Investigation of the Protective and / or Therapeutic Effects of Lycopene on Diethylnitrosamine-Induced Chronic Hepatotoxicity Using Biochemical, Molecular and Histopathological Data

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ABSTRACT

The aim of the study is to investigate the role of lycopene on diethylnitrosamine (DEN)-induced chronic hepatotoxicity using biochemical, molecular and histopathological approaches. Thirty five male Wistar albino rats were assigned into five groups of 7 rats each. Groups were formed as control, lycopene, DEN, lycopene+DEN and DEN+lycopene. Lycopene was applied to rats every other day at 10 mg/kg/bw, gavage for 10 days. DEN was applied intraperitoneally to rats at a single dose, 200 mg/kg/bw for 90 days. Lycopene administration was started 10 days before the DEN administration in lycopene+DEN group, together with the DEN administration in DEN+lycopene group. The study was terminated 90 days after DEN administration. DEN caused the oxidative stress by the increased malondialdehyde level and the decreased reduced glutathione level, antioxidant enzyme activities (p<0.001). Lycopene administration improved the biochemical indices of both blood and liver tissue compared to the DEN group. RT-PCR analysis revealed that the catalase enzyme in the DEN group increased expression levels. Histopathologically, many histopathologic changes such as karyomegaly, necrosis and hydropic degeneration were observed in the liver tissues of the DEN and lycopene+DEN groups. Both biochemical and histopathological results showed that healing of DEN+lycopene group was better than lycopene+DEN group. These results suggest that besides the protective effects, the therapeutic effect of lycopene is due to its antioxidant effects on DEN-induced hepatotoxicity.

Keywords: Antioxidant, Diethylnitrosamine, Lycopene, Malondialdehyde, CAT Enzyme Expression

ÖZ

Biyokimyasal, Moleküler ve Histopatolojik Veriler Kullanılarak Likopenin Dietilnitrozamine Bağlı Kronik Hepatotoksiteste Üzerine Koruyucu ve / veya Tedavi Edici Etkilerinin Araştırılması

Çalışmanın amacı, likopenin dietilnitrozamin (DEN) kaynaklı kronik hepatotoksitisti üzerindeki rolünü biyokimyasal, moleküler ve histopatolojik yaklaşımları kullanarak araştırmaktır. 35 adet erkek Wistar albino rat, her grupta 7 rat olarak şekilde 5 gruba ayrılmıştır. Gruplar, kontrol, likopen, DEN, likopen+DEN ve DEN+likopen şeklinde oluşturulmuştur. Likopen, gün aşırı olarak 10 mg/kg/vücut ağırlığı dozunda gavag ve 10 gün uygulanmıştır. DEN, ratlara 200 mg/kg/vücut ağırlığı dozunda intraperitoneal olarak tek doz 90 gün uygulanmıştır. Lycopene uygulaması, likopen+DEN grubunda DEN uygulamasından 10 gün önce, DEN+likopen grubunda ise DEN uygulaması ile birlikte başlamıştır. DEN uygulamasından 90 gün süre sonra çalışma sonlandırılmıştır. DEN, dokularda malondialdehid düzeylerinde artışa, redukte glutatyon düzeyi ve antioksidan enzim aktivitelerinde düşüşe sebep olarak oksidatif stres neden olmuştur (p<0.001). Lycopene uygulaması hem kan hem de karaciğer dokusunda DEN grubuna lâyısla biyokimyasal endekslere de iyişleme sağlar. RT-PCR analizlerinde DEN grubundaki katalaz enziminin ekspresyon düzeylerini artırdığı belirlenmiştir. Histopatolojik olarak, DEN ve likopen+DEN gruptuların karaciğer dokularında karyomegal, nekroz ve hidroplik dejenersiyon gibi birçok histopatolojik değişiklikler gözlenmiştir. Hem biyokimyasal hem de histopatolojik sonucarda DEN+likopen grubundaki iyileşmesinin likopen+DEN grubundan daha iyi olduğu gözlenmiştir. Bu sonuçlar, likopenin koruyucu etkisinden ziyade tedavi edici etkisinin DEN’e bağlı hepatotoksitistede likopenin антиоксидан etkisine bağlı kaynaklanan göstermektedir.

Anahtar Kelimeler: Antioksidan, Dietilnitrozamin, Likopen, Malondialdehid, CAT Enzim Ekspresyonu

*This article is a part of the first author’s doctoral thesis named “The Effect of Lycopene on Oxidative Stress and DNA Damage in Diethylnitrosamine Administered Rats.”
INTRODUCTION

Nitrosamines and nitroso compounds are one of the most important chemicals and adversely affect human health and can cause tumor formation. They exhibit this activity mainly by interacting with nucleic acids and proteins (Jahan et al. 2007). The dose, frequency of exposure, and method of administration of the nitrosamines may affect different organs (e.g., lung, kidney or liver) and may lead to tumor formation (Lijinsky and Kovatch 1989; Lijinsky 1992). Toxic effects of nitrosamines, particularly diethylnitrosamine (DEN), are primarily seen in the blood and liver. Most drugs lead to nitrosamine formation and increase of cancerous cells in the liver when used with sodium nitrite. Furthermore, other organs where blood flow is high, such as kidneys, are also affected at a lower dose than the liver (Yılmaz et al. 2008).

DEN is a carcinogenic substance, widely used in experimental animal models. DEN causes degenerative, proliferative, and neoplastic lesions in the liver (Gayathri et al. 2009). DEN is hydroxylated in the liver by cytochrome P-450 isoenzymes and becomes bioactivated by alkylation mechanism. Activation results in the production of radical metabolites, mainly ethyl radical (CH3CH2) (Lijinsky 1992). DEN increases reactive oxygen species (ROS), and by this way leads to cell damage and oxidative stress. The generation of ROS may be responsible for the carcinogenic effects of DEN (Yamada et al. 2006; Gayathri et al. 2009). High intracellular levels of ROS lead to mitochondrial damage, DNA modification, and lipid peroxidation (LPO), resulting in many diseases, including cancer (Yamada et al. 2006; Pradeep et al. 2007). The metabolites of DEN mediate the binding of tumor promoters by covalently binding to the DNA with one or two oxidation-providing electrons. The tumor promoter causes the formation of reactive oxygen molecules and hydrogen peroxide (H2O2) by acting as a superoxide radical (O2−) inducer (Yamada et al. 2006; Pradeep et al. 2007).

Lycopene have anticarcinogenic, anti-inflammatory and antioxidant effects. The relationship between free oxygen radicals and lycopene is as follows: adding a new radical to the free radical, neutralizing the radical by removing a hydrogen atom from its structure, and neutralizing the radical by transferring an electron from structure of the free radical (El-Agemy et al. 2004). Lycopene can protect cells against oxidative stress by acting together with various mechanisms (e.g. singlet oxygen scavenger) as an antioxidant. Before being reduced, a molecule of lycopene can bind thousands of singlet oxygen molecules (Krinsky 1988). Its effectiveness in capturing singlet oxygen depends on conjugated double bonds number it mainly contains (Ukai et al. 1994).

The aim of the study was to determine the possible protective and / or therapeutic effects of lycopene on DEN-induced oxidative damage and CAT expression in the blood and liver tissues using biochemical, molecular and histopathological approaches in rats.

The experimental applications were conducted conveniently with the conditions for the care and use of laboratory animals (12 hours light; 12 hours darkness; 24±3°C). During the experimental applications, the rats were provided the commercial rat feed (pellet feed) and tap water ad libitum.

The animals were randomly divided into five experimental groups, including seven rats in each. These groups were arranged as follows: control group (not treated), lycopene group (lycopene was administered), DEN group (a single dose of DEN was administered), lycopene+DEN (lycopene was started 10 days before the DEN administration), DEN+lycopene group (DEN and lycopene were administered together).

Lycopene was suspended in corn oil and it was administered to the animals by gavage every other day at the dose of 10 mg/kg body weight for 10 days. DEN was administered the animals a single intraperitoneal (i.p.) dose at the dose of 200 mg/kg body weight dissolved in serum physiological (0.9% NaCl) for 90 days. The doses of DEN (El-Shahat et al. 2012) and lycopene (Kumar ve Kumar 2009) used in this study was selected according to previous studies.

Sample collection

Following the applications, rats were sacrificed and tissue samples were obtained from blood and liver. Blood samples were collected in tubes containing anticoagulant (ethylenediaminetetraacetic acid [EDTA]) and centrifuged for 10 min at 3000 rpm at +4°C to separate plasma. Autoanalyser (Advia 1800 Chemistry Analyser, Siemens Healthineers, Germany) was used to determine the aspartate transaminase (AST), alanine transaminase (ALT), alkaline phosphatase (ALP), lactate dehydrogenase (LDH) enzyme activities. Before starting the analyses, liver tissues were washed with saline and then diluted 1:10 with distilled water and homogenized in a Potter-Elvehjem homogenizer. Homogenates were centrifuged at 3500 rpm for 15 min for malondialdehyde (MDA), reduced glutathione (GSH), CAT, glutathione-S-transferase (GST) and superoxide dismutase (SOD) analysis and at 14000 rpm for 55 min for glutathione peroxidase (GSH-Px) analysis.

For the reverse transcription-polymerase chain reaction (RT-PCR) analyses, blood and liver samples were taken into tubes containing RNA stabilization solution and stored at -80°C until the study was performed.

Biochemical analysis

MDA and GSH levels, CAT, GSH-Px, GST, SOD activities, protein, hemoglobin levels were determined by spectrophotometric methods in tissue samples.

The MDA level was tested according to the method described by Placer et al. (1966). This method was based on the reaction of thiobarbituric acid (TBA) with MDA, one of the aldehyde products of lipid peroxidation. The GSH level was determined by the method of Ellman et al. (1961). This method was a spectrophotometric method based on the formation of highly stable yellow colour of sulphydryl groups when 5,5’dithiobis-2-nitrobenzoic acid (DTNB) was added.

The CAT activity was carried out by using Aebi’s method (1984). It was determined by measuring the resolution of H2O2 at 240 nm. The GSH-Px activity was determined by the Beutler method (1984). GSH-Px catalyses the oxidation of GSH to oxide glutathione (GSSG) using H2O2.

MATERIALS and METHODS

Animals and experimental groups

A total of 35 male Wistar-Albino rats (weighing 250–300 g, aged 3 months) were used in this study. Ethics committee approval was obtained from Firat University Local Ethics Committee for Animal Experiments (2014/18). The animals to be used in the study were obtained from the Firat University Experimental Research Center.
The method described by Habig et al. (1974) was utilised to test GST activity. The enzyme activity was determined by measuring the amount of enzyme catalysing 1 μmol of 1-(5-glutathionyl)-2,4-dinitrobenzene per minute at 340 nm at 37°C using GSH and 1-chloro-2,4-dinitrobenzene. The SOD activity was tested by quantifying O2− generated by xanthine and xanthine oxidases reacting with nitroblue tetrazolium (Sun et al. 1980). In the determination of hemoglobin level, Frankel et al. (1970) method is used. The protein concentration was determined using the method described by Lowry et al. (1951).

**Total RNA isolation and gene expression analysis**

Both blood and liver were processed based on the method of Rotimi et al. (2017). Blood samples were placed into RNAProtect® animal blood tubes containing stabilization reagent; whereas, liver tissues were preserved in RNAlater. The gene expression levels of CAT enzyme was assessed using semi-quantitative reverse transcriptase polymerase chain reaction (RT-PCR). RNA was extracted from the blood using an RNAprotect® animal blood kit (Qiagen, Germany). RNA was extracted from liver using an RNaseasy® Plus Mini Kit according to the manufacturer’s protocol. cDNA was obtained from RNA samples via a Reverse Transcription System Kit (First Strand Kit, Qiagen, Germany) as well as an Applied Biosystem, 7500 Fast RT-PCR System.

The RNA samples were incubated for 15 minutes at 42°C, as well as for 5 minutes at 95°C for cDNA synthesis in accordance with the manufacturer’s protocol. The gene specific primers (rat CAT primer (NM_012520)) for RT-PCR were obtained from Qiagen. Reactions were prepared in 25 μl volumes by the SYBR Green Master Mix (Qiagen, Germany). The cycling conditions were performed according to the manufacturer’s protocol as follows: 15 minutes at 95°C, 15 seconds at 95°C, and 1 minute at 60°C (repeated for 40 cycles). The RT-PCR analysis results were then normalized against the expression level of the internal control β-Actin (NM_031144). The values of all the samples were expressed as percentages with respect to the control.

**Histopathological examination**

Following necropsies, liver tissue samples were fixed in 10% neutral buffered formalin, were embedded in paraffin and cut at a thickness of 5 μm. Then, they were stained with Hematoxylin and Eosin (H&E). After that, 10 random microscopic areas were examined at x40 magnification. The changes were rated to be none (0), mild (1), moderate (2), and severe (3) (Luna, 1968).

**Statistical analysis**

Shapiro-Wilk normality test was applied to determine whether the raw values of all the measured parameters showed normal distribution and as a result of the test, it was determined that the values in all parameters showed normal distribution. Based on the result of this test, one-way analysis of variance (ANOVA) was used to determine differences between groups, and post hoc Tukey test for paired comparisons.

All statistical analyses were performed using SPSS version 22.0 software (IBM Corporation, Armonk, NY, USA). The data obtained as a result of the study are expressed as mean and standard error. A p value of <0.05 was considered statistically significant.

**RESULTS**

**MDA, GSH levels, and antioxidant enzyme activities**

Tables 1 and 2 show the MDA, GSH levels and CAT, GSH-Px, GST and SOD activities in blood and liver tissues. When the DEN treated group was compared with the control group, an increase was observed in the MDA levels and a decrease in GSH levels, CAT, GSH-Px, GST and SOD activities in blood and liver tissue samples (p<0.001). When the DEN+lycopene group was compared with the group in which lycopene was started before DEN administration (lycopene+DEN group) and group in which lycopene was started together with DEN administration (DEN+lycopene group), MDA levels were found to decrease whereas GSH levels, CAT, GSH-Px, GST and SOD activities were found to increase in the blood tissue and the values of the DEN administered groups approached the values of the control group (p<0.001). Similar changes were observed in the liver tissue and the changes were found to be more significant in the group in which lycopene was given with DEN administration (DEN+lycopene group).

**CAT enzyme gene expression levels**

Figures 1 and 2 show CAT enzyme gene expression levels in blood and liver tissues. The CAT enzyme gene expression levels were found to be significantly higher in the liver tissues of the DEN treated group compared to those of the control group, however, there was no statistically significant change in blood tissue compared to the control group. A significant decrease was observed in the expression levels in the liver tissue in the lycopene supplemented groups (lycopene+DEN and DEN+lycopene groups) compared to the DEN group and this decrease was found to be higher in DEN+lycopene group.

**AST, ALT, ALP and LDH activities**

Plasma AST, ALT, ALP, LDH activities are given in Table 3. A statistically significant increase was observed in all values in the DEN group compared to the control group (p<0.05 and p<0.001). When the lycopene+DEN group and DEN group were compared, no difference was observed in the lycopene+DEN group in terms of AST and ALP activities whereas ALT, LDH activities decreased and approached to the control group values. In the DEN+lycopene group, there was a decrease in AST, ALT, LDH activities compared to the DEN group, but there was no change in ALP activity.

**Histopathological results**

Microscopic lesions and scoring values of control and experimental groups are given in Table 4. While the severity and distribution varied according to groups, the most significant microscopic changes were generally as follows: cloudy swelling, necrotic changes characterized by karyopyknosis, karyomegaly and apoptosis in hepatocytes and disorganization, sinusoidal and portal congestion, periportal cell infiltration, intra- and extra-cellular bile pigment accumulations, and Kupffer cell hyperplasia on the hepatic cord. The most severe microscopic changes were observed in DEN and lycopene+DEN groups. In the DEN+lycopene group, it was noted that there was a decrease in the severity of degenerative changes in the liver compared to the DEN group.
### Table 1. Effects of Lycopene on MDA and GSH levels and CAT, GSH-Px, SOD activities in blood tissue of DEN treated rats.

|                | Control         | Lycopene       | DEN             | Lyco.+DEN                   | DEN+Lyco.  |
|----------------|-----------------|----------------|-----------------|-----------------------------|------------|
| MDA (nmol/ml)  | 8.34±0.04ab     | 7.00±0.10a     | 11.50±0.17c     | 8.57±0.10b                  | 8.79±0.14b |
| GSH (μmol/ml)  | 47.21±0.22      | 47.01±0.40a    | 36.03±0.84c     | 45.58±0.38b                 | 45.60±0.38a|
| CAT (k/g Hb)   | 65.33±0.79a     | 65.22±0.52a    | 44.37±0.63d     | 62.81±0.99a                 | 61.84±1.40a|
| GSH-Px (U/g Hb)| 60.7±1.74a      | 82.4±2.11a     | 60.63±2.74b     | 78.55±2.43a                 | 79.16±0.41a|
| SOD (U/g Hb)   | 72.93±0.49a     | 72.01±0.44a    | 65.84±0.67b     | 71.52±0.41a                 | 72.34±0.33a|

The data are expressed in mean±S.E. for seven animals per group. Within rows, means with different letters (a, b, c, and d) are significantly different.

### Table 2. Effects of lycopene on MDA and GSH levels and CAT, GSH-Px, GST, SOD activities in liver tissue of DEN treated rats.

|                | Control         | Lycopene       | DEN             | Lyco.+DEN                   | DEN+Lyco.  |
|----------------|-----------------|----------------|-----------------|-----------------------------|------------|
| MDA (nmol/g doku) | 0.63±0.01abc  | 0.61±0.01a     | 0.93±0.02d      | 0.68±0.01c                  | 0.65±0.01abc|
| GSH (μmol/ml)  | 17.12±0.23a     | 17.08±0.10a    | 15.10±0.07c     | 16.39±0.05b                 | 16.93±0.05a|
| CAT (k/mg protein) | 0.37±0.01a    | 0.38±0.01a     | 0.18±0.01c      | 0.32±0.02a                  | 0.335±0.01a|
| GSH-Px (U/g rotein) | 34.19±1.31a   | 35.07±0.88b    | 28.43±1.11b     | 32.61±1.10a                 | 32.28±1.59a|
| GST (U/mg protein) | 20.92±0.28a   | 20.56±0.46a    | 17.44±0.26c     | 19.37±0.23b                 | 19.96±0.30ab|
| SOD (U/mg protein) | 0.64±0.01a    | 0.64±0.01a     | 0.62±0.01c      | 0.64±0.01a                  | 0.639±0.01ab|

The data are expressed in mean±S.E. for seven animals per group. Within rows, means with different letters (a, b, c, and d) are significantly different.

### Table 3. Effects of lycopene on AST, ALT, ALP, LDH activities in plasma of DEN administrated rats.

|                | Control         | Lycopene       | DEN             | Lyco.+DEN                   | DEN+Lyco.  |
|----------------|-----------------|----------------|-----------------|-----------------------------|------------|
| AST (U/L)      | 171,00±6,77ab   | 146,33±2,87a   | 225,00±5,97d    | 242,50±5,71d                | 189,66±12,51bc|
| ALT(U/L)       | 78,25±6,10a     | 83,75±2,32ab   | 96,00±2,22bc    | 88,71±5,62ab                | 87,50±6,63ab|
| ALP(U/L)       | 6,50±0,22a      | 7,00±0,28a     | 15,00±0,88b     | 14,33±2,92b                 | 13,50±2,46b|
| LDH(U/L)       | 794,00±49,07ab  | 724,57±63,34a  | 1363,20±73,63c  | 980,66±27,79b               | 853,00±28,98b|

The data are expressed in mean±S.E. for seven animals per group. Within rows, means with different letters (a, b, c, and d) are significantly different.

### Table 4. Microscopic lesions observed in liver of DEN treated rat.

|                | Control         | Lycopene       | DEN             | Lyco.+DEN                   | DEN+Lyco.  |
|----------------|-----------------|----------------|-----------------|-----------------------------|------------|
| Karyomegaly    | 0.14±0.0a       | 0.14±0.0a      | 2.00±0.31bc     | 2.14±0.34c                  | 1.86±0.14b |
| Sinusoidal Congestion | 0.57±0.20a    | 1.00±0.00c     | 2.00±0.00c      | 1.71±0.18b                  | 2.00±0.00c |
| Karyopyknosis  | 0.14±0.0a       | 0.14±0.0a      | 1.57±0.20b      | 2.00±0.00c                  | 1.29±0.18b |
| Cloudy Swelling (Periasiner) | 0.0±0.0a     | 0.44±0.20b     | 3.00±0.00c      | 3.00±0.00c                  | 1.71±0.18bc|
| Single Cell Necrosis | 0.0±0.0a      | 0.57±0.20a     | 2.57±0.20b      | 2.71±0.18b                  | 1.43±0.18bc|
| Fatty Degeneration | 0.14±0.14a    | 0.0±0.00a      | 1.00±0.00ab     | 1.00±0.00ab                 | 1.00±0.00ab|
| Bile Duct Proliferation | 0.0±0.0a      | 0.14±0.0a      | 2.00±0.00c      | 1.00±0.00b                  | 1.00±0.00b |
| Apoptotic Bodies | 0.14±0.0a       | 0.14±0.0a      | 2.00±0.00c      | 1.00±0.00b                  | 1.00±0.00b |
| Kupffer Cell Activation | 0.0±0.0a     | 0.0±0.00a      | 0.43±0.20ab     | 1.00±0.00b                  | 1.00±0.00b |
| Periportal Cell Infiltration | 0.57±0.20a    | 0.71±0.18a     | 1.43±0.20b      | 1.29±0.18b                  | 1.00±0.00ab|
| Hepatic Cord Disorganization | 0.29±0.18a    | 0.29±0.18a     | 2.57±0.20c      | 2.00±0.31bc                 | 2.00±0.21bc|
| Portal Congestion | 1.00±0.0a      | 1.00±0.0a      | 1.29±0.18b      | 1.43±0.20b                  | 1.43±0.20b |
| Bile Pigment Accumulations | 0.0±0.0       | 0.0±0.0        | 1.00±0.00       | 1.00±0.00                   | 1.00±0.00  |

The data are expressed in mean±S.E. for seven animals per group. Within rows, means with different letters (a, b, and c) are significantly different (p<0.001).
Figure 1. The effect of lycopene supplementation on blood CAT expression level.

![Graph showing the effect of lycopene supplementation on blood CAT expression level.](image)

Figure 2. The effect of lycopene supplementation on liver CAT expression level (*P < 0.001).

![Graph showing the effect of lycopene supplementation on liver CAT expression level.](image)

Figure 3. Light microscopic view of the liver of DEN-treated rats. A. Control Group, normal appearedance liver structure. B. Lycopene Treated Group, normal appearedance liver structure. C. DEN Treated Group, karyomegaly (arrowhead), necrosis-necrotic changes (*), single cell necrosis (thick arrow), bile duct proliferation (thin arrow). D. Lycopene + DEN Treated Group, cloudy swelling (*), periportal cell infiltration (thick arrow), apoptotic bodies (thin arrow). E. DEN + Lycopene Treated Group, karyomegaly (arrowhead), fatty degeneration (thin arrow). (H&E staining, magnification x40).

DISCUSSION

The free radical formation may be suggested as the main mechanism of hepatotoxicity although many opinions have been reported in the pathogenesis of DEN-induced toxicity. If the generation of free radicals exceeds the protective effects of the defense system, they adversely affect metabolism. The increase in the MDA level is an important sign of oxidative stress (Gaweł et al. 2004). In the present study, the increase in MDA levels in plasma and liver tissues obtained from rats in the DEN group can be considered as an indicator of the DEN-induced damage. This increase may be further related to the metabolism of DEN, which reacts with cellular components, particularly with polyunsaturated fatty acids, in the cell membrane and causes the release of iron ions that increases the MDA levels. GSH prevents the peroxidation, protects cell membranes and allows the removal of free radicals (Farombi and Fakoya 2005). In the present study, the decrease observed in the GSH levels in blood and liver tissues of the rats in the DEN group is attributed to DEN or its metabolites forming a conjugate with GSH to facilitate its excretion from the body. Furthermore, GSH plays a critical role in protecting tissues against the harmful effects of DEN and therefore, GSH consumption can also lead to significant decreases in GSH-Px activity. The decrease in GSH-Px activity following the DEN administration may be due to decreased levels of the relevant substrate (GSH) and ROS-induced changes in the protein structure (Farombi and Fakoya 2005; Halliwell and Gutteridge 2015).

In the present study, MDA levels in plasma and liver tissues were found to be significantly higher in the DEN group than in the control group. This indicates that DEN causes high levels of free radical formation that cannot be tolerated by the cellular antioxidant defense system. However, lycopene administration decreased the MDA level to the level in the control group, suggesting that lycopene may be effective in clearing free radicals, inhibiting LPO, and protecting membrane lipids against oxidative damage in plasma and liver of rats.

Antioxidant enzymes such as CAT and GSH-Px are the first line antioxidant defense system against ROS and their activities have been observed to decrease by DEN. The decrease in the CAT and GSH-Px activities due to the DEN-induced oxidative stress may be due to the possible inhibitory effect of DEN-induced ROS on these enzymes, and decreased availability of GSH as a substrate while GSH-Px catalyzes the conversion of lipid peroxide to hydroxy acids in the presence of GSH (Farombi and Fakoya 2005; Halliwell and Gutteridge 2015). In the study, lycopene supplementation increases blood antioxidant activities and thus prevents the detrimental effects of peroxidation products.

In a study by Kim et al. (1997), in which different nitrosamines (DEN, N-methyl-N-nitrosourea, and 1,2-dimethylhydrazine) were used in mice, the combination of DEN (10 mg/kg i.p.) with other nitrosamines has been reported to increase the neoplasia and oxidative stress in the lungs of mice at the end of 32 weeks. In a study by Pradeep et al. (2007), the effects of silymarin on DEN-induced hepatotoxicity (200 mg/kg i.p.) has been investigated and authors have reported an increase in
some liver-specific enzymes and LPO whereas there is a decrease in the antioxidant enzyme levels in the DEN group. In another study by the same authors investigating the effects of Cassina fistula on DEN-induced hepatotoxicity (100 mg/kg i.p.), they have reported that some liver-specific enzymes and LPO have increased whereas antioxidant enzyme levels have decreased, similar to their previous study (Pradeep et al. 2010).

In a study published in the literature, a significant decrease has been reported in gene expression levels of GST, CAT and GSH-Px enzymes in the DEN group in parallel with the decrease in antioxidant enzyme activities. The decrease in antioxidant enzyme activities and CAT gene expression after DEN administration has been attributed to the increase in the production of free radicals during DEN metabolism (Sayed-Ahmed et al. 2010). In a study by Bingül et al. (2015), a decrease has been reported in the CAT activity and expression levels in DEN treated rats. However, CAT enzyme expression levels were found to increase in groups treated with DEN in our study, contrary to other studies in the literature. The difference between CAT activity and expression level is thought to be due to the procedures performed after the post-translational transformations. There are studies, in which antioxidant compounds were used against oxidants in different tissues in oxidative stress, showing that these oxidant compounds increase (Kim et al. 2009) or decrease (Fustinoni-Resi et al. 2016) the CAT gene expression levels in particular among the antioxidant enzymes. Changes in the present study are thought to be caused by DEN-induced oxidative DNA damage in rat liver. Metabolites formed during the metabolism of DEN via cytochrome P-450 lead to oxidative, mutagenic and genotoxic damages in tissues.

In the present study, CAT enzyme activity in blood and liver tissues decreased significantly but the expression level of the same enzyme increased in DEN treated groups compared to the control group. The difference between CAT activity and gene expression level is thought to be due to variation in protein and mRNA transformation. Furthermore, the relationship between genotype and mRNA/protein levels can be also influenced by various factors such as environment and transcriptional regulation.

There are studies reporting that lycopene significantly changes LPO levels and antioxidant enzyme activities. In a study by Gupta et al. (2013) involving mice, the group received pretreatment with lycopene was compared with the DEN treated group and pretreatment with lycopene has been reported to have no significant effect on the activity of phase I enzymes, to decrease the LPO and GSH-Px, GST and SOD activities. Blanche et al. (2013) have investigated the effects of Apo-10′-lycopenoic acid (10 mg/kg diet), a lycopene metabolite, in mice that were given DEN initially and fed with a high-fat diet for 24 weeks. They have observed a decrease in both the development of hepatic oxidative, mutagenic and genotoxic damages in tissues. The difference between CAT activity and expression level is thought to be more evident than its protective effect. Other studies have shown that lycopene in DEN-induced hepatic inflammation and tumorigenesis of the study and this article is a part of the first author’s doctoral thesis named “The Effect of Lycopene on Oxidative Stress and DNA Damage in Diethylnitrosamine Administered Rats”.

CONCLUSION

Increased LPO levels and liver-specific enzyme activities and decreased enzymatic and non-enzymatic antioxidant levels are the signs of a liver injury. Moreover, elevated levels of CAT expression due to the increased oxidative stress are supportive. Considering the results of our study, it can be concluded that oxidative stress plays an important role in DEN-induced liver injuries. Furthermore, the effect of lycopene given together with DEN administration has been observed to be higher than the effect of lycopene given before DEN administration. Based on the biochemical, molecular and histopathological results of the study, the therapeutic effect of lycopene can be said to be more evident than its protective effect.

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