ANTIOXIDANT AND ANTIAPOPTIC EFFECTS OF COMBINED SIDR HONEY AND NIGELLA SATIVA OIL AGAINST PARACETAMOL-INDUCED HEPATO-NEPHROTOXICITY IN RATS

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ABSTRACT

Background: Acetaminophen (Paracetamol; PCM), commonly utilized as analgesic and antipyretic drug in many painful and febrile disorders, has been found to induce liver and renal disorders in both animals and humans. Its administration in a high dose causes hepatic and renal toxicities and results in hepato-renal cell deaths by activating multiple stress pathways. Objective: This study aimed to investigate and compare effects of Sidr honey (SH), Nigella sativa oil (NS) and their combination on the paracetamol (PCM)-induced hepato-renal toxicities in rats.

Methods: Forty male adult albino rats were divided into five groups and treated for 4 weeks (n= 8 each): (1) the control group; receiving distilled water orally, (2) PCM-treated; receiving single high dose PCM (SHDP) of 1 g/kg once orally, (3) NS- and PCM-treated (NS/P); receiving NS in the dose of 2 ml/kg/day orally, and at the end of NS-treatment, the same single PCM dose is given, (4) SH- and PCM-treated (H/P); receiving SH in the dose of 1 g/kg/day, and at the end of SH-treatment, the same PCM dose is given, (5) SH-, NS- & PCM-treated (H&NS/P) group; receiving SH and NS orally in the same doses, and at the end of treatments, the same PCM dose is given. Serum alanine transaminase (ALT), aspartate transaminase (AST), blood urea nitrogen (BUN), creatinine, total antioxidative capacity (TAC) and Fas ligand (Fas L), and liver tissue TAC were measured after sacrificing the rats at the end of experiment.

Results: Compared to control group, the SHDP-treated rats developed significant increases in serum ALT, AST, BUN, Creatinine and Fas L, and decreases in serum and liver tissue TAC. SH, NS or more effectively the combined H/NS/P treatments produced significant decreases in serum ALT, AST, BUN, Creatinine and Fas L, and increases in serum and liver tissue TAC when compared with SHDP-treated rats.

Conclusion: These findings suggested that oral combined SH and NS administration is more protective against PCM-induced hepato-renal toxicity in rats than using each of them alone. The collective data demonstrated that SH and NS have considerable ability to protect against oxidation, apoptosis, and other harmful effects of PCM in rats.

Keywords: Sidr honey, Nigella sativa, Paracetamol, Apoptosis, Oxidation.

INTRODUCTION

Drug-induced hepatic injury is the commonest cause of acute hepatic failure1. The direct toxic injury is dose-dependent and predictable and can be experimentally-reproducible. The idiosyncratic damage is supported by the innate and the adaptive immune system. Unfortunately, drug-induced liver and kidney injuries are significant and still unresolved clinical problem. Paracetamol (PCM)-induced toxicity is a common cause of acute hepatic and renal failure. It is predominantly metabolized through conjugation with sulfate and glucuronide, but a small amount is degraded by CYP2E1 to the highly reactive (toxic) metabolite N-acetyl-p-benzoquinoneimide (NAPQI)2,3. Then, this toxic metabolite NAPQI is detoxified by binding with glutathione (GSH). If an extra amount of paracetamol reaches the liver, the liver conjugation capacity is overwhelmed and the remaining unbound NAPQI covalently binds to cellular and mitochondrial proteins, leading to cell death1,4.

Al-Jabri stated that honey is a natural product which is widely used for its therapeutic effects5. It has been reported that honey contains about 200 compounds. Chinese, ancient Egyptians, Assyrians, Greeks and Romans used honey for wound healing and treatment of intestinal diseases. Also, all Muslims believed that Honey is a curative agent from nearly all diseases according to what confirmed in Tibb
3. The experimental design and animal groups: The rats were randomly divided into five groups of 8 animals each, as follows:

i. Normal control group: The rats received distilled water in the dose of 1 mL/kg body weight/day orally by gavage for 4 weeks.

ii. Single high dose PCM-treated (SHDP) group: The rats received PCM orally by gavage (given in 0.2% gum tragacanth) \([10]\) in the dose of 1 g/kg once \([11]\).

iii. NS and PCM-treated (NS/P) group: The rats received NS oil in the dose of 2 mL/kg/day \([12]\) for 4 weeks, and at the end of this NS-treatment duration, they received the same above dose of PCM.

iv. SH- and PCM-treated (H/P) group: The rats received SH (dissolved in distilled water) in the dose of 1 g/kg/day \([6]\) for 4 weeks, and at the end of this SH-treatment, they received the above dose of PCM.

v. SH-, NS- & PCM-treated (H&NS/P) group: The rats received orally SH and NS in the same above doses for 4 weeks, and at the end of the duration of treatments, they received the above dose of PCM.

4. Experimental procedures: The duration of SH and or NS treatment was four weeks. The drugs were administered orally with an intragastric tube (using a Portex 4FG cannula, Portex Ltd., Hythe, UK). At the end of the experiment period (4 weeks), the rats underwent the following procedures:

i. Collection of blood samples and separation of serum: After finishing the experiment, the rats were fasted overnight and then held in a glass chamber to be anesthetized with diethyl ether. The venous blood samples were collected by heparinized microcapillary tubes from the retro-orbital plexus (24 hours after the last drugs administration). The samples were incubated at 37°C until blood clotted and then centrifuged (5000 g, 10 min) for separation of serum which was stored at -20°C till used for biochemical assays as described below.

ii. Tissue sampling: After collection of blood samples, rats were then sacrificed and their livers were excised, weighed and rapidly washed in cold normal saline and then kept in ice-cold isotonic potassium chloride solution \((1.15% \text{ KCl w/v})\) containing 0.1 mM EDTA. The livers were then divided into 5 volumes of
50 mM phosphate buffer (pH= 7.4) and homogenized by a homogenizer fitted with a Teflon pestle. The homogenate was then centrifuged for 10 min at 3000 g, the lipid layer was removed and the resulting supernatant was further centrifuged for 60 min at 15,000 g at 4°C. The supernatant was stored at -80°C till use [13].

iii. Biochemical assays: The serum were used to measure aspartate transaminase (AST), alanine transaminase (ALT), blood urea nitrogen (BUN), creatinine, total antioxidative capacity (TAC) and Fas ligand (Fas L). Also, the TAC in liver homogenate was investigated. The determination of the values of the investigated parameters was achieved with an automated analyzer (Hitachi, Japan) and commercial kits (Technichon, Germany) [14].

iv. FasL assay: The protein levels of FasL in the serum were measured by dot blot assay. Serum samples (1 µl) were put on nitrocellulose (NC) paper and were blocked using the blocking buffer [5% non-fatty milk in TBST (a mixture of Tris-Buffered Saline and Tween 20) (10 mm Tris–HCl pH 8.0, 0.15 mm NaCl, 0.05% Tween 20] at room temperature for 1 hour. The NC paper was then incubated with primary antibody (FasL) for 1 hour at room temperature. After 3 washes in the TBST buffer, the NC paper was transferred to a secondary antibody (anti-rabbit IgG alkaline phosphatase conjugate) for 1 h at room temperature. After 3 washes in phosphate-buffered saline (PBS), the NC paper was developed with an alkaline phosphatase substrate (Sigma) for 10 min in the dark. Several concentrations of rat recombinant FasL as described above were used as standards. The target dots were analyzed and quantified densitometrically using a GS-700 Imaging Densitometer (BioRad). The experiments were repeated more than twice [15].

v. The total antioxidant capacity (TAC) assay in the serum and tissue samples was assayed by commercially available kits (Randox labs, Grumlin, UK). The assay principle was based on the ability of antioxidants to quench the absorbance of the radical cation that is formed by the reaction of a chromogen with the peroxide and H2O2 [16].

5. Statistical analysis: The recorded parameters were expressed as means and standard error of means (mean ± SEM) for all groups and statistically analyzed using SPSS (version 16) software, for the one-way analysis of variance (one-way ANOVA) followed by Post Hoc and least significant difference (LSD) tests. The level of acceptance for statistical significance was considered to be P < 0.05.

RESULTS

1. The effects of SHDP, NS/P, H/P and H&NS/P treatments on the serum AST and ALT in rats (Table 1 and Figure 1): Single high oral PCM dose increased significantly (P < 0.001) the serum ALT and AST in SHDP-treated group when compared to control group. NS, H, or combined H/NS treatments, when administered before PCM, decreased significantly (P < 0.001) the PCM-induced elevations in the serum ALT and AST in the NS/P-treated, H/P-treated and H&NS/P-treated rats when compared to SHDP-treated rats. While, the combined H/NS treatment more effectively decreased these liver enzymes to nearly the control ranges.

Table 1: The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on the serum AST (s.AST) and ALT (s.ALT) in rats

<table>
<thead>
<tr>
<th>Animal group</th>
<th>s.AST (U/L)</th>
<th>s.ALT (U/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group</td>
<td>44.63 ± 0.26</td>
<td>39 ± 0.46</td>
</tr>
<tr>
<td>SHDP-treated group</td>
<td>94.38 ± 1.28#</td>
<td>61.38 ± 0.38#</td>
</tr>
<tr>
<td>NS/P-treated group</td>
<td>81 ± 0.46 # $</td>
<td>57.25 ± 0.88 # $</td>
</tr>
<tr>
<td>H/P-treated group</td>
<td>55.13 ± 0.4 # $</td>
<td>45.5 ± 0.93 # $</td>
</tr>
<tr>
<td>H&amp;NS/P-treated group</td>
<td>45.25 ± 0.37 $</td>
<td>39.75 ± 0.49 $</td>
</tr>
</tbody>
</table>

- Values are expressed as mean ± SEM.
# : Significant in comparison to control group.
$: Significant in comparison to SHDP-treated group.
Figure 1: The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on the serum AST and ALT in rats

2. The effects of SHDP, NS/P, H/P and H&NS/P treatments on the serum BUN and creatinine in rats (Table 2 and Figures 2 & 3): Single high oral PCM dose increased significantly ($P<0.001$) the serum BUN and creatinine in the SHDP-treated rats when compared to control group. NS, H, or combined H/NS treatments, when administered prior to PCM, decreased significantly ($P<0.001$) the PCM-induced elevations in these parameters in the NS/P-treated, H/P-treated and H&NS/P-treated rats when compared to SHDP-treated rats. Furthermore, the combined H/NS treatment resulted in more effective decrements in these parameters to be nearly equal to the control ranges.

Table 2: The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on the serum BUN and creatinine in rats

<table>
<thead>
<tr>
<th>Animal group</th>
<th>s.BUN (mg/dl)</th>
<th>s.Creatinine (mg/dl)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group</td>
<td>18.36 ± 0.57</td>
<td>0.64 ± 0.018</td>
</tr>
<tr>
<td>SHDP-treated group</td>
<td>42.13 ± 0.93 #</td>
<td>1.12 ± 0.043 #</td>
</tr>
<tr>
<td>NS/P-treated group</td>
<td>34.75 ± 0.59 # $</td>
<td>0.79 ± 0.016 # $</td>
</tr>
<tr>
<td>H/P-treated group</td>
<td>27.38 ± 0.42 # $</td>
<td>0.71 ± 0.009 # $</td>
</tr>
<tr>
<td>H&amp;NS/P-treated group</td>
<td>19.38 ± 0.41 $</td>
<td>0.65 ± 0.006 $</td>
</tr>
</tbody>
</table>

- Values are expressed as mean ± SEM.
- #: Significant in comparison to control group.
- $$: Significant in comparison to SHDP-treated group.

Figure 2: The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on the serum BUN in rats
3. The effects of SHDP, NS/P, H/P and H&NS/P treatments on the serum TAC and tissue TAC in rats (Table 3 and Figures 4 & 5): Single high oral PCM dose decreased significantly ($P < 0.001$) the serum TAC and tissue TAC in SHDP-treated group when compared to control group. NS, H, or combined H/NS treatments when administered before PCM, increased significantly ($P < 0.001$) the PCM-induced decreases in these parameters in the NS/P-treated, H/P-treated and H&NS/P-treated rats when compared to the SHDP-treated rats. In addition, the combined H/NS treatment more effectively decreased the serum TAC and liver tissue TAC to be nearly equal to the control ranges.

**Table 3:** The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on the serum and tissue TAC in rats

<table>
<thead>
<tr>
<th>Animal group</th>
<th>s.TAC (µmol/L)</th>
<th>t.TAC (nmol/mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group</td>
<td>2.12 ± 0.022</td>
<td>0.48 ± 0.005</td>
</tr>
<tr>
<td>SHDP-treated group</td>
<td>1.4 ± 0.005 #</td>
<td>0.19 ± 0.004 #</td>
</tr>
<tr>
<td>NS/P-treated group</td>
<td>1.6 ± 0.006 # $</td>
<td>0.4 ± 0.004 # $</td>
</tr>
<tr>
<td>H/P-treated group</td>
<td>1.82 ± 0.005 # $</td>
<td>0.46 ± 0.003 # $</td>
</tr>
<tr>
<td>H&amp;NS/P-treated group</td>
<td>2.03 ± 0.02 # $</td>
<td>0.48 ± 0.004 $</td>
</tr>
</tbody>
</table>

- Values are expressed as mean ± SEM.
- #: Significant in comparison to control group.
- $$: Significant in comparison to SHDP-treated group.

**Figure 4:** The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on the serum TAC in rats
Figure 5: The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on tissue TAC in rats

4. The effects of SHDP, NS/P, H/P and H&NS/P treatments on the serum FasL in rats (Table 4 and Figure 6): Single high oral PCM dose increased significantly ($P < 0.001$) the serum FasL in SHDP-treated group when compared to control group. NS, H, or combined H/NS treatments when administered before PCM, decreased significantly ($P < 0.001$) the serum FasL in the NS/P-treated, H/P-treated and H&NS/P-treated rats when compared to the SHDP-treated rats. In addition, the combined H/NS treatment more effectively decreased the serum FasL more than the decrease by either drug alone.

Table 4: The effects of SHDP, NS/P, H/P and H&NS/P treatments on the serum Fas L in rats

<table>
<thead>
<tr>
<th>Animal group</th>
<th>s.Fas L (ng/ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control group</td>
<td>3.28 ± 0.045</td>
</tr>
<tr>
<td>SHDP-treated group</td>
<td>25.93 ± 0.25 #</td>
</tr>
<tr>
<td>NS/P-treated group</td>
<td>15.33 ± 0.11 # $</td>
</tr>
<tr>
<td>H/P-treated group</td>
<td>10.41 ± 0.11 # $</td>
</tr>
<tr>
<td>H&amp;NS/P-treated group</td>
<td>3.84 ± 0.08 # $</td>
</tr>
</tbody>
</table>

- Values are expressed as mean ± SEM.
- #: Significant in comparison to control group.
- $$: Significant in comparison to SHDP-treated group.

Figure 6: The effects of SHDP, NS/P, H/P and H&NS/P treatments orally for 4 weeks on serum Fas L in rats
DISCUSSION

The studies for treating the toxic liver injury lead us to perform many investigations regarding the hepato- and nephrotoxicities induced by different drugs and the possible hepato- and nephro-protective effects of therapeutic strategies from the alternative or complementary medicine. The present work was designed to investigate effects of Sidr honey (SH) and Nigella sativa oil (NS) on paracetamol (PCM)-induced hepato-renal toxicity in rats. In this study, single high paracetamol (PCM) dose caused significant elevations in the serum ALT and AST liver enzymes, and the serum BUN, creatinine and Fas L, and caused significant reductions in the serum TAC and liver tissue TAC in SHDP-treated rats. While NS or SH treatment, when given before PCM, prevented the PCM-induced increases in the serum ALT, AST, BUN, creatinine and Fas L, and prevented the PCM-induced decreases in the serum TAC and tissue TAC in SHDP-treated rats. The combined NS and H treatment decreased the liver enzymes, improved the renal functions, increased the tissue TAC to nearly the control rats ranges, and, they decreased the serum Fas L more than the decrease done by either drug alone.

At the outset, liver and renal toxicities could be experimentally-examined by administering the compound at increasing doses, in presence of metabolic inducers or inhibitors, with depletion of protective systems, or similarly by co-administering the drug with a known toxic substance. First, Lee and Kass stated that cellular organelles with their functions are usually the primary targets of liver toxicity. Likewise, Jaeschke and Bajt explained that reactive metabolite formation, protein alkylation, and antioxidant depletion are the general mechanisms of hepatotoxicity. In the same line, necrotic death occurs following antioxidant depletion and oxidation of intracellular proteins, which lead to increased mitochondrial membrane permeability, decreased ATP synthesis, inhibition of Ca\(^{2+}\) dependent ATPase and decreased ability to sequester Ca\(^{2+}\) in mitochondria. The main intracellular mechanisms that lead to apoptosis are due to activation of nucleases with energy participation of mitochondria. In the same way, Grattagliao et al. reported that intracellularly-generated signaling can activate B-cell/lymphoma 2 (Bcl-2) members which promotes the release of intramembranous proteins, chromatin condensation and DNA fragmentation through formation of pores in the outer mitochondrial membrane. Reactive oxygen species (ROS) and peroxynitrite formation, triggering decreased membrane permeability and membrane potential with decreased production of energy are the causative mechanisms for mitochondrial dysfunction. The deleterious effects of paracetamol on liver and kidney in the present study are in accordance with many authors who reported that paracetamol is predominantly metabolized by conjugation. Only a small amount is metabolized by CYP2E1 to its toxic metabolite NAPQI which is detoxified through binding with GSH. If the total amount of paracetamol reaching the liver exceeds 12-15 g, the conjugating capacity of the liver is exceeded and unfortunately, the remaining unbound NAPQI covalently binds to cellular and mitochondrial proteins, leading to necrotic cell death. Thus, in the presence of CYP2E1 hypertrophy and/or decreased GSH availability (for examples as during malnutrition, chronic alcoholism, and prolonged intake of barbiturates), NAPQI formation is increased even at therapeutic doses, to overwhelm the stores of GSH, and then it may result in severe liver damage.

Furthermore, Naguib et al. reported that acetaminophen-induced toxicity is the predominant cause of liver and kidney failure. In humans and rodents, the toxicity starts with the reactive metabolite that binds to proteins, leading to dysfunction of the mitochondria and DNA fragmentation in the nucleus resulting in death of the cell. In addition, Knight et al. clarified that the necrotic process starts with disturbed Ca\(^{2+}\) homeostasis with increased cytosolic Ca\(^{2+}\) levels, oxidative changes in mitochondria and accumulation of oxidized GSH and peroxynitrite. Then induction of membrane permeability, with drop of mitochondrial membrane potential, no synthesis of ATP, and release of mitochondrial proteins, cytochrome C and endonucleases will happened. Also as showed by Watkins and
Seeff [23], the deficiency of ATP prevents caspase activation but induces DNA damage, and activates intracellular proteases that lead to liver cell membrane rupture and necrosis. Grattagliano et al. [1] referred the liver cell death and failure observed after paracetamol poisoning to these intracellular events.

It might be argued that the intrinsic properties of the drug, its dose, its metabolites and the local O2 supply are the factors which detect the cellular and intracellular targets. These targets of drug-induced hepatocellular injury include non-parenchymal hepatic cells, mitochondria, and nuclear receptors. Besides, the release of inflammatory mediators and reactive oxygen species (ROS) may be due to activation of Kupffer cells. Also, administration of TNF-α antagonists or inhibition of macrophage activation protects liver cells against paracetamol toxicity [1]. Correspondingly, Kaplowitz [17] and Jaeschke and Bajt [20] found that the drug-induced hepatotoxicity is dose-dependent. Apoptosis and necrosis initially may follow common metabolic pathways. Apoptosis occurs when hepatocyte injury affects the maintenance of cell functions. Necrosis generally begins at the cytoplasm and involves mitochondria. Hence, drug-induced toxicity results mainly from the reactive metabolite formation, then depletion of GSH and alkylation of proteins with mitochondrial dysfunction. Also, the essential steps in hepatocyte death are opening of pores in the outer mitochondrial membrane, release of proteins and cytochrome C, and intracellular imbalance of Ca2+ homeostasis, and accumulation of Na+ [20]. Therefore, detoxification of ROS and maintenance of membrane protein sulfhydryls, including the ATP synthase and the Ca2+-dependent ATPase depends mainly, as confirmed by Lauterburg [4], on the maintenance of mitochondrial GSH pool.

Thus from this concept, our results are in accordance with the reports of many authors [1,17] who found that the drug-induced hepatotoxicity is dose-dependent. The hepatocytes and other cells in the liver with the cellular organelles and their functions are the primary targets of hepatotoxicity, and necrosis is the mechanisms of drug-induced liver injury [19].

In another perspective, Ahmad et al. [9] stated that the N. sativa seeds and their oil are used in the treatment of various diseases. So it is recommended for use on regular basis in Tibbe-Nabwi (Prophetic Medicine) [24]. Indeed, Goreja [25] explained that N. sativa was showed to possess wide range of activities as gastroprotective, hepatoprotective, anti diabetic, anticancer, immunomodulatory, analgesic, antimicrobial, anti-gastritis and anti-inflammatory, bronchodilator, nephroprotective and antioxidant properties. Likewise, as stated by Abel-Salam [8] and Aggarwal and Kunnammakara [26], N. sativa seeds pharmacologically showed to be used widely in the treatment of various diseases like bronchitis, asthma, diarrhea, and rheumatism and skin disorders. It is also used as liver tonic, diuretic, digestive, appetite stimulant, and to support immune system. Furthermore, Khazdair [27] showed that N. sativa has been widely used in treatment of various CNS disorders such as Alzheimer disease, epilepsy and neurotoxicity. Ahmad et al. [9] referred most of its therapeutic properties to thymoquinone (TQ) which is the major active ingredient of the essential oil. TQ, also the methanol extracts of the shoots roots and seeds inhibited the oxidative stress and exhibited antioxidant activity as evidenced by normalizing glutathione (GSH), catalase (CAT), SOD and NO [28-30]. Also, in agreement with our study, Zaheer et al. [31] reported that N. sativa treatment protects the liver in rat against hepatic ischemia reperfusion injury through improving the serum AST, ALT and lactate dehydrogenase (LDH) levels, total oxidative status (TOS), and oxidative stress index (OSI). In addition, Saleem et al. [32] showed that TQ has a protective role on cadmium-induced hepatotoxicity and induces modulatory effect on the antioxidant defense system. Moreover, Ahmad et al. [9] found that both vitamin C and N. sativa oil produced nephro-protective effect as evidenced from lowering the values of serum creatinine, blood urea nitrogen (BUN), and increasing the antioxidant activity as indicators of gentamicin-associated nephrotoxicity in rabbits, and these two antioxidants combination proved to have synergistic nephroprotective effect.
Furthermore, Abul-Nasr et al. [33] revealed that N. sativa oil, via reducing serum urea and creatinine levels and increasing TAC levels in kidney tissue and blood, protected against methotrexate-induced nephrotoxicity, and also against renal ischemia-perfusion injury in rat kidneys. Also, oral treatment of N. sativa oil by different doses, in gentamicin (GM)-induced nephrotoxicity in rats, ameliorated the biochemical and histological indicators of GM-induced kidney toxicity; increased plasma total antioxidant status (TAS) and reduced renal cortex GSH concentrations. Hence, as reported by Yildiz et al. [34], in rats, N. sativa oil prevents GM-induced acute nephro-toxicity. TQ stimulated the resistance to oxidative stress. Kim et al. [35] stated that TQ elevated the lowering of mitochondrial transmembrane potential ($\Delta \psi _{M}$) and attenuated the elevated cytosolic Ca$^{2+}$ caused by drug toxicity and activated the release of mitochondrial cytochrome-c, increased the expression of Bax and decreased the expression of an anti-apoptotic protein (Bcl-2). Cherian et al. [36] and Peng et al. [37] added that TQ stabilizes mitochondrial membrane potential and inhibits apoptotic cascade by decreasing DNA damage. Indeed, it has been reported that TQ exhibits antioxidant, anti-inflammatory and anticancer activities (mediated via peroxisome proliferator-activated receptor gamma, p53-dependent and p53-independent pathways), against many types of malignancy, with minimal toxicity in normal cells [38–41]. In addition, it has been mentioned that the pharmacological effects of N. Sativa are referred to its strong antioxidant effect which opposes the mitochondrial dysfunction and oxidative stresses associated with drug overdose, and also to antagonizing the free radical-generating agents by reducing reactive oxygen species (ROS), and maintaining the mitochondrial integrity [42–44].

In another perspective, Alvarez-Suarez et al. [45] reported that Aasal, the Arabic name for honey, is a naturally sweet product produced by honeybees from the nectar of blossoms or from the exudates of trees and plants giving the nectar honey. Again, it is a natural, unprocessed and easily digested food and has been shown to have many nutritional and biological effects which include antibacterial, antioxidant, antiviral, antiparasitic, anti-inflammatory, anticancer, and immunosuppressive activities. Its antioxidant effects, as confirmed by Jaganathan et al. [46], may be the key in understanding its preventive effect on paracetamol induced hepato-renal toxicity. In the same way it is used for the treatment of burns, faster healing of wound, asthma, and gastrointestinal, skin and eye diseases. Honey is considered to be the first in the line to treat jaundice in traditional medicine of different countries [47]. Besides, Chow [48] explained the beneficial composition of honey; it is composed primarily of fructose and glucose but also contains fructo-oligosaccharides and many amino acids, vitamins, minerals, flavonoids phenolic acids, ascorbic acid, tocopherols, catalase (CAT), superoxide dismutase (SOD), reduced glutathione (GSH), Millard reaction products, peptides and enzymes. Hence, in accordance to our study as mentioned by Ahmad et al. [49], honey has been shown to prevent the reactive oxygen species (ROS)-induced oxidation of low-density lipoprotein (LDL). Also, Erejuwa et al. [50] reported that honey regulates cell cycle hence it can oppose the toxic effect of drug on cell mitochondria and stops the cascade of depletion of GSH pool in hepatic and renal cell mitochondria.

CONCLUSIONS

The current study indicated that single high paracetamol dose caused a liver toxicity; elevating the serum ALT and AST enzymes, and a kidney toxicity; elevating the serum BUN and creatinine and Fas L, with reductions in the serum and tissue total antioxidant capacity (TAC). Also, this study confirmed that combination of N sativa oil and Sidr honey protect against these hepatic and renal tissues changes via through their strong antioxidant effects opposing the oxidative damages accompanying paracetamol-induced hepatonephrotoxicity in rats, and thus, could be used as an effective protector against these paracetamol-induced liver and kidney toxicities. Our present issue included the use of sider honey and Nigella sativa oil combination, a strategy directed at factors that cause liver and kidney damage.

More researches in this area are recommended to investigate the therapeutic effects of this
natural combination and also to evaluate the prophylactic effect of them on other drug-induced toxicities in other organs. It is of prior importance to carry out experiments to ensure their prophylactic and therapeutic effects in humans.

**ABBREVIATIONS**


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