Antioxidant and DNA damage protecting activities of *Eulophia nuda* Lindl.

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**A B S T R A C T**

**Introduction:** In recent years, natural antioxidants have seen an unprecedented importance and demand in bio pharmaceuticals, nutraceuticals besides their use as food additives. Antioxidants act as potential prophylactic and therapeutic agents against various diseases caused by free radicals. Plants offer tremendous source of antioxidants and are therefore being evaluated for their potentials. *Eulophia nuda* is an important medicinal plant used by local healers in India; however its antioxidant properties have not yet been investigated.

**Methods:** Aqueous (AqE), methanol (ME), aqueous–methanol (AqME) and acetone (AE) extracts of shade dried tubers were obtained and were concentrated in vacuo. Total phenols, flavonoids, ascorbic acid and carotenoids were estimated from all extracts using standard methods. Antioxidant activities of extracts were determined by total antioxidant activity, FRAP, ABTS, DPPH, and OH radical scavenging assays besides lipid peroxidation inhibition. Extracts were evaluated for protection of Fenton’s reagent induced DNA damage.

**Results:** The results confirmed the plant as a rich source of phenols, flavonoids, vitamin C and carotenoids. Among four extracts, AqME showed highest antioxidant activities as evidenced by maximum scavenging of ABTS (98%), DPPH (87%), and OH radicals (99%) at 1 mg ml$^{-1}$ concentration and showed maximal inhibition of lipid peroxidation. All extracts protected the DNA from hydroxyl-radical-induced damage. Again, AqME was proved to be best in providing protection to DNA against damage caused by free radicals.

**Conclusion:** The results provide scientific basis for its traditional usage as natural antioxidant and phytotherapeutic agent. The plant possesses high amount of phenolic compounds and showed a broad-spectrum antioxidant properties including DNA protection.

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1. Introduction

Oxidative stress, induced by the generation of reactive oxygen species (ROS) consisting of both free radical compounds such as superoxide anions ($O_2^-$), hydroxyl radicals ($OH$), as well as non-free-radical compounds hydrogen peroxide ($H_2O_2$), organic peroxide (ROOH), ozone ($O_3$) and singlet oxygen ($O_2^*$) are considered as major causative factors of many of today’s diseases including diabetes and cardiovascular diseases. These ROS are known as exacerbating factors in DNA damage and mutations, cellular injury, oncogenesis (as many mutagens and carcinogens acts through the ROS) and ultimately the aging process. Further, ROS are considered to cause cancer and several neurodegenerative diseases such as Alzheimer’s disease, Parkinson’s disease, Down’s syndrome, inflammation, viral infection and various other digestive disorders including ulcer and gastrointestinal disorders.

Antioxidants are considered to play an imperative role in providing protection against ROS-driven oxidative damage and associated lipid peroxidation, and DNA strand breaking. Even though many antioxidants of synthetic origin are available and used quite frequently, especially in food industry for preservation and prolonging the shelf-lives of food products, however, they are often being associated with quality deterioration, nutritional losses and off-flavor development. Furthermore, available synthetic antioxidants have been reported to exhibit toxic and mutagenic effects. On the other hand, their natural counterparts have an edge over them for being less- or non-toxic and hence can serve as potential drug and dietary molecules. Therefore, in recent years the

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antioxidants of phyto-origin have seen an unprecedented demand in bio-pharmaceuticals, nutraceuticals besides their use as food additives.

Medicinal plants offer an excellent source of various phytochemicals such as phenols, flavonoids, vitamins, tannins, many of which have potent antioxidant activities and therefore can be exploited in drug discovery programs as well as in the preparation of foods and pharmaceutical products.\textsuperscript{10}

*Eulophia nuda* Lindl. (*Orchidaceae*), a medicinally important perennial orchid with underground tubers, is found in central and Southeast Asian regions. In India, this plant is found in tropical Himalayas, from Nepal to Assam, and in Deccan from Konkan southwards. The tubers are reported to be used against tumors, scrofulous glands of the neck, bronchitis, blood diseases and as vermifuge.\textsuperscript{12–15} Raw tubers are eaten for curing rheumatoid arthritis.\textsuperscript{16} Earlier our group has reported anti-proliferative activities of a phenanthrene derivative compound 9,10-dihydro-2,5-dimethoxyphenanthrene-1,7-diol isolated from this plant against human cancer cells.\textsuperscript{17} The tubers are eaten raw and therefore make it a perfect candidate for its exploration as potential antioxidative source.

However, in spite of the fact that various medicinal values have been described by folklore and is extensively used by local healers in different parts of India, till date no scientific validation has been evidenced for antioxidant potential of this plant. This is the first report dealing with the phytochemical analyses, antioxidant potentials and oxidative DNA damage preventive activities of various organic and aqueous extracts of *E. nuda* tubers.

2. Materials and methods

2.1. Plant material

Fresh tubers of *E. nuda* Lindl. (synonym: *Eulophia spectabilis*) were collected from Belgaum region (N 15° 41.099', E 74° 25.026', elevation 723 m) of Western Ghats, India. The botanical identification of the plant was carried out at the Botanical Survey of India, Pune 411001, India (Ref. No. BSI/WC/Tech/2012/244).

2.2. Preparation of plant extracts

Shade dried tubers of *E. nuda* were finely powdered with automix blender. One kilogram dry powder of bulbs were soaked in 3 L acetone, methanol, aqueous methanol (1:1) solvents (Merck, India) or distilled water separately. The crude extract was prepared by cold percolation for 24 h at room temperature (26 ± 2 °C). The filtrate was concentrated in vacuo at 40, 40, 56 and 60 °C to get acetone, methanol, aqueous methanol and aqueous extracts respectively. This process was repeated thrice to get total extracts. The extracts were labeled as AE (Acetone extract), ME (Methanol extract), AQME (Aqueous Methanol extract) and AQE (Aqueous extract), and were obtained as reddish/brown solid residues with 2.44%, 1.84%, 4.20% and 3.14% yield, respectively.

2.3. Chemicals

All the chemicals were of analytical grade and obtained from HiMedia, Merck or Fisher while standard antioxidants were procured from Sigma–Aldrich.

2.4. Determination of total phenols

The total soluble phenolic content was determined by Folin–Ciocalteu (FC) method.\textsuperscript{18} In brief, 10 μl of extract was taken and the final volume was made 2 ml with distilled water. To this 0.5 ml of FC reagent was added and sample was incubated for 3 min, followed by addition of 2 ml of Na2CO3 and samples were placed in boiling water for 1 min. The reaction mixture was allowed to cool and then absorbance was recorded at 765 nm on Chemito Spectrascan UV-2600 spectrophotometer. The total phenolic content was expressed as mg gallic acid equivalents (GAE) g\textsuperscript{-1} extract calculated using standard gallic acid calibration graph.

2.5. Determination of total flavonoids

Total flavonoids were estimated using modified Marinova et al\textsuperscript{19} method. Briefly, 10 μl plant extract was added to distilled water to make a final volume of 2 ml and kept at room temperature for 3 min. To this, 3 ml of 5% NaNO2 and 0.3 ml of AlCl3 was added. After 6 min incubation, 2 ml 1 M NaOH was added and volume was made 10 ml with distilled water. Absorbance was taken at 510 nm and the concentration of flavonoid compounds was expressed as mg quercetin equivalents per g extract.

2.6. Determination of total ascorbic acid

Total ascorbic acid was estimated by 2,4-dinitrophenylhydrazine (DNPH) method as described earlier.\textsuperscript{20} Ten microliter of each extract was separately taken and total volume was made to 2 ml with distilled water. To this 2 ml of DNPH and 1 drop of 10% thiourea was added, the mixture was heated in a boiling water bath for 15 min and cooled to room temperature. Five microliter of 80% (v/v) H2SO4 was added to the mixture at 0 °C in an ice bath. Absorbance was taken at 521 nm and ascorbic acid was used as standard.

2.7. Determination of total carotenoid content

For estimation of total carotenoids, 10 microliter plant extract was added to distilled water to make a final volume of 2 ml. To this, 3.75 ml of 10% methanolic KOH and 3.75 ml diethyl ether was added. The reaction mixture was washed with 5% ice cold saline and dried over anhydrous Na2SO4 for 2 h. The absorbance of the filtrate was taken at 450 nm and concentration of carotenoids was expressed as mg β-carotene equivalents per g extract.\textsuperscript{21}

2.8. Antioxidant activities

2.8.1. Total antioxidant activity (TAA)

Total antioxidant activity was determined using modified phosphomolybdenum method.\textsuperscript{22} The assay is based on the reduction of Mo(VI) to Mo(V) by sample compound and formation of green colored phosphate/Mo(V) complex at acidic pH (4.0). 0.1 ml of extract from varying concentrations (200–1000 μg ml\textsuperscript{-1}) was added to 1 ml reagent solution (0.6 M H2SO4, 28 mM sodium phosphate and 4 mM ammonium molybdate). The mixture was incubated at 95 °C for 90 min and the absorbance was measured at 695 nm after cooling the samples. Total antioxidant capacity was expressed as GAE.

2.8.2. FRAP (ferric reducing antioxidant power)

The antioxidant capacity of *E. nuda* extracts was estimated spectrophotometrically.\textsuperscript{23} The method is based on reduction of Fe\textsuperscript{3+}–tetra (2-pyridyl) pyrazine (TPTZ) complex to Fe\textsuperscript{2+}–tripyrildftiazine formed by action of electron donating antioxidants at low pH. FRAP reagent was prepared by the addition of 300 mM acetate buffer 10 ml TPTZ dissolved in 40 mM HCl and 20 mM FeCl3.6H2O in the ratio 10:1:1. Five hundred microliter of standard was added to 1 ml of FRAP and the mixture was incubated at 37 °C for 30 min. Absorbance was recorded at 593 nm against blank. The
values of FRAP were expressed as GAE for varying concentrations of the extracts (200–1000 µg ml⁻¹).

2.8.3. ABTS radical scavenging assay

Free radical scavenging activity of plant extracts was determined by ABTS radical decolorization assay.26 In brief, ABTS⁺ (2,2'-azino-bis-(3-ethylbenzthiazoline-6-sulphonic acid) cation radical) was produced by the reaction between 7 mM ABTS and 2.45 mM potassium persulfate in water (1:1). This reaction mixture was stored in dark at room temperature for 16–20 h. This ABTS⁺ solution was then diluted with methanol to get absorbance of 0.7 at 734 nm. Five microliter of plant extract was added to 3.995 ml of ABTS⁺ solution and incubated for 30 min and absorbance was measured at 734 nm. The results were expressed as percent scavenging effect of plant extracts.

2.8.4. DPPH radical scavenging activity

The antioxidant activity of the plant extract was examined on the basis of the scavenging effect on the stable 1,1-diphenyl-2-picrylhydrazyl (DPPH) free radical activity as described by Braca et al.27 Ethanolic solution of DPPH 0.05 mM (300 µl) was added to 40 µl of extract of different concentrations (200–1000 µg ml⁻¹). After 5 min, absorbance was measured spectrophotometrically at 517 nm against extract blank. The radical scavenging activity of the plant extract, expressed as percent inhibition was calculated according to the following equation:

Percent inhibition of DPPH radical = [(absorbance control – absorbance test)/absorbance control] × 100

2.8.5. Hydroxyl radical scavenging activity

Hydroxyl radical scavenging activity was measured by studying the competition between deoxyribose and test extract for hydroxyl radical generated by Fenton’s reaction.26 The reaction mixture contained deoxyribose (2.8 mM in KH₂PO₄–KOH buffer, pH 7.4), FeCl₃ (0.1 mM), EDTA (0.1 mM), H₂O₂ (1 mM), ascorbate (0.1 mM), and various concentrations of the sample extracts (200–1000 µg ml⁻¹) in a final volume of 1.0 ml. The reaction mixture was incubated for 1 h at 37 °C. Deoxyribose degradation was measured using thiobarbituric acid (TBA) assay. One ml of TBA (1%) and 1 ml of TCA (2.8%) was added to above mixture and incubated at 100 °C for 20 min. The development of pink color was measured at 532 nm and percent inhibition was calculated.

2.8.6. Determination of inhibition of lipid peroxidation

Lipid peroxidation inhibition potential of E. nuda extracts were evaluated in vitro by following the modified Halliwell and Gutteridge27 method. Briefly, freshly excised goat liver was minced using glass Teflon homogenizer in cold phosphate buffered saline, pH 7.4. Ten percent homogenate was prepared and then was filtered to obtain a clear homogenate. The process of homogenization and filtration was carried on ice. Different concentrations of the extracts (200–1000 µg ml⁻¹) in water were added to the liver homogenate. Lipid peroxidation was initiated by adding 100 µl of 15 mM ferrous sulfate solution to 3 ml of the tissue homogenate. After 30 min, 100 µl of this reaction mixture was taken in a tube containing 1.5 ml of 10% TCA. After 10 min, tubes were centrifuged and supernatant was separated and mixed with 1.5 ml of 0.67% TBA in 50% acetic acid. The mixture was heated for 30 min in a boiling water bath. The intensity of the pink colored complex was measured at 535 nm. The degree of lipid peroxidation was asayed by estimating the TBARS (TBA-reactive substances) content. The results were expressed as percentage inhibition using the formula:

Percent inhibition of lipid peroxidation = [(absorbance control – absorbance test)/absorbance control] × 100

2.9. DNA protection activity

The ability of different extracts to protect DNA (pBR322, Merck, India) from damaging effects of hydroxyl radicals generated by Fenton’s reagent was assessed by DNA nicking assay28 with minor modifications. The reaction mixture contained 2.5 µl of DNA (0.25 µg) and 10 µl Fenton’s reagent (30 mM H₂O₂, 500 µM ascorbic acid and 800 µM FeCl₃) followed by the addition of 5 µl extract and the final volume of the mixture was brought up to 20 µl with distilled water. The reaction mixture was then incubated for 45 min at 37 °C and followed by addition of 2.5 µl loading buffer (0.25% bromophenol blue, 50% glycerol). The results were analyzed on 0.8% agarose gel electrophoresis by staining with ethidium bromide. Quercetin was used as positive control.

2.10. Statistical analyses

All experiments were conducted in triplicate to check the reproducibility of the results obtained. The graphs were plotted using Microcal Origin 6.0. The results are presented as means ± standard error (SE) and means were compared using Duncan’s Multiple Range Test (DMRT) at P ≤ 0.05, using MSTAT-C statistical software.

3. Results and discussion

Quantitative evaluation of phytochemicals, known for their roles in providing antioxidant properties to plants, from tuber extracts of E. nuda indicated the plant as a rich source of phenolics, flavonoids, ascorbic acid and carotenoids, though with solvent dependent variations in their contents (Table 1). Several reports have shown a correlation between higher amounts of total polyphenols in plants and correspondingly higher antioxidant potential.18,28–32 Our results also largely supported these conclusions. In the present study, overall, methanol extract (ME) showed maximum amounts of total phenols, flavonoids, ascorbic acid and carotenoids closely followed by aqueous methanol (AqME) extract. Total phenol content in tuber extracts expressed as gallic acid equivalent (GAE, Table 1). Total flavonoids were comparably in lesser quantities than total phenols in all the extracts, where ME showed highest concentrations (22.2 mg AAE g⁻¹) followed by AqME with 19.9 mg AAE g⁻¹. Similar patterns were seen in case of ascorbic acid and carotenoids as evidenced from the results presented in Table 1.

Even though free radical generation and oxidation process are intrinsic in energy management of all living organisms and are kept under strict control by several cellular mechanisms,33 however, if
the supply of oxygen is in excess or its reduction is insufficient, ROS are generated that causes cellular injuries and initiate peroxidation of polyunsaturated fatty acids in biological membranes. In recent years, the scientific community has strongly advocated exploration and use of plant-origin natural effective antioxidants, especially from edible plants and their parts as they have less or no side effects than their synthetic counterparts. Therefore, in the present study, E. nuda known for its enormous medicinal properties in folklore was selected for the first time for evaluating its antioxidant properties.

Antioxidant activity is generally attributed to phytochemicals present, a number of mechanisms in plants and the synergies between them. Thus, antioxidant activity of plant extracts cannot be evaluated by a single method. Hence, in order to explore and understand these possible mechanisms, several antioxidant assays including TAA, FRAP, ABTS, DPPH and OH radical scavenging assays were performed for evaluating antioxidant activities of E. nuda. The results confirmed that this plant has a broad range of antioxidant properties, including substantial inhibition of lipid peroxidation. The results of TAA and FRAP scavenging activity are summarized in Table 2. Amongst all the four extracts, again ME showed highest antioxidative potential (in terms of FRAP measured as GAE) of all the extracts of E. nuda tubers, which was gradually increased with increasing concentrations of the samples. The FRAP assay provides a reliable method for evaluation of antioxidant activities of various plant extracts and compounds as antioxidant capacity is directly correlated with its reducing capacity and our results are in conformity of these findings.

Fig. 1 depicts the results of ABTS radical scavenging activities of all the extracts as well as gallic acid (standard antioxidant compound). AQME showed highest capacity to scavenge ABTS cation radical amongst all the extracts, however the activities were inferior to gallic acid, and therefore the standard was used at comparably low range (20–100 µg ml⁻¹) than the extracts (200–1000 µg ml⁻¹), respectively. Proton radical scavenging is considered as an important attribute of antioxidants. ABTS, a protonated radical, has characteristic absorbance maxima at 734 nm which decreases with the scavenging of the proton radicals. The magnitude of free-radical quenching was dose-dependent and steadily increased with increase in plant sample concentrations (Fig. 1).

DPPH is a stable free radical and the DPPH assay based on its characteristic absorbance maximum at 517 nm which decreases on reaction with free radicals. The results showed noticeable antioxidant potential (in terms of FRAP measured as GAE) of all the extracts of E. nuda tubers, which was gradually increased with increasing concentrations of the samples. The FRAP assay provides a reliable method for evaluation of antioxidant activities of various plant extracts and compounds as antioxidant capacity is directly correlated with its reducing capacity and our results are in conformity of these findings.

Among the four extracts used, acetone extract did not show inhibition of OH radical generation at concentrations ranging from 200 to 1000 µg ml⁻¹ (Fig. 3). AQME again showed notably higher hydroxyl radical scavenging activity (IC₅₀ Value: 230 µg ml⁻¹) than other extracts as well as the standard (ascorbic acid, IC₅₀ Value: 617 µg ml⁻¹). The radical scavenging activity was found to be concentration dependent and was increased with concentrations of all the extracts and standard antioxidant as well (Fig. 3). Hydroxyl radical is an extremely reactive ROS which initiates auto-oxidation and attacks almost every biological molecule causing damage to DNA, protein and lipids leading to mutagenesis, carcinogenesis and aging. The plants with higher OH radical scavenging holds great importance as their consumption can result in controlling and mitigating the devastating effects of oxidative stresses.

The current results are of great importance as the extracts especially AQME exhibited great potential in scavenging the OH radicals.

Table 2 FRAP and total antioxidant activities (TAA) of E. nuda extracts.

<table>
<thead>
<tr>
<th>Conc. of extracts (µg ml⁻¹)</th>
<th>TAA of various extracts (GAE) (mean ± SE)</th>
<th>FRAP of various extracts (GAE) (mean ± SE)</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>AE</td>
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<tr>
<td>200</td>
<td>64 ± 2.1a</td>
<td>70 ± 3.3a</td>
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<td>400</td>
<td>66 ± 2.4a</td>
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<td>132 ± 4.7c</td>
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<tr>
<td>800</td>
<td>70 ± 3.1d</td>
<td>138 ± 4.4c</td>
</tr>
<tr>
<td>1000</td>
<td>75 ± 3.4</td>
<td>160 ± 5.1</td>
</tr>
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</table>

GAE: gallic acid equivalents. Each value represents the mean of three replications ± SE. Means within a column for each extract followed by different superscript letters were significantly different from each other according to Duncan’s Multiple Range Test (DMRT) at P ≤ 0.05.
investigators. In the present investigation, in vitro lipid peroxidation was induced in goat liver by FeSO₄ which takes place through OH radical generation by Fenton’s reaction. E. nuda tuber extracts showed significant inhibition of lipid peroxidation rate measured in terms of TBARS, an indicator of malondialdehyde (MDA) content which is a degradation product of lipid peroxidation (Fig. 4). The results hold great significance as AqME showed tremendous potential in terms of inhibition percentage of lipid peroxidation, and showed 100% inhibition at 1000 µg ml⁻¹ concentration followed by ME (96% inhibition) against standard ascorbic acid with 90% inhibition at similar concentration (Fig. 4). AqME thus offered a good degree of protection against the biological end-point of oxidative damage. Lipid peroxidation induces cellular damage that eventually leads to many human diseases. Antioxidant may offer resistance against the oxidative stress by scavenging the free radicals, inhibiting the lipid peroxidation and thus prevent diseases.

All the extracts were evaluated for their oxidative damage protective activity against a model DNA pBR322. Hydroxyl radicals generated by Fenton’s reaction are known to cause DNA damage, as DNA band is absent in Fig. 5, lane 3 and only a smear of degraded DNA can be observed. Even though all the extracts effectively mitigated the oxidative stress and protected the DNA from hydroxyl radicals generated by Fenton’s reaction, as confirmed by the presence of DNA bands, aqueous methanol extract (Fig. 5, lane 6) seems to be comparably most effective in maintaining the DNA intact followed by aqueous (Fig. 5, lane 4), acetone (Fig. 5, lane 7) and methanol extract (Fig. 5, lane 5), respectively. Standard antioxidant compound quercetin was also used for comparison with plant extracts (50 µg ml⁻¹ each) for DNA protection efficacy. Free radicals are known for DNA strand breaking and damage which eventually contributes to carcinogenesis, mutagenesis and cytotoxicity. Various researchers have reported the similar results and used plant extracts and fractions for DNA protection against oxidative damage.

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**Fig. 2.** DPPH radical scavenging activities of various extracts of *E. nuda*. Each value represents the mean of three replications ± SE. The bars with different letters are significantly different from each other at *P* ≤ 0.05 according to Duncan’s Multiple Range test.

**Fig. 3.** Hydroxyl radical scavenging activities of various extracts of *E. nuda*. Each value represents the mean of three replications ± SE. The lines with the same color and symbol with different letters are significantly different from each other at *P* ≤ 0.05 according to Duncan’s Multiple Range test.

**Fig. 4.** Inhibition of lipid peroxidation in goat liver by various extracts of *E. nuda*. Each value represents the mean of three replications ± SE. The lines with the same color and symbol with different letters are significantly different from each other at *P* ≤ 0.05 according to Duncan’s Multiple Range test.

**Fig. 5.** DNA damage protecting effect of aqueous, aqueous methanol, methanol and acetone extracts of tubers of *Eulophia nuda* against hydroxyl radicals induced DNA damage of pBR322. Lane 1: native pBR322 plasmid DNA; Lane 2: DNA + Fenton’s reagent + quercetin (50 µg ml⁻¹, positive control); Lane 3: DNA + Fenton’s reagent (DNA damage control); Lane 4: DNA + Fenton’s reagent + AqE; Lane 5: DNA + Fenton’s reagent + ME; Lane 6: DNA + Fenton’s reagent + AqME; Lane 7: DNA + Fenton’s reagent + AE.
4. Conclusion

It can be concluded that *E. nuda* contains considerable amount of total phenols, flavonoids, vitamin C and carotenoids and showed solvent-dependent variations in their quantities. Various *in vitro* antioxidant assays clearly indicated that the plant extracts exhibited broad spectrum of antioxidant properties mediated by effective scavenging of various free radicals and subsequently inhibited the lipid peroxidation. Amongst all the extracts, AqME showed considerably higher antioxidant activities than other extracts. The plant extracts successfully protected the DNA from damage caused by free radicals. Overall, methanol and aqueous methanol extracts showed great promise and should therefore be used further for fractionation and isolation of pure compound responsible for antioxidant activities.

**Conflicts of interest**

All authors have none to declare.

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10. Salar R, Seasotiya L. Free radical Scavenging activity, phenolic contents and antioxidant assays clearly indicated that the plant extracts exhibited broad spectrum of antioxidant properties mediated by effective scavenging of various free radicals and subsequently inhibited the lipid peroxidation. Amongst all the extracts AqME showed considerably higher antioxidant activities than other extracts. The plant extracts successfully protected the DNA from damage caused by free radicals. Overall, methanol and aqueous methanol extracts showed great promise and should therefore be used further for fractionation and isolation of pure compound responsible for antioxidant activities.