Protective Effect of Captopril and Nigella Sativa Oil against Carbon Tetrachloride Induced Nephrotoxicity in Male Rats

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Abstract Carbon tetrachloride (CCl4) is considered an environmental pollutant. Exposure to CCl4 is commonly associated with nephrotoxicity. Objective: The aim of this study was to demonstrate the possible preventive role and underlying mechanism of the single and combined administration of captopril and nigella sativa oil against CCl4 induced nephrotoxicity in male rats. Methods: This study was conducted on sixty four adult healthy male albino rats and divided into 8 groups (8 rats/group). Group I, rats which served as negative control, group II, rats which given liquid paraffin and saline (positive control), groups III, rats which given captopril in a dose of 100 mg/kg daily by gavage (positive control), group IV, rats which given nigella sativa oil in a dose of 4 ml/kg daily by gavage (positive control), group V, rats which injected i.p. with CCl4 three times/ week in a dose of 20 μL/100 gm body weight, group VI, rats which given captopril concurrently with CCl4 in the same mentioned dose and route, group VII, rats which given nigella sativa oil concurrently with CCl4 in the same mentioned dose and route; group VIII, rats which given captopril and nigella sativa oil daily concurrently with CCl4 in the same mentioned dose and route. All rats were given treatment for 6 weeks then sacrificed by decapitation. Biochemical parameters were assessed in all rat groups and statistically analyzed. In addition, histological and immunohistochemical investigations of renal tissues were done. Results: In CCl4 treated rats, the biochemical assays revealed significant statistical increase in the level of serum urea, creatinine, uric acid and nitric oxide as well as increase level of lipid peroxides, reduced glutathione and superoxide dimutase in the kidney tissue. Histological examination showed degeneration of glomeruli and tubules. High immunolabelling for activated caspase-3 in the kidney was detected on immunohistochemical examination. Treatment with captopril and nigella oil singly or in combination with CCl4 resulted in improvement of the biochemical, histological and immunohistological changes and this effect is more evident on their combined administration. Conclusion: Captopril and nigella sativa oil can be used as therapeutic agents in protection against CCl4 induced nephrotoxicity. This effect is attributed to decrease free radicals generation and apoptosis through inhibition of caspase-3 activity in renal tissue. It is recommended to conduct further experimental studies on large numbers and to investigate this effect on nephrotoxic patients.

Keywords CCl4, oxidative stress, captopril, nigella sativa oil and kidney.

Introduction Carbon tetrachloride (CCl4) is an environmental pollutant. It is one of the aliphatic hydrocarbons that have widespread uses in different industrial sectors as in production of chlorofluorocarbon refrigerants, foam blowing agents, cleaning compounds and organic solvents industry (Kalaf et al., 2006).

Carbon tetrachloride causes cellular damage in the kidney and other tissues such as the liver, heart, lung, testis, brain and blood (Pirinccioğlu et al, 2012). This harmful effect is attributed to oxidative stress that results in generation of free radicals which involved in lipid peroxidation, accumulation of dysfunctional proteins and upregulation of nitric oxide (NO) production (Khan et al., 2009). Free radicals are neutralized by an elaborate antioxidant defense system of enzymes such as glutathione and superoxide dimutase (Abdel Moneim and El-Khadragy, 2013).
NO is produced from L-arginine by the action of nitric oxide synthase (NOS) which exists in 3 isoforms, the neuronal NOS (nNOS), endothelial NOS (eNOS) and inducible NOS (iNOS). All the three NOS isoforms are present in the renal cortex. The nNOS is expressed predominantly in tubular epithelial cells of macula densa. The eNOS is found in vascular endothelial cells. The iNOS is abundant in tubular epithelium, including proximal tubules, thick ascending limbs of Henle, distal convoluted tubule and intercalated cells of the collecting duct (Tojo et al., 1997 and Dautzenberg et al., 2011).

Chronic kidney disease represents a serious public health problem because of increased its prevalence. Continuous studies for finding a novel renoprotective agents are required (Hrenák et al., 2013).

Captopril is an angiotensin-converting enzyme (ACE) inhibitor containing sulphydryl (SH) group. It was reported that it has a renoprotective action against renal damage induced by nephrotoxic agents such as cisplatin, gentamycin, doxorubicin and lead. This may be related to its free radicals scavenging due to its antioxidant effect (El-Sayed et al 2008, Rahman et al., 2009, Hrenák et al., 2013 and Ng et al. 2013).

Before establishing captopril as a therapeutic protective agent against nephrotoxicity in human, further studies should be conducted to confirm its renoprotective effect in exposure to different toxins (Rahman et al., 2009).

In addition it was mentioned that captopril, angiotensin converting enzyme inhibitors and angiotensin receptor blockers were used to attenuate hepatic damage in both animal and human studies (Abbas et al., 2011).

The nigella sativa seeds contain fixed oil, volatile oil, alkaloids and saponin. The fixed oil is composed mainly of unsaturated fatty acids. The volatile oil has been shown to contain thymoquinone and many monoterpene such as p-cymene, and α-pinene (Raza et al., 2008).

The nigella sativa seeds/oils are used in folk (herbal) medicine all over the world as anti-inflammatory, analgesic, antipyretic, antimicrobial, antineoplastic, anti asthmatic, antidiarrheal and dyslipidaemic. Both volatile oil and thymoquinone have been reported to give protection against nephrotoxicity and hepatotoxicity induced by chemicals. The beneficial effects are related to their antioxidant and cytoprotective actions by effect on mediators of inflammations (Ali and Blunden, 2003).

No available data are existing describing the role of captopril and nigella sativa oil in protection against CCl4 induced nephrotoxicity in rats.

**Aim of work**

This study was designed to evaluate the possible protective effect of single and combined administration of captopril and nigella sativa oil against carbon tetrachloride induced nephrotoxicity. In addition, the effects of these nephroprotective agents on oxidative stress markers and caspase-3 were investigated.

**Materials and methods**

**A (Chemicals)**

Carbon tetrachloride (analar grade) was obtained from ADVIC (Egypt). Captopril, reduced glutathione, Ellman's reagent [5,5-Dithiobis (2-nitrobenzoic acid), DTNB]) were purchased from Sigma Aldrich Co. (USA). All other chemicals were of analytical grades.

The nigella sativa oil was obtained from Cap Pharm Company, Egypt. Immunohistochemical kits for caspase-3 (CPP 32; AB-4 Rabbit Polyclonal Antibody) and Universal detection kit were obtained from Thermo Fisher Scientific (Fermont, California, USA).

**B (Animals and Experimental design)**

A total number of sixty four healthy adult male albino rats were used in this study and their weight ranged between (220-250) gms. They were obtained from the Animal House of Faculty of Medicine, Assiut University. The animals were housed in stainless steel cages with appropriate ventilation and temperature, fed on standard rodent pellet diet manufactured by the Egyptian Company for Oil and Soap (Cairo, Egypt) and tap water available ad libitum. These rats were maintained at constant daily light/dark periods for 12 hours. The rats were acclimatized to lab conditions prior to starting dosing for a period of 2 weeks. Dealing with the animals followed the regulations of the Animal house of Faculty of Medicine, Assiut University. Rats were weighed weekly in order to adjust the dose of chemicals/drugs according to their body weight. The duration of exposure to drugs/chemicals was 6 weeks, this was the time suggested by (Ma et al., 2014-b) to develop toxic effect on the kidneys.

The rats were divided into 8 groups (each group consisted of 8 rats).

- **Group I:** (negative control). Rats were given ordinary rat diet and did not receive any drug. They were living in the same environmental conditions as the other groups to determine the normal values of the tested parameters.

- **Group II:** Rats were used as positive control, given paraffin oil by i.p. injection and saline (0.9% NaCl) by gavage for 6 weeks to test the effect of the vehicle (solvent) on the rats (Tuncer et al., 2003 and Ahmed et al., 2011).

- **Group III:** Rats were used as positive control, given captopril in a dose of 100 mg/kg daily by gavage for 6 weeks to test the effect of the treating drug on the rats.

- **Group IV:** Rats were given nigella sativa oil in a dose of 4 ml/kg by gavage for 6 weeks to test the effect of the treating drug on rats (Abdel-Zaher et al., 2011).

- **Group V:** Rats were injected with CCl4 three times/ week for 6 weeks in a dose of 20 μL/100 gm b.w. i.p. (represents 1/8 LD50 of CCl4) (Choi et al., 2011), diluted 1:6 in paraffin oil to induce nephrotoxicity (Ahmed et al., 2011).

- **Group VI:** Rats were given captopril in a dose of 100 mg/kg daily by gavage concurrently with CCl4 i.p. for 6 weeks. This dose is the therapeutic dose
which mentioned to give protection against nephrotoxicity induced by agents induces oxidative stress (Hrenák et al., 2013).

Group VII: Rats were given nigella sativa oil in a dose of 4 ml/kg by gavage in combination with CCl₄ i.p. for 6 weeks. This was found to be the dose which gives protection against oxidative stress (Abdel-Zaher et al., 2011).

Group VIII: Rats were given captopril in a dose of 100 mg/kg along with nigella sativa oil in a dose of 4 ml/kg by gavage concurrently with CCl₄ i.p. for 6 weeks.

(C) Blood and tissue samples collection
Blood samples were collected from all rat groups 24 hours after the last dose at the end of 6th weeks from the retro-orbital plexus of veins using a capillary pipette. The collected blood samples in glass tubes were centrifuged at 3000 rpm for obtaining sera which stored at -20 until required for estimation of biochemical parameters.

The rats were sacrificed by decapitation after light ether anesthesia and dissected at the end of 6th weeks. The kidneys were removed and divided into two parts; the first part was homogenized in ice cold saline. The kidney tissue homogenates were centrifuged at 3000 rpm for 15 minutes. The resultant supernatant was stored at -70 for biochemical analysis. The second part divided into 2 portions, part was fixed in 10% formalin and processed for light microscopic examination and other part fixed in 5% gluteraldehyde for electron microscopic examination (Khan et al., 2009).

(D) Biochemical Measurements
(I) Assessment of renal function tests
Measurement of serum urea, creatinine and uric acid was done by spectrophotometer using colorometric diagnostic kits (Biodiagnostic, Giza, Egypt).

- Serum urea was measured according to the method that described by Coulombe and Favreau (1963).
- Serum creatinine was measured according to the method that described by Larsen (1972).
- Serum uric acid was measured according to the method that described by Whitehead et al. (1991).

(II) Measurement of oxidative stress markers

1. Serum nitric oxide (NO) level assay
It is measured in the serum by assaying nitrate, one of the stable and non volatile end-products of NO oxidation. Serum nitrite concentration was measured using Griess reagents as described by Green and his coworker (1982). The absorbance measured at 550 nm using UV-visible spectrophotometer and calculate the serum NO level in samples from the standard curve and expressed as μmol/ml.

2. Lipid peroxidation assay
Measurement of lipid peroxidation in kidney tissue was carried out by the method described by Ohkawa, et al. (1979). Lipid peroxidation is an oxidative stress marker, was determined by measuring the level of the Malonic aldehyde (MDA) in the kidney tissue homogenates. The MDA is an end product of lipid peroxidation and its level was determined by use of thiobarbituric acid reactive substances (TBARS) method. The TBARS which formed in each of the samples was assessed spectrophotometrically by measuring optical density of the supernatant at 532 nm against a reagent blank and expressed as μmol/mg protein.

(3) Reduced glutathione (GSH) assay
Estimation of intracellular GSH content of neutralized supernatant renal tissue was assayed using Ellman's reagent (Ellman, 1959) which composed of [5,5-Dithiobis (2-nitrobenzoic acid), DTNB solution] which measured spectrophotometrically according to the method described by Griffith (1980). A standard reference curve was prepared for each assay. The optical density of the supernatant measured at 412 nm. Results were expressed as μmol/g protein. The protein contents were determined by using the method described by Lowry, et al. (1951).

4. Superoxide dimutase (SOD) assay
The assay of SOD activity in the supernatant of renal homogenate was determined spectrophotometrically by using commercial kit utilizing a dye that produces a water soluble formazan upon reduction with a superoxide anion. The rate of the reduction with a superoxide anion is inhibited by SOD. The higher the activity of SOD is the less formation of formazin and the lower the absorption value. The optical density of the supernatant measured at 480 nm. Results were expressed as μ/mg protein for tissue supernatant according to the method described by Kono (1978).

(E) Histopathological analysis
(1) Light microscopic examination
The specimens were taken from the kidneys of all groups and were fixed in formalin 10%. Paraffin sections were prepared and stained with Haematoxylin and Eosin and PAS (Periodic-acid Schiff) stains (Durary and Wallington, 1980).

(2) Immunohistochemical examination
Thick sections (3-5µ) were cut, mounted on poly L-lysine coated slides. Deparaffinization was done in xylene and rehydration in descending grades of alcohol. Hydrogen peroxide was added for 15 minutes to reduce non specific background staining due to endogenous peroxidase then the specimens were washed 2 times in phosphate buffer saline (PBS) for 5 minutes. Ultra V Block was applied and incubated for 5 minutes at room temperature to block non specific background staining. The slides were rinsed in PBS then in citrate buffer in a microwave oven for 5 minutes. The slides were left to cool down for 20 minutes then washed 4 times in PBS, Caspase-3 (CPP32) Ab 4 rabbit polyclonal antibody, Thermo scientific-USA) was added for one hour at room temperature in a dilution of 1:100, after that washed 4 times in PBS. Next step, Biotinylated Goat Anti-Polyvalent (secondary antibody) was applied for all the slides and incubated for 30 minutes at room temperature, afterward they washed 4 times in PBS. Streptavidin Peroxidase was applied and incubated for 10 minutes at room temperature, after that rinsed in
PBS. One to two drops of DAB chromogen (Diaminobenzidine) was added for 5-15 minutes and washed in distilled water for 5 minutes. Counterstaining by Meyer’s Hx was followed by washing in tap water. Dehydration by ascending grades of alcohol, clearing for few seconds, mounting using coverslip and a permanent mounting media (Cerson, 1990). For negative control staining, some sections were incubated with PBS instead of the primary antibody. No immunoreactivity was present in these sections. Positive control sections for caspase-3 was from tonsil.

(3) Transmission electron microscopy

Immediately after the rats being sacrificed, 10-12 small pieces of the kidney cortex were fixed in 5% gluteraldehyde for 24 hours. The specimens were after that washed in 3-4 changes of cacodylate buffer (PH 7.2) for 20 minutes in every change and post fixed in cold 1% osmium tetroxide for two hours. Next they were washed in four changes of cacodylate buffer for 20 minutes each. Dehydration was done by using ascending grades of alcohol (30, 50, 70, 90 and absolute alcohol) each for two hours. Clearing in propylene oxide followed by embedding in Epon. These samples were kept in incubator at 35 degree for one day, then at 45 degree for another day and lastly for three days at 60 degree (Gupta, 1983). Semithin sections (0.5-5um) were prepared by using LKB ultra microtome. The sections were stained by Toluidine blue, examined by light microscope " JEM-100 CXII, Japan " in the electron microscope unit of Faculty of Medicine, Assiut University.

(F) Statistical Analysis

The data were presented in the form of mean± standard deviation (SD). Statistical analysis was done using one way analysis of variance (one-way ANOVA) was done to compare between means of all different rat groups using SPSS computer program, Version 20. The difference was non significant at P≥ 0.05, significant at P < 0.05 and highly significant at P ≤ 0.01) (Kirkwood and Sterner, 2003).

Results

(A) Biochemical assays

(I) Kidney function tests

Table (1): shows one way ANOVA statistical analysis of serum urea, creatinine and uric acid levels in all rat groups.

The mean of serum levels of urea, creatinine and uric acid (mg/dl) were highly significantly increased in CCl₄ treated group (61.2 ± 2.4), (0.622 ± 0.04) and (8.17 ± 0.44) respectively when compared to control group I (23.8 ± 1.9), (0.338 ± 0.01) and (4.96 ± 0.24) and other control groups (P. value ≤ 0.01).

There were highly significant decrease in the mean of serum urea, creatinine and uric acid levels on administration of captopril with CCl₄ (47.1 ± 2.5), (0.458 ± 0.02) and (5.85 ± 0.29) respectively (P. value ≤ 0.01). There were highly significant decrease in the mean of serum urea, creatinine and uric acid levels on administration of nigella sativa oil with CCl₄ (45.3 ± 2.4) (0.452 ± 0.02), and (5.95 ± 0.29) respectively (P. value ≤ 0.01). A highly significant decrease in the elevated serum level of urea, creatinine and uric acid was found in combination of nigella sativa oil+ captopril treated group (35.5 ± 2.3), (0.364 ± 0.014) and (5.32 ± 0.15) respectively when compared with CCl₄ treated group (P. value ≤ 0.01). There was no significant statistical difference between the mean of serum urea, creatinine and uric acid levels in all control rat groups (I, II, III and IV) (P. value ≥ 0.05).

(II) Serum nitric oxide (NO) level

Table (2): shows one way ANOVA statistical analysis of serum NO levels in all rat groups. The mean of serum NO level (µmol/L) was highly significantly increased in CCl₄ treated group (28 ± 2.1) as compared to control group (16.3 ± 1.3) (P. value ≤ 0.01). There was a highly significant decrease in the elevated serum NO level to near control value in rats which administrated of CCL₄ concomitant with captopril alone (22.3 ±2) or nigella sativa oil alone (21.4 ± 1.8) and captopril combined with nigella sativa oil (19.2 ± 1.6) when compared to CCl₄ treated group (28 ± 2.1) (P. value ≤ 0.01). There was no significant statistical difference between the mean of in serum NO level of all control rat groups (I, II, III and IV) (P. value ≥ 0.05).

(III) Serum lipid peroxidation (TBARS) level

Table (3): shows one way ANOVA statistical analysis of TBARS level (nmol/mg protein) in kidney tissues of rats. The mean of TBARS level was a highly significantly increased in CCl₄ treated group (0.912 ± 0.07) as compared to control group (0.262 ± 0.01). There was a highly significant decrease in the mean of serum TBARS level in rat groups treated with nigella sativa oil alone (0.572 ± 0.02), captopril alone (0.632± 0.01) and when nigella sativa oil combined with captopril (0.486 ± 0.01). There was no significant statistical difference between the mean of TBARS level in all control rat groups (I, II, III and IV) (P. value ≥ 0.05).

(IV) Kidney reduced glutathione (GSH) level

Table (4): shows one way ANOVA statistical analysis of GSH level (µmol/g protein) in the kidneys of all rat groups. There was a highly significantly decrease in the mean of GSH level in CCl₄ treated group (1.9 ± 0.07) as compared to control group (4.8 ± 0.01). There was a highly significantly increase in the mean of decreased serum GSH level in rat group treated with nigella sativa oil alone (3.21 ± 0.01), captopril alone (2.9 ± 0.01) and when nigella sativa oil combined with captopril (3.75 ± 0.01) when compared to CCl₄ treated group (P. value ≤ 0.01). There was no significant statistical difference between the mean of kidney GSH level in all control rat groups (I, II, III and IV) (P. value ≥ 0.05).

(V) Kidney superoxide dimutase level (SOD)

Table (5): shows one way ANOVA statistical analysis of SOD level (U/mg protein) in kidneys of all rat groups.
groups. The mean of SOD level was a highly significantly decreased in CCl4 treated group (0.16 ± 0.07) as compared to control group (0.63 ± 0.01) (P. value ≤ 0.01). There was a highly significant increase in the mean of decreased serum GSH level in rat groups treated with nigella oil alone (0.49 ± 0.01), captopril alone (0.45 ± 0.01) and when nigella sativa oil combined with captopril (0.57 ± 0.01) when compared to CCl4 treated group (P. value ≤ 0.01). There was no significant statistical difference between the mean of control rat groups (I, II, III and IV) in kidney SOD level (P. value ≥ 0.05).

Table (1): One Way ANOVA statistical analysis of serum urea, creatinine and uric acid levels in all rat groups. Each group composed of 8 rats.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Urea (mg/dl) Mean ± SE</th>
<th>Creatinine (mg/dl) Mean ± SE</th>
<th>Uric Acid (mg/dl) Mean ± SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. I: negative control</td>
<td>23.8 ± 1.9</td>
<td>0.338 ± 0.01</td>
<td>4.96 ± 0.24</td>
</tr>
<tr>
<td>G. II: paraffin oil (positive control)</td>
<td>25.1 ± 1.6</td>
<td>0.342 ± 0.02</td>
<td>4.86 ± 0.25</td>
</tr>
<tr>
<td>G. III: captopril</td>
<td>24.3 ± 1.7</td>
<td>0.339 ± 0.01</td>
<td>4.96 ± 0.24</td>
</tr>
<tr>
<td>G. IV: nigella sativa oil</td>
<td>24.11 ± 1.6</td>
<td>0.337 ± 0.015</td>
<td>4.86 ± 0.25</td>
</tr>
<tr>
<td>G. V: CCl4</td>
<td>61.2 ± 2.4 oo</td>
<td>0.622 ± 0.04 oo</td>
<td>8.17 ± 0.44 oo</td>
</tr>
<tr>
<td>G. VI: CCl4 + captopril</td>
<td>47.1 ± 2.5 **</td>
<td>0.458 ± 0.02 **</td>
<td>5.85 ± 0.29 **</td>
</tr>
<tr>
<td>G. VII: CCl4 + nigella sativa oil</td>
<td>45.3 ± 2.4 **</td>
<td>0.452 ± 0.02 **</td>
<td>5.95 ± 0.29 **</td>
</tr>
<tr>
<td>G. VIII: CCl4 + captopril + nigella sativa oil</td>
<td>35.5 ± 2.3**</td>
<td>0.364 ± 0.01**</td>
<td>5.32 ± 0.15**</td>
</tr>
</tbody>
</table>

oo Highly significant, in group V versus groups II, III and IV (P. value ≤ 0.01), ** Highly significant groups VI, VII and VIII versus group V (P. value ≤ 0.01).

Table (2): One Way ANOVA statistical analysis of serum nitric oxide levels in all rat groups. Each group composed of 8 rats.

<table>
<thead>
<tr>
<th>Groups</th>
<th>Serum NO (µmol/ml) Mean ± SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. I: negative control</td>
<td>16.3 ± 1.3</td>
</tr>
<tr>
<td>G. II: paraffin oil (positive control)</td>
<td>16.6 ± 1.2</td>
</tr>
<tr>
<td>G. III: captopril</td>
<td>16.5 ± 1.1</td>
</tr>
<tr>
<td>G. IV: nigella sativa oil</td>
<td>16.4 ± 1.1</td>
</tr>
<tr>
<td>G. V: CCl4</td>
<td>28 ± 2.1 oo</td>
</tr>
<tr>
<td>G. VI: CCl4 + captopril</td>
<td>22.3 ± 2 **</td>
</tr>
<tr>
<td>G. VII: CCl4 + nigella sativa oil</td>
<td>21.4 ± 1.8 **</td>
</tr>
<tr>
<td>G. VIII: CCl4 + captopril + nigella sativa oil</td>
<td>19.2 ± 1.6 **</td>
</tr>
</tbody>
</table>

oo Highly significant, in group V versus groups II, III and IV (P. value ≤ 0.01), ** Highly significant groups VI, VII and VIII versus group V (P. value ≤ 0.01).

Table (3): One Way ANOVA statistical analysis of lipid peroxidation (TBARS) level (nmol/ml) in kidneys of all rats groups. Each group composed of 8 rats.

<table>
<thead>
<tr>
<th>Groups</th>
<th>TBARS (nmol/mg protein) (Mean ± SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. I: negative control</td>
<td>0.262 ± 0.01</td>
</tr>
<tr>
<td>G. II: paraffin oil (positive control)</td>
<td>0.266 ± 0.01</td>
</tr>
<tr>
<td>G. III: captopril</td>
<td>0.270 ± 0.01</td>
</tr>
<tr>
<td>G. IV: nigella sativa oil</td>
<td>0.268 ± 0.01</td>
</tr>
<tr>
<td>G. V: CCl4</td>
<td>0.912 ± 0.07 oo</td>
</tr>
<tr>
<td>G. VI: CCl4 + captopril</td>
<td>0.572 ± 0.02 **</td>
</tr>
<tr>
<td>G. VII: CCl4 + nigella sativa oil</td>
<td>0.632 ± 0.01 **</td>
</tr>
<tr>
<td>G. VIII: CCl4 + captopril + nigella sativa oil</td>
<td>0.486 ± 0.01 **</td>
</tr>
</tbody>
</table>

oo Highly significant, in group V versus groups II, III and IV (P. value ≤ 0.01), ** Highly significant groups VI, VII and VIII versus group V (P. value ≤ 0.01).

Table (4): One Way ANOVA statistical analysis of reduced glutathione level (GSH) (nmol/ml) in kidneys of all rats groups. Each group composed of 8 rats.

<table>
<thead>
<tr>
<th>Groups</th>
<th>GSH (µmol/g protein) (Mean ± SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. I: negative control</td>
<td>4.6 ± 0.01</td>
</tr>
<tr>
<td>G. II: paraffin oil (positive control)</td>
<td>4.9 ± 0.01</td>
</tr>
<tr>
<td>G. III: captopril</td>
<td>4.8 ± 0.01</td>
</tr>
<tr>
<td>G. IV: nigella sativa oil</td>
<td>4.7 ± 0.01</td>
</tr>
<tr>
<td>G. V: CCl4</td>
<td>1.9 ± 0.07 oo</td>
</tr>
<tr>
<td>G. VI: CCl4 + captopril</td>
<td>3.21 ± 0.01</td>
</tr>
<tr>
<td>G. VII: CCl4 + nigella sativa oil</td>
<td>2.7 ± 0.01</td>
</tr>
<tr>
<td>G. VIII: CCl4 + captopril + nigella sativa oil</td>
<td>3.75 ± 0.01</td>
</tr>
</tbody>
</table>

oo Highly significant, in group V versus groups II, III and IV (P. value ≤ 0.01), ** Highly significant groups VI, VII and VIII versus group V (P. value ≤ 0.01).
Table (5): One Way ANOVA statistical analysis of superoxide dimutase level (SOD) (nmol/ml) in kidneys of all rats groups. Each group composed of 8 rats.

<table>
<thead>
<tr>
<th>Groups</th>
<th>SOD (U/mg protein) (Mean ± SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>G. I: negative control</td>
<td>0.63 ± 0.01</td>
</tr>
<tr>
<td>G. II: paraffin oil (positive control)</td>
<td>0.61 ± 0.01</td>
</tr>
<tr>
<td>G.III: captopril</td>
<td>0.62 ± 0.01</td>
</tr>
<tr>
<td>G.IV: nigella sativa oil</td>
<td>0.64 ± 0.01</td>
</tr>
<tr>
<td>G. V: CCl₄</td>
<td>0.16 ± 0.07 oo</td>
</tr>
<tr>
<td>G. VI: CCl₄ + captopril</td>
<td>0.49 ± 0.01 **</td>
</tr>
<tr>
<td>G. VII: CCl₄ + nigella sativa oil</td>
<td>0.44 ± 0.01 **</td>
</tr>
<tr>
<td>G.VIII: CCl₄ + nigella sativa oil + captopril</td>
<td>0.57 ± 0.01 **</td>
</tr>
</tbody>
</table>

oo Highly significant, in group V versus groups I, II, III and IV (P. value ≤=0.01), ** Highly significant groups VI, VII and VIII versus group V(P. value ≤=0.01).

(B) Histopathological Examination

I-Light Microscopic Examination

(Fig.1): A photomicrograph of a section of the renal cortex of control rat groups (I, II, III and IV). The renal corpuscle shows the glomerulus (G) with its capillary tufts surrounded by a narrow capsular space (S) and Bowman's capsule (arrow). Proximal convoluted tubules (P) have a narrow lumen and a highly acidophilic cytoplasm. Distal convoluted tubules (D) have a wider lumen and less acidophilic cytoplasm. H&E x400

(Fig.2-a): A photomicrograph of the renal cortex of control rat groups (I, II, III and IV), showing a part of a renal corpuscle. Glomerular capillaries are defined by the prominent glomerular basement membrane (C). Podocytes (P) have pale nuclei and pale stained cytoplasm. Note the mesangial cell (M) and the squamous cells of Bowman's capsule (arrow).

(Fig.2b): A photomicrograph of the renal cortex of control rat groups (I, II, III and IV). The cells of proximal convoluted tubules (P) have apical brush border (bb) and well-formed basal striations (arrow). Their nuclei are central and pale with prominent nucleoli. The distal convoluted tubules (D) have pale apical nuclei with no brush border. Toluidine blue x1000

(Fig.3): (a)-A photomicrograph of the renal cortex of group V rats, showing obliteration of the capsular space (arrow).

(Fig.3): (b): A photomicrograph of group III rats, showing dilatation of some tubules (arrow) with obliteration of the lumen of others by exfoliated cells (asterisk). H&E x400

(Fig.4): (a)- A photomicrograph of the renal cortex of group V rats, showing a renal corpuscle with atrophic glomeruli (G). Note congestion of glomerular capillaries (C).
(Fig.4): (b)- A photomicrograph of group IV rats, showing vacuolation of the cells of some kidney tubules (asterisk). Note: thickening of the basement membrane of some kidney tubules (arrow). Toluidine blue x1000

(Fig.5): A photomicrograph of the renal cortex of group VI rats, showing normal appearance of the renal corpuscle. Note: Focal cellular infiltration (arrow) with congested blood vessels (c). (H & E)x400

(Fig.6): a- A photomicrograph of the renal cortex of group VI rats, showing normal appearance of the renal corpuscle. Note: the proximal tubules show many deeply stained granules (arrow).

(Fig.6): b- A photomicrograph of the same group showing some tubules with vacuolated cytoplasm (asterisk) and thickened basement membrane (arrow). Note: a congested blood vessel is seen (C). Toluidine blue x 1000

(Fig.7): A photomicrograph of the renal cortex of group VI rats, showing normal appearance of the renal cortex. H&E x 400

(Fig.8): - A photomicrograph of the renal cortex of group VII rats, showing a normal appearance of the renal cortex. Toluidine blue x 1000

(Fig.9): A photomicrograph of the renal cortex of group VIII, showing normal appearance of the renal cortex. H&E x400

(Fig.10): A photomicrograph of the renal cortex of group VIII, showing normal appearance of the renal cortex. Toluidine blue x1000
(Fig. 11): A photomicrograph of a section of the renal cortex of control rat groups (I, II, III and IV rats showing PAS-positive reaction along the basement membranes of the renal tubules (asterisk), the parietal layer of Bowman’s capsule (arrow) and glomerular capillaries (g) and the apical brush border of the proximal convoluted tubules (b).

   (PAS x400).

(Fig. 12): A photomicrograph of a section of the renal cortex of group V rats showing thickening in the basement membrane of many kidney tubules (arrow). (PAS x400).

(Fig. 13): A photomicrograph of a section of the renal cortex of group VI rats showing decrease in thickening of basement membrane of kidney tubules compared with group III.

   (PAS x400).

(Fig. 14): A photomicrograph of a section of the renal cortex of group VII rats showing mild thickening in the basement membrane of some kidney tubules compared with group III. (PAS x400).

(Fig. 15): A photomicrograph of a section of the renal cortex of group VIII showing normal appearance of kidney tubules. (PAS x400).
(II) Immuno-histochemical Examination

(Fig.16): A photomicrograph of the renal cortex of control rat groups (I, II, III and IV rats). The caspase-3 immunostaining is virtually absent in the renal corpuscle and the tubules.
(Caspase-3 immunostaining x1000)

(Fig.17): A photomicrograph of the renal cortex of group V rats. The caspase-3 immunostaining is observed in the renal corpuscle and in the kidney tubules (arrows).
(Caspase-3 immunostaining x1000)

(Fig.18): A photomicrograph of the renal cortex of group VI rats. Decrease in the reaction for caspase-3 immunostaining in the renal corpuscle and kidney tubules.
(Caspase-3 immunostaining x1000)

(Fig.19): A photomicrograph of the renal cortex of group VII rats. A noticeable reduction of caspase-3 immunostaining in the renal corpuscle and kidney tubules.
(Caspase-3 immunostaining x1000)

(Fig.20): A photomicrograph of the renal cortex of group VIII rats: showing marked decrease in caspase-3 immunostaining in the renal corpuscle and kidney tubules.
(Caspase-3 immunostaining x1000)

(III) Transmission Electron Microscopy Examination

(Fig.21): Electron micrograph of the renal cortex of control rat groups (I, II, III and IV) showing a podocyte with central nucleus (N) with characteristic well-formed primary (arrow head) and secondary foot processes (arrow) resting on the glomerular basement membrane. Note, the glomerular filtration barrier (astick) with its three components (fenestrated endothelium, glomerular basement membrane, and slit diaphragms) with uniform thickness. X 8000.

(Fig.22): Electron micrograph of the proximal convoluted tubule of control rat groups (I, II, III and IV) showing a spherical basal euchromatic nucleus (N) and elongated radially arranged mitochondria (m), Note the long apical microvilli (mv) and electron dense lysosomes (arrow). X 4000.

(Fig.23): Electron micrograph of the distal convoluted tubule of control rat groups (I, II, III and IV) showing regular basal infoldings (arrow), numerous elongated mitochondria (m) and euchromatic nuclei (N). X 4000.
(Fig.24): 24a-Electron micrograph of the renal cortex of group V rats, showing part of the podocyte with broadening of foot processes (arrow) with irregular spacing between them. Also, focal thickening of the basement membrane is also observed (arrow head). X 4800.

(Fig.24): b-Magnification of the previous micrograph, showing focal thickening and irregularity of the glomerular membrane (arrow). X 7200.

(Fig.25): Electron micrograph of the proximal convoluted tubule of group V rats showing short and irregular apical microvilli (mv). The mitochondria (m) are swollen and irregularly arranged in between large cytoplasmic vacuoles (v). X 3600.

(Fig.26): Electron micrograph of the proximal convoluted tubule of group V rats showing thickened irregular basement membrane (arrow) with electron dense cytoplasm (astrick). X 3600.

(Fig.27): Electron micrograph of the distal convoluted tubule of group V rats showing swollen cells with apical cytoplasmic bleb (astrick) and vacuolated apical cytoplasm (v). Note the preservation of basal infoldings (arrow) and organized mitochondria (m). X 3600.

(Fig.28): Electron micrograph of the distal convoluted tubule of group V rats showing few irregular basal infoldings (arrow). X 4800

(Fig.29): Electron micrograph of the renal cortex of group VI rats showing focal broadening of the foot processes with irregularity and thickening of the glomerular basement membrane (arrow). X4800

(Fig.30): Electron micrograph of the proximal convoluted tubule of group VI rats showing many irregular electron dense lysosomes (arrow). X 4800.

(Fig.31): Electron micrograph of the distal convoluted tubule of group VI rats showing irregular basement membrane (arrow) with electron dense mitochondria (m). X 4800.
Captopril inhibits the oxidative stress damage by generation of free radicals is implicated in the pathogenesis of kidney injury (Hismiogullari et al., 2015).

The metabolic conversion of CCl₄ by cytochrome P-450 give rise to formation of the reactive metabolite trichloromethyl radical (CCl₃) and trichloromethyl peroxide radical (CCl₂O₂). When the O₂ tension rises, a greater fraction of CCl₄ present in the system reacts very rapidly with O₂ and more reactive free radicals, such as CCl₂O₂ is generated. The free radicals initiate the peroxidation of membrane polyunsaturated fatty acids, cell necrosis, GSH depletion, membrane damage and loss of antioxidant enzyme activity (Ganie et al., 2011).

Carbon tetrachloride (CCl₄) has been used in rat experimental models to investigate the oxidative stress induced in various organs (Haghi et al., 2014).

Exposure to CCl₄ is commonly associated with acute and chronic renal damage (Mann et al., 2006) so this study aimed to investigate effect of CCl₄ on the kidney by biochemical, histological and immunohistochemical investigations.

Male rats were used in this work because they were particularly suitable animal model for evaluating tubular lesions because their intrarenal enzyme distribution is similar to that present in man (Weichert-Jacobsen et al., 1999).

In this work CCl₄ induced kidney damage which evidenced by biochemical assays of renal function tests by presence of increase level of serum urea, creatinine and uric acid. Also elevated serum level of the oxidative stress markers; malondialdehyde and NO. In addition, reduced antioxidant enzyme activity in kidney as shown by decrease level of GSH and SOD. Histopathological and immunohistochemical investigations confirmed occurrence of kidney damage. This was recorded by study of El-Gengaiehi et al., (2013).

The present study was designed to evaluate the possible protective effect of captopril and nigella sativa oil against the CCl₄ induced oxidative stress and nephrotoxicity by biochemical, histopathological and immunohisto-chemical.

It was reported that captopril inhibits the conversion of angiotensin I into angiotensin II. The peptide angiotensin II (AngII) plays a prominent role in oxidative stress-related disorders through actions on the AngII-AT1 receptors. AngII, is a potent inducer of reactive oxygen species (ROS), oxidative stress and inflammation in many cells (Bernstein et al., 2013).

It was found that captopril has renoprotective effective. Thus, it could be useful in conditions associated nephrotoxicity such as hypertension with acute glomerulonephritis, acute and chronic renal failure (Rahman et al., 2009).

The sulphhydryl-containing captopril appears to act as a scavenger of oxygen derived free radicals. Captopril decreased MDA concentration and activity of both glutathione reductase and peroxidase (Ackerman et al., 2008).

In this work the NO level as an oxidative stress marker was increased in CCl₄ treated rats which was reported by Malekinejad et al. (2012), in addition to histopathological and immuno-histochemical changes. The NO level is decreased and reach near normal level in single or combined captopril and nigella sativa oil administration.

It was reported that the CCl₄ induced upregulation of inducible nitric oxide synthase expression (Annadura et al., 2011). The NO level is corrected near normal by captopril. Also captopril can normalize renal morphological and functional alterations in rats. The protective effect of captopril is related to regulation of endothelial NO synthase expression and to a normalization of the oxidative stress parameters due to the inhibition of angiotensin II (García-Estahil et al., 2006).

In harmony with the present study the results of Anand et al., (2011) who reported that the expression of the iNOS gene appeared to be up-regulated in the liver and kidney samples of CCl₄ exposed untreated rats, whereas in CCl₄ exposed chrysin-treated rats, the mRNA transcript levels of iNOS approximated to normal levels.

**Discussion**

Carbon tetrachloride (CCl₄) is a potent nephrotoxic chemical. The target of CCl₄ is the brush border-bearing proximal tubule cells (Kang et al., 2011). Oxidative stress damage by generation of free radicals is implicated in the pathogenesis of kidney injury (Hismiogullari et al., 2015).

(Fig.32): Electron micrograph of the kidney cortex of group VII rats showing less broadening of foot processes with decrease in the thickening and irregularity of the basement membrane. X4800

(Fig.33): Electron micrograph of the proximal convoluted tubule of group VII rats, showing elongated radially arranged mitochondria (m). X4800.

(Fig.34): Electron micrograph of the distal convoluted tubule of group VII rats showing well developed basal infolding (arrow) but still there are electron dense mitochondria (m). X4800.
Also in agreement with the present work the results of EL Denshary et al., (2012) who reported that sider honey and/or Korean gingseng extract as an antioxidant resulted in a significant improvement in all evaluated kidney function tests and oxidative stress parameters which disturbed by CCl4 administration.

In agreement with results of the present study that was reported in other studies that captopril was used to suppress lipid peroxidation, oxidative stress and improve the antioxidant enzymes level (GSH and SOD) reaching values closer to normal and correct the hepatic damage induced by CCl4. This hepatoprotection could be attributed, at least in part, to the free radical scavenging properties of the drug (El-Katib and Mansour 2001, Tuncer et al., 2003 and Mansour et al., 2011). Also study of El-Sayed et al. (2008) reported that captopril has protective effect against cisplatin-induced nephrotoxicity in rats.

In this study the nigella sativa oil decrease the elevated renal function tests, TBARS and decrease the elevated NO level to reach normal value and correct histopathological alteration in kidneys.

In agreement, many studies mentioned that the nigella sativa seed extracts and oil or pure thymoquinone were found to suppress oxidative stress, improve antioxidant enzyme levels and correct renal damage which induced by many nephrotoxic agents such as cyclosporine (Uz et al., 2008), amakcin (Abdelaziz and Kandeel, 2011), cisplastin (Hadjadihdeh et al., 2012) and garamycin (Saleem et al., 2012).

It was found that nigella sativa seed extracts and oil corrected biochemical and oxidative stress damages induced by CCl4 in the liver (Mansour et al., 2001 and Kantor et al., 2005) and the blood (Meral and Kaner 2003). In agreement, the results of studies which reported that decrease the elevated NO production by CCl4 by many substances which has antioxidant properties as ghrelin (Cetin et al. 2010), Mushroom insoluble polysaccharides (Nada et al. 2012), yam peel extract (Yeh et al. 2013)

It was mentioned that nigella sativa seed and thymoquinone corrected the decreased level of antioxidant enzymes (El Sayed, 2011).

In harmony the other studies which reported antioxidant activity of some herbal extracts that have potential protective effects against CCl4 toxicity similar to nigella sativa such as terminalia arjuna (Manna et al., 2006), digera muricata (Khan et al., 2009), sonchus asper (Khan et al., 2010), podophyllum hexandrum (Ganie et al., 2011), punica granatum (Abdel Moneim and El-Khadragy, 2013), oxalis corniculata (Khan and Zehra, 2013). These studies revealed that these substances decreased the elevated serum urea, creatinine and uric acid levels, diminished the increased lipid peroxidation, corrected the altered NO level, enhancing the decreased antioxidant enzymes activities (such as SOD, GSH) and corrected the histopathological finding of kidney damage.

The key enzyme involving CCl4 induced nephrotoxicity is cytochrome P450, which is localized in the cortical tubule cells, and the increased lipid peroxidation is evident in the renal brush border. CCl4 also affects renal mitochondrial function including calcium flux across mitochondrial membranes (Natarajan et al., 2006).

It was found that nigella sativa oil has effect on cytochrome P450 which is involved in nephrotoxicity and hepatotoxicity (Ibrahim et al., 2008)

In agreement that was reported by Hwang et al., (2013) who mentioned that CCl4 induced renal damage through differential regulation of the caspase-3. Renal damage was manifested by rise of blood urea nitrogen and serum creatinine, histologically showed decreased diameter of the glomerulus, a lower number of capillaries, and a convoluted tubule in the kidney section. This damage was corrected by administration of gomisin A which has antioxidant properties.

In addition, the present study revealed presence of histopathological changes in the kidneys by light and electron microscopic examination such as degeneration of renal glomeruli, thickening of basal membrane of renal tubules, dense mitochondrial, cytoplasm and lysosymes damage. The immunohistochemical examination showed high immunolabelling for activated caspase-3 in the kidney in CCl4 treated rats. The concomitant administration of captopril and nigella sativa oil with CCl4 resulted in improvement of all mentioned changes.

In agreement with the current work was the study of Taskin et al., (2014) who reported that captopril and other inhibitors of angiotensin-II were effective against acute adriamycin induced nephrotoxicity via restoration of mitochondrial membrane potential, adenosine triphosphate (ATP) production and prevention of mitochondrial damage in vivo.

In harmony the results of Hermenean et al., (2013) who reported that CCl4 resulted in increase of MDA concentration and decrease in GSH and SOD levels. These were accompanied by glomerular and tubular degenerations, vascular congestion, necrosis, Ultrastructural examination showed proximal and distal tubular epithelial cells alterations including dense numerous lysosomes, altered mitochondria and basal enfolding dilatation. Pre-treatment with naringenin which has antioxidant effect resulted in return of the changed biochemical and histopathological parameters to normal.

CCl4-induced profound elevation of reactive oxygen species production which generated from oxidative stress, as evidenced by increased level of lipid peroxidation and depleted the level of total antioxidant enzymes. Also it induced activation of apoptotic related proteins including caspase-3 (Haghi et al., 2014).

It was found that CCl4 caused increase in kidney MDA levels and prominent histopathological kidneys damage. Glomerular and tubular degeneration, interstitial mononuclear cell infiltration and fibrosis, and vascular congestion in the peritubular blood vessels were observed in the renal cortex. These histopathological changes were disappeared in rats treated with CCl4 plus caffeic acid phenethyl ester due to its antioxidant effect (Ogeturk et al., 2005).
In this work the concomitant administration of captopril and nigella oil with CCl₄ resulted in improvement of histological and immunohistochemical changes in the kidneys which may attributed to decrease apoptosis.

It was reported sesamin, L-theanine and curcumin attenuated CCl₄-induced apoptosis through marked inhibition of caspase-3 activities. This inhibition is attributed to the antioxidant activities of these compounds, modulation of apoptosis signaling pathway or inhibiting metabolic activation of CCl₄ (Jiang et al., 2012, Ma et al., 2014-a and Hismiogullari et al. 2015).

Conclusions and Recommendations

Captopril and nigella sativa oil have protective effects against renal damage induced by CCl₄. The renoprotective effect of captopril is related to its antioxidant effect because it contains sulfhydryl (-SH) group and inhibition of caspase-3 activity which involved in apoptosis pathway.

The renoprotective effect of nigella sativa oil is related to its cytoprotective and antioxidant actions, suppression of cytochrome p450 in renal cortex which results in metabolic conversion of CCl₄ to toxic metabolites and its effect on caspase-3.

Both decrease the elevated NO level to reach near normal level and protect the mitochondrial from damage. Their combined administration gives more potent renoprotective effect than single drug administration which is of therapeutic value against nephrotoxicity.

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الخلاصة

التأثير الوقائي للكابتوبريل وزيت حبة البركة المضاد للتسمم الكلوي

العنوان:
التأثير الوقائي للكابتوبريل وزيت حبة البركة المضاد للتسمم الكلوي

الخلاصة

راثع كلوريد الكربون يعتبر من الملوثات البيئية، ومن الشائع أن تؤدي نزيفه إلى تضرر السكري في الكبد، وتفاقم حالة التسمم الكلوي عند التعرض له. هدفت هذه الدراسة إلى توضيح الدور الوقائي المحتمل للكلوريد الفعال في الوقاية من التسمم الكلوي في ذكور الحمامات عند تناولها مفردة ومجموعة. أجريت هذه الدراسة على أربعة وأربعون من ذكور الحمامات البالغة، والتي تم تقسيمهم إلى ثمانية مجموعات سطوعية (تحتوي كل منها ٢ حمار).

المجموعة الأولى هي جراثيم الأمول 알려كم المقصورة. المجموعة الثانية هي جراثيم الأمول اللازمة الموجبة التي تم علاجها بالكابتوبريل نويلي ١٠٠ مجم / كجم يومياً، والمجمعة الثالثة هي جراثيم الأمول اللازمة الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٤ مل / كجم يومياً، والمجمعة الرابعة هي جراثيم الأمول اللازمة الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٣ مل / كجم يومياً، والمجمعة الخامسة هي جراثيم الأمول اللازمة الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٢ مل / كجم يومياً، والمجمعة السادسة هي جراثيم الأمول اللازمة الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ١ مل / كجم يومياً، والمجمعة السابعة هي جراثيم الأمول اللازمة الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٠.٥ مل / كجم يومياً، والمجمعة الثامنة هي جراثيم الأمول اللازمة الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٠ مل / كجم يومياً.

تم جمع الفحوصات الكيميائية والهستوغرافيميا (بالميكروسكوب) وتحليلها إحصائياً، وكذلك تم عمل الفحوصات الهستوغرافيميا والهستوكيماوية وقياساتها الأكسيدية في الجراثيم البالغة، وذلك لقياس عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند معالجة الجماع بالكابتوبريل أو زيت حبة البركة مما مفردة من التسمم الكلوي، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند علاج مجموعة الأمول بالكابتوبريل أو زيت حبة البركة مما مجوعة، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد.

الخلاصة

تهدف الدراسة إلى توضيح الدور الوقائي المحتمل للكلوريد الفعال في الوقاية من التسمم الكلوي عند تناولها مفردة ومجموعة. أجريت هذه الدراسة على أربعة وأربعون من ذكور الحمامات البالغة، والتي تم تقسيمهم إلى ثمانية مجموعات سطوعية (تحتوي كل منها ٢ حمار). المجموعة الأولى هي جراثيم الأمول المقصورة، والمجمعة الثانية هي جراثيم الأمول الموجبة التي تم علاجها بالكابتوبريل نويلي ١٠٠ مجم / كجم يومياً، والمجمعة الثالثة هي جراثيم الأمول الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٤ مل / كجم يومياً، والمجمعة الرابعة هي جراثيم الأمول الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٣ مل / كجم يومياً، والمجمعة الخامسة هي جراثيم الأمول الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٢ مل / كجم يومياً، والمجمعة السادسة هي جراثيم الأمول الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ١ مل / كجم يومياً، والمجمعة السابعة هي جراثيم الأمول الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٠.٥ مل / كجم يومياً، والمجمعة الثامنة هي جراثيم الأمول الموجبة التي تم علاجها بالكابتوبريل وزيت حبة البركة نويلي ٠ مل / كجم يومياً.

تم جمع الفحوصات الكيميائية والهستوغرافيميا (بالميكروسكوب) وتحليلها إحصائياً، وكذلك تم عمل الفحوصات الهستوغرافيميا والهستوكيماوية وقياساتها الأكسيدية في الجراثيم البالغة، وذلك لقياس عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند علاج مجموعة الأمول بالكابتوبريل أو زيت حبة البركة مما مجوعة، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند علاج مجموعة الأمول بالكابتوبريل أو زيت حبة البركة مما مجوعة، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند علاج مجموعة الأمول بالكابتوبريل أو زيت حبة البركة مما مجوعة، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند علاج مجموعة الأمول بالكابتوبريل أو زيت حبة البركة مما مجوعة، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند علاج مجموعة الأمول بالكابتوبريل أو زيت حبة البركة مما مجوعة، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد. وعند علاج مجموعة الأمول بالكابتوبريل أو زيت حبة البركة مما مجوعة، نتج عن ذلك تقليل عدد الخلايا المبكرة من المادة (الكازبيز -٣) في أنسجة الكبد.

توصيات

توصيات البحث حول استخدام الكابتوبريل وزيت حبة البركة كعلاجاتاً إضافيتين في حالات التسمم الكلوي مع كلوريد الكربون. وتوصيات البحث حول استخدام الكابتوبريل وزيت حبة البركة كعلاجاتاً إضافيتين في حالات التسمم الكلوي مع كلوريد الكربون. وتوصيات البحث حول استخدام الكابتوبريل وزيت حبة البركة كعلاجاتاً إضافيتين في حالات التسمم الكلوي مع كلوريد الكربون. وتوصيات البحث حول استخدام الكابتوبريل وزيت حبة البركة كعلاجاتاً إضافيتين في حالات التسمم الكلوي مع كلوريد الكربون.