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In Vitro Lipid Peroxidation Inhibitory and Antimicrobial Activity of Phyllanthus niruri (Euphorbiaceae) Extract

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10 ABSTRACT

11 The present work was designed to evaluate the in vitro lipid peroxidation inhibitory and antimicrobial ac-12 tivities of the methanol extract of Phyllanthus niruri (MEPN) (Family: Euphorbiaceae). Lipid peroxidation 13 was measured by the optical density of the prepared solutions (10-320 μg/ml) and then the percent inhibi-14tion was calculated. Ascorbate/FeSO4-induced peroxidation was inhibited by standard antioxidants such 15as L-ascorbic acid, quercetin and MEPN. Moreover, the percent inhibition of the methanol extract was 16 increased in a concentration-dependent manner. IC50 value for the MEPN, L-ascorbic acid and guercetin 17 for lipid peroxidation was found to be 62.5 µg/ml, 41 µg/ml and 19.75 µg/ml respectively. The antimicro-18bial activity of MEPN was determined by disc diffusion method with various gram-positive and gram-19 negative microorganisms. The MEPN showed strong antimicrobial activity against Bacillus pumillus 8241, 20 Bacillus cereus, Escherichia Coli 54B and Vibrae Cholera at a concentration of 750 μg/ml/disc. However, 21 its activity against Staphylococcus aureus ML 152 and Vibrae cholera 14035 was less significant. The 22 antimicrobial activity of the extract was compared with the standard drug, chloramphenicol at a concentra-23 tion of 10µg/ml/disc. The results obtained in the present investigation clearly suggest that MEPN can be a 24potential source of natural antioxidant and antimicrobial agent.

25 Keywords: Phyllanthus niruri, In vitro lipid peroxidation inhibitory activity, Antimicrobial activity

There has been growing interest in the investigation 46 PN (family: Euphorbiaceae) is a perennial herb dis-27 of the natural products from plants for the discovery of 47 tributed throughout the tropical and subtropical regions 28 new antimicrobial and antioxidant agents as well as an 48 of both hemispheres. In India, it is widespread in drier 29 alternative route for the substitution of synthetic chemi- 49 tropical areas of Andhra Pradesh, Tamil Nadu, Kerala 30 cals, side effects of which are always in question. For 50 and Karnataka states of South India. It is named the 31 this, the essential oils and the extracts of many plants 51 'stone breaker' by the indigenous people. Whole plant, have been prepared and screened for their antimicrobial 52 fresh leaves and fruits are used to treat various ailments 33 and antioxidant activities leading to the accumulation of 53 like dysentery, influenza, vaginitis, tumors, diabetes, 34a large number of reports in the literature concerning the 54diuretics, jaundice, kidney stones, dyspepsia, antihepa-35 above-mentioned properties of plants [1-5]. Much atten- 55 tototoxic, antihapatitis-B, antihyperglycemic and also as 36tion has been paid to the plant extracts and the isolated 56antiviral and antibacterial [8]. Antitumor and anticar-37 compounds because of their less side effects and the 57 cinogenic activities of Phyllanthus amarus have also 38 strong resistance towards various microorganisms [6]. 58 been reported [9]. Other medicinal properties such as 39 Plant-based antimicrobials represent a vast untapped 59 hypolipidemic [10] and antiviral [11, 12] activities of 40 source for medicines and further exploration of plant 60 Phyllanthus niruri have also been shown. Several bioac-41 antimicrobials is needed as antimicrobials of plant ori- 61 tive molecules, such as lignans, phyllanthin, hypophyl-42gin have enormous therapeutic potential. They are effec- 62lanthin, flavonoids, glycosides and tannins, have been 43 tive in the treatment of infectious diseases while simul- 63 shown to be present in the extracts of PN [9]. The phy-44taneously mitigating many of the side effects that are 64tochemicals from PN and their pharmacological proper-45 often associated with synthetic antimicrobials [7].

65 ties were studied by Bagalkotkar et al [13]. Using a rat

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67 have shown that phyllanthin and hypophyllanthin pro-122 Laboratory, Calcutta University, Kolkata, India. 68 tected cells against carbon tetrachloride cytotoxicity 69 whereas *triacontanal* was protective against galactosa-123 *In vitro lipid peroxidation* 70 mine toxicity. PN is used as one of the components of a124 78treating viral and bacterial diseases [21].

80 activity of MEPN. In the present study, we have tested 134 ml was removed and treated with 0.2 ml sodium dode-81 the in vitro lipid peroxidation and antimicrobial activity 135 cyl sulphate (SDS) (8.1%); 1.5 ml thiobarbituric acid 82 (against various Gram positive and Gram negative bac-136(TBA) (0.8%); and 1.5 ml acetic acid (20%, pH 3.5). 83teria) of the methanol extract of PN.

MATERIALS AND METHODS

85 Chemicals

90(A.P), India.

91 Extraction procedure

The plant PN was obtained from the tribal area of 93 Karimnagar District, Andhra Pradesh, India. The plant 150 94 was identified taxonomically by Dr. Alok Bhattacharya 95 of the Botanical Survey of India (BSI), Shibpur, Kol-96kata, India. A voucher specimen (No. GPS-2) has been 151 98extract, the whole plant was dried in shade and pow-153ple of the extracts. 99 dered in a mechanical grinder. The powder of PN was 100 initially defatted with petroleum benzene (60-80°C) 101 followed by 1 liter of methanol by using a Soxhlet ex-155 107 antimicrobial activity.

108 Previously isolated classes of compounds

110 contained alkaloids, flavonoids, saponins and cou-165 the medium suitably spaced apart and the plates were 111 marins, polyphenols, tannins, terpenoids, lipids and lig-166 incubated at 5°C for 1 h to permit good diffusion and 112nans [13].

113 Microorganisms utilized for antimicrobial activity

115 Staphylococcus aureus ML 174, Staphylococcus aureus 171 activity. All data on antimicrobial activity are the aver-116ML 152, Bacillus pumillus 8241, Bacillus cereus, Es-172age of triplicate analyses. In order to determine the anti-117 cherichia coli 51, Escherichia coli 54B, Vibrea cholera 173 bacterial effect of the MEPN, chloramphenicol 11814035, Vibrea cholera 1353, and Vibrea cholera 174(10µg/ml/disc) were used as positive control. Inhibition 119226101) were obtained from the stock culture of Central 175 diameters were measured after incubation for 24 h at 120 Drugs Laboratory, Kolkata; Indian Institute of Chemical 17637°C.

66hepatocyte primary culture, Shamasundar et al [14]121Biology, Kolkata and Mycology and Plant Pathology

Lipid peroxidation induced by Fe²⁺-ascorbate system 71 multiherbal preparation for treating liver ailments [15]. 125 in rat liver homogenate by the method of Bishayee and 72Liver damage is followed by complex disturbances in 126Balasubramaniyam [24] was estimated as thiobarbituric 73the lipolytic activity of the vascular space which often 127acid reacting substances (TBARS) by the method of 74appeared with hyperlipoproteinemia in patients [16].128Ohkawa et al. [25]. The reaction mixture contained rat 75 Abnormalities with lipid metabolism have been reported 129 liver homogenate 0.1 ml (25% w/v) in Tris-HCl buffer 76in cholesteosis [17], alcoholism [18], chemical intoxica-130(20mM, pH 7.0); KCl (30mM); FeSO₄(NH₄)₂SO₄.7H₂O 77tion [19] and hepatitis [20]. The plant is also useful in 131(0.06mM); and various concentrations of PN extract in a 132 final volume of 0.5 ml. The reaction mixture was incu-Previously, we reported the antihyperglycemic [22] 133 bated at 37°C for 1 h. After the incubation period, 0.4 137 The total volume was made up to 4.0 ml with distilled 138 water and then kept in a water bath at 95 to 100°C for 1 139h. After cooling, 1.0ml of distilled water and 5.0 ml of 140*n*-butanol and pyridine mixture (15:1 v/v) were added to 141 the reaction mixture, shaken vigorously and centrifuged 142at 4000 rpm for 10 min. The butanol-pyridine layer was 87(TBA) were purchased from Sigma Chemicals Co. (St. 144 quantify TBARS. Inhibition of lipid peroxidation was 88 Louis, MO, USA). All other chemicals and reagents 145 determined by comparing the optical density (OD) of 89 used were purchased from SD-Fine Chem, Hyderabad 146 treatments with that of the control. Quercetin and L-147 ascorbic acid were used as the standard controls. The % 148 inhibition of lipid peroxidation was calculated by using 49the following formula:

$$\% inhibition = \frac{[A_{blank} - A_{test}]}{A_{blank}} \times 100$$

where A_{blak} is the absorbance of the blank reaction 97 preserved in our laboratory for future purposes. For the 152 and A test is the absorbance in the presence of the sam-

54 Determination of antimicrobial activity

Antimicrobial activity was measured using the stan-102 tractor for 72 h at a temperature not exceeding the boil-156 dard method of disc diffusion plates on agar [26]. Then 103ing point of the solvent [23]. The extract was filtered 1570.1 ml of each culture of bacteria was spread on agar 104 using Whatman filter paper (No. 1) and then concen-158 plate surfaces. For antibacterial assays, all bacterial 105 trated in vacuum and dried. The methanol extract was 159 strains were grown in Mueller Hinton Broth medium 106 used in the assay of lipid peroxidation inhibitory and 160 (Merck) for 24 h at 37°C. The concentration of bacterial 161 suspensions was adjusted to 10^8 colony forming units 162(108cfu/ml) in Mueller Hinton Agar. Paper discs (6 mm 163 in diameter) were impregnated on the agar to load 10µl The phytochemical study revealed that the MEPN₁₆₄ of each sample. The impregnated discs were placed on 167then transferred to an incubator at 37°C for 24 h. The 168 results were recorded by measuring the zones of growth 169 inhibition surrounding the disc. Clear inhibition zones Microorganisms (Staphylococcus aureus 8531,170 around the discs indicate the presence of antimicrobial

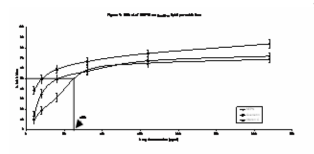


Fig 1. Effect of methanol extract of *Phyllanthus niruri* (MEPN) on lipid peroxidation.

177 Statistical Analysis

All treatments were performed in triplicate. Statisti-179 cal analysis was performed using Graphpad prism, 3.0 180 version (Graphpad Software Inc., San Diego, CA,217 181 USA). The statistical significance of a treatment effect 182 was evaluated by student's t-test and ANOVA. The val- 218 183 ues were expressed as mean \pm SD. IC₅₀ values for all the ²¹⁹ riety of diseases. Some of the medicinal usages have 184 above experiments were determined by linear regres-220 been proven in experimental models, which suggest that 185 sion. Probability limit was set at p < 0.05.

RESULTS

187 Effect of MEPN on lipid peroxidation

189 tioxidants namely L-Ascorbic acid and quercetin on the 228 optical density (OD) at 532nm. The results of the inves-190 in vitro inhibition of lipid peroxidation is shown in 229 tigations reveal that MEPN has no potent lipid peroxida-191 Fig 1. The generation of lipid peroxidase by Fe²⁺-230 tion inhibition activity. 192ascorbate in rat liver homogenate appears to be inhibited 231 193 by MEPN with IC50 value of 62.5 µg/ml. Though, the 232 by the disc diffusion method against various microor-194 inhibitory activity was observed, but it was found not so233 ganisms. Disc diffusion methods are used extensively to 195 remarkable when compared to L-Ascorbic acid and 234 investigate the antibacterial activity of natural sub-196 Quercetin. The percentage inhibition of lipid peroxida-235 stances and plant extracts [27]. These assays are based 197 tion of MEPN at 320 µg/ml was found to be 68.88% 236 on the use of discs as reservoirs containing solutions of 198 and for L-ascorbic acid and Quercetin the percentage 237 the substances to be examined. In case the activity is 199 inhibition was found to be 72.11% and 84.09%, respec-238 low, higher concentrated solutions are used. Because of

200 tively.

11 Effect of MEPN on antimicrobial activity

The data presented in Table 1 indicate that the 3 methanol extract of Phyllanthus niruri (MEPN) inhibit 14the growth of some of the tested microorganisms (Gram 5 positive and Gram negative) to various degrees. The)6MEPN at a concentration of 500 μg/ml/disc showed 17 moderate activity and 750 µg/ml/disc exhibited moder-)8 ate to strong antimicrobial activity against all the tested 99 microorganisms. The extract showed strong antibacteorial activity against Bacillus pumillus 8241, Bacillus 11 cereus, Escherichia Coli 54B and Vibrae Cholera. 212However, its activity against Staphylococcus aureus ML 213152 and Vibrae cholera 14035 was found to be less. The 214 antimicrobial activity of the extract was compared with 215the standard Chloramphenicol at a concentration of 21610µg/ml/disc.

DISCUSSION

PN has many effective traditional uses for a wide va-221 the extracts of the plant possess various pharmacologi-222 cal actions. Unsaturated lipids in liver tissue are very 223 susceptible to peroxidation when they are exposed to 224reactive oxygen species (ROS). In the present investiga-225 tion, we have incubated the liver tissue in presence of a 226ROS generating system, ascorbate/FeSO₄, and exam-The effect of MEPN and commercially available an egzined the effect on tissue homogenate by measuring the

The antimicrobial activity of the MEPN was studied

Table 1: Effect of methanol extract of Phyllanthus niruri (MEPN) on selected microbial strains

Microorganism	10% DMSO/ ml/disc	MEPN		Chloramphenicol
		500µg/ ml/disc	750µg/ ml/disc	(10µg/ml/disc)
Staphylococcus aureus 8531	9	6	10	16
Staphylococcus aureus ML 174	6	6	11	19
Staphylococcus aureus ML 152	6	6	7	22
Bacillus pumillus 8241	7	7	23	21
Bacillus cereus	6	10	16	14
Escherichia coli 51	6	7	10	24
Escherichia coli 54B	6	12	15	17
Vibrea cholera 14035	6	7	9	22
Vibrea cholera 1353	6	11	16	11
Vibrea cholera 226101	6	10	14	21

6-9mm: low activity; 10-14mm: moderate activity; ≥15mm: strong activity.

All the values were the mean of three experiments.

The values given are the diameter of zone of inhibition (mm) including disc diameter of 6mm.

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240erably used [27]. MEPN showed a broad spectrum of ³⁰⁴₃₀₅ 241 activity against all the microorganisms employed as 306 242 shown in Table 1. Chloramphenicol at a concentration 30712. 243 of 10µg/ml/disc was used as a positive control.

On the basis of the results obtained in the present in-245 vestigation, it is revealed that MEPN has no in vitro 311 246 lipid peroxidation inhibitory but has significant antim-312 247 icrobial activity. The phytoconstituents responsible for 313 248the inhibition of lipid peroxidation may be due to the 31414. 249 presence of flavonoids such as rutin, quercetin, quer-316 250 citrin, etc. and the antimicrobial activity of MEPN may 31715. 251 be due to the presence of p-cymene, a monoterpenoid, 318 252 present in the plant extract [13]. P-cymene was also 319 253 tested for antimicrobial properties using the paper disc 32116. 254diffusion method, in which it revealed a good anti-3 255 microbial activity [28]. More importantly, there have 323 256 been no side effects or toxicity reports from many years 32417. 257 on this plant. Although there has been extensive re-325 258 search on this plant, there is still a lot of scope for fur-32718. 259ther research, especially towards the mechanism of bio-328 260 logical activity of phytochemicals from this plant.

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