

1 **PRELIMINARY PHYTOCHEMICAL ANALYSIS AND *IN VITRO* ANTIMICROBIAL**
2 **STUDY OF THE ROOT AND STEM BARK EXTRACTS OF *FICUS SYCOMORUS LINN***

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9
10 **Abstract**

11 This study was conducted to carryout preliminary phytochemical analysis and *in vitro*
12 antimicrobial activities of aqueous and ethanolic root and stem bark extracts of *Ficus sycomorus*.
13 Qualitative phytochemical analysis for tannins, saponin, terpenoids, flavonoids, alkaloids,
14 glycosides, steroids, phenols, and reducing sugar was done using standard **methods**. The
15 antimicrobial activities of the extracts were tested against four micro- organisms; *Escherichia*
16 *coli*, *Staphylococcus aureus*, *Shigella dysenterae*, and *Salmonella typhi*. Agar well diffusion
17 method was used for the antimicrobial studies. Phytochemical screening of both root and stem
18 bark aqueous extracts showed the presence of tannin, saponin, terpenoid, flavonoid , alkaloids,
19 glycoside, steroid, reducing sugar, and phenol. Glycoside was not detected in both the aqueous
20 and ethanolic extracts of the root bark. The result of the antimicrobial studies showed that the
21 aqueous root extract have higher antimicrobial activity ranging from (2-12 mm) on the tested
22 microorganisms than aqueous stem bark extract (3-9 mm), while for ethanol extract both stem
23 and root bark extract has almost the same effect or antimicrobial activity on the tested pathogens
24 ranging from (2-15 mm) which is having higher activity compared to the aqueous extracts. The
25 Minimum inhibitory concentration (MIC) and Minimum bactericidal concentration (MBC) of
26 both the extracts were found to be 50 mg/mL and 100 mg/mL respectively. From this study, it
27 can therefore be concluded that, the root and stem bark extract is a potential antimicrobial agents
28 which support the claim of the traditional users of this plant in herbal medicine for the treatment
29 of diseases that are of microbial origin.

30 **Key words:** *Ficus sycomorus*, *Escherichia coli*, *Staphylococcus aureus*, *Shigella dysenterae*,
31 and *Salmonella typhi*, phytochemical **screening, antimicrobial** activity.

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34 INTRODUCTION

35 Medicinal plants besides therapeutic agents are also a big source of information due to a
36 variety of chemical constituents which could be developed as drugs with precise selectivity.
37 They are reservoirs of potentially useful chemical compounds which could serve as newer leads
38 and clues for modern drug design [1]. Among the most important of these bioactive constituents
39 of plants are alkaloids, tannins, flavonoids and phenolic compounds [2]. Correlation between
40 the phytochemical constituents and the bioactivity of plant is desirable to know for the synthesis
41 of compounds with specific activities to treat various health ailments and chronic diseases as
42 well [3].

43 In the developing countries, the use of herbal medicine is drawing the attention of
44 researchers as a result of resistance posed by microbes to synthetic drugs. These synthetic drugs
45 are mostly expensive and with so many adverse side effects. Due to their unlimited therapeutic
46 benefits, the support for the use of medicinal plants by the World Health Organization (WHO) is
47 quite encouraging [4].

48 Micro-organisms, affect man's life in many ways, from the day of his birth to the day of
49 his death. Micro-organisms are always ready to help us or destroy us and only circumstances
50 decide which it shall be. Therefore, micro-organisms are often classified as useful ones or
51 harmful ones. The latter organisms are the causes of numerous infectious diseases which are
52 great enemies of man and his flocks and crops.

53 Antimicrobial activity is a property of a wide variety of compounds. This activity may be
54 bactericidals, fungicidals or virucidals which is concerned with the killing of bacteria, fungi or
55 viruses respectively. On the other hand, the activity may be only growth inhibiting i.e.
56 bacteriostatic or fungistatic. However, this classification is not sharp since a bacteriostatic agent
57 may only inhibit the organism if it is used in low concentration, or when the exposure time is
58 limited. Therefore, a substance is usually considered germicidal when its effective concentration
59 range is low and its killing rate is rapid. Only antimicrobial substances with a selective action on
60 the parasite would be suitable. The ideal therapeutic agent would be entirely selective, having no
61 action whatsoever on the hosts tissues.

62 In recent years, an intensive effort has been made to find new antimicrobial agents. The
63 major part of the reported investigations was concerned with lower plants, with special attention
64 being paid to different species of streptomycetes and some fungi. A total of 428 extracts of plants
65 from 43 families, encompassing 100 species and selected on the basis of literature data and
66 medicinal folkloric reports were evaluated for antimicrobial, antiviral, antiparasitic and
67 pharmacological activities.

68 In Nigeria like many African countries, several plants are still being used for the treatment
69 of various ailments. Nigeria is naturally blessed with both savannah and tropical rainforests
70 vegetation and these offer a wide distribution of plants believed to possess secondary
71 metabolites which are responsible for treating or curing various diseases [4]. Quite a number of
72 plants are used as medicines virtually in all cultures of the world. A good number of these
73 medicinal plants are in common use in African traditional medicine. Most of the plants grow
74 near houses and are easily overlooked, especially by urban dwellers [5].

75 This research work was carried out to study the Phytochemical screening and *in vitro* anti-
76 microbial activities of root and stem bark extract of *Ficus sycomorus* on some selected micro
77 organisms.. *Ficus sycomorus* is a common savannah tree that grows or can be found almost
78 everywhere. It is called in English Language as “Wild fig” “sycamore fig”, or common cluster
79 fig. Spanish call it “sicomoro”. The Sukur people call it “Dashakwai” , Tiv people called it
80 “Tur”, in Hausa it is known as “Baure”, Kilba and Marghi people called it “Kamda” , in Fali
81 Language is called “Boduven” and Gude call it “Bodeva”. It grows in high water table areas, it
82 can be found along water courses such as streams, rocky places, swamps and water holes [6].
83 The sycamore fig is sensitive to frost but can withstand some cold. The relevance of this plant in
84 traditional medicine is as a result of the secondary metabolites such as glycosides, reducing
85 sugar, phenols, saponins, steroids, tannins, alkaloids, terpenoids and flavonoids which they have
86 been screened to contain. Also referred to as phytochemicals, they are reported to possess
87 inhibitory activities against the growth and disease inducing activities of some pathogenic
88 microorganisms [7, 8, 9, 10, 11].

89 The root and stem-bark of *Ficus sycomorus* are said to be used as herb in Northern
90 Nigeria for treatment of diseases like diarrhea, dysentery, cough, sore throat, chest diseases, and
91 infertility and as antidote for snake. Therefore, this study was conducted to carry out the

92 phytochemical screening and to evaluate antimicrobial activity of root and stem-bark of *Ficus*
93 *sycomorus* in order to validate the claims of the traditional users of this plant.

94

95 **MATERIALS AND METHODS**

96 **Sample Collection and identification of plant material.**

97 Plant roots and stem-barks of the plant *Ficus Sycomorus* were collected from Sukur
98 Kingdom in Madagali Local Government Area, Adamawa State, Nigeria. It was identified and
99 authenticated by a Botanist from the Department of Biological Sciences, Adamawa State
100 University, Mubi. A voucher number AD170023 was assigned. Sampling was carried out from
101 March to July, 2016.

102 **Sample preparation**

103 The root and Stem-barks (cut into small pieces) washed with water and rinsed with
104 distilled water and then dried in the shade for two weeks. The dried samples was grinded by
105 wooden mortar and pestle and sieve using clean Kitchen sieve to obtain a fine powder and was
106 stored in a tight container until required for use.

107 **Extraction**

108 **Aqueous Extract**

109 For the water extraction was done by cold maceration method according to the procedure
110 described by [12, 13] with little modification. Two hundred grams (200 g) of each of the stem
111 and root barks powder was weighed and soaked in 1000 mL of distilled water in a beaker for 48
112 h to obtain aqueous extracts. The aqueous extracts were filtered using sterile filter paper
113 (Whatman No.1) into a clean conical flask. The filtrate was concentrated with a rotary
114 evaporator. The extracts were then stored in a refrigerator.

115 Percentage yield was calculated as: $\text{weight of extract} / \text{weight of dried powdered sample} \times 100$

116 **Preparation of ethanol extracts**

117 Maceration method of extraction as described by [12, 13] was adopted in this study. Two
118 hundred grams (200g) each of the root and stem bark powdered material was weighed and
119 soaked in 1000 mL of 70% ethanol and left for 24 h .Thereafter, it was decanted. The procedure
120 was repeated with another 1000 mL to ensure complete extraction of the active ingredient .The

121 extract was filtered and evaporated to dryness with rotary evaporator. The dried extract was then
122 weighed and stored in tightly closed bottle in a refrigerator until required.

123 Percentage yield was calculated as: $\text{weight of extract} / \text{weight of dried powdered sample} \times 100$

124

125 **Qualitative Phytochemical analysis.**

126 The qualitative phytochemical screening of the samples was carried out as described by [14, 15,
127 16] with slight modification. The root or stem bark extracts were screened for carbohydrates,
128 alkaloids, flavonoids, steroids, phenols and tannins, saponin, glycosides, and proteins.

129

130 **Preparation of stock solution**

131 Two grams (2g) each of root or stem bark extracts were dissolved in 10 mL of water or ethanol
132 to make a concentration of 200 mg/mL

133 **Test for Tannins**

134 One milliliter (1 mL) of the extracts was taken in a test tube and 2 mL of 5 % ferric chloride was
135 added. Formation of blue –black, green or blue – green precipitate was taken as evidence for the
136 presence of tannins.

137 **Test for Saponins**

138 One milliliter (1 mL) of the extracts was shaken with 5 mL of distilled water in a test tube for 5
139 min. Frothing which persists on warming was taken as evidence for the presence of Saponins.

140 **Test for Terpenoids**

141 Five milliliters (5mL) of aqueous extract of each plant sample was mixed with 2mL of CHCl_3 in
142 a test tube and then 3mL of concentrated H_2SO_4 was carefully added to the mixture to form a
143 layer. An interface with a reddish brown coloration was considered as indication for the presence
144 of terpenoids.

145 **Test for Flavonoids**

146 A little amount of magnesium powder and a few drops of concentrated hydrochloric acid were
147 added to 3 mL of the extracts. A red or intense coloration indicated the presence of flavonoids.

148 **Test for Alkaloids**

149 To 2 mL of plant extracts, 2 mL of concentrated hydrochloric acid was added. The mixture was
150 filtered and then 3 drops of Mayer's reagent was added. Presence of green colour or white
151 precipitate indicated the presence of alkaloids.

152 **Test for glycosides**

153 Two milliliter (2 mL) of the extracts was hydrolyzed with HCl solution and neutralized with
154 NaOH solution. A few drops of Fehling's solution A and B were added. Presence of red
155 precipitate indicates the presence of glycosides.

156

157 **Test for Steroids (Salkowski's test)**

158 To 1 mL of plant extract, equal volume of chloroform and 3 drops of concentrated sulphuric acid
159 was carefully added to form a lower layer. Formation of brown ring indicates the presence of
160 steroids.

161 **Test for phenols**

162 Five drops of 10% ferric chloride was added to 1 mL of the extracts in a test tube. Formation of
163 green or dirty green precipitate indicated the presence of phenols.

164 **Test for reducing Sugar**

165 To 2 mL of plant extract, 1 mL of Molisch reagent and 4 drops of concentrated sulphuric acid
166 was added. Formation of purple or reddish ring indicates the presence of carbohydrates.

167

168 **Antimicrobial Analysis**

169 *Escherichia coli*, *Shigella dysenteriae*, *Salmonella typhi* and *Staphylococcus aureus* were used in
170 this study. The microorganisms were obtained at the Microbiology Laboratory of Modibbo
171 Adama University of Technology, (MAUTECH) Yola, Nigeria.

172 **Standardization of Isolates:**

173
174 Test organisms were sub-cultured onto fresh plates of MacConkey agar and incubated
175 aerobically at 37°C for 24 h. Colonies from these plates were suspended in Mueller- Hinton
176 broth to a turbidity matching 0.5 McFarland standard (108cfu/ml). Mueller-Hinton agar was then
177 used for antimicrobial assay. All the broth cultures were incubated at 37°C.

178

179 **Preparation of the Extract for Antimicrobial Study**

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181 Two grams (2g) each of aqueous and ethanol root or stem bark extracts were separately
182 dissolved in 10 mL of dimethylsulfoxide (DMSO) to obtain a concentration of 200mg/mL.
183 This was the initial concentration of each of the extracts used.

184

185 **Antimicrobial Test:**

186

187 The method described by the National committee for Clinical Laboratory Standard [17] was
188 used.

189 Suspensions of the bacteria obtained contained approximately 1×10^8 cfu/mL. Each labeled
190 plate was uniformly seeded with a test organism by means of sterile swab stick rolled in the
191 culture medium. Five wells, 4mm each in diameter were created using cork borer. Aliquots were
192 dropped in each well to fullness at various concentrations of 100, 50, 25 and 12.5 mg/mL for
193 both the root and stem bark extracts on different plates. Each plate was kept in the refrigerator
194 for 1 hour to allow the extracts to diffuse into the culture medium while the immediate growth of
195 the organism was stopped from taking place. These plates were then incubated at 37°C for 24 h.
196 The zones of inhibition around the wells were measured in millimeter (mm). Control antibiotic
197 (tetracycline capsule 100 µg/mL) was placed in a well on each plate along with the test extracts
198 as control.

199 **Determination of the Minimum Inhibitory Concentration (MIC)**

200 The minimum inhibitory concentration of the extract was evaluated by the method described by
201 [18].

202 The extract concentration were serially diluted with distilled water to various concentrations of
203 100, 50, 25and 12.5mg/mL. The extract and the nutrient agar broth were mixed in the sterile test
204 tube; the cultured medium was added to each test tube and incubated for 24hrs at 37°C .The
205 lowest zones of inhibition for all the tested organisms showing no visible growth of **bacteria** was
206 taken as the MIC.

207 **Minimum Bactericidal Concentration (MBC).**

208 The minimum bactericidal concentration (MBC) was determined after the minimum
209 inhibitory concentration (MIC) was obtained. This was **carried out** by selecting the test tube that
210 shows no growth during the MIC determination. A loopful from the test tube containing the
211 media and the extract were inoculated into a sterile nutrient broth media. This was further
212 incubated for another 24-48 hrs at 37°C for bacteria, after which was examined for bacteria for

213 any microbial growth. The lowest concentration at which no growth was observed on the plate
214 was taken as the MBC [18].

215 **RESULTS AND DISCUSSION**

216 This study was undertaken to investigate the antimicrobial activity and phytochemical
217 screening the aqueous and ethanolic root and stem bark extracts of *Ficus sycomorus* Linn. Due
218 to the side effects of the current drugs and the resistance that pathogenic microorganisms build
219 against antibiotics, much attention has led to the study of biologically active compounds isolated
220 from plant species used in herbal medicine [19]. Different scientific studies provided evidence
221 that medicinal plants might indeed be potential sources of new antibacterial agents even against
222 some antibiotic-resistant strains [20].

223 The yield of the plant extracts is presented in Table 2. It was observed that Ethanol stem bark
224 extract (ESB) gave the highest yield 16.00g (8.0%) followed by Ethanol root bark extract (ERB)
225 14.14 g (7.07 %) then Aqueous root bark extract (ARB) 12.23 g (6.12%) and the lowest is Aqueous
226 stem bark extract (ASB) 11.16 g (5.58 %). From the result it is generally observed that the solvent,
227 ethanol gave higher yield irrespective of the plant part than the aqueous solvent.

228 The result of this study shows the presence of phytochemicals considered as active
229 medicinal chemical constituents as shown in table 2. Phytochemicals such as tannins, saponin,
230 terpenoids, flavonoids, alkaloids, glycosides, steroids, phenols and reducing sugars were all
231 found to be present in both the ethanol extracts of roots and stem bark of *Ficus sycomorus*.
232 However, glycosides was the only constituent not detected in Aqueous extracts of the root and
233 stem bark. The result is contrary to the findings of [21] who reported the presence glycoside in
234 the methanolic stem bark extract of *Ficus sycomorus* obtained from Zaria city of Kaduna State,
235 Nigeria. The absence of some of these constituents that have been reported in the previous
236 studies and are reported to be present in this study may be due to geographical location which
237 has been reported to affect the chemical constituents of plant extracts of the same genus found in
238 different environments and also differences is polarity of the solvents used for extraction . This
239 could therefore be the reason why glycoside was not detected in the aqueous root and stem bark
240 extract of *Ficus sycomorus* in this present work. Similar report has also been documented [22],
241 where they reported that phytochemical screening of methanolic stem bark extract showed the

242 presence of tannins, saponins, terpenoids, flavonoids, phenols, steroids, except glycosides and
243 proteins.

244 The various phytochemical compounds detected are known to have beneficial importance in
245 industrial and medicinal sciences. These secondary metabolites exert antimicrobial activity
246 through different mechanisms. Plant phenolic compounds especially flavonoids are currently of
247 growing interest owing to their supposed properties in promoting health (anti-oxidants) [23].
248 Flavonoids have been demonstrated to have antiinflammatory, antiallergenic, anti-viral, anti-
249 aging, and anti-carcinogenic activity. The broad therapeutic effects of flavonoids can be largely
250 attributed to their antioxidant properties. In addition to an antioxidant effect, flavonoid
251 compounds may exert protection against heart disease through the inhibition of cyclooxygenase
252 and lipoxygenase activities in platelets and macrophages[24].Tannins are reported to possess
253 physiological astringent and haemostatic properties, which hasten wound healing and ameliorate
254 inflamed mucus membrane and also inhibit the growth of microorganisms by precipitating
255 microbial proteins and making nutritional proteins unavailable for them; they form irreversible
256 complexes with proline rich proteins, resulting in the inhibition of the cell protein synthesis.
257 They have important roles such as stable and potent antioxidants [24, 25]. They act as binders
258 and for treatment of diarrhea and dysentery [26] Tannins also reported to exhibit antiviral,
259 antibacterial, anti-tumor activities. It was also reported that certain tannins are able to inhibit
260 HIV replication selectivity and is also used as diuretic [24].

261 The results of the zones of inhibition of the different extracts (ARB, ASB, ESB and ERB)
262 against the tested pathogens are exhibited in Tables 3 – 6. It showed that the extracts have dose
263 dependent antimicrobial activities against the pathogens at various concentrations used in this
264 study. It was noticed that the extract was more effective at concentration of 100 mg/mL, but the
265 effectiveness increases as the concentration increases. The highest activity was shown by the
266 ESB and ERB at 100 mg/mL (15mm) against *E. coli*. Although most of the extracts at the
267 various concentrations used showed activity against the pathogens, it was observed on the
268 general that the extracts are more effective at 100 mg/mL on *E. coli*, which showed similar
269 activity with the standard drug (Tetracycline at 100µg/mL) used. At lower concentrations, the
270 extracts seem to show more activity against shigella dysenteriae as seen in tables 3 - 6.

271 From table 3, it is revealed that the zones of inhibitions of the extract (ARB) against
272 the tested pathogens showed that the extract has antimicrobial activities against the pathogens at
273 various concentrations respectively. It was noticed that the extract was very effective at a
274 concentration of 100 mg/mL, the effectiveness increases as the concentration increases. The
275 control was more effective on *E. coli* with zone of inhibition up to 20 mm. Table 4 shows the
276 zones of inhibitions of the aqueous stem bark extract (ASB) on the microorganisms. The result
277 shows that the extract was effective at different concentrations with various zones of inhibitions
278 as the concentration increases. However, *E. coli* was resistant against the extract at higher
279 concentration of 100 mg/mL and 50 mg/mL but effective at lower concentration 25 mg/mL and
280 also the control which has the highest zone of inhibition (11mm) on *E. coli*. From table 5, the
281 ethanol stem bark extract (ESB) also showed considerable antimicrobial activities on the tested
282 clinical isolates at various concentrations used. The result shows that at a higher concentration
283 the extract was active against the clinical isolates or pathogens but more effective on *Shigella* at
284 lower concentration (25 mg/mL) with zone of inhibition 10 mm, also the control was more
285 effective with the highest zone of inhibition 16 mm. This extract show more activity against *E.*
286 *coli* than the control drug at 100 mg/mL with 15 mm zone of inhibition. From table 6 the results
287 of ethanol root extract (ERB) against the pathogens also shows that the antimicrobial potential of
288 the extract increases considerably as the concentration increases.

289 The result of the antimicrobial activity of root and stem bark extracts in this study is
290 similar to that of [27,28,29] who asserted that many plants have been reported for therapeutic
291 purposes because of the chemical compounds synthesized in these plants. The antibacterial
292 activities of the ethanolic extracts of the leaves and stem bark of *F. sycomorus* have been previously
293 reported [28]. The present study suggests that *F. sycomorus* may serve as a potential source of
294 antibacterial and/or antimicrobial agents of plants origin. Hence, the observed antimicrobial
295 activity of the root and stem bark extracts against the test organisms in this study may be due to
296 the presence of phytochemical components. The findings demonstrated that the stem and root
297 bark extract were sensitive to all the tested organisms and thus showed that the extract contained
298 potential antimicrobial agents such as tannin, saponin, alkaloid, glycosides as secondary
299 metabolite responsible for curing various sicknesses .The presence of tannin in all the extract
300 could be probably responsible for the observed antimicrobial activity. The claim of literature
301 that *F. sycomorus* has antimicrobial activity is hereby verified. The anti-microbial activity of the

302 extracts, both the ethanol and aqueous of root and stem have shown a reasonable zone of
 303 inhibition to the concentration from 12.5 – 100 mg/mL and the control drug (Tetracycline) at 100
 304 µg/mL concentration. However, the ASB extracts of *F. sycomorus* was observed to be less potent
 305 against the tested clinical isolate respectively.

306 **The Minimum Inhibitory Concentration (MIC) and Minimum Bactericidal Concentration**
 307 **(MBC) of the extracts are shown in Tables 7 and 8.** The result has shown that the MIC for all
 308 extracts of root and stem bark was 50 mg/mL. At this concentration, the extract was able to
 309 inhibit the growth of microorganisms. The result also revealed that the MBC was at 100 mg/mL
 310 these means that at this concentration the extract was able to kill the bacteria completely. This
 311 result is similar to the work of [27] who reported that the Minimum Inhibitory Concentration
 312 (MIC) and Minimum Bactericidal Concentration (MBC) of the aqueous and ethanolic root and
 313 stem bark extracts of *Ficus sycomorus* extracts ranged from 3.125 mg/mL to 100 mg/mL. [21].
 314 also reported that the minimum inhibitory concentration (MIC) of methanol root bark extract of *F.*
 315 *sycomorus* was observed within the range of 2.5 – 5.0mg/ml against *E. faecalis*, *E. coli*, *S. typhi*, *S.*
 316 *dysenteriae* and *C. albicans*. This result therefore suggests that the extracts are more of
 317 bacteriostatic.

318 **Table 1:** Percentage yield of the root and stem extracts

Extract	Initial weight	YIELD(g)	%
ERB	200.00g	14.14g	7.07
ESB	200.00g	16.00g	8.00
ARB	200.00g	12.23g	6.12
ASB	200.00g	11.16g	5.58

319 Key: ERB-----Ethanol Root Extract, ESB-----Ethanol Stem Bark Extract, ARB-----Aqueous
 320 Root Extract, ASB-----Aqueous Stem Bark Extract

321
 322 **Table 2: Qualitative Phytochemical analysis of the root and stem bark extract of**
 323 *Ficus sycomorus*

TEST	Aqueous extract		Ethanol extract	
	Root	Stem bark	Root	Stem bark
Tannins	+	+	+	+
Saponin	+	+	+	+

328	Terpenoid	+	+	+	+
329	Flavonoids	+	+	+	+
330	Alkaloids	+	+	+	+
331	Glycosides	-	-	+	+
332	Steroids	+	+	+	+
333	Phenols	+	+	+	+
334	Reducing sugar	+	+	+	+

335 + = Present - = Absent

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Table 3: Zone of Inhibition in (mm) Aqueous Root bark Extract (ARB) Against Opportunistic Pathogens.

S/No.	Name of Organism	Concentration mg/mL				
		100	50	25	12.5	Tetracycline(Control)
	<i>S. aureus</i>	7	6	5	2	13
	<i>Escherichia coli</i>	10	8	7	4	20
	<i>Salmonella spp</i>	12	7	5	R	13
	<i>Shigella spp</i>	10	9	7	4	13

351 Key:

352 **Resistant---- R**
353 **Aqueous Root bark Extract----- ARB**
354

Table 4: Zone of inhibition in (mm) of Aqueous stem bark extract (ASB) against Opportunistic Pathogen

Name of Organism	Concentration mg/mL
------------------	---------------------

S/No.	Name of Organism	Concentration mg/MI				
		100	50	25	12.5	Tetracycline(Control)
	<i>S. aureus</i>	6	5	4	2	10
	<i>Escherichia coli</i>	15	9	3	2	12

		100	50	25	12.5	Tetracycline(Control)
	<i>S. aureus</i>	9	6	4	3	7
	<i>Escherichia coli</i>	R	R	10	4	11
	<i>Salmonella spp</i>	7	5	4	3	8
	<i>Shigella spp</i>	9	6	5	4	10

357 **Key: Resistant----- R Aqueous stem bark extract ----ASB**

358 **Table 5: Zone of Inhibition in (mm) of Ethanol stem bark extract (ESB) against**
359 **Opportunistic Pathogens**

Name of Organism	Concentration mg/mL				
	100	50	25	12.5	Tetracycline(Control)
<i>S. aureus</i>	6	5	4	2	10
<i>Escherichia coli</i>	15	9	3	2	12
<i>Salmonella spp</i>	10	6	5	3	11
<i>Shigella spp</i>	5	4	10	5	16

360 **Key: Ethanol stems bark extract----- ESB**

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362

363 **Table 6: Zone of Inhibition (mm) of Ethanol root bark Extract (ERB) against**
364 **Opportunistic Pathogens.**

<i>Salmonella spp</i>	10	5	6	3	11
<i>Shigella spp</i>	10	5	5	4	16
	100	50	25	12.5	

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Key: Ethanolic root bark extract----- ERB

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368 **Table 7: The Result of Minimum Inhibitory Concentration (MIC) of both aqueous and**
369 **ethanol extracts of root and stem bark of *Ficus sycomorus***

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371 **Microorganism** **MIC (mg/mL)**

372 **+ = Growth ; - = No growth**

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	100	50	25	12.5
<i>Staphylococcus aureus</i>	-	-	+	+
<i>Escherichia coli</i>	-	-	+	+
<i>Salmonella spp</i>	-	-	+	+
<i>Shigella spp</i>	-	-	+	+

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388 **Table 8: The Result of Minimum Bactericidal Concentration (MBC) of both aqueous and**
389 **ethanol extracts of root and stem bark of *Ficus sycomorus***

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391 **Microorganism** **MBC (mg/mL)**

<i>Staphylococcus aureus</i>	-	+	+	+
<i>Escherichia coli</i>	-	+	+	+
<i>Salmonella spp</i>	-	+	+	+
<i>Shigella spp</i>	-	+	+	+

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393 + = Growth ; - = No growth

394 **CONCLUSION**

395 Phytochemicals such as tannins, saponin, terpenoids, flavonoids, alkaloids, glycosides, steroids,
 396 phenols and reducing sugars were all found to be present in both the aqueous extracts of roots
 397 and stem bark of *Ficus sycomorus*.

398 From the studies of the antimicrobial activities, the research revealed that, for aqueous stem and
 399 root bark, **ARB** had more antimicrobial potentials against the selected pathogens than the **ASB**,
 400 but for ethanol stem and root bark both have almost the same inhibitory activities on the tested
 401 pathogens.

402 From the research, it was noticed that both the root and stem bark may serve as potential
 403 antimicrobial agents. This validates the claim of the traditional users who used it to treat
 404 diseases of microbial origin. Therefore, it can be used for therapeutic purposes.

405

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410

411 **Ethical approval and consent are not applicable.**

412

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