was significantly reduced in 100mg/kg MSG-vitamin C treated rats (6.50±0.84) compared with 4gm/kg MSG-vitamin C treated group (11.2±1.36).

**D. Histopathological results**

Figure (4) showed some of these abnormal sperm forms compared to the normal morphology of sperm in the form of normal head shape with characterized hook and straight tail (A). They include sperms with bent necks (B), bent tails (C) or bent neck and tail (D).

H&E stained sections of the testes in adult male rats of the control and vitamin C groups (group I,III) showed normal morphology of the testis where there was normal testicular histology and normal spermatogenic progression with active seminiferous tubules and interstitial spaces (figures 5, 6).

MSG treated rats with 100mg/kg were less affected than those treated with 4gm/kg as illustrated in figures (7,8). There was damage of spermatogonia and elongation of primary spermatocytes with few damaged late spermatids in the tubules in 100mg/kg MSG-treated animals (figure 7), while in 4gm/kg MSG-treated group there was severely damaged seminiferous tubules with few primary spermatocytes in the tubules (figure 8).

As regards to the MSG-vitamin C treated rats, there was marked improvement in 100mg/kg MSG-vitamin C treated animals as illustrated in figure (9) in which there was some normal seminiferous tubules with normal spermatogenesis admixed with degenerated tubules. While vitamin C with 4gm/kg MSG-treated animals was less effective as demonstrated in figure (10) where there was few preserved normal seminiferous tubules with normal spermatogenesis admixed with severely damaged tubules.

**Immunohistochemical results**

The sections of the control and vitamin C treated animals (group I,II) showed that spermatogenic cells had a positive survivin reaction in the nuclei and the cytoplasm. Survivin immunostaining showed strong expression in most of the nuclei and cytoplasm of spermatogenic cells of the control and vitamin C treated animals (figure 11).

As regards to MSG-treated rats with 100mg/kg, there was mild nuclear and cytoplasmic expression in spermatogenic cells of most seminiferous tubules (figure 12), while 4gm/kg MSG-treated animals showed marked reduction in survivin expression in spermatogenic cells of most seminiferous tubules (figure 13).

Concerning the group treated with vitamin C along with MSG (group IV), there was strong effect of vitamin C on rats treated with 100mg/kg as illustrated in figure (14) in which there was strong nuclear and cytoplasmic survivin immunostaining of the spermatogenic cells, while moderate nuclear and cytoplasmic surviving immunostaining of spermatogenic cells was found in 4gm/kg MSG-vitamin C treated animals (figure 15).

Table (4) and figure (16) show significant decrease in survivin expression in rats on MSG alone (group III) compared with other studied groups (I, II, IV), while treatment with vitamin C along with MSG in group IV significantly improved this decrease. There was no significant difference between 100mg/kg MSG and 4gm/kg MSG-treated animals. Moreover, survivin expression was significantly improved in 100mg/kg MSG-vitamin C treated animals (2.6±0.63) compared with 4gm/kg MSG-vitamin C treated group (2±0.42).
Table (1): ANOVA one way statistical analysis of initial and final body weights (gm) among all studied groups

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<tr>
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<td>MSG 100mg/kg</td>
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**Sheffe test**

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**Sheffe test**

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<th>Control&amp; MSG 100mg/kg + vitamin C</th>
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<th>vitamin C &amp; MSG 4gm/kg</th>
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NO= number, B.M= body weight; *p<0.05 significant; **p<0.001 highly significant.
Table (2): ANOVA one way statistical analysis of serum testosterone levels (ng/ml) among all studied groups.

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<td>Positive</td>
<td>Vitamin C</td>
<td>MSG 100mg/kg</td>
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Sheffe test

Control& vitamin C
Control& MSG 100mg/kg
Control& MSG 4gm/kg
Control& MSG 100mg/kg + vitamin C
Control & MSG 4gm/kg + vitamin C
vitamin C& MSG 100mg/kg
vitamin C& MSG 4gm/kg + vitamin C
vitamin C& MSG 100mg/kg + vitamin C
vitamin C& MSG 4gm/kg + vitamin C

NO = number; *p<0.05 significant; **p<0.001 highly significant.

Table (3): ANOVA one way statistical analysis of cauda epididymal sperm reserves, sperm count (X10⁶), sperm motility (%) and incidence of abnormal forms (%) among all studied groups.

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Sheffe test

Control& vitamin C
Control& MSG 100mg/kg
Control& MSG 4gm/kg
Control& MSG 100mg/kg + vitamin C
Control & MSG 4gm/kg + vitamin C
vitamin C& MSG 100mg/kg
vitamin C& MSG 4gm/kg + vitamin C
vitamin C& MSG 100mg/kg + vitamin C
vitamin C& MSG 4gm/kg + vitamin C

0.322
0.001**
0.058
0.095
0.001**
0.257
0.067
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**Sheffe test**

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**Abnormal forms (%)**

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NO = number; *p < 0.05 significant; **p < 0.001 highly significant.
Table (4): ANOVA one way statistical analysis of survivin expression among all studied groups.

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Shefe test

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<th>Control&amp; MSG 4gm/kg</th>
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NO= number; *p <0.05 significant; **p <0.001 highly significant.

Figure (1): Mean values of the initial and final body weights (gm) of adult male rats of all studied groups (B.W= body weight).
Figure (2): Mean values of serum testosterone level (ng/ml) in all studied groups.

Figure (3): Mean values of cauda epididymal sperm reserves as regard sperm count (X10^6), sperm motility (%) and incidence of abnormal forms (%) among different studied groups.
Figure (4): Photomicrograph showing normal and abnormal forms of sperms. A: normal sperm. B: sperm with bent neck. C: sperm with bent tail. D: sperm with bent neck and tail (H&E x 400).

Figure (5): A photomicrograph of a cross section in a testis of an adult male albino rat of the control group (group I) showing normal testicular histology and normal spermatogenic progression. Note the active seminiferous tubules (T) and interstitial spaces (S). (H&E stain X200).
Figure (6): A photomicrograph of a section in the seminiferous tubule of an adult male albino rat of the vitamin C treated group (group II) showing unremarkable histopathological changes. The seminiferous epithelium shows active spermatogenesis. (H&E stain X400).

Figure (7): A photomicrograph of a section in the testis of an adult male albino rat treated orally by MSG in a dosage of 100mg/kg showing damage of spermatogonia and elongation of primary spermatocytes. Few damaged late spermatids are still present in the tubules. (H&E stain X400).
Figure (8): A photomicrograph of a section in the testis of an adult male albino rat treated orally by MSG in a dosage of 4gm/kg showing a severely damaged seminiferous tubule. Few primary spermatocytes (arrows) are still seen in the tubule (H&E stain X400).

Figure (9): A photomicrograph of a section in the testis of an adult male albino rat treated orally by 100mg/kg MSG and 100mg/kg vitamin C showing some normal seminiferous tubules with normal spermatogenesis admixed with few degenerated tubules (arrows) (H&E stain X200).
Figure (10): A photomicrograph of a cross-section of seminiferous tubules of a testis of an adult male albino rat treated orally by 4gm/kg MSG and 100mg/kg vitamin C showing few preserved normal seminiferous tubules with normal spermatogenesis (arrows) admixed with severely damaged tubules (H&E stain X200).

Figure (11): A photomicrograph of a section in the seminiferous tubule of an adult male albino rat of control and vitamin treated rats (group II) shows strong survivin immunostaining in most of the nuclei and cytoplasm of spermatogenic cells (IHCx400).
Figure (12): A photomicrograph of a section in the seminiferous tubule of an adult male albino rat treated orally by MSG in a dosage of 100mg/kg rat showing mild nuclear and cytoplasmic survivin immunostaining of the spermatogenic cells (IHCx400).

Figure (13): A photomicrograph of an adult male albino rat treated orally by MSG in a dosage of 4gm/kg (IHCx400) showing marked reduction in survivin expression in the nuclei and cytoplasm of spermatogenic cells of most seminiferous tubules.

Figure (14): A photomicrograph of a section of seminiferous tubule of an adult male albino rat treated orally by 100mg/kg MSG and 100mg/kg vitamin C showing strong nuclear and cytoplasmic surviving immunostaining in the spermatogenic cells (IHCx400).
Figure (15): A photomicrograph of a section in the seminiferous tubule of an adult male albino rat treated orally by 4gm/kg MSG and 100mg/kg vitamin C showing moderate nuclear and cytoplasmic surviving immunostaining in spermatogenic cells. (IHCx400).

![Image](image_url)

**Figure (16):** Mean values of survivin expression in testes sections among all studied groups.

<table>
<thead>
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<th>Group</th>
<th>Mean Value</th>
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<td>GII</td>
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<td>GIII</td>
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**Discussion**

Glutamate was reported to have many functions in the human body as a metabolic intermediate and protein constituent (Zai et al., 2009). Despite its taste stimulation and improved appetite enhancement, reports indicate that MSG is toxic to human and experimental animals (Belluardo et al., 1990). MSG leads to generation of free radicals, activation of proteases, phospholipases and endonucleases, transcriptional activation of apoptotic programs and genotoxicity in mouse and rat (Goldsmith, 2000 and Farombi and Onyema, 2006).

MSG has high excitotoxic potential, which is mediated at the level of second messenger by calcium anions (Gao et al., 1994 and Lin et al., 1995). Bojanic et al., (2009) hypothesized that high concentration of MSG could alter permeability of neural membrane for calcium which could be involved in the mechanisms of MSG toxicity. Calcium ion influx into cells accompanies cell injury caused by a variety of stimuli, including oxidative stress and calcium also appears to be involved in the production of reactive oxygen species (ROS). Hydrogen peroxide (which can produce ROS) induces raised intracellular calcium levels, the response being suppressed by antioxidants (Feng and Hart, 1996). Being rich in membrane bound polyunsaturated fatty acids (PUFA), mammalian spermatozoa are more susceptible to ROS attack (Lenzi et al., 2000).

In the present study, toxicity of MSG in low dose (100mg/kg) and high dose (4gm/kg) was
evaluated on adult male albino rat testis and epididymis and possible ameliorative effect of vitamin C. The rat model is probably the best choice in the experimental study because purchase and harvesting are inexpensive. In addition, spermatogenesis observed in the rat is similar to that in humans (Turner, 2001).

There was a significant increase in the body weight of rats treated with MSG compared with control or vitamin C treated animals. Moreover, there was more significant increase in the body weight in 4gm/kg than in 100mg/kg MSG- treated animals.

These results were in agreement with Bloom, (2001) who suggested that MSG affected hypothalamus which is responsible for appetite, body temperature, water balance and sleep. These results are also in line with the study carried out by Nakagawa et al. (2000) and Ortiz et al. (2006).

Other studies have also shown that (MSG) causes reduction in the secretion of growth hormones, leading to stunted growth and irreversible obesity, essentially due to the accumulation of excess fats in adipose tissue (Ikonomidou et al., 1995; Eskes, 1998 and Rodriguez et al., 1998). Through its stimulation of the orosensory receptors and by improving the palatability of meals, MSG influences the appetite positively, and induces weight gain (Rogers and Blundell, 1990). Macho et al. (2000) had cited that obesity induced by MSG was due to chemical lesions in arcuate nucleus structure that resulted in endocrine disorders such as obesity and glucose intolerance.

The current work demonstrated a significant decrease in the level of serum testosterone hormone in MSG treated groups compared to the control group and vitamin C treated group. This decrease in serum testosterone level was dose dependent. The testosterone level in 4gm/kg MSG-treated animals was significantly reduced than in 100mg/kg. This was in agreement with Igwebuike et al., (2011) who reported marked reduction of serum testosterone level in both young and adult rats treated with MSG.

Neuronal losses in the hypothalamus can result in disruption of the hypothalamic-pituitary-testis regulatory axis that controls the steroidogenesis of testicular Leydig cells (McLachlan et al., 1996). This will ultimately impact serum testosterone levels, because 95% of the androgen is produced by Leydig cells of the testes, while cells in the cortex of the adrenal gland produce about 5% (Urban, 1999). Such alterations in reproductive hormone secretion may adversely affect the reproductive capacity of the affected animals.

Because testosterone plays a crucial role in male reproductive system, disruption of its production may impair male reproductive health (McLachlan et al., 2002 and O’Shaughnessy et al., 2010). Testosterone is needed for the continued production of different generation of germ cells in the seminiferous tubules. Therefore, reduction of testosterone level may lead to the separation of germ cells from the epithelium of the seminiferous tubules, decreased daily sperm production and morphological abnormality of testis (Blanco-Rodriguez and Martinez-Garcia, 1998 and Zitzmann, 2008). This was evidenced in the present study by histological examination of testis treated with MSG where there was damage of spermatogonia and elongation of primary spermatocytes with few damaged late spermatids in the tubules in 100mg/kg MSG-treated animals. While in 4gm/kg MSG- treated group there was severely damaged seminiferous tubules with few primary spermatocytes in the tubules.

Similarly, Eweka and Om’iniaibohs, (2011) reported that male rats treated with 6gm of MSG showed cystic degenerative changes, with some intercellular vacuolations appearing in the stroma of the seminiferous tubules when compared to the control sections. Also, Mohamed, (2012) observed structural changes of testes of MSG-treated rats which were dosage-duration-dependent. These changes ranged from slight to moderate damage in case of the short-term treatment however, severe damage of spermatogenic and Sertoli cells was recorded in case of the long-term treatment.

This was in contrast with Igwebuike et al., (2011) who reported the absence of any overt pathological lesions in the histology of the testes of the young and adult rats that received varied doses of monosodium glutamate suggests that MSG did not have any direct toxic effect on the testes.

The present study revealed dose-dependent reduction in cauda epididymal sperm counts in the MSG-treated animals compared to control and vitamin C treated groups. Reduction in sperm count was more significant in 4gm/kg treated animals. This was in agreement with Igwebuike et al., (2011). The dose-dependent reduction in cauda epididymal sperm counts following oral administration of monosodium glutamate to both young and adult male rats may be an indication that the normal process of spermatogenesis was adversely affected in the rats that consumed monosodium glutamate. Spermatogenesis is a complex cyclic process in which germ cells undergo series of mitotic and meiotic cell divisions, followed by morphological differentiation in a delicately regulated spatiotemporal fashion in the seminiferous epithelium. In mammals, spermatogenesis is totally dependent upon testosterone (Pakarainen et al., 2005 and Wang et al., 2009); however, only total abolition of testicular androgen action resulted in consistent azoospermia (Zhang et al., 2003).

As regards, androgens act to stimulate spermatogenesis through androgen receptors (ARs) located on Sertoli cells and peritubular myoid cells of the testis (De Gendt et al., 2004). It has been demonstrated that androgen stimulation of spermatogenesis requires direct androgen action on the Sertoli cells (O’Shaughnessy et al., 2010), and this was necessary for postmeiotic spermiogenesis (Zhang et al., 2003). Thus, appropriate regulation of androgen activity via the hypothalamic-pituitary-testis axis is necessary for initiation and maintenance of spermatogenesis. The indication is that the reduced cauda epididymal sperm counts observed in the MSG-treated rats in the present study may be the end result of a considerable decline in the influence of testosterone on spermatogenesis in these rats.
Furthermore, there was a significant increase in the abnormal sperm morphology and a significant decrease in sperm motility in MSG treated groups compared to control and vitamin C treated groups. This effect was more noticed in 4gm/kg MSG- treated animals than in 100mg/kg MSG- treated animals. The toxic effect of MSG on the testes was reported by other studies where a significant oligozoospermia and increase abnormal sperm morphology in a dose-dependent fashion in male Wistar rats were observed (Onakworo et al., 1998). Also, it has been established that MSG may be implicated in cases of male infertility as it causes testicular hemorrhage, degeneration and alteration of sperm cell population and morphology (Oforoflio et al., 1997).

The primary mechanism for glutamate induced cell death is apoptosis in different cell types (Martin et al., 2000 and Schelman et al., 2004). Apoptosis of selected germ cells occurs normally in the testis and is essential for the normal maintenance of spermatogenesis (Hauoui et al., 2008). Among the mammalian apoptosis regulators, the inhibitor of the apoptosis (IAP) protein family is just beginning to be elucidated for its role in spermatogenic function.

Survivin is an IAP protein regulating apoptosis at cell division. Survivin has been studied extensively in cancer cells, but has only recently been studied extensively in male germ cells. In the rodent testis, survivin expression has been localized to germ cells, especially mature spermatocytes. Testicular survivin was found to be one of the highly expressed transcripts in adult human testis, specially in the nuclei of mature spermatocytes, and its down-regulation is associated with spermatogenic failure (Weikert et al., 2004; Weikert et al., 2006; Absalan et al., 2010 and Al-Magrebi et al., 2011). The fact that survivin regulates cell proliferation, cell cycle, and programmed cell death makes it a candidate gene that may be involved in the regulation of germ cell production (Fukuda et al., 2002).

Survivin expression in the cytoplasm could be associated with its function in controlling cell survival (inhibitor of apoptosis), whereas nuclear staining may rather promote cell proliferation (Taubert et al., 2010). As an inhibitor of apoptosis, survivin is believed to block apoptosis by inhibiting caspases and antagonizing mitochondria-dependent apoptosis (Marusawa et al., 2003 and Sah et al., 2006). In contrast, survivin activity is blocked through its interaction with the mitochondrial protein; S-mac, which is released into the cytoplasm as a response to apoptotic stimuli, rendering survivin unable to inhibit caspases (Ceballos-Cancino et al., 2007). This finding suggested that suppression of survivin expression could promote the apoptotic stimuli. The potential mechanism of survivin down-regulation has been suggested by several studies to be secondary to ROS generation and ROS-induced apoptosis (Seo et al., 2007). Also, McCrann et al. (2009) have shown that over-expression of Nox4, a member of the NADPH oxidase family and a major source for ROS production has resulted in down-regulation of survivin mRNA. These findings led to the hypothesis that one of the manifestations of deregulated ROS signaling could be the down-regulation of survival genes.

In the present study, there was a significant down-regulation of survivin in MSG-treated animals compared to control and vitamin C treated groups. Reduction in survivin expression in MSG-treated rats was dose dependent. It was more marked in animals treated with 4gm/kg that indicated more liability for cell death with higher dose.

Vitamin C (ascorbic acid) has a well-known antioxidant protective role as it is considered the front line of defense against free radicals through; reactive oxygen species scavenging, reduction of peroxides and repair of peroxidized biological membranes and sequestration of iron (Loo et al., 2003).

In the male reproductive system, vitamin C is known to protect spermatogenesis. It increases gamete mobility since it reduces oxidative stress level (Hsu et al., 1998). Also, it plays a major role in semen integrity and fertility both in men and animals (Agarwal et al., 2005 and Eskenazi et al., 2005) increases testosterone levels (Sőnmez et al., 2005) and prevents sperm agglutination. Moreover, it is an important chain-breaking antioxidant, contributing up to 65% of the total antioxidant capacity of seminal plasma found intracellular and extracellular (Makker et al., 2009).

In the present study, protective effect of vitamin C was observed in MSG- vitamin C treated groups. There was significant reduction in the body weight in MSG-vitamin C treated groups compared to MSG –treated groups which was more significant in 100mg/kg MSG-vitamin C than in 4gm/kg MSG-vitamin C treated animals.

Also, there was a significant improvement in serum testosterone levels, sperm count and sperm motility with decreased abnormal forms in MSG-vitamin C treated animals compared to MSG-treated animals. These effects were more significant in 100mg/kg MSG treated animals than in 4gm/kg MSG treated ones.

Histological examination has confirmed this protective effect of vitamin C on MSG –treated animals. This protection was more marked in animals treated with 100mg/kg MSG where some normal seminiferous tubules with normal spermatogenesis admixed with few degenerated tubules were observed, while in 4gm/kg MSG –vitamin C treated animals showed few preserved normal seminiferous tubules with normal spermatogenesis admixed with severely damaged tubules. As regards survivin expression, there was strong nuclear and cytoplasmic expression in the spermatogenic cells in animals given vitamin C with MSG in a dosage of 100mg/kg while moderate nuclear and cytoplasmic immuno-staining of spermatogenic cell was observed in animals given vitamin C with MSG in a dosage of 4gm/kg.

These findings were in accordance with Farombi and Onyema, (2006) who suggested that antioxidants such as vitamin C have protective potential against oxidative stress induced by MSG. Furthermore, Shrilatha and Muralidhara, (2007) have
reported the protective effect of vitamin C on testicular oxidative stress, sperm oxidative stress and genotoxic effects using a diabetic mice model. Similarly, Nazıroğlu (2003) concluded that vitamin C acted as an antioxidant in reproductive milieu. Also, Duarte and Lunec, (2005) and Li and Schellhorn, (2007) have reported that vitamin C attenuated the reduction of seminiferous tubular diameters probably because this vitamin may have a role in the synthesis of extracellular matrix components.

Conclusions

The present work has demonstrated that oral administration of monosodium glutamate resulted in lowered serum testosterone levels and reduction in the cauda epididymal sperm reserves of adult male rats and cause overt pathological lesions in their testes. These effects were dose dependent and oral therapeutic administration of vitamin C along with MSG provided a significant improvement in the biochemical and pathological changes in testes of rats administrated MSG only at low dose (100mg/kg) in comparison with high dose (4gm/kg) MSG treated group. Thus, vitamin C holds a promise as an agent that can potentially reduce MSG induced toxic effects in testes.

Recommendations

Extra-antioxidant supplementation through diets can be helpful as a defense mechanism and might be helpful in prevention of oxidative stress and allied ailments. Further studies on animals and/or human are recommended to study the protective and preventive effects of different anti-oxidants, also to know if this toxic effect is reversible or not.

References


Furst P and Stehle P (2004): What are the essential elements needed for the determination of amino acid requirements in humans? J Nutri 34: 1558S-1565S.


Taubert H, Heidenreich C, Holzhausen H-J, et al., (2010): Expression of survivin detected by immunohistochemistry in the cytoplasm and in the nucleus is associated with prognosis of
leiomyosarcoma and synovial sarcoma patients. BMC Cancer 10:65

The effect of levonorgestrel in infertile patients with sperm defects on the production of normal spermatocytes and spermatozoa. Workshop on the outcomes of this study (December 2008) and its clinical implications. The effects of long-acting progesterone on sperm production and quality. Workshop on the outcomes of this study (December 2008) and its clinical implications.

We investigated the effect of levonorgestrel in infertile patients with sperm defects on the production of normal spermatocytes and spermatozoa. The results of this study highlight the potential of levonorgestrel as a treatment option for infertile patients with sperm defects.