The Antioxidant Flavonoids, Orientin and Vicenin Enhance Repair of Radiation-Induced Damage

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Abstract
Orientin (Ot) and vicenin (Vc), two water soluble flavonoids isolated from the Indian Holy Basil, are potent radioprotectants, but the mechanisms of protection are not fully understood. In the present study, we investigated the effects of these flavonoids as antioxidants in chemical systems and plasmid DNA and radioprotectants in bacterial and mammalian cells. The radical scavenging activities of Ot and Vc were assessed in two chemical systems, diphenylpicrylhydrazyl (DPPH) and 2,2´-azino-bis(3-thylbenz-thiazoline-6-sulfonic acid) (ABTS) by estimating absorbance at 517 and 734 nm, respectively. Protection by the flavonoids against H2O2 (30 mM) damage to plasmid pGEM-7Zf was assessed by DNA electrophoresis. Effect of flavonoid pre-treatment on survival of irradiated (30-150 Gy γ-rays) repair proficient (AB1157) and repair deficient (MD2130) E.coli cells was assayed. Furthermore, induction and repair of DNA damage to 4 Gy irradiated mouse splenocytes was analyzed using the comet assay. Ot and Vc significantly scavenged DPPH and ABTS. Both flavonoids protected plasmid pGEM-7Zf in the nicked circular form following H2O2 treatment. Pretreatment with 200 µM of either flavonoid significantly increased clonogenic survival of repair proficient cells at all radiation (RT) doses; in repair deficient strain, protection was significant at the lower RT doses (30 and 50 Gy, p<0.01). Ot and Vc significantly reduced 4 Gy-induced DNA damage and allowed for faster repair of ex-vivo irradiated mouse splenocytes.

These studies underscore the potent antioxidant activity of orientin and vicenin; in addition, both flavonoids appear to facilitate repair of radiation-induced injury.

Keywords: Orientin; Vicenin; Radical scavenging; DNA protection; Bacterial survival; Comet assay

Introduction
Irradiating radiations cause damage to the DNA, lipids and proteins, resulting in perturbation of cell signaling pathways, cell-cycle disruption, apoptosis, mitotic death, or necrosis. The role of radioprotective compounds in mitigating the structural and functional damage to normal tissues in the event of planned exposures or radiation accidents has been studied extensively for more than half a century [1-3]. To date, there are no FDA-approved countermeasures to prevent or treat acute radiation syndrome.

Ocimum sanctum (also known as the Indian Holy Basil) is used by medicinal practitioners for its antidiabetic, antifungal, antimicrobial, analgesic, and anticancer potential [4]. Orientin (Ot) and vicenin (Vc) are water soluble c-glycoside flavonoids isolated from the aqueous leaf extract of Ocimum sanctum. Ot and Vc have demonstrated potent radioprotective activity in vivo at low non-toxic doses with dose reducing factors of 1.3 and 1.37, respectively [5] and excellent hydroxyl radical scavenging activity in vitro [5,6]. Further, we have reported that treatment of pregnant mice with 50 µg/kg body weight of either Ot or Vc prior to irradiation significantly reduced radiation-induced chromosomal anomalies in fetal and adult hematopoietic cells, and restored blood indices to normal range in the off-spring; while radiation-induced solid tumor incidence was significantly reduced [7]. On the basis of the authors’ own studies and other publications, several mechanisms such as radical scavenging, metal chelation,
An ideal radioprotector should be effective against acute- and long-term radiation effects, non-toxic, affordable, and be chemically stable to permit ease of handling and storage. Our goal was to use a battery of different assays that complemented each other to further understand the role of Ot and Vc in protecting against chemical- and radiation-induced damage. Free radical scavenging was confirmed by the DPPH and ABTS assays and in plasmid DNA; DNA strand break and clastogenicity was studied in DNA plasmid and bacterial cells. The efficacy of Ot and Vc in repairing radiation-induced DNA strand breaks in mouse splenocytes was measured by the comet assay.

Materials & Methods

Materials

Materials were purchased from the following sources: Eagles minimum essential medium, fetal calf serum, low melting point and normal agarose, DPPH and ABTS from Sigma-Aldrich, St Louis, MO, USA. Luria broth, Luria agar and Hank’s balanced salt solution from Himedia, Mumbai, India.

Plasmid pGEM-7Zf and E. coli were provided by Dr. Sathyamoorthy, Department of Cell and Molecular Biology, Kasturba Medical College, Manipal, India.

The bacterial strains AB1157 (repair proficient) and MD2130 (repair deficient) were kindly provided the Radiation Biology Division, Bhabha Atomic Research Center (BARC, Mumbai, India).

Isolation of Ocimum flavonoids

Fresh leaves of the dark variety of Ocimum sanctum (Sanskrit: Tulasi, English: Indian Holy Basil) were collected locally, washed and shade-dried. An aqueous extract was prepared by refluxing the powder with double-distilled water at 80°C and condensed in a vacuum drier (Speedvac SC 110 A, MN, USA). The active components were isolated from the extract by chromatographic separation and their identity confirmed using Nuclear Magnetic Resonance (NMR) spectroscopy [5].

Preparation of Drug Solution

Both Ot (Mol. wt. 448) and Vc (Mol. Wt. 594) are stable at room temperature. The drug solutions were freshly prepared by dissolving appropriate quantities of the two compounds in double-distilled water (DDW) under sterile conditions shortly before use.

Animals

Healthy, 8-12 weeks old BALB/c male mice, weighing ~29 g, from the inbred colony at the Institute of Nuclear Medicine and Allied Sciences (INMAS), Delhi, India were used. The mouse splenocytes were collected as described under methods [13].

Irradiation

Bacteria were exposed on ice in sterile 2 mL stoppered tubes (Nunc, Germany) in a field size of 10 x 20 cm² at a source to sample distance of 60 cm and dose rate of 3.6 Gy/min using a cobalt-60 teletherapy unit (Siemens, Germany) in the Department of Radiotherapy and Oncology, Shirdi Sai Baba Cancer Hospital and Research Center, Manipal.

Mouse splenocytes were exposed to gamma rays on ice in sterile Petri dishes (Nunc, Germany) using a 60Co teletherapy unit (Eldorado, Theratron 780C, AECL, Canada) at a source to sample distance of 80 cm. The dose rate was 1.4 Gy/min. All animal studies were conducted according to the institutional regulations and national criteria for animal experimentation.

Experimental Procedures

The following experiments were conducted

Radical scavenging activity in chemical systems: These following two chemical tests were conducted to validate the antioxidant activity of the flavonoids isolated from the Indian Holy Basil.

Scavenging of 1,1-diphenyl-2-picrylhydrazyl (DPPH): DPPH, a stable free radical characterized by a deep violet color, is reduced to a pale yellow to colorless form, dipicrylhydrazine, in the presence of antioxidants [14], with a concomitant decrease in absorbance at 517 nm. DPPH (96 µM) in DMSO was mixed well with Ot or Vc at different concentrations (0-250 µM) to give a final volume of 1.0 mL and the reaction mixture was placed in the dark at room temperature for 20 min. Absorbance was recorded at 517 nm. The antiradical activity is measured by the percentage of inactive DPPH:
Inactive percentage = [(Ab-Aa)/Ab] X 100; where Ab is the absorption of a blank specimen at t = 0 min, Aa is the absorption of Ot or Vc in the reaction mixture.

Scavenging of 2,2'-azino-bis(3-ethylbenzthiazoline-6-sulfonic acid) cation (ABTS): The ABTS assay is based on the oxidation of ABTS to an intensely-colored nitrogen-centered radical cation, ABTS⁺ by antioxidants, which has a characteristic long-wavelength absorption maxima at 734 nm [15].

The reaction mixture contained ABTS radical [ABTS (2 mM), potassium persulphate (0.17 mM)], various concentrations of Ot or Vc (0–80 µM) in PBS buffer (pH 7.4, 20 mM) in a total volume of 3.5 mL. Absorbance was recorded at 734 nm.

The antiradical activity was calculated by applying the same formula as for the DPPH assay. Both assays were repeated and percent inhibition of DPPH and ABTS calculated. The data presented are mean±SEM (standard error of mean) of three separate experiments. The IC₅₀ values for both assays (the concentration of flavonoids required to inhibit either DPPH or ABTS activity by 50%) were calculated from the concentration-inhibition linear regression curves [14].

DNA nicking assay for radical scavenging activity: All experiments were conducted in potassium phosphate buffer (pH 7.4, 50 mM). Two µL each of ethylenediaminetetraacetic acid (EDTA), phosphate buffer, H₂O₂ (30 mM), FeSO₄ (16 mM), Ot or Vc (250 µM) and plasmid pGEM-7Zf DNA (0.1 µg/µL) were mixed in a 500 µL tube. The final volume of the reaction mixture was brought to 12 µL with deionized DDW and incubated for 1 h at 37°C. Following incubation, 2 µL loading dye (6XESB-20XTEA buffer, glycerol, bromophenol blue, xylene cyanole and water) was added and 12 µL of this mixture was loaded onto a 0.7% agarose gel. Electrophoresis was conducted at 60 volts in Tris, boric acid, EDTA (TBE) buffer for 2 h. The gel was stained with ethidium bromide (0.5 µg/mL in DDW) for 20 min and DNA bands visualized under UV light and photographed using a gel documentation system (Amersham Biosciences, GE Healthcare, UK).

Cell survival (macrocolony) assay in E. coli strains

Selection of optimum drug dose: Ot or Vc at concentrations of 0, 50, 100, 150, 200, or 250 µM of were added to 10⁶ AB 1157 bacterial cells (optical density at 600 nm: 0.4-0.45) in PBS and incubated for 30 min at 37°C. The cells were irradiated on ice with 100 Gy gamma rays, mixed thoroughly, and serially diluted. Dilutions of 10⁶ and 10⁵ cells were plated on Luria agar in triplicate, incubated at 37°C for 24 h and colonies manually counted under a bacterial colony counter with 1.5 x magnification. Based on the data from this experiment and an earlier study [10], a dose of 200 µM of Ot and Vc was used for studying the modification of radiation dose-response in both AB1157 and MD2130 strains.

Radio-modifying effects of Ot and Vc in AB1157 and MD2130 bacteria: AB1157 and MD2130 E.coli were incubated with DDW, or 200 µM of Ot or Vc for 30 min at 37°C prior to irradiation with 0, 30, 50, 100, or 150 Gy γ-rays on ice. The cells were then serially diluted; dilutions of 10⁴ and 10³ were plated in triplicate in Luria agar plates. The cells were allowed to grow for 24 h at 37°C and the visible colonies counted as above.

Surviving fraction (SF) was calculated as the ratio of the average number of colonies in the irradiated group to the average number of cells in the sham-treated control. The dose reduction factor (DRF) was calculated as the ratio of radiation dose resulting in 50% survival of the vehicle group compared to the radiation dose resulting in 50% survival in the Ot or Vc treated groups. The data are expressed as mean±SEM obtained from three independent experiments.

DNA damage and repair in mouse splenocytes (comet assay)

BALB/c mouse was killed by cervical dislocation and spleen flushed through a nylon gauze (mesh size 45 µm) under sterile conditions to obtain a debris free, single cell suspension. The splenocytes (1x10⁹ cells) were incubated with DDW or 17.5 µM (based on a previous work [9]) Ot or Vc for 30 min before exposure on ice to 4 Gy. Approximately 3x10⁵ cells were withdrawn at 0, 15, 30, 60, 180, and 360 mins after irradiation and each aliquot mixed with 0.75% ultralow gelling agarose (BDH Electran; gelling temperature 17°C) on microscopic slides pre-coated with 0.1% agarose. The slides were cooled on a cooling plate and immersed in pre-cooled neutral lysis buffer (2.5% sodium dodecyl sulphate [SDS], 1% sodium sarcosine, 25 mM EDTA, pH 9.5) for 15 min at room temperature and electrophoresed in TBE buffer [90 mM Tris, 90 mM boric acid and 2 mM EDTA; pH 8.3] at 2 V/cm for 5 min at 10°C in a horizontal electrophoresis unit (Multiphor-II, Pharmacia Biotech, Uppsala, Sweden). Slides were air dried at 45°C and stored in a refrigerated humid chamber. Comets were stained with 25 µM propidium iodide and examined with a fluorescent microscope (BX60, Olympus) coupled with a CCD camera. Images were acquired and analyzed using a standard Comet assay program (Optimas, UK). Fifty comets per group were analyzed after rehydrating the slides. Tail length, tail movement, percentage of DNA in tail and head of the comet were parameters acquired using the Optimas image analysis software [16]. The percentage of total DNA in comet tail is used as a measure of DNA damage. To quantify the efficiency of Ot and Vc in repairing radiation-induced strand breaks, the cellular DNA repair index (CRI) was applied. CRI for percent tail DNA is defined as the decrease from initial values of percent tail DNA. This relation is utilized to monitor the rate of decrease in DNA damage parameter in irradiated cells under different treatment conditions [17].

CRI = [1-(percent tail DNA at time 't'/percent tail DNA at initial time t₀)]x100

Data from three separate experiments are expressed as mean±SEM.
Statistical analysis

The data for free radical scavenging activity, survival of *E. coli* and repair kinetics in mouse splenocytes were analyzed by one-way ANOVA and Student's t-test. Each experiment was repeated in triplicate. A value of *p*<0.05 was considered significant. Homogeneity of variance was tested. Radiation dose–response curves were fitted with linear (*Y* = *C* + α*D*) and linear-quadratic (*Y* = *C* + α*D* + α*D^2*) models using Microsoft excel, version 2007.

Results

Radical scavenging in chemical systems

Scavenging of DPPH: In the concentration range used in the present studies (62.5 to 250 µM) both Ot and Vc decreased the absorbance of DPPH at 520 nm (Figure 1A). At all concentrations used, radical scavenging activity of Ot was higher than that of Vc, although the difference was statistically significant only at 125 µM (*p*<0.001). The IC_{50} values for DPPH inhibition by Ot and Vc were 115.48 µM and 196.44 µM, respectively (Table 1).

Scavenging of ABTS: There was a steep decrease in the absorbance of ABTS at 734 nm induced by Ot and Vc at concentrations of 10-80 µM (Figure 1B). Radical scavenging by Ot was significantly higher than that by Vc at all equimolar concentrations (*p*<0.001) and was associated with an IC_{50} of 5.49 µM, compared to 16.47 µM for Vc (Table 1).

<table>
<thead>
<tr>
<th>Assay</th>
<th>Orientin IC_{50}±SE (µM)</th>
<th>Vicenin IC_{50}±SE (µM)</th>
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<tbody>
<tr>
<td>DPPH Assay</td>
<td>115.48±5.2</td>
<td>196.44±7.5</td>
</tr>
<tr>
<td>ABTS assay</td>
<td>5.49±0.5</td>
<td>14.55±1.2</td>
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Table 1: Values are mean±SEM of 3 separate experiments.

*p*<0.001 compared to Vc.

Hydroxyl radical scavenging by the DNA nicking assay

DNA is prone to oxidation in the presence of reactive oxidative species and will undergo base oxidation and strand scissions in the presence of the radicals generated by Fenton reaction. The antioxidant assay used here is based on the principle that the undamaged supercoiled DNA, which is more compact, will move faster in the gel compared to the plasmid DNA that is attacked by the free radicals and becomes less compact, linear, and which moves slower in the electrophoretic gel. Representative electrophorogram for the plasmid DNA nicking assay is shown in Figure 2, where the migration profile of plasmid DNA reflects the degree of breakage from supercoiled to linear form. The radical species produced by Fenton process disintegrated the native DNA leaving a faint band of Form II DNA (lane 2). Ot or Vc alone at either of the doses used did not have any adverse effect on the plasmid pGEM-7Zf (lanes 3, 4). Neither flavonoid protected the plasmid in the supercoiled form (Form I), but both Ot (lanes 5) and Vc (lanes 6) protected DNA in the nicked circular form (Form II). These results demonstrate the protective effect of the flavonoids against oxidative damage.

Cell survival (macrolony) assay in *E. coli* strains

a. The preliminary experiment to select the optimum drug dose was performed in the AB1157 strain, exposed to a single radiation dose of 100 Gy. Pretreatment with Ot or Vc increased the survival of 100 Gy γ-irradiated AB1157 cells in a drug dose-dependent manner. Survival increased with dose (*r^2*=0.98) up to 200 µM with Ot and Vc. Further increase in drug dose to 250 µM did not increase protection (Figure 3). There was no significant difference in clonogenic cell survival with either Ot or Vc.
Figure 2: Modulation by Ot or Vc of non-site-specific DNA strand cleavage of plasmid pGEM-7Zf treated with 30 mM hydrogen peroxide. (A) Form I represents the native supercoiled form of the plasmid, Form II is the nicked circular form.

Lane 1: Original DNA
Lane 2: DNA+EDTA+H$_2$O$_2$+FeSO$_4$+Phosphate buffer
Lane 3: DNA+ 250 μM Ot
Lane 4: DNA + 250 μM Vc
Lane 5: DNA+EDTA+H$_2$O$_2$+FeSO$_4$+250 μM Ot
Lane 6: DNA+EDTA+H$_2$O$_2$+FeSO$_4$+250 μVc

Figure 3: Survival of AB1157 E. coli pretreated with 0–250 μM Ot or Vc before exposure to 100 Gy γ-rays. The surviving fraction is plotted as a function of drug dose. All values represent mean values of surviving fractions ±SEM from 3 independent experiments.

b. Ot and Vc (200 μM) did not alter the number of colonies in the sham-exposed group indicating that they were non-toxic to the bacterial strain at the applied dose. Irradiation induced a linear dose-dependent decrease in the survival of AB1157 cells resulting in an LD$_{50}$ of ~40 Gy ($r^2=0.98$; Figure 4A). Pretreatment with Ot or Vc significantly increased survival at all doses of radiation (p<0.01-0.001). The survival benefit by the two flavonoids was nearly identical with LD50 of ~85 Gy, resulting in a dose reduction factor (DRF) of 2.12 (95% CI=2.10-2.14).

c. Ot and Vc by themselves had no impact on the number of colonies in the sham-irradiated repair deficient cells. Irradiation decreased the survival of MD2130 cells at all doses and the dose response was linear-quadratic ($r^2=0.99$), with survival reducing steeply as the RT dose increased from 30 to 100 Gy. Pretreatment with Ot or Vc significantly (p<0.05, Figure 4B) increased the survival at the lowest RT doses (30 and 50 Gy). At higher RT doses (100 and 150 Gy) however, increase in survival was not statistically significant (p=0.4-0.8). The LD$_{50}$ for the repair deficient strain at ~22 Gy, was much lower than LD50 for the AB1157 strain. The LD50 increased to 39 and 42 Gy for Ot and Vc, respectively, resulting in DRFs of 1.77 (CI=1.75 and 1.79 for Ot) and 1.9 (CI=1.88-1.92 for Vc).
Figure 4: Effect of pretreatment with 200 µM Ot or Vc on clonogenic survival of γ-irradiated bacterial cells. (A) Survival of AB1157 E. coli cells. Curve fitting for radiation alone: polynomial ($R^2$ = 0.98); Ot+RT and Vc+RT: exponential ($R^2$=0.99). (B) Survival of MD2130 E. coli cells. All survival data were fitted using exponential function; RT alone ($R^2$=0.98), Ot+RT ($R^2$=0.98) and Vc+RT ($R^2$=0.95). All values represent mean values of surviving fractions ±SEM of 3 independent experiments normalized to control values and plotted as a function of radiation dose. *$p$<0.05, **$p$<0.01 compared to the irradiated group.

DNA strand break and repair analysis using neutral comet assay

Exposure of mouse splenocytes to gamma irradiation ex vivo resulted in cellular damage evidenced by increase in tail length, percent DNA in tail, and tail movement of comets. Here, we focus on the percentage of DNA in the comet tail to express damage and subsequent repair of radiation-induced strand breaks. Percent DNA in the comet tail at time 0 (in relation to time of exposure) was used to access induction of initial DNA damage following 4 Gy. The highest increase in percent tail DNA was observed in splenocytes irradiated with 4 Gy immediately after exposure (0 min) (25.37, 95% CI:21.1-30.3) (Figure 5A). Pretreatment with either Ot or Vc reduced percent DNA in comet tail (16.73, 95% CI: 12.7-19.4 for Ot and 18.8, CI: 15.8-20.2 for Vc) (Figure 5A), and was significant for Ot ($p$<0.01) and Vc ($p$<0.05) compared to 4 Gy-treated splenocytes alone. Although the % tail DNA for Ot-treated cells was lower than the Vc-treated group, the difference was not significant.
The rates of DNA repair was quantified in the irradiated comets as functions of time post-irradiation. The percentage of DNA in the tail of 4 Gy irradiated comets declined at 15 and 30 min compared to the initial damage, yet remained significantly higher than untreated controls even 180 min post-irradiation, and only returned to baseline values 360 min after exposure. Ot- or Vc-pretreated splenocytes demonstrated an earlier decrease in percent tail DNA, reaching baseline values within 60 min after exposure to 4 Gy. These results indicate faster repair in the flavonoid-treated comets compared to slower rejoining of strand breaks in the irradiation alone group.

In order to compare the kinetics of DNA repair in the control and flavonoid treated groups, we estimated the Cellular DNA Repair Index (CRI). The CRI is a measure of reduced strand breaks and therefore, DNA repair. Our analyses of CRI are indicative of accelerated repair in the Ot and Vc-treated splenocytes. CRI for 4 Gy, Ot+4 Gy and Vc+4 Gy groups were determined at 15, 30, 60, 180, and 360 min after exposure to gamma rays (Fig 5B). Vc-treated comets demonstrated the highest CRI, followed by Ot-treated comets, while 4 Gy - irradiated controls had the lowest CRI.

Discussion

Ionizing radiation causes ionization and excitation of water, leading to formation of water radiolysis products such as hydrated electron (e\textsuperscript{-}), ionized water (H\textsubscript{2}O\textsuperscript{+}), hydroperoxyl radical (HO\textsubscript{2}•), hydroxyl radical (•OH), hydrogen radical (H•), and hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}) in a very short period of time (~10\textsuperscript{-8} s) in a biological system [18]. These highly reactive free radicals interact with biomolecules resulting in disruption of signal transduction pathways, DNA damage, decreased proliferation and mortality. Flavonoids can prevent injury caused by hydroxyl and other free radicals by directly scavenging the radicals resulting in a stable, inactive form [19]. Data from the present study confirm the antioxidant property of Ot and Vc, and indicate that both flavonoids may have a role in repair of DNA damage.

Two different radical species, DPPH and ABTS, were selected to examine free radical scavenging abilities of the Ocimum flavonoids, Ot (8-C-b-D-glucopyranosyl-luteolin) and Vc (6-C-b-D-xylopyranosyl-8-C-b-D-glucopyranosyl-apigenin). DPPH is a stable free radical with a maximum absorption at 517 nm, which has frequently been used to estimate the antiradical activity of antioxidants [20-22]. The free radical character of DPPH is neutralized in presence of molecules having ability to donate a hydrogen atom. The scavenging of DPPH by Ot and Vc indicate that both flavonoids act by donating hydrogen atom. Similarly, ABTS can also be neutralized by either hydrogen or electron donation. Furthermore, ABTS radicals are more reactive than DPPH radicals; the reactions with ABTS radicals generally take place in less than a millisecond [20]. Both Ot and Vc reduced absorbance of ABTS. The fast reactivity implies that the flavonoids will function better if present at critical sites at the time of radical formation. The most successful radioprotectors studied- aminothiol compounds- are also effective if present in the system at the time of exposure [23], but not when administered after irradiation.

In both DPPH and ABTS assays, Ot demonstrated superior radical scavenging activity to Vc. Protection against hydroxyl radical-induced plasmid DNA damage by Ot was similarly higher than Vc (discussed next). Luteolin, the parent structure of Ot, is associated with higher antioxidant activity than apigenin, the parent molecule of Vc [24]. Apigenin, with three hydroxyl groups at 5, 7 and 4´ positions, was associated with a small antioxidant activity, while luteolin, with an additional hydroxyl group at 3´ position, was significantly more protective against oxidative DNA damage in lymphocytes than apigenin [25]. We hypothesize that the presence of a similar hydroxyl group in Ot explains the higher radical scavenging activity observed.
Protection of plasmid DNA (pGEM-7Zf) in the nicked circular form by both flavonoids appears to be a direct consequence of radical scavenging activity. Under the reaction conditions, hydroxyl radicals generated by Fenton reaction attack DNA guanosine residues, resulting in strand breakage and transformation from native super-coiled form to relaxed circular or linear forms and complete degradation of DNA [26]. Neither Ot nor Vc protected DNA in the super-coiled native form, while both were effective in protecting circular, nicked DNA. This observation is in keeping with direct scavenging action by these flavonoids reported earlier in a chemical system [6,9] and in E. coli [10]. Collectively, DPPH and ABTS radical scavenging assays, and plasmid protection demonstrate that orientin and vicenin are good antioxidants.

In addition, our results on survival of irradiated repair proficient and repair deficient E. coli and comet formation in irradiated murine splenocytes indicate that Ot and Vc may also facilitate repair of DNA damage.

Both repair proficient (AB1157) and repair deficient (MD2130) strains demonstrated a radiation dose-dependent increase in cellular mortality. However, MD2310 cells were more sensitive to the clastogenic effects of radiation as evidenced by a steep reduction in surviving fractions at RT doses of 50 Gy and above, while AB1157 cells showed higher survival at the same radiation doses. Enhanced radiosensitivity of repair deficient strain can be attributed to decreased ability of injured cells to rejoin DNA strand breaks. Pretreatment with Ot or Vc significantly increased survival in both strains; the effect was more pronounced in AB1157 cells (Figure 4A). Increase in survival fraction of MD2310 E. coli by the flavonoids at lower radiation doses (30 and 50 Gy; Figure 4B) demonstrated that Ot and Vc are effective in protecting bacterial cells against induction of initial lesions, which, if unrepaired, could result in cell death. Neither Ot nor Vc protected MD2310 cells at higher RT doses (100 Gy and above). If these flavonoids functioned only via antioxidant mechanism, there would be no difference in the survival trend for either bacterial strain. In the absence of a competent repair system, the flavonoids could not protect repair deficient bacteria to the same extent as the repair proficient strain. It was reported that radioprotection by cysteamine, an aminothiol, on E. coli cells was almost absent if the cells had genetic defects in the repair systems [27]. The authors inferred that radioprotection by radical scavenging activity for cysteamine was invalid and that specific repair mechanisms may be involved. In contrast, using wild type and radiosensitive (lacking repair pathway) strains of yeast, the radioprotective activity of caffeine was attributed solely to reduction of DNA damage, rather than by interference with repair processes [28]. Here, radioprotection by the ocimum flavonoids appear to be partly by radical scavenging to reduce initial damage and partly by protection/stimulation of inherent repair processes in cells containing active repair machinery.

The dual role of Ot and Vc in reducing initial damage and enhancing repair processes is further established by DNA damage and repair analysis of mouse splenocytes using the comet assay. The comet assay provides a highly sensitive indicator of DNA strand breaks in individual mammalian cells [29]. The presence of Ot and Vc during exposure of the splenocytes reduced DNA damage and facilitated faster repair. Comet profile at later post-irradiation times as well as CRI indicate that Vc and Ot were more effective in repairing RT-induced damage, while both flavonoids increased repair capacity of the splenocytes compared to 4 Gy-irradiated group.

Conclusions

The current results demonstrate the efficacy of Ot and Vc as efficient free radical scavengers and modulators of radiation injury. In addition to an antioxidant activity, enhancement of DNA repair by these flavonoids is also indicated. These findings highlight the potential for development of Ot and Vc as radiation protectants.

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