Research Article

**Phyllanthus wightianus** Müll. Arg.: A Potential Source for Natural Antimicrobial Agents

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**Phyllanthus wightianus** belongs to Euphorbiaceae family having ethnobotanical importance. The present study deals with validating the antimicrobial potential of solvent leaf extracts of *P. wightianus*. 11 human bacterial pathogens (*Bacillus subtilis*, *Streptococcus pneumoniae*, *Staphylococcus epidermidis*, *Proteus vulgaris*, *Pseudomonas aeruginosa*, *Klebsiella pneumoniae*, *Salmonella typhimurium*, *Escherichia coli*, *Shigella flexneri*, *Proteus vulgaris*, and *Serratia marcescens*) and 4 fungal pathogens (*Candida albicans*, *Cryptococcus neoformans*, *Mucor racemosus*, and *Aspergillus niger*) were also challenged with solvent leaf extracts using agar well and disc diffusion methods. Further, identification of the active component present in the bioactive extract was done using GC-MS analysis. Results show that all extracts exhibited broad spectrum (6–29 mm) of antibacterial activity on most of the tested organisms. The results highlight the fact that the well in agar method was more effective than disc diffusion method. Significant antimicrobial activity was detected in methanol extract against *S. pneumoniae* (29 mm) with MIC and MBC values of 15.62 μg/mL. GC-MS analysis revealed that 29 bioactive constituents were present in methanolic extract of *P. wightianus*, of which 9,12-octadecaenioic acid (peak area 22.82%; RT-23.97) and N-hexadecanoic acid (peak area 21.55% RT-21.796) are the major compounds. The findings of this study show that *P. wightianus* extracts may be used as an anti-infective agent in folklore medicine.

1. Introduction

Plants play an important role in human life, primarily, as a source of food and medicine. Man is continuously faced with several lethal infectious diseases caused by pathogenic microorganisms [1]. In recent years, several pathogenic microorganisms have gained resistance to currently available synthetic antimicrobial agents and also caused many health problems [2]. Hence, there is an urgent need to discover an alternative new, broad spectrum, more active, and safer antimicrobial agent. Plant materials remain an important resource to combat serious diseases in the world. Especially, the plants from Pinaceae, Cupressaceae, Apiaceae, Burseraceae, Anacardiaceae, Palmaceae, Euphorbiaceae, Dracenaceae, and Fabaceae families are rich source for antimicrobial agents [3, 4]. Plants have an exceptional ability to synthesize *de novo* antimicrobial agents, in response to microbial attack for its protection [5]. Plant derived natural compounds (such as flavonoids, terpenoids, and steroids) have received considerable attention due to their diverse pharmacological properties including antibacterial and antifungal activities [6].

Antimicrobial components from plants which are mainly secondary metabolites act as inhibitors of bacterial growth, bacterial adherence, exopolysaccharide synthesis, DNA gyrase, cytoplasmic membrane function, and energy metabolism [7, 8]. Berberine, an isoquinoline alkaloid, which is present in roots and stem-bark of *Berberis* species, shows antimicrobial potential against bacteria, fungi, protozoa, and viruses [9]. Diterpene alkaloids, commonly isolated from the plants of the Ranunculaceae family, have antimicrobial properties [10]. Several phenolic compounds such as, caffeic acid, catechol, and pyrogallol are effective antimicrobial
agents. The antibacterial activity of some monoterpenes, diterpenoids, sesquiterpenes, triterpenoids, and their derivatives isolated from plants was recently reported [11]. Nowadays, search for plants with antimicrobial activity has evolved [12]. Importance of plants in drug discovery is growing due to vast diversity of the secondary metabolites which possess varied biological activities and act as main source of molecule leading the discovery of new, effective, and safer drugs [13]. Recent attention has been paid to extraction and isolation of biologically active compounds from plant species which are used in herbal medicines [14]. Pharmacognostic investigations of plants or plant extracts were needed to ascertain their biological activities which lead to the discovery of novel drugs or templates for the development of new therapeutic agents [15].

*Phyllanthus wightianus* (Euphorbiaceae) is a monoecious glabrous shrub which grows up to 4.5 m high and is found in the hills (750 to 1000 m) of peninsular India, on the floor and border of shoals, low altitudes in sandveld, hot dry deciduous, mopane woodlands, along banks of seasonal streams, and rivers [16]. It exhibits various biological properties, such as antimicrobial [17–19], larvicidal [20, 21], analgesic [22], wound healing [23], and antioxidant properties [24]. On the basis of the above information, the present investigation was focused on antimicrobial properties of different solvent leaf extracts and GC-MS analysis of bioactive extract of *P. wightianus* Müll. Arg.

### 2. Materials and Methods

#### 2.1. Plant Material

Fresh, matured, uninfected leaves of *P. wightianus* were collected from higher altitudes (900–1100 m) of Kolli Hills (latitude 10° 12′–11° 7′ N, longitude 76°–77° 17′ E), Namakkal district, Tamil Nadu, India. The plant material was authenticated by Botanical Survey of India (BSI) (reference number: BSI/SRC/5/23/2013-14/Tech/2081) Coimbatore, Tamil Nadu, India. The voucher specimen (specimen number: PU/BT/NDRDL/2010/05) has been deposited in the Natural Drug Research Laboratory, Department of Biotechnology, Periyar University, Salem, Tamil Nadu, India.

#### 2.2. Preparation of Extracts

Collected leaves were washed and shade-dried for three weeks and then powdered. The powdered plant material (500 g) was extracted in increasing polarity order (successively) with hexane, chloroform, acetone, ethyl acetate, and methanol in a Soxhlet apparatus for 72 hours. The extracts were filtered through Whatman number 1 filter paper and evaporated under vacuum at 40°C to yield crude extracts.

#### 2.3. Used Microorganisms

Three gram positive bacteria, namely, *B. subtilis* (MTCC 441), *S. pneumoniae* (MTCC 655), *S. epidermidis* (MTCC 435), and eight gram negative bacterial strains, such as *P. vulgaris* (MTCC 426), *P. aeruginosa* (MTCC 741), *K. pneumoniae* (MTCC 109), *S. typhimurium* (MTCC 98), *E. coli* (MTCC 739), and *S. flexneri* (MTCC 1457), with two clinical isolates (*P. vulgaris* and *S. marcescens*) and 4 fungal pathogens (*C. albicans*, *C. neoformans*, *M. racemosus*, and *A. niger*) were used in this investigation. The fungal strains were obtained from clinical laboratories of Salem District, Tamil Nadu. Each test organism was prepared by inoculating a loop-full of mother culture in a 5 mL of broth (Muller-Hinton broth for bacteria and Sabourud Dextrose broth for fungal cultures) and incubated at appropriate temperature and time for bacterial pathogens (37°C for 16 hours) and fungal strains (room temperature (28°C) for 72 hours). The culture turbidity was adjusted to 0.5 McFarland equivalence (1.5 × 10^8 CUf) prior to use.

#### 2.4. Agar Well Diffusion Method

The agar well diffusion method was employed to determine antibacterial activity of extracts as per the modified method of Natarajan et al. [19]. The standardized test cultures (50 μL) were swabbed on the per-molten Müller Hinton Agar (MHA) for bacteria and Sabourud Dextrose Agar (SDA) for fungus using aseptic cotton swab. Six wells were made in the seeded plates using sterile cork borer (5 mm diameter). Then, each extract (50 μL = 50 μg) was separately introduced into wells and allowed to diffuse at room temperature. Equal volume of DMSO was served as negative control. About 25 μL of standard antibiotics like fluconazole (fungus) and ciprofloxacin (bacteria) was used as positive control (each 1 μg/μL). The bacterial and fungal plates were incubated at 37°C for 24 hours and at room temperature for 72 hours, respectively. After the incubation period, the zone of growth inhibition was measured (in mm).

#### 2.5. Disc Diffusion Method

The disc diffusion test was performed by the method of NCCLS [25] with minor modifications. Test microbial suspension culture (50 μL) was spread on the MHA for bacteria and SDA for fungus by a sterile cotton swab. Sterile discs (5 mm diameter) were loaded with each extract (50 μL) and allowed to dry at room temperature. The dried discs were placed aseptically on the seeded plates. Standard antibiotic discs were used as positive control (gentamicin, vancomycin, and ampicillin for bacteria and fluconazole for fungus (10 mcg/disc)). The plates were incubated as the conditions mentioned in the well diffusion methods and the diameter (in mm) of clear zone of growth inhibition was recorded.

#### 2.6. Minimum Inhibitory Concentration (MIC)

The MIC was determined by broth microdilution bioassay method using the modified method of Eloff et al. [26]. MIC was carried out on the basis of antimicrobial results, the extracts which exhibited considerable antimicrobial activity against tested organisms. 100 μL of different concentrations (1–1000 μg/mL) of extracts was introduced into 96-well microplates containing 200 μL of Muller-Hinton broth and 20 μL bacterial cultures were added to each well. The microplate was closed with lid and incubated for 24 h at 37°C. After incubation period, 40 μL of 10 μM sodium fluorescein (INT) (0.2 mg/mL) was added to the wells to serve as an indicator of bacterial growth and incubated at 37°C for 1 hour. The minimum inhibitory concentration (MIC) was taken as the lowest concentration of the extract that completely inhibited bacterial growth.
2.7. Minimum Bactericidal Concentration (MBC). The MBC was determined as per method of Khan et al. [27]. MIC test dilutions (5µl) which showed no color change was subcultured on freshly prepared Mueller Hinton Agar plates and incubated at 37°C for 24 h. The lowest concentration in which no bacterial growth occurred was taken as the Minimum Bactericidal Concentration.

2.8. GC-MS Analysis. GC-MS analysis of bioactive extract was carried out on GC Clarus 500 Perkin Elmer system comprising AOC-20i auto sampler. The spectrums of unknown components present in the bioactive extract were identified by compared with known components spectrum which stored in the NIST and WILEY libraries. The name, molecular weight and structure of the components present in the bioactive extract were ascertained. The GC-MS analysis was carried out at Sophisticated Analytical Instrument Facility (SAIF), Indian Institutes of Technology, Chennai, India.

2.9. Statistical Analysis. All determinations were done at least in triplicate and averaged. Values were expressed as mean ± standard deviations. Statistical analyses were conducted using SPPS software (16.0 Version). Analysis of variance (ANOVA) in a completely randomized design and Tukey's multiple range tests were used to compare any significant differences between samples. The confident limits used in this study were based on 95% (P < 0.05).

3. Results and Discussion

The color of extracts and extractive yield of the plant material are presented in Table 1. The results of antimicrobial activity of P. wightianus were given in Table 2. All the extracts of P. wightianus show broad spectrum antibacterial activity in the range between 6 and 29 mm. The results of agar well diffusion method show that methanolic extract has significant activity against S. pneumoniae (29 mm) followed by S. epidermidis (17 mm). The considerable amount of antibacterial activity was observed in acetone extract against S. pneumoniae (28 mm) followed by S. flexneri (12 mm). In disc diffusion method, methanol extract exhibits good antimicrobial activity and the maximum growth inhibition was observed in S. pneumoniae (18 mm) followed by S. epidermidis (17 mm). Acetone extract having well to moderate antimicrobial activity and maximum activity was detected in S. epidermidis (18 mm) followed by S. pneumoniae (14 mm). Most of the tested clinical fungal pathogenic strains exhibit no sensitivity on the tested extracts in both agar well and disc diffusion methods. However, high antifungal activity was recorded in methanol extract against C. neoformans (10 mm) followed by hexane extract against M. racemosus (10 mm) in well diffusion method (Table 2). The overall results highlight that methanol extract exhibits significant activity against most of the tested pathogens compared with other extracts.

The MIC and MBC results (Tables 3 and 4) indicate that the crude extracts of P. wightianus inhibited the growth of selected bacterial species. The lowest MIC and MBC value were detected in methanolic extract against S. pneumoniae (15.62 µg/mL) and S. epidermidis (31.25 µg/mL). The rest of extracts showed moderate to high MIC and MBC values (500–1000 µg/mL) against most of the tested bacterial pathogens. Plant extracts are considered as having a good inhibitory activity, if they present MICs ≤ 100 µg/mL, a moderate inhibitory activity, if they present MICs ranging from 100 to 500 µg/mL, a weak inhibitory activity, if they present MICs ranging from 500 to 1000 µg/mL, and no inhibitory activity, if they present MICs > 1000 µg/mL [28, 29]. While considering these reports, the MIC and MBC values recorded from antibacterial activity of present investigation might be good to moderate.

The results highlights that significant antibacterial activity was observed in methanol extract against S. flexneri, S. pneumoniae, and S. epidermidis. Similar observations were made with P. wightianus methanolic extract [17]. Our previous findings, report that acetone extract expresses significant activity against most of the clinical bacterial pathogens compared to methanol extract [19]. Whereas, the present results show MTCC bacterial strains to be more sensitive to the methanol extract than acetone extract of P. wightianus. This difference in observation may be due to the high concentration (5 mg/well/disc) of extracts being used in the earlier study, whereas, in the present findings, low concentrations of extracts did not inhibit bacterial growth.

Studies have shown that methanolic extracts of many Phyllanthus species, such as P. acidus [30], P. muellerianus [31], P. amarus, P. maderaspatensis [32], P. debilis [33], P. amarus, P. emblica [34], and P. niruri [35], harbor promising antimicrobial activity which strengthens the present findings. Several reports stated that methanol is potent solvent for extracting variety of important phytoconstituents, like alkaloids, phenols, tannins, fatty acids, and flavonoids, which harbor antimicrobial potential which support the findings of present investigation [36, 37].

GC-MS analysis shows the presence of 29 compounds which were identified based on their retention time (RT), molecular formula, molecular weight (MW), and concentration (%) (Figure 1 and Table 5). Methnolic extract of P. wightianus has 9,12-octadecadienoic acid (with the peak area 22.82% and retention time 23.970) and N-hexadecanoic acid (with the peak area 21.55% and retention time 21.796) acting as major components. A variety of compounds, such as aliphatic ether, aliphatic carboxylic acid, aliphatic ester, alkene, and phenolic compounds were identified. Some of the compounds were present only in low quantities (ranging from 0.6 to 4%).

| Name of extract | Color          | Yield (%) |
|-----------------|----------------|-----------|
| Hexane          | Yellowish green| 6.24      |
| Chloroform      | Dark green     | 0.67      |
| Acetone         | Pale green     | 1.33      |
| Ethyl acetate   | Pale brown     | 2.49      |
| Methanol        | Brown          | 4.61      |
N-Hexadecanoic acid and 9,12-octadecadienoic acid are common secondary metabolites present in several plants [38–43] and are reported as having many biological properties, like antimicrobial, anti-inflammatory, hypcholesterolemic, cancer preventive, hepatoprotective, and antioxidant [44,45]. 9,12-Octadecadienoic acid was also stated as a model compound of unsaturated fatty acids, which selectively inhibits FabI enzyme in S. aureus and E. coli, catalyzing the final and rate-limiting step of the chain elongation process of the type II fatty acid synthesis (FAS-II) in bacteria [46]. Several fatty acids and phenolic compounds were identified in GC-MS analysis of methanol extract which may be the responsible for the antimicrobial activity. The mechanisms of antimicrobial action of fatty acids are non-specific modes of action [47]. However, antimicrobial effects of fatty acids were observed to form mostly either by a complete inhibition of oxygen uptake or stimulating uptake of amino acids into the cells, which occurs in a dose dependent manner [48]. Fatty acids intercalate in the phospholipid bilayer of microbes due to their lipophilicity, which increases the permeability of the cell membrane, dissipation of the proton-motive force, and leakage of inorganic ions, leading to cell death [49,50].

Studies have shown that phenolic compounds have bactericidal action by interfering with bacterial cell adhesins,
enzymes, cell envelope, and transport proteins [51]. They also increase the free radical concentration within the bacterial protoplasm and irreversibly complex with nucleophilic amino acids in microbial proteins determining loss of their function [52]. As a result, this causes bacterial cell lysis [53]. The antibacterial activity of methanol extract is not only caused by their major compounds, but it could be due to a synergism among their other components present in it. Hence, the presence of these components in higher quantity in methanol extract of P. wightianus may be responsible for better bioactivity.

4. Conclusions
The findings of present investigation show that agar well diffusion method is ideal for determining the antimicrobial activity of P. wightianus extracts. Methanolic extract of P. wightianus contributed significant activity against most of the

\begin{table}
\centering
\begin{tabular}{|l|c|c|c|c|c|c|}
\hline
Organisms & Methanol & Acetone & Ethyl acetate & Chloroform & Hexane & Ciprofloxacin \\
\hline
Bacillus subtilis & >1000^a & >1000^a & >1000^a & >1000^a & >1000^a & 3.9^b \\
(MTCC 441) & & & & & & \\
\hline
Escherichia coli & >1000^a & >1000^a & >1000^a & >1000^a & >1000^a & 15.62^b \\
(MTCC 739) & & & & & & \\
\hline
Klebsiella pneumoniae & 500^b & >1000^a & >1000^a & >1000^a & >1000^a & 7.81^c \\
(MTCC 109) & & & & & & \\
\hline
Proteus vulgaris & >1000^a & >1000^a & >1000^a & >1000^a & >1000^a & 15.62^b \\
(MTCC 426) & & & & & & \\
\hline
Pseudomonas aeruginosa & 250^c & 250^c & 500^b & >1000^a & >1000^a & 31.25^d \\
(MTCC 741) & & & & & & \\
\hline
Salmonella typhimurium & 500^b & >1000^a & >1000^a & >1000^a & >1000^a & 15.62 \\
(MTCC 98) & & & & & & \\
\hline
Shigella flexneri & 500^b & >1000^a & >1000^a & >1000^a & >1000^a & 31.25^c \\
(MTCC 1457) & & & & & & \\
\hline
Streptococcus pneumoniae & 15.62^d & 31.25^c & 250^b & >1000^a & >1000^a & 3.9^e \\
(MTCC 655) & & & & & & \\
\hline
Staphylococcus epidermidis & 31.25^d & 62.5^c & 500^b & >1000^a & >1000^a & 3.9^c \\
(MTCC 435) & & & & & & \\
\hline
\end{tabular}
\caption{MIC (µg/mL) for antibacterial activity of P. wightianus against some pathogens.}
\end{table}

\begin{table}
\centering
\begin{tabular}{|l|c|c|c|c|c|c|}
\hline
Organisms & Methanol & Acetone & Ethyl acetate & Chloroform & Hexane & Ciprofloxacin \\
\hline
Bacillus subtilis & >1000^a & >1000^a & >1000^a & >1000^a & >1000^a & 15.62^b \\
(MTCC 441) & & & & & & \\
\hline
Escherichia coli & >1000^a & >1000^a & >1000^a & >1000^a & >1000^a & 31.25^b \\
(MTCC 739) & & & & & & \\
\hline
Klebsiella pneumoniae & >1000^a & >1000^a & >1000^a & >1000^a & >1000^a & 15.62^b \\
(MTCC 109) & & & & & & \\
\hline
Proteus vulgaris & >1000^a & >1000^a & >1000^a & >1000^a & >1000^a & 31.25^b \\
(MTCC 426) & & & & & & \\
\hline
Pseudomonas aeruginosa & 500^b & 500^b & 500^b & >1000^a & >1000^a & 31.25^c \\
(MTCC 741) & & & & & & \\
\hline
Salmonella typhimurium & 500^b & >1000^a & >1000^a & >1000^a & >1000^a & 31.25^c \\
(MTCC 98) & & & & & & \\
\hline
Shigella flexneri & 250^c & 250^c & 500^b & >1000^a & >1000^a & 62.5^d \\
(MTCC 1457) & & & & & & \\
\hline
Streptococcus pneumoniae & 15.62^d & 62.5^c & 500^b & >1000^a & >1000^a & 3.9^e \\
(MTCC 655) & & & & & & \\
\hline
Staphylococcus epidermidis & 31.25^d & 125^b & 1000^a & >1000^a & >1000^a & 3.9^d \\
(MTCC 435) & & & & & & \\
\hline
\end{tabular}
\caption{MBC (µg/mL) for antibacterial activity of P. wightianus against some pathogens.}
\end{table}
Table 5: Compounds detected in methanol leaf extract of *P. wightianus* using GC-MS analysis.

| Peak | R. time | Peak area (%) | Molecular formula | Molecular weight | Compound name |
|------|---------|---------------|-------------------|------------------|---------------|
| 1    | 4.726   | 0.94          | C₆H₁₂O₂           | 116              | 2-Pentanone, 4-hydroxy-4-methyl-3-2-hydroxy-2-methyl-4-pentanone-2-  |
| 2    | 9.358   | 0.28          | C₆H₁₂O₃           | 126              | Levoglucosenone, 6,8-oxabicyclooct-2-en-4-one  |
| 3    | 10.907  | 0.67          | C₆H₁₂O₂           | 120              | 4-Vinylphenol  |
| 4    | 11.198  | 0.26          | C₉H₁₄O₂           | 174              | T-Butyldimethylsilyl acetate  |
| 5    | 12.387  | 0.08          | C₆H₁₀O₂           | 204              | 1-(4-Methoxyphenyl)-5-hexen-1-one  |
| 6    | 13.436  | 1.60          | C₆H₁₂O₂           | 126              | 1,2,3-Benzenetriol (pyrogallol)  |
| 7    | 15.093  | 6.07          | C₁₀H₂₄O₂          | 236              | Butanoic acid, 3-methyl-3,7-dimethyl-2,4,6-octatrienyl ester  |
| 8    | 15.241  | 3.56          | C₁₀H₁₈Br₂O₂       | 390              | 3,8-Dioxabicyclononane, 6-bromo-4-(1-bromopropyl)-2-[2-penten-4-ynyl]  |
| 9    | 15.525  | 1.04          | C₁₀H₁₈O           | 206              | Phenol, 2,4-bis(1,1-dimethylethyl)-2,4-ditert-butylphenol  |
| 10   | 16.037  | 2.46          | C₁₀H₁₈O₂          | 180              | 2-Benzo furanone, 5,6,7,7a-tetrahydro-4,4,7a-trimethyl-(2,6,6-trimethyl-2-hydroxy cyclohexylidene)acetic acid lactone  |
| 11   | 16.586  | 0.55          | C₁₂H₁₄O₄          | 222              | 1,2-Benzenedicarboxylic acid, diethyl ester phthalic acid  |
| 12   | 17.658  | 0.77          | C₁₃H₁₈O₃          | 332              | 2-Dodecahydro  |
| 13   | 19.062  | 0.71          | C₁₃H₁₈O₃          | 196              | 2(4H)-Benzo furanone, 5,6,7,7a-tetrahydro-6-hydroxy-4,4,7a-trimethyl-, (6S-cis)-(++)-loliolide  |
| 14   | 19.716  | 3.03          | C₁₂H₂₀O₂          | 208              | Pluchidiol  |
| 15   | 20.264  | 10.65         | C₂₀H₃₈           | 278              | 2,6,10-Trimethyl-14-ethylen-14-pentadecane  |
| 16   | 20.347  | 1.19          | C₁₈H₃₆O           | 268              | 2-Pentadecanone, 6,10,14-trimethyl-hexahydrofarnesyl acetone  |
| 17   | 20.591  | 3.16          | C₂₀H₄₀           | 296              | 3,7,11,15-Tetramethyl-2-hexadecen-1-ol (2e)-3,7,11,15-tetramethyl  |
| 18   | 20.831  | 5.85          | C₂₀H₄₀           | 296              | 3,7,11,15-Tetramethyl-2-hexadecen-1-ol (2e)-3,7,11,15-tetramethyl  |
| 19   | 20.925  | 0.36          | C₁₃H₁₆O₂          | 446              | Butanal  |
| 20   | 21.674  | 0.58          | C₁₉H₃₂            | 156              | 6-Octen-1-ol, 3,7-dimethyl-  |
| 21   | 21.796  | 21.55         | C₁₀H₁₈O₂          | 256              | N-Hexadecanoic acid, palmitic acid, pentadecanecarboxylic acid  |
| 22   | 23.300  | 0.53          | C₁₉H₃₂O           | 270              | Bis-(3,5,5-trimethylhexyl) ether  |
| 23   | 23.808  | 0.40          | C₂₀H₃₈           | 278              | 2,6,10-Trimethyl-14-ethylen-14-pentadecane neophytadiene  |
| 24   | 23.881  | 3.81          | C₁₀H₁₂O₂          | 186              | Nonanoic acid, 7-methyl ester methyl 7-methyl nonanoate  |
| 25   | 23.970  | 22.82         | C₁₀H₁₈O₂          | 280              | 9,12-Octadecadienoic acid-linoleic acid grape seed oil linoleic linole  |
| 26   | 24.199  | 5.93          | C₁₀H₂₀O₂          | 278              | 9,12,15-Octadecatrienoic acid, linolenic acids alpha-linolenic acid  |
| 27   | 24.308  | 0.68          | C₁₀H₂₀O₂          | 284              | Octadecanoic acid (stearic acid)  |
| 28   | 26.055  | 0.46          | C₆H₁₂O             | 126              | (3Z)-6-methyl-3,6-heptadien-1-ol  |

tested bacterial pathogens with least MIC and MBC values. Hence, methanol is the best solvent system for extracting the bioactive principles from leaves of *P. wightianus* which possesses promising antimicrobial principles which may be used in the treatment of infectious diseases caused by pathogenic microbes. The antimicrobial principles from the bioactive extracts may need further purifications to have its synthetic analogues which will be carried out in the future.
Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

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