

*Original Research Paper

Comparative Assessment of Antibacterial and Antifungal Activity of Dried Leaves of
Acalypha wilkesiana

Comment [t1]: Antibacterial and Antifungal
Activity of *Acalypha wilkesiana*

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Abstract

Comparative studies on the antimicrobial activities of the leaves of *A. wilkesiana* were carried out. Methanol was used as the extraction solvent. The crude methanolic extracts and four other derivative fractions were tested against human pathogenic bacteria namely strains of *S.aureus*, *S.pyogenes*, *E. faecalis*, *P. aeruginosa*, *P.vulgaris* and *E.coli* and fungi; *Aspergillusniger*, *A. flavus*, *A. carbonerium*, *Trichophytonmetagrophytes* and *Candida albicans*. 200mg/ml of each of the extract and the fractions were tested on the bacteria and fungi using the disc diffusion method. Results showed broad spectrum antimicrobial activity against the Gram-negative and Gram-positive bacteria but same cannot be said about its activity against the fungi. The result further showed that the ethyl acetate fraction was the most potent, closely followed by the aqueous while hexane fraction demonstrated the least antimicrobial activity. The extract and fractions were potent against some of the bacteria which standard antibiotics were not able to inhibit. Methanolic extracts of *A.wilkesiana* leaves showed a better antibacterial activity than antifungal activity. The demonstration of antimicrobial activity against the test organisms is an indication that there is possibility of sourcing alternative antibiotic substances in this plant for the development of newer antibacterial agents.

Keywords: *Acalypha wilkesiana*, antimicrobial, antibacterial, antifungal, resistance, zone of inhibition.

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63 Introduction.

64 Plants have been a source of medicine in the past centuries and today scientists and the general
65 public recognize their value as a source of new or complimentary medicinal products.
66 (Premanath, R., Lakshmideri, N. 2010). This plant-based, traditional medicine system continues
67 to play an essential role in health care, with about 80% of the world's inhabitants relying mainly
68 on traditional medicines for their primary health care (Owolabiet al., 2007). Long before
69 mankind discovered the existence of microbes, the idea that certain plants had healing potential,
70 indeed, that they contained what we would currently characterize as antimicrobial principles, was
71 well accepted. Since antiquity, man has used plants to treat common infectious diseases and some
72 of these traditional medicines are still included as part of the habitual treatment of various
73 maladies. For example, the use of bear-berry (Arctostaphylosuvaursi) and cranberry juice
74 (Vacciniummacrocarpon) to treat urinary tract infections is reported in different manuals of
75 phytotherapy, while species such as lemon balm (Melissa officinalis), garlic (Allium sativum)
76 and tea tree (Melaleucaalternifolia) are described as broad-spectrum antimicrobial agents (R'io,
77 JL., Recio, MC. 2005).

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78 During the last two decades, there has been a considerable increase in the study and use of
79 medicinal plants all over the world especially in advanced countries. Medicinal plants have been
80 used in Africa before the introduction of antibiotics and other modern drugs (Kabiret al., 2005)

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81 According to World Health Organization, medicinal plants would be the best source to obtain a
82 variety of drugs. Therefore, such plants should be investigated to better understand their
83 properties, safety and efficacy (Nascimentoet al., 2000)

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84 The success story of chemotherapy lies in the continuous search for new drugs to counter the
85 challenge posed by resistant strains of microorganisms. The investigation of certain indigenous
86 plants for their antimicrobial properties may yield useful results. Many studies indicate that in
87 some plants there are many substances such as peptides, unsaturated long chain aldehydes,
88 alkaloidal constituents, some essential oils, phenols and water, ethanol, chloroform, methanol
89 and butanol soluble compounds. These plants then emerged as compounds with potentially
90 significant therapeutic application against human pathogens, including bacteria, fungi or virus
91 (El astalet al., 2005).

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92 Medicinal plants are used by 80% of the world population as the only available medicines
93 especially in developing countries (EL-Kamali, HH., EL-amir, MY 2010). More importantly in
94 Africa, particularly West Africa, new drugs are often beyond the reach of the poor such that up
95 to 80% of the population use of medicinal plants as remedy against infections and diseases
96 (Kirby,G.C. 1996; Hostettmann, K., and Maston, A. 2002). Nigeria has a great variety of natural

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97 vegetation, which is used in trado-medicine to cure various ailments.(Egwaikhinde, P.A., Gimba,
98 P.C.2007). Among the plants use for medicinal purpose in Africa, particularly in Nigeria is
99 Acalyphawilkesiana.

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100 The genus “Acalypha” comprises about 570 species(Riley,H.P.1963). Acalyphawilkesiana
101 belongs to the family euphorbiaceae and grows as an annual bedding plant (Oladunmoye, M.K
102 2006).This large ,fast growing, evergreen shrub provides a continuous splash of colour in the
103 landscape with the bronze red to muted red, 4 to 8 inch long, hear-shaped leaves available in
104 varying mottled combinations of green, purple, yellow, orange, pink or white, depending upon
105 cultivar (Gilman, E.F 1999). Investigation is ongoing on almost all the available cultivars within
106 Nigeria with respect to their phytochemicals and antimicrobial action against medically inclined
107 and agriculturally related pathogens (Adeşinaet al., 2000; Akinde, B.E., Odeyemi, O.O 1987;
108 Alade,P.I.,Irobi, O.N., 1993;Ezekiel et al., 2009;Ogbo,E.M.,Oyibo, A.E 2008;Oladunmoye, M.K
109 2006).Consequently, this plant has been reported to have antibacterial and antifungal properties
110 (Alade, P.I., Irobi, O.N.,1993) as the expressed juice or boiled decoction is locally used within
111 Nigeria and some other parts of West Africa for the treatment of malaria, dermatological and
112 gastrointestinal infections (Akinde, B.E., Odeyemi, O.O 1987).

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113 Seeds from Acalyphawilkesiana are essential components of a complex plant mixture used by
114 traditional healers in southwest Nigeria in the treatment of breast tumors and inflammation
115 (Udobanget al., 2010).

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116 The aim of this study was to compare the antibacterial and antifungal potency of
117 Acalyphawilkesiana against bacteria and fungi of medical importance.

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119 MATERIALS AND METHODS.

120 Collection and Identification of Plant Samples

121 Healthy and matured fresh plant leaf samples of Acalyphawilkesiana were collected from the
122 horticulture garden of Babcock University, Ilishan Remo, Ogun state in May 2011 and Identified
123 by a botanist from the botanical unit of the same institution. The leaves were thoroughly rinsed
124 twice in running tap water and then in sterile water before being air-dried for 2weeks. The dried
125 leaves were grounded into fine texture using an electric blender, then stored in sealed and labeled
126 containers for use.

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127 Extraction

128 200mg of the dried and powdered A. wilkesiana leaves were extracted at room temperature with
129 absolute methanol. The crude methanol extract obtained was redissolved in methanol and made
130 aqueous with distilled water. The aqueous solution was extracted with hexane in a separating
131 funnel to obtain the hexane fraction. The aqueous layer was further partitioned with chloroform
132 to obtain the chloroform layer. Finally, the remaining aqueous layer was partitioned with

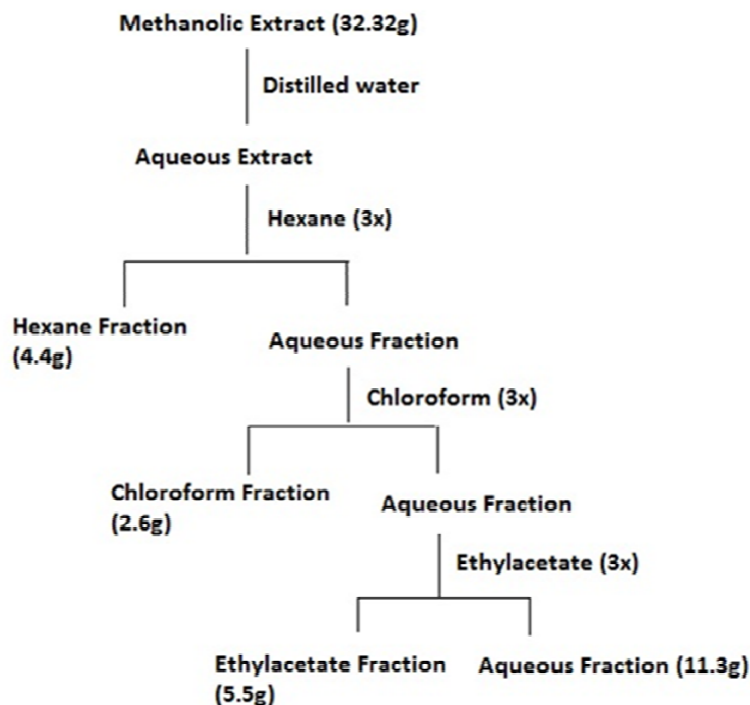
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ethylacetate to obtain ethylacetate fraction and aqueous fraction. A portion of the crude extract was also kept for analysis. All the fractions were concentrated in the rotary evaporator at 45°C and stored at 4°C till use. (Fig.1).



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138 Figure 1: Extraction yield of all the fractions of methanolic extract of *A. wilkesiana*.

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139 Collection and maintenance of test organisms

140 The test organisms used were standard strains of pathogenic bacteria and clinical isolate of
 141 fungi. They include five strains of Gram-positive bacteria; which are three strains of
 142 *Staphylococcus aureus* - *S. aureus* (ATCC 29213), *S. aureus* (ATCC 55620) and *S. aureus*
 143 (ATCC 25923), *Streptococcus pyogenes* (ATCC 8662) and *Enterococcus faecalis* (ATCC
 144 29212). Six strains of Gram-negative bacteria namely; three strains of *Escherichia coli* - *E. coli*
 145 (ATCC 23922), *E. coli* (ATCC 25922) and *E. coli* (ATCC 35218) others are
 146 *Klebsiella pneumoniae* (ATCC 700603), *Pseudomonas aeruginosa* (ATCC 27853) and *Proteus*
 147 *vulgaris* (ATCC 13315). The fungi isolates include: *Aspergillus niger*, *A. flavus*, *A. carbonarius*,
 148 *Trichophyton metagrophytes* and *Candida albicans*. They were obtained from the Department of

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149 Medical Laboratory Sciences, Babcock University, Ilisan-Remo, Ogun state. Biochemical
150 analysis was carried out on each of the test organisms for confirmation.

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151 Stock cultures were maintained at 4°C on slopes of nutrient agar. Active cultures for experiments
152 were prepared by transferring a loopful of cells from the stock cultures to test tubes of Mueller-
153 Hinton broth (MHB) and were incubated without agitation for 24hours at 37°C. The cultures
154 were diluted with Mueller-Hinton broth to achieve optical densities corresponding to 2.0×10^{-6}
155 colony forming units (CFU/ml)

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156 Antimicrobial Assay of the Crude metanolic extract and the various fractions.

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157 Antimicrobial susceptibility test for Bacteria

158 Stock cultures were maintained at 4°C on slopes of nutrient agar. Active cultures for experiments
159 were prepared by transferring a loopful of cells from the stock cultures to test tubes of Mueller-
160 Hinton broth (MHB) and were incubated without agitation for 24hours at 37°C. The cultures
161 were diluted with Mueller-Hinton broth to achieve optical densities corresponding to 2.0×10^{-6}
162 colony forming units (CFU/ml). The disc diffusion method was used to determine the
163 antibacterial activity of the crude methanol extracts and the other four fractions. In vitro
164 antibacterial activity was screened by using Mueller Hinton Agar (MHA) (LAB, United
165 Kingdom). The MHA plates were prepared by pouring 15 ml of molten media into sterile petri
166 dishes. The plates were allowed to solidify for 10 minutes and a standard loopful of each of the
167 eleven bacteria strain was streaked uniformly on the different plates and incubated at room
168 temperature for 10mins after which sterile cork borer of 5mm diameter was used to make two
169 ditches (wells) on each inoculated plate and filled with 200 mg/ml of the crude methanol extract
170 of the plant and the same was done for each of the eleven bacteria strain using the other four
171 fractions. These were carried out in duplicate for each organism. They were left on the bench for
172 30 minutes to ensure adequate diffusion of the fractions of extract and thereafter were incubated
173 at 37°C for 24 hours and the diameter of all resulting zones of inhibition around the ditches were
174 measured to the nearest millimeter along two axis and the mean of the two measurement was
175 calculated. The duplicate cultures were used for confirmation.

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176 Antibiotic susceptibility test was carried out on the test bacteria as control. A multi-sensitivity
177 disc bearing different antibiotics of GBMTS-NEG (Lot: NH05/P) (AbtekBiologicals Ltd.
178 Liverpool L9 7AR, UK) with their concentrations; Amoxycillin(25µg), Cotrimoxazole(25µg),
179 Notrofurantoin(300µg), Gentamicin(10µg), Nalidixic Acid(30µg), Ofloxacin(30µg), Augmentin(30
180 µg), Tetracycline(30µg) and DT-POS (Lot: JB04/P) with their concentrations; Ampicillin(10µg),
181 Chloramphenicol(10µg), Cloxacillin(5µg), Erythromycin(5µg), Gentamicin(10µg), Penicillin(1
182 i.u), Streptomycin(10µg), Tetracycline(10µg) were used against each of the test bacteria
183 inoculated on Mueller Hinton agar plates. These were incubated at 37°C for 24hours. After
184 incubation, the diameter of the zone of inhibition around each ditch was measured to the nearest
185 millimetre along two axis and the mean of the two readings was then calculated.

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186 Antimicrobial susceptibility test for Fungi

187 Stock fungi were maintained at room temperature on Potatoes Dextrose Agar (Oxoid, UK).
188 Active fungi for experiments were prepared by seeding a loopful of fungi into Potatoes dextrose
189 broth and incubated without agitation for 48 hours at 25°C . The broth was diluted with Potatoes
190 dextrose broth to achieve optical densities corresponding to 2.0×10^{-5} spore/ml for the fungal
191 strains

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192 The disc diffusion method was also used to screen for antifungal properties. In vitro antifungal
193 was screened by using Potatoes Dextrose Agar (PDA). The PDA plates were prepared by
194 pouring 15ml of molten media into sterile petri plates. The plates were allowed to solidify for
195 10minutes and 1ml of the test culture was introduced into agar and allowed to spread while the
196 excess was drained off. The plate was incubated at room temperature for 10 minutes. A sterile
197 cork borer of 5mm diameter was used to make two ditches (wells) on each plate and filled with
198 200mg/ml of the crude methanol extract. The same was repeated for each fungus strain using the
199 different fractions of the extract. The plates were incubated at 25°C for 96hrs and the resulting
200 zone of inhibition around the ditches were measured to the nearest millimeter along two axis and
201 the mean of the two measurement was calculated. The duplicate seeded agar were used for
202 confirmation. Control test was carried out using 10mg/ml of Fluconazole

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204 Determination of Minimum Inhibitory Concentration (MIC) :

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205 In determining the antimicrobial activity of *Acalypha wilkesiana*, the minimum bacterial growth
206 inhibition was accessed using the crude methanol extract and other fractions used in this study.

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207 2mls of nutrient broth was prepared into test tubes for the crude methanol extract and the four
208 fractions of the extract. 0.5ml of 25g/ml, 50g/ml, and 100g/ml of each extract fraction was added
209 to different test tubes containing the nutrient broth. This was prepared for each organism and
210 done in duplicate. A colony of 24hrs cultured organism was inoculated into test tube containing
211 1ml of normal saline to form a turbidity of 0.5 McFarland standard and was thereafter dispense
212 into the test tube containing the suspension of nutrient broth and the various fractions of the
213 extract . This was done for all the organisms at the varying concentrations. All test tubes were
214 properly corked and incubated at 37°C for 24hrs and at 25°C for 96hrs for bacteria and fungi
215 respectively . After which they were observed for absence or present of visible growth . The
216 lowest concentration without visible growth (turbidity) of organisms was regarded as the
217 Minimum inhibitory concentration (MIC). It was further standardized in terms of absorbance at
218 600 nm in a visible spectrophotometer. Positive and negative controls were set up alongside this
219 experiment.

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221 RESULT

222 Susceptibility of the test bacteria to crude and fractions of *A.wilkesiana* extract.

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223 The result of this study revealed the in vitro susceptibility of some bacteria to the crude extracts
 224 and other fractions of *A. wilkesiana*. Table 1 shows the mean \pm standard deviation of the zone of
 225 inhibition in the various agar plates of bacteria exposed to the extract fractions. It was noticed
 226 that all the fractions of *A. wilkesiana* extract used inhibited the growth of *S. aureus* (ATCC
 227 25923). However, the crude methanol extract, aqueous, ethyl acetate and hexane fractions
 228 inhibited *S. aureus* (ATCC 29213), while crude, aqueous and ethyl acetate fractions inhibited
 229 *S. aureus* (ATCC 55620). This study showed that all the *S. aureus* strains were the only organisms
 230 susceptible to the crude methanol extract while aqueous and ethyl acetate fractions were the only
 231 fractions that inhibited *P. vulgaris* (ATCC 13315), *P. aeruginosa* (ATCC 27853) and *S.*
 232 *pyogenes* (ATCC 8662). The *E. coli* strains and *Enterococcus faecalis* were resistant to the crude
 233 extract and two of the fractions except ethyl acetate and chloroform which inhibited *E. coli*
 234 (ATCC 35218) and *E. faecalis* (ATCC 29212) respectively. *Klebsiella pneumoniae* (ATCC
 235 15380), *E. coli* (ATCC 25922) and *E. coli* (ATCC 23922) were not susceptible to any of the
 236 fractions used in this study.

237 The aqueous fraction against *S. pyogenes* (ATCC 8662) yielded the highest inhibition value
 238 while ethyl acetate fraction gave the greatest number of inhibition, i.e. more test bacteria were
 239 susceptible to ethyl acetate fraction.

240 Table 1: The mean \pm S.D (mm) of zone of inhibition observed on bacteria cultured plates of
 241 isolates exposed to different fractions of *A. wilkesiana* extract

Organisms	Crude	Aqueous	Ethyl acetate	Hexane	Chloroform
<i>S. aureus</i> (ATCC 25923)	5.0 \pm 0.00	6.5 \pm 0.29	7.50 \pm 2.88	5.0 \pm 0.00	6.0 \pm 0.0
<i>S. aureus</i> (ATCC 29213)	7.5 \pm 0.86	5.0 \pm 0.0	7.0 \pm 0.41	3.0 \pm 0.0	0.00
<i>S. aureus</i> (ATCC 55620)	7.5 \pm 0.28	7.25 \pm 0.5	9.5 \pm 2.28	0.00	0.00
<i>P. aeruginosa</i> (ATCC 27853)	0.00	5.0 \pm 0.0	8.0 \pm 0.41	0.00	0.00
<i>P. vulgaris</i> (ATCC 13325)	0.00	7.0 \pm 0.0	6.5 \pm 0.29	0.00	0.00
<i>S. pyogenes</i> (ATCC 8662)	0.00	10.0 \pm 0.0	8.75 \pm 0.49	0.00	0.00
<i>E. faecalis</i> (ATCC 29212)	0.00	0.00	0.00	0.00	6.5 \pm 0.29
<i>E. coli</i> (ATCC 35218)	0.00	0.00	9.5 \pm 0.29	0.00	0.00
<i>E. coli</i> (ATCC 23922)	0.00	0.00	0.00	0.00	0.00
<i>E. coli</i> (ATCC 25922)	0.00	0.00	0.00	0.00	0.00
<i>K. pneumoniae</i> (ATCC 15380)	0.00	0.00	0.00	0.00	0.00

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244 Susceptibility of the test fungi to crude and fractions of *A. wilkesiana* extract.

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The susceptibility of the clinical fungi isolates used is shown in Table 2, which revealed that four of the fungi were completely resistant to all fractions of the extract. *A.niger* was susceptible to the ethyl acetate fraction while *C. albicans* was susceptible to the aqueous, ethyl acetate and chloroform fractions, with the plate treated with aqueous fraction producing the highest of zone of inhibition observed.

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Table 2: The mean± S.D (mm) of zone of inhibition observed on fungi seeded plates of isolates exposed to different fractions of *A.wilkisiana* extract

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Organisms	Crude	Aqueous	Ethyl acetate	Hexane	Chloroform
<i>Aspergillusniger</i>	0.00	0.00	6.5±0.29	0.00	0.00
<i>A.flavus</i>	0.00	0.00	0.00	0.00	0.00
<i>A. carbonarius</i>	0.00	0.00	0.00	0.00	0.00
<i>C. albicans</i>	0.00	7.5± 0.28	7.0 ±0.41	0.00	5.0± 0.00
<i>Trichophytonmetagrophytes</i>	0.00	0.00	0.00	0.00	0.00

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Antibiotics sensitive test

Antibiotics sensitivity test were carried out on all the test organisms. All test organisms expressed various resistant pattern as shown in table 3 for bacteria while, table 4 shows the resistant pattern of the fungi to Fluconazole used.

Table 3: Antibiotic Resistant Pattern of the Test Bacteria

Bacteria	Antibiotic Resistance
<i>Staphylococcus aureus</i> (ATCC 55620)	AMP, CHL, CXC, ERY, GEN, PEN, STR and TET
<i>S. aureus</i> (ATCC 29213)	Resistant to none
<i>S. aureus</i> (ATCC 25923)	AMP, CXC, ERY, GEN, PEN and STR
<i>Streptococcus pyogenes</i> (ATCC 8662)	AMP, CHL, CXC, ERY, GEN, PEN, STR and TET
<i>Enterococcus faecalis</i> (ATCC 29212)	AMP, CHL, CXC, ERY, PEN, STR and TET
<i>Pseudomonas aeruginosa</i> (ATCC 27853)	AMX, COT, NIT, GEN, NAL, AUG and TET
<i>Proteus vulgaris</i> (ATCC 13315)	AMX, COT, NIT, NAL and AUG
<i>Escherichia coli</i> (ATCC 35218)	AMX, COT, NAL, and AUG
<i>E. coli</i> (ATCC 23922)	AMX and AUG
<i>E. coli</i> (ATCC 25922)	AMX and AUG
<i>Klebsiella pneumonia</i> (ATCC 700603)	AMX, COT, NIT, NAL and AUG

Key:

OFL = Ofloxacin GEN = Gentamicin STR= Streptomycin

TET = Tetracyclin AUG = Augumentin

262 NIT = Nitrofurantoin AMX = Amoxicillin
 263 COT = Cotrimoxazole CHL = Chloramphenicol
 264 NAL = Nalidixic acid ERY = Erythromycin
 265 AMP= Ampicillin CXC= Cloxacillin
 266 GEN= Gentamicin PEN= Penicillin

267

268 Table 4: Antifungal Susceptibility pattern to Fluconazole

Fungi	Susceptibility pattern
<i>Aspergillusniger</i>	Resistant
<i>A.flavus</i>	Resistant
<i>A.carbonarius</i>	Resistant
<i>C.albicans</i>	Susceptible
<i>Trichophytonmetagrophytes</i>	Susceptible

269

270 Discussion

271 Many studies (Banso, A., Mann, A 2006; El-Mahmood, A.M., Ameh, J.M 2007; Falodun et al.,
 272 2006) have established the usefulness of medicinal plants as a great source for the isolation of
 273 active principles for drug formulation.

274 Several species of the genus *Acalypha* has been studied and it has been demonstrated that they
 275 present antioxidant, wound healing, post-coital antifertility, neutralization of venom,
 276 antibacterial, antifungal and antitrypanosomal activities (Perez Gutierrez, R.M., Vargas, S.R. 2006;
 277 Marwahet al., 2007; Shirwaikar et al., 2004). The result of this study support the antibacterial and
 278 antifungal activities of *A.wilkesiana* as a broad spectrum antimicrobial agent since it inhibited the
 279 growth of gram positive (*S. aureus*, *S. pyogenes*, *E. faecalis*) and gram negative bacteria (*E. coli*,
 280 *P. aeruginosa*, *P. vulgaris*) as well as some fungi (*A. niger*, *C. albicans*)

281 However the effectiveness of its antimicrobial potency seems to be more of antibacteria than
 282 antifungi. This study revealed that only *A.niger* and *C.albicans* were inhibited among the fungi
 283 used which support the work of Onocha and Olusanya 2010 which showed that the methanolic
 284 extracts of *A.wilkesiana* inhibited only *A.niger* and *C.albicans*. Also to the report of
 285 Oladunmoye 2006 which revealed that *A.niger* was inhibited by methanolic extracts. It is note
 286 worthy to see that *A.niger* which was resistant to the fluconazole was susceptible to the ethyl
 287 acetate fraction. The resistance of fungi to the tested extract may be due to the presence of more
 288 complex cell wall with rigidity than the thin cell membrane of bacteria. Also, this may be due to
 289 their ability to produce extracellular enzymes that helps them to degrade and metabolize
 290 substrate such that the extract becomes a source of food to the fungi instead of inhibiting their
 291 growth after they have been rendered non toxic due to degradation (Tortora et al., 2002).

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292 The result also showed that the extract and its fraction was potent against *S. aureus* (ATCC
293 29213) and *Streptococcus pyogenes* (ATCC 8662) which were resistant to standard antibiotics.
294 The disparity between the activities of the extract and the standard antimicrobial drug may be
295 due to the mixtures of bioactive compounds present in the extract compared to the pure
296 compound contained in the standard antibiotics (D.Gatsinget al.,2010) This demonstration of
297 activity against such test bacteria may form the scientific bases for the local dependent on this
298 plant in the treatment of various ailments.

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299 This present study also revealed that the ethylacetate fraction of the extract was the most potent
300 of all the fractions used. It is the only fraction which inhibited the highest number of bacteria and
301 fungi. The only exception to this is *E. faecalis* which characteristically was only susceptible to
302 the Chloroform fraction.

303 In conclusion, the search for new drugs to counter the challenge posed by resistant strains of
304 bacteria and some fungi might have started yielding results as the investigation of this plant has
305 demonstrated enormous therapeutic and preferential potential. They can serve the desired
306 purpose with lesser side effects that are often associated with synthetic antimicrobial agents.

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Comment [t125]: ?????????????

References

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