Role of ginger (zingiber officinale) against metalaxyl induced hepatotoxicity in male albino rats: a histological and immunohistochemical study

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Abstract

Background: Metalaxyl is a fungicide used to control soil-borne fungal diseases on fruits, cotton, soy bean, peanuts and grasses. Food contamination with metalaxyl is a worldwide problem especially in developing countries. Ginger has antioxidant properties.

Aim of the work: To elucidate the histological alterations that may occur in the liver tissue of rat by metalaxyl and to investigate the role of ginger supplementation against these alterations.

Materials and methods: Thirty healthy adult male albino rats were used in this study. They were equally divided into three groups:
Group I served as control group. Group II (metalaxyl-treated) received 130 mg/kg body weight/day of metalaxyl dissolved in distilled water using a gastric tube for 3 times/week for continuous 4weeks. Group III (metalaxyl-ginger treated group) received metalaxyl as group II in addition to ginger given orally at a dose of 100 mg/kg three times/week for successive 4 weeks. At the end of experiment, all rats were anaesthetized and liver specimens were taken and processed for light and electron microscope examination. Immunohistochemical staining was carried out for detection of the proapoptotic marker Bcl-2 antagonist-X (BAX) protein in hepatocytes. Area % of BAX protein was measured and statistically analyzed.

Results: Examination of the liver sections of metalaxyl-treated group II showed different degrees of focal lobular affection. Central veins and blood sinusoids were dilated and congested. The hepatic lobules lost their normal architecture and the hepatocytes showed cytoplasmic vacuolization with darkly stained nuclei. Enlarged portal areas with numerous bile ductules and cellular infiltration were detected. Ultrastructurally, hepatocytes had condensed heterochromatin nuclei and electron lucent areas were observed in their cytoplasm. Significant increase in area % of positive immunoreaction for BAX was detected in comparison with control group and group III. Examination of liver sections of group III revealed preservation of nearly normal histological structure with slight congested blood vessels and few cellular infiltration.

Conclusion: Metalaxyl had pathological effects on the structure of rat’s liver and these effects were decreased by ginger.

Keywords: Metalaxyl, ginger, liver histology, ultrastructure, rat

Introduction

Metalaxyl is a benzenoide fungicide that widely used in developing countries; it is used to protect crops against soil-borne pathogens [1]. The problems resulting from metalaxyl come from their high residual level in agriculture crops especially vegetables cultivated under greenhouse conditions and during storage and germination of seed grain [2,3]. Dasgupta et al., [4] reported that residues of buprafezin, chlorpyriphas, metalaxyl and mychobutanil were detected in cured grapes and wine samples.

Metalaxyl was reported to have cytogenetic effects on human and animal chromosomes in vitro [5] and Demsia et al., [6]
found that imidaclopride and metalaxyl have genotoxic effects as they induced micronucleus formation in human lymphocytes in vitro and in polychromatic erythrocytes of the rat’s bone marrow in vivo.

AL-Amoudi [7] detected hematological effects and co-carcinogenic potential of metalaxyl in Swiss albino mice. Reproductive toxicity of metalaxyl and mancozeb in adult male albino rat was detected by Rao et al., [8] and Mehra et al., [9]. Recent published studies have reported that metalaxyl toxicity may be associated with the enhanced production of reactive oxygen species (ROS). The production of ROS is caused by a mechanism in which xenobiotics, toxicants and pathological conditions may produce oxidative stress. If ROS formation exceeds the capacity of antioxidants, ROS react with macro-molecules such as lipid, protein and DNA causing cell dysfunction and damage [10].

Ginger (Zingiber officinal Roscoe) is an example of botanicals which gains popularity amongst modern physicians and its underground rhizomes are medicinally useful parts [11]. One of the most popular uses of ginger is to relieve the symptoms of nausea and vomiting associated with motion sickness, surgery and pregnancy [12]. Ginger extracts had different pharmacological effects such as anti-inflammatory [13], anti-oxidant [14], anti-tumor [15] and anti-nephrotoxic effects [16].

Liver is an important organ in the body as it is responsible for removal of toxins and poisons and it is greatly affected by pollutants that cause increase in liver enzymes [17]. The histological study of the liver and immunohistochemical detection of proapoptotic marker bcl-2 antagonist-X protein (BAX) in hepatocytes after metalaxyl administration to albino rats may clarify metalaxyl hazardous effects on liver.

So, the aim of the present study was detection of the histological and immunohistochemical changes that may occur by metalaxyl on liver of adult rat and role of ginger against these changes.

Materials and methods

Animals

In this study, we used thirty healthy adult male albino rats aged 3-4 months and weighed 180-200gm. They were housed in stainless steel cages at Animal House, Faculty of Medicine, Zagazig University at room temperature (25±1°C) and illumination (normal light and dark cycle), fed standard balanced diet and water ad- libitum. One week after acclimatization rats were randomly divided into three equal groups (10 rats in each group). The experiment was performed in accordance to the “Guide for the Care and Use of Laboratory Animals” [18].

Experimental design

Group 1 (control group)

Animals were further subdivided into two equal subgroups: Subgroup 1a: Included animals that received ordinary food and distilled water (the vehicle of metalaxyl) by gastric tube three times/week for successive 4 weeks. Subgroup 1b: included rats that received 100 mg/kg body weight of ginger orally three times per week by gastric tube for successive 4 weeks.

Group II (Metalaxyl-treated)

Animals were given metalaxyl at a dose of 130 mg/kg body weight. Metalaxyl was dissolved in distilled water as a vehicle and given orally by means of gastric tube three times/week for continuous 4 weeks [19]. Metalaxyl was supplied from Central Agricultural Pesticides Laboratory, ARC, Egypt.

Group III (Metalaxyl and ginger treated group)

Rats were given the same dose of metalaxyl given to animals of group II (130 mg/kg body weight three times/week) and after 1 hour they received ginger dissolved in distilled water by gastric tube at a dose of 100 mg/kg body weight [16] three times/week for successive 4 weeks. Ginger was purchased from Mepaco-Medifood, (Enshas, Sharkeya, Egypt, Ministry of Health Reg No, 65006).

At the end of the experiment, the rats were anesthetized with 50 mg/kg body weight sodium pentobarbital by intraperitoneal injection. Intra-cardiac perfusion was done by 2% glutaraldehyde for partial fixation. Liver specimens from right lobes were dissected out and processed for light, transmission electron microscope and immunohistochemical study.

Histological study

For light microscope, specimens were fixed in 10% buffered formalin for 24 hours and processed to prepare 5um sections stained with hematoxylin and eosin (H&E) to verify the histological structures [20]. Immunohistochemical staining was done for detection of proapoptotic marker bcl-2 antagonist X protein (BAX) [21,22].

For transmission electron microscope, small pieces (1mm³) from liver were immediately fixed in 2.5% glutaraldehyde in 0.1M cacodylate buffer at pH 7.4 for 24hours at 4°C then the specimens were washed with the buffer, post fixed in 1% osmium tetroxide in distilled water for 2h at 4°C. Specimens were dehydrated with ascending grades of ethanol and embedded in epoxy resin. After staining the ultrathin sections by uranyl acetate and lead citrate according to Hayat [23], they were examined by using a JEOL JEM 1010 and JEOL JEM 1200 EXI (Japan) electron microscopes (JEOL, Ltd, Tokyo, Japan). Preparation and examination of ultrathin sections were done in the Electron Microscope laboratory of the Histology and cell Biology Department, Faculty of Medicine, Zagazig University, Egypt and in the Electron Microscope Research Laboratory, Faculty of Science, Ain Shams University, Egypt.

For immunohistochemical study, liver sections were deparaffinized in xylene, rehydrated in descending grades of alcohol and immersed in 0.1% hydrogen peroxide to block the endogenous peroxidase activity. Mouse monoclonal antibody (Ab-14 Golden, Lab Vision Clone B-9, Santa Cruz
Biotechnology Inc., Santa Cruz, California, USA) for BAX was put on each section, the dilution used was 1:50. The antibody was detected by using a biotin-streptavidin system with diaminiobenzide (DAB) that applied for 20 min at room temp as a chromogen (Dako Carpentaria California USA) for BAX (Dako Corp., code no, k0673, lot 07110). Slides were counterstained with Mayer’s hematoxylin (Park Scientific Limited, Northampton, UK), dehydrated and covered by cover slips. In negative control slides, the same system was applied but the primary antibody was not added. The (BAX) cytoplasmic site of reaction was stained brown and nuclei stained blue. This immunohistochemical technique was carried out in the Department of Pathology, Faculty of medicine, Cairo University.

Quantitative morphometric study
At this study, Leica Qwin 500 Image Analyzer computer system (Leica Microsystem Imaging Solution Ltd., Cambridge, UK) in the image analyzing unit at Pathology Department Faculty of Dentist, Cairo University was used to estimate the area % for positive BAX immunoreaction in the hepatocytes of all studied groups. Measurements were done from ten non-overlapping fields for each section at x-400 magnification to determine the area % of BAX immunoreaction for detection of apoptosis. For each specimen the mean values and SD were calculated automatically using the image analyzer.

Statistical analysis
BAX immunoreactions were expressed as mean±standard deviation (X±SD). The data were subjected to SPSS program (SPSS Inc., Chicago, 111inois, USA). Statistical analysis was carried out using T test for comparison between subgroups la and lb of the control group, one-way analysis of variance (ANOVA) and the least significant difference test (LST) for comparison among groups. The results were considered statistically significant, highly significant and non significant when the P values were <0.05, <0.001 and more than 0.05 respectively.

Results
Histological results
Group I (control group)
Light microscope examination of liver sections of control subgroups showed the same findings. Hence, we chose the results of subgroup 1a to represent the control group. The liver lobules consisted of anastomosing cords of hepatocytes radiating from the central vein. The hepatocytes appeared acidophilic with central rounded vesicular nuclei. Few cells were binucleated. Blood sinusoids with their kupffer and endothelial cells lining were noticed between the hepatocyte plates (Figure 1). Each portal area contained branches of portal vein and hepatic artery. Bile duct with cuboidal epithelial lining was also seen. Some hepatocytes with pale acidophilic cytoplasm and pale vesicular nuclei were observed (Figure 2). Weak positive granular immunoreaction for BAX was diffusely scattered in the cytoplasm of hepatocytes (Figure 3).

Liver sections of control group examined by electron microscope showed hepatocytes with euchromatic nuclei, mitochondria, cisternae of rough (rER) and smooth endoplasmic reticulum (sER) (Figure 4).
Figure 3. A photomicrograph of a section in a control adult male albino rat’s liver showing weak positive granular immunoreactions for BAX diffusely scattered in the cytoplasm of hepatocytes [Immunoperoxidase technique for BAX x400].

Figure 5. A photomicrograph of a section in the liver of metalaxyl treated group II showing dilated congested central vein (CV) and blood sinusoids (s) [H&E ×400].

Figure 6. A photomicrograph of a section in the liver of metalaxyl treated group II showing that most of the peripheral hepatocytes (thin arrows) appear with cytoplasmic vacuolization. Centrilobular hepatocytes (thick arrows) contain deeply stained acidophilic cytoplasm and darkly stained nuclei. The central vein (CV) is seen [H&E ×400].

Immunohistochemical stained sections of metalaxyl treated group revealed strong positive immunoreaction for BAX in nearly all hepatocytes (Figure 8).

Electron microscope examination of the liver sections of metalaxyl treated group II showed hepatocytes with condensed heterochromatin nuclei and irregular nuclear envelopes. Electron lucent areas within the cytoplasm, lipid droplets, mitochondria, rough and smooth endoplasmic reticulum were seen. Apoptotic nuclei were also noticed (Figures 9 and 10).

Group III (metalaxyl and ginger)
Examination of the liver sections of group III, received ginger concomitantly with metalaxyl showed preservation of nearly
Figure 7. A photomicrograph of a section in the liver of metalaxyl treated group II showing enlarged portal area with branches of dilated portal vein (PV), congested hepatic artery (A), cellular infiltration (*) and numerous bile ductules (thin arrows). One of those bile ductules shows multiple epithelial layers (thick arrow) [H&E ×400].

Figure 8. A photomicrograph of a section in the liver of metalaxyl treated group II showing strong positive immunoreaction for BAX in nearly all hepatocytes. Part of dilated central vein (CV) is also seen [Immunoperoxidase technique for BAX ×400].

Figure 9. An electron micrograph of a section in the liver of metalaxyl treated group II showing hepatocytes with condensed heterochromatin nuclei (N) and irregular nuclear envelop (arrow). The cytoplasm contains electron lucent area (*), mitochondria (m), lipid droplet (L) and rough endoplasmic reticulum (r). Apoptotic nucleus (n) is also seen [Mic. Mag ×5000].

normal hepatic lobular architecture with the presence of slightly dilated congested central veins and blood sinusoids with few cellular infiltration (Figure 11). Portal area contained normal bile duct and congested portal vein. Hepatocytes appeared with acidophilic cytoplasm and vesicular nuclei. Slightly dilated congested blood sinusoids were seen (Figure 12).

Immunohistochemical examination of liver section of group III showed weak positive immunoreaction for BAX in the cytoplasm of most hepatocytes and few cells showed strong reaction (Figure 13).

Electron microscope examination of the liver sections of the group III showed preserved hepatocyte ultrastructure with euchromatic nucleus and two prominent nucleoli (Figure 14). Some hepatocytes were binucleated (Figure 15). Their cytoplasm contained aggregations of rough and smooth endoplasmic reticulum and many mitochondria (Figures 14 and 15). Fat droplets were seen (Figure 15).

Statistical results
Statistical comparison between subgroups 1a and 1b as regard to the area % of BAX immunoreaction revealed no significant difference (P>0.05). Therefore, the control group 1a was used for comparison with other groups (Table 1 and Histogram 1).

Metalaxyl treated group (II) showed a statistically significant increase in area % of BAX immunoreaction when compared to other groups. Also, there is statistical significant difference between group I and II and between groups II and III (p<0.05) but there is no statistical significant difference between group I and III (p>0.05) (Table 2 and Histogram 2).

Discussion
Liver is the primary target organ of metalaxyl toxicity [19,24].
Results obtained in the present study showed that metalaxyl induced histopathological alterations in the liver tissue such as congestion of blood vessels, cytoplasmic vacuolization of hepatocytes, apoptosis and necrosis.

### Table 1. The mean area % of BAX immunoreaction of subgroups 1a and 1b of control rats.

<table>
<thead>
<tr>
<th></th>
<th>X±SD</th>
<th>Range</th>
<th>T test</th>
<th>P-Value</th>
</tr>
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<tbody>
<tr>
<td>Control a</td>
<td>6.56±2.58</td>
<td>4.18-10.06</td>
<td>0.95</td>
<td>&gt;0.05 N.S</td>
</tr>
<tr>
<td>Control b</td>
<td>9.08±5.43</td>
<td>11.17-26.03</td>
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</tr>
</tbody>
</table>

N.S: Non significant; T: Independent sample T test
This table shows no significant difference between subgroups 1a and 1b of control rats (P>0.05).
Histological examination of the liver sections of group II revealed different forms of focal lobular affection with congested dilated central vein and blood sinusoids. These changes were also previously observed with metalaxyl [24,25] and other fungicides [26,27]. These previous studies referred dilatation to increased levels of prostaglandins that induce smooth muscle relaxation with subsequent vasodilatation. Congestion might be due to loss of fluid from the blood and the vessels engorged with RBC’s [28-30].

In liver sections of group II, cytoplasmic vacuolizations were observed in hepatocytes in peripheral regions of liver lobules while those around central veins were less affected. Sigala

Table 2. The mean area % of BAX immuno-reaction in different studied groups.

<table>
<thead>
<tr>
<th>Group</th>
<th>X±SD</th>
<th>Range</th>
<th>ANOVA</th>
<th>P-Value</th>
<th>LSD</th>
</tr>
</thead>
<tbody>
<tr>
<td>ІІ</td>
<td>6.56±2.58</td>
<td>4.18-10.06</td>
<td>4.64</td>
<td>&lt;0.05*</td>
<td>&gt;0.05 N.S</td>
</tr>
<tr>
<td>ІІІ</td>
<td>17.80±8.94</td>
<td>10.03-32.01</td>
<td>--</td>
<td>--</td>
<td>&lt;0.000**</td>
</tr>
<tr>
<td>І</td>
<td>11.10±4.11</td>
<td>5.94-16.72</td>
<td>--</td>
<td>--</td>
<td>&lt;0.05*</td>
</tr>
</tbody>
</table>

ANOVA: Analysis of variance test; LSD: Least significant difference

Group І: control; Group ІІ: Metalaxyl-treated; Group ІІІ: Metalaxyl-ginger-treated
N.S: Non significant *significant **highly significant

This table shows high significant difference between group II and other groups by ANOVA test. By LSD test, there is statistical significant difference between group І and ІІ and also between group ІІ and ІІІ but there is no statistical significant difference between group І and ІІІ.
et al., [31] attributed the affection of periportal hepatocytes to rapid uptake of toxins from portal vessels and fast rate of lipid peroxidation.

Lange and Gartzk [32] revealed that cytoplasmic vacuolizations caused by the disruption in microvilli which are normally act as diffusion barrier which inhibit the entrance of hydrophilic and lipophilic xenobiotics into the cytoplasm. Sakr and Shalaby [33] attributed similar findings to the oxidative activity of fungicides with subsequent generation of superoxide anions that cause lipid peroxidation. Lipid accumulation leads to alteration and damage of cellular lipid membranes with paralysis of Na-K pump and hepatocytes edema.

In the current study, numerous bile ductules were detected in the portal areas of group II. The same results were observed by other researchers [34,35] and attributed this proliferation to trans-differentiation of hepatocytes into biliary cells or the mitogenic effect of bile salt stasis on biliary epithelial cells. Cants et al., [36] and Zaghoul [37] reported that endogenous stem cells located at the junction between hepatocytes and the terminal bile ductules may play role in bile duct proliferation.

Cellular infiltration was noticed in the portal areas in liver sections of group II in this study. These observations were in accordance with the results obtained by other studies [38,39] that referred this cellular infiltration to ROS production which indirectly regulate chemokine receptor expression and promote cytokine IL-6 and IL-8 which are key modulators of inflammatory response. Other investigators considered that cellular infiltration as a prominent immune response of the body tissues by movement of fluids and leukocytes from the blood into the extravascular tissues [28,30].

Electron microscope study of liver sections of group II in the current work, showed that some hepatocytes contained nuclei with condensed heterochromatin and irregular nuclear envelop which most probably indicating apoptosis. Electron lucent areas in their cytoplasm were also seen revealing necrosis. Similar results were observed by Hashem [10] who correlated apoptosis and necrotic cell death of hepatocytes in metalaxyl treated rats with high blood and tissue levels of malondialdehyde which is a marker for oxidative stress.

In the current study, ginger treatment simultaneously with metalaxyl in the group III leads to preservation of the liver histological structure with slight dilatation and congestion of blood vessels and few cellular infiltration. Ultrastructurally, hepatocytes showed normal appearance of their nuclei and cytoplasmic organelles. The hepato-protective effect of ginger may be due to its volatile oils which had anti-inflammatory, anti-analgesic and immunomodulatory effects [11,13]. Previous reports have documented the ability of ginger ingredients as gingerols to inhibit prostaglandins and leukotrienes synthesis [16,19]. It was reported that ginger have hepatoprotective effect due to its antioxidiant activity to reduce the activity of free radicals [19]. Some lipid droplets were noticed in the cytoplasm of some hepatocytes of group III. This was explained as a defense mechanism by which hepatocytes attempt to collect all toxic compounds invading the cells prior to excretion [10].

Immunohistochemical study showed strong positive immunoreaction of BAX in nearly all hepatocytes in the affected areas of group II compared to the control group indicating that these cells are susceptible to apoptosis. Statistical analysis revealed significant increase in the area % of BAX-in group II when compared with other groups. Other studies reported the same results of BAX immunoreaction in mouse liver [22] and thymus [40].

Apoptosis is a mode of cell death which occurred physiologically [22]. Chen et al., [41] noticed high expression of BAX protein in chronic hepatitis. Other researchers referred apoptosis of hepatocytes to activation of tumor protein (p53) and release of cytochrome c from the damaged mitochondria to the cytoplasm of hepatocytes [42].

The organophosphorus pesticides induced apoptosis by activation of intracellular cysteine-aspartic acid protease (caspase-3) [43] or by oxidative stress and ROS which lead to DNA damage [44,45].

Fungicides induced a significant decrease in the serum antioxidant enzymes activities, superoxide dismutase and increase in lipid peroxidation in albino rats [27,46]. Marked reduction in antioxidant enzymes activities and tissue glutathione contents resulted in oxidative damage of tissues and leads to apoptosis in cellular systems, especially hepatocytes [7,47].

In the current study, immunodetection of BAX protein in liver sections of group III showed weak positive reaction in most of hepatocytes compared to the strong positive reaction observed in group II and significant decrease in the mean area % of BAX when compared with group II suggesting that ginger reduced apoptosis. These findings were supported by previously performed clinical and experimental investigations in liver of mice [19], testis [48] and kidney [16] that have shown
that ginger has a protective effect against oxidative damage and apoptosis through its antioxidant properties.

Lamfon [19] reported that the underground rhizome of ginger contained many flavonoids with anti-oxidant activity which prevents free radicals generation. Also, previous investigators demonstrated that ginger oil had a dommative protective effect on DNA damage and might act as a scavenger of oxygen radicals [49,50].

Conclusion
Metalaxyl causes histological and immunohistochemical changes in liver probably through oxidative stress. Ginger therapy could ameliorate these changes in liver and this may be attributed to its antioxidant and free radicals scavenging properties. This study showed that ginger supplementation may minimize the hazardous effects of metalaxyl.

Competing interests
The author declares that he has no competing interests.

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