Protective effect of thymoquinone, the active constituent of *Nigella sativa* fixed oil, against ethanol toxicity in rats

Sayed Masoud Hosseini 1, Elahe Taghiabadi 2, Khalil Abnous 3, Alireza Timcheh Hariri 4, Hamed Pourbakhsh 5, Hossein Hosseiniazadeh 6*

1 Department of Pharmacodynamics and Toxicology, School of Pharmacy, Shahid Beheshti University of Medical Sciences, Tehran, Iran
2 Department of Pharmacodynamics and Toxicology, School of Pharmacy, Mashhad University of Medical Sciences, Mashhad, Iran
3 Pharmaceutical Research Center, Department of Medicinal Chemistry, School of Pharmacy, Mashhad University of Medical Sciences, Mashhad, Iran
4 Medical Toxicology Research Center, Faculty of Medicine, Mashhad University of Medical Sciences, Mashhad, Iran
5 Food Control Laboratory, Food and Drug Administration, Shiraz University of Medical Sciences, Shiraz, Iran
6 Pharmaceutical Research Center, Department of Pharmacodynamics and Toxicology, School of Pharmacy, Mashhad University of Medical Sciences, Mashhad, Iran

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**ABSTRACT**

Objectives: Long term consumption of ethanol may induce damage to many organs. Ethanol induces its noxious effects through reactive oxygen species production, and lipid peroxidation and apoptosis induction in different tissues and cell types. Previous experiments have indicated the antioxidant characteristics of thymoquinone, the active constituent of *Nigella sativa* fixed oil, against biologically dangerous reactive oxygen species. This experiment was planned to evaluate the protective effect of thymoquinone against subchronic ethanol toxicity in rats.

**Materials and Methods:** Experiments were performed on six groups. Each group consisted of six animals, including control group (saline, gavage), ethanol-receiving group (3 g/kg/day, gavage), thymoquinone (2.5, 5, 10 mg/Kg/day, intraperitoneally (IP)) plus ethanol and thymoquinone (10 mg/Kg/day, IP) groups. Treatments were carried out in four weeks.

**Results:** Thymoquinone reduced the ethanol-induced increase in the lipid peroxidation and severity of histopathological alteration in liver and kidney tissues. In addition it improved the levels of proinflammatory cytokines in liver tissue. Furthermore, thymoquinone corrected the liver enzymes level including alanine transaminase, aspartate transaminase and alkaline phosphatase in serum and glutathione content in liver and kidney tissues. Other experiments such as Western blot analysis and quantitative real-time RT-PCR revealed that thymoquinone suppressed the expression of Bax/Bcl ratio (both protein and mRNA level), and caspases activation pursuant to ethanol toxicity.

**Conclusion:** This study indicates that thymoquinone may have preventive effects against ethanol toxicity in the liver and kidney tissue through reduction in lipid peroxidation and inflammation, and also interrupting apoptosis.

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**Introduction**

Ethanol has played an important role in medical practice and alcoholic beverages include wines, beers and spirits (1). Hence ethanol is a widely consumed organic solvent with toxic properties both systemically and to the central nervous system (CNS). Its toxic properties can be expressed following an acute overdose or after extended chronic consumption. A relatively high LD50 value of ethanol, suggests that this agent acts relatively non-specifically on a widespread range of cellular targets rather than at a single critical site. Therefore, the noxious effects of ethanol toxicity perhaps represent a summation of a series of adverse metabolic modulations (2). Ethanol toxicity in different tissues subsequent to acute and chronic consumption has been proved in human and animals. Ethanol changes fat and carbohydrate metabolism. Hepatic lipoprotein release is decreased, while hepatic *de novo* lipogenesis, mobilization of fat stores for use by peripheral tissues, and hepatic absorption of circulating lipids are increased. The changes in NADH/NAD+ ratio prevent the function of the Krebs cycle and decelerate fatty acid oxidation. These effects increase serum triglycerides and cause steatosis (fatty liver), a condition in which triglycerides accumulate in hepatocytes (3). Liver disorders such as adipose infiltration, alcoholic hepatitis, and fibrosis are associated with ethanol consumption (4). Ethanol can...

*Corresponding author: Hossein Hosseiniazadeh. Pharmaceutical Research Center, Department of Pharmacodynamics and Toxicology, School of Pharmacy, Mashhad University of Medical Sciences, Mashhad, Iran. Tel: +98-51-38819042; Fax: +98-51-38823251; email: hosseiniazadeh@sums.ac.ir*
cause hepatotoxicity and nephrotoxicity as indicated by histopathological changes in addition to increased levels of liver enzyme like alanine transaminase (ALT), aspartate transaminase (AST) and alkaline phosphatase (ALP). Some investigation demonstrated that the levels of malondialdehyde (MDA) and inflammatory mediators such as tumor necrosis factor-α (TNF-α) and interleukin-6 (IL-6) were significantly increased in the liver and kidney of ethanol treated rats. In addition glutathione (GSH) content reduction in such tissues was reported (5, 6).

It was shown that ethanol causes toxic effects by production of reactive oxygen species (ROS) and lipid peroxidation induction in different tissues and cells. It was indicated that excess microsomal ROS production has been started during ethanol metabolism. By means of spin-trapping techniques, it was shown that the hydroxyl ethyl radical, an active and relatively short-lived oxidant species, is the main deleterious species. This radical is not as reactive as the hydroxyl and thereafter can diffuse toward, and impair more distant cells and target molecules. In addition, ROS activity could be monitored via an intracellular fluorescent probe produced by the oxidation of a non-fluorescent precursor. Using this probe (2',7'-dichlorodihydrofluorescein), the evidence of ethanol-induced hepatic and cerebral ROS production was found (2). Besides to oxidative stress, ethanol can stimulate apoptosis by initiation of both intrinsic and extrinsic signaling pathway (5, 6). It is pointed that oxidative stress and subsequent depletion of antioxidant defense of cells and tissues may lead to apoptotic cell death. So that this procedure can be blocked by antioxidant agents (7).

Antioxidants have a fundamental role in preventing free radical induced damage through scavenging them. Thymoquinone (TQ) (2-isopropyl-5-methyl-1,4-benzquinone) is the foremost component of the volatile oil of *Nigella sativa* seeds. The black seed of *N. sativa* L., family Ranunculaceae, comprises a fixed oil (>30%) and a volatile oil (0.40%-0.45%), which contains of 18.4% to 24% TQ. Previous studies suggest that TQ or *N. sativa* oil might have different pharmacological actions including protective effects in metabolic syndrome, chronic neuropathic pain, ischemia-reperfusion induced kidney and skeletal muscle damage, and epileptic seizures (8-15). In previous studies, it was found that *N. sativa* fixed oil (NOS) has neuroprotective, antimicrobial, antibacterial, antitussive, anticonvulsant, analgesic and anti-inflammatory effect, and also has protective effects against ethanol induced toxicity in liver and kidney through lessening the oxidative degradation of lipids (lipid peroxidation), increasing antioxidant capacity (glutathione levels), reducing some liver enzyme levels like ALT, AST, and ALP, and some inflammatory cytokine levels (IL-6), attenuating histopathological changes, and also inhibiting apoptosis (16-23). In 1960 after the first extraction of TQ, the active constituent of NOS, it was investigated for its antioxidant, anti-inflammatory and anticancer activities in both in vitro and in vivo models. Its antioxidant/anti-inflammatory effects have been described in numerous disease models, including encephalomyelitis, diabetes, asthma, carcinogenesis, and other conditions such as acetaminophen induced hepatotoxicity or paraparabindolence induced lung fibrosis (24-26). According to the Ismail et al. results (27), thymoquinone rich fraction (TQRF) and TQ at dosages ranging from 0.5 to 1.5 g/kg and 20 to 100 mg/kg body weight, respectively, for 8 weeks demonstrated considerable inhibitory activity against the OH⁺ development in comparison to untreated rats. In line with that, Kanter et al. (28) have reported that TQ could partially protect gastric mucosa against mucosal injury induced by acute ethanol exposure, and these gastroprotective effects may be due to their antiperoxidative and antioxidant effects. A few studies have also shown that TQ may have protective effects against lipid peroxidation process (7). El Mezyen et al. (29) implied that TQ may have an anti-inflammatory effect during the allergic response in the lung by inhibition of prostaglandin D₂ synthesis and Th2-driven immune response. Recent investigation demonstrated the protective effect of metformin and TQ against ethanol-induced neuronal apoptosis in primary rat cortical neurons, and also against acrylamide-induced neurotoxicity in Wistar rats (30, 31).

Hence, this study was designed to evaluate the protective effect of TQ, the active constituent of *N. sativa*, against toxic effects of ethanol in rats.

### Materials and Methods

#### Chemicals

Thymoquinone (2-isopropyl-5-methyl-1,4-benzquinone) purchased from Sigma, St Louis, Mo, absolute ethanol, malondialdehyde, thiobarbituric acid (TBA), sodium dodecyl sulphate (SDS),β-mercaptoethanol (β-ME), and N,N,N-Tetramethyl-ethylenediamine (TEMED) were purchased from Merck. Complete protease inhibitor cocktail, Phenyl-methanesulfonyl fluoride (PMSF), and reduced glutathione were provided by Sigma-Aldrich Chemical Company. Rabbit monoclonal antibody for detection of caspase-3, caspase-8, caspase-9, Bcl-2, and Bax protein, and also mouse monoclonal β-actin antibody, anti-rabbit secondary antibody, and anti-mouse secondary antibody were obtained from Cell Signaling. Coomassie (Bradford) Protein Assay Kit and polyvinylidene difluoride (PVDF) membrane were provided by Bio-Rad. Enhanced chemiluminescence Western Blotting Substrate (ECL) and BCA protein assay kit were purchased from Pierce. Express one-step SYBR R Green ER™kit and Rat ELISA Kit for IL-6 and TNF-α detection were obtained from Invitrogen.
**Animals**

Thirty six adult male Wistar rats (weight 200-250 g) were provided by Animal Center of School of Pharmacy, Mashhad University of Medical Sciences. Rats were housed in plastic cages, on a 12 hr light/dark cycle and kept in a conditioned atmosphere at 25 °C and fed standard laboratory pellets with tap water *ad libitum* throughout the experimental period. All experimental procedures were carried out in accordance with Mashhad University of Medical Sciences, Ethical Committee Acts.

**Experimental design**

The rats were randomly assigned into six groups each consisted of six rats. The first group received normal saline orally by gavage once for four weeks, the second group was administered ethanol 3 g/kg/day, the third, fourth, and fifth groups were treated with ethanol 3 g/kg/day plus TQ at 2.5, 5 and 10 mg/kg/day doses respectively, the sixth group received TQ 10 mg/kg/day. Ethanol was diluted in normal saline to a final concentration of 40% (v/v) and was administered orally by gavage for four weeks. TQ was dissolved in 0.8% Tween 80 (in normal saline) and the solution was prepared fresh just before intraperitoneal administration. The ethanol and TQ doses were determined according to the results of pilot test (the data are not shown).

After four weeks, fasting rats were killed by decapitation. Immediately after decapitation, blood samples were collected in dry tubes and allowed to coagulate. Clotted blood samples were subjected to centrifuge at 2000×g for 15 min to segregate the serum that were stored at -80 °C for biochemical analysis. The liver and kidney tissues were immediately dissected out and washed in normal saline. Quadrade lobe of liver and left kidney preserved in 10% neutral buffered formalin for histopathology examination. Three other lobes of liver and right kidney were kept at -80 °C for several other analyses.

**Biochemical blood tests**

Serum levels of some liver enzyme like AST, ALT and ALP were assessed employing commercial colorimetric kits.

**Inflammatory biomarker measurement**

The inflammatory biomarkers like TNF-α and IL-6 were evaluated in the liver tissue employing Invitrogen Elisa kits.

**Assessment of lipid peroxidation in the liver and kidney tissues**

The extent of oxidative stress was evaluated by measuring malondialdehyde (lipid peroxidation biomarker) concentration using spectrophotometry. For this purpose the liver and kidney tissues from different groups were homogenized for 2 min at 4 °C in 1.15% KCl in order to provide a 10% homogenate (w/v). Homogenates were subjected to centrifuge (Hettich, Germany) at 3000 g for 10 min to acquire supernatants. MDA levels were determined in accordance with the method of Niehaus and Samuelsson (32). Briefly, 0.5 ml of the sample was mixed with 3 ml of 1% phosphoric acid and 1 ml of 0.6% TBA solution and the mixture was incubated for 45 min in a boiling water bath. The tested tubes were cooled at room temperature. Then, 4 ml of n-butanol was added, and vortex-mixed for 1 min followed by centrifugation at 3000 g for 10 min. Butanol phase (supernatant) was transferred to a fresh tube and its absorbance measured at 532 nm. Tissue MDA content was expressed as nmol/g tissue.

**Measurement of reduced glutathione (GSH) content in the liver and kidney tissues**

Hepatic and nephrotic antioxidant capacity were assessed through GSH content measurement in accordance with the method of Moron et al (33). To obtain 10% homogenate (w/v), the liver and kidney tissues were homogenized in ice cold phosphate buffered saline (PBS, pH 7.4). Homogenates were centrifuged at 3000 g for 10 min. reduced glutathione content was measured in supernatants employing 5,5’-dithiobis (2-nitrobenzoic acid) (DTNB) which produced a yellow-colored 5-thio-2-nitrobenzoic acid (TNB). Briefly, homogenates were immediately precipitated with the equal amount of 10% TCA and the precipitate was removed after centrifugation at 3000 g for 5 min. Then 0.5 ml of 0.04% DTNB reagent was added to 0.5 ml of supernatants plus 2 ml sodium phosphate buffer (0.1 M, pH 8.0). Finally, the absorbance of yellow colored TNB was read at 412 nm using a UV–VIS spectrophotometer. Tissue GSH contents were expressed as nmol/g tissue.

**Histopathological evaluation**

For histopathological examination specimens from liver and kidney were taken from all rat groups after scarification. The tissues were fixed in 10% buffered formalin for at least 24 hr and then embedded in paraffin; sectioned at 6 μm and finally stained with hematoxylin and eosin stain (H&E stain) for microscopical assay by a standard protocol. Histopathological criteria such as severe congestion were determined semi-quantitatively as mild (+), moderate (+++) and severe (++++).

**Western blot analysis**

For Western blot analysis, liver and kidney tissues were homogenized in the homogenization buffer containing 50 mM Tris pH 7.4, 10 mM NaF, 2 mM EDTA, 10 mM β-glycerol phosphate, 1 mM Na$_2$VO$_4$, 0.2% W/V sodium deoxycholate, 1 mM phenylmethylsulfonyl fluoride (PMSF), and complete
protease inhibitor cocktail (Sigma, P8340) employing polytron homogenizer (POLYTRON, PT 10–35, Kinematica, Switzerland) in ice. The lysates were clarified by centrifugation at 4 °C for 15 min at 10,000 g. Protein concentration of the supernatants was determined using the Bradford protein assay kit (Bio-Rad). Levels of Bax, Bcl-2, caspase-3, caspase-8, caspase-9 and β-actin were measured by immune blotting analysis. Briefly, prepared samples were mixed with loading buffer and heated for 8 min at 95 °C, separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis on a 12% gel, and then transferred to polyvinylidene fluoride (PVDF) membrane. The blots were incubated in blocking buffer TBS-T (5% nonfat milk and 0.1% Tween-20 in Tris-buffered saline) for 2 hr at room temperature. The primary antibodies were rabbit monoclonal anti-serum against Bcl-2 (Cell Signaling, #2870), caspase-3 (Cell Signaling, #9665), caspase-8 (Cell Signaling, #4790), caspase-9 (Cell Signaling, #9506) and rabbit polyclonal anti-serum against Bax (Cell Signaling, #2772), and mouse monoclonal anti-serum against β-actin (Cell Signaling, #3700). All antibodies were used at a dilution of 1:1000. Anti-rabbit IgG labeled with horseradish peroxidase (Cell Signaling, #7074) and anti-mouse IgG labeled with horseradish peroxidase (Cell Signaling, #7076) were used as secondary antibodies. The blots were incubated with the primary antibodies at 4 °C overnight and then washed three times for 10 min each in TSB with 0.1% Tween-20. Finally, the membranes were incubated with the corresponding secondary antibody and washed as described above. Protein bands were visualized using an enhanced chemiluminescence (Pierce ECL Western blotting substrate) and Alliance gel doc (Alliance 4.7 Gel doc, UK). UV Tec software (UK) was used to semi-quantify protein bands. All protein bands were normalized against β-actin protein.

**Extraction of RNA and real-time quantitative polymerase chain reaction**

Total RNAs were extracted from frozen liver and kidney tissues using High pure RNA tissue kit (Roche, #12033674001) in accordance with the manufacturer instruction. Nanodrop (NanoDrop™ 2000, USA) was employed to determine the quality (260/280 and 260/230 ratios) and quantity of isolated RNAs and samples were stored at -80 °C until use. Quantitative RT-PCR was performed with step one thermal cycler (ABI) and Express one-step SYBR Green ER™kit (Invitrogen, #11780–200). Primer pairs to measure Bax, Bcl-2 and β-actin mRNA expression, were chosen according to previous Design® software (BioSoft) (Table 1).

**Table 1. Sequences of different primers used for real-time PCR reactions**

| Gene   | Primer     | Sequence               |
|--------|------------|------------------------|
| Bcl-2  | Forward    | 5'-GGTGAGGAAGCTTCTCAGGGA-3' |
|        | Reversed   | 5'-GGTCAGGTACTCAGTCACTCA-3' |
| Bax    | Forward    | 5'-TGCTGATGGCAACTCTACT-3' |
|        | Reversed   | 5'-ATGATGGTCTGATCAGTGG-3' |
| β-actin| Forward    | 5'-GGGAAATCGTGGTGA-3'    |
|        | Reversed   | 5'-GCCGCGATGGCCATCTC-3'  |

Melting curve analysis was done to analyze quality of primers and products. The relative quantitation values of considered genes were normalized against β-actin. Delta-delta CT procedure was used to determine fold increase of genes in compare to control group.

**Statistical analysis**

Data are expressed as mean ± SEM. All statistical comparisons were made by one-way analysis of variance (ANOVA) followed by Tukey–Kramer test. Statistical analysis was performed using GraphPad InStat v. 6.07 (GraphPad Software, Inc, La Jolla, CA, USA). P-values less than 0.05 were considered significant.

**Results**

**Biochemical blood tests**

ALT, AST and ALP were increased significantly in the ethanol treated group in comparison to the control group. TQ prevented ethanol induced increase in ALT, AST and ALP levels (P<0.001) (Figures 1A, 1B and 1C, respectively).

**Inflammatory biomarker measurement**

The ethanol treated group showed significant increase in hepatic levels of IL-6 and TNF-α, when compared with the control group (P<0.001 and P<0.01, respectively). The augmentation of IL-6 and TNF-α levels by ethanol was significantly decreased by TQ (10 mg/kg) (P<0.01 and P<0.05, respectively) (Figures 2A and 2B).

**Assessment of lipid peroxidation and reduced GSH content in the liver and kidney tissues**

Results demonstrated that ethanol increased lipid peroxidation (MDA level) but reduced the antioxidant...
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Figure 1. Effects of ethanol and TQ treatment (4 weeks) on serum levels of ALT (A), AST (B) and ALP (C) in rats. TQ was administered intraperitoneally, once a day. Ethanol was administered via gavage to rats once a day. Data showed as mean±SEM, # Comparison with control, * Comparison with ethanol treated group. *** or ### P<0.001, Tukey-Kramer test, n=6

Figure 2. Effects of ethanol and TQ treatment (4 weeks) on IL-6 (A) and TNF-α (B) levels in the liver. TQ was administered intraperitoneally, once a day. Ethanol was administered via gavage to rats once a day. Data showed as mean±SEM, # Comparison with control, * Comparison with ethanol-treated group. * P<0.05, ** P<0.01 or ## P<0.01 and ### P<0.001, Tukey-Kramer test, n=6

capacity (GSH content) in the liver and kidney of rats (P<0.001), which could be reversed by TQ treatment. MDA levels were significantly decreased in TQ plus ethanol (group 3, 4 and 5) in the liver and kidney (P<0.001) (Figures 3A and 3B, respectively) in comparison to ethanol group (group 2). TQ plus ethanol also significantly augmented the GSH content compared to ethanol treated rats in the liver and kidney (P<0.001) (Figures 4A and 4B, respectively).

Histopathological evaluation

Histopathological alteration of liver and kidney tissues are presented in Table 2, Figure 5 and Figure 6. The control and TQ (10 mg/kg) treated groups (group 1 and group 6 respectively) presented livers with
Table 2. Effect of thymoquinone (TQ) and ethanol (Et) on histopathological changes in liver and kidney tissues of rat after 4 weeks treatment. Histopathological criteria were determined semi-quantitatively from mild (+) to moderate (+++) and severe (+++), n=6

| Groups                  | Liver                                      | Kidney                                      |
|-------------------------|--------------------------------------------|---------------------------------------------|
|                         | Severe congestion (+++), steatosis (++) and | Severe congestion (+++), hematuria (++) and  |
|                         | infiltration of inflammatory focal portal   | infiltration of inflammatory focal adjacent |
|                         | space (+++)                                | glomerular (+++)                            |
| Control                 | normal                                     | normal                                      |
| Ethanol                 | severe congestion (+++), steatosis (++) and | severe congestion (+++), hematuria (++) and  |
|                         | infiltration of inflammatory focal portal   | infiltration of inflammatory focal adjacent |
|                         | space (+++)                                | glomerular (+++)                            |
| Et + TQ (2.5 mg/kg)     | normal                                     | normal                                      |
| Et + TQ (5 mg/kg)       | moderate congestion (+++)                  | moderate congestion (+++)                   |
| Et + TQ (10 mg/kg)      | mild congestion (+)                        | mild congestion (+)                         |

Figure 3. Effects of ethanol and TQ treatment (4 weeks) on MDA level in the liver (A) and kidney (B) in rats. TQ was administered intraperitoneally, once a day. Ethanol was administered via gavage to rats once a day. Data showed as mean ± SEM, # Comparison with control, * Comparison with ethanol-treated group. *** or ### \( P<0.001 \), Tukey-Kramer test, n=6

Figure 4. Effects of ethanol and TQ treatment (4 weeks) on GSH content in the liver (A) and kidney (B) in rats. TQ was administered intraperitoneally, once a day. Ethanol was administered via gavage to rats once a day. Data showed as mean ± SEM, # Comparison with control, * Comparison with ethanol-treated group. *** or ### \( P<0.001 \), Tukey-Kramer test, n=6

normal architecture (Figures 5A and 5G). Histological changes were observed in ethanol treated group (group 2) in the liver. Liver damage caused by ethanol exposure included severe congestion (+++), steatosis (++) and infiltration of inflammatory focal portal space (++) (Table 2 and Figures 5B, 5C). Although these outcomes were also observed in the ethanol plus TQ groups (group 3, 4 and 5), the incidence and severity of histopathological lesions were less than those in the ethanol group (group 2) (Table 2 and Figures 5D, 5E and 5F).

Similarly, all histological features were normal in the control group (group 1) and TQ alone at dose (10 mg/kg) (group 6) in the kidney (Figures 6A and 6G). Kidney tissues from all rats treated with ethanol (group 2) showed extensive histopathological changes,
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Figure 5. (A) Normal liver of control rats. Hematoxylin and eosin, ×100. (B) and (C) Ethanol treated rats show severe congestion, steatosis and infiltration of inflammatory focal portal space. Hematoxylin and eosin, ×100. (D) Liver sections of rats received TQ 2.5 mg/kg plus ethanol showing moderate congestion. Hematoxylin and eosin, ×100. (E) Mild congestion was observed in rats received TQ 5 mg/kg plus ethanol. Hematoxylin and eosin, ×100. (F) and (G) Liver tissues of rats received TQ 10 mg/kg plus ethanol and TQ alone at dose 10 mg/kg were normal. Hematoxylin and eosin, ×100.

Figure 6. (A) Normal kidney of control rats. Hematoxylin and eosin, ×100. (B) Ethanol treated rats show severe congestion and hematuria. Hematoxylin and eosin, ×100. (C) Kidney sections of rats received TQ 2.5 mg/kg plus ethanol showing mild congestion. Hematoxylin and eosin, ×100. (D) Mild congestion was observed in rats received TQ 5 mg/kg plus ethanol. Hematoxylin and eosin, ×100. (E) and (F) Kidney tissues of rats received TQ 10 mg/kg plus ethanol and TQ alone at dose 10 mg/kg were normal. Hematoxylin and eosin, ×100.

characterized by moderate or severe congestion (+++), hematuria (+++) and infiltration of inflammatory focal adjacent glomerular (+++) (Table 2 and Figures 6B and 6C). The kidney lesions were attenuated in all ethanol plus TQ groups in comparison to ethanol treated group (Table 2 and Figure 6D, 6E and 6F).

Western blot analysis

To determine whether TQ exerts its effects by inhibiting apoptosis, we investigated BAX, BCL-2, caspase-3, caspase-8 and caspase-9 protein levels by Western blot analysis. Results showed that ethanol treatment may up-regulated the expression of Bax/Bcl-2 in the liver and kidney (P<0.001) (Figures 7 and 8 respectively). Similarly, expression of caspase-3, caspase-8 and caspase-9 were up-regulated by ethanol in the liver (Figures 9, 11 and 13 respectively) and kidney (Figures 10, 12 and 14 respectively). In contrast, co-treatment of ethanol with TQ (10 mg/kg) (group 5) significantly reduced the ratio of Bax/Bcl-2 in liver and kidney (P<0.001 and P<0.001, respectively) (Figures 7 and 8 respectively). Furthermore, TQ (10 mg/kg) reduced the ethanol induced activation of caspase-3 (19 KDa), and subsequently decreased the protein level of cleaved caspase-3 (17 KDa) in the liver.

Figure 7. Effect of TQ (10 mg/kg) and ethanol on the protein levels of Bax and Bcl-2 in the rat liver tissue. (A) Representative Western blots showing specific bands for Bax, Bcl-2 and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole liver homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean ± SEM. ** P<0.01 vs ethanol treated group, ### P<0.001 vs control group

Figure 8. Effect of TQ (10 mg/kg) and ethanol on the protein levels of Bax and Bcl-2 in the rat kidney tissue. (A) Representative Western blots showing specific bands for Bax, Bcl-2 and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole kidney homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean±SEM. *** P<0.001 vs ethanol treated group, ### P<0.001 vs control group

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Figure 9. Effect of TQ (10 mg/kg) and ethanol on the protein level of caspase-3 (cleaved caspase-3) in the rat liver tissue. (A) Representative Western blots showing specific bands for cleaved caspase-3 (17 KDa) and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole liver homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean±SEM. *** P<0.001 vs ethanol treated group, ### P<0.001 vs control group

Figure 10. Effect of TQ (10 mg/kg) and ethanol on the protein level of caspase-3 (cleaved caspase-3) in the rat kidney tissue. (A) Representative Western blots showing specific bands for cleaved caspase-3 (17 KDa) and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole kidney homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean±SEM. * P<0.05 vs ethanol treated group, # P<0.05 vs control group

Figure 11. Effect of TQ (10 mg/kg) and ethanol on the protein level of caspase-8 (cleaved caspase-8) in the rat liver tissue. (A) Representative Western blots showing specific bands for cleaved caspase-8 (10 KDa) and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole liver homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean±SEM. * P<0.05 vs ethanol treated group, # P<0.05 vs control group

Figure 12. Effect of TQ (10 mg/kg) and ethanol on the protein level of caspase-8 (cleaved caspase-8) in the rat kidney tissue. (A) Representative Western blots showing specific bands for cleaved caspase-8 (10 KDa) and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole kidney homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean±SEM. * P<0.05 vs ethanol treated group, # P<0.05 vs control group

Effects of TQ on BAX/BCL-2 related gene expression in the liver and kidney tissues

To further confirm whether BAX/BCL-2 related genes are induced by ethanol, we inspected the mRNA levels of these genes by quantitative RT-PCR. Our results demonstrated that ethanol may up-regulate Bax/Bcl-2 mRNA expression ratio in the liver and kidney. Bax/Bd-2 mRNA expression ratio was diminished in simultaneous administration of ethanol and TQ (10 mg/kg) in the liver and kidney (P<0.05) (Figures 15A and 15B respectively).
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Figure 13. Effect of TQ (10 mg/kg) and ethanol on the protein level of caspase-9 (cleaved caspase-9) in the rat liver tissue. (A) Representative Western blots showing specific bands for cleaved caspase-9 (35 KDa) and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole liver homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean±SEM. *** P<0.001 vs ethanol treated group, ### P<0.001 vs control group.

Figure 14. Effect of TQ (10 mg/kg) and ethanol on the protein level of caspase-9 (cleaved caspase-9) in the rat kidney tissue. (A) Representative Western blots showing specific bands for cleaved caspase-9 (35 KDa) and β-actin as an internal control. Equal amounts of protein sample (50 μg) obtained from whole kidney homogenate were applied in each lane. (B) Densitometric data of protein analysis. Data are expressed as the mean±SEM. * P<0.05 vs ethanol treated group, # P<0.05 vs control group.

Figure 15. Effect of ethanol and TQ (10 mg/kg) on Bax/ Bcl-2 mRNA expression in the rat liver (A) and kidney (B) after 4 weeks treatment by real time PCR. The transcript level of each sample was normalized against β-actin transcript level. Data are expressed as the mean±SEM. * P<0.05 vs ethanol treated group and ## P<0.01 or # P<0.05 vs control group.

Discussion

Alcohol ingestion is linked with a number of alteration in cell function and the oxidant–antioxidant system (35). Chronic ethanol exposure is associated with constant presence of its byproducts such as acetaldehyde and acetate in the body that cause direct and indirect toxic effects to mammals and humans (35) and induces its toxic effects by generation of reactive oxygen species (ROS) and lipid peroxidation in different tissues and cell types. Furthermore, ethanol can stimulates apoptosis initiation in various cells and tissues (5, 6).

Chronic ingestion of ethanol significantly increased MDA levels (an end-product of lipid peroxidation) in liver and kidney tissues (36). Lipid peroxidation is due to oxidative stress, in which the reciprocal action...
between free radicals of varied origin and cell membrane unsaturated fatty acids occurs. This process proceeds by a free radical chain reaction mechanism and leads to production of some reactive aldehydes, like malondialdehyde (MDA). Thus, the level of MDA is considered as lipid peroxidation biomarker (37). Also chronic ingestion of ethanol significantly decreased glutathione peroxidase activity and GSH content in liver and kidney. It is suggested that reduced GSH, as an antioxidant defense, can protect liver and kidney tissues against oxidative stress through scavenging free radicals, so that oxidative damage decrease the reduced GSH content in tissues (38). Our data indicated that following ethanol administration, hepatic and nephrotic levels of MDA were significantly increased, while hepatic and nephrotic content of reduced GSH was significantly decreased (Figures 3 and 4 respectively). It has been shown that subchronic ethanol exposure provokes oxidative stress damages through free radicals generation and subsequent lipid peroxidation in various tissues like liver and kidney (3). These findings are congruent with other studies showing that ethanol significantly increases MDA and decreases reduced GSH in some tissues (6, 8, 39-41).

The histopathological findings revealed that subchronic ethanol exposure caused severe steatosis, central vein congestion, and infiltration of inflammatory factors in focal portal space in the liver (Figures 5B and 5C). Also, ethanol caused severe congestion, infiltration of inflammatory factors in focal adjacent glomerular, and hematuria in the kidney (Figures 6B and 6C). It has been indicated that ethanol induced fatty liver, necrosis, inflammation, and accumulation of collagenous fibers (6, 7).

Biochemical blood tests like serum AST, ALT and ALP activities measurement are mainly monitored for the assessment of liver damage. Although these biochemical blood tests are not essentially specific, rise in such enzyme activities are associated with active liver damage. It is suggested that ethanol ingestion may induce increase in AST, ALT and ALP activities (6, 7). Subsequent to hepatocyte plasma membrane destruction due to lipid peroxidation caused by ethanol exposure, the cytosolic enzymes released into the circulatory system. As a result of liver injury, ALP generally is excreted through the bile which results in increased levels of ALP in serum (42). Our data indicate that ethanol induced a considerable increase in AST, ALT and ALP activity in rats (Figure 1). These findings imply that increased activity of these enzymes reflects hepatic deterioration and it is likely that ethanol caused such biochemical alterations in liver (43).

TNF-α, a pivotal cytokine involved in inflammation, is produced primarily by Kupffer cells in the liver. The main event in the process of alcohol-induced liver damage is activation of Kupffer cells by gut-derived endotoxin. Activated Kupffer cells yield various mediators, including cytokines, eicosanoids, proteases and oxygen radicals that participate in inflammation, immune responses, and modulation of hepatocyte metabolism. Plasma levels of TNF-α, IL-1 and IL-6 were raised in patients with severe alcoholic hepatitis (44). Inflammatory cytokines such as IL-6, IL-1 and TNF-α together, induce the acute phase response. The acute phase response is a rapid inflammatory response that provides protection against infection, tissue injury (e.g. due to ethanol toxicity), trauma etc. Typically, it is associated with an increase in inflammatory factors (such as pro-inflammatory cytokines like TNF-α and IL-6) (45). Our data implicate that ethanol treated group shows an increase in IL-6 and TNF-α levels compared to control group (Figure 2). These results were in agreement with other experiments that demonstrated increased levels of TNF-α and IL-6 due to ethanol exposure in some tissues of rats (6, 7).

Intracellular reactive oxygen species and increased levels of lipid peroxidation products have been signified as being associated with apoptosis (46). Various experiments indicated that ethanol toxicity could be result in apoptosis (39). Subsequent to cell exposure to programmed cell death (apoptosis) stimulator, cytochrome c release, from mitochondria into the cytoplasm, occurred. Cytochrome c in cytoplasm activates proteolytic molecules known as caspases that specifically cleave the amino acid sequence Asp-Glu-Val-Asp (DEVD), and are crucial for the execution of apoptosis. The protein Bcl-2 prevents the cytochrome c release and therefor inhibits the activation of caspases, oppositely the protein Bax induces the cytochrome c release, caspase-3 activation, membrane blebbing, nuclear fragmentation, and cell death (47). Once cytochrome c release occurred, it activates caspase-9, and then active caspase-9 cleaves and activates the effector caspase-3 (48). The initiator caspase-8 may be stimulated by cell membrane death receptors. These apoptotic pathway key factors were investigated in this study, to comprehend whether subchronic exposure to ethanol can trigger apoptosis in liver and kidney tissues. Our data demonstrated that subacute ethanol toxicity may increase the both mRNA and protein ratio of Bax and Bcl-2 (Bax/Bcl-2) in the liver and kidney tissues (Figures 7, 8 and 15). Rise in Bax/Bcl-2 ratio leads to apoptosis in the liver and kidney tissues. To investigate the activation mechanism of apoptosis by ethanol, the levels of caspase-3, caspase-8, and caspase-9 were also estimated. Our finding implied that ethanol exposure induced caspase-3, caspase-8 and caspase-9 activation and elevated the level of cleaved caspase-3, caspase-8 and caspase-9 in the liver and kidney tissues (Figures 9-14 respectively). Thus it is likely that both intrinsic and extrinsic pathways of apoptosis have a major effect in the hepatic and nephrotic toxicity of ethanol.
N. sativa L. (family Ranunculaceae), generally identified as black seed or black cumin, is an annual plant that has been conventionally used in the Indian subcontinent (49), Arabian countries (50) and Europe (51) for culinary and medical purposes as a natural therapy for several diseases and disorders like asthma, hypertension, diabetes, cough, inflammation, bronchitis, eczema, headache, fever, influenza and dizziness (52). N. sativa seeds include 36%-38% fixed oils, proteins, alkaloids, saponin and 0.4%-2.5% essential oil (51). Unsaturated fatty acids are the main component of N. sativa fixed oil. Many components were described, but the foremost one was thymoquinone (52). Thymoquinone (TQ) was intensively studied and ascribed to possess antioxidant and anti-inflammatory properties (29, 53-55), and to decrease the nephrotoxicity of some chemotherapeutic agents (56). Furthermore the anticonvulsant (57), antiinflammatory (58), anticancer, antiproliferative (24), antia apoptotic (48) and antibacterial (59) activity of TQ has been investigated.

According to our results, simultaneous administration of TQ and ethanol reduced the severity of lipid peroxidation (MDA levels) and increased antioxidant capacity (reduced GSH content) in the liver and kidney tissues in rats (Figures 3 and 4 respectively). In addition, the protective effect of TQ against ethanol-induced hepatotoxicity indicated by the significant reduction of liver enzyme (AST, ALT and ALP) activity (Figure 1), along with considerable decrease in inflammatory cytokine (IL-6 and TNF-α) in liver tissue (Figure 2). Also, TQ through antioxidative (60) and anti-inflammatory effects (53, 61) may defend the liver (Figures 5D, 5E, 5F) and kidney (Figures 6D, 6E, 6F) tissues from histological change stimulated by ethanol. So, in this experiment it was indicated that the lipid peroxidation inhibition characteristic of TQ could be ascribed to the protective effect which prevents plasma membranes destruction and hepatic enzymes release.

It was reported that antioxidants are involved in gene expression and signal transduction pathways, and can impede the incidence of apoptosis. TQ exerts an antiapoptotic effect through attenuating oxidative stress and inhibiting TNF-α induced NF-κB activation. Furthermore, it regulates the Bax/Bcl-2 ratio and inhibits downstream caspas (48). Our results revealed that administration of TQ clearly reduced the both protein and mRNA ratio of Bax and Bcl-2 (Bax/Bcl-2) in the liver and kidney of ethanol treated groups. Furthermore, TQ diminished the activation of caspase-3, caspase-8 and caspase-9 triggered by ethanol in liver and kidney tissues.

Conclusion

In summary, our findings indicated that ethanol-induced hepatotoxicity and nephrotoxicity could be prevented by TQ administration. TQ exerts its protective effect through reducing lipid peroxidation, increasing antioxidant defense (reduced GSH content), reducing liver enzyme (AST, ALT and ALP) and specific inflammatory cytokine release, attenuating histopathological changes, and reducing the severity of apoptosis by decreasing Bax/Bcl-2 ratio and inhibition of caspase-3, caspase-8 and caspase-9 activation.

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Conflict of interest

The authors declare that there are no conflicts of interest.

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