Intercalation of Rhodamine B into Calf Thymus DNA in presence and absence of olive leaf extract: A fluorescence study

Abstract
Aqueous olive leaf extract (AOLE) of the Mediterranean tree Olea europaea is stated to have several therapeutic properties including: antifungal, antimicrobial, hypotensive and hypoglycemic actions as well as to use in treatment of some cancer cases as breast, skin and pancreatic cancers. Experimentally, the olive leaf extract is capable to protect nuclear calf thymus and human genomic DNA in vitro to Rhodamine B toxicity. Thus, this study was aimed to evaluate the protective effect of AOLE against toxic DNA damage induced by Rhodamine B using different techniques as UV–VIS Spectroscopy and fluorometry. Genomic DNA was used in a concentration of 75μg/ml. DNA was extracted from peripheral lymphocytes of anticoagulated blood (EDTA) samples by Proteinase K digestion and phenol/chloroform extraction. Dried leaves of Olea europaea (from West part of Libya) were grinded with distilled water; the resulting aqueous solution was then filtered and used. The protective effect of the AOLE was evaluated for its ability to decrease the amount of Rhodamine B available for binding with double stranded DNA. Thus, DNA resulted in a significant decrease in fluorescence intensity of Rhodamine B. Comparing to control samples, pre- and post-treatment with AOLE, significantly attenuated the binding ability of Rhodamine B to the DNA (p<0.01). Treatment with 5g/15 ml of AOLE was enough to prevent Rhodamine B toxicity to the DNA. The cytoprotective and antioxidant properties were detected in AOLE by a fluorescence method. The antioxidant activity of AOLE was evaluated using the 2,2-diphenyl-1-picrylhydrazyl (DPPH) assay and the IC50 was 65±2.3μg/ml. In conclusion, this study suggests that AOLE induced a genoprotective effect by increasing the antioxidant capacity and the chemical interaction between Rhodamine B and constituents of AOLE.

Keywords: Rhodamine B, olive leaf extract, ctDNA, genomic DNA, genoprotective

Abbreviations: RB, Rhodamine B; AOLE, aqueous olive leaf extract

Introduction
Interaction of small organic molecules with DNA has become an essential subject of medical research. Numerous attempts have been established by research groups to discover agents with a new therapeutic potential. Binding studies of various polyaromatic ligands to DNA have become a very promising exploration due to its relation to genetic effects and carcinogenicity. DNA interacting agents have exceptional been valuable in cancer biology due to their ability of therapeutic use. Many of presently used planar molecule anticancer agents exert their effect by interacting with DNA. Because of such reasons, characterization of the interaction of small molecular ligands as Rhodamine B (RB) with DNA has been the subject to some studies. RB is widely used for industrial purposes including printing and coloring of textiles, papers, paints, leathers etc. In addition, RB, a common dye of the Rhodamine family, has extensively been used by researchers due to its biological importance. In molecular biology, RB is known as an intercalating agent and has been used as a nucleic acid stain. Due to its exceptional structure, it can easily intercalate into the DNA double strand. When RB is exposed to light, it can form reactive oxygen compounds. There are two types of reactions resulting in the formation of reactive oxygen compounds. The first increases the energy of RB being transferred to biomolecules to form reactive oxygen compounds. In a second reaction, the energy is transferred to molecular oxygen to form singlet oxygen. When RB enters the body together with food, it may trigger oxidative stress on ovarian follicles and decreases the number of primary, secondary and de Graaf follicles. The main three-ring part of RB resembles a flat purine-pyrimidine base pair and can slide into DNA between adjacent co-planar base pairs (intercalation). In terms of thermodynamics, the formation constant of the Rhodamine B–DNA complex occurs spontaneously which has negative enthalpy and positive entropy. The negative enthalpy change is due to the binding of RB to the groove of DNA and the positive entropy change is due to release of the bound water from the DNA that is replaced by the binding of RB to DNA.13

In in vitro studies, Oleuropein, the major phenolic secoiridoid in Olea europaea L. (Oleaceae), has been described as a potent antioxidant and in vivo14 studies. Olive leaves or oleuropein have been shown to modulate cardiac, hepatic and metabolic conditions in high-carbohydrate, high-fat-fed rats14 and protect heart function against ischemia and reperfusion.13 Cumaoğlu et al. showed that ethanolic extracts of olive leaves, oleuropein or hydroxytyrosol have protective and regenerative effects on redox balance and related signaling pathways in insulin releasing pancreatic β-cells exposed to hydrogen peroxide or a cytokine cocktail. In addition, Bali et
al.\textsuperscript{18} have studied the ability of olive leaf extracts or polyphenolic compounds (oleuropein, hydroxytyrosol and quer cetin) to protect H\textsubscript{9c} cardiomycocytes \textit{in vitro} against 4-hydroxynonenal stress-induced toxicity and apoptosis. In the present study, it was aimed to examine the oligoprotective (genoprotective) properties of aqueous olive leaf extract (AOLE), which is a cost-effective herbal treatment with well-known benefits, hoping to be used either alone or in combination with other medication such as N-acetylcystein and pirfenidone to protect the DNA from damage by harmful agents.

**Materials and methods**

Genomic DNA was used in a concentration of 75μg/ml. DNA was extracted from peripheral lymphocytes of anticoagulated blood (EDTA) samples by Proteinase K digestion and phenol/chloroform extraction. The purity was determined by measuring the absorbance at 260/280nm indicating that the sample is free from protein contamination.\textsuperscript{19} Calf thymus (CT) DNA was obtained from Sigma-Aldrich (St. Louis, MO, USA). The concentration was assayed spectrophotometrically using 66000 M\textsuperscript{-1} cm\textsuperscript{-1} as a molar extinction coefficient at 260 nm. The results were expressed in terms of base-pair equivalents per dm\textsuperscript{3}.\textsuperscript{20} RB was obtained from Hopkin & Williams Ltd (London, UK) and was used without further purification. All the experiments were conducted in Tris buffer (0.01M Tris, 0.1M NaCl, at pH 7.34). Glass-distilled deionized water and analytical grade reagents were used throughout all the experiments. The pH values of solutions were measured with a calibrated Jenway pH-meter model 3510 (Staffordshire, UK). All buffer solutions were filtered through Millipore filters (Millipore, UK) of 0.45 mm pore diameter.

**Absorbance spectra:** Absorbance spectra were measured on a Jenway UV-visible spectrophotometer, model 6505 (London, UK) using quartz cells of 1.00cm path length. A concentration of 0.24μM for RB was chosen for all absorption and emission measurements. This concentration was high enough to get good fluorescence signal strength in all the samples, and low enough to avoid possible aggregating of molecules, or other molecular effects that might affect the absorption and emission characteristics. In addition, RB has a somewhat a weaker signal which photo bleaches more easily and 0.24μM was found experimentally to be good enough to give a good signal with limited noise.

**Fluorescence spectra:** Fluorescence emission and excitation spectra were measured using Jasco FP-6200 spectrophotometer (Tokyo, Japan) using fluorescence 4-sided quartz cuvettes of 1.00cm path length. The automatic shutter-on function was used to minimise photo bleaching of the sample. System A formation was induced by sequential addition of aliquots of 1790μl Tris buffer, 10μl RB (stock solution of 48μM), 100μl ct-DNA or genomic DNA from stock solutions (1.5mg/ml) and finally 100μl of AOLE (5g/15ml). System B formation was induced by sequential addition of aliquots of 1790μl Tris buffer, 10μl RB (stock solution of 48 μM), 100μl of AOLE, and finally 100μl ct-DNA or genomic DNA from stock solutions (1.5mg/ml). Emission spectra were recorded for each system using excitation wavelengths of maximum fluorescence intensity determined for the systems to be 550 nm using a slit width of 5nm to examine alterations in emission spectra resulting from the complex construction of both systems. On construction of the full systems, the system was allowed to equilibrate for 2-5 minutes at room a temperature and emission spectra (560-650 nm) were recorded to monitor changes in RB intensity.

**Stoichiometric studies of RB and AOLE:** The binding stoichiometry was determined using fluorescence techniques.\textsuperscript{21} Fluorescence emission intensity of RB was monitored at 578nm after exiting at λ\textsubscript{ex} nm at room temperature. The fluorescence intensities were recorded in solutions where the concentration of RB was kept constant (0.24μM)\textsuperscript{11} and the concentration of AOLE varied. Differences in fluorescence intensity at 576nm of RB in the absence and presence of AOLE were plotted against different volumes of AOLE. Data reported are averages of at least three experiments.

**Preparation of aqueous Olea europaea leaf extracts**

Leaves of Olea europaea were collected from the Novellien zone, Tripoli, Libya in November 2017. The leaves (5gm) were cleaned and washed with distilled water and dried at a room temperature of 25°C for about 20 minutes. Dried leaves were grinded in a homogenizer (HO4A Edmund Buhler GmbH, UK) along with 15ml of distilled water, the resulting aqueous solution was then filtered using a Millipore filter (0.45μm GHD Acrodisc GF, UK).

**Determination of the total phenolic content of Olea europaea**

The total phenolic contents of the aqueous Olea europaea was determined using the Folin Ciocalteau reagent.\textsuperscript{22} A calibration curve was plotted by mixing 1 ml aliquots of 25, 50, 100, 150, 225, 250, 290, 350, 390 and 450 mg/ml Gallic acid solutions with 5.0ml of Folin Ciocalteau reagent (diluted tenfold) and 4.0ml of sodium carbonate solution (75g/l). The absorbance was measured after thirty min at 765nm. After one hour, the absorbance was measured to determine the total phenolic contents.

**Free Radical Scavenging Activity**

Free radical scavenging activity of AOLE extracts was measured by 2,2-diphenyl-1-picrylhydrazyl (DPPH) assay. In brief, 0.1mM solution of DPPH in ethanol was prepared. This solution (1ml) was added to 3ml of AOLE in ethanol at different concentration. The mixture was shaken vigorously and allowed to stand at room temp for 30 min. then, absorbance was measured at 517nm by using spectrophotometer (Jenway UV-visible spectrophotometer).\textsuperscript{23} A reference standard compound being used was ascorbic acid and experiment was done in triplicate. The IC50 value of the AOLE, which is the concentration of sample required to inhibit 50% of the DPPH free radical, was calculated.\textsuperscript{0} The percent DPPH scavenging effect was calculated by using following equation: DPPH scavenging effect (%) or Percent inhibition = \( A_s - A_r \times 100 \). Where \( A_s \) was the Absorbance of control reaction and \( A_r \) was the Absorbance in presence of test or standard sample.\textsuperscript{23}

**Melting curve experiments of Olea europaea leaf extracts components with DNA**

\( T_m \) experiments were performed spectrophotometrically (using Peltier-thermostatted cuvette holder in a Cary 4000 UV-visible spectrophotometer) by measuring the change in absorbance at 260nm with temperature. The spectrophotometrically determined melting curves were based on monitoring the change in A260nm with
temperature over a range of 5°C to 110°C at a rate of 5.0°C increment. The melting curve obtained was used determine \( T_m \) values. Melting temperatures \( (T_m) \) was determined by taking the point at half the curve height.\(^{24,25}\) The DNA experiment was performed by heating ctDNA 100µl (75µg/ml) in 0.01M Tris, 0.1 M NaCl at pH 7.34 to 100°C and then the 5µL of rhodamine B (0.24µM) was added and the \( T_m \) was determined. Similarly, the experiment was performed again by heating the DNA 100µl (75µg/ml) to 100°C and then 5µL of rhodamine B (0.24µM) and 100µl AOLE (17mg/ml) of AOLE were added and the \( T_m \) was determined.\(^{26}\)

### Molecular docking

The starting geometry some AOLE contents was constructed using chem3D Ultra (version 8.0, Cambridge soft Com., USA). The optimized geometry of AOLE contents with the lowest energy was used in the molecular docking. Crystal structures of two DNA fragments with the B-form were downloaded from the Protein Data Bank (https://www.rcsb.org/structure/102D). The molecular dockings of AOLE contents with B-DNAs were accomplished using Auto Dock 4.2 software from the Scripps Research Institute (TSRI) (http://autodock.scripps.edu/). Firstly, the polar hydrogen atoms were added into B-DNA molecules. Then, the partial atomic charges of the B-DNA and AOLE contents molecules were calculated using Kollman methods.\(^{27}\) In the process of molecular docking, the grid maps of dimensions (62Å X 62Å X 62Å) with a grid-point spacing of 0.376Å and the grid boxes is centered. The number of genetic algorithm runs and the number of evaluations was set to 100. All other parameters were default settings. Cluster analysis was performed on the results of docking by using a root mean square (RMS) tolerance of 2.0Å, and this was dependent on the binding free energy. Lastly, the dominating configuration of the binding complex of AOLE contents and B-DNA fragments with minimum energy of binding can be determined.

### Results and discussion

#### Fluorometric findings

Similar fluorescence emission data were obtained by either using genomic DNA or ctDNA. In Figure 1, the fluorescence spectrum of RB (0.24µM) in the wavelength range of 560–650nm excited at 550nm in the presence of increasing AOLE concentrations is shown. On addition of AOLE, a hypochromatic effect was observed by titration with no clear isosbestic point. Complex formation between RB and AOLE was confirmed by a decrease in fluorescence intensity with increasing AOLE concentration. This significant fluorescence change suggests a strong association between RB and the AOLE. In addition, data obtained from the fluorometric titrations are presented in inset of Figure 1 clearly demonstrating a stoichiometry titration pattern between RB and AOLE. Safitri et al.\(^{28}\) have shown that RB increases lipid peroxidation and can be used as an oxidative stress marker. Olive leaves contain many polyphenolic compounds such as oleuropein, hydroxytyrosol and flavonoids, including luteolin, rutin, caffeic acid, catechin and apigenin. Oleuropein, together with other closely related compounds such as 10-hydroxyoleuropein, ligstroside and 10-hydroxyligstroside, are tyrosol esters of elenolic acid.

The total phenolic contents of the olive leaves aqueous extract was expressed and presented in terms of mg gallic acid/g extracts. The obtained results showed that the amount of phenolic compounds of the olive leaf was 122.2±1.20mg gallic acid/g. Most of these compounds are powerful antioxidant agents present abundantly in olive leaves and hence can reduce the oxidative stress produced by RB in a ratio of 1:1 as shown by stoichiometry titration. The antioxidant activity of aqueous olive leaf extract was evaluated using the 2,2-diphenyl-1-picrylhydrazyl (DPPH) assay and the IC\(_{50}\) was 65±2.3µg/mL, lower than the value of vitamin C (59.45±µg/mL).\(^{31}\) The reducing ability of the AOLEs serve as an important indicator of its potential antioxidant activity, because its associated with their ability to donate electron to free radical species, reducing them into more stable and un-reactive form. The reducing power of the AOLEs increased with concentration but it was below the reducing power capacity of ascorbic acid. The obtained result is in a good agreement with Ferreira et al.\(^{32}\) who investigated the reducing power of olive leaf extract using the ferricyanide assay. Based on Phan et al.\(^{33}\) studies the chemical structure of polyphenols and RB, the distribution of hydroxyl groups on polyphenols as well as the presence of a carboxyl group in RB can influence interactions through the formation of hydrogen bonds and the partitioning of polyphenols in a non-polar core of RB will facilitate the hydrophobic interaction.\(^{34}\)

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Figure 1 Plot of fluorescence emission of RB (0.24µM) vs wavelength from 562-650nm using excitation \( \lambda_{ex} \) in the presence of different volumes of AOLE (5g/15ml) in 0.01M Tris, 0.1 M NaCl at pH 7.34. Inset: Fluorescence intensity at 576 nm plotted versus different volumes of AOLE (5g/15 ml) in 0.01M Tris, 0.1 M NaCl at pH 7.34 and R\(_0\) was 0.9931. The volumes of AOLE were 1-µl, 2-20µl, 3-40µl, 4-60µl, 5-80µl and 6-100µl, respectively.

Figure 2 shows a decrease in fluorescence intensity of RB (0.24µM) in 0.01 M Tris, 0.1 M NaCl at pH 7.34 on addition of ctDNA (system A). This change in fluorescence intensity of RB in presence of ct-DNA is indicative of a strong association of RB to ct-DNA and suggesting the location of the bound RB in a hydrophobic environment. As described in the literature, RB can more easily bind if their association to double stranded DNA is facilitated by positive charges. DNA attracts counter ions to its surface to screen polyionic charges on the phosphodiester backbone. RB may compete with sodium ions for binding to DNA phosphates which further stabilizes the RB-DNA complex. It is observed that in high salt concentrations the binding of RB to the DNA phosphate backbone decreases and the binding constant decreases moderately.\(^{34}\) In addition, the binding of RB to the minor groove of DNA was further indicated by circular dichroism, fluorescence quenching, and molecular docking studies.\(^{25,34}\) On addition of 100µl of AOLE (stock: 5g/15ml), a further reduction in RB fluorescence intensity was seen due to the interaction between free unbound RB and AOLE as mentioned above. The inset of Figure
2 shows differences in the fluorescence intensities of system A and system B. Fluorescence intensity of the system B is higher than that of the system A because the AOLE did not allow the RB to interact with the ct-DNA and more free (unbound) RB is exposed to the excitation light and then produces more emission fluorescence light.

The antigenotoxic effect obtained in this study is in agreement with many literature studies. Cabarkapa et al. found that the excessive release of stress hormone adrenaline is accompanied by generation of reactive oxygen species which may cause disruption of DNA integrity leading to cancer and age-related disorders. Phenolic-rich plant product dry olive leaf extract was reported to attenuate formation of DNA lesions induced by adrenaline. It has also been found that the antigenotoxic effect of the extract was more pronounced at smaller concentrations and post-treatment with 0.125mg/ml dry olive leaf extract was the most effective against adrenaline genotoxicity. Topalovic et al. have also reported that thyroid hormones change the rate of basal metabolism, modulating the consumption of oxygen and causing production of reactive oxygen species, which leads to the development of oxidative stress and DNA strand breaks. Olive (Olea europaea L.) leaves contain many potentially bioactive compounds, making it one of the most potent natural antioxidants. These findings indicate that L-thyroxine exhibit genotoxic effects and that olive leaf extract was the most effective against adrenaline genotoxicity. It has also been found that the AOLE did not allow the RB to interact with ct-DNA and more free (unbound) RB is exposed to the excitation light and then produces more emission fluorescence light.

The melting temperature ($T_m$) of ctDNA (88.0±0.2°C), which is dissociated into single strands, is strictly related to the stability of the DNA. In addition, the binding of RB to ctDNA has increased the $T_m$ by 4.0±0.06°C due to increase the intercalation ability between the two DNA strands and this consistent what has been noticed by Masun et al. The intercalation of RB into the double helix has increased the helix melting temperature and stabilize the natural structure of DNA, and thus lead to the $T_m$ of DNA increase of about 4.0±0.06°C, while the intercalative binding of AOLE components lead to the $T_m$ of ctDNA increase of about 0.52±0.03°C. It can be seen that the $T_m$ of DNA with RB and AOLE components has a $T_m$ of 88.5°C which is nearly similar to the $T_m$ of ctDNA alone (88.0±0.2°C) and this suggesting that the AOLE components has abolished the RB intercalated ability.

**Molecular docking strategy**

Table 1 shows the energies of binding of the some AOLE contents of with DNAs fragments obtained by using molecular docking strategy. In this study, the molecular dockings of AOLE contents with two B-DNA fragments were performed using Auto Dock 4.2 to understand the binding mode of AOLE contents with B-DNA and to obtain the information about interaction forces between Rhodamine, AOLE contents and DNA. Rhodamine and AOLE contents were kept as flexible molecule and was docked into two types of rigid B-DNA fragments to obtain the preferential binding site AOLE contents on B-DNAs and the molecular docking results were shown in Table 1. It can be found from Table 1 that the more adenine (A) and thymine (T) base pairs in DNA sequence is, the more negative the binding energy (DG) is to some extent, indicating that AOLE contents prefers to bind on the minor groove of A–T rich DNA fragments as shown in Table 1. This is because the narrower and deeper shape of the minor groove can offer several sites of action during the binding process of AOLE contents with DNAs, resulting in the close contact with the surface of AOLE contents through van der Waals forces and hydrogen bonding interaction. Meanwhile, it can be found that the significant change of conformation of AOLE contents be present in the AOLE contents–DNA complexes to orient easily along the minor groove.

The conformation change of AOLE contents is close relation with the structure of minor groove of DNA fragments as shown in Table 1. It is implied that the flexibility of AOLE contents molecule also plays an important role in the binding process of Rhodamine and AOLE contents with DNA. In addition, it can be found that there are van der Waals, hydrogen bonding and electrostatic interactions between Rhodamine, AOLE contents and DNAs and the contribution of van der Waals and hydrogen bonding interaction is much greater than that of the electrostatic interaction because the sum of van der Waals energy, hydrogen bonding energy and desolvation free energy is larger than the electrostatic energy (Table 1), which is consistent with the experimental results. As indicated in the literature, the binding mode of Rhodamine B on DNA is groove binding as the interaction mode. The fluorescence emission intensity of system A (in which the formation of the system was formed by sequential addition of aliquots of Tris buffer, RB, ct-DNA and finally AOLE, Figure 2) did not change on further addition of AOLE and this suggesting that there is no competitive binding interaction between RB and AOLE on ct-DNA. Therefore, it can be concluded that the binding mode of Rhodamine on ct-DNA is major groove binding interaction but the AOLE is mainly minor groove binding interaction.

**Structural features of duplex DNA with Olea europaeae leave extracts**

The melting temperature ($T_m$) of ctDNA (88.0±0.2°C), which is defined as the temperature at which half of the double-stranded DNA is dissociated into single strands, is strictly related to the stability of the DNA. In addition, the binding of RB to ctDNA has increased the $T_m$ by 4.0±0.06°C due to increase the intercalation ability between the two DNA strands and this consistent what has been noticed by Masun et al. The intercalation of RB into the double helix has increased the helix melting temperature and stabilize the natural structure of DNA, and thus lead to the $T_m$ of DNA increase of about 4.0±0.06°C, while the intercalative binding of AOLE components lead to the $T_m$ of ctDNA increase of about 0.52±0.03°C. It can be seen that the $T_m$ of DNA with RB and AOLE components has a $T_m$ of 88.5°C which is nearly similar to the $T_m$ of ctDNA alone (88.0±0.2°C) and this suggesting that the AOLE components has abolished the RB intercalated ability.

**Molecular docking study**

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**Figure 2** Plot of fluorescence emissions from 562-650nm vs wavelength using excitation λ550 nm of 1- 10µl RB (0.24µM) and 1990µl 0.01M Tris, 0.1M NaCl at pH 7.34. 2- System B formation was induced by sequential addition of aliquots of Tris buffer, RB, AOLE and finally ct-DNA. 3- System A formation was induced by sequential addition of aliquots of Tris buffer, RB, ct-DNA and finally AOLE.

**Table 1**

| AOLE Contents | RNA (%) | DNA (%) |
|---------------|---------|---------|
| AOLE 1        | 95.84   | 50.23   |
| AOLE 2        | 96.52   | 51.43   |

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Table 1 Various energies in the binding process of the some AOLE contents of with DNAs obtained from molecular docking. The unit of all energies was kJ/mol

| Ligand          | DNA PBD ID(102d) d(CGCAAAATTTGGCG) | DNA PBD ID(1bna) d(CGCGAASSCGCG) | DNA PBD ID(102d) d(CGCAAAATTTGGCG) | DNA PBD ID(1bna) d(CGCGAASSCGCG) |
|-----------------|-----------------------------------|---------------------------------|-----------------------------------|---------------------------------|
|                 | Binding energy ΔG* Binding site  | Binding energy ΔG* Binding site | Binding energy ΔG* Binding site  | Binding energy ΔG* Binding site |
| Rhodamine       | -3.37 Major G                     | -3.46 Major G                  | Nuzhenide                         | -3.74 Major G                  |
| Catechin        | -7.85 Minor G                     | -7.64 Minor G                  | Hydroxytyros                      | -4.46 Minor G                  |
| 5-Caffeoylquinic| -5.05 Minor G                     | -4.46 Minor G                  | Homovanillic acid                 | -3.84 Minor G                  |
| acid            |                                   |                                |                                   | Minor G                         |
| Cynadin-3       | -7.21 Minor G                     | -7.51 Minor G                  | Vanillic acid                     | -2.89 Minor G                  |
| glucoside       |                                   |                                |                                   | Minor G                         |
| Dimethyloleuropein | -4.69 Minor G                     | -2.91 Minor G                  | Tyrosol                           | -4.68 Minor G                  |
| 3,4-DHPEA-EDA   | -3.79 Minor G                     | -1.87 Minor G                  | Rutin                             | -7.23 Minor G                  |
| Hesperidin      | -7.98 Minor G                     | -7.73 Minor G                  | Oleurosides                       | -5.07 Minor G                  |
| Luteolin        | -8.69 Minor G                     | -7.74 Minor G                  | Ligrostoside                      | -5.36 Minor G                  |
| Quercetin       | -8.02 Minor G                     | -7.54 Minor G                  | Apigenin                          | -8.16 Minor G                  |
| Verbascoside    | -5.09 Minor G                     | -2.62 Minor G                  | Caffeic acid                      | -4.02 Minor G                  |

ΔG is the binding free energy change in the binding process.

Conclusion

The present study created convincing evidence that AOLE has a protective effect against chemically induced genotoxicity and oxidative damage. The genoprotective activity could be exhibited through an increase in antioxidant activity, scavenging of free radicals and inhibition of oxidative stress. These findings indicate genoprotective and antioxidant properties in olive leaf extract, strongly supporting further explorations of its underlying mechanisms of action.

Acknowledgement

None.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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