

Original Research Article

Antibacterial activity of bioflavonoid from fruit pulps of *Acacia nilotica* Willd ex Delile.

Abstract

Emergence The emergence of multi-drug resistance in bacteria has led to call for research and development of new leads as antibiotics from medicinal plants. *Acacia nilotica* (Linn) is a plant of multipurpose medicinal uses, three bioactive flavonoids (methyl gallate, gallic acid, and catechin) were isolated from its fruit pulps through a bioassay guided fractionation technique and characterized based on **High High-**Performance Liquid Chromatography, Liquid Chromatography-Mass Spectra and Nuclear Magnetic Resonance spectra. Antibacterial activity of these compounds was determined by microplate tetrazolium dye assay of broth microdilution technique against *Staphylococcus aureus*, *Pseudomonas aeruginosa*, *Escherichia coli*, and clinical isolates of *Salmonella typhi*, *Klebsiella pneumonia*, *Candida albicans*, and *Bacillus subtilis*. Catechin, methyl gallate, and gallic acid at 19.5, 39, and 39 µg/ml respectively caused a significant bio-reduction in cells of test organisms. Time kill kinetic study of the extract shows that there was a percentage of growth reduction in test organisms at 2, 4, 6, 8 and 12 hrs of contact. The extent and rate of killing of the organism by the extract at 2 x MIC followed the same trend as the rate of killing was time dependent. Antibacterial effects of these compounds are within the breakpoint of control drug chloramphenicol and could serve as leads in new drug development.

22 Keywords: *Acacia nilotica*, antibacterial, catechin, methyl gallate and gallic acid, bioactive
23 compound, high through put techniques.

24 **1.0 Introduction**

25 *Acacia nilotica* Willd. Ex Delile is commonly known as Gum Arabic tree, Egyptian
26 mimosa, Egyptian thorn, red thorn, Babool, babul in Indian [1]. It is recognized by the following
27 names: Acacia, Acacia Arabica, Babbul – Hindi and Nepalese, Babool Baum – German etc
28 [Steve, 2004]. It has naturalized in several countries where it has been introduced as a
29 medicinal, forage and fuel wood plant [2; 1]. It is proverbial, medium sized tree and is broadly
30 scattered in tropical and subtropical countries. It has an inspiring range of medicinal uses with
31 potential antioxidant activity. It contains a profile of a variety of bioactive components such as
32 gallic acid, (+)-catechin and methyl gallate. It also contains L-arabinose, catechol, galactan,
33 galactoaraban, galactose, N-acetyl djenkolic acid, sulphoxides and pentosan. The seeds contain
34 crude protein (18.6%), ether extract (4.4%), fiber (10.1%), nitrogen-free extract (61.2%), ash
35 (5.7%), and silica (0.44%); phosphorus (0.29%) and calcium (0.90%) of dry matter (17%) [3].
36 Phytochemical analysis of stem bark of *A. nilotica* showed the presence of carbohydrates,
37 saponins, tannins, and cardiac glycosides [4].

38 As a multipurpose medicinal plants, leaves, flowers, seeds, roots, fruits and gum act as
39 anti-tuberculosis [5], anti-hypertensive and anti-spasmodic, , antibacterial and antifungal [6],
40 antiplasmodial [7], and antioxidant [8, 9, 10]. Toxicologically, *A. nilotica*, at 2 and 8% levels,
41 has a low toxicity potential [11]. In a survey of potentially allergenic plants in Pondicherry, it
42 was reported likely to cause pollen allergy [12]. *A. nilotica* has a wealth of medicinal uses for
43 stomach upset and pain, the bark is chewed to protect against scurvy, an infusion is taken as
44 therapy for dysentery and diarrhea [13]. In Nigeria, it is one of the customary drugs for treating

45 diarrhea and it has been authenticated to have an antidiarrheal property [3] and antimicrobial
46 activity against neuro-pathogenic *E. coli* K1, MRSA and *Klebsiella pneumonia* [14].

47 This study describes the inhibitory effect of bioactive molecules of *A. nilotica* as leads for drug
48 development against pathogenic organisms using tetrazolium salts reduction as indicators of cell
49 inhibition.

50 **2.0 Methodology**

51 **2.1 Chemicals/Reagents/equipment:** Tetrazolium dye, Nutrient broth No 1, Muller Hinton
52 Agar from Fluka Analytical, Sigma-Aldrich Co., USA; Nutrient Agar from [HiMedia](#) [HiMedia](#)
53 Laboratories Pvt. Ltd., India; chloramphenicol disk ([HiMedia](#) [HiMedia](#) Laboratories Pvt. Ltd,
54 India), [UvUV/vis Vis](#) spectrophotometer (Jenway 6405, Britain), nuclear magnetic resonance
55 (300 MHz, Varian machine), HPLC₂ and LCMS.

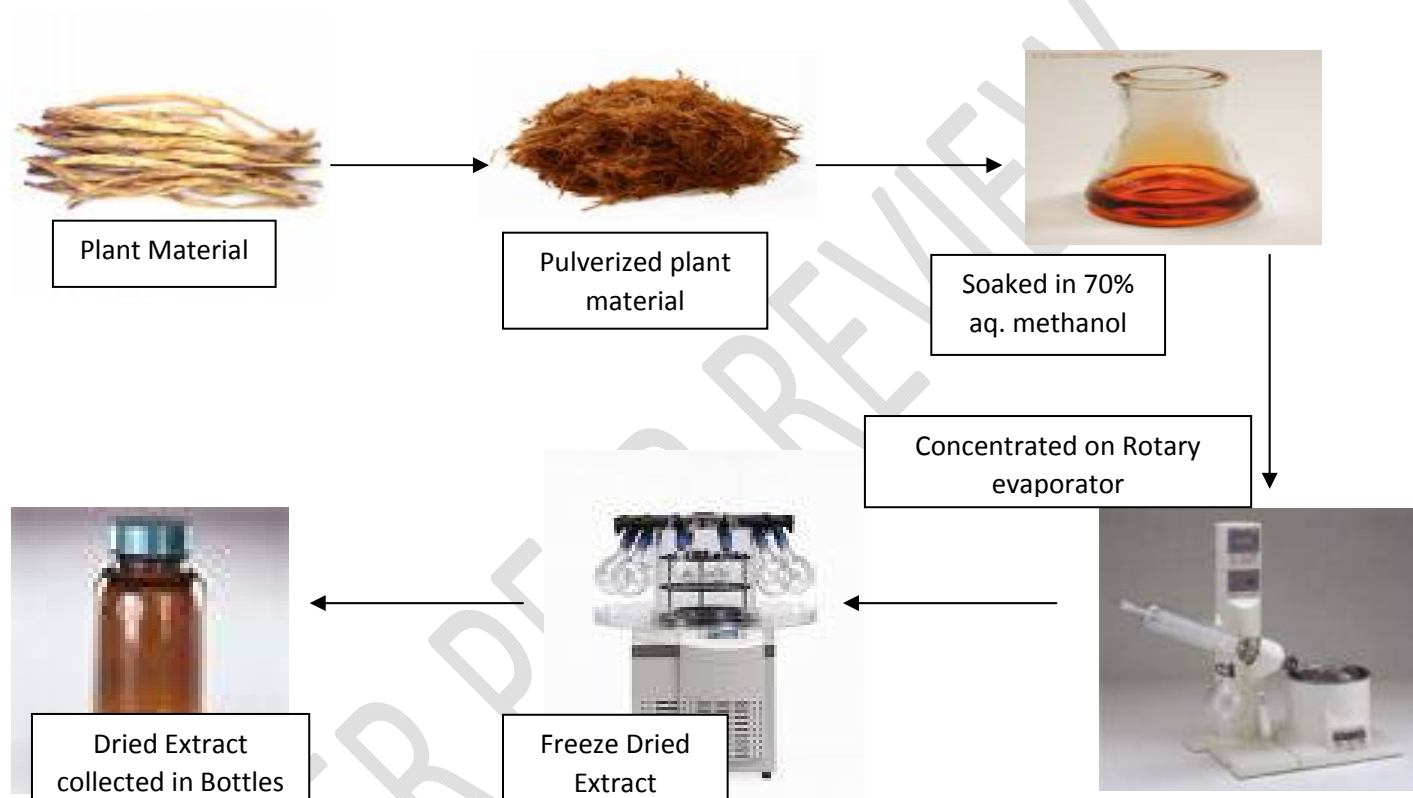
56 **2.2 Extract preparation and isolation:** The pulp of *A. nilotica* fruit was collected from Suleja,
57 Niger state, Nigeria, identified at Herbarium Unit of National Institute for Pharmaceutical
58 Research and Development, Abuja, Nigeria and herbarium specimen was deposited. This part
59 was air dried to constant weight at room temperature and pulverized. Extraction of both polar
60 and non-polar components of the plants using 70% aqueous methanol was carried out. One
61 hundred grams of dried plant part was soaked in 70% aqueous methanol (1000 ml) for 24 hrs.
62 The extract was filtered, concentrated on [a](#) rotary evaporator at 40°C and [freeze](#) [freeze](#)-dried. A
63 portion of the crude extract was used to determine the preliminary antibacterial effect of the
64 plant, while the other portion was subjected to [bioassay](#) [bioassay](#)-guided isolation of the
65 bioactive molecules (Figure 1).

66

67 Fractionation of the crude extract of *A. nilotica* (seeds) was carried out according to [15].
68 Activated silica gel (50 g) was packed by a wet method into a column. The extract (2.6g) which
69 was absorbed on silica gel (60 – 120 mesh) and dried was loaded on the column. Gradient
70 **elution elution** was performed with 100 ml of each mobile phase mixture in a series. The **elusion**
71 **elution** was performed in a flash chromatography. The mobile phase consisted of hexane, ethyl
72 acetate, methanol, and water, starting from 100% hexane and 10% increment in the next polar
73 component. The final **elusion elution** was performed with 70% methanol in water until the
74 column appeared exhausted with a sign of colourless silica gel. The eluates were monitored by
75 thin layer chromatography (TLC) using normal phase pre-coated silica gel K5 TLC plates. The
76 TLC mobile phase consists of a mixture of ethyl acetate: hexane (7:3). The eluates were
77 combined based on the similarity of TLC fingerprint to give six fractions labelled as FR1- FR6.
78 The pooled fractions (0.8 g) were absorbed on Merck – Kiesegel and introduced into a column
79 packed with the same absorbent. It was eluted with petroleum ether and an increasing gradient
80 of ethyl acetate. A total of 6 fractions of 100 ml each were collected. Fractions with **the** same
81 similarity (2-4) were pooled together into 3 portions viz; P1, P2, and P3 based on their TLC
82 profile (Figure 2). On staining with ferric chloride solution showed **blue blue**-black colouration
83 which is an indication of phenolic compounds. These fractions were purified further by HPLC
84 technique by separating on a reverse phase column, eluting with a gradient of acetonitrile 5% -
85 98% in water for a period of 40 minutes. Some 100 mg of extract was dissolved in 1 ml of
86 methanol, and water was added gradually to the point of precipitation before injecting on the
87 HPLC reversed phase (C-18) column. Fractions were collected at intervals of one minute. The
88 structural elucidation of the compounds was done using Nuclear Magnetic Resonance (300
89 MHZ, Varian machine) and LC-MS techniques. [2008]. The pure fractions from TLC analysis

90 were dissolved in deuterated methanol (CD₃OD) and transferred to NMR tube for measurement
91 of the proton (¹H) and ¹³C (carbon 13). Masses were acquired using liquid chromatography- Mass
92 Spec (LC-MS).

93



94

95 Figure.1. Extraction procedure of fruit pulps of *A. nilotica*

96

97 **2.3 Phytochemical analysis:** Phytochemical analysis of fruit pulp of *A. nilotica* was carried out
98 according to Trease and Evans [16]. Metabolites tested for include alkaloid, saponins, tannins,
99 anthraquinone, cardiac glycosides, and flavonoids.

100

101 **2.4 Antibacterial activity of the crude extract**

102 **2.4.1 Extract solution preparation:** The extract was dissolved in dimethylsulphoxide (DMSO)
103 and further diluted to 40 mg/ml concentration. The extract was sterile-filtered with 0.22 µm
104 syringe filter (Fisher brand). From this, a 2 mg/ml concentration of extract in agar was prepared
105 by dispensing 1ml of the extract solution (40 mg/ml) into 19 ml of molten Mueller Hinton Agar,
106 maintained in a water bath at 45°C. The molten agar/extract was poured into sterile Petri dish,
107 allowed to gel and dried to remove moisture. Overnight broth cultures of the test organisms viz;
108 *S. aureus* (ATCC 28923), *P. aeruginosa* (ATCC 27853), *E. coli* (ATCC 25922) and clinical
109 isolates of *S. typhi*, *K. pneumonia*, *C. albicans*, and *B. subtilis* were standardized by diluting to
110 10^7 cfuCFU/ml using a uv/vis-spectrophotometer as described by [17]. Two to three colonies of
111 20 hr growth on Mueller Hinton agar of the organisms to be studied were suspended on 50 ml
112 pre-warmed (37°C) Mueller Hinton broth. The suspension was incubated overnight at 37° C,
113 diluted 1/2500 in the same pre-warmed medium and incubated in a water bath with agitation (50
114 rpm). The optical density of the culture was monitored at 450 nm until absorbance of 0.1 was
115 reached (equivalent $2.5-3.0 \times 10^7$ cfuCFU/ml for *E. coli* and *P. aeruginosa*, $1.8-2.0 \times 10^7$ for
116 *S. aureus* and *B. subtilis*). The agar containing extract was inoculated by dropping 50 µl of
117 standardized organism on the surface of the agar, allowed to stand for 2 hrs. The plate was
118 incubated for 24 hrs at 37°C. The experiment was done in duplicate. Control plates containing 30
119 µg/ml concentration of chloramphenicol was used as drug control to confirm the susceptibility of
120 the organisms. Agar containing solution of the extract served as extract sterility control. Post
121 incubation, the presence or absence of growth of colonies of test organisms on agar plates
122 indicated positive or negative results of the crude extract.

123

124 **2.4.2 Minimum inhibitory concentration of molecules from *Acacia nilotica* (P1, P2, and P3)**

125 The minimum inhibitory concentration of molecules of *A. nilotica* was determined by broth
126 microdilution method according to British Society of Antimicrobial Susceptibility [18]
127 recommendation with a slight modification using tetrazolium colorimetric assay as
128 growth/inhibition indicator [19; 20]. Ten milligrams of each molecule was dissolved in Muller
129 Hinton broth. 100 µl of each of the solutions was transferred to the first column of micro well
130 plate, from where 50 µl was transferred to 2nd well already containing 50 µl Muller Hinton broth,
131 mixed thoroughly and the procedure was repeated through to well 11 where 50 µl was
132 discarded. All the wells were inoculated with 50 µl of overnight diluted cultures of each of *S.*
133 *aureus*, *Pseudomonas aeruginosa*, *E. coli*, *S. typhi*, *K. pneumonia*, *C. albicans*, and *B. subtilis*
134 prepared as described above. The plates were incubated for 24 hrs at 37°C. Post incubation the
135 plates were stained with tetrazolium dye; 3-(4, 5-dimethylthiazol-2-yl)-2, 5 diphenyltetrazolium
136 (MTT). MICs were defined as [the](#) lowest concentration of antimicrobial agents that red formazan
137 of MTT was not observed.

138 **2.4.3 Time kill Kinetic antibacterial study of fruit extract of *Acacia nilotica***

139 One hundred milligrams per ml concentration of the crude extract was prepared in sterile
140 water and diluted to 2 mg/ml concentration (being 2 X [mic](#) [MIC](#) of the crude extract) filtered by
141 centrifuging for 10 mins at 4500 rpm to remove woody sediments. The filtrate was inoculated
142 with 10⁵ [cfu](#)[CFU](#)/ml of *S. aureus* (ATCC 28923) and *E. coli* (ATCC 25922) and incubated at
143 37°C, percentage growth decrease was measured at 2 hrs interval. Exactly 0.5 ml of each
144 suspension was withdrawn at the appropriate time intervals and transferred to 4.5 mL nutrient
145 broth recovery medium containing 3% Tween 80 to neutralize the carry-over effects of the
146 antimicrobial compounds from the test suspensions. The suspension was shaken properly then
147 serially diluted in sterile physiological saline. Exactly 0.5 mL of the final dilution of the test

148 organism was transferred into pre-sterile Nutrient agar at 45°C and plated out. The plates were
149 allowed to set and incubated upside down at 37°C for 72 h. **Optical** The optical density of
150 growing culture in extract solution was measured using uvUV-spectrophotometer (Jenway 6405
151 uvUV/visVis, UK) at 520 nm to determine decrease in cells growth. **Control** The control
152 experiment which was set up without the inclusion of antimicrobial agent (i.e. excluding the
153 crude extract). Viable counts were made in triplicates for each sample. Depression in the viable
154 counts indicated killing by the antimicrobial agent.

155

156 **3.0 Results and Discussion**

157 Phytochemical analysis of fruit pulp of *A. nilotica* showed that the plant contains
158 alkaloid, saponins, tannins (phlobatannins), anthraquinone, and flavonoids. The presence of these
159 phytochemicals could be responsible for the observed activity. For instance, tannins
160 therapeutically have antiseptic properties and their precipitating quality is used in detecting
161 gelatin, proteins and alkaloids [21]. Flavonoids and phenolic compounds are plants metabolites
162 with at least one hydroxyl group [22] and are often found effective *in vitro* as antimicrobial
163 substance against a wide array of microorganisms [23]. The presence of these phytochemicals
164 could be responsible for the antibacterial activity recorded in this study.

165 **Bioassay** Bioassay-guided fractionation of the crude extract of fruit pulp of *A. nilotica*
166 yielded 3 distinct spots on TLC (Fig. 2). The compounds were analyzed on Liquid
167 Chromatography- Mass Spectrometer (LCMS) for purity and mass. Three compounds were
168 isolated from the fractions by the preparative. The pure compounds were fractions 5, 9, and 12.
169 **Fractions** The fractions in between were mixtures of these compounds as indicated by NMR
170 spectra.

171

172 The ^1H NMR of fractions 5 revealed it to be methyl gallate, 9 to be gallic acid and 12 to be
173 catechin through a library search [24, 15].



174

175 **Figure 2. Thin Layer Chromatography plate of active fraction of *A. nilotica* (showing 3**
176 **distinct spots P1, P2, and P3).**

177

178 **3.1 Characterization of Isolated Compounds:**

179 The first compound P1 (fraction 5) was a white solid with a melting point of 202-204 °C

180 (Lit. 201-204 °C). The ^1H NMR revealed the presence of aromatic protons at δ 7.2, which

181 integrated for two protons at positions 2 and 6 and that of methoxyl at δ 3.3 which integrated for

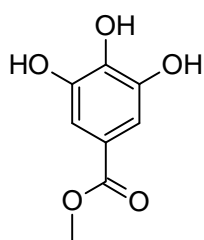
182 3 protons at position 8. This indicated a simple aromatic system. The broad band was decoupled

183 as; ^1H NMR: 7.20 (2H, H-2 and H-6), 5.0 (OH), 3.3 (3H, CH₃), ^{13}C NMR: 167.9 (C-7), 146.6

184 (C-3), 146.6 (C-5), 139.4 (C-4), 122.2 (C-1), 110.3 (C-2), 110.3 (C-6), 52.5 (OMe). ^{13}C NMR

185 revealed the presence of 8 carbons at δ (167.9 for one C=O, 110.3 for two CH, 52.5 for OCH₃,
 186 and four quaternary aromatic carbons at 122.2, 138.4, 146.6, 146.6) (Table 1). HRMS of this
 187 fraction gave 184.03717 amu, which was consistent with molecular formula C₈H₈O₅. A library
 188 search revealed it to be methyl gallate (methyl 3, 4, 5-trihydroxybenzoate) [15] and [24]. The
 189 sample was also compared with an authentic sample by TLC and was confirmed to be methyl
 190 gallate: Methyl 3, 4, 5-trihydroxybenzoate (Fig 3). Molecular formula; C₈H₈O₅

191



192 Methyl gallate Figure 3. Chemical structure of methyl gallate with [the](#) molecular formula
 193 C₈H₈O₅

194

195

196 **Table 1. Carbon 13 and Proton NMR spectra of methyl gallate (P1) and gallic acid (P2)**

Position	Group	Compound P1		Compound P2	
		¹³ C Chemical Shift (δ _C)	Proton Chemical Shift (δ _H)	¹³ C Chemical Shift (δ _C)	Proton Chemical Shift (δ _H)
1	C	122.2	-	122.2	-
2	CH	110.3	7.2	110.3	7.0
3	C-OH	146.6	6.8	146.6	4.789
4	C-OH	139.4	5.0	140.0	4.789
5	C-OH	146.6	6.8	146.6	4.789
6	CH	110.3	7.2	110.3	7.0
7	C=O	167.9	-		
8	OCH ₃	52.5	3.30		

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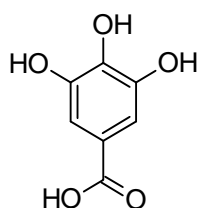
199 The second compound (P2) was a solid, light brownish in colour, m. p. = 237-239°C. ¹H NMR:

200 7.0 (2H, H-2 and H-6), 4.789 (OH), ¹³C NMR: 170 (C-7), 146.6 (C-3), 146.6 (C-5), 139.4 (C-4),

201 122.2 (C-1), 110.3 (C-2), 110.3 (C-6). The ¹H NMR of this compound revealed the presence of

202 aromatic protons at δ 7.0, which integrated for two protons at positions 2 and 6 and the OH
 203 proton at position 7 at δ 3.212. The broad band decoupled (Fig 4.) ^{13}C NMR revealed a total of 7
 204 carbons at δ (170 for one C=O, 110.3 for two CH, and 4 quaternary aromatic carbons at 122.2,
 205 138.4, 146.6, [and](#) 146.6). These were almost identical with that of methyl gallate except for the
 206 absence of the OCH_3 ^{13}C NMR peak. As a result, this compound was subsequently identified as
 207 gallic acid [15]. (Fig 4).

208
 209



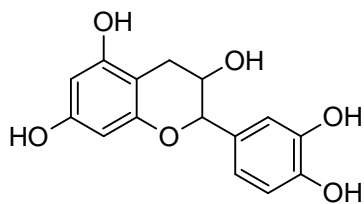
210 Gallic acid
 211 | Fig. 4 Chemical structure of gallic acid with [the](#) molecular formula $\text{C}_7\text{H}_6\text{O}_5$
 212

213 **Table 2. Carbon 13 and Proton NMR spectra of Catechin (P3)**

Position	Carbon 13 Chemical Shift (δ_{C})	Proton Chemical Shift (δ_{H})
1	-	
2	80.9	4.56
3	66.3	4.00
4	27.7	5.58
5	156.4	
6	93.9	
7		
8	95.1	
9		
1'		
2'	114	6.89
3'		
4'		
5'	115.1	6.79
6'	18.4	18.4

214
 215
 216 ^1H -NMR spectra of the third compound (P3) showed peak at δ 4.56 (H-2, d), 4.01 (H-3, ddd),
 217 2.54 (H-4, dd), 2.90 (H-4, dd), 5.87 (H-6, d), 6.01 (H-8, d) 6.89 (H-2', d), 6.79 (H-5', d) , 6.73

218 (H-6', dd) and 8.00 (phenolic protons, m). ¹³C-NMR, Carbon atoms showed peaks at δTMS 80.9
219 (C-2), 66.3 (C-3), 27.7 (C-4), 93.9 (C-6), 95.1 (C-8), 114.5 (C-2'), 115.1 (C-5'), 18.4 (C-6')
220 (Table 2). The NMR chemical shifts correlate well with those available in [the](#) literature for
221 catechin [25; 26]. The compound was identified as catechin (Fig 5).



224 Catechin

225 **Fig 5. Chemical structure of catechin with [the](#) molecular formula C₁₅H₁₄O₆**

227 3.2 Antibacterial activity

229 Antibacterial activity of the crude fractions and compounds from fruit pulps of *A. nilotica*
230 are as shown in [table Table 3](#). The result shows that the crude extract was significantly active
231 against all the test organisms but at varying degree of concentration. **Minimum inhibitory**
232 **concentration** MIC against *B. subtilis* and *S. aureus* were 500 and 750 ± 0.0 µg/ml respectively
233 while the susceptibility of **Gram Gram-negative** organisms (*E. coli*, *P. aeruginosa*, *S. typhi*, [and](#)
234 *K. pneumonia*) and *C. albicans* were slightly higher at 1000 – 2000 µg/ml respectively). These
235 concentrations are higher than the **mic MIC** breakpoint for chloramphenicol (30 ± 0.1 µg/ml) the
236 control drug. Hence, the need for further purification of the extract to compound level for better
237 activity. The fractions (F1-F6) produced by column chromatography with TLC finger printing
238 yielded better minimum inhibitory concentrations (MICs) at 78 – 156 ± 0.0 µg/ml, respectively.
239 Fraction F1 had no antibacterial effect, fraction F5 had **mic MIC** of 156 µg/ml against all the test

240 organisms while F6 was only active against *S. aureus* and *B. subtilis* at 156 µg/ml. Fractions F2-
241 F4 produced the best antibacterial effects at 78 µg/ml against **Gram Gram**-positive organisms
242 and 156 µg/ml against **Gram Gram**-negative organisms. These fractions have a lot in common;
243 TLC profile, positive test for phenolic and antibacterial effects. Microplate tetrazolium assay of
244 isolated molecules against the pathogenic organisms as shown in **table Table** 1, showed that
245 methyl gallate, gallic acid at 39 µg/ml, and catechin at 19.5 µg/ml concentration are significantly
246 ($p < 0.05$) more effective against *E. coli*, *S. aureus*, and *B. subtilis* than against *P. aeruginosa*, *S.*
247 *typhi*, and *K. pneumonia* (78 µg/ml). Catechin has shown to be the most effective of the
248 molecules with MIC of 19 ± 0.5 µg/ml against *S. aureus* and *B. subtilis* and 39 µg/ml against other
249 **Gram Gram**-negative organisms.

250 This result shows that the purer the fractions the better the antibacterial activity. The use
251 of microplate tetrazolium assays to measure cell proliferation has increased exponentially since
252 their introduction [19]. Nevertheless, these assays do not actually measure the number of viable
253 cells in a culture or their growth but rather, an integrated set of enzyme activities that are related
254 in various ways to cell metabolism. They utilize the cofactor nicotinamide adenine dinucleotide
255 (NADH) and the dye 3-(4,5-dimethylthiazol-2-yl)-2,5 diphenyltetrazolium (MTT), and other
256 substrates like succinate and pyruvate which may also contribute to their reduction. Depending
257 on the particular dye chosen, reduction will be linked in various ways to cofactor/substrate
258 production, utilization, and compartmentalization, and can be associated with the plasma
259 membrane, intracellular membranes, organelles, and cytosol. **Reduction** The reduction can vary
260 widely within and between cell populations depending on the cell growth conditions, phase of
261 cell growth and stage of the cell cycle [20].

262 In the case of antibacterial activity of methyl gallate, the result is in agreement with a
 263 similar study [27] that methyl gallate from the leaves of *Toona sureni* had antibacterial activity
 264 against *E. coli*, *S. aureus*, and *B. subtilis*. Other similar studies have reported that gallic acid by
 265 mechanism of action in *P. aeruginosa*, *E. coli*, *S. aureus*, and *Listeria monocytogens* led to
 266 irreversible changes in membrane properties (charge, intra, and extracellular permeability, and
 267 physicochemical properties) through hydrophobicity changes, decrease of negative surface
 268 charge, and occurrence of local rupture or pore formation in the cell membranes with consequent
 269 leakage of essential intracellular constituents [28]. Antibacterial activity of gallic acid of *A.*
 270 *nilotica* in this study corroborates the report that gallic acid inhibited the growth of
 271 *Campylobacter jejuni* and *E. coli* strains at 15.63-250 µg/ml. [12].

272 Prolonged incubation of the microplates was used to determine the end point assay of the
 273 molecules as bacteriocidal. Wells with bacteriostatic effect at concentrations lower than 78 µg
 274 /ml showed tetrazolium reduction after 48 hours of incubation while bacteriocidal concentrations
 275 of 78 µg /ml upward remained colourless. This assay has been used to distinguish between
 276 dormant and metabolically active microbial cells (Tab. 3). The test organisms were able to
 277 reduce tetrazolium dyes in their electron transport chain, generating results within hours.

Molecules µg/ml	MICROORGANISMS						
	<i>S. aureus</i>	<i>P. aeruginosa</i>	<i>E. coli</i>	<i>B. subtilis</i>	<i>S. typhi</i>	<i>K. pneumonia</i>	<i>Ca</i>
Crude extract	750 ± 0.0	1000 ± 0.0	1000 ± 0.0	500 ± 0.0	2000 ± 0.0	2000 ± 0.0	1000 ± 0.0
F1	NA	NA	NA	NA	NA	NA	NA
F2	78	156	156	78	156	156	156
F3	78	156	156	78	156	156	156
F4	78	156	156	78	156	156	156

F5	156	156	156	156	156	156	156
F6	156	NA	NA	156	NA	NA	156
Methyl gallate	39 ± 0.0	*78 ± 0.0	78 ± 0.0	39 ± 0.0	78	78	39
Gallic acid	39 ± 1.0	78 ± 0.0	78 ± 0.0	39 ± 0.0	78±0.0	78±0.0	39±0.0
Catechin	19.5 ± 0.0	39 ± 0.0	39 ± 0.0	19.5 ± 0.0	39±0	39±0.0	39±0.0
Chloramphenicol	30± 0.0	30 ± 0.00	30 ± 0.0	30±0.0	30±0.0	30±0.0	

278

279 **Table 3. Antibacterial activity of crude, fractions and compounds of fruit pulp of *A. nilotica***

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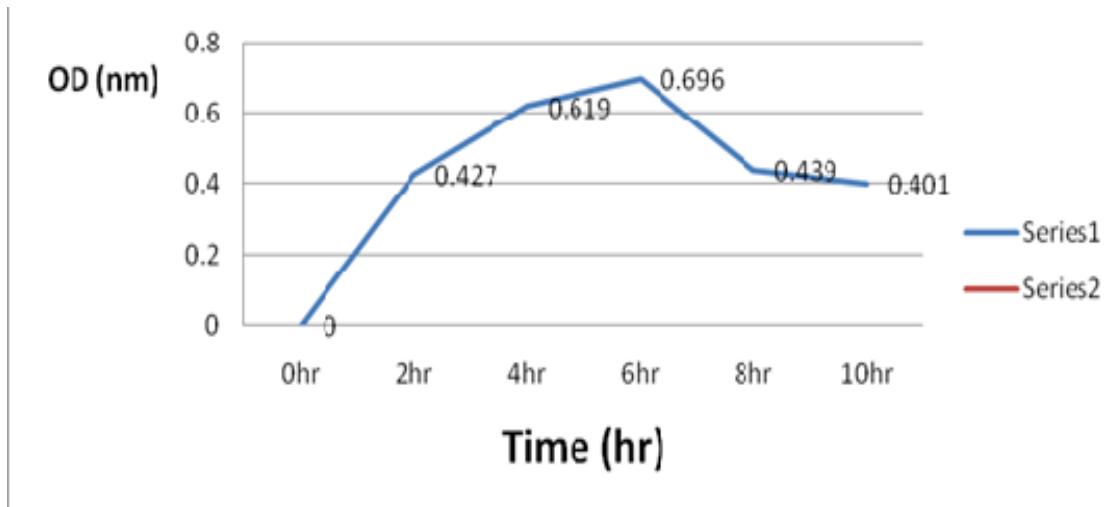
281 **Key:** F= Fractions, NA = No activity, no growth inhibition

282

283 **3.3 Time kill kinetic antibacterial of crude extract**

284 The time kill kinetic antibacterial of [the](#) crude extract against *S. aureus* and *E. coli* are as
285 shown in Figs. 6 and 7. The percentage of growth reduction in *S. aureus* at 2, 4, 6, 8, and 12 hrs
286 of contact were 52.2, 60.8, 70, 80, and 99.2%, respectively. Percentages of *E.coli* growth
287 reduction for each corresponding time were; 45.7, 51.7, 64.16 and 99.9% respectively. The
288 extent and rate of killing of the organism by the extract at 2 x MIC followed the same trend as
289 the concentrations of the extract increased with increase in contact time, the percentage of the
290 organisms killed also increased.

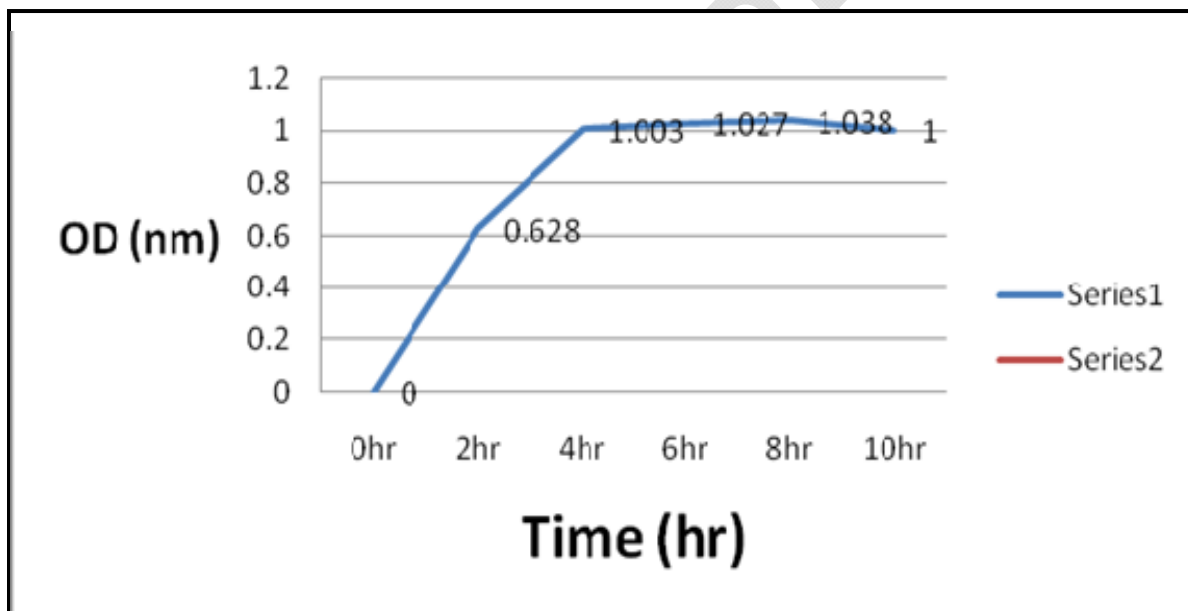




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298 **Fig. 6. Effect of extract of *A. nilotica* on the growth of *S. aureus* at different time**

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300

301 **Fig 7. Effect of extract of *A. nilotica* on the growth of *E. coli* at different time**

302

303

304

305 **4.0 Conclusion**

306 In conclusion, this study has demonstrated the antibacterial properties of bioflavonoids
307 from fruit pulps of *A. nilotica*. The compounds isolated could serve as leads in the search for new
308 and potent antibacterial compounds from indigenous medicinal plants.

309

310

311 **Conflict of interest:** Authors have declared that no competing interests exist

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