Comparing the protective effects of L-carnitine and *Silybum marianum* aqueous extract after diazinon-induced hepatotoxicity in male rat liver

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**ABSTRACT**

Diazinon (DZN), as an organophosphorus pesticide, induces oxidative stress leading to the generation of free radicals, and causes some pathological changes in the body. The purpose of this study was to assess the protective effects of L-carnitine (LC) and *Silybum marianum* aqueous extract (SMAE) against DZN-induced hepatotoxicity in male rat liver. Rats were assigned in 9 groups and were subjected to different combinations of DZN, SMAE, and LC. Thirty days after the treatment by oral gavage, blood samples were taken and serum levels of aspartate aminotransferase (AST), alanine aminotransferase (ALT), alkaline phosphatase (ALP), gamma-glutamyl transferase (GGT), Albumin (Alb) and total protein (TP) were measured using photometric method. The liver samples were also evaluated histopathologically. The DZN treatment significantly increased the AST, ALT, ALP and GGT levels (p < 0.05) and conversely decreased the Alb and TP levels (p < 0.05). Moreover, administration of the DZN resulted in lymphocytic inflammation, congestion, hepatocytes apoptosis, and lesser sinusoidal space. However, administration of SMAE and LC along with DZN treatment stabilized the ALT, ALP, GGT, Alb and TP levels (p > 0.05), but increased the AST levels significantly compared to the control (p < 0.05). Besides that, lesser structural alterations and improvement in the liver tissue were observed. These findings suggest that co-administration of SMAE and LC could reduce DZN-induced hepatic tissue damages and improve the hepatic biochemical parameters in male rats.

**Keywords**

*Diazinon, L-carnitine, Silybum marianum, Hepatotoxicity, Rat*

**Abbreviations**

DZN: Diazinon  
OP: Organophosphorus pesticide  
LC: L-carnitine  
SAME: *Silybum Marianum* Aqueous Extract  
SM: *Silybum Marianum*  
AST: Aspartate aminotransferase  
ALT: Alanine aminotransferase  
ALP: Alkaline phosphatase  
GGT: Gamma-glutamyl transferase  
Alb: Albumin  
TP: Total protein
Introduction

Liver is the largest gland in the body which performs a variety of functions, including protein synthesis, storing required substances, metabolic functions, disinfecting harmful substances, producing and secreting bile as well as controlling metabolism [1]. Organophosphorus pesticide (OP) refers to various toxic compounds that contain phosphorus in their basic structure. They have been introduced to the world market since 1960s. Since they were more potent than other toxins such as organochlorine, OP compounds replaced other toxins rapidly. In 1970, due to their environmental and food chain contamination they were only used in certain cases in developed countries, however, they are still used in some countries [2].

One of the most important OPs used to control insects in agriculture is DZN. This compound is easily and quickly (within a few hours) absorbed by the intestine [3,4]. This compound inhibits acetylcholine esterase through the phosphorylation of amino acid serine at its active site, resulting in the accumulation of acetylcholine in cholinergic synapses, and causing cholinergic seizure and brain injury and death in acute cases [5,6]. DZN is also absorbed through skin and respiratory tract, and rapidly metabolized to diazoxide in the liver. Most OP compounds are converted to active toxic metabolites in the liver by the cytochrome P450 system through oxidative desulphurization [7]. It should be noted that the side effects of these compounds depend on the type of toxin, dose, duration of exposure and also the tissue. The main mechanism of OP action, especially DZN, is inhibiting acetylcholine esterase [8]. They are alkylated agents that react with cellular macromolecules, such as proteins, nucleic acids and lipids, and alter their functions [9]. Some researchers believe that DZN induces oxidative stress by producing free radicals and reactive oxygen, and induces cell death in organisms by increasing lipid peroxidation [10]. Indeed, exposure to DZN causes severe histopathologic damages in the kidney and liver [11].

Antioxidant compounds play an important role in preventing damages and pathological changes caused by free radicals. For example, LC prevents oxidative stress and regulates nitric acid, cellular respiration, and the activity of enzymes involved in oxidative stress [12,13]. The antioxidant system consists of three enzymes: glutathione peroxidase, catalase and superoxide dismutase. As an antioxidant compound, LC can protect these enzymes against oxidative damage; it is also very effective in modulating age-related changes [14]. Food sources of amino acid carnitine are very important. Approximately 75% of total carnitine is obtained from carnitine, lysine, and methionine food sources [15]. The condition of carnitine in the human body depends on the body composition, sex and diet. There are two sources for these amino acids: (1) diet and (2) degradation of endogenous proteins. Micronutrients such as iron, vitamin C, pyridoxine and niacin are essential for the synthesis of carnitine [16].

The plant Silybum marianum (SM) is an annual or biennial plant of the Asteraceae family [17]. SM has global vegetation and is native to Iran as well [18]. SM contains silybin, isosilybin, silydianin and taxifolin, which are collectively called flavonoids silymarin. Silymarin is found in all parts of the plant and is considered as the main and effective compound of SM. The main pharmacological properties of this plant include: antioxidant, anti-inflammatory and anti-cancer effects and also protecting hepatic cells against many liver toxins [19,20].

Since LC and SM show antioxidant and anti-inflammatory effects and are used in the treatment of liver disorders, in this study, we tried to investigate their possible hepatoprotective effects in DZN-induced hepatic damage in male rats.

Results

Biochemical analysis findings

Table 1 and 2 represent the mean and standard deviation (SD) of the serum levels of ALT, AST, ALP, and GGT as well as Alb and TP among different groups, respectively. The findings of this study showed that there was no significant difference in the serum levels of ALT, AST, ALP, GGT, Alb and TP between the control and sham groups (p > 0.05). Compared to the control and sham groups, the serum levels of ALT, AST, ALP, and GGT increased significantly (p < 0.05) in the DZN15 group, while the serum levels of Alb and TP decreased significantly (p < 0.05).

Statistical comparison of the results showed that there were no significant differences in the serum levels of ALT, AST, ALP, GGT, Alb and TP in the SMAE100, LC300 and SMAE100 + LC300 groups, when compared to the control and sham groups (p > 0.05).

In the DZN15+MS100 and DZN15+LC300 groups, the serum levels of ALT, AST, ALP, and GGT increased significantly compared to the control and sham groups (p < 0.05); however, they showed a significant decrease compared to the DZN15 group (p < 0.05). In addition, the serum levels of Alb and TP decreased significantly in the DZN15+MS100 and DZN15+LC300 groups compared to the control and sham groups (p < 0.05); while, showed a significant increase compared to the DZN15 group (p < 0.05).

In the DZN15 + SMAE100 + LC300 group, the se-
rum levels of ALT, AST, ALP, and GGT decreased significantly compared to the DZN group (p < 0.05) but the serum levels of Alb and TP increased significantly (p < 0.05). Although, there was no significant difference in the serum levels of ALT, ALP, GGT, Alb and TP (p > 0.05), the serum levels of AST increased significantly in the DZN15 + SMAE100 + LC300 group compared to the control and sham groups (p < 0.05).

**Histopathological findings**

Histopathological findings on the liver tissues indicated that the liver parenchymal cells, sinusoid spaces and central vein in the control and sham groups were completely normal and were not damaged (Figure 1A and 1B). In the DZN15 group, lymphocytic inflammation, congestion, hepatocyte apoptosis, and lesser sinusoidal space were observed (Figure 1C). There were no signs of liver damage in the LC300, SMAE100, and SMAE100 + LC300 groups (Figure 1D, 1E and 1F). In the DZN15 + LC300 and DZN15 + SMAE100 groups, lymphocytic inflammation and vacuolization of cytoplasm were detected (Figure 1G and 1H). Histological cross-sectional study in the DZN15 + SMAE100 + LC300 group revealed lesser pathological changes and improvement in the liver tissue compared to the DZN group (Figure 1I).

### Table 1
Comparison of serum levels of aspartate aminotransferase (AST), alanine aminotransferase (ALT), alkaline phosphatase (ALP), and gamma glutamyl transferase (GGT) in different groups.

<table>
<thead>
<tr>
<th>Groups</th>
<th>AST (u/l)</th>
<th>ALT (u/l)</th>
<th>ALP (u/l)</th>
<th>GGT (u/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>359.87 ± 10.46</td>
<td>82.37 ± 5.18</td>
<td>777.62 ± 16.26</td>
<td>5.37 ± 0.406</td>
</tr>
<tr>
<td>Sham</td>
<td>365.12 ± 09.49</td>
<td>84.87 ± 4.51</td>
<td>769.75 ± 13.62</td>
<td>5.27 ± 0.541</td>
</tr>
<tr>
<td>DZN15</td>
<td>782.25 ± 11.81 *</td>
<td>180.25 ± 9.06 *</td>
<td>1278.62 ± 37.30</td>
<td>11.43 ± 0.462 *</td>
</tr>
<tr>
<td>SMAE100</td>
<td>362.25 ± 10.12 b</td>
<td>84.62 ± 7.81 b</td>
<td>779.00 ± 18.60 b</td>
<td>5.26 ± 0.434 b</td>
</tr>
<tr>
<td>LC300</td>
<td>356.00 ± 09.05 b</td>
<td>85.42 ± 4.57 b</td>
<td>773.00 ± 16.69 b</td>
<td>4.87 ± 0.281 b</td>
</tr>
<tr>
<td>SMAE100 + LC300</td>
<td>352.25 ± 08.27 b</td>
<td>81.00 ± 4.24 b</td>
<td>769.75 ± 14.95 b</td>
<td>4.91 ± 0.285 b</td>
</tr>
<tr>
<td>DZN15 + LC300</td>
<td>533.00 ± 07.65 * ,b</td>
<td>115.125 ± 5.56 * ,b</td>
<td>1001 ± 55.79 * ,b</td>
<td>7.68 ± 0.318 * ,b</td>
</tr>
<tr>
<td>DZN15 + SMAE100</td>
<td>513.28 ± 07.52 * ,b</td>
<td>105.42 ± 10.57 * ,b</td>
<td>949.00 ± 73.07 * ,b</td>
<td>6.87 ± 1.17 * ,b</td>
</tr>
<tr>
<td>DZN15+ SMAE100 + LC300</td>
<td>411.87 ± 13.95 * ,b</td>
<td>90.25 ± 4.13 * ,b</td>
<td>804.12 ± 19.88 * ,b</td>
<td>5.22 ± 0.310 * ,b</td>
</tr>
</tbody>
</table>

Data is shown as mean ± SD.
*a*, p < 0.05 as compared to the control and sham groups
*b*, p < 0.05 as compared to the DZN15 group

### Table 2
Comparison of serum levels of Albumin (Alb) and Total protein (TP) in different groups.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Alb (g/dl)</th>
<th>TP (g/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>3.55 ± 0.244</td>
<td>6.58 ± 0.473</td>
</tr>
<tr>
<td>Sham</td>
<td>3.53 ± 0.277</td>
<td>6.52 ± 0.337</td>
</tr>
<tr>
<td>DZN15</td>
<td>1.75 ± 1.92 a</td>
<td>3.85 ± 0.272 a</td>
</tr>
<tr>
<td>SMAE100</td>
<td>3.60 ± 0.261 b</td>
<td>6.28 ± 0.485 b</td>
</tr>
<tr>
<td>LC300</td>
<td>3.52 ± 0.269 b</td>
<td>6.20 ± 0.258 b</td>
</tr>
<tr>
<td>SMAE100 + LC300</td>
<td>3.97 ± 0.361 b</td>
<td>6.00 ± 0.392 b</td>
</tr>
<tr>
<td>DZN15 + LC300</td>
<td>2.68 ± 0.247 * ,b</td>
<td>5.30 ± 0.272 * ,b</td>
</tr>
<tr>
<td>DZN15 + SMAE100</td>
<td>3.24 ± 0.403 * ,b</td>
<td>5.23 ± 0.297 * ,b</td>
</tr>
<tr>
<td>DZN15 + SMAE100 + LC300</td>
<td>3.47 ± 0.265 b</td>
<td>6.67 ± 0.310 b</td>
</tr>
</tbody>
</table>

Data is shown as mean ± SD.
*a*, p < 0.05 as compared to the control and sham groups
*b*, p < 0.05 as compared to the DZN15 group
Discussion

In the present experimental study, the effects of LC, SMAE and DZN toxin on the serum levels of hepatic enzymes (ALP, ALT, AST and GGT), Alb and TP in adult male rats were investigated. Since these hepatic enzymes are intracellular, and enter bloodstream in cases where cell damage occurs, it can be concluded that DZN is able to damage hepatocytes.

In the present study, the elevated level of ALP in the DZN15 group is possibly the result of the cholestasis, and the increase in the levels of ALT and AST in this group is likely to result from liver cell necrosis. Similarly, it has been shown that the intake and absorption of DZN by the liver increases the serum levels of ALT, ALP and AST in mice [21]. Also, in their experiments on rabbits, Solati et al., reported that DZN increases activities of ALT, ALP and AST [22], which is consistent with the results of this study. Gökçimen et al., reported that DZN caused pathological changes including hepatocyte necrosis and infiltration of inflammatory cells due to the activity of detoxification of the liver and the production of high levels of free radicals in the liver cells in rats [23]. OP damage the cells and tissues of the body by increasing lipid peroxidation, cell apoptosis and free radical production as well as inhibition of the activity of antioxidant enzymes such as superoxide dismutase, glutathione peroxidase and catalase [24,25].

In the SMAE100, LC300 and SMAE100+LC300 groups, administrating SMAE and LC alone as well as co-administration of SMAE+LC did not make a significant difference in the serum levels of AST, ALT, ALP, GGT, Alb and TP in comparison with the control and sham groups. However, in all three groups compared to the DZN group, the serum levels of liver enzymes decreased significantly, while the serum levels of Alb and TP increased significantly. In accordance with the results of biochemical parameters, histopathologic study in all three groups indicated that the liver tissue structure was completely normal and similar to the control and sham groups. It has been reported that the use of natural antioxidant compounds can play an important role in improving and decreasing oxidative stress damages. The safety of synthetic antioxidants has always been questioned, therefore, using natural

![Figure 1](https://example.com/figure1.png)

**Figure 1**
Histopathological analysis of liver in different groups. H&E 100× (A-I). A) Control, B) Sham, C) DZN15, D) SMAE100, E) LC300, F) SMAE100+LC300, G) DZN15+LC300, H) DZN15+SMAE100, I) DZN15+SMAE100+LC300.
It can be argued that SMAE prevents the progression of liver damage and inhibits liver cirrhosis by its numerous properties such as antioxidant, anti-lipid peroxidase, anti-fibrotic, anti-inflammatory, immune-regulation and liver cell regeneration. The significantly decreased AST and ALT levels in the DZN15+SMAE100 group confirms that the compounds present in SMAE can protect outer membrane of the hepatocytes, preventing penetration of toxic substances as well as increasing the ribosomal protein synthesis and thus, improving liver function [34].

In an in vitro study, Gulchin showed that LC has an antioxidant effect against free radicals of superoxide and hydrogen peroxide [35]. Hence, as an antioxidant compound capable of destroying free radicals, LC is able to reduce the damage caused by DZN in the liver tissue.

According to our results, the serum levels of Alb and TP significantly declined in the DZN15 groups. Alb is one of the most important circulating proteins that accounts for more than half of the plasma protein. It is made by the liver and secreted in the bloodstream. Blood Alb level is a sensitive and valuable parameter of liver function and its decline is a sign of liver failure [36]. A recent study by Yehia et al., on rabbits poisoned with DZN reports a decrease in Alb and TP levels; this decline was considered to be the result of an increase in proteolytic activity due to post-poisoning stress [37].

Since the levels of Alb and TP significantly declined in the DZN15 group while their levels increased significantly in the groups receiving mixture of DZN, LC and or SMAE, it can be argued that DZN toxin damages hepatic tissues leading to failure in producing enough Alb, but when the SMAE and LC are used (especially in the DZN15+SMAE100+LC300 group) the liver functions are restored. In other words, the concomitant use of the toxin, extract or the drug may have effective protective influences on the liver cells against the damage caused by DZN. These beneficial effects may essentially be the result of antioxidant properties of phenolic compounds present in SM, which allows them to act as reducing agents (hydrogen donor and oxygen inactivator). In general, the results of this study showed that the combination of SMAE and LC can have beneficial effects on restoring serum levels of hepatic enzymes and proteins like Alb in rats intoxicated by DZN. This combination may have antioxidant properties capable of eliminating free radicals leading to a reduction in DZN-induced hepatic injuries that provides a suitable therapeutic approach to use in such toxin poisoning.
Material and methods

Animals

Seventy-two adult male Wistar rats weighing 220 ± 20 g and 2 months old were provided from the animal house of Kazerun Islamic Azad University and were kept at the same place. The animals were supplied with water and food ad libitum. During the study, animals were exposed to standard conditions at 22 ± 2 °C, 12-hour light and 12-hour darkness cycles and 70% humidity in polycarbonate cages. The protocol of this study was approved by the Ethics Committee of Islamic Azad University of Kazerun, Iran, in relation to working with laboratory animal care (No. IR.Kiau 15230599971001).

Medications

LC was purchased from Merck Company (Germany) and technical DZN with 95 percent purity was obtained from Sam Gol Company (Iran). Normal saline was used to prepare LC solution.

Preparing SMAE

The aerial part of the SM contains the stem and seeds dried and then powdered. 100 g of the powder was added to 500 ml of distilled water and mixed well, and then kept at room temperature for 24 hours. The mixture was mixed using magnetic heating stirrer at 60 °C for 1 hour. The extract was centrifuged for 20 minutes at 10,000 rpm and then filtered. The extract was stored in the refrigerator until it was used [38].

Study design

The animals were assigned in 9 groups of 8 each including: the control group (untreated and only received water and food), the sham group (received normal food, water and 1 ml distilled water as the drug solvent), the DZN group (received 15 mg/kg DZN), the SMAE100 group (received 100 mg/kg SM aqueous extract), the LC300 group (received 300 mg/kg LC), the SMAE100+LC300 group (received 100 mg/kg SM aqueous extract and 300 mg/kg LC in the afternoon), the DZN+LC300 group (received 15 mg/kg DZN in the morning and 300 mg/kg LC in the afternoon), the DZN+SMAE100 group (received 15 mg/kg DZN in the morning and 100 mg/kg SM aqueous extract), and the DZN+SMAE100+LC300 group (received 15 mg/kg DZN in the morning and 100 mg/kg SM aqueous extract as well as 300 mg/kg LC in the afternoon). The extract and the drugs were administered in all groups by oral gavage for 30 days. The acute oral LD50 (the dose to kill half the population of laboratory animals) of DZN have been determined to be 1250 mg/kg body weight for the rats [39,40]. The basis for selecting the dose of DZN and LC was based on the previous studies (40,41).

Biochemical analysis

At the end of the treatment period, animals were anesthetized with diethyl ether (Merck, Germany), and blood samples were taken from their left heart ventricle using a 5 ml sterile syringe. Blood samples were transferred into the test tubes without anticoagulant, and were centrifugated at 5000 rpm for 15 minutes. The supernatant (serum) was isolated by sampler, transferred to new tubes, covered by Parafilm, and stored at -20 °C to be tested later. The serum levels of AST, ALT, ALP, GGT, Alb and TP were evaluated spectrophotometrically (Pars kits, Pars Co., Iran) using the Technicon RA-1000 machine (USA).

Histopathological analysis

After blood sampling, and euthanasia, the animal’s abdominal cavity was opened and the livers of all rats were removed. The tissue samples were washed with physiological normal saline and fixed in 10% formalin buffer solution. After the dehydration in alcohol with increasing concentrations of 60% to absolute, they became transparent using xylene and then were blocked in paraffin. 10 sections of 5-micron thickness of each liver tissue were stained with hematoxylin and eosin using a light microscope. In each section, five different areas were studied. A total of 10 sections of 5-micron thickness of each liver tissue were stained with hematoxylin and eosin using a light microscope. In each section, five different areas were studied. The liver tissues in each individual were evaluated for the following pathological features of apoptosis, congestion, sinusoidal spaces, and vacuolization of cytoplasm were studied.

Statistical analysis

The results were analyzed using SPSS software version 20 (SPSS Inc., Chicago, IL, USA). The results normalization were confirmed using the Kolmogorov-Smirnov test. The significance level was defined as $p < 0.05$, and was applied to examine the significant difference between the experimental and control groups by one-way ANOVA and post-hoc Tukey’s tests. The results are shown as mean ± standard deviation (SD) in the tables.

Acknowledgment

We thank Department of Biology, Kazerun Branch of Islamic Azad University for providing the facilities of physiological laboratory.

Author Contributions

Designed the experiments, performed statistical analysis and revised the manuscript: M.S., M.M. Performed histopathological analysis and wrote the manuscript: F.M. All authors approved the final version of the manuscript.

Conflict of Interest

None declared.

References

RESEARCH ARTICLE


مقایسه اثرات محافظتی ال کارنیتین و عصاره آبی خار مریم پس از سمیت کبدی القا شده با دیازینون در کبد موش صحرایی نر

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گروه زیست شناسی، واحد کازرون، دانشگاه آزاد اسلامی، کازرون، ایران

چکیده
دیازینون (DZN) یک کت کش ارگانوفسفور (OP) یا ایجاد استرس اکسیدتیو منجر به تولید رادیکال های آزاد و تغییرات پاتولوژیک در بدن است. هدف این مطالعه بررسی اثرات محافظتی ال کارنیتین (LC) و عصاره آبی خار مریم (SMAE) در برابر سمیت کبدی (DZN15 + SMAE100 + LC300) یا کنترل (n=8) شامل کنترل و SMAE100 در کبد موش صحرایی نر است. موش ها در ۹ روز، برای دیازینون (DZN) اقدام کردند. نمونه‌های خون در هفت روز بعد از درمان تحت گاواش دهانی گرفته شد. سطح سرمی آسپارتات آمینو ترانسفراز (AST)، آلانین آمی نو ترانسفراز (ALT)، گاما گلوتامیل ترانسفراز (GGT) و آلکالین فسفاتاز (ALP) با استفاده از روش فوتومتریک اندازه‌گیری شدند. همچنین، کبد های دیازینون (DZN15) تثبیت گردید و مورد ارزیابی هیستوپاتولوژیک قرار گرفتند. درمان با لفنوسیتی، سرای خون، آپوپتوسیتی، در سطح AC300 با اندازه‌گیری بالا حرکت می‌کرد و میزان قابل توجهی سطح SMAE100 با اندازه‌گیری بالا حرکت می‌کرد. افزایش داد (5.05 < ρ< 0.05) و تغییر نداد (5.05 < ρ< 0.05) با ایجاد استرس اکسیدتیو منجر به تولید رادیکال های آزاد و تغییرات پاتولوژیک در بدن است.

واژگان کلیدی
دیازینون، آل کارنیتین، خار مریم، سمیت کبدی، موش صحرایی