INTRODUCTION

Acute renal failure from ischemic damage to the kidney cause morbidity and mortality. Tubular epithelial cell death is the most common cause of this renal failure [1]. Even after reperfusion of extended ischemia of kidney, injury occurs [2]. Although reperfusion is essential for the survival of ischemic renal tissue, it may cause additional damage to kidney [3]. Renal damage caused by ischemia reperfusion injury (IRI) occurs in surgical procedures in which the kidney remains without blood supply for some time. IRI is the most common cause of acute renal failure after renal transplantation, major abdominal and vascular surgery, coronary bypass graft surgery, trauma, and sepsis [4]. IRI seen in kidney transplantation effects the short- and long-term graft survival [5]. IRI results in persistent intrarenal vasoconstriction, injury of microvascular endothelial and tubular epithelial cells [6]. Reperfusion of ischemia causes a rapid burst of free radicals, which is responsible for endothelial injury and edema. Reperfusion damages endoplasmic reticulum, leading to autophagosome formation and cellular disintegration [7].

Medical protection of the kidney against IRI is very important for nephron sparing surgery and renal transplantation, so many studies continue in this area. Many agents have been used for the protection of ischemia reperfusion renal tissue injury, such as recombinant human manganese superoxide dismutase, n-acetylcysteine and desferrioxamine, telmisartan, exendin-4, and sitagliptin [8, 9, 10].

In this study we aimed to evaluate the radioprotective effect of lycopene on the kidney after unilateral ischemia due to renal artery temporary clamping through the use of histopathological evaluation, malondialdehyde (MDA), myeloperoxidase (MPO) and total nitrite level analysis and 99mTc-dimercaptosuccinic acid (DMSA) scintigraphy uptake.
METHODS

This experimental study was approved by the Ankara Training and Research Hospital Local Ethics Committee of Animal Experiments and conducted in Hüsnü Sakal Experimental and Clinical Practice Center at the same hospital. The study group consists of total of 21 Wistar male albino rats (260 ± 45 grams, three to five months old). The rats were acclimated for at least one week prior to the study and housed under standard laboratory conditions (constant temperature (21°C ± 2°C) with relative humidity of 50–60%, with 12-hour light and dark cycles). During the study, the animals had ad libitum access to water and standard food.

Three groups of rats were formed and evaluated. Group 1 (n = 9) was administered lycopene [5 mg per kg of body weight per day (LYC-O-MATO®, GNC Holdings Inc., Pittsburgh, PA, USA)] starting three days before right renal ischemia reperfusion injury and this was continued for 15 days. Group 2 was the control group (n = 9), to which right renal ischemia reperfusion injury was applied without any medication. Group 3 (n = 3) was the sham group, to which neither right renal ischemia reperfusion injury nor medication were applied.

For static renal scintigraphic examination, 99mTc-DMSA images were obtained preoperatively and on postoperative day 15 for each group. Surgery and 99mTc-DMSA scintigraphy were applied under anesthesia of 40 mg/kg ketamine hydrochloride (Ketalar, Parke-Davis/Eczacıbaşı, Istanbul, Turkey) and 5 mg/kg xylazine (Rompun, Bayer, Istanbul, Turkey) applied intramuscularly. The subjects were sacrificed by decapitation under 50 mg/kg, intraperitoneally, with propofol anesthesia (Abbott Laboratory Corporation, Istanbul, Turkey). After the sacrifice for each group, left and right kidneys were removed surgically for histopathological examination.

Surgical method

DMSA renal scintigraphy was performed in all rats before the surgical procedure. General anesthesia was administered intramuscularly as a combination of 40 mg/kg ketamine hydrochloride (HCl) (Ketalar, Parke-Davis/Eczacıbaşı) and 10 mg/kg xylazine HCl (Rompun, Bayer). The incision area was shaved and the surgical site was cleaned preoperatively with soap and povidone-iodine. A sterile environment was prepared. By laparotomy via midline incision, the right kidney was reached and the right renal pedicle was isolated. The subjects were sacrificed by decapitation under 50 mg/kg, intraperitoneally, with propofol anesthesia (Abbott Laboratory Corporation, Istanbul, Turkey). After the sacrifice for each group, left and right kidneys were removed surgically for histopathological examination.

Histopathological evaluation

Kidney tissues obtained after sacrificing the rats were fixed immediately in 10% formaldehyde and were processed in paraffin tissue blocks and macroscopic sections were taken to include the renal cortex and pelvis. Sections of 5 μm thickness cut from formalin fixed paraffin-embedded blocks were stained with hematoxylin and eosin. Histopathological examination was performed in 40–100–200–400 × original magnification with light microscopy (Figure 2). For the histopathological score the following criteria were used:

- Normal histology: 0 points;
- Swelling of tubule cells, loss of brush border and nuclear condensation of ≤ 1/3: 1 point;
- In addition to the changes in 1 point, 1/3–2/3 tubule changes: 2 points;
- Tubular changes more than 2/3: 3 points.

All kidneys were examined in 100 areas with a maximum score of 300.
Biochemical Evaluation

Tissue samples were stored at -80°C until analysis. Kidney samples were homogenized with 0.15M KCl at a rate of 1/10 (weight per volume). MDA, MPO, and total nitrite levels were measured in the tissue samples.

Determination of MPO level

MPO activity was obtained spectrophotometrically by determining the decomposition of hydrogen peroxide using o-dianisidine as the hydrogen donor. Tissue samples of approximately 50 mg were taken, weighed and homogenized three times for 30 seconds at 4°C in 1 ml of ice-cold 0.5% hexadecyltrimethylammonium bromide in 50 mmol/L phosphate buffer (pH 6). The homogenate was subjected to three freeze/thaw cycles and centrifuged for 15 minutes at 40,000 × g. MPO activity was determined by the addition of 0.1 ml of the supernatant to 2.9 ml of 50 mmol/L phosphate buffer containing 0.167 mg/mL o-dianisidine dihydrochloride and 0.0005% hydrogen peroxide. The change in absorbance at 460 nm over a five-minute period was measured at 25°C. The data are expressed as the change in absorbance per minute per gram of tissue (Δabs/min/g of tissue) [11].

Determination of MDA level

MDA levels were calculated by the fluorometric method, developed by Wasowicz et al. [12]. After the reaction of thiobarbituric acid with MDA, the reaction product was extracted in butanol and was measured spectrofluorometrically at wavelengths of 525 nm for excitation and 547 nm for emission. As standard, 0–5 μmol/L 1,1’,3,3’-tetraethylicrylylamine solutions were used. For the measurement of tissue MDA, 50 μl of homogenate was added and introduced into 10 mL glass tubes containing 1 ml of distilled water. After the addition of 1 ml of the solution containing 29 mmol/L thiobarbituric acid in acetic acid and mixing, the samples were placed in a water bath and heated for one hour at 95–100°C. After the samples were cooled, 25 μL of 5 mol/L HCL was added and the reaction mixture was extracted by agitation for five minutes with 3.5 mL n-butanol. After separation of the butanol phase by centrifugation at 1,500 × g for 10 minutes, the fluorescence of the butanol extract was measured with F-2500 fluorometer (Hitachi Ltd., Chiyoda, Tokyo, Japan) at wavelengths of 525 nm for excitation and 547 nm for emission. As standard, 0–5 μmol/L 1,1’,3,3’-tetraethylylcrylylamine solutions were used. MDA levels are given as μmol per gram of wet tissue.

Determination of total nitrite level

A total of 300 μL of homogenate was added to 300 μL of KH2PO4/K2HPO4 buffer (pH 7.5), 50 μl of 2 mmol/L NADPH, 50 μl of 50 μmol/L FAD, and 50 μl of 1 unit/mL aspergillus nitrate reductase. This was incubated at room temperature for one hour followed by the addition of 500 μl of 0.2 mol/L ZnSO4 and 70 μl 2 mol/L NaOH to deproteinate the sample. After centrifugation, 0.75 mL of the supernatant was added to 1 mL of 1% sulphanallic acid (in 4 mol/L HCl). After 10 minutes at room temperature, 0.75 mL of freshly prepared 1% N-(1-Naphthyl)ethylene-diamine was also added. The resultant color change was measured at 548 nm using a spectrophotometer. Nitrite concentration was calculated from 5, 12.5, 25, 50 μmol/L sodium nitrite standards [13]. Total nitrite levels are given as nmol per liter per gram of wet tissue.

99mTc-DMSA scintigraphy evaluation

Preoperatively and on postoperative 15th day of the renal injury, adequate hydration with sterile saline and induction of anesthesia were carried out before the scintigraphic study. A commercially available kit of DMSA (MON. DMSA KIT, Monrol Eczacıbaşı, Istanbul, Turkey) was labeled with 99mTc in accordance with the manufacturer's leaflet for use. A 37 MBq (1 mCi) dose of 99mTc-DMSA in a 0.1 ml volume was administered to the tail vein. Two hours after injection, static renal images were taken under anesthesia. Renal static scintigraphies were performed using a single-head eCam gamma camera (Siemens Healthcare, Erlangen, Germany) equipped with pinhole collimator, energy peak adjusted to 140 keV ± 20% with 256 × 256 matrix and 2.67 zoom factor for five minutes in posterior position. ROIs (regions of interest) were drawn on the anterior and posterior images and the geometric mean of these values was accepted as the relative uptake percentage of the kidneys.

Statistical analysis

Statistical analysis was performed with SPSS Statistics for Windows, Version 20.0 (IBM Corp., Armonk, NY, USA). Continuous and categorical variables are expressed as the mean value ± standard deviation and frequency (%) of numbers. Group comparisons for each parameter were done using the nonparametric Mann–Whitney U-test and the Kruskal–Wallis tests. A p–value less than 0.05 was considered to be a statistically significant difference.

RESULTS

The histopathological score was 45.00 ± 16.58 in the lycopene group, 95.56 ± 40.34 in the control group, and 30.00 ± 5.00 in the sham group (Table 1). The histopathological score was lower in the lycopene group than in the control group, 95.56 ± 40.35 in the control group, and 30.00 ± 5.00 in the sham group (p = 0.012). The histopathological score of the

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**Table 1. Histopathological scores of the groups; values are given as mean ± standard deviation (SD)**

<table>
<thead>
<tr>
<th>Group</th>
<th>Minimum score</th>
<th>Maximum score</th>
<th>Mean score ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lycopene group</td>
<td>10</td>
<td>60</td>
<td>45 ± 16.58</td>
</tr>
<tr>
<td>Control group</td>
<td>75</td>
<td>200</td>
<td>95.56 ± 40.35</td>
</tr>
<tr>
<td>Sham group</td>
<td>25</td>
<td>35</td>
<td>30 ± 5</td>
</tr>
</tbody>
</table>
lycopene group was higher than that of the sham group, but it was not statistically significant (p = 0.093) (Table 2).

In biochemical analysis, the mean value of MPO level (Δabs/min/g tissue) was 0.08 ± 0.09 in the lycopene group, 0.11 ± 0.03 in the control group, and 0.06 ± 0.03 in the sham group. The MPO level was lower in the lycopene group than in the control group, and higher in the control group than in the sham group, but not statistically significant. The mean value of the MDA level (µmol/g of wet tissue) was 4.61 ± 2.10 in the lycopene group, 6.85 ± 1.68 in the control group and 13.36 ± 3.56 in the sham group. The MDA level was lower in the lycopene group than in the control group (p = 0.009) and the sham group (p = 0.013), and lower in the control group than in the sham group (p = 0.021). The mean value of the nitrite level (nmol/l/gm wet tissue) was 0.05 ± 0.02 in the lycopene group, 0.09 ± 0.02 in the control group, and 0.04 ± 0.01 in the sham group. The nitrite level was lower in the lycopene group than in the control group (p = 0.003) and higher in the control group than in the sham group (p = 0.013). The mean value of urea level (mg/dL) was 175.56 ± 52.94 in the lycopene group, 144.44 ± 27.89 in the control group and 50 ± 10 in the sham group. The urea level was higher in the lycopene group than in the control group but not statistically significant, and higher in the control group than in the sham group (p = 0.012). The mean value of creatine level (mg/dL) was 1.46 ± 0.47 in the lycopene group, 1.24 ± 0.19 in the control group and 0.8 ± 0 in the sham group. The creatine level was higher in the lycopene group than in the control group but not statistically significant, and higher in the control group than in the sham group (p = 0.011) (Tables 2 and 3).

In the scintigraphic analysis, the preoperative mean uptake values were 50.16 ± 1.49 in the lycopene group, 50.56 ± 1.38 in the control group and 50.18 ± 0.34 in the sham group. Preoperative 99mTc-DMSA uptake values were not statistically significant between the groups (p = 0.767). The postoperative mean uptake values were 44.82 ± 1.84 in the lycopene group, 38.92 ± 1.17 in the control group and 50.21 ± 1.35 in the sham group. There was statistically significant difference between the lycopene and control groups, higher in the lycopene group (p < 0.001). The uptake values of the lycopene and control groups were lower than those of the sham group, as expected (p = 0.013, both) (Figure 3).

In our study, lycopene showed histopathological protective effect in IRI of the kidney. The histopathological evaluation, the histopathological score was lower in the lycopene group than in the control group, and histopathological score of the control group was higher than that in the sham group. MPO level in the lycopene group was lower than in the control group, but not statistically significant. MDA and nitrite levels in the lycopene group were lower than in the control group. These biochemical parameters indicate that lycopene has protective effect against IRI. 99mTc-DMSA uptake of the effected kidneys were higher in the lycopene group than in the control group, which indicates lycopene has protective effect on renal parenchyma.

In animal models, reperfusion of the ischemic kidney is followed by tissue destruction and many morbidities ensue. Neutrophils accumulate in the area of ischemic tissues after reperfusion and tissue damage is mediated through neutrophil-mediated oxidative stress. Reactive oxygen species are produced at the sites of inflammation by neutrophils and cause formation of lipid peroxides, damage...
of cell membrane and destruction of antioxidative defense mechanism. Reperfusion may increase the damage after ischemia in tissues. After reperfusion of ischemic tissue, inflammatory and metabolic damage occurs as a result of disruption of cellular integrity. When the cascade begins, various distant organs, such as lungs, liver and heart, are affected by many activated system and toxic mediators [14].

Effects of IRI on kidneys have been studied by various methods such as biochemical assays, histopathological examination, and scintigraphic imaging, but there is no consensus for the best method for the evaluation of impaired renal functions [15]. After reperfusion of ischemic kidney, MDA and MPO levels increase and injure the renal tissue to a greater extent. Reactive oxygen species released by the neutrophils increase the tissue damage further during reperfusion [14, 16]. MDA, which is a marker of lipid peroxidation, increases during reperfusion after an ischemic renal episode [17]. MPO activity increases if neutrophil infiltration into the tissues occurs as in IRI, and total nitrite level is the marker of total lipid membrane damage [18].

Lycopene is the most common studied compound on the preventive effects of dietary intake of tomato products [19, 20]. Lycopene is the most potent antioxidant among various carotenoids. Lycopene has the best relative radical scavenging abilities among the carotenoids. Carotenoids prevent damage to DNA, proteins, cell membranes, lipids, and other structures, which occurs after oxidative injury. Lycopene has higher singlet oxygen quenching ability than β-carotene and α-tocopherol by its high number of conjugated double bonds [21]. Lycopene has antiproliferative properties to protect the development of prostate cancer and inhibits cholesterol synthesis and enhances low density lipoprotein degradation [22]. Lycopene has also many other functions in the immune system, metabolic pathways, and cell–cell communication. Lycopene normalizes the change of intrathymic Th-cell differentiation seen in tumorogenesis [23].

Due to antioxidative properties, lycopene mediates free radical scavenging activity and results in the reduction of infarct volume in ischemia reperfusion brain injury [22]. Lycopene ameliorated the ischemia reperfusion induced tissue damage and was found to protect the germ cells after testicular torsion [24]. However, in another study, lycopene was not effective for testicular torsion in the long term, there was no improvement in the groups treated with lycopene for therapeutic purposes [25]. Liu et al. [26] identified the pro-regenerative, anti-apoptotic and anti-oxidant properties of mesenchymal stromal cells in IRI. Lycopene ameliorated lysosomal membrane damage as well as alterations in cardiac enzymes, lipid profile, and oxidative stress markers. Yue et al. [27] thought that lycopene protects myocardium against hypoxia reoxygenation induced apoptosis by maintaining the mitochondrial function. Lycopene was also found effective in pancreatitis by inhibition of neutrophil infiltration and lipid peroxidation [20]. Bayramoglu et al. [21] used lycopene in IRI of liver in different doses of 2.5 and 5 mg/kg body weight. Improvements of alanine aminotransferase, aspartate aminotransferase, lactate dehydrogenase, and MDA levels were partial and dose dependent. Lycopene showed the protective effect as a decrease in MDA level, nitrite level, and histopathological score, and an increase in DMSA uptake of the kidney.

Pektaş et al. [17] evaluated the effectiveness of lycopene in IRI with biochemical and histopathological parameters and found that lycopene may have a protective effect on IRI. Kaya et al. [28] also used high dose of lycopene (100 mg/kg) in a single dose in IRI of the kidney. They also used biochemical and histopathological parameters and mentioned that lycopene administered prior to renal IRI prevented renal damage. A few studies used 99mTc-DMSA scintigraphy for the evaluation of IRI of the kidney. DMSA scintigraphy was found to be an effective non-invasive method in the evaluation of kidney restoration after IRI injury [29]. In our study, we also used DMSA scintigraphy in addition to biochemical and histopathological parameters.

**CONCLUSION**

In conclusion, lycopene seems to be an effective agent for protection of the kidney in reperfusion injury after renal ischemia, as demonstrated by the histopathological, biochemical, and scintigraphic parameters. However, further larger studies are necessary for clinical use.

**REFERENCES**


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Сцинтиграфска, патохистолошка и биохемијска процена ефекта ликопена код реперфузионих оштећења бубrega после исхемије код пацова

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САЖЕТАК
Увод/Циљ Защита бубрега од реперфузионих оштећења после исхемије је врло значајна. За ову заштиту су коришћене многи средства. Циљ рада је процена радиопротективног ефекта ликопена на реперфузионих оштећења бубrega после исхемије. Методе Група од 21 мужјака вистар албино пацова подеље на је у следеће три групе: ликопенску, контроверну и псевдогрупу. Пацијенти у ликопенској групи ликопен је даван три дана пре и 15 дана после реперфузионих оштећења бубрега, контроверној групи нису даван ликопен после оштећења, а пацови псевдогрупе нису имали исхемију бубrega и није им даван ликопен. После 15 дана урађена је сцинтиграфија са 99mTc-DMSK (димер каптосукцинска киселина), а потом су пацови жртвовани и урађена су патохистолошка и биохемијска истраживања. Резултати Патохистолошки сарт кас је био нижи у ликопенској групи. Биохемијска анализа је показала да је ниво миелопе- роксидазе био нижи у ликопенској групи него у контроверној, али не статистички значајан. Нивои мононаправа је био нижи у ликопенској групи у сравнености са контролном групом.

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