Comparative Evaluation of Ethyl Acetate, Hexane and Methanol Dr.K. S. Manjunatha

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Abstract

Cellular damage arising from free radical is one of the fundamental mechanism underlying a number of human neurodegenerative disorder like diabetes, inflammation, Alzheimer's disease, autoimmune pathologic and digestive system disorder. Thus antioxidant plays an important role in the treatment of such disease. The present study aims at a comparative evaluation of ethyl acetate, hexane and methanol extract of Vitex leucoxylon Linn. bark for antioxidant activity. Vitex

Keywords:

Vitex leucoxylon;

DPPH, Nitricoxide

Superoxideandhydroxylradical scavenging

leucoxylon Linn. a medicinal plant of the verbenaceaefamily, used in traditional medicine for relieving headache and catarrh. HIME was 1.8964 μ g/ml after 48 h of incubation. In this study, it was observed that HIME induces a concentration dependent inhibition of HT29 cells, with an IC50value of 1.8964 μ g/ml after 48 h of incubation

Introduction:

Vitex leucoxylon Linn. (Verbenaceae) commonly known as Songarbhi (Marathi) an excellent herbal crude drug in the nature which has composition of the entire essential constituent that are required for normal and good health of human. It is small to large tree with a sort thick trunk and a spreading crown and almost throughout the Deccan peninsula of India up to an altitude 900 metres, it extends northwards up to Jhansi and part of Bihar. The trees are generally found on the river bank, stream & ponds. The root and the bark are astringent and roots are used as a febrifuge. The leaves are smoked for reliving headache and catarrh and are also used for medicinal baths in fever andanaemia 1.

revealed General pharmacological studies anti-psychotic, anti- depressant, analgesic, antiinflammatory, anti-parkinsonian and anti-microbial activities of aqueous **2.** Sarmaet al ³have studied and ethanolic extracts of leaves of V. Leucoxylon and wound healing properties of the crude alcoholic extract of the leaves anti-inflammatory in acute inflammation model3. The roots and bark are astringent and the roots are reported to be used as a febrifuge. β- Sitosterol, dimethyl terphthalate, vitexin, and aucubinwere isolated from the leaves or barks of V. isovitexin, agnuside Leucoxylon⁴. Majority of the diseases/disorders mainly linked to oxidative stress are due to free radicals **5**. Free radicals are fundamental to process anv biochemical and an part of aerobic life and metabolism antirepresent essential common reactive oxygen species (ROS) include superoxide (O2) anion, hydrogen peroxyl (ROO) radicals and reactive hydroxyl (OH) radicals. The nitrogen derived andperoxynitrite free radicals are nitric oxide (NO) anion. ROS have of diseases states implicated in over а hundreds which range tissue disorders to carcinogenesis, and connective aging, physical injury, infection and acquired immunodeficiency syndrome 7.

In treatment of these diseases, antioxidant therapy has gained importance. Current research is now finding naturally occurring directed towards origin. antioxidants of plant Antioxidants have been reported to prevent oxidative damage by free radical and ROS, and may prevent the occurrence of disease, cancer and aging. It can interfere with the oxidation process by reacting with free radicals, chelating, and catalytic metals and also by acting as oxygen scavengers 8-9. Plant and plant products are being used as a source of medicine since long. The medicinal properties of plants have been investigated in the recent scientific developments throughout the world, due to their potent antioxidant activities, no side effects 10. Flavonoids and phenolic compound widely distributed in plants economic viability multiple biological effect, including antioxidant, exert which have been reported to free radical scavenging abilities, anti inflammatory, anticarcinogenicetc $^{f 11}$. They were chelator 12-13. Novel (2 L). The extract was suggested to be a potential iron concentrated in a rotary flash evaporator and dried in desiccators.

Hydroxyl Radical Scavenging Activity: The scavenging capacity for hydroxyl radical was determined according to the natural antioxidants from some plants modified method.

The assay was have been extensively studied in the past few years for their antioxidant and radical scavenging properties. In view of this and the present understanding about ROS-induced multiple diseases, we have selected one of such ayurvedic herb—Vitex—leucoxylon Linn.—The objective of this investigation was to ascertain the scientific basis for the use of this plant in the treatment of antioxidant, using different antioxidant models.

Materials and Methods:

Vitex leucoxylon Linn. (Verbenaceae) were collected in flowering stage during late September from the natural population of Jhansi (U.P.) and authenticated by Dr. P.B Singh, Head of regional Research Institute Jhansi, shade dried and powdered then passed from

40# mesh size.

Preparation of Various Extracts of Vitex Leucoxylon:

(750)of V. leucoxylon bark, was extracted with hexane (2 material g) L), ethyl acetate (1.75 L) and methanol (1.75 L) using a Soxhlet apparatus and the spent material was then successively extracted with aqueous methanol (80%, 2 L) and water performed by adding 0.1 ml of EDTA, 0.01 ml of ferric chloride, 0.1 ml of hydrogen of deoxyribose, 1.0 ml of test solutions (5-100 µg/ml) in distilled water, 0.33 ml of phosphate buffer (50 mM, pH 7.4) and 0.1 ml of ascorbic acid were then incubated at 37°C for 1 hr and dissolved in sequence. The mixture was 1.0 ml portion of the incubated mixture was mixed with 10% TCA and 1.0 ml of 0.5% TBA to develop the pink chromogen and measured at 532 nm.

DPPH Radical Scavenging Activity:

The free radical scavenging activity was measured in terms of hydrogen donating or radical scavenging ability, using the stable radical, DPPH 14. A 0.1 mM solution of DPPH in methanol was prepared and 1 ml of this solution was added to 3 ml of control i.e. standard butylated hydroxyl toluene (BHT) at different concentration (25-100 μ g/ml) and test solutions at different concentrations (5-100 μ g/ml) in different test tubes. Thirty minutes later, the absorbances were measured at 517 nm.

Nitric Oxide Scavenging Activity:

Nitric oxide scavenging activity was measured by the spectrophotometric

method 15. Sodium nitroprusside (5 mM) in phosphate-buffer saline was mixed with a control without the test compound, but with an equivalent amount of methanol. Test solutions at different concentrations (5-100 µg/ml) were dissolved in and incubated at 25°C for 30 min. After 30 min, to 1.5 ml of the incubated solution was diluted with 1.5 ml of Griess reagent (1% sulphanilamide, 2% phosphoric naphthylethylenediamine dichloride). 0.1% The absorbance of the chromophore formed during the diazotization of the nitrile with sulphanilamide the subsequent coupling with naphthyethylenediaminedihydrochloride was measured at 546 nm.

Superoxide Scavenging:

Superoxide scavenging was carried out using the alkaline dimethyl sulfoxide (DMSO) method ¹⁶. Solid potassium superoxide was allowed to stand in contact with dry DMSO for at least 24 hrs and the solution was filtered immediately before use; filtrate (200 μl) was added to 2.8 ml of an aquous solution containing nitrobluetetrazolium (56 μM), EDTA (10 μM) and potassium phosphate buffer (10 μM, pH 7.4). different concentrations Test at (5-100)ug/ml) were added andabsorbances were recorded at 560 nm against the control.

Statistical Analysis:

The results are presented as mean \pm SEM. All parameters were analysed using Student's t-test. P<0.05 was considered as significant.

Results:

Inhibition of DPPH Radical:

The potential decrease in the concentration of DPPH radial due to scavenging property of ethyl acetate extract of Vitex leucoxylon Linn and BHT showed significant free radical scavenging activity viz. 88.52 and 86.73 %, respectively at $100 \, \mu g/ml$, whereas Hexane and Methanol extract of Vitex leucoxylon Linn. did not show any significant activity (Table 1).

Nitric Oxide Scavenging Activity:

The scavenging of nitric oxide by ethyl acetate extract of Vitex leucoxylonLinn and BHT was concentration dependent. There was a moderate inhibition of nitric oxide formation with the maximum inhibition being 74.00 and 82.24% respectively at $100\mu g/ml$ ethyl acetate extract of Vitex leucoxylon Linn and BHT. Similar results were not found in case of Hexane and Methanol extract of Vitex leucoxylon Linn (Table 1)

Table 1: Free radical scavenging activity of various extracts of Vitex leucoxylon Linn.

Drug	Concentration	DPPHradical	Nitricoxide
Ethyl	(µg/ml)	inhibition(%)	
acetate	5		42.70±0.5411
extractof		10.60±0.2698	51.67±0.5457*
Vitex	10	17.24±1.396	62.29±1.0380**
Leucoxylon	25	54.21±2.191**	71.00±0.9290***
Linn.	50	84.29±0.1402***	74.00±1.7698***
(ECVL)	100	88.52±0.3861***	
II.			37.57±0.6910
Hexane	5	09.53±0.5543	41.28±0.5382
extractof	10	10.02±1.029	44.18±0.4970
Vitex	25	17.74±0.4495	47.24±0.6458*
Leucoxylon	50	20.99±0.5698	49.11±0.2250*
Linn. (HEVL)	100	26.74±1.6920	
Methanol	5		02.54±0.103
extractof	10	06.24±0.109	06.88±0.142
Vitex	25	12.43±0.122	13.99±0.005
leucoxylon	50	20.26±0.002	26.28±0.008
Linn.	100	22.26±0.009	33.81±0.029
(MEVL)		25.59±0.004	77.13±0.6458
Butylated	25	06 7210 2015	79.23±1.7770
	50	86.73±0.3915	82.24±0.4976
hydroxyl	100	88.47±0.1520	
toluene		91.45±0.1782	
(BHT)			

Valuesaremean±SEM,6independentanalysis,P<0.05*, P<0.01**,P<0.001***ascomparedtostandard(Student's t-test).

Scavenging: The ethyl acetate extract of Vitex leucoxylon Linn Superoxide Radical showed a moderate inhibition of the superoxide radical 74.22 and 81.76% respectively at 100 µg/ml. There significant inhibition was no by Hexane and Methanol extract of Vitex leucoxylon Linn. (Table superoxide radical 2).

Hydroxyl Radical Activity: The effect of ethyl acetate extract of Vitex leucoxylon Linn and BHT on hydroxyl radical and iron (II)-dependent deoxyribose damage was protected significantly at all concentrations; the percentage of inhibition of

hydroxyl radical being 79.04 and 73.03 % respectively at 100 μ g/ml. No significant inhibition of superoxide radical by Hexane and Methanol extract of Vitex leucoxylon Linn. (Table 2)

Discussion:

The antioxidative system protects the organism against ROS-induced oxidative damage. There are restrictions on the use of synthetic antioxidants such as BHT, as they are suspected to be carcinogenic Natural antioxidants therefore have gained importance. DPPH is a stable free radical at room temperature and accepts an electron or hydrogen radical to form stable diamagnetic molecule. The reduction capability а DPPH radicals was determined bv the decrease in its absorbance at 517 nm, which is induced by antioxidants. .The ethvl acetate extract of Vitex leucoxylon radical scavenging effects in different in-vitro antioxidant Linn has potent and free systems, Hexane Methanol extract of Vitex leucoxylon Linn. Showed no but and significant effects as compared to standard BHT.

Table 2: Free radical scavenging activity of various extracts of Vitex leucoxylon Linn.

Ethylacetateextract	 	25 C5 10 0100*	46.00+0.7004*
ofVitexleucoxylon	5	35.65±0.9198*	46.99±0.7081*
Linn.(ECVL)	10	57.05±1.2561***	52.37±0.5575**
	25	68.70±0.7579***	61.71±0.3296***
	50	71.50±0.8742***	67.15±0.6439***
	100	74.22±0.5889***	79.04±0.6439***
Hexaneextractof	5	28.56±1.6000	42.83±0.6519
VitexleucoxylonLinn.	10	39.61±1.8190	49.36±0.8242*
(HEVL)	25	38.40±1.7762	52.81±0.6751*
	50	38.49±1.8220*	62.83±0.4191*
	100	43.46±1.6551**	67.77±0.3100
Methanolextractof			
VitexleucoxylonLinn.	5	05.12±0.748	04.11±0.529
(MEVL)	10	08.50±0.539	05.66±0.549
	25	23.28±0.649	19.91±0.639
	50	28.26±0.674	32.35±0.458
	100	35.26±0.229	43.88±0.367
		35.20±0.229	43.8810.307
Butylatedhydroxyl	25	74.82±0.8156	
toluene(BHT)	50	77.06±0.8905	57.77±0.3100
	100		70.58±0.7873
		81.76±1.6011	73.03±0.3610

 $Values are mean \pm SEM, 6 in dependent analysis, P<0.05^*, P<0.01^{***}, P<0.001^{****} as compared to standard (Student's t-test)$

References:

- 1. Nandkarni KM: Indian MateriaMedica, Popular prakashan Mumbai, edition 3, Vol 1, 1976: 1278-1280
- 2. Makwana HG, Ravishankar B, Shukla VJ,Vijayan NP, Sasikala CK, Saraswathy VN: General pharmacology of Vitex leucoxylon linn leaves. Indian J. Physiol. Pharmacol 1994; 38: 95-100.
- 3. Sarma, SP, Aithal KS, Srinivasan KK, Udupa AL, Kumar V, Kulkarni D.R. andRajagopal PK: Antiinflammatory and wound healing activities of the crude alcoholic extract and flavonoids of Vitex leucoxylon. Fitoterapia 1990; 61: 263-265.
- 4. Rao RVK, Satyanarayana T. and Jena R: Phytochemical studies on Vitex leucoxylon L. Indian Drugs 1997, 34: 50-51.
- 5. Gutteridgde JMC, Free radicals in disease processes- A complication of cause and consequence. Free Radic. Res. Comm 1995; 19: 141-158.
- 6. Tiwari A: Imbalance in antioxidant defence and human diseases-Multiple approach of natural antioxidants therapy. Current Science 2001; 81: 1179-1187.
- 7. Joyce DA, Oxygen radicals in disease. Advance Drug Research Bulletin 1987; 127:476-79.
- 8. Buyukokuroglu ME, Gulcin I, Oktay M, and Kufrevioglu OI: In vitro antioxidant properties of dantrolene sodium. Pharmacology Research 2001; 44:491-495.
- 9. Shahidi F and Wanasundara PD:Phenolic antioxidants. Cri. Rev. Food. Sci. Nutrition 1992; 32: 67-103
- 10. Auudy BF, Ferreira L, Blasina F, Lafon F, Arredondo R and Tripathi PC, Screening of antioxidant activity of three Indian medicinal plants, traditionally used for the management of neurodegenerative diseases. J. Ethnopharmacology 2003; 84:131-138
- 11. Miller AL, Antioxidant flavonoids: structure, function and clinical usage. Alt. Med. Review 1996; 1:103-111.
- 12. Boyer RF, Clark HM and LarocheAP: Reduction and release of ferritin iron by plant phenolics. J. Inorganic. Biochemistry 1988; 32:171-181.
- 13. Havsteen B: Flavonoids a class of natural products of high pharmacological potency. Biochemistry Pharmacology 1983; 30: 1141-1148.
- 14. Madan MP, Raghavan G, Singh AK and Palpu P, Free radical scavenging potential of Saussareacostus. Acta. Pharma 2005; 55:297-304.
- 15. Rajeshwar Y, Senthil GP, Malay AG and Mazumder UK: Studies on in vitro antioxidant activities of methanol extract of Mucunapruriens (Fabaceae) seeds. Eur. Bull. Drug Research 2005;13:131-138.
- 16. Sreejayan N. and MN Rao: Free radical scavenging activity of curcuiminoids. Drug Research 1996; 46: 169-171