Investigation of the Genotoxic, Cytotoxic, Apoptotic, and Oxidant Effects of Olive Leaf Extracts on Liver Cancer Cell Lines

Zeytin Yaprığı Ekstrelerinin Karaciğer Kanseri Hücre Hatları Üzerine Genotoksik, Sitotoksik, Apoptotik ve Oksidan Etkilerinin Araştırılması

ABSTRACT

Objectives: Hepatocellular carcinoma (HCC) is the seventh most common cancer and the third leading cause of tumor-related deaths worldwide. Mechanisms underlying tumor onset, progression, and metastasis in the case of HCC have not been adequately studied. In this study, we aimed to investigate the genotoxic, cytotoxic, apoptotic and oxidant effects of olive leaf extract (OLE) on HCC cells.

Materials and Methods: H4IIE Rattus norvegicus hepatoma cells and Rattus norvegicus healthy liver clone-9 cells were treated with the increasing concentrations of OLEs (250-2000 ppm) in ethanol, acetone, dichloromethane, and methanol. ATP cell viability, intracellular reactive oxygen species generation levels, double staining test with acridine orange/ethidium bromide, comet assay, levels of interleukin 1-beta (IL-1β), IL-6, and tumor necrosis factor alpha were measured. Significance was determined using ANOVA test.

Results: Apoptotic, genotoxic, cytotoxic, and oxidative effects of OLEs increased with the increasing concentrations as compared to controls in H4IIE cells (p<0.001).

Conclusion: This is the first study to show a significant and selective cytotoxic activity of OLEs in the selected H4IIE cancer cell lines. OLEs could selectively increase the apoptotic damage and show anti-proliferative and pro-apoptotic properties against the H4IIE cells. They could be recommended as potential nutraceuticals in the prevention of cancer.

Key words: Olive leaf, hepatocellular carcinoma, oleuropein, oxidative stress, genotoxicity, apoptosis

ÖZ

Amaç: Hepatoselüler karsinoma (HSK), en yaygın yedinci kanser türüdür ve kanserle ilişkili ölümlerde üçüncü sıradadır. HSK’de tümör başlangıcına, ilerlemesine ve metastazlara neden olan mekanizmalar tam olarak bilinmemektedir. Bu çalışmada zeytin yaprağı ekstresinin (OLE) HSK hücreleri üzerine genotoksik, sitotoksik, apoptotik ve oksidan etkilerinin araştırılması amaçlanmıştır.

Gereç ve Yöntemler: H4IIE Rattus norvegicus hepatoma hücrelerine ve Rattus norvegicus sağlıklı karaciğer klon-9 hücrelerine etanol, aseton, diklorometan ve metanol içinde OLE’leri (250-2000 ppm) uygulanmıştır. Ekstrelerin sitotoksikitesi ATP testiyle, hücre içi reaktif oksijen tüplerinin oluşumu florometrik yöntemle, genotoksik etkileri alkalen tekli hücre jel elektroforez (comet assay) yöntemiyle, apoptotik etkileri akırdın turuncusu/etidyum bromür yöntemiyle ölçülmüştür. İnterlökin 1 beta (IL-1β), IL-6 ve tümör nekroz faktörü alpha düzeyleri ELISA yöntemi ile belirlenmiştir. İstatistiksel test olarak ANOVA testi kullanılmıştır.

Bulgular: OLE’nin apoptotik, genotoksik, sitotoksik ve oksidatif etkileri H4IIE hücrelerinde kontrol göre artan konsantrasyonlarla istatistiksel olarak anlamlı şekilde yükselemiştir (p<0.001).
INTRODUCTION

Hepatocellular carcinoma (HCC) is the seventh most common cancer and the third leading cause of cancer-related mortality worldwide. Despite significant advances in the diagnosis and treatment, HCC remains a terminal disease. The most common cause for mortality in HCC is progression and the metastasis. However, the molecular mechanisms underlying tumor onset, progression, and metastasis in the case of HCC have not been adequately studied. The neoplastic development of HCC is related to many histological incidents. Hyperplastic nodules formed in the hepatocyte regeneration observed in the liver in response to the cell deaths due to risk factors are cytologically of normal findings, which will be the first step in HCC formation. These lesions may transform into premalignant dysplastic nodules and cause nuclear aggregation with cytologically well-detected cell changes. As a result of molecular analysis in the HCCs, various genetic and epigenetic changes have been detected.

Epidemiological studies and animal experiments have been revealed that some nutritional compounds play a role in the incidence rates of various cancers, including HCC. One-third of all the known human cancers may be related to specific components of nutrients. Many research groups are investigating the effects of nutritional components on cancer processes and state that dietary fats increase the risk of brain, colon, breast, and prostate cancer. In addition, the incidence of the brain, cardiovascular diseases, and colon and breast cancers are lower in the Mediterranean Region than in Europe. Around the Mediterranean and European territories olive tree leaf (Olea europaea L.) is commonly used as traditional medicine. Olive leaf extract (OLE) has been reported to show anti-aging, immune system enhancing, and antimicrobial effects. In studies, olive leaf antioxidants have been shown to be effective in the treatment of cancers like liver, prostate, and breast. Olive leaf contains substantial secoiridoids, flavonoids, phenolic acids, and the lignans. Of these, oleuropein is an important phenolic compound in the secoiridoid group. Oleuropein and its metabolite hydroxytyrizole (HT) have antioxidant activity, showed in both in vivo and in vitro experimental models. In addition to in vivo and in vitro activities, the antioxidant potential of oleuropein and HT has also been investigated on experimental cell line models and animal models. In recent epidemiological studies, it was shown that OLEs rich in phenolic compounds correlated with a decreased cardiovascular risk, neurodegenerative disease, and cancer. Besides the different phenolics and flavonoids olive leaves contain the most oleuropein content when compared with other parts of the plan such as root, bark, fruit. In this study, it was aimed to investigate the in vitro genotoxic, cytotoxic, apoptotic, and oxidant effects of OLE on the HCC cell lines.

MATERIALS AND METHODS

Collection of plant samples

The plants have been purchased commercially as medical drugs with a batch number of 26/0/2019. Plant leaves of Olea europaea L. were dried at room temperature, then shredded with a plant shredder and stored at room temperature until they were used in the study.

Preparation of methanol, acetone, dichloromethane, and aqueous-ethanol extracts

The preparation of OLEs was performed for the methanol extract (O-MeOH), acetone (O-ACE), dichloromethane (O-DCM), and aqueous-ethanol extract (O-EtOH), a total of 70 g of the plant sample was ground with a laboratory blender, extracted in the solvent, and after the extraction, the extract has been filtered, followed by the evaporation of the solvent. The amount of oleuropein was expressed as milligram per gram of OLE. The OLE was dissolved in dimethyl sulfoxide (0.1%) prior to the analysis.

Determination of total phenolic and flavonoid content

The total amount of phenolic compounds found in the extracts was determined with the Folin-Ciocalteu Reagent. Gallic acid (GA) was used as a standard phenolic compound. The total amount of phenolic compounds found in all OLEs was evaluated according to the study conducted by Gülcin et al. The findings were given in GA equivalent (GAE) and micrograms.

Determination of free radical activity

The free radical scavenging activities of the extracts were determined using the 1,1-Diphenyl-2-picrylhydrazyl (DPPH) free radical. The absorbance was measured at 517 nm, values of the samples were evaluated against the control. As standard, bybutylated hydroxy anisole (BHA), BH toluene (BHT), and α-tocopherol (α-Toc) were used.

Determination of ABTS cation radical scavenging activity

This method was used to assess the potential of the extracts to sweep the ABTS cation radical. The absorbance of each solution was detected at 734 nm, and the inhibition % was calculated.

Determination of reduction force

The cupric ion (Cu²⁺) reduction capacities of the extracts were measured with the CUPRAC method. As standard, BHA, BHT, and α-Toc were used.
**High-performance liquid chromatography (HPLC) analysis**
The amount of oleuropein in leaf extracts was measured by HPLC according to the method developed by Al-Rimawi. HPLC analysis was performed using the Shimadzu (Japan) LC-20A model, and the peak areas of automatic injections of 20 μL with ultraviolet (UV) detector were calculated using the LC solution computer software. Chromatographic separation was attempted to be optimized using C
\textsubscript{18} (5 μm, 150×4.6 mm) and C
\textsubscript{8} (5 μm, 150×1.6 mm) columns.

The calibration coefficient was calculated from the calibration curve created by the concentration-dependent linear mathematical equation of the field data, and linearity was calculated (Figure 1G). The UV detection of the separation was performed using the mobile phase of the water (pH: 3): acetonitrile (70:30) at 1 mL min
\textsuperscript{-1} flow over the C
\textsubscript{18} ODS-3 (150 mm, 4.6×5 mm) column at 280 nm. The linear coefficient of the linear calibration curve created in the range of 100-1000 μg/mL was found to be 0.99.

**Cell culture**
H4IIE [American Cell Cultures Collection (ATCC)\textsuperscript{R} CRL-1548\textsuperscript{TM}] Rattus norvegicus hepatoma cells were obtained from the (ATCC, Middlesex, UK). Cells were grown in the Eagle's minimum essential medium containing 10% FBS and 1% P/S in a humidified incubator with 5% CO
\textsubscript{2} at 37°C.

Healthy rat liver cell clone-9 cells (ATCC\textsuperscript{R} CRL-1439\textsuperscript{TM}) were obtained from ATCC (Middlesex, UK). Cells were grown in an F12K medium containing 10% FBS, and 1% P/S in a humidified incubator with 5% CO
\textsubscript{2} at 37°C. Samples were prepared at different doses and left for 24-hour incubation in cells. Each dose was analyzed with a minimum of four replicates.

**Cytotoxicity assay**
To measure cytotoxicity in cancer and healthy cells, 1.5×10
\textsuperscript{4} cells per well were added to 96 opaque white plates. Samples of different concentrations were added and incubated for 24 hours. After incubation, the ATP solution was added to the wells and measured luminometrically using the Cell Titer-Glo\textsuperscript{R} Luminescent Cell Viability Test Kit.

**Intracellular reactive oxygen generation**
The intracellular reactive oxygen species (ROS) levels were measured using the H
\textsubscript{2}DCF fluorescence probe. Cancer and healthy cells seeded on the opaque black plates, and samples of different concentrations (250-2000 ppm) were incubated for 24 hours. After the incubation, cells were washed three times with dPBS and incubated at 37°C in the dark with 5 μM H
\textsubscript{2}DCFH-DA. After the cells were washed, fluorometric measurements were taken at Ex: 482 nm/Em: 512 nm. Results were normalized according to the ATP levels.

**Apoptosis**
Acridine orange (AO)/ethidium bromide (EB) dye was used to measure the apoptosis.\textsuperscript{17} After mixing AO/EB and cell pellet in a 1:1 ratio, images were taken under a fluorescence microscope (Leica DM 1000, Solms, Germany). The separation of apoptotic cells performed according to the morphological characteristics of the nuclei. Apoptotic cell nuclei take up only EB. On the other hand, AO is taken up by both living and dead cells. Green fluorescence at 480-490 nm was flared by living cells. Healthy cells flared green fluorescence while apoptotic cells luminating yellow-orange and necrotic cells were seen red. Briefly, 2×10
\textsuperscript{5} H4IIE and clone-9 cells/well were seeded in 6-well plates and incubated for 24 hours. Cells seeded in 6-well plates under IC
\textsubscript{50} doses were washed for 24 hours after incubation with the samples. The preferred solution for collection of the cells and washing was PBSR, then cells were stained with 1:1 mixture of AO/EB (100 μg/mL). Triplicate samples of 100 cells each were counted and scored for the incidence of apoptotic chromatin condensation using a fluorescent microscope (Leica DM 1000, Solms, Germany).\textsuperscript{17,18} Apoptotic cells were calculated relative to the total number of cells.\textsuperscript{18}

**Genotoxicity**
Genotoxicity analysis was performed according to the comet assay method in our previous study. DNA damage in cells was given as tail intensity percentage. Comet assay (alkaline single cell gel electrophoresis assay) with a slight modification of Singh et al.'s methods was carried out to assess the genotoxic effects of OLEs on H4IIE and clone-9 cells. The 6-well cell culture containing the cell culture medium plates were used. For the evaluation of genotoxicity approximately 2×10
\textsuperscript{5} cells per well plated on wells at 37°C. Then, the OLE samples < IC
\textsubscript{50} concentrations were added and incubated for 24 hours. Trypsin-EDTA (0.25%) used for collection of the cells and, phenol red for 2-3 min in the incubator. The cells were centrifuged at 1500x rpm for 5 min at 4°C. The supernatant was withdrawn, and the cell density of 2×10
\textsuperscript{5} cells/mL was adjusted with cold dPBS. Total of 15 μL cell suspension and 85 μL of 0.6% low melting point agarose were mixed and placed on 1% normal melting point agarose precoated slides. The gel mixture was solidified on a cold tray for 2 mins, and the slides were treated for overnight with lysis buffer, pH 10.0 (1% Triton X-100, 2.5 M NaCl, 10 mM Tris, 0.1 M EDTA, Sigma-Aldrich). In an alkaline solution, the incubation of the slides performed (0.3 M NaOH, 1 mM EDTA, Sigma-Aldrich) for 40 min in the dark in the presence of cooling blocks to unwind the DNA. At 0.72 V/cm (26 V, 300 mA) for 25 min in an electrophoresis tank at 4°C, 0.72 V/cm (26 V, 300 mA) electrophoresis was performed for 25 min. The slide was again treated with tris buffer (0.4 M Tris, pH: 7.5) for 5 mins. Before staining the slides, ethanol was used for dehydrating the slides. A staining procedure with EB (2 μg/mL in distilled H
\textsubscript{2}O, 70 μL/slide) applied to the slides and coated with a coverslip and scored under fluorescence microscope (Leica DM 1000, Solms, Germany) using the Comet assay IV software (Perceptive Instruments, Suffolk, UK).

**Inflammation assays**
Cancer and healthy cells were incubated at concentrations < IC
\textsubscript{50} for 24 hours. The levels of interleukin-1β (IL-1β), IL-6, and tumor necrosis factor-α (TNF-α) in the medium were measured by commercially available ELISA kits (Elabscience, Texas, USA) by photometric methods. Levels of IL-1β, IL-6, and TNF-α in a cell medium after 24-hour incubation were measured by the ELISA.
method according to the protocol provided by the manufacturer. All ELISA kits were purchased from Elabscience (Texas, USA). IL1β, IL6, and TNF-α levels were read at 450 nm using an ELISA plate reader (Thermo Scientific, Massachusetts- USA), and the concentrations were calculated according to the standard curves obtained from the external standards.

Statistical analysis
All experiments were repeated for the minimum number of 4 times. Results are given as mean ± standard deviation. Variance analysis was done with One-Way ANOVA. p<0.05 level was considered as statistically significant. Regression analysis was done for IC50 values. All analyzes were done with the Statistical Package Program for Social Sciences (IBM, Inc version 25).

RESULTS
Flavonoid and phenol content, free radical activity, ABTS cation radical scavenging activity, reduction force, and oleuropein level analysis
Polyphenolic compounds, namely flavonoids, phenolic acids, and their derivatives, are typical phytochemicals for olive leaves. Total flavonoid levels in O-DCM, O-ACE, O-MeOH, and O-EtOH were found 8.52, 7.18, 3.29, and 1.73 mg QUE/mL, respectively (Figure 1A), while total phenolic levels in O-DCM, O-ACE, O-MeOH, and O-EtOH were 11.24, 6.92, 9.96, and 9.81 mg GAE/mL, respectively (Figure 1B).

The free radical scavenging activities of the extracts are determined using the DPPH free radical (Figure 1C). O-DCM showed inhibitions of 0.13%, 0.13%, 18.72%, and 18.85% with different concentrations. O-ACE showed inhibitions of 9.49%, 18.72%, 41.54%, and 81.85% with different concentrations. O-MeOH showed inhibitions of 6.63%, 19.76%, 42.78%, and 81.85% with different concentrations. BHA was used as a standard.

ABTS cation radical scavenging activity of the extracts was shown in Figure 1D. O-DCM showed inhibitions of 9.96%, 0.13%, 18.72%, and 18.85% with different concentrations. O-ACE showed inhibitions of 30.32%, 65.82%, 70.54%, and 87.14% with different concentrations. O-MeOH showed inhibitions of 18.13%, 50.43%, 85.49%, and 85.93% with different concentrations. O-EtOH showed inhibitions of 42.30%, 69.89%, 87.36%, and 86.92% with different concentrations. The determination of reducing ability was showed as absorbance levels of extracts (Figure 1E). O-DCM showed absorbance of 0.14, 0.24, 0.31, and 0.52 with different concentrations. O-ACE showed absorbance of 0.20, 0.43, 0.70, and 1.24 with different concentrations. O-MeOH showed absorbance of 0.17, 0.39, 0.60, and 1.04 with different concentrations. BHA was used as a standard. Oleuropein level determination was conducted with HPLC as three parallels (Figure 1E). Mean values of these three were 19.56 mg/mL (O-DCM), 97.97 mg/mL (O-ACE), and 72.20 mg/mL (O-MeOH).

Inhibition of cell viability
Cytotoxicity results of OLEs to liver hepatoma (H4IIE) and healthy liver clone-9 cells are presented in Figure 2A. These results showed a robust concentration dependent response relationship with cytotoxicity of OLEs. In Figure 2A, a gradual decrease was seen in the viability of H4IIE cells, a statistically significant difference obtained p<0.001 with increased concentrations.

Intracellular iROS generation levels
The results of the iROS of OLEs to liver carcinoma (H4IIE) and healthy liver clone-9 cells are presented in Figure 2B. There is a gradual increase in the iROS level of H4IIE cells, with all concentrations of O-MeOH and O-EtOH of 250, 500, 1000, and 2000 ppm showed increased iROS compared to control cells (p<0.001) (Figure 2B).

Induction of apoptosis
OLEs induce Apoptosis of H4IIE cells in a concentration-dependent manner, showing a gradual increase of EB-positive cells in OLEs-treated cells compared to the control, and a statistically significant difference was obtained p<0.001 (Figure 2C, D).

Induction of DNA damage
Extracts at all concentrations lead to the breaking of DNA single-strand in H4IIE cells, and there is a gradual concentration-response relationship and a statistically significant difference of p<0.001. Comet assay images of control and OLEs-treated H4IIE cells are presented in Figures 3A, B.

Anti-inflammatory activity of OLEs and its phenolic compounds
Results showed that all extracts at the concentrations of 250, 500, 1000, and 2000 ppm causes elevated levels of IL-1β, IL-6, and TNF-α in H4IIE and clone-9 cells (Figure 4A-C). The OLEs had a significant activity on IL-1β, IL-6, and TNF-α production in a concentration-dependent manner.

DISCUSSION
Studies on Olea europaea L. generally includes data on the chemical and physical properties of the oil extracted from the fruits and their samples. There are few studies on the antioxidant activities of leaf extracts in the literature since the oil of the fruits is very valuable. Ethanol and acetone have been found to be highly effective solvents in extracting the phenolic compounds. Phenolic compounds are one of the main secondary metabolites of olive leaves.20-23 It has also been found that water/aqueous-ethanol/acetone mixtures are generally more effective solvents to extract the polyphenolic compounds of olive leaves.20-23
The antioxidant activity of OLEs may directly be related to the polyphenol content.21,23 Oleuropein, is the major secondary metabolite and has been extensively studied, but available information is limited for different OLEs.21,23 Oleuropein is an important metabolite for the antioxidant ability of OLEs.9 For the OLEs, acetone and ethanol extracts exhibit higher antioxidant capacity in both methods (DPPH and ABTS). Our results were consistent with the literature, and acetone and ethanol extracts with high oleuropein content were effective as antioxidant scavengers.23,26 In a study using solid-liquid extraction technique and three different extraction solvents
[petroleum ether, water, methanol (80%)], suggested that flavonoids (from 3.33-17.64 mg catechin equivalents/g), phenolics (from 3.64-21.47 mg GAE/g) were abundant in the content of OLEs, and their analysis correlated with antioxidant capacity.\textsuperscript{24} These data are consistent with our results presented in this study.

OLEs caused cytotoxicity in many different cancer cells such as human breast cancer cells (MCF7) and colorectal cancer cells (HT29).\textsuperscript{25,26} Kimura and Sumiyoshi\textsuperscript{27} investigated the effects of OLE and oleuropein on the skin cancer animal model. According to their investigations, oleuropein and OLE were superior to the control group in skin thickness and tumor incidence. Barbaro et al.\textsuperscript{28} suggested that olive leaves and oleuropein antitumor activity may be related with ROS scavenging effect, antiproliferative effect, apoptosis induction, angiogenesis inhibition, and antimigration effect. Our study is coherent with the previous studies, and we found that OLEs at different concentrations inhibits H4IIE cancer cells more.

**Figure 1.** Evaluation of the flavonoid and phenol content, free radical activity, ABTS cation radical scavenging activity, reduction force, and oleuropein level in Olea leaf extracts. (A) Total phenolic compounds of extracts; (B) total flavonoid compounds of extracts; (C) free radical activity of extracts for different concentrations; (D) ABTS cation radical scavenging activity of extracts for different concentrations; (E) reduction force of extracts; (F) oleuropein concentration in extracts. (G) Oleuropein calibration curve and extract spectrums of acetone, dichloromethane, and methanol. GAE: Gallic acid, QUE: Quercetin, O-DCM: OLE with dichloromethane, O-ACE: OLE with acetone, O-MeOH: OLE with methanol, O-EtOH: OLE with ethanol, BHA: Butylated hydroxy anisele
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than clone-9 healthy cells (Figure 2). In fact, at 0.25 mg/mL concentrations, acetone and methanolic extracts positively affected cell viability in terms of Relative Luminescence Units in the healthy clone-9 cell line. This indicates that OLE causes selective cytotoxicity in rat hepatoma cell lines, but not in healthy hepatocytes. This selectivity could be a result of the susceptibility of cancer cells to ROS formation. Cancer cells have increased ROS producing capacity, and therefore the selective activity of OLE on H4IIE may be due to their increased susceptibility to the ROS. The effects of OLE on H4IIE cells may offer opportunities for new chemoprevention studies for liver cancer.

On the other hand, the higher concentration of OLE was induced cytotoxicity and apoptosis for both healthy and cancer cell lines. These results indicate that the dose of the extracts have a crucial role in the pharmacological effect. According to concentration, OLE may induce or inhibit cytotoxicity and apoptosis. According to Park et al., mice fed with an oleuropein-supplemented diet attenuated hepatic steatosis in microscopic and macroscopic scale. According to the literature and our results, OLE or oleuropein-supplemented diet has protective effects on liver diseases.

In literature, via various mechanisms, olive leaves may induce apoptosis and inhibit cell proliferation. Oleuropein stops cell growth and promotes the apoptosis in HT29 colorectal cancer cells via a p53-dependent pathway. This apoptotic effect with a similar mechanism was also reported in breast cancer cells. In addition, on doxorubicin-induced cardiomyopathy, oleuropein, major phenolic of olive leave showed a protective effect through AMPK activation and iNOS suppression. This study revealed that OLE, which contains oleuropein as major secondary metabolites has antitumor activity in the hepatoma cell lines. These results are partly consistent with the apoptosis mechanism described in MCF-7 cells treated with oleuropein. Oleuropein is thought to be related the activation of AKT. Oleuropein may suppress the phosphorylation of AKT. Apoptotic effects of OLE may be related PI3K/AKT pathway, which is a critical pathway for H4IIE cells.

Antigenotoxic agents often show expected therapeutic effects that may be effective to control the cancer. Our results were
promising that OLE reduced cell viability in H4IIE cells and were consistent with the literature showing that OLE may inhibit the proliferation of malignant cells in vivo. This feature of OLE may be explained by its capacity to act as a strong free radical scavenger, as shown in Figure 1.

Our results pointed out, OLE have induced the IL-1β, IL-6, and TNF-α production in a concentration-dependent manner. A significant decrease in the proinflammatory cytokine levels was found in cells treated with different OLE. Our results were also consistent with the literature in terms of anti-inflammatory effects. In comparison with the healthy and cancer cell lines, anti-inflammatory effects were significantly higher in healthy cell lines. On the other hand, the increased inflammatory response of cancer cell lines is desirable, which starts apoptosis cascades. The possible anti-inflammatory mechanism may be related with arachidonic acid pathway, mitogen-activated protein kinase (MAPK), and MAPK enzymes such as p38, extracellular signal-related kinase, and c-Jun N terminal kinase induce transcriptional and post-transcriptional, activation of COX enzymes. Anti-inflammatory effects of O. europaea L. may be accomplished through NFκB activation. Inhibition of this proinflammatory cytokines production could be related either with the oleuropein or other secondary metabolites. Oleuropein is the fundamental OLE component that exhibits anti-inflammatory effects at a concentration of 20 μg/mL. In another study, OLE reduced mRNA expression of E-selectin in human coronary artery endothelial cells stimulated with serum amyloid-A, as well as decreased IL-6 and IL-8 protein levels and reduced matrix metalloproteinase 2 levels in non-stimulated cells. In both studies, the amount of OLE used was 20 μg/mL and 0.5-1 μg/mL, respectively. Compared with our OLE (0.25-2 mg/mL) concentrations, these results show that high amounts of OLE show pro-oxidant activity.

CONCLUSION

To the best of our knowledge, this is the first study in the literature investigating the cytotoxic, genotoxic, and apoptotic activity of OLEs on liver cancer cell lines. OLEs activated cytotoxic and apoptotic mechanisms on cancer cells. For this reason, OLEs may have a potential for the cancer treatment. More detailed in vivo studies are needed in the future.
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