**Morphometrical, Histopathological, and Cytogenetical ameliorating Effects of Green tea Extract on Nicotine Toxicity of the Testis of Rats**

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**Abstract:** Nicotine is a major toxic component of cigarette smoke and it is a major risk factor in the development of functional disorder of several organ systems. The natural diet contains a variety of compounds that exhibit protective effects towards different toxicities of nicotine as green tea. Four groups of male Swiss albino mice were divided: untreated control group; Nicotine–treated group (2.5 mg/kg/day); Green tea-treated group (40 mg/kg/day); and Nicotine and green tea treated group interperitoneal administration for successive 28 days. Results showed that disorganization of the seminiferous tubules associated with reduction of spermatogenic cells, leading to widening of lumen of tubules upon nicotine toxicity. Many of seminiferous tubules exhibited degenerative phases of spermatocytes and spermatides as well as missing of sperms and hypo-spermatogenesis. The recorded data in nicotine intoxicated group showed significant and gradual decrease of number of leydig cells throughout all intervals of experiment. In the last, cytogenetically examination demonstrated significant increased in the number of nucleated polychromatic erythrocytes (MnPCE) and decreased in number of polychromatic erythrocytes (PCE) in bone marrow of nicotine-treated animals using micronucleus assay. Green tea treatment reduced number of nucleated polychromatic erythrocytes (MnPCE) and restored number of polychromatic erythrocytes (PCE) to nearly normal. In conclusions, intake of green tea might suppress the toxicity and mutagenic activity of nicotine.

**Keywords:** Smoking - Nicotine – Fertility – Antioxidants – Green tea

1. **Introduction:**

Smoking has enormous negative health consequences worldwide. Nicotine is a naturally occurring alkaloid found in tobacco plant (*Nicotiana tabacum*) and it is the major constituent of tobacco responsible for the compulsive use of tobacco (Wu et al., 2002). Nicotine and its metabolites are also being investigated and researched for the treatment of a number of disorders as Alzheimer's diseases (Hecht, 2003). Okamoto *et al.*, (1994) recorded the LD50 of nicotine as (50 mg/kg) for rats and (3 mg/kg) for mice.

Nicotine administration induces changes in gonadal functions and deficiency in sperm maturation and spermatogenesis and has a detrimental effect on the sperm-fertilizing potential of male rats (Reddy *et al.*, 1998 and Yamamoto *et al.*, 1998). Also nicotine was reported to have toxic effects on gonadal functions in males in addition to its role in the lowering testosterone and estradiol levels in the serum (Kavitharaj *et al.*, 1999). Nicotine has been associated with decrease in number of germ cells, germinal cells and increased chromosomal abnormalities in sperm and with increased the risks of birth defects and neonatal death and genetic mutations in sperm, reduced sperm fertilizing capacity and decreased embryonic implantation rates (Polyzos *et al.*, 2009). Nicotine has a deleterious effect of nicotine on sperm membrane intactness and DNA and apoptosis in mouse leydig cells treated with nicotine were recorded by (Arabi, 2004 and Kim *et al.*, 2005).

From human studies; nicotine smoke contains harmful mutagens and carcinogens metabolites; that may induce defective semen quality, nuclear DNA damage of spermatozoa (Arif *et al.*, 2000, De Flora *et al.*, 2003 and Elshal *et al.*, 2009). Nicotine had been shown to increase the frequency of micronuclei in human gingival fibroblasts and induce DNA strand breaks in human spermatozoa (Arabi *et al.*, 2004). Information on the *in vivo* genotoxicity of nicotine have shown that nicotine induces aneuploidy and polyploidy (Bishun *et al.*, 1972), sister chromatid exchange and chromosome aberrations in bone-marrow cells of mice (Sen *et al.*, 1991). High doses of nicotine
increase the frequencies of premature centromere separation and premature anaphase and reduce the number of oocytes ovulated; however, the results of this study suggested that nicotine does not elevate aneuploidy levels in mouse oocytes (Mailhes et al., 2000). The increased generation of ROS can produce a condition of oxidative stress that can result in the oxidation of lipids, induction of DNA single-strand breakage, inactivation of certain proteins, and disruption of biological membranes (Durak et al., 2002). Increased oxidative stress has been suggested to play a major role in the pathogenesis of several smoking-related diseases such as cancer, cardiovascular and oral diseases (Reibel, 2003 and Sudheer et al., 2007).

Natural antioxidants as polyphenols of green tea extracts have received much attention for treatments of oxidative-stress-related pathological conditions (Park et al., 1998 and Yokozawa et al., 2004). Green tea can suppress the DNA adduction, and hence act as inhibitors of cancer and it is a rich source of polyphenols, which are antioxidants in natural and their ameliorating effect on genital organs were recorded (Ogura et al., 2008). Study on acute effect of green tea extract and its polyphenols constituents, in vitro showed stimulated testosterone production by rat Leydig cells and reduced the testicular tissue content of total cholesterol (TC), triglycerides (TG), phospholipids (Fabiano et al., 2009).

Tea polyphenol could inhibit the mutagenicity of chemical mutagens and chromosome aberrations (Shim et al., 1995). The frequencies of sister-chromatid exchange (SCE) in peripheral lymphocytes were study were significantly higher among smokers who were non tea drinkers, than those of non-smokers and smokers who consumed green tea at least two to three cups per day during the past 6 months (Lee et al., 1997). It is possible that consuming high levels of green tea over a long period may reduce the DNA damage caused by tobacco smoking (Liang et al., 2007). Our aim of work is the evaluation of the role of green tea in the protection from the effects of cigarette smoking.

2. Materials and methods

1-Experimental animals:

The experimental animals used in this study were male Swiss albino mice. Eighty Male Swiss albino mice aged 9 – 12 weeks and weighing 25 -30 gm were used throughout the study. Animals were fed a commercially prepared diet and had free access to tap water. All mice were kept under the same experimental condition, feeding standard diet, and water was available Adlibitum.

After one-week acclimatization period, the selected animals of nearly a similar weight were divided into 4 experimental groups so as to keep more or less the same mean body weight within the individual groups. The selected animal groups treated as follows:

1-Nicotine – Treated group: Each mouse in this group was given intraperitoneally (i.p.) dose of 3 nicotine 2.5mg/ kg / day for successive 28 days.

2-Green Tea – Treated group: For successive 28 days each mouse in this group was injected i.p. with freshly prepared green tea extract (40 mg / kg body weight / day).

3-Nicotine-Green Tea -Treated group: Each animal in this group was given i.p. of nicotine (2.5mg/kg/day) at the same dose as group1 concomitantly with green tea extract (40 mg / kg body weight) at the dose as second group for successive 28 days.

4-Control -Treated group: Each animal in this group was injected i.p. with distilled water (1ml/ day) for successive 28 days and handling on the same conditions exactly similar to that of the previously mentioned groups.

II-Chemicals:

The treated elements in the experiment were nicotine ((S)-3-(1-Methyl-2-pyrroli- dinyl) pyridine) and green tea extract. Nicotine was supplied as colorless liquid, obtained from faculty of pharmacy, Cairo University, Egypt. The mean LD 50 for intraperitoneal administration of nicotine to 8-week-old mice was reported as 12.5 mg (Favaro et al., 2003). Green tea extract was supplied in form of tablets obtained from Technomad Groups Company, Egypt, they were soluble in water.

III- Experimental design:

Five mice from each group were scarified by cervical dislocation at end of 1, 2, 3, and 4 weeks (7th, 14th, 21th, 28th days) of the experimental period and decapitation, two femurs are removed and stripped clean of muscle for cytogenetically examination. Also, lung, liver and testes were also sampled and kept in aqueous Bouin for histological, histochemical and morphometricall examinations.

Experimental analysis:

1--Histopathological examination:

For histopathology and morphometric evaluation, testes of all groups were collected deception and fast dissection. They were fixed in aqueous Bouin solution, dehydrated through alcohols, cleared in xylene and then embedded in paraffin wax. According to the method described by

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Bancroft and Stevens, (2002). 5µ thickness paraffin sections were prepared and mount on clean slides. For histopathological study sections were stained with Ehrlich's haemotoxylin and counterstained with eosin. A number of photomicrographs were taken at known magnification.

2- Image analysis:

The data were obtained using Leica Qwin 500 image analyzer computer system (England). The image analyzer consisted of a colored video camera, colored monitor, hard disc of IBM personal computer connected to the microscope, and controlled by Leica Qwin 500 software. The image analyzer was first calibrated automatically to convert the measurement units (pixels) produced by the image analyzer program into actual micrometer units. These measurements were done using an objective lens of magnification 40, i.e. of total magnification 400. Ten readings were obtained in each specimen. Regarding the no of Leydig cells using an objective lens of magnification 40, i.e. of total magnification 400, they were counted using interactive measurement in 10 fields in each specimen. The data obtained were subjected to statistical analysis using Duncan's Test and ONE WAY ANOVA.

3- Micronucleus assay

The most attractive features of the micronucleus assay are rapidly and easily with which in vivo genetic activity can be demonstrated. The first test to be developed and the most commonly used is carried out in mouse bone marrow by scoring micronuclei polychromatic erythrocytes (Heddle, 1973). In this study the micronucleus polychromatic erythrocyte were prepared according to an improved method adopted by (Salamone et al., 1980).

a. Harvesting Marrow.

Animals are killed by cervical dislocation, and two femurs are removed and stripped clean of muscle. The marrow is removed by making a small opening at the iliac end of the femur and introducing the pointed shaft of a 2.5-cm safety pin into the femur at the epiphyseal end. As the pin is slowly pushed and twisted into the marrow canal, the marrow exudes out the hole at the iliac end. The marrow is placed directly on a slide, and then a drop of fetal calf serum is added. With the aid of the edge of a clean slide, the marrow is mixed with the serum until homogeneous and then is spread as a smear; additional slides from a given animal can be prepared by simply transferring some of the mixed preparation onto other slides. Prepared slides are air-dried, fixed for 5 min in absolute methanol.

b- Slide Staining:

After air-drying, the smear were stained for 20 min in May-Grunwald stain in conjunction with the Gimsa stain (May-Grunwald/Gimsa (Schmid, 1975)) , this combination consisted of: 3 ml Gimsa (stock), 12 ml May-Grunwald (0.25gm MayGrunwald/100ml methanol), 2 ml 0.1M Na2Hpo4, -4.5 ml 0.1M KH2po4, and 51 ml D. Stain was freshly prepared and mixing well before used After the staining, the slides were washed thoroughly in distal water and left to dry overnight. All glass slides were coded before observation. Examination under oil lence.

c- Scoring

1000 polychromatic erythrocytes (PCEs) were scored per animal and the numbers of micronucleated PCEs were recorded. The results were expressed as the average number of micronucleated PCEs / 100 PCEs. For each sampling time, bone marrow smears from five animals per four / group were used for evaluation. From each animal, 1000 polychromatic erythrocytes (PCE) were scored under the microscope (1000-1250x; Optech, Germany) for the incidence of micronucleated polychromatic erythrocytes (MnPCEs). In addition, the number of PCEs among 1000 total erythrocytes (PCE + NCE) per animal was recorded to evaluate bone marrow cytotoxicity. The ratio of polychromatic to normochromatic erythrocytes (PCE / NCE) was calculated, based on 1000 erythrocytes (PCE / NCE) scored per slide. These ratios were used as a measure of toxicity of test materials.

IV- Statistical analysis:

Statistical analyses were carried out using analysis of variance, Duncan's Test and One Way ANOVA (Senedecor and Cochran, 1980). To evaluate the effect in between groups and give a change for multiple comparison in between.

3. Results:

1- Histopathological results:

The control untreated male mice testes tissues showed an outer capsule of fibroblastic connective tissue bound normal seminiferous tubules microscopy (Fig 1).

Testis of Swiss albino mice due to interperontial and daily injected nicotine treatment at the dose level 2.5mg/kg showed more or less normal testis architecture with complete spermatogenesis and focal decreased in population of leydig cells at the first week of nicotine treatment (Fig2). Along three weeks of nicotine treatment, some seminiferous tubules shows disorganization and atrophy of
Seminiferous tubules with widening of its lumen and reduction in sperm count. Other seminiferous tubules had arrest in spermatogenesis. Some spermatogenic cells showed apoptosis especially in 1ry and 2ry spermatocytes. Also, few vacuoles were appeared in some seminiferous tubules. Leydig cells were more reduction compared with previous stages and control stage (Fig. 3). Compared with normal group, the animal treated with nicotine for four weeks revealed most of seminiferous tubules had reduction in more than one stage of spermatogenesis such as reduced in number of spermatocytes and spermatid (marked arrest in spermatogenesis) and marked degeneration of the Leydig cells and decrease in this cell population (Fig. 4). Histological examination of testis sections of mice treated with green tea at different interval of experimental showed normal architecture (Fig. 7), complete spermatogenesis and maturation of germinal epithelium. Also, animals treated with green tea exhibited normal appearance, regarding size and arrangement of the tubules (Fig8).

2-Morphometric results:

Number of Leydig cells in testis:

Table (1) and Figure (9) represent the variation in the number of Leydig cells of testis due to nicotine intoxication and/or green tea protection. In normal group, all Swiss albino mice had normal distribution in nucleus area of liver and no significant changed presented along all interval of experimental (23.40 ± 1.784; 23.70 ± 1.726; 23.90 ± 1.940; 24.60 ± 2.242). The recorded data in nicotine intoxicated group showed significant (P<0.05) and gradual decrease of number of Leydig cells throughout all intervals of treatment experimental (18.10±2.282), (15.30±1.783), (12.40±1.979) and (9.50±1.462). As compared to control level, nicotine intoxicated group obtained insignificant (P>0.05) decrease of number of Leydig cells after first week of treatment (18.10±2.282 vs., 23.40 ± 1.784) but the value recorded being highly significant decreased till reached to minimal value at last week of experimental (15.30±1.783 vs., 23.70 ± 1.726; (12.40±1.979 vs., 23.90 ± 1.940); (9.50±1.462 vs., 24.60± 2.242). Treatment with green tea prevented the decrease in number of Leydig cells in nicotine treated mice with different degree and showed general increase in these cells. As compared to nicotine intoxicated group, the recorded data showed insignificant increased in number of Leydig cells after first and second weeks of treatment. With continues injection of green tea, till four weeks of experimental showed showed significant increased in number of Leydig cells and highly significant (P< 0.05) increased at three and last week of experimental (19.50±2.591vs.12.40±1.979 and 21.60±2.390 vs., 9.50±1.462) were observed compared to control values.
Fig (3): Micrograph of testis section treated with nicotine for three weeks showing apyknosis among spermatogenic cells especially necrosis in 1ry & 2nd spermatocytes (thin arrow). (H&E., 400X)

Fig (4): Micrograph of testis section treated with nicotine for four weeks showing different degree of damage of seminiferous tubules such as a disorder of systemic arrangement of the stages of spermatogenesis, loss of one or more stages of spermatogenesis (S.T.) and few number of Leydig cells are appeared (L.C.). (H&E., 400X)

Fig (5): Micrograph of testis section treated with green tea concomitantly with nicotine for a week showing almost normal appearance of seminiferous tubules (S.T.) with normal spermatogenic cells and arrangement Leydig cells are more increased, (H&E., 200X)

Fig (6): Micrograph of testis section treated with green tea concomitantly with nicotine for three weeks showing most of somniferous tubules

Fig (7): Micrograph of testis section treated with green tea concomitantly with nicotine for four weeks showing a little change in architecture of somniferous tubules with normal and order spermatogenesis (S.T.). (H&E., 400X)

Fig (8): Micrograph of testis section treated with green tea showing a normal testis structure, complete spermatogenesis (S.T.) and normal appearance of Leydig cells (arrow). (H &E , 400 X).
Improvement and protective effect of green tea increased significant and gradually with increased time of administration on nicotine intoxicated group. Individual green tea injection to albino mice showed no significant difference in the number of Leydig cells throughout all the tested experimental period or over all intervals of experimental were attained compared to control level. In conclusion, green tea had obviously improved on nicotine toxicity and Leydig cells back to normal number in male albino mice. As well as, it is showed an amelioration and obviously improvement to the nicotine toxicity effect on area of nucleus in liver, elastic fiber and leydig cells of testis of male albino mice.

3-Cytogenetical Results:
The cytogenetic damage induced by intraperitoneal injection of nicotine at ( 1/5 LD50 = 0.6 mg /kg to mice ) as well as the antimutagenic effects of green tea at ( 40 mg /kg ) treated intraperitoneal for one, two, three and four weeks with or without nicotine were investigated in bone marrow of male mice utilizing micronucleus assay. Micronucleus test were performed with nicotine and green tea. The % MnPCE in each treatment group as well as the PCE / NCE ratio are shown in tables 1 and 2. Normal polychromatic erythrocytes (PCE), normal normochromatic erythrocytes (NCE) and micronucleated polychromatic erythrocytes (MnPCE) showing in photos (1) a, b, and c.

In the present study, The ameliorative effect of green tea is gradual, increased with increased the time of adminstration of green tea with nicotine when study polychromatic etherocyttes (PCE), PCE/ NCE ratio and Mn-PCEs as shown in Tables 1&2. Treatment of green tea with nicotine was effective in reduction the frequencies of MnPCE in response to the time treatment. When green tea treated with nicotine for one, two or three weeks, the reduction in MnPCE come down to the same range of control.
group and the percent of the reduction in group 1, 2, 3 and 4 was 62.5%, 57.1 %, 77.35 % and 81.8 % respectively. Whereas treatment of green tea with nicotine, significantly (p<0.05) improved the number of PCEs and the ratio of PCEs to NCEs were significant (p<0.05) enhanced in compared to the control group.

Table (1): Effect of nicotine and / or green tea on the number of Leydig cells of testis section of Swiss albino mice.

<table>
<thead>
<tr>
<th>Groups of experiment</th>
<th>Time Of Treatment</th>
<th>Control groups (Mean ± S.E.)</th>
<th>Nicotine groups (Mean ± S.E.)</th>
<th>Green Tea groups (Mean ± S.E.)</th>
<th>Nicotine -Green Tea groups (Mean ± S.E.)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Week 1</td>
<td></td>
<td>23.40±1.784^BAA</td>
<td>18.10±2.282^Ba</td>
<td>24.80±1.836^Aa</td>
<td>21.30±1.880^BAA</td>
</tr>
<tr>
<td>Week 2</td>
<td></td>
<td>23.70±.726^BAA</td>
<td>15.30±1.785^Ca</td>
<td>25.40±1.507^Aa</td>
<td>19.70±2.290^Bca</td>
</tr>
<tr>
<td>Week 3</td>
<td></td>
<td>23.90 ± 1.940^Aa</td>
<td>12.40±1.979^Bba</td>
<td>25.50±1.607^Aa</td>
<td>19.50±2.591^Aa</td>
</tr>
<tr>
<td>Week 4</td>
<td></td>
<td>24.60± 2.242^Aa</td>
<td>9.50±1.462^Bb</td>
<td>25.80±1.576^Aa</td>
<td>21.60±2.390^Aa</td>
</tr>
</tbody>
</table>

(P< 0.05) is significant; (P< 0.01) is highly significant
Numbers of experimental animals were 5 in all groups
Compared between control treated group, nicotine treated group, nicotine-green tea treated group and green tea treated group tacked capitals latter but compared within group of each group tacked small latter.

Fig (10): Effect of green tea administration on number of leydig cells in testis of nicotine intoxicated male albino mice.

Figur (11): Histogram showing: (a) frequency of MnPCEs in nicotine and nicotine-green tea groups (b) Percentage of the reduction in MnPCEs in different treatments
Par each column (small letters) or each raw (capital letters) are significant at 5 % level. = Micronucleated polychromatic.

Statistical analysis of results were done according to Dun can's multiple rang test. Means with different letters within each column (small letters) or each raw (capital letters) are significant at 5 % level. = Micronucleated polychromatic.

electrical dysfunction, erectile dysfunction, and male factor

testis (Aydos, 2001). Destruction of Leydig cells may cause testicular atrophy; gonadal dysfunction, erectile dysfunction, and male factor

Table (2): Represent the effect of Nicotine and Green Tea Treatment on Frequencies of MnPCE.

<table>
<thead>
<tr>
<th>Groups of Experimental</th>
<th>Control Groups (Mean ± S.E.)</th>
<th>Nicotine groups (Mean ± S.E.)</th>
<th>Green Tea Groups (Mean ± S.E.)</th>
<th>Nicotine -Green Tea Groups (Mean ± S.E.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Week 1</td>
<td>23 4.60 ± 0.748&lt;sup&gt;a&lt;/sup&gt;Aa</td>
<td>31 6.20 ± 0.860&lt;sup&gt;a&lt;/sup&gt;Aa</td>
<td>20 4.00 ± 0.707&lt;sup&gt;a&lt;/sup&gt;Aa</td>
<td>6.80 ± 0.663&lt;sup&gt;b&lt;/sup&gt;Bb</td>
</tr>
<tr>
<td>Week 2</td>
<td>4.00 ± 0.837&lt;sup&gt;b&lt;/sup&gt;Ba</td>
<td>9.60 ± 1.166&lt;sup&gt;a&lt;/sup&gt;Aa</td>
<td>3.80 ± 0.860&lt;sup&gt;a&lt;/sup&gt;Ba</td>
<td>6.40 ± 1.208&lt;sup&gt;b&lt;/sup&gt;Bb</td>
</tr>
<tr>
<td>Week 3</td>
<td>4.40± 0.678&lt;sup&gt;c&lt;/sup&gt;Ba</td>
<td>15.00± 1.224&lt;sup&gt;ab&lt;/sup&gt;Ba</td>
<td>3.60 ± 0.509&lt;sup&gt;c&lt;/sup&gt;Ca</td>
<td>8.90 ± 1.140&lt;sup&gt;b&lt;/sup&gt;Ba</td>
</tr>
<tr>
<td>Week 4</td>
<td>4.20 ± 0.583&lt;sup&gt;a&lt;/sup&gt;Ba</td>
<td>30.60± 4.523&lt;sup&gt;a&lt;/sup&gt;Aa</td>
<td>3.20 ± 0.583&lt;sup&gt;a&lt;/sup&gt;Ba</td>
<td>9.00 ± 1.140&lt;sup&gt;b&lt;/sup&gt;Ba</td>
</tr>
</tbody>
</table>

4. Discussion:

Our result showed that disorganization of the seminiferous tubules associated with reduction of spermatogenic cells, leading to widening of lumen of seminiferous tubules upon nicotine toxicity. Many of seminiferous tubules exhibited degenerative phases of spermatocytes and spermatides as well as missing of sperms and hypo-spermatogenesis.

Fávaro et al. (2006); Ahmadnia et al. (2007) and Yamamoto et al. (2007) reported many alterations attributed to the direct cytotoxic effects of nicotine leading to the inhibition of prostaglandin's synthesis and decrease of testosterone which play a functional role in reproduction system of the male mice. Other investigation displayed that nicotine was a CNS depressor that can inhibit the neural stimulus essential for the release of pituitary gonadotrophine (Reddy et al., 1998), which lead to a lack of pituitary gonadotrophins essential for initiating and completing spermatogenesis and steroidogenesis in the testis (Aydos et al., 2001). Destruction of Leydig cells may cause testicular atrophy; gonadal dysfunction, erectile dysfunction, and male factor infertility were showed by Kim et al., (2005). Other studies explained that metabolites of nicotine as cotinine produced dose response inhibition of luteinizing hormone and 3α-hydroxysteroid dehydrogenase, enzyme stimulated testosterone production (50-70%) (Fávaro et al., 2006).

Aydos et al., (2001) and Ahmadnia et al., (2007) explained that nicotine toxicity may be due to change in the proportion of collagen fibers and contractile myofibroblastic cells, which may prevent the appropriate release of spermatozoa from the germinal epithelium into the lumen.

The oxidative stress of nicotine may cause a peroxidant/antioxidant imbalance in blood cells, blood plasma, and tissues resulting in a decrease in the activity of endogenous antioxidant enzymes (Suleyman et al., 2002). Other researchers reported disruption of spermatogenesis in nicotine treated animals' testes tissues may be modulated by free radical toxicity (El-Sweedy et al., 2007). Also nicotine metabolite cotinine generates free radicals / ROS in tissues in the liver and testes (Kalpan and Menon, 2004), and induces oxidative tissue injury
(Husain et al., 2001 and Argentine & Cicchetti, 2004).

Direct genotoxic effects have been shown in human gingival fibroblasts (Sassen et al., 2005 and Kleinsasser et al. (2006) and DNA strand breaks in human spermatoza in nicotine aneuploidy and polyplody, sister chromatid exchanges and chromosome aberrations in bone marrow cells of mice (Sen and Sharma, 1991 and Yauk et al., 2007). High doses of nicotine increase the frequencies of premature centromere separation and premature anaphase and reduce the number of oocytes ovulated (Attia, 2007 and Sudheer et al., 2007).

Results concerning the mutagenicity of nicotine in several test systems are contradictory. In the present study, nicotine intoxicated group obtained highly significant and gradual increased of MnPCEs throughout all the tested experimental periods. The statistical analysis of data is also provided. Nicotine treated mice, for one, two, three and four weeks resulted in highly significant alterations of all monitored parameters, but with different intensity and distinctive time trends. Nicotine intoxicated group obtained highly significant and gradual decreased of PCEs and PCEs /NCEs but gradual increased in NCEs throughout all the tested experimental periods.

The differences between the results can be attributed to different drug concentrations and the different genotoxic endpoints considered in the test systems. Different repair capacities of the various cell types used may also explain the discrepancies. The micronucleus technique has been proposed as a useful tool for measurement of genotoxicity (Madhumita et al., 2003) induced by nicotine as it is able to assess both the clastogenic and aneugenic properties of a test compound. The genotoxic effects of nicotine have been shown in human gingival fibroblasts and spermatoza and damage to DNA may result in mutations and altered cell functions (Kleinsasser et al., 2005 and Ogura et al., 2008).

In the present investigation a significant increase in the levels of NCE, and increased micronuclei frequency in nicotine-treated groups, as indicative of DNA damage, when compared with control group. The results were in a gradient with those reported by Villard et al. (1998) who mentioned that nicotine causes in vivo DNA single-strand breaks (SSB) in lung and liver of mice and those reported by Sassen et al. (2005) who reported that nicotine increases the DNA fragmentation in mini organ cultures. Hassan and Ahmed 2004, proved that green tea were inhibit the induction of Mn-PCEs in bone marrow of mice by 66% and 90% when treated for one and two weeks respectively. Ayako (2004) suggests that tea catechins are not genotoxic but rather have a preventive effect against reactive oxygen species ROS -induced chromosomal damage at their physiological condition. These useful effects of tea catechins against ROS-induced chromosomal damage may support the cancer-preventing effects of tea constituents (Fujiki et al., 2002).

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