ABSTRACT

INTRODUCTION: Cypermethrin toxicity has been a subject of extensive studies because of its world widespread distribution. The major source of cypermethrin in the human body is through intake of food and water. Aim of the Work: This work was performed to study the protective role of garlic extract and vitamins C on the toxic effects of cypermethrin on the liver and kidneys of adult albino rats. Materials and Methods: This study was carried out on 126 adult albino rats for 12 weeks and gavaged 6 days / weeks. The rats were divided into 7 groups each of 18 rats; Group I (control group), Group II (garlic extract group): each rat was gavaged orally with 500 mg/kg vitamin C dissolved in (1 ml) distilled water once daily and Group III (vitamin C group): each rat was gavaged orally with 20 mg/kg vitamin C dissolved in (1 ml) distilled water once daily. Group IV (cypermethrin group): Each rat was gavaged orally with 32.5 mg/kg in (1 ml) distilled water once daily. Group V (Cypermethrin and garlic extract group): Each rat was gavaged orally with in the previous dose once daily. Group VI (Cypermethrin and vitamin C group): Each rat was gavaged orally with cypermethrin and vitamin C in the previous doses once daily. Group VII (Cypermethrin, vitamin C and garlic extract group): Each rat was gavaged orally with cypermethrin, garlic extract and vitamin C in the previous doses. Results: there was a significant increase in the mean values of AST, ALT, ALP, urea and creatinine in cypermethrin treated group when compared with control group (I). Upon supplementation with garlic extract or vitamin C alone or combined to cypermethrin treated rats there was a significant decrease in the mean values of AST, ALT, ALP, urea and creatinine, when compared with cypermethrin treated group. Microscopic examination of cypermethrin treated groups of liver revealed vaculated hepatocytes, focal necrosis, and aggregates of inflammatory cells, dilated congested central vein, enlarged portal tract with dilated congested blood vessels and bile duct proliferation, areas of hemorrhage and disturbed hepatic lobular architecture. Microscopic examination of cypermethrin treated groups of kidneys revealed congestion and hemorrhage; dilated tortuous tubules lined by vaculated epithelium and destructed renal corpuscles with wide Bowman's space. Immunohistochemical examination for detection of BAX protein in hepatic and renal tissues showed strong positive brown reaction in cypermethrin treated group when compared with control group (I) showed negative reaction. Upon supplementation with garlic extract or vitamin C alone or combined to cypermethrin treated rats there was a significant improvement with moderate to faint brown reaction respectively. Conclusion: Cypermethrin toxicity induced liver and kidneys damage in adult albino rats, and concomitant administration of either garlic extract or vitamin C or both was efficient in protecting liver and renal tissues.

Recommendation: it is recommended to increase public awareness regarding the health impact of cypermethrin and the protective role of garlic extract or vitamins C on its toxicities.

Key Words: Cypermethrin; liver; kidneys; hepatotoxicity; nephrotoxicity; oxidative stress; garlic extract and vitamin C.

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Mobile: 01141452918

INTRODUCTION

Pesticide poisoning is an important cause of morbidity and mortality in developing countries. Every year there are three million cases of severe poisoning and 220,000 deaths. The toxicity of pyrethroid insecticides to mammalian animals has received much attention in recent years because they have a widespread use in agriculture, domestic and veterinary applications and also the animals exposed to these insecticides exhibit changes in their physiologic activities beside other pathologic changes (Abdel-Rahim et al., 2009).
Synthetic pyrethroids comprise more than 1000 powerful broad spectrum insecticides and they are environmentally compatible due to their moderate persistence, low volatility and poor aqueous mobility in soil. They represent approximately one fourth of the worldwide insecticide market due to their high efficacy and enhanced stability. Cypermethrin belongs to the group of pyrethroids, classified by the world health organization as class II, moderately hazardous insecticides (Rodriguez et al., 2009).

Cypermethrin is the most widely used type II synthetic pyrethroid pesticide. Consistent with its lipophilic nature, cypermethrin has been accumulated in body fat, skin, brain, liver, kidneys, adrenals, testicles and ovaries. The cypermethrin toxicity is due to interaction with sodium channels in the nerve cells, making these channels in open configuration, thus generating repeated nerve impulses in the affected organs (Kakko et al., 2008).

Chemoprevention against oxidative stress can be obtained by the administration of one or more chemical entities, either as individual drugs or as naturally occurring dietary constituents such as vitamin C and garlic extract providing anti-oxidant cellular protection against cellular membrane oxidative toxicity derived from cypermethrin administration (Wilson, 2004).

The aim of this study was to demonstrate some changes in livers and kidneys of adult albino rats intoxicated sub chronically with cypermethrin and to evaluate the possible protective role of garlic extract and vitamin C supplementation against these changes.

1. **Material:**

1) **Cypermethrin:** It was purchased from Sigma Chemical Co. (St Louis, Mo, USA). It was provided in a white powder form.

2) **Garlic extract:** It was purchased from SEKOM Herbal Company marketed as yellowish powder.

3) **Vitamin C:** In the form of white powder manufactured by EPICO pharmaceutical company, Egypt.

4) **Distilled water:** All the previous material were dissolved in this distilled water.

5) **Detection kit:**

The primary antibodies for the specific localization of BAX receptors in frozen sections, as well as, in formalin fixed and paraffin embedded tissue. They were delivered from Medco Pharma Trade Egypt (Ab-4(6A7), Code no. (Ms-714-Po).

6) **Animals:**

This study was carried on 126 adult albino rats, each weighing 150-200 gm. The animals were obtained from Animal House of the Faculty of Veterinary Medicine, Zagazig University. The study had been designed in the Faculty of Medicine, Zagazig University.

**Study design:**

The study was done or 3 months and the rats were divided into 7 groups each of 18 rats:-

- **Group I (control group):** Each rat received only regular diet and distilled water to determine the basic values of performance. These rats were left without intervention to measure the basic parameters.

- **Group II (Garlic extract (G) group):** Each rat was gavaged orally with 500 mg/kg body weight garlic extract dissolved in 1ml of distilled water once daily 6 days/ week (Singh et al., 2008).

- **Group III (Vitamin C (C) group):** Each rat was gavaged orally with 20 mg/kg body weight vitamin C dissolved in 1 ml of distilled water once daily 6 days/ week (Khan and Sinha, 2008).

- **Group IV (cypermethrin (CYP) group):** Each rat was gavaged orally with 32.5 mg/kg body weight (1/20 of LD50) of cypermethrin dissolved in 1ml of distilled water once daily 6 days/ week. Grayson (2012) reported Ld50 650mg/kg.

- **Group V (Cypermethrin and garlic extract (CYP-G) group):** Each rat was gavaged orally with (500mg/kg body weight garlic extract) 15min before (32.5 mg/kg body weight cypermethrin) once daily 6 days/ week.

- **Group VI (Cypermethrin and vitamin C (CYP-C) group):** Each rat was gavaged orally with (20 mg/kg body weight vitamin C 15 min
before oral gavage with (32.5 mg/kg body weight cypermethrin) once daily 6 days/ week.

-Group VII (Cypermethrin, vitamin C and garlic extract (CYP-G-C) group): Each rat was gavaged orally with (20 mg/kg body weight vitamin C+ 500 mg/kg body weight garlic extract 15 min before oral gavage with + 32.5 mg/kg body weight cypermethrin) once daily 6 days /week.

II- Methods:

At the end of 12th week of the study period, rats from each group were subjected to blood sample collection from the retro-orbital plexuses. The blood samples will be used for estimating the following tests Liver function tests {serum aspartame aminotransferase (AST), serum alanine aminotransferase (ALT) and alkaline phosphatase (ALP)}, kidney function tests (urea and creatinine). Then the anesthetized rats were sacrificed & specimens from the liver and kidneys were taken for histopathological study.

(A) Liver enzymes:
Serum Alanine aminotransferase (ALT), Serum Aspartate aminotransferase (AST) and Serum Alkaline phosphatase (ALP) (IU/L): These enzymes were assayed as (Reitman and Frankel, 1957).

(B) Kidney functions:
(i) Blood Urea Nitrogen (BUN) (mg/dL): (Kaplan, 1965).
(ii) Serum creatinine (mg/dL): (Popper et al., 1937).

(C) Histopathological study by H & E:
The liver and kidneys specimens were routinely processed and sectioned at 4-5 Um thickness. The obtained sections were stained by Haematoxylin and Eosin (H&E) according to Horobin and Bancroft (1998) then examined by light microscope.

(D) Immunohistochemical staining:
A mouse monoclonal antibody of IgG immunoglobulin type, designed for the specific localization of BAX receptors in frozen sections, as well as, in formalin fixed and paraffin embedded tissue. They were delivered from Medco Pharma Trade Egypt (Ab-4(6A7), Code no. (Ms-714-Po). The sensitive universal immunostaining kit contained the following reagent and material (Shimizu et al., 2014).

(E) Statistical analysis:
The results were analyzed by ANOVA-test: for comparison of means of different groups (Dean et al., 2004).

Results:
(A) Biochemical results:
The biochemical findings of control groups were within normal values, in addition there were non significant difference in the levels of all tested biochemical parameters between control group (I), garlic extract group (II) and vitamin C group (III), so we used group (I) for comparison with other treated groups (Tables 1 & 6).

There were highly significant differences in groups by (ANOVA) study (P<0.001) as regard (alanine aminotransfersae (ALT), aspartate aminotransferase (AST) and Alkaline phosphatase (ALP) values (Table 2).

The least significant difference (LSD) of (ALT, AST & ALP) among groups I, CYP (IV), G + CYP (V), C + CYP (VI) and G + C+CYP (VII) revealed that there was highly significant elevation in (ALT, AST & ALP) level in all groups when compared with control group (I) (P<0.001). There was highly significant elevation in ALT in group CYP (IV) when compared with other groups of the study (P<0.001). There was no significant difference in (ALT, AST & ALP) value between groups G+ CYP(V) and C + CYP (VI) groups G+ C+ CYP(VII) (P>0.05) (Table 3, 4, 5) (Fig. 1).

There were highly significant differences in groups by (ANOVA) study (P<0.05) as regard Blood urea nitrogen (BUN) and Serum creatinine (Creat) (Table 7).

The least significant difference (LSD) of blood urea nitrogen (BUN & Creat) among control group (I), CYP (IV), G + CYP (V), C + CYP (VI) and G + C+CYP (VII) revealed that there was highly significant elevation in (BUN & Creat) level in all groups when compared with control group (I) of the study (P<0.001). There was highly significant elevation in BUN in group CYP (IV) when
compared with other groups of the study (P<0.001). There was no significant difference in BUN value between groups G+ CYP(V) and C + CYP (VI) and G+ C+ CYP (VII) (P>0.05) (Table 8, 9) (Fig. 2,3).

(B) Histopathological study:
(1) Histopathological changes of the liver:
Light microscopic examination of H&E stained sections from the livers of the control groups (Groups I, II and III) revealed that; the liver showed multiple polygonal classic hepatic lobules with tightly packed cords of hepatocytes radiating from central veins toward the periphery of the lobules where the portal area is noticed. It contains the pre terminal branches of the portal vein, hepatic artery, and bile duct. Polygonal hepatocytes with rounded vesicular nuclei and acidophilic cytoplasm. Some cells are binucleated. Blood sinusoids and their lining of endothelial and kupffer cells are observed between hepatocyte cords (Fig. 4)

In cypermethrin treated group the sections showed disturbed hepatic lobular architecture with hepatocytes showed vacuolated cytoplasm, focal necrosis, aggregates of chronic inflammatory cells, dilated congested central vein, enlarged portal tract with dilated congested blood vessels, bile duct proliferation and areas of hemorrhage (Fig. 5).

Examination of H&E stained sections of the liver specimens of the rats treated with garlic extract or vitamin C alone showed partial improvement of the changes that occurred after cypermethrin administration with almost normal lobular architecture and normal hepatocytes with average sized nuclei (Fig. 6).

While microscopic examination of the liver specimens of the rats treated with combined garlic extract and vitamin C showed near complete improvement of the changes that occurred after cypermethrin administration with almost normal lobular architecture. Congested blood vessels still present (Fig. 7).

Immunohistochemistry examination for detection of BAX protein apoptotic markers that appears as a cytoplasmic reaction brown in color. The stained sections from the livers of the groups I, II and III showed no immunoreactivity within hepatocytes. Central vein (CV) and sinusoids (S) were noticed (Fig. 8).

In cypermethrin treated group the sections showed most hepatocytes with granular strong +ve brown reaction within cytoplasm (arrows) around central vein (CV) (Fig. 9).

Examination of sections of the liver specimens of the rats treated with garlic extract or vitamin C alone showd little restoration of the normal cellular structure. Most hepatocytes had granular nucleus and granular moderate brown reaction within cytoplasm (arrows) around central vein (CV) (Fig. 10).

While microscopic examination of the liver specimens of the rats treated with combined garlic extract and vitamin C showed restoration of the normal cellular structure. Most hepatocytes had granular nucleus and granular faint brown reaction within cytoplasm (arrows) around central vein (CV) (Fig. 11).

(2) Histopathological changes of the kidneys:
Light microscopic examination of H&E stained sections from the kidneys of the control groups (Groups I, II and III) revealed that; the kidneys showed normal cortex, medulla, interstitial tissue and blood vessels allover the period of the study. The cortex is formed of malpigian renal corpuscles which are formed of tuft of capillaries (Glomerulus) surrounded by Bowman's capsule. Proximal and distal tubules were normal; medulla was mostly formed of loop of Henle and collecting tubules (Fig. 12).

In cypermethrin treated group the sections showed congestion and hemorrhage, dilated tortuous tubules lined by vaculated epithelium and destructed real corpuscles with wide Bowman's space (Fig. 13).

Examination of sections of the liver specimens of the rats treated with garlic extract or vitamin C alone showed partial improvement of the changes that occurred after cypermethrin administration with showed minor congestion, dilated tortuous tubules lined by vaculated epithelium, and shrinkage real
corpuscles with wide Bowman's space (Fig. 14).

While microscopic examination of the kidney specimens of the rats treated with combined garlic extract and vitamin C showed near complete improvement of the changes that occurred after cypermethrin administration almost normal glomeruli. Congested blood vessels still present. It showed fewer distorted Glomeruli and Bowman's capsule, and fewer vacuolar spaces (Fig. 15).

Immunohistochemistry examination for detection of BAX protein apoptotic markers revealed that stained sections from the kidneys of the control groups (Groups I, II and III) showed no immunoreactivity within nephrocytes. Glomeruli (G) and tubules (T) are noticed (Fig. 16).

In cypermethrin treated group the sections showed a nephrocyte showing granular strong +ve brown reaction within cytoplasm of most nephrocytes (arrows) glomeruli (G) and tubules (T) are noticed (Fig. 17).

Examination of sections of the kidney specimens of the rats treated with garlic extract or vitamin C alone showing little restoration of the normal cellular structure. The nephrocyte has granular nucleus and granular moderate brown reaction within cytoplasm of most nephrocytes (arrows) around glomeruli (G) and tubules (T) are noticed (Fig. 18).

While microscopic examination of the kidney specimens of the rats treated with combined garlic extract and vitamin C showing restoration of the normal cellular structure. The nephrocyte has granular nucleus and granular faint brown reaction within cytoplasm of most nephrocytes (arrows) around glomeruli (G) and tubules (T) are noticed (Fig. 19).

### Table (I): A statistical comparison between control group “I”, (G) group “II” and (C) group “III” as regard mean values of liver enzymes (ALT, AST and ALP) after 12 weeks in adult albino rats by ANOVA test.

<table>
<thead>
<tr>
<th></th>
<th>Control (I)</th>
<th>Garlic extract (II)</th>
<th>Vitamin C (III)</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean ± SD 12 weeks</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALT (U/L)</td>
<td>24.5 ± 1.5</td>
<td>24 ± 3.4</td>
<td>29 ± 5.3</td>
<td>3.25</td>
<td>0.067#</td>
</tr>
<tr>
<td>AST (U/L)</td>
<td>43.3 ± 7.9</td>
<td>44.1 ± 5.8</td>
<td>40 ± 6.3</td>
<td>0.626</td>
<td>0.548#</td>
</tr>
<tr>
<td>ALP (U/L)</td>
<td>60 ± 4.1</td>
<td>57 ± 4</td>
<td>55.7 ± 3.3</td>
<td>2</td>
<td>0.169#</td>
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</table>

SD = Standard Deviation.
# p>0.05 = non significant
Table (2): A statistical comparison between control group “I” and treated group CYP “IV”, CYP+G “V”, CYP+C “VI”, CYP+G+C “VII” as regard mean values of liver enzymes (ALT, AST and ALP) after 12 weeks adult albino rats along the period of the study by ANOVA test.

<table>
<thead>
<tr>
<th></th>
<th>Control (I)</th>
<th>CYP (IV)</th>
<th>CYP-G (V)</th>
<th>CYP-C (VI)</th>
<th>CYP-G-C (VII)</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>ALT (U/L)</td>
<td>24.5 ± 1.5</td>
<td>97.33 ± 6.71</td>
<td>30.50 ± 6.15</td>
<td>26.00 ± ±</td>
<td>23.83 ± ±</td>
<td>187.91</td>
<td>&lt;0.001**</td>
</tr>
<tr>
<td>AST (U/L)</td>
<td>43.3 ± 7.9</td>
<td>145.50 ± 32.52</td>
<td>47.00 ± 4.00</td>
<td>55.66 ± ±</td>
<td>40.66 ± ±</td>
<td>30.60</td>
<td>&lt;0.001**</td>
</tr>
<tr>
<td>ALP (U/L)</td>
<td>60 ± 4.1</td>
<td>586.66 ± 83.10</td>
<td>74.16 ± 14.74</td>
<td>75.66 ± ±</td>
<td>60.50 ± ±</td>
<td>175.84</td>
<td>&lt;0.001**</td>
</tr>
</tbody>
</table>

SD = Standard Deviation.  
G= garlic extract.  
** = highly significant.  
CYP= cypermethrin.

Table (3): Least significance difference (LSD) among the control (I), (CYP) IV, (CYP+ G) V, (CYP+ C) VI and (CYP+ G+ C) VII groups as regard alanine aminotransferase enzyme ALT values after 12 weeks of the study.

<table>
<thead>
<tr>
<th>ALT (IU/L)</th>
<th>I</th>
<th>(CYP) IV</th>
<th>V(CYP–G)</th>
<th>VI(CYP-C)</th>
<th>VII(CYP–G-C)</th>
<th>Mean ± SD 12 weeks</th>
</tr>
</thead>
<tbody>
<tr>
<td>24.5 ± 1.5</td>
<td>97.33 ± 6.71</td>
<td>30.50 ± 6.15</td>
<td>26.00 ± ±</td>
<td>± ± 23.83 ± ±</td>
<td>187.91</td>
<td>&lt;0.001**</td>
</tr>
<tr>
<td>0.00**</td>
<td>0.43#</td>
<td>0.14#</td>
<td>0.79#</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IV</td>
<td>0.00**</td>
<td>0.00**</td>
<td>0.00**</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V</td>
<td>0.11#</td>
<td>0.08#</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VI</td>
<td>0.40#</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

** = highly significant (P<0.001).  
# = Non–significance (P>0.05).  
CYP= cypermethrin.  
G= garlic extract.  
C= vitamin C.
Table (4): Least significance difference (LSD) among the control (I), (CYP) IV, (CYP+ G) V, (CYP+ C) VI and (CYP+ G+ C) VII groups as regard aspartate aminotransferase enzyme AST values after 12 weeks of the study.

<table>
<thead>
<tr>
<th>AST (IU/L)</th>
<th>I (CYP) IV</th>
<th>V(CYP–G)</th>
<th>VI(CYP–C)</th>
<th>VII(CYP–G–C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± SD 12 weeks</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>43.3 ± 7.9</td>
<td>145.50 ± 32.52</td>
<td>47.00 ± 4.00</td>
<td>55.66 ± 8.84</td>
</tr>
<tr>
<td>I</td>
<td>0.00**</td>
<td>0.33#</td>
<td>0.29#</td>
<td>0.61#</td>
</tr>
<tr>
<td>IV</td>
<td>0.00**</td>
<td>0.00**</td>
<td></td>
<td>0.00**</td>
</tr>
<tr>
<td>V</td>
<td>0.054#</td>
<td></td>
<td>0.17#</td>
<td></td>
</tr>
<tr>
<td>VI</td>
<td></td>
<td></td>
<td>0.19#</td>
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</table>

** = highly significant (P<0.001).
#= Non–significance (P>0.05).
CYP= cypermethrin.
G= garlic extract.
C= vitamin C.

Table (5): Least significance difference (LSD) among the control (I), (CYP) IV, (CYP+ G) V, (CYP+ C) VI and (CYP+ G+ C) VII groups as regard alkaline phosphatase enzyme ALP values after 12 weeks of the study.

<table>
<thead>
<tr>
<th>ALP (IU/L)</th>
<th>I (CYP) IV</th>
<th>V(CYP–G)</th>
<th>VI(CYP–C)</th>
<th>VII(CYP–G–C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± SD 12 weeks</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>60 ± 4.1</td>
<td>586.66 ± 83.10</td>
<td>74.16 ± 14.74</td>
<td>75.66 ± 8.82</td>
</tr>
<tr>
<td>I</td>
<td>0.00**</td>
<td>0.47#</td>
<td>0.30#</td>
<td>0.50#</td>
</tr>
<tr>
<td>IV</td>
<td>0.00**</td>
<td>0.00**</td>
<td></td>
<td>0.00**</td>
</tr>
<tr>
<td>V</td>
<td></td>
<td>0.83#</td>
<td>0.08#</td>
<td></td>
</tr>
<tr>
<td>VI</td>
<td></td>
<td></td>
<td>0.16#</td>
<td></td>
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</table>

** = highly significant (P<0.001).
#= Non–significance (P>0.05).
CYP= cypermethrin.
G = garlic extract.
C = vitamin C.

Fig. (1): Bar chart showing comparative magnitude of mean values of liver enzymes (ALT, AST & ALP) at different groups control group (I), cypermethrin treated group (IV), garlic extract & cypermethrin group (V), vitamin C & cypermethrin group (VI) and garlic extract & vitamin C and cypermethrin group (VII) after 12 weeks of the study.

Table (6): A statistical comparison between control group “I”, G group “II” and C group III as regard mean values of kidney function (BUN & serum creatinine) after 12 weeks in adult albino rats by ANOVA test.

<table>
<thead>
<tr>
<th></th>
<th>Control (I)</th>
<th>Garlic extract (II)</th>
<th>Vitamin C (III)</th>
<th>F</th>
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<tr>
<td></td>
<td>Mean ± SD</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>12 weeks</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BUN (mg/dl)</td>
<td>16.5 ± 1.5</td>
<td>16.1 ± 1.4</td>
<td>16 ± 1.04</td>
<td>0.238</td>
<td>0.791#</td>
</tr>
<tr>
<td>serum creatinine</td>
<td>0.43 ± 1.77</td>
<td>0.46 ± 1.9</td>
<td>0.47 ± 1.01</td>
<td>0.001</td>
<td>0.999#</td>
</tr>
</tbody>
</table>

SD = Standard Deviation.
# p > 0.05 = non significant
Table (7): Statistical comparison among the control (I), cypermethrin (IV/CYP), cypermethrin and garlic extract (V/ CYP-G), cypermethrin and vitamin C (VI/CYP- C) and cypermethrin, garlic extract and vitamin C (CYP- G- C) groups as regard blood urea nitrogen and serum creatinine levels along the periods of the study by ANOVA test.

<table>
<thead>
<tr>
<th></th>
<th>Control (I)</th>
<th>CYP (IV)</th>
<th>CYP-G (V)</th>
<th>CYP-C (VI)</th>
<th>CYP-G-C (VII)</th>
<th>F</th>
<th>P</th>
</tr>
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<tbody>
<tr>
<td>BUN (mg/dl)</td>
<td>16.5 ± 1.5</td>
<td>68.16 ± 9.53</td>
<td>17.33 ± 3.07</td>
<td>29.00 ± 5.09</td>
<td>26.50 ± 1.76</td>
<td>91.11</td>
<td>&lt;0.001**</td>
</tr>
<tr>
<td>Serum creatinine (mg/dl)</td>
<td>0.43 ± 1.77</td>
<td>0.70 ± 3.12</td>
<td>0.31 ± 0.07</td>
<td>0.23 ± 1.08</td>
<td>0.18 ± 1.07</td>
<td>39.96</td>
<td>&lt;0.001**</td>
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</table>

SD = Standard Deviation.
** = highly significant.
CYP= cypermethrin.
G= garlic extract.
C= vitamin C.

Table (8): Least significance difference (LSD) among the control (I), (CYP) IV, (CYP- G) V, (CYP-C) VI and (CYP-G - C) VII groups as regard BUN values after 12 weeks of the study.

<table>
<thead>
<tr>
<th>BUN (mg/dL)</th>
<th>I</th>
<th>I (CYP) IV</th>
<th>V(CYP–G)</th>
<th>VI(CYP-C)</th>
<th>VII(CYP–G-C)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Mean ± SD 12 weeks</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>16.5 ± 1.5</td>
<td>68.16</td>
<td>17.33 ± 3.07</td>
<td>29.00 ± 5.09</td>
<td>26.50 ± 1.76</td>
<td></td>
</tr>
<tr>
<td>0.00**</td>
<td>0.56#</td>
<td>0.27#</td>
<td>0.31#</td>
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** = highly significant (P<0.001)
# = Non–significance (P>0.05).
CYP= cypermethrin.
G= garlic extract.
C= vitamin C.
Table (9): Least significance difference (LSD) among the control (I), (CYP) IV, (CYP-G) V, (CYP-C) VI and (CYP-G-C) VII groups as regard serum creatinine values after 12 weeks of the study.

<table>
<thead>
<tr>
<th>Serum creatinine (mg/dL)</th>
<th>I (CYP) IV</th>
<th>V(CYP–G)</th>
<th>VI(CYP-C)</th>
<th>VII(CYP–G–C)</th>
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<tbody>
<tr>
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<td>Mean ± SD 12 weeks</td>
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<td></td>
<td>0.43 ± 1.77</td>
<td>0.70 ± 3.12</td>
<td>0.31 ± 0.07</td>
<td>0.23 ± 1.08</td>
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</tbody>
</table>

** = highly significant (P<0.001)
# = Non–significance (P>0.05)

CYP= cypermethrin.
G= garlic extract.
C= vitamin C.

Fig. (2): Bar chart showing comparative magnitude of mean values of kidney functions (BUN) at different groups in control group (I), cypermethrin treated group (IV), garlic extract & cypermethrin group (V), vitamin C & cypermethrin group (VI) and garlic extract & vitamin C and cypermethrin group (VII) after 12 weeks of the study.
The Evaluation Of Protective Role Of Garlic

Fig. (3): Bar chart showing comparative magnitude of mean values of kidney functions (serum creatinine) at different groups in control group (I), cypermethrin treated group (IV), garlic extract & cypermethrin group (V), vitamin C & cypermethrin group (VI) and garlic extract & vitamin C and cypermethrin group (VII) after 12 weeks of the study.

Fig (4): A photomicrograph of a section of liver of control adult albino rat showing a hexagonal hepatic lobule, consisting of cords of hepatocytes radiating from the central vein (CV) toward portal area (PA) kupffer cells (arrows) (H&E X 200).

Fig (5): A photomicrograph of a section of liver from adult albino rat orally gavaged by 32.5 mg/ kg cypermethrin daily for 12 weeks showing distorted liver archticher severely congested central vein (CV), mononuclear inflammatory cellular infiltration (I) of portal area, dilated congested portal vein (PV) and proliferated bile ducts (BD) with dilated sinusoids (S) (H&E X 200).
Fig (6): A photomicrograph of a section of liver from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin followed by 20 mg/kg vitamin C daily for 12 weeks showing pericentral hepatocytes radiating from the central vein (CV) with vesicular nuclei (arrow) and granular cytoplasm. Few hepatocytes (V) are still vacuolated. Blood sinusoids (S) are also noticed (H&E X 400).

Fig (7): A photomicrograph of a section of liver from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin followed by 500mg/kg garlic extract and 20 mg/kg vitamin C daily for 12 weeks showing pericentral hepatocytes with vesicular nuclei (arrow) and granular cytoplasm radiating from dilated central vein (CV). Blood sinusoids (S) are also noticed (H&E X 400).

Fig (8): A photomicrograph of a section of liver of negative control adult albino rat showing no immunoreactivity within hepatocytes. Central vein (CV) and sinusoids (S) are noticed. (BAX immunostaining X 400).

Fig (9): A photomicrograph of a section of liver from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin daily for 12 weeks showing granular strong +ve brown reaction within cytoplasm of most hepatocytes (arrows) around central vein (CV)and sinusoids (S) are noticed (BAX immunostaining X 400).
Fig (10): A photomicrograph of a section of liver from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin followed by 20mg/kg vitamin C daily for 12 weeks showing granular faint +ve brown reaction within cytoplasm of most hepatocytes (arrows) around central vein (CV) and sinusoids (S) are noticed (BAX immunostaining X 400).

Fig (11): A photomicrograph of a section of liver from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin followed by 500mg/kg garlic extract and 20mg/kg vitamin C once daily for 12 weeks showing -ve reaction within cytoplasm of most hepatocytes around central vein (CV) (BAX immunostaining X 400).

Fig. (12): A photomicrograph of a section in the kidney of adult albino rat from the negative control group showing normal kidney structure, the glomeruli (G) and tubules (T) appears normal (H & E x 200).

Fig. (13): A photomicrograph of a section in the kidney of adult albino rat orally gavaged by 32.5 mg/kg cypermethrin daily for 12 weeks showing shrunken glomeruli (G) with wide Bowman's space (B), associated with dilated tortuous tubules (T) lined with vacuolated epithelium (V) with apoptotic nuclei (arrows) (H&E x 400.)
Fig. (14): A photomicrograph of a section of kidney from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin followed by 20mg/kg vitamin C same result for that administrated 500mg/kg garlic extract after cypermethrin daily for 12 weeks showing shrunken glomeruli (G) dilation of tubules (T) with vesicular nuclei (arrow) (H&E X 400).

Fig. (15): A photomicrograph of a section of kidney from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin followed by 500 mg/kg garlic extract and 20 mg/kg vitamin C daily for 12 weeks showing venous congestion (*) and slightly shrunken glomruli (G) with tubules are noticed (T) (H&E X 400).

Fig. (16): A photomicrograph of a section of kidney of negative control adult albino rat showing no immunoreactivity within glomerular (G) and tubular (T) cells (BAX immunostaining X 400).

Fig. (17): A photomicrograph of a section of kidney from adult albino rat orally gavaged by 32.5 mg/kg cypermethrin daily for 12 weeks showing granular strong +ve brown reaction within cytoplasm of most glomerular (G) and tubular (T) cells (arrows) (BAX immunostaining X 400).
DISCUSSION

The cypermethrin incorporation to humans takes place via drinking water, food, with a minor contribution from the air causing various health problems in man. It is easily degraded on soil and plants but can be effective for weeks when applied to indoor inert surfaces. Exposure to sunlight, water and oxygen will accelerate its decomposition. Cypermethrin is highly toxic to fish, bees and aquatic insects, according to the National Pesticides Telecommunications Network (NPTN) (NPTN, 2009).

(A) Hepatotoxicity:

In the present study, there was a significant increase in liver enzymes (AST, ALT and ALP) mean values in cypermethrin treated group as compared to their corresponding values in control group. These results confirmed by obvious histopathological changes in the general architecture of the liver stained with H&E associated with perportal inflammatory cells infiltration, blood vessels dilatation and congestion. The most striking histological feature was vacuolar degeneration in the cytoplasm of hepatocytes, pyknotic nuclei and infiltration with mononuclear leukocytes. Bile duct proliferation was observed. Also immunohistochemical staining for detection of BAX protein (apoptotic marker) in hepatic tissues appear as cytoplasmic strong brown reaction.

These results also report by Yanpallewar et al. (2003); Choudhary et al. (2003) and Nair et al. (2011) who stated that high serum concentrations of ALT, AST and ALP indicated cellular leakage due to disintegration of cell membrane in liver. The increase in transaminase activities is probably due to the cypermethrin-induced pathological changes in liver and is an indication of liver damage.

Further more, liver is a prime organ associated with xenobiotic metabolism (Hinton and Grasso 2000). Perhaps, production of metabolically toxic intermediates capable of
causing hepatocellular damage occurs during processing of cypermethrin in liver, causing respective leakage of these enzymes in blood (Bhushan et al., 2010).

The histopathological changes in the liver of rats exposed to cypermethrin obtained also by El-Toukhy and Girgis (2003) and Grewal et al. (2010a) in the form of necrotic areas in hepatocytes, cell swelling, cytoplasmic hypertrophy and intracytoplasmic vacuoles were seen.

These histopathological changes explained by that cypermethrin induced oxidative stress in various organs such as liver and kidneys (Grewal et al., 2010b).

Khan et al. (2009) who stated that, pathologic changes in hepatocytes due to cypermethrin can be related to its inhibitory effect on total adenine triphosphate activity in the hepatocytes disturbin g active transport of Na+, K+ and Ca2+ ions, thus injuring hepatocytes.

The cytoplasmic vacuolization that were observed in hepatocytes of cypermethrin treated animals is due to accumulation of ions and water in cytosol. Massive accumulation of fluids in the vacuoles may finally lead to cell lysis (Gores et al., 2000). These vacuoles are due to the free radicals that facilitate the release of lysosomal enzymes into the cytosol with subsequent oxidation of the protein architecture of the cells causing their fragmentation (Kulisz et al., 2002).

Azeez et al. (2011) and Jin et al. (2011) stated that the observed vascular dilatation may represent an adaptive process as an attempt to overcome oxygen deficiency. The cypermethrin exposure producing induction of hepatic oxidative stress is responsible for many deleterious effects in cell, including DNA damage and the alteration of gene expression related to apoptosis in adult albino rats.

The BAX protein suppresses the ability of Bcl-2, which enhances cell survival by inhibiting apoptosis induced under various circumstances (Oltvai et al., 1993).

The DNA contents increase in most of the tissues in response to cypermethrin exposure. This is due to activation of some dormant regulating factors controlling DNA synthesis. The drastic increase in the level of DNA in liver in response to the cypermethrin might be due to increased thymidine uptake in the hepatic DNA, as reported by Holbrook (2008).

In the present study, upon the administration of garlic with cypermethrin in group (V) and vitamin C with cypermethrin in group (VI) or together with cypermethrin in group (VII) result in improvement of the enzyme levels a significant improvement in histopathological changes; almost normal lobular architecture and normal hepatocytes with average sized nuclei except of few central veins and sinusoids that showed congestion. The immunohistochemistry examination showed hepatocyte with faint to negative immuno-reactivity when compared with cypermethrin treated group (IV)

These findings are reported by El-Banna et al. (2009) who stated that antioxidant garlic extract and vitamin C prevent the accumulation of toxins within the cell and promote the production of energy improvement of antioxidants enzymes levels with reduced Thiobarbituric acid-reactive substances (TBARS) levels which promotes the apoptosis.

However, hepa-toprotective effect of vitamin C tends to increase synergistically when co administered with other agents precisely antioxidants and the administration of garlic and vitamin C can modulate the oxidative stress and improve the antioxidant system through the direct cytoprotective effect of garlic extract constituents; diallyldisulphide, allyl propyl disulphide, allicin and diallyltriisulphide by modulating lipid peroxidation and enhancing antioxidant status in the liver and blood (Mirunalini et al., 2004; Pari et al., 2007; Hassan et al., 2010 and Assayed et al., 2010).

Free radicals, known to cause oxidative stress, can be prevented or reduced by dietary natural antioxidants through their capacity to
scavenge these products. Garlic extract or vitamin C may protect lipids and lipoproteins in cellular membranes against this oxidative damage, thus may prevent certain types of hepatic cellular damage (Aruoma, 2008).

Sushma and Devasena (2010) also reported increased oxidative stress (MDA, thiobarbituric acid reactive substances, or “oxidation index”) in erythrocytes, livers, or kidneys caused by cypermethrin exposure.

(B) nephrotoxicity:

In the present study, there was a significant increase in BUN and serum creatinine mean values in cypermethrin treated group as compared to their corresponding values in control group (I). This confirmed by obvious changes microscopic examination of the kidney shows hemorrhages in renal tubules, different stages of degeneration, cast deposition and increased urinary spaces were observed. Also immunohistochemical staining for detection of BAX protein (apoptotic marker) in renal tissues appear as cytoplasmic strong brown reaction.

The increased blood urea concentration in rats treated with cypermethrin are also obtained by Yousef et al. (2003) who recorded that cypermethrin induced increase in urea level observed may be due to its effect on liver function, as urea is the end product of protein catabolism and conversion of ammonia to urea because of increased synthesis of enzyme involved in urea production is more efficient.

An increase in serum creatinine was recorded in cypermethrin-treated rats by Sakr et al. (2001) and Abu-El-Zahab et al., (2003). Creatinine is more specific to kidneys since renal damage is the only significant parameter that increases serum creatinine in mammals (Garba et al., 2007). Like many other waste products of metabolism, most (> 75%) of the creatinine is removed from the body through glomerular filtration and the rest (<25%) through tubular secretion (Ravel, 2005). The increased serum concentrations of creatinine thus might be a result of alteration in these two mechanisms. Urea is also excreted by kidneys, so impaired kidney function causes diminished ability to excrete urea from the blood into urine (Aslam et al., 2010).


Due to various degenerative changes in the kidneys caused by cypermethrin, epithelial cells are detached from the tubular basement membranes. When detached epithelial cells are mixed with leaked proteins, the resultant mixture appears in the form of epithelial casts (Khan et al., 2009).

While, the pathological changes in kidney make sure that cypermethrin a hyperactivity and detoxication and the dilated cortical tubules due to the filling with protein casts, as the same with (Khan et al., 2003).

Sangha et al. (2011) reported these histological examination of kidney of cypermethrin -intoxicated birds revealed many alterations such as tubular degeneration, atrophy of glomeruli, hyperemia in glomeruli with focal area of necrosis and cytoplasmic vacuolation, cloudy swelling of the cells this is similar to (Ullah et al., 2006) in rabbits and (Grewal et al., 2010b) in mice. Cypermethrin induced oxidative stress, food and water intake of cypermethrin intoxicated changes in cell morphology and tissue injury (El-Demerdash, 2011) in animals. The oxidative stress indicates diminished ability of the kidneys to filter these waste products from the blood.

Braun et al. (2008) found that the kidney of cypermethrin treated adult albino rat showing apoptotic nucleus and +ve brown cytoplasmic reaction in cypermethrin treated rats. The protein level of the protective Bcl-2 was decreased, whereas BAX was increased, indicating that the protective activity of Bcl-2 might have been further reduced by up-regulation of the endogenous inhibitor (BAX). The family Bcl2 prevents apoptosis by inhibiting the generation or action of reactive oxygen species (Hockenberg et al., 2013).

The antioxidants combination of vitamin C and garlic extract prevent the dysregulation of Bcl-2 and BAX. Recent findings
demonstrate the activation of several proteases such as cysteine proteases and aspartic proteases in the apoptotic signal transduction pathway. The activation of one of these proteases possibly may also account for the decreased Bcl-2 protein levels. This study suggests that lipopolysaccharide induced apoptosis is mediated by oxidative stress and leads to a reduction of the endogenous protection potential of the cells by disturbance of the balance between Bcl-2 and BAX (Enari et al., 1996).

In the present study, upon the administration of garlic with cypermethrin in group (V) and vitamin C with cypermethrin in group (VI) or together with cypermethrin in group (VII) result in a significant decrease in the mean values of BUN and serum creatinine level a significant improvement in histopathological changes; negative immunoreactivity when compared with cypermethrin treated group (IV).

The use of natural antioxidants for curing pesticide induced kidney toxicity or injury is being studied extensively. It exhibited concentration-dependent inhibitory effects on hydroxyl radical, hydrogen peroxide, reducing oxidative stress and lipid peroxidation. Cypermethrin induces oxidative stress as well as alter the defense mechanisms of detoxification and scavenging enzymes producing cytotoxic changes through generation of ROS (Rasoul et al., 2012).

Banerjee et al. (2009) observed morphological improvement in the kidney of rats following administration of garlic extract and vitamin C as slight degeneration, few vacular spaces, few merging of distal convoluted tubules, and slight distortion of cells and basement membrane indicating significant reduction in the severity of damage caused by cypermethrin.

It can be concluded that subchronic exposure to cypermethrin is hepatotoxic and nephrotoxic. Concomitant administration of garlic extract or vitamin C with cypermethrin protect liver and kidneys from cypermethrin’s induced injury & both together are more protectors than each of them alone.

Recommendations:

Improvement of health education programs for the purpose of increasing public awareness regarding the health impact of cypermethrin and their sources, reading the label on consumer products and instructing farmers and workers about the handling of cypermethrin and using the simplest hygienic and protective measures Administration of garlic extract or vitamins C may be of immense prophylactic and therapeutic values in exposed individuals.

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The Evaluation Of Protective Role Of Garlic.


Cypermethrin has the potential to induce hepatic oxidative stress, DNA damage and apoptosis in adult zebrafish (Danio rerio). *Journal of Environmental Biology;* 12(1): 18–20


تقييم الدور الوقائي لمستخلص الثوم وفيتامين (ج) ضد التأثير السام دون المزمن للمبيد الحشرى السايرمثريين على الكبد والكلى

داليا محمد أمين

-65-

يركز البحث: تضخس سمية السايرمثريين للدراسات المستقبلية بناءً على التوزيع على نطاق واسع على مستوى العالم ويعتبر المصدر الرئيسي للسایرمثريين في الجسم البشري من خلال تناول الغذاء والقوارض. هجع البحث: يجري هذا البحث لدراسة تأثير إعطاء فيتامين (ج) والمستخلص الثوم على الكبد والكلى اثناء الاستخدام المتكرر للمبيد الحشرى السايرمثريين على الغلاف الجوي في جرذان التجارب البيضاء البالغة.

المجموعات: مجموعة الضابطة: لم يتم إعطاء أي علاج. المجموعة الأولى (المجموعة السايرمثريين): تم إعطاء الجرذان السايرمثريين والمستخلص الثوم مذابًا في الماء المقطوع عن طريق الفم مرة واحدة في مجموعات 32.5/كمجم و 500/كمجم.

النتائج: أدت المياجات الكيميائية واختبارات الفحص المجهري باستخدام صبغة الهيماتوكسلين والأيوسين لشرائح الكبد والكلى إلى مشاهدة أضرار مؤلمة واضحة. في المجموعة السايرمثريين، وجدت الدراسات أن إنزيمات الكبد وبيوماBAD تظهر انخفاضًا في مستوى إنزيمات الكبد (أنزيم ألانين أمينو ترانسفراز) والكيراتينين والكيراتينين أو وانزيم الأسبرتيت أمينو ترانسفيراز، بالإضافة إلى أن وجود إرتفاع في مستوى الباكس بروتين في أنسجة الكبد والكلى. وتظهر مشاهدات تلوث الكبد والكلى من خلال الجمعيات الهيستولوجية والمناعية بصفة عامة، وتظهر مشاهدات تلوث الكبد والكلى من خلال الجمعيات الهيستولوجية والمناعية بصفة عامة، وتظهر مشاهدات تلوث الكبد والكلي.