Effects of Grape seed Extract and Origanum Onites Essential Oil on Cisplatin-Induced Hepatotoxicity in Rats

Aysun CETIN1, Umit ARSLANBAS1, Berkay SARAYMEN1, Ozlem CANOZ3, Ahmet OZTURK4, Osman SAGDIC5

1 Erciyes University, Faculty of Medicine, Department of Biochemistry and Clinical Biochemistry
2 Erciyes University, Faculty of Pharmacy, Department of Biochemistry
3 Erciyes University, Faculty of Medicine, Department of Pathology
4 Erciyes University, Faculty of Medicine, Department of Bioistatistics
5 Erciyes University, Faculty of Engineering, Department of Food Engineering, Kayseri, TURKEY

ABSTRACT
Cisplatin is a widely used anticancer drug but, it can produce undesirable effects such as hepatotoxicity even in therapeutic doses. The underlying mechanism in hepatotoxicity has been attributed to free oxygen radicals. The present study was designed to determine the possible protective effects of grape seed extract (GSE) and Origanum onites essential oil (OOEO) on liver toxicity induced by cisplatin. Ninety-six male Wistar albino rats were divided into eight groups, twelve in each (Control, GSE, OOEO, GSE+OOEO, Cisplatin, GSE+Cisplatin, OOEO+Cisplatin, GSE+OOEO+Cisplatin) and followed up for 10 days. Cisplatin and OOEO were injected intraperitoneally. GSE was administered with gavage. The histopathological examination of liver tissues was performed by light microscope. Superoxide dismutase (SOD), glutathione peroxidase (GSH–Px) activities and malondialdehyde (MDA) levels were determined in liver tissues. Alanine aminotransferase (ALT) and aspartate aminotransferase (AST) levels were measured in serum. Cisplatin-induced considerable hepatocyte damage under microscopic examination, and an increase in MDA levels as well as a decrease in the activities of antioxidant enzymes, including SOD, GSH–Px in liver tissues and an increase in serum ALT and AST (p< 0.001). The addition of GSE and/or OOEO significantly prevented microscopic tissue damage, reversed oxidative stress parameters and biochemical values compared to the cisplatin group (p< 0.001). In conclusion GSE and OOEO may be used in adjuvant therapy to prevent cisplatin–induced hepatotoxicity, but further studies using various doses, different time intervals, and a larger number of animals need to be carried out.

Keywords: Cisplatin, Hepatotoxicity, Free Radicals, Grape, Thyme

ÖZET
Sisplatin, kanser tedavisinde yaygın olarak kullanılan fakat tedavi dozlarında dağı karaciğer hasarı gibi iyi istemeyen yan etkileri sebep olabilen bir ilaçtır. Sisplatinin karaciğer dokusunda yaptığı hasarın altında yatan mekanizma büyük oranda serbest oksijen radikallerine bağlıdır. Bu çalışmada sıçanlarda sisplatin tedavisine bağlı oluşan karaciğer hasarında üzüm çekirdeğinin özütünün (ÜÇO) ve esansiyel kekik yağınnı (EKY) muhtemel koruyucu etkilerinin araştırılması amaçlanmıştır. Doksanaltı adet Wistar albino cinsi erkek sıçan; biri oniki hayvanın olmak üzere sekiz gruba bölünerek (Kontrol, ÜÇO, EKY, ÜÇO+EKY, Sisplatin, ÜÇO+Sisplatin, EKY+Sisplatin, ÜÇO+EKY+Sisplatin), on gün süreyle takip edildi.
INTRODUCTION

Cisplatin (cis-diamminedichloroplatinum II) is one of the most widely used antineoplastic agents in the treatment of solid tumour and haematological malignancies. Despite its significant antitumour activity, the clinical use of cisplatin is often limited by its undesirable side-effects e.g. nephrotoxicity and neurotoxicity, which are examples of the most severe and dose limiting effects. Nevertheless, other less frequent toxic effects, such as hepatotoxicity, which is frequently seen after the administration of high doses of cisplatin, can alter the clinical situation of patients. Although cisplatin-induced nephrotoxicity has been very well documented in clinical oncology, hepatotoxicity has been rarely characterized and less studied. El-Sayyad et al. showed that cisplatin induced drastic abnormalities in liver tissue histology and the ultrastructure with respect to direct toxic effect on hepatocyte organelles. Like other chemotherapeutics, cisplatin causes excessive reactive oxygen species (ROS) generation and also induces a decrease in plasma antioxidant levels, which may reflect a failure of the antioxidant defence mechanism against oxidative damage.

Superoxide dismutase (SOD) and glutathione peroxidase (GSH-Px) are endogenous antioxidants and their natural supplements play a role in the prevention of cisplatin-induced oxidative injury in cancer patients. When an oxidant is administered as an adjuvant therapy in rats receiving cisplatin, a significant clinical improvement is obtained. It has been demonstrated that a high dose of vitamin E, or a combination of antioxidants, restored GSH-Px activity with a concomitant reduction in malondialdehyde (MDA) levels for prevention of cisplatin side effects. In another study, cisplatin induced mitochondrial sensitivity prevented by the thiol group of protective antioxidant agents, namely glutathione, dithiothreitol, N-acetyl-L-cysteine and cysteine, whereas SOD, catalase and ascorbate.

Grape seed extract (GSE) contains a number of polyphenols, including procyanidins and proanthocyanidins, which are powerful free radical scavengers. Grape seed proanthocyanidin extract is a rich source of polyphenolic antioxidants, a naturally occurring family of oligomeric proanthocyanidins found in a wide range of fruit and vegetables. A number of studies have demonstrated the superior free radical scavenging ability of GSE as compared to vitamins C, E and β-carotene.

Turkish oregano (Origanum onites L.), a member of the mint family, Lamiaceae, is well known for its antioxidative activity. Carvacrol and thymol, the two main phenols that constitute about 78-82% of the essential oil of oregano, are principally responsible for its activity. Origanum also contains phe-nolic terpenoids, flavonoids, tannins, phenolic glycosides and sitosterol. Authors have reported a broad spectrum of pharmacological and therapeutic effects provided by origanum including antioxidant, antispasmodic, antitumoural, antimicrobial, antifungal, diuretic and analgesic activities.

The aim of this study was to investigate the role of oxidative stress on cisplatin-induced hepatotoxicity and the possible protective roles of antioxidant food supplementation on this toxicity through tissue oxidative/antioxidant parameters and light microscopic evaluation.

MATERIALS and METHODS

Animals: Ninety-six adult male Wistar albino rats, initially weighing 230-310 g at 16 weeks old, were purchased from The Animal Laboratory of the Faculty of Medicine at Erciyes University. The animals were fed with a commercial standard diet and...
water ad libitum during the experimental period, and were kept at an ambient room temperature of 20-22°C, with a relative humidity of 50±5%; 12:12 h light and dark cycles were maintained. All animals were housed for acclimatization for 1 week before the experiments. This research was performed in accordance with the National Guide for the Care and Use of Laboratory Animals and approved by the Local Ethics Committee at the University.

**Experimental design:** The animals were randomly divided into eight groups (Control, GSE, OOEO, GSE+OOEO, Cisplatin, GSE+Cisplatin, OOEO+Cisplatin, GSE+OOEO+Cisplatin), each group consisting of 12 rats. As a preventive supplement GSE (100 mg/ kg-bw solution was administered with gavage by a curved 3-inch long 16-gauge gavage tube inserted after applying a proper restraint) and OOEO (1 mg/ kg-bw, intraperitoneally), and as a placebo distilled water (orally with gavage) were given to each group throughout the 10 days. The GSE+OOEO group was administered with two preventive supplements in the same dose and in a similar manner. As a therapeutic implementation physiological saline was given to the control, GSE, OEO and GSE+OEO groups (ip, as a placebo) and cisplatin (7 mg/ kg, single dose, intraperitoneally) was given to the cisplatin, GSE+cisplatin, OOEO+cisplatin and GSE+OOEO+cisplatin groups on the 6th day. On the 11th day liver tissue and serum samples were obtained from the rats under deep anaesthesia. The liver samples were then removed from the animals and immediately stored at -30°C until analysis.

**Preparation of Grape Seed Extract:** Ripened grapes (*Vitis vinifera* L) from the most popular wine making grape cultivars grown in Turkey, Okuzgozu (red grape cultivar), were obtained from the Tokat region of Turkey. The grape seed powder (100 g) was extracted in a Soxhlet extractor with petroleum ether (60°C for 6 hours) to remove the fatty materials. The defatted grape seed powder was re-extracted in a Soxhlet apparatus for 8 hours with 200 ml ethanol. The extract was then concentrated in a rotary evaporator (Rotavator Evaporator R 200, Buchi, Switzerland) under vacuum at < 40°C to lyophilize the crude extracts (*Labconco Freezone 2.5, Missouri, US*). It was then stored in a dark bottle at 4°C until use. The concentration of total phenolic compounds in the seed extract was determined by the Folin-Ciocalteu colourimetric method. The total phenolics were expressed as gallic acid equivalents (milligram GAE per gram extract).20 The content of total phenolics was found as 587.4±5.6 mg GAE/g in the grape seed extracts.

**Preparation of wild oregano essential oils:** *Origanum onites* samples were collected from Izmir (Turkey). The wild oregano (*Origanum onites*) essential oils were produced by the Clevenger hydrodistillation method. Plant materials (500 g), cut into small pieces, were placed in a distillation apparatus with double distilled water and hydrodistilled for 3 h. The essential oil was dried over anhydrous sodium sulfate, filtered and stored at -60°C until analysis.21 The chemical compositions of the wild oregano essential oils were found as 0.92% myrceane, 0.73% α-terpinene, 3.02% α-terpinene, 2.91% p-cymene, 5.18% linalool, 6.21% thymol and 80.23% carvacrol.

**Chemicals:** All chemicals were obtained from Sigma Chemical Inc. (St. Louis, MO, USA) and all organic solvents from Merck Chemical Inc. (Darmstadt, Germany) except for cisplatin. Cisplatin (Cisplatinum Ebewe 1 mg ml⁻¹) was obtained from the Liba Drug Company, Istanbul, Turkey. All reagents were analytical grade. All reagents except for the phosphate buffers were prepared each day and stored in a refrigerator at +4°C. The reagents were equilibrated at room temperature for 0.5 h before use, either at the start of analysis or when reagent containers were refilled. Phosphate buffers were stable at +4°C for 1 month.

**Biochemical determinations:** The livers were excised immediately and homogenized in a 10-fold volume of phosphate buffer solution at pH 7.4, using a homogenizer (Ultra-Turrax T 25, IKA; Werke 24.000 r.p.m, Germany). All preparation procedures were performed at +4°C. The homogenates were centrifuged at 10,000 x g for about 60 min and the resulting supernatants were stored at -80°C until for MDA, SOD and GSH-Px assays.

**Determination of MDA Level:** The levels of MDA in liver tissue were assessed according to the method of Ohkawa et al.22 The assay procedure to determine the MDA levels in the rat livers was set up as follows: to samples less than 0.2 ml of 10% (w/v) tissue homogenate, 0.2 ml of 8.1% sodium dodecyl sulphate and 1.5 ml of 20% acetic acid so-
olution were added. The pH was adjusted to 3.5 with NaOH and 1.5 ml of 0.8% aqueous TBA solution. The final volume was brought to 4 ml by distilled water and then heated in an water bath at 95°C for 60 min using a glass ball as a condenser. After cooling with tap water, 1.0 ml of distilled water and 5.0 ml of an n-butanol and pyridine (15:1 v/v) mixture were added and the mixture was shaken vigorously. After centrifugation at 4.000 rpm for 10 min, the organic layer was removed and its absorbance was measured at 532 nm. Tissue MDA levels were expressed in nanomoles MDA per milligram of protein in tissue homogenates (nmol/mg protein). Protein concentrations were measured by the Lowry method.

**Determination of SOD Activity:** Total (Cu-Zn and Mn) SOD (EC 1.15.1.1) activity was determined according to the method of Sun et al. The principle of this method is based on the inhibition of NBT reduction by the xanthine–xanthine oxidase system as a superoxide generator. Activity was assessed in the ethanol phase of the supernatant after a 1.0 ml ethanol/chloroform mixture (5/3, v/v) was added to the same volume of sample and centrifuged at 4000 g. One unit of SOD was defined as the amount of enzyme causing 50% inhibition in the NBT reduction rate. SOD activity was also expressed as units per milligram liver protein.

**Determination of GSH-Px Activity:** GSH-Px (EC 1.6.4.2) activity was measured by the method of Paglia et al. The enzyme reaction in a tube containing NADPH, reduced glutathione (GSH), sodium azide, and GSH reductase was initiated by the addition of H2O2 and the change in absorbance at 340 nm was monitored with a spectrophotometer. Activity was given as units per gram protein in liver tissue and all samples were assayed in duplicate.

**Determination of Liver Functions:** Serum ALT and AST levels were measured to evaluate liver function by using reagents (Thermo Electron CO, USA) for Olympus Analysers. All liver enzyme assays were performed spectrophotometrically using an Olympus AU 2700 autoanalyzer (Olympus CO, Tokyo, Japan). ALT, AST levels were also expressed as units per litre.

**Histopathological Examinations:** For the histopathological examinations, the liver tissue samples taken from the rats were fixed in 10% phosphate buffered formalin and embedded in paraffin. Then, 5 µm sections were cut from the paraffin blocks and stained with hematoxylen-eosin. Liver sections were blindly observed under a light microscope and scored to assess histopathological changes according to the following grading system: 0, normal; 1, mild hydropic degeneration, no Kupffer cell proliferation and no necrosis; 2, moderate hydropic degeneration, Kupffer cell proliferation and no or little necrosis; 3, severe hydropic degeneration, Kupffer cell proliferation and necrosis.

**Statistical Analysis:** Data were expressed as mean±standard deviation (±SD) (for normally distributed data), or as median with 25%-75% percentile (for skewed data). Comparisons of MDA, SOD, GSH-Px, AST and ALT between the groups were made by using a One Way Analysis of Variance (ANOVA). Post-hoc comparisons on parameters were performed using the Tukey procedure. Comparisons of groups for histopathological examinations were made by using a Kruskal-Wallis analysis of variance (Post-hoc comparisons: Dunn’s test). All analyses were performed with the statistical package for scientists (SIGMASTAT) Windows version 3.5 and p values less than 0.05 were regarded as statistically significant.

**RESULTS**

Cellular damage was observed in liver the samples of cisplatin-injected animals. The liver morphology was characterized by severe activation of Kupffer cells, degenerated hepatocytes and moderate enlargement of the sinusoids (Figure 1). The microscopic damage score of the liver samples was significantly higher in the cisplatin-injected group (median score: 2, range 1.75-2.25) than those of the other groups (p < 0.05); Cisplatin: 2 (range 1.75-2.25). The addition of GSE and OOEO effectively improved microscopic appearance and also significantly reduced the histopathological scores (median scores: GSE+Cisplatin: 1, range 1-2; OOEO+Cisplatin: 1, range 1-2; GSE + OOEO + Cisplatin: 1, range 1-1.5) as compared to cisplatin-treated groups (p < 0.05). The biochemical oxidation parameter results are expressed in Figure 2. All groups were compared with each other statistically. GSE and GSE+OOEO administration significantly decreased MDA levels.
as compared to the control group (p < 0.001), while the addition of OOEO decreased the MDA level to a lesser degree but this decrement was not remarkable (p > 0.05). The MDA levels in the rats receiving cisplatin were significantly higher than in the other groups (p < 0.001). However, the addition of GSE and/or OOEO to the treatment significantly decreased the cisplatin-induced MDA increment (p < 0.001; Figure 2a)

GSE and GSE+OOEO administration significantly increased SOD and GSH-Px activities as compared to the control group (p < 0.001), while OOEO addition did not change antioxidant activity compared to the controls (p > 0.05). While cisplatin treatment significantly decreased liver SOD and GSH-Px activities (p < 0.001; Figure 2b and Figure 2c), GSE and/or OOEO treatment reversed these reductions in SOD and GSH-Px activities in the cisplatin group (p < 0.001).

As expected, cisplatin caused significant increases in serum ALT and AST levels when compared to control levels (Figure 3a - 3b; p < 0.001). Treatment with GSE and/or OOEO markedly reversed cisplatin-induced alterations in serum hepatocyte enzymes levels (p < 0.001).

DISCUSSION

Cisplatin, one of the most active cytotoxic agents against cancer, has several toxicities. Hepatotoxicity occurs during high dose treatment with cisplatin.27,28 El-Sayyad et al showed many histopathological abnormalities including inflammatory infiltration, hyperplasia, periportal fibrosis, marked disruption of hepatic cords and dilated blood sinusoids in the rats livers of receiving cisplatin. These changes were confirmed at ultrastructural level, including vesiculated rough endoplasmic reticulum and atrophied mitochondria.7 The data in our study also revealed that even a single dose of cisplatin (7 mg kg⁻¹) causes considerable damage in the liver, as assessed microscopically. The liver morphology was characterized by severe activation of Kupffer cells, degenerated hepatocytes and moderate enlargement of the sinusoids. The present study also showed also clinical evidence of cisplatin-induced liver injury, demonstrated by elevated activities of serum enzymes ALT and AST.

In various studies, it has been shown that cisplatin administration is associated with increased formation of free radicals and generates active oxygen species, such as superoxide anion and the hydroxyl ra-
In the present study, an increment in MDA, a highly reactive compound and remarkable marker for oxidative stress and a decrement in GSH-Px and SOD activities, which are key antioxidants, were found in the liver of rats treated with cisplatin. After understanding the possible role of free radicals in cisplatin-induced oxidative liver damage, several agents were added to the therapy as an adjuvant supplement and they attempted to protect and/or to prevent the side effects of chemotherapeutics. Recent studies have shown that different natural antioxidants such as vitamin C, vitamin E, ellagic acid, nigella sativa, lycopen, caffeic acid phenethyl ester prevent cisplatin-induced toxicity. In this study, we investigated the effects of the natural antioxidants GSE and OOEO in cisplatin-treated rats by microscopic examination and biochemical assays.

Most grape products are rich sources of dietary chemical constituents that may have a potential for the prevention and treatment of human malignancies. These products have generated remarkable interest based on positive reports of their antioxidant properties and their ability to serve as free radical scavengers. Several studies have indicated that GSE inhibits the oxidation of LDL, inhibits the enzyme systems that are responsible for the production of free radicals is antimutagenic, and anticarcinogenic. Hence, GSE is widely consumed as a dietary supplement and could be useful in synergizing the efficacy of cancer chemotherapeutic agents in cancer treatment. The protective effect of GSE against chemotherapy-induced toxicity may be attributed to its potential to act as a hydrogen-donating free radical scavenger and singlet oxygen quencher and to their antioxidant activity. Similarly, in this study, the combined treatment of cisplatin and GSE resulted in a significant decrease in the MDA levels of liver tissue, a significant elevation in the antioxidant GSH-Px and SOD activities of tissues indicating the protection offered by GSE against cisplatin injury.

Carvacrol, the main constituent of OOEO, is an oxygenated monoterpene with multiple pharmacological actions including antioxidant, antitumoural, hepatoprotective activities. It is also well known that essential oils which are rich in carvacrol possess strong antioxidant properties equivalent to those of ascorbic acid, butyl hydroxy toluene and vitamin E. In the present study OOEO ameliorated the side effects of cisplatin by inhibiting lipid peroxidation. Contrary to our expectations, the
use of OOEO with GSE did not have a synergical antioxidant effect on cisplatin-induced oxidative stress. This additive effect may have a similar antioxidant mechanism but further in-vivo and in-vitro experimental sets up are needed to explain the exact mechanism.

In conclusion, we showed that the experimental administration of cisplatin is accompanied by increased lipid peroxidation in rats. We therefore suggest that oxidative stress is a cause of cisplatin-induced pathophysiology. Simultaneously, we studied the co-administration of GSE and/or OOEO as an approach to ameliorate cisplatin-induced hepatotoxicity. These agents may protect against cisplatin-induced toxicity by overcoming the inactivation of antioxidant enzyme systems induced by cisplatin, and upregulating of SOD and GSH-Px activities in the liver. Our findings could provide a more promising strategy for the prevention of hepatotoxicity in cisplatin-based chemotherapy. However, additional studies with various doses, different time intervals, and a larger number of animals are necessary.

REFERENCES

Correspondence
Dr. Aysun ÇET‹N
Erciyes Üniversitesi Tıp Fakültesi
Biyokimya ve Klinik Biyokimya Anabilim Dalı
38039, Kayseri / TURKIYE
Tel: (+90.352) 437 93 48
e-mail: aysuncetin@yahoo.com