ISOLATION AND CHARACTERIZATION OF ANTIOXIDANT (FLAVONE-3-RUTINOSIDE, 3, 3', 4', 5, 7-PENTAHYDROXY) FROM LEAVES OF MELIA DUBIA

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ABSTRACT

Objective: Various antioxidants are extensively found at different levels in many medicinal plants. This study mainly focuses on identification and separation of antioxidant from the leaf extract of Melia dubia.

Materials and Methods: Soxhlet extraction adopted for extraction using solvents, namely hexane, chloroform, ethyl acetate, 70% Ethanol, methanol, and water. The antioxidant capacity of six extracts was analyzed by quick and convenient 1, 1-diphenyl-2-picrylhydrazyl method. The compound responsible for high scavenging activity was isolated and separated by chromatography techniques such as thin-layer chromatography and high-performance liquid chromatography. Antioxidant was characterized by UV-visible, Fourier-transform infrared (FT-IR), H1-nuclear magnetic resonance (NMR), C13-NMR, and mass spectroscopy.

Results and Discussion: On extraction, water extract shows the highest yield percentage followed by methanol and 70% ethanol. Based on our results, IC50 value among the extracts, methanol and 70% ethanol, showed high antioxidant activity. Antioxidant which is commonly present in the methanol and 70% ethanol extract is flavone-3-rutinoside, 3, 3', 4', 5, 7-pentahydroxy.

Conclusion: The occurrence of the antioxidant, i.e. rutin flavonoid was identified in the study, and the biological activity of this compound will determine in future work.

Key words: Soxhlet extraction, 1, 1-diphenyl-2-picrylhydrazyl assay, Thin-layer chromatography and high-performance liquid chromatography, Characterization, flavone-3-rutinoside, 3, 3', 4', 5, 7-pentahydroxy.

INTRODUCTION

An antioxidant is a compound that inhibits the process of oxidation or delays some types of cell damage. Oxidation is a chemical reaction that can create free radicals, leading to chain reactions that may damage cells [1]. Molecules, atoms, or ions with unpaired electrons are known as free radicals. They are highly unstable and active toward chemical reactions with other molecules and from oxygen, nitrogen, and sulfur, thus creating reactive oxygen species (ROS), reactive nitrogen species, and reactive sulfur species. ROS include free radicals such as the superoxide anion (O2-). These free radicals react rapidly with the membranes eventually causing cellular degeneration and finally lead to death [2]. The generation of free radicals through lipid peroxidation is caused due to the continuous usage of the same vegetable oil which is not even properly stored and by reusing the already fried oil (rancid). The reason sometimes could be economic, but then it is highly damaging to the health. Nowadays, smoking and chronic alcoholism are creating health problem, and it reduces many important antioxidants in the serum which is detrimental to the health. Reports say that the proper intake of antioxidants will be quenching all these inevitably free radicals present in the body and thus improving the health by lowering the risk of various diseases [3,4]. The importance of the antioxidant constituents of plant materials in the maintenance of health and protection from diseases and also it raising interest among scientists, food manufacturers, and consumers as the trend of the future are moving toward functional food with specific health effects [5].

Potential sources of antioxidant compounds have been searched in several parts of plant materials such as vegetables, fruits, leaves, oilseeds, cereal crops, barks and roots, spices and herbs, and crude plant drugs. Beta-carotene, lutein, lycopene, selenium, and Vitamins A, C, and E are the available antioxidant in recent days. High-dose supplements of the above antioxidants may be linked to health risks. For example, high doses of beta-carotene may increase the risk of lung cancer in smokers. High doses of Vitamin E may increase the risks of prostate cancer and one type of stroke. Antioxidant supplements may also interact with some medicines. For the health risk instead of using this already available anti-oxidant, flavonoid and phenolic compounds can provide an alternate source for antioxidant [6,7].

Flavonoids and other phenolics have been suggested to play a preventive role in the development of diseases such as cancer and heart disease. Flavonoids and other plant phenolics, such as phenolic acids, stilbenes, tannins, lignans, and lignin, are, especially, common in leaves, flowering tissues, and woody parts such as stems and barks [8]. Flavonoids also partly provide plant colors present in flowers, fruits, and leaves and it generally occurs as glycosylated derivatives in plants, although conjugation with inorganic sulfate or organic acid. The antioxidant activity of phenolics is mainly due to their reductox properties that act as reducing agents, hydrogen donors, and singlet oxygen quenchers and also have a metal chelation potential [9].

Antioxidant uses are being extensively studied in pharmacology, more specifically in the treatment of cancer, stroke, cardiovascular and neurodegenerative diseases, and certain diabetic complications. Antioxidants are also helping in protecting the skin from sun exposure roughness, winklike depth, ultraviolet-induced skin cancer, and skin swelling from sunlight. Hence, these antioxidants are used in body lotions, creams, so as to protect the skin from sunlight. Antioxidants are used as food additives to help guard against food deterioration. A common use is as stabilizers in fuels and lubricants to prevent oxidation and in gasoline to prevent the polymerization that leads
Table 1: Result of the extracts yield

| S.No | Extracted solvent | Yield (g)      | Yield (%) |
|------|-------------------|---------------|-----------|
| 1    | Hexane            | 0.3810±0.0152 | 1.52      |
| 2    | Chloroform        | 0.8242±0.0145 | 3.30      |
| 3    | Ethyl acetate     | 0.9091±0.0274 | 3.64      |
| 4    | Methanol          | 1.6942±0.0061 | 6.78      |
| 5    | Water             | 2.9477±0.0415 | 11.79     |
| 6    | 70% Ethanol       | 1.4235±0.1181 | 5.69      |

Table 2: Result of the antioxidant activity on different extracts of MD leaves

| S.NO | MD leaves | Sample concentration and the percentage of inhibition (µg/mL) | IC₅₀ value µg/mL |
|------|-----------|--------------------------------------------------------------|-----------------|
| 1    | Hexane    | 25  50  100  200  400                                       | 120.78±0.445    |
| 2    | Chloroform| 1.99  7.87  10.89  34.51  66.23  120.78±0.445             | 74.84±0.350     |
| 3    | Ethyl acetate | 12.60  19.42  24.28  48.12  65.05  53.89±0.890            | 33.70           |
| 4    | 70% Ethanol | 26.80  38.12  58.43  85.08  91.71  17.47±29.315          | 65.05           |
| 5    | Methanol  | 15.60  47.97  84.77  94.33  93.52  15.61±0.1845          | 85.08           |
| 6    | Water     | 14.22  19.55  23.48  31.29  46.90  86.30±0.4897         | 91.71           |

MD: Melia dubia

Fig. 1: Extraction by Soxhlet method

Fig. 2: Isolation of antioxidant by column chromatography

Fig. 3: Before spraying 1,1-diphenyl-2-picrylhydrazyl reagent

Fig. 4: After spraying 1,1-diphenyl-2-picrylhydrazyl reagent

Fig. 5: High-performance liquid chromatography spectrum of antioxidant

to the formation of engine-fouling residues. They are widely used to prevent the oxidative degradation of polymers such as rubbers, plastics, and adhesives that causes a loss of strength and flexibility in these materials [10,11].

Melia dubia is the medicinal plant, which is available in all seasons and possesses very good medicinal values in the Indian system. Recent days, it was used in the treatment of dengue fever in Tamil Nadu, India. Most of the people are employ number of methods for the isolation and characterization of compounds from plants or crude extracts, but this study aimed to isolate only bioactive compounds, especially antioxidant. This study mainly focuses on the identification and separation of antioxidant from the leaves of M. dubia, and before isolation process, the antioxidant activity of the all the extracts will be evaluated by
testing with 1,1-diphenyl-2-picrylhydrazyl (DPPH). Extracts which show high antioxidant potential will choose for further isolation and characterization process.

METHODS

Chemicals and reagents
All the chemicals and glassware (hexane, chloroform, methanol, ethanol, iodine, dimethylsulfoxide (DMSO), double-distilled water, precoated thin-layer chromatography (TLC) silica gel 60 F, and DPPH) used in extraction, antioxidant activity, and isolation are the analytical grade purchased from Ponmani and Co Chemicals, Tiruchirappalli, Tamil Nadu, India.

Plant material collection
The leaves of *M. dubia* were collected from in and around areas in Tiruchirappalli, and the plant is authenticated in the Rapinat Herbarium.

Table 3: MSMS data of m/z 611(+ve) under different collision energies using ESI-positive ionization

| Collision energy | MS-MS fragmentation of M/Z 611 (+VE) |
|------------------|--------------------------------------|
| −5               | 611 465 345 303                     |
| −10              | 611 465 345 303                     |
| −15              | 611 465 345 303                     |
| −25              | 464 345 303 85                      |
| −35              | 303 85 71                           |
| −45              | 303 85 71                           |
| −55              | 345 303 85 71                       |
| −60              | 303 85 71                           |

ESI: Electrospray ionization

Table 4: MSMS data of m/z 303(+ve) under different collision energies using ESI-positive ionization

| Collision energy | MS-MS fragmentation of M/Z 303 (+VE) |
|------------------|--------------------------------------|
| −5               | 303                                  |
| −10              | 303 285                              |
| −15              | 303 285 195                          |
| −25              | 303 285 229 207 195 153 105          |
| −35              | 303 229 195 153 105                 |
| −45              | 303 229 206 195 167 115 43           |
| −55              | 303 206 195 167 115 43              |
| −60              | 303 206 167 115 43                  |

ESI: Electrospray ionization

Table 5: The effect of shifting reagent on isolated antioxidant compound, i.e., rutin

| S.No | CD With shifting reagents | Spectral maxima (nm) |
|------|--------------------------|----------------------|
|      |                          | Band-I  | Band-II | Band-III |
| 1    | CD-1 in Methanol         | 256     | 281     | 355      |
| 2    | CD-1+2 drops of 2M NaOH  | 275     | 324     | 422      |
| 3    | CD-1+2 drops of 5% AlCl₃ | 278     | 305     | 431      |
| 4    | CD-1+NaOAc              | 271     | 328     | 399      |
| 5    | CD-1+NaOAc+H₃BO₃        | 268     | 300     | 383      |
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% of DPPH Scavenged=(Ab of Control-Ab of test⁄(Ab of control) ×100)

The IC₅₀ values were calculated by linear regression of plots, where the abscissa represented the concentration of the tested sample and ordinated the average percentage of radical scavenging activity.

Isolation and separation of antioxidant

Extracts (70% ethanol and methanol extracts) which showed a high percentage of scavenging activity were further taken for the isolation process. Selected extracts were spotted at 2 cm from the edge of the sheet. The chromatogram is developed in a mixture of a suitable solvent system (trial and error method) and dried at room temperature. The spots were visualized in UV and iodine chamber. The Rf values of the spots were recorded. After that, the TLC plate was dried and sprayed with DPPH reagent, and color changes were noted. The compound with similar Rf value identified in two extracts was pooled together. The active compound (CD-1) was separated by performing column chromatography and purified by preparative high-performance liquid chromatography (HPLC) [15,16].

Characterization of antioxidant

A separated compound was identified by the phytochemical screening test and further taken for spectral studies for the characterization purpose. The UV-visible absorption spectrum of CD-1 was taken, and methanol was used as a reference solvent. Functional groups are identified by FT-IR, and it was conducted by the minimal amount of CD-1 mixed with spectroscopic grade KBr and then well-grounded before preparing the pellet. Proton NMR (H¹-NMR) and Carbon-13 NMR (C¹³-NMR) were analyzed in Bucker NMR 200MHz spectrophotometer. DMSO and TMS were used as the solvent and internal standard, respectively.

RESULTS AND DISCUSSION

The yield of the extraction

In the extraction process, hexane extract was yellow in color, chloroform, and ethyl acetate in dark green color. 70% Ethanol, methanol, and water extracts were brown in color. Regarding yield percentage, water extract shows a high percentage of extract with 11.79% followed by methanol and 70% ethanol in the percentage of 6.78% and 5.69%.
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respectively. Least yield percentage showed by hexane is 1.52%. The Soxhlet extraction setup was showed in Fig. 1 and the yield percentage is explained in Table 1.

Antioxidant activity by DPPH method
The effect of radical scavenging potential is examined by discoloration of DPPH reagent, i.e. violet color of the DPPH was changed into yellow after adding extract. Radical scavenging activity on the different extracts of M. dubia leaves shows noticeable potential against stable DPPH radical, and it is shown in Table 2. 70% Ethanol and methanol extracts show greater antioxidant activity compared to others. Beyond the concentration, the 70% ethanol and methanol extracts show a decrease in the percentage of inhibition, i.e., at 400 µg/mL, and it indicates the insufficiency of DPPH reagent due to higher activity of particular extracts. The IC50 value of methanol with 15.61 µg/mL and 70% ethanol with 17.47 µg/mL are minimum in comparison with the other solvents namely Hexane, Chloroform, Ethylacetate, and Water. 70% Ethanol and methanol extracts were further taken into further identification and separation process.

Isolation and separation of antioxidant
Ethanol and methanol extracts were subjected to TLC for the isolation of antioxidant from extracts. In the TLC, the spots were developed in ethyl acetate:acetic acid:water:n-butanol with RF value of 0.36, and after spraying the DPPH reagent, the compound responsible for scavenging activity shows bright yellow in the violet background. The spot was collected, and a purified form of antioxidant was obtained by preparative HPLC. The Figs. 2-6 have shown the TLC, column, and HPLC report of antioxidant. The report of HPLC reveals that the retention time of antioxidant is 15.46, and peak area and the height of the predominant peak of antioxidant are 20,358,456 and 2,618,390, respectively.

Fig. 12: H1 nuclear magnetic resonance spectra of rutin

Fig. 13: C13 nuclear magnetic resonance spectra of rutin
CHARACTERIZATION OF ANTIOXIDANT

Screening test for flavonoid

The antioxidant shows a positive result for flavonoid test and is discussed as follows.

Alkaline reagent test

An aqueous solution of the antioxidant compound was treated with 10% ammonium hydroxide solution. Yellow fluorescence indicates the presence of flavonoids.
Magnesium and hydrochloric acid reduction
The antioxidant compound was dissolved in 5 ml of alcohol, and a few fragments of magnesium ribbon and concentrated hydrochloric acid (dropwise) are added. A color change from pink to crimson red indicates the presence of flavonol glycosides. Figs. 7 and 8 show the positive result for flavonoids.

UV-visible and FT-IR spectra
Isolated antioxidant exhibits signs of flavonoid. In general, most flavonoids consist of two major absorption maxima, one of which occurs in the range 240–285 nm (band-II) and the other in the range 300–400 nm (band-I). Similarly, a strong absorption spectrum of the compound at 256 nm (band-II) and 355 nm (band-I) reveals the presence of the flavonoid nucleus. In FT-IR spectra, peak at 3365 cm⁻¹ gives strong evidence of functional activity of the free OH group in the isolated compound and 1612.49 cm⁻¹ responsible for C=O stretching frequency.

Proton NMR spectra of antioxidant compound
Proton NMR spectra of the isolated compound give peak at δ-12.602 responsible for OH proton in C (singlet) similarly 6.196 and 6.191 ppm responsible for glycosides linkage in flavonoid nucleus. The antioxidant compound was dissolved in 5 ml of alcohol, and a few fragments of magnesium ribbon and concentrated hydrochloric acid (dropwise) are added. A color change from pink to crimson red indicates the presence of flavonol glycosides. Figs. 7 and 8 show the positive result for flavonoids.

C¹³ and H¹ NMR spectra of antioxidant compound
C¹³ and H¹ NMR spectra of antioxidant compound give peak at δ-12.602 responsible for OH proton in C (singlet) similarly 6.196 and 6.191 ppm responsible for glycosides linkage in flavonoid nucleus. For the G ring peak at 5.353 and 5.355 (dd, 1H in C5'), 4.414 and 4.379 (dd, H and OH in C2'), 3.717 (td) and 3.692 (dt) (H and OH in C3'), peak at 3.093 (dd) to 3.047 (dd) (H and OH in C4'), 3.308 (td, C5'), 3.293 and 3.271 (dd, 2H in C6'), for the H ring peak at 3.382 and 3.355 (dd, 1H in C1'), 3.217 and 3.207 (dd, H and OH in C2'), peak at 3.250 (td) to 3.2225 (t) (H and OH in C3'), peak at 2.507 to 2.499 (dd, H and OH in C4'), 3.263 (q, 1H in C5'), and 1.070 and 1.055 (d, 3H in C6').

In the case of C¹³ spectra solvent peak obtained at 40 ppm and peak at 156.853 responsible for C2 carbon in the same manner 133.740 (C3), 177.804 (C4), 157.039 (C5), 99.5 (C6), 164.490 (C7), 94.014 (C8), 161.655 (C9), 104.408 (C10), 122.021 (C11), 115.654 (C12), 145.188 (C13), 148.844 (C14), 116.700 (C15), 121.611 (C16) for the Grignard 10.1186 (C1), 74.503 (C2), 70.994 (C3), 72.722 (C4), 76.346 (C5), and 67.423 (C6) for the H ring 70.808 (C2'), 70.994 (C3'), 72.436 (C4'), 68.678 (C5'), and 18.177 (C6'). The structure and C¹³ and H¹ NMR spectra of isolated antioxidant compound (rutin) are shown in Figs. 11-13.

Conclusions
This research work mainly investigates the isolation of active antioxidant from the leaf extract of M. dubia. On extraction, water and methanol have high yield; similarly, methanol and 70% ethanol reveal superior antioxidant activity. In the point of the isolation process, 70% ethanol and methanol extract have a similar compound with the same RF and characteristics properties. The active compound was found to be flavone-3-rutinoside, 3,3',4',5,7-pentahydroxy, and its structure was confirmed by spectral analysis.

Effect on CD-1 using shifting reagents
Shifting reagents such as sodium acetate, boric acid, and alcoholic aluminum chloride were used for the structural diagnosis of flavone-3-rutinoside, 3,3',4',5,7-pentahydroxy. Table 5 undoubtedly explains the structure and its substitution in isolated antioxidant, i.e., rutin.

From Table 5 when adding sodium hydroxide, spectrum was shifted toward the bathochromic region and it is because of free OH group in C3'. Similarly, adding AlCl₃ bathochromic shift due to free OH in C5 and for NaOAc, the band -1 was shifted because of C7 free OH. Finally, adding a mixture of NaOAc and H₃BO₃, the spectrum resulted in a bathochromic shift for complexation at C3' and C4' OH and it is shown in Fig. 16 [17]. The above effect on shifting reagents confirms the structure and active sides of the antioxidant, i.e., rutin.

Mass spectrum of isolated antioxidant
Mass fragmentation spectrum of CD-1 is shown in Fig. 14, and the energy required for the fragmentation is shown in Table 3. It clearly explains the major parent molecular ion (M+1) peak at m/z = 611.16 responsible for flavone-3-rutinoside, 3,3',4',5,7-pentahydroxy (Rutin). Furthermore, the flavonoid nucleus was fragmented, and its fragments and energy required for the fragmentation electrospray ionization are listed in Table 4. Based on the MSMS fragments obtained from m/z 611[+ve] and m/z 303[+ve], the proposed fragmentation pathway of the compound is displayed in Fig. 15. From the above shreds of evidence, the molecular formula of antioxidant (CD-1) is C₂₇H₃₀O₁₆ and accurate molecular weight is found to be 610.5175.
3,3′,4′,5,7-pentahydroxy can be used in the food processing industry and as preventive for many diseases in the field of medicine. In general, after isolation, the biological activity of the compounds was tested, but in our work initially, the antioxidant activity of the different extracts was identified after identifying best extract, and further isolation and characterization process was carried out. This work is one of the ways for finding and isolating useful bioactive compounds in a similar manner for the rest of the antioxidants present in the leaves. This approach findings may lead to a different area of research in this field.

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CONFLICTS OF INTEREST

The authors have declared no conflicts of interest.

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