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EVALUATION OF ANTIMICROBIAL ACTIVITIES OF FRACTIONS OF PLANT PARTS OF Pterocarpus santalinoides

ABSTRACT

Aims: The study aims to investigate the antimicrobial activities of the leaves, seeds, bark and root of Pterocarpus santalinoides plant.

Study Design: Agar well diffusion and Agar well dilution methods were used to test the preliminary antimicrobial and minimum inhibitory/bactericidal/fungicidal concentrations respectively of Pterocarpus santalinoides plants.

Place and Duration of Study: Department of Pharmaceutical Microbiology and Biotechnology, Faculty of Pharmaceutical Sciences, Nnamdi Azikiwe University, Agulu Campus, Nigeria, between February -

Methodology: Primary extraction and fractionation was done on the plant parts with methanol, butanol, ethyl acetate and n-hexane. Agar diffusion method for the primary antimicrobial screening on Muller-Hinton agar (Bacteria) and Sabouraud dextrose agar (Fungi) were used to test the antimicrobial activities of the sixteen (16) samples on some microorganisms; Salmonella typhi, Escherichia coli, Candida albicans, Aspergillus niger, Microsporon canis and Trichophyton rubrum. The minimum Inhibitory concentration (MIC), minimum bactericidal/fungicidal concentration (MBC/MFC) and percentage inhibition diameter growth (PIDG) of the samples that gave activity were also evaluated.

Results: Twelve (12) samples showed inhibitory activity on at least one or more of the test organisms. The MIC range observed for the extracts and fractions that gave activity was 12.5 - 100 mg/ml. The nhexane fraction of the plant root gave the best value of 12.5 mg/ml against Microsporon canis. The best MBC/MFC value of 25 mg/ml was observed with the ethyl acetate fraction of the bark (against E. coli and M. canis) and the n-hexane fraction of the root (against M. canis). The result showed S. typhi to be the most sensitive organism to the metabolites of P. santalinoides. Extended spectrum activity was observed with the ethyl acetate fraction of the bark against three (3) of the test organisms; S. typhi, E. coli and M. canis. The determination of PIDG values for the test organisms against the plants' extracts/fractions showed that crude methanol extract (28.57%) and ethyl-acetate fraction (0.14%) of the leaves, Butanol fraction (0.14%) of the root (all against Salmonella typhi) were the most potent test samples.

Conclusion: The result indicates that the plants may have potential medicinal values and suggests its use in traditional medicine.

Keywords: Pterocarpus santalinoides; Antimicrobial; Methanol; n-hexane; Ethyl acetate; Butanol.

1. INTRODUCTION

Medicinal plants have been of great value to mankind. The use of herbs as complementary and alternative medicine has increased dramatically in the last 20 - 25 years [1]. Higher plants have been used for centuries as remedies for human diseases [2, 3]. Different plant parts have also been used for various forms of diseases and infections. This has encouraged research into screening of plants for antibacterial and antifungal activities [4].

The increased material worth of medical treatment and their strong physiological or chemical effects, contribute to the reason why individuals make use of herbal therapy. The increasing demand of plant extracts used in the cosmetic, food and pharmaceutical industries suggests that systematic studies of medicinal plants are very important in order to find active compounds and their use as a medicine for curing various diseases.

 2.52.4 Extraction of Plant Materials

Nnamdi Azikiwe University, Awka, Nigeria.

2.42.3 Microbial Test Organisms

The different plant parts of *Pterocarpus santalinoides* were prepared using the method of [11]. They were dried at room temperature (25 °C), grounded into a fine powder using laboratory grinding mill and stored in a cool and dry place. Total extraction of each plant materials were prepared by mixing with solvent (methanol) in the ratio of 1:10 (plant material/solvents) in a cold maceration system. The plant materials

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This research is aimed at investigating the antimicrobial activities of various parts of the plant against some indicative organisms implicated in their traditional use.

Pterocarpus santalinoides is a tree 9 - 12 m tall, 1 m diameter at breast height (DBH), with low straggling

branches. Bark thin and flaking in small patches, slash vellowish-white exuding drops of red gum [5]. It is

In Nigeria many indigenous plants including Pterocarpus Santalinoides are used as food or medicine. The

tender leaves are used as vegetable in soup preparing, while the stem bark is used in making pepper

soup. Pterocarpus santalinoides is a plant believed to possess potent antibacterial properties in ethno

medicine. The plants are used in treating rheumatism, diarrhea, dysentery, cough, ashma, diabetes, malaria, elephantiasis, cold and others [7]. The use of the plant leaves in treating skin disease such as

eczema, candidiasis, and acne have been reported [8]. The use of the concoction made from its root in

treating asthmatic patients have also been reported [9]. It is also used in treating diarrhea which is a

major cause of death as it has a proven anti enteropoling activity in traditional medicine [10].

a shade tolerant tree commonly found along riverine forests in Africa and tropical South America [6].

2. MATERIALS AND METHODS

2.1 Plants Materials

The fresh leaves, seeds, bark and roots of *Pterocarpus santalinoides* were sourced in September 2016 from Okofia - Otolo Nnewi, Anambra state, Nigeria, latitude 5⁰58'48.86" and longitude 6⁰54'30.78". They were identified and authenticated by a Curator in the Department of Pharmacognosy, Faculty of Pharmaceutical Sciences, Nnamdi Azikiwe University, Nigeria.

2.2 Culture Media and Reagents

Culture media used were Nutrient Broth, Mueller-Hinton agar (Oxoid Limited, England) and Sabouraud dextrose agar (Titan Biotech, India). Culture media were prepared according to the instructions of the manufacturers.

Reagents used include McFarland 0.5 turbidity standard (prepared from Barium chloride, Sulfuric acid and water), sodium hypochlorite solution and dimethyl sulphoxide (Triveni Chemicals, India).

2.3 Equipment

Equitron partially automatic autoclave (Medica Instrument Manufacturing CO., India), hot air oven (Genlab, UK), incubator (Genlab, UK), electronic weighing balance (Ohaus Corp., USA).

The microorganisms used in this study were two bacteria isolates (Escherichia coli and Salmonella typhi)

stored at 4 - 8 °C in Mueller-Hinton Agar slants and four fungi isolates (*Candida albicans*, *Aspergillus niger*, *Trichophyton rubrum* and *Microsporon canis*). These were clinical isolates previously purified and

standardized to McFarland. Their susceptibility to commonly used antibiotics was already established. The isolates were obtained from the Department of Pharmaceutical Microbiology & Biotechnology,

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were soaked in 1000 ml solvent in conical flasks for 12 hours. The extracts were then filtered using Whatman No. 1 filter paper and solvents removed through evaporation under reduced pressure at $45\,^{\circ}\text{C}$ using a rotary evaporator (Yamato, USA). The extracts were kept in stoppered sample vials at $4\,^{\circ}\text{C}$ until they were used.

2.62.5 Fractionation of Crude Methanol Extracts

The crude methanol extracts of the different parts of the plant were fractionated using solvent-solvent method with n-hexane, ethyl acetate and butanol in order of increasing polarity.

2.72.6 Percentage Yield Determination

The percentage yield (%, w/w) from all the dried crude methanol extracts was calculated using the formula outlined below:

Percentage (%) yield = Weight of extract X 100......Equation

Weight of dried plant material

2.8 Preliminary Antimicrobial Assay

The Preliminary antimicrobial assay for each of the crude extracts and their respective fractions was carried out using the agar well diffusion assay as described by [12]. The antimicrobial activity of the plant parts (i.e. crude extracts and fractions) were tested against standard clinical isolates (two bacteria and four fungi isolates) as listed above. A 0.5 McFarland standard bacterial/fungal suspension of each of the test isolates were prepared and these formed both the bacterial and fungal stock solutions used in the agar well diffusion assays as outlined below.

2.9 Agar Well Diffusion Assay

Briefly, the media, Mueller-Hinton agar, MHA (Oxoid, USA) and Sabouraud dextrose Agar, SDA, were prepared and treated according to the manufacturer's specification. MHA (38 g) and SDA (65 g) respectively were mixed with 1L of sterile distilled water and sterilized at 121 of for 15 mins. The media was allowed to cool to 50 °C and later transferred into 90 mm sterile agar plates and left to set. The sterile MHA and SDA plates were inoculated with the test culture from each of the test suspensions, thereafter, 20 ml of the sterile molten agar cooled to 50 °C was added then to the plate and was rocked clockwise and anti-clockwise to ensure even distribution of the test organism. This was done to obtain uniformity of the inoculums. A sterile cork borer was used to make wells (6 mm in diameter) on each of the MHA and SDA plates respectively. Aliquots of 60 µl of the stock concentration (100 mg/ml) of each extracts and fractions, reconstituted in dimethyl sulphoxide (DMSO) were applied in each of the wells in the culture plates previously seeded with the test organisms. Ciprofloxacin (5 µg) and miconazole (50 µg/ml) served as the positive controls for the bacteria and fungi respectively, while DMSO served as the negative control. The cultures were incubated at 37 °C for 18 - 24 hours for the bacterial plates and 25 – ^oC for 48 hours for the fungal plates respectively. The antimicrobial potential for each extract and fraction was determined by measuring the zone of inhibition around each well (excluding the diameter of the well). For each extract and fraction, three replicates were conducted against each organism. Each extract and fraction was tested against all the bacterial and fungal isolates.

Furthermore, based on the recorded activity, the extracts and fractions that showed activity were subjected to minimum inhibitory concentration (MIC) and maximum biocidal concentration (MBC) Determinations.

2.10 Anti-dermatophyte Activity

The anti-dermatophyte activity was conducted using the method as described by [13]. A sterile swab was used to aseptically inoculate each of the fungal suspension (*Trichophyton rubrum and microsporon canis*)

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on the surface of sterilized Sabouraud dextrose agar. The tests were carried out using a stock concentration of 100 mg/ml prepared by dissolving 200 mg of the crude extracts and fractions into 2 ml of DMSO. A well of 6 mm diameter was made in the agar plate then loaded with 60 μ l of each of the stock concentration of the crude extracts and fractions of the test samples and incubated at 28 \pm 2 °C for 15 - 20 days. The inhibition zone was observed and then recorded in millimeters using a transparent metre rule. The test was conducted in triplicate and results presented as mean. Miconazole 50 μ g/ml served as the standard positive control against the *dermatophytic species*.

2.11 Determination of Minimum Inhibitory Concentration (MIC)

The MIC was interpreted as the lowest concentration of the test samples (extracts and fractions) that inhibited visible growth. The MICs of the active samples were determined by agar dilution method as described by [13, 14] for the antimicrobial and anti-dermatophytic activities respectively. The MIC was determined for the micro-organisms that showed reasonable sensitivity to the test crude extracts and fractions. In this test, a stock solution of the crude extracts and fractions (2,000 mg/ml) was made, then 2-fold serial dilution was done to get graded dilutions (1000, 500, 250, 125, and 62.5 mg/ml) of each of the crude extract and fraction. Then 1 ml of each of these concentrationthese concentrations was transferred into a sterile petri dish and properly mixed with 9 ml of molten Mueller-Hinton agar and Sabouraud dextrose agar, then cooled to 45 - 50 ° C. After the mixing was done the final concentrations becomes 100, 50, 25, 12.5, and 6.25 mg/ml respectively. Finally, the different test organisms were streaked on the solidified agar properly labeled and incubated at 37 °C for bacteria and 28 ± 2 °C for fungi. The bacteria where incubated for 18 – 24 hours, while *Candida albicans* and *Aspergillus niger* were incubated for 48 hours and the dermatophytes incubated for 15 - 20 days. Each experiment was performed in triplicate.

2.12 Determination of the Minimal Bactericidal/Fungicidal Concentration of the Crude Extracts and Fractions (MBC)

This is the minimal concentration which kills off the cells in a microbial population. The MBC/MFC is the lowest concentration producing no evidence of growth [14]. It is an extension of the MIC procedure (stated above) carried out. Here, the plates that show no visible growth in the MIC test were selected. Then, they are incubated for 48 more hours. Thereafter, the plates were examined for signs of microbial growth.

2.13 Determination of the Percentage Inhibition Diameter Growth (PIDG)

Following the observation for the antimicrobial evaluation, the percentage inhibition diameter growth (PIDG) values were determined according to the equation as below.

Diameter of control

2.14 Data Analysis

The results were expressed as mean. Statistical analysis was carried out using one way analysis of variance (ANOVA) and SPSS (version 20) statistical program. The obtained results were considered significant at P = 0.05.

3. RESULTS

3.1 Extraction, Fractionation and Percentage Yield

Each of the leaves, seeds, barks and roots of *Pterocarpus santalinoides* were extracted with methanol as primary solvent. This yielded four (4) crude methanol extracts of the plant samples. The leaves (8.27 %)

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Plant Parts

Leaves

Seeds

Barks

Roots

the Test Isolates

species - Candida albicans, Aspergillus niger, Microsporon canis and Trichophyton rubrum. The IZDs were variable ranging from 2 - 9 mm. Broad spectrum antimicrobial agents (ciprofloxacin and miconazole) were used as a positive control while Dimethylsulfuroxide (DMSO) was used as negative control. A total of 16 crude methanol extracts and fractions of the plant parts were tested for their antimicrobial

activity among which 12 samples showed inhibitory activity on at least one or more of the test pathogenic microorganisms. In contrast, 4 samples (PSL n-hexane, PSB Crude, PSB Butanol, PSB n-hexane) showed no activity against any of the selected microorganism at the tested concentration. The result showed that ethyl acetate fraction of the plant bark (PSB Ethyl acetate) has antimicrobial activity against 3 of the tested organisms (Salmonella typhi, E. coli and Microsporon canis).

3.2 Antimicrobial Activity of Pterocarpus sanatalinoides Extracts and Fractions against

The antimicrobial activities of the plant extracts and fractions were tested against six (6) pathogenic cultures comprising of two (2) bacteria strains - Escherichia coli and Salmonella typhi, and four (4) fungi

gave the highest percentage yield as depicted in (Table 1). This was followed by the bark (5.97 %). The

The solvent-solvent fractionation of the four (4) crude methanol extract of the plant, Pterocarpus santalinoides with n-hexane, ethyl acetate and butanol in their order of increasing polarity gave a total of

four (4) fractions each of the three (3) secondary solvents giving a total sample of sixteen ((4) each of the

Extract weight (g)

45.15

9.87

26.87

15.16

Percentage yield (%)

8.27

1.97

5.97

4.92

crude methanol extracts, n-hexane, ethyl acetate and butanol fractions respectively) samples.

Table 1: The percentage yield of crude extracts obtained from Pterocarpus santalinoides

Plant

least value was observed with the seeds (1.97 %).

Powdered

546.10

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materials (Wt. in g)

The Ciprofloxacin (5 µg/ml) conventional drug used as control was active against the bacterial isolates with mean inhibition zone diameter (IZD) of 7 mm against S. typhi and 5 mm against E. coli. The Miconazole (50 µg/ml) used as control for fungal isolates was active against all the fungal isolates with mean inhibition zone diameter (IZD) of 18 mm for Candida albicans and Microsporon canis and 9 mm for Aspergillus niger and Trichophyton rubrum respectively. This reveals that the tests organisms used in the course of this work were susceptible strains. The DMSO serving as the negative control had no inhibition activity on any of the test organisms. The results of the antimicrobial activity of the plant parts are summarized in Tables 2.

Table 2: Mean Inhibition Zone Diameter (IZD) mm of the preliminary antimicrobial activity of the crude extracts and fractions of Pterocarpus santalinoides against some selected clinical isolates.

Samples	(100 mg/ml)	ST	EC	CA	AN	MC	TR
Leaves	PSL Crude	9	0	0	0	0	0
	PSL Butanol	3	0	0	0	0	0
	PSL Ethyl acetate	7	0	0	0	0	0
	PSL n-hexane	0	0	0	0	0	0

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Seeds	PSS Crude	0	5	0	0	0	0
	PSS Butanol	2	0	0	0	0	0
	PSS Ethyl acetate	5	0	0	0	0	0
	PSS n-hexane	4	0	0	0	0	0
Barks	PSB Crude	0	0	0	0	0	0
	PSB Butanol	0	0	0	0	0	0
	PSB Ethyl acetate	4	3	0	0	4	0
	PSB n-hexane	0	0	0	0	0	0
Roots	PSR Crude	0	2	0	0	0	0
	PSR Butanol	7	0	0	0	0	0
	PSR Ethyl acetate	6	0	0	0	0	0
	PSR n-hexane	0	0	0	2	6	0
Control	Ciprofloxacine 5 µg/ml	7	5	nd	nd	nd	nd
	Miconazole 50 μg/ml	Nd	Nd	18	9	18	9
	DMSO	0	0	0	0	0	0

Key: nd: not determined;ST: Salmonella typhi; EC: Escherichia coli; CA: Candida albican; AN: Aspergillus niger, MC: Microsporon canis,TR: Trichophyton rubrum,PSLCrude: Pterocarpus santalinodes Leaf (Crude extract); PSLButanol: Pterocarpus santalinodes Leaf (Butanol fraction); PSLEthyl acetate: Pterocarpus santalinodes Leaf (Ethyl acetate fraction); PSLn-hexane: Pterocarpus santalinodes Leaf (nhexane fraction), PSSCrude: Pterocarpus santalinodes Seed (Crude extract); PSSButanol: Pterocarpus santalinodes Seed (Butanol fraction); PSSEthyl acetate: Pterocarpus santalinodes Seed (Ethyl acetate fraction); PSSn-hexane: Pterocarpus santalinodes Seed (n-hexane fraction), PSBCrude: Pterocarpus santalinodes Bark (Crude extract); PSBButanol: Pterocarpus santalinodes Bark (Butanol fraction); PSBEthyl acetate: Pterocarpus santalinodes Bark (Ethyl acetate fraction); PSBn-hexane: Pterocarpus santalinodes Bark (n-hexane fraction), PSRCrude: Pterocarpus santalinodes Root (Crude extract); PSRButanol: Pterocarpus santalinodes Root (Butanol fraction); PSREthyl acetate: Pterocarpus santalinodes Root (Ethyl acetate fraction); PSRn-hexane: Pterocarpus santalinodes Root (n-hexane fraction); Standard error: ST: 0.7.

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The comparison of the activity of the extracts and fractions of Pterocarpus santalinoides according to the parts showed that the extracts and fractions significantly (p=0.05) inhibited some of the test organisms. For the leaves; the crude methanol extract (PSL Crude) had better antimicrobial activity to the only sensitive organism which is Salmonella typhi followed by ethyl acetate fraction (PSL Ethyl acetate) and Butanol fraction (PSL Butanol) respectively. The n-hexane fraction (PSL n-hexane) had no inhibition activity. This result indicates that the active agents in the plant leaves against this organism are chiefly polar compounds. For the Seeds; the ethyl acetate fraction (PSS Ethyl acetate) had the best inhibition activity against S. typhi followed by n-hexane (PSS n-hexane) and butanol (PSS Butanol) fractions respectively. The crude methanol extract (PSS Crude) was not active against S. typhi, however, was the only sample of the plant seeds active against E. coli. No part of the plant seeds has inhibition activity against C. albicans, A. nig r, M. canis and T. rubrum. From this result, the phytochemicals in the seeds active against S. typhi are mainly non-polar compounds, however, the compounds active against E. coli are polar compounds. It also showed that after fractionation of the crude methanol extract (PSS Crude), Formatted: Space Before: 0.3 line, After: 0.3

the active principles in the seed against *E. coli* was lost, which indicates an antagonistic effect of the fractionation process on the active principles. No part of the bark had inhibition activity on any of the tested organisms except for ethyl acetate fraction (PSB ethyl acetate) which had activity against *S. typhi*, *E. coli* and *M. canis*. This result showed that the active principles in the plant are bipolar compounds. This is logically in accordance to some work done [15] that tested the antimicrobial activities of the ethanol and water extracts of *P. santalinoides* bark against organisms including *S. typhi* and *E. coli*, of which the water extracts (a polar solvent) did not inhibit these test isolates at the concentration used. The ethanol extracts only marginally inhibited the organisms. For the roots, the crude methanol extract (PSR Crude) and the n-hexane fraction (PSR n-hexane) had no activity against *S. typhi*. The butanol fraction (PSR Butanol) had the best activity against *S. typhi*. Only the crude methanol extract (PSR Crude) showed inhibition activity against *E. coli*. The best antimicrobial activity was observed with the n-hexane fraction (PSR n-hexane).

Comparison of the susceptibility of all the test organisms to the various extracts/fractions of the plant parts, indicated that *Salmonella typhi* appeared to be the most sensitive organism to metabolites from *Pterocarpus santalinoides* with 9 of the extracts/fractions showing inhibitory activity and the best activity was observed with the crude methanol extract of the plant leaves (PSL Crude) were IZD of 8 mm, MIC of

Pterocarpus santalinoides with 9 of the extracts/fractions showing inhibitory activity and the best activity was observed with the crude methanol extract of the plant leaves (PSL Crude) were IZD of 8 mm, MIC of 50 mg/ml and MBC of 100 mg/ml were recorded. In contrast, Candida albicans and Trichophyton rubrum were found to be the most resistant organisms where none of the extracts/fraction showed antimicrobial effect at tested concentrations.

304 effect at tested concentration305 |

Three samples (PSS Crude, PSB Ethyl acetate and PSR Crude) showed antimicrobial activities against *Escherichia coli* with the best activity observed for crude methanol extract of the plant seed (PSS Crude) where IZD of 5 mm and MIC of 100 mg/ml were record. No part of the plants' leaves extract/fractions had activity against *E. coli*. This is similar to work done by [16], who evaluated the antimicrobial activities of the methanol extracts of the plant leaves against *S. aureaus* and *E coli*, of which *E, coli* showed no activity at all concentrations (up-to a maximum concentration of 50 mg/ml). *Microsporon canis* was inhibited by only two fractions (PSB Ethyl acetate and PSR n-hexane), while *Aspergillus niger* was inhibited by only one fraction (PSR n-hexane).

The ethyl acetate fraction of the plant bark (PSB Ethyl acetate) showed activities against 3 of the tested isolates producing IZD of 4 mm against *Salmonella typhi*, 3 mm against *Escherichia coli* and 4 mm against *Microsporon canis* showing an extended activity than the other extracts/fractions with activities against 2 Gram negative and 1 fungi isolates. This is similar to a work done by [17], were a broad spectrum activity of the plant bark was reported as was observed in this study. However, the best antifungal activity was observed for n-hexane fraction of the plant root (PSR n-hexane) where an IZD of 2 mm against *Aspergillus niger* and 6 mm against *Microsporon canis* were recorded.

The result of this study showed that the various extracts and fractions of the plant have antimicrobial activity against *Salmonella typhi*. Similarly, [18] reported the activity of this plant to this organism.

3.3 Minimum Inhibitory Concentration (MIC) and Minimum Bactericidal/Fungicidal Concentration (MBC/MFC) of *Pterocarpus santalinoides* Extracts and Fractions

The lowest concentration of the extracts and fractions of the plant parts at which no growth of microorganism was observed upon visual observation after incubation was considered the MIC value. The range of MIC of the plants extracts and fractions recorded was 12.5 – 100 mg/ml. The n-hexane fraction of the root gave the lowest MIC value (12.5 mg/ml) against *Microsporon canis*, followed by ethyl acetate fraction of the bark with 25 mg/ml against *E. coli* and *Microsporon canis*.

The lowest concentration of the extracts and fractions of the plant parts at which no growth of microorganism was observed upon visual observation after further incubation of the MIC plates were recorded as the Minimal Bactericidal Concentration (MBC) for bacteria isolates and Minimum Fungicidal Concentration (MFC) for fungal isolates. This is shown in Table 4. The ethyl acetate fraction of the bark

 (PSBE) and the n-hexane fraction of the plant root (PSRn) gave the lowest value (25 mg/ml) against E. coli and $Microsporon\ canis$ respectively.

3.4 Determination of the Percentage Inhibition of Diameter Growth (PIDG)

The determination of the PIDG for all the test organisms showed the most potent samples using the samples with higher Inhibition Zone Diameter (IZD) (Table 5) compared to the positive control used (5 μ g/ml Ciprofloxacin) as threshold/standard.

Table 3: Minimum Inhibitory Concentration (mg/ml) of the crude extracts and fractions of *Pterocarpus santalinoides* against tests isolates

Isolate	PSLC	PSLB	PSLE	PSLn	PSSC	PSSB	PSSE	PSSn	PSBC	PSBB	PSBE	PSBn	PSRC	PSRB	PSRE	PSRn ·
ST	50	100	100	-	-	-	50	100	-	-	100	-	-	100	50	-
EC	50	-	-	-	100	-	-	-	-	- 2	25	-	-1	-	-	-
CA	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
AN	-	-	-	-	-	-	-	-	-	-	- V	-	_	-	-	100
TR	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
MC	-	-	-	-	-	-	-	-	-	-	25	-	-	-	-	12.5

Key: ST: Salmonella typhi; EC: Escherichia coli; CA: Candida albicans; AN: Aspergillus niger, MC: Microsporon canis, TR: Trichophyton rubrum, PSLC: Pterocarpus santalinodes Leaf (Crude extract); PSLB: Pterocarpus santalinodes Leaf (Butanol fraction); PSLE: Pterocarpus santalinodes Leaf (Ethyl acetate fraction); PSLn: Pterocarpus santalinodes Leaf (n-hexane fraction), PSSC: Pterocarpus santalinodes Seed (Crude extract); PSSB: Pterocarpus santalinodes Seed (Butanol fraction); PSSE: Pterocarpus santalinodes Seed (Ethyl acetate fraction); PSSn-hexane: Pterocarpus santalinodes Seed (n-hexane fraction), PSBC: Pterocarpus santalinodes Bark (Crude extract); PSBB: Pterocarpus santalinodes Bark (n-hexane fraction), PSRC: Pterocarpus santalinodes Bark (Crude extract); PSRB: Pterocarpus santalinodes Root (Butanol fraction); PSRE: Pterocarpus santalinodes Root (Ethyl acetate fraction); PSRn: Pterocarpus santalinodes Root (n-hexane fraction).

Table 4: Minimum Bactericidal/Fungicidal Concentration (mg/ml) of the crude extracts and fractions of *Pterocarpus santalinoides* against tests isolates

1	Isolate	PSLC	PSLB	PSLE	PSLn	PSSC	PSSB	PSSE	PSSn	PSBC	PSBB	PSBE	PSBn	PSRC	PSRB	PSRE	PSRn
	ST	100	100	100	_ `	-	-	50	100	-	-	-	-	-	-	100	-
	EC	100	-		-	-	-	-	-	-	-	25	-	-	-	-	-
	CA	-		-	•	-	-	-	-	-	-	-	-	-	-	-	-
	AN	-	-	- -	-	-	-	-	-	-	-	-	-	-	-	-	-
	TR	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
	MC	-	-		-	-	-	-	-	-	-	25	-	-	-	-	25

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Key: ST: Salmonella typhi; EC: Escherichia coli; CA: Candida albicans; AN: Aspergillus niger, MC: Microsporon canis, TR: Trichophyton rubrum, PSLC: Pterocarpus santalinodes Leaf (Crude extract); PSLB: Pterocarpus santalinodes Leaf (Butanol fraction); PSLE: Pterocarpus santalinodes Leaf (Crude extract); PSSB: Pterocarpus santalinodes Seed (Crude extract); PSSB: Pterocarpus santalinodes Seed (Crude extract); PSSB: Pterocarpus santalinodes Seed (In-hexane fraction); PSSC: Pterocarpus santalinodes Seed (In-hexane fraction); PSBC: Pterocarpus santalinodes Bark (Crude extract); PSBB: Pterocarpus santalinodes Bark (Butanol fraction); PSBE: Pterocarpus santalinodes Bark (In-hexane fraction); PSRC: Pterocarpus santalinodes Bark (In-hexane fraction); PSRC: Pterocarpus santalinodes Bark (In-hexane fraction); PSRC: Pterocarpus santalinodes Root (In-hexane fraction).

The best MIC value was observed with n-hexane fraction of the plant root (PSRn) against *Microsporon canis*, followed by ethyl acetate fraction of the plant bark (PKBE) with MIC of 25 mg/ml against *E. coli and Microsporon canis*. The best MBC value was observed with ethyl acetate fraction of the bark (PSBE) and n-hexane fraction of the roots (PSRn) were 25 mg/ml was observed against *E. coli* and *Microsporon canis* respectively.

The Research conducted by [6], on the preliminary phytochemical analysis of methanolic extract of *Pterocarpus santalinoides* leaves showed the presence of alkaloids, anthocyanins, carotenoids, flavonoids, resins, saponins, steroids, terpenoids and tannins, while [19] revealed the presence of leucoanthocyanins, coumarins, flavonoids, mucilage, saponins and tanins. The presence of alkaloids, flavonoids, tannins, saonins, phenolics and cyanogenic glycosides were revealed by [15] in the ethanoloc extracts of the plant stem bark. No study was seen on the phytoconstituents of the plant's seed and root, but relatively will have the compounds as listed above, though environmental factors affects plants phytoconstituents. The antibacterial activities observed with these plant extracts and samples can be attribiuted to these secondary metabolites [15, 20]

Table 5: PIDGs of test organisms towards different extracts and fractions of *Pterocarpus santalinoides*

Samples	(100 mg/ml)	ST (%)	EC (%)	CA (%)	AN (%)	MC (%)	TR (%)
Leaves	PSL Crude	28.57	*	*	*	*	*
	PSL Butanol	-57.14	*	*	*	*	*
	PSL Ethyl acetate	0.14	*	*	*	*	*
	PSL n-hexane	*	*	*	*	*	*
Seeds	PSS Crude	*	0	*	*	*	*
	PSS Butanol	-71.43	*	*	*	*	*
	PSS Ethyl acetate	-28.57	*	*	*	*	*
	PSS n-hexane	-42.86	*	*	*	*	*
Barks	PSB Crude	*	*	*	*	*	*
	PSB Butanol	*	*	*	*	*	*
	PSB Ethyl acetate	-42.86	-40	*	*	-77.78	*
	PSB n-hexane	*	*	*	*	*	*
Roots	PSR Crude	*	-60	*	*	*	*
	PSR Butanol	0.14	*	*	*	*	*
	PSR Ethyl acetate	-14.29	*	*	*	*	*
	PSR n-hexane	*	*	*	-77.78	-66.67	*

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Key: *: Not applicable;ST: Salmonella typhi; EC: Escherichia coli; CA: Candida albican; AN: Aspergillus niger, TR: Trichophyton rubrum, MC: Microsporon canis, PSLCrude: Pterocarpus santalinodes Leaf (Crude extract); PSLButanol: Pterocarpus santalinodes Leaf (Butanol fraction); PSLEthyl acetate: Pterocarpus santalinodes Leaf (Ethyl acetate fraction); PSLn-hexane: Pterocarpus santalinodes Leaf (n-hexane fraction), PSSCrude: Pterocarpus santalinodes Seed (Crude extract); PSSButanol: Pterocarpus santalinodes Seed (Butanol fraction); PSSEthyl acetate: Pterocarpus santalinodes Seed (Ethyl acetate fraction); PSSn-hexane: Pterocarpus santalinodes Seed (n-hexane fraction), PSBCrude: Pterocarpus santalinodes Bark (Crude extract); PSBButanol: Pterocarpus santalinodes Bark (Butanol fraction); PSBEthyl acetate: Pterocarpus santalinodes Bark (Ethyl acetatefraction); PSBn-hexane: Pterocarpus santalinodes Root (Crude extract); PSRButanol: Pterocarpus santalinodes Root (Butanol fraction); PSREthyl acetate: Pterocarpus santalinodes Root (In-hexane fraction).

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The determination of PIDG values for the test organisms against the plant extracts/fractions showed that crude methanol extract (28.57%) and ethyl-acetate fraction (0.14%) of the leaves, Butanol fraction (0.14%) of the root (all against $Salmonella\ typhi$) were the most potent test samples using the samples that outstrips the positive control used, which was $5\mu g/mL$ Ciprofloxacin as the threshold/standard. Additional information from further studies on the mechanism of action of these extracts and fractions might contribute to its usage as an alternative to the conventional antibacterial drugs for the management of $Salmonella\ typhi$ and $E.\ coli\ implicated\ diseases$.

4. CONCLUSION

The study revealed that the various parts of the plant has antibacterial potentials especially organisms implicated in diarrhea as was seen with *S. typhi* and *E. coli*. The plant can be said to have negligible antifungal activities. It also shows that the solvent used for extraction or fractionation can affect the antimicrobial effect of the extracts/fractions. There is need for more studies towards isolation, identification and characterization of the bioactive agents and also test for activities against wilder spectrum of organisms.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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