

1 **Nutrient, bioactive components and effects of ethanol extracts of the leaves *Annona***
2 ***muricata* and roots of *Fagara zanthoxyloide* on zidovudine-induced oxidative stress in**
3 **wistar rats**

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6 **ABSTRACT**

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10 ethanol extracts of the leaves of *Annona muricata* (*AM*) and the roots of *Fagara zanthoxyloide*
11 (*FZ*) on zidovudine-induced oxidative stress in wistar albino rats. Wistar albino rats were divided
12 into **four groups of five** rats each. Groups 2-4 were induced with 100g/ml/Kg bw of zidovudine
13 (*ZDV*) and varying concentrations of the extracts (group 3 and 4); while group 1 served as the
14 control. The results of the proximate composition of both plants showed the following ranges:
15 moisture (10.32-18.30%), ash (0.65-9.45%), crude protein (1.38-10.54%), crude fat (2.35-
16 9.73%), crude fibre (3.00-15.53%) and carbohydrate (50.19-65.23%). Iron was the highest
17 mineral present in all the samples followed by zinc and calcium for *FZ* and *AM* respectively;
18 while folate and ascorbic acid were the highest vitamins present in both samples. Phytochemical
19 composition results revealed higher concentrations of alkaloids, flavonoid and phenols in the
20 leaves and roots of both samples. Acute toxicity study revealed no short term toxicity below
21 6g/ml/Kg bw for the leave extract of *Annona muricata* and 4g/ml/Kg bw for the root extract of
22 *Fagara zanthoxyloide*. Administration of zidovudine to albino rats resulted in a significant
23 increase ($p \leq 0.05$) in biomarkers of oxidative stress; while subsequent treatment with ethanol
24 extracts of the leaves of *AM* and roots of *FZ* reduced the activities of superoxide dismutase,
25 catalase and glutathione. The splenic histology revealed atrophy, early onset necrosis and
26 reduction in sinusoidal pore size in the negative control group which were absent in the extract
27 treatment groups indicating a protective effect conferred by extracts against oxidative stress. The
28 study, therefore suggests that these plants may play some key roles in alleviating salient
29 nutritional, physiological and oxidative stress related challenges.

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33 **Keywords: *Annona muricata*, *Fagara zanthoxyloide*, nutrients, bioactive components,**
34 **oxidative stress, zidovudine**

35 INTRODUCTION

36 Several plants have been used by rural dwellers within Nigeria as a source of medicine and
37 nutritional nourishments in periods of famine, drought, and civil unrest. With the increased
38 interest in alternative medicines and healthy feeding observed in the past decades, urban dwellers
39 have widened their scope to embrace the possible nutritional and medicinal value attached to
40 several plants observed around cities, towns and villages. The growing concern for the
41 alternatives have spurred research into several plants to further broaden the genetic diversity and
42 suggest varieties of nutritionally viable alternatives to the increasing medical and malnutrition
43 problems observed in South-southern Nigeria (Adisa *et al.*, 2014). Of the vast array of plants
44 which surround this region, two plants (*Annona muricata* and *Fagara zanthoxyloide*) have been
45 used by 65% of traditional medicine dispensers to prepare medicinal concoctions for treatment
46 and general body well being.

47 *Annona muricata* is one plant with widely acclaimed historical benefits to human beings and
48 commonly referred to as sour soup in Nigeria. Its fruit is the main portion consumed by most
49 people. This plant has been acclaimed to possess antihypertensive and antiplasmodic potentials;
50 as well as been used in the treatment of neuralgia, palpitation, parturition, rashes, rheumatism,
51 ringworm, spasms, tumors and ulcers with little mention of its nutritional potential. Notably, the
52 leaves have been used traditionally in the treatment cystitis, diabetes, insomnia and headaches
53 (Mishra *et al.*, 2013), coughs, skin diseases and pains (Moghadamtousi *et al.*, 2015). According
54 to Abdul-Wahab *et al.*, (2018), the roots have been acclaimed to have anti-inflammatory and
55 anthelmintic potentials (Adewole and Ojewole, 2009). They leaves, fruits and roots have also
56 been used as insecticides and pesticide agents among Africans as well as insect repellants
57 (Abdul-Wahab *et al.*, 2018).

58 *Fagara zanthoxyloide* is another ethnomedicinal plant which belongs to the family *Rutaceae*. It
59 is an indigenous south-southern Nigeria plant that is widely used as chewing stick for tooth
60 cleaning in West Africa. *Fagara zanthoxyloide* has been acclaimed to possess antiplasmodial
61 activity and used in the treatment of elephantiasis, impotence, sexually transmitted infections
62 such as gonorrhoea, abdominal pain and malaria (Adefisoye *et al.*, 2012).

63 Zidovudine, a potential inhibitor of HIV replication has been reported to increase mitochondrial
64 lipid peroxidation and mitochondrial glutathione probably due to the releases of free radicals as
65 well as reactive oxygen species in zidovudine medicated individuals (De la Asuncion et *al.*,
66 2004). Oxidative stress **occurs** when the free radicals produced during normal cellular activities
67 exceed the anti-oxidative capacity of the systems resulting in an increase in the radicals which
68 leads to attack of the system's proteins, lipids and nucleic acids. The resulting cellular damage as
69 a result of oxidative stress has been implicated to play a role in the pathogenesis of several
70 diseases. In a bid to provide data to back up the belief displayed by traditional medicine
71 dispensers on these plants, this study was carried out to investigate the nutrient, bioactive and
72 effects of the extracts of *Annona muricata* leaves and *Fagara zanthoxyloide* roots on zidovudine
73 induced oxidative stress in wistar rats.

74 **MATERIALS AND METHOD**

75 ***Collection of plant samples***

76 The leaves and roots of *Annona muricata* were obtained from Alakahia community
77 (3.92°N,7.80°E) in Obio/Akpor Local government Area of Rivers state; while leaves and roots of
78 *Fagara zanthoxyloide* were obtained from Opoo community (8.28°N,3.67°E) in Itesiwaju Local
79 Government Area of Oyo State. The plant materials were identified by Dr. B. Chikezie in the
80 Department of Plant Science and Biotechnology, University of Port-Harcourt and the Chief
81 Taxonomist, Dr. A. Olatunji University of Ibadan Herbarium (UIH) with a voucher copy
82 **(UIH/034/8212)** placed in the herbarium for reference.

83 ***Preparation of plant samples***

84 The leaves and the roots were sorted, washed with distilled water and air dried at room
85 temperature (25°C) until constant weight was obtained and milled with a Thomas-Willey (model
86 3383L40) mechanical grinder into a uniform coarse powder. The sample was stored in an air
87 tight container until analysis.

88 ***Plant sample extraction***

89 Three hundred grams (300g) of each of the dried powdered samples were placed in a conical
90 flask and extraction was performed using 3 litres of absolute ethanol for a period of 1 week. The
91 mixture was agitated at an interval of 48 hours on a rotary shaker. The extracts were centrifuged
92 twice at 1500rpm for 15 min in a Wilten-bioteknika Microspin-12 LCM-3000 centrifuge,

93 filtered with Whatman No. 1 filter paper, evaporated to dryness at 40°C with a rotary evaporator
94 and lyophilized to recover the residue as sticky pastes which were stored at 4°C in a refrigerator
95 for further use.

96 ***Laboratory Animals***

97 ***Acute toxicity study***

98 The toxicity study was carried out using wistar albino rats (200g - 237g) divided into six groups
99 with **five rats** each (one control group and 5 treatment groups) performed according to the
100 **Organization for Economic Cooperation and Development (OECD, 2011) as described in**
101 **Ogbuehi *et al.*, 2015.** They were acclimatized for seven days while on standard feed and water
102 *ad libitum*. Treatment group were administered leaf extract of *Annona muricata* and *Fagara*
103 *zanthoxyloide* at 2,4,6,8 and 10 g/ml/Kg BW while the control group was administered only
104 distilled water (2.5ml/kg orally). A graph of dose to experimental response was plotted for each
105 extract from which the LD₅₀ of the various extract was determined.

106 ***Experimental design***

107 Healthy albino rats were purchased from the animal house of the Department of Biochemistry,
108 University of Port Harcourt, Rivers state, Nigeria and divided into four groups (with 5 rats each)
109 and housed in Griffin and George modular cage system. The extracts of the leaves of *Annona*
110 *muricata* and roots *Fagara zanthoxyloide* were used for animal studies. All animals were treated
111 in a manner that complied with the National Institute of Health (NIH) guidelines for the care and
112 use of laboratory animals (NIH, 1985). Zidovudine was used to induce stress *in vivo* for two
113 weeks in groups 2-4, prior to experimental treatment with extracts while treatment with extracts
114 was performed for a period of 6 weeks.

115 **Group 1 (Normal control):** received no zidovudine or extract treatment

116 **Group 2 (Negative control):** received 100mg/ml of Zidovudine per Kg bw,

117 **Group 3 (AM+ZDV):** Served as *Annona muricata* treatment group which received
118 4.5g/ml/Kg bw of *Annona muricata* leaves extract.

119 **Group 4 (FZ+ZDV):** Served as *Fagara zanthoxyloide* treatment group which received
120 3.8g/ml/Kg bw of *Fagara zanthoxyloide* root extract.

121 **METHODS**

122 ***Proximate analysis***

123 The proximate analysis was performed according to the method described by the Association of
124 Official Analytical Chemists (AOAC, 2010).

125 ***Mineral analysis***

126 The evaluation of the potassium, calcium, magnesium, copper, iron, sodium and potassium
127 concentrations were performed according to the procedures by AOAC (2010) and Achi *et al.*,
128 (2017).

129 ***Vitamin analysis***

130 The concentrations of retinol, α -tocopherol, thiamine, niacin, riboflavin, vitamin K and folate
131 were performed by the method of AOAC (2010) and Okwu and Josiah (2006).

132 ***Phytochemical Analysis***

133 The concentrated extract samples were screened for phytochemical constituents according to the
134 methods described by Sofowora (2008) and the quantitative constituents according to the method
135 described by Ogunnka-Nnnoka *et al.*, (2019).

136 ***Biomarkers of oxidative stress***

137 Splenic homogenate was used for the assay of biomarkers of oxidative stress. Superoxide
138 dismutase, catalase activity and reduced glutathione in spleen were determined by the method
139 described by Fridovich (1989) and Aebi *et al.*, (1983). Lipid peroxidation assay and splenic H₂O₂
140 concentration were performed by the methods of De Las Heras *et al.*, (2003).

141 ***Histological analysis***

142 Histological examination of the excised spleen was performed by the method of Al-Hasawi and
143 Al-Harbi (2014)

144 **STATISTICAL ANALYSIS**

145 Results were expressed as Mean \pm Standard error of mean with analysis of variance and Student
146 t-test performed using SPSS software version 20 for Windows (SPSS Inc. USA). The significant
147 level during the test was set at $p \leq 0.05$.

148 **RESULTS AND DISCUSSIONS**

149 The result of the proximate analysis of the roots and leaves of *Annona muricata* as well as
150 *Fagara zanthoxyloide* (Table 1) showed that they contained high crude fibre, fat and protein. The

151 leaves of *Annona muricata* was observed to possess high carbohydrate while the roots of *Fagara*
 152 *zanthoxyloide* possessed high ash content.

153

154 **Table 1: Proximate composition of the leaves and roots of *Annona muricata* and *Fagara***
 155 ***zanthoxyloide***

Proximate parameter	<i>Annona muricata</i>		<i>Fagara zanthoxyloide</i>	
	Leaves	Roots	Leaves	Roots
Crude protein (%)	1.38 ± 0.03 ^a	7.73 ± 0.09 ^{a,c}	9.85 ± 0.02 ^{c,d}	10.54 ± 0.021 ^b
Crude fat (%)	9.73 ± 0.11 ^{a,c}	6.37 ± 0.04 ^{a,d}	2.35 ± 0.015 ^d	5.80 ± 0.005 ^{b,c}
Crude ash (%)	0.65 ± 0.01 ^c	1.94 ± 0.04 ^d	8.31 ± 0.011 ^{a,b}	9.47 ± 0.015 ^c
Crude fiber (%)	3.00 ± 0.02 ^{b,c}	8.27 ± 0.08 ^c	15.53 ± 0.005 ^a	10.63 ± 0.011 ^{a,d}
Moisture (%)	18.30 ± 0.01 ^b	13.38 ± 0.18 ^{c,d}	10.32 ± 0.011 ^c	12.85 ± 0.036 ^{c,d}
Carbohydrate (%)	65.23 ± 0.12 ^{a,b}	52.76 ± 0.33 ^{a,c}	50.19 ± 0.011 ^{c,d}	55.32 ± 0.011 ^{b,d}

156 Values expressed as Mean ± SEM of triplicate determinations. Values with same superscript are
 157 statistically significant (p≤0.05).

158

159 The high fibre content present in the roots and leaves of *FZ* may aid digestion of food thus
 160 preventing constipation. It also results in reduction of cholesterol levels in the serum
 161 (Shemishere *et al.*, 2018). The high crude protein observed in the roots and leaves of *Fagara*
 162 *zanthoxyloide* and roots of *Annona muricata* may play a key role in transmission of neuro-
 163 informations and genetic traits. The crude fat also observed to be present in all samples may
 164 indicate the possibility of samples to act as alternative energy sources. The carbohydrate values
 165 in all the samples may suggest that they may serve as good sources of energy. The ash content of
 166 the roots and leaves of *Fagara zanthoxyloide* may suggest the possibility of an appreciable
 167 amount of minerals present in these samples (Shemishere *et al.*, 2018).

168 Mineral analysis revealed significant (p≤0.05) levels of iron in all the samples followed by zinc
 169 and calcium for *FZ* and *AM* respectively. Significantly (p≤0.05) high levels of magnesium and
 170 copper were recorded for *AM* leaves and roots of *FZ* respectively.

171 **Table 2: Mineral content of the leaves and roots of *Annona muricata* and *Fagara***
 172 ***zanthoxyloide***

	<i>Annona muricata</i>	<i>Fagara zanthoxyloide</i>
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	Leaves	Roots	Leaves	Roots
Calcium (%)	3.67 ± 0.06 ^{a,c}	1.59 ± 0.01 ^{c,d}	0.19 ± 0.020 ^a	1.03 ± 0.015 ^{b,d}
Magnesium (mg/100g)	3.04 ± 0.01 ^{b,d}	2.18 ± 0.005 ^{b,d}	0.27 ± 0.01 ^{c,d}	0.47 ± 0.040 ^{a,c}
Sodium (%)	0.36 ± 0.38 ^{a,d}	1.08 ± 0.015 ^a	0.27 ± 0.350 ^{b,c}	0.167 ± 0.011 ^{b,d}
Potassium (%)	0.47 ± 0.021 ^{c,e}	1.68 ± 0.040 ^c	0.28 ± 0.005 ^{a,e}	0.57 ± 0.012 ^{c,d,e}
Zinc(mg/100g)	0.34 ± 0.040 ^c	1.35 ± 0.010 ^{d,e}	5.16 ± 0.02 ^d	5.32 ± 0.011 ^{d,e}
Iron (mg/100g)	20.23 ± 0.01 ^{b,d}	5.21 ± 0.02 ^{a,d,e}	10.01 ± 0.01 ^{b,c,d}	15.02 ± 0.02 ^{c,e}
Copper (mg/kg)	2.17 ± 0.011 ^a	0.16 ± 0.01 ^{a,b}	2.53 ± 0.011 ^{b,c}	7.38 ± 0.017 ^{a,c}

173 Values expressed as Mean ± SEM of triplicate determinants. Values with same superscript are
174 statistically significant (p≤0.05).
175

176 The samples studied recorded appreciable amount of iron. Iron has been known to play a part in
177 haemoglobin formation as well as aid in the oxidation of biomolecules. In synergy with copper
178 and cobalt, iron as observed in *Moringa oleifera* may stimulate bone marrow activity and
179 enhance red blood cell production and maturation. Thus, their presence in these plants studied
180 may suggest their usefulness in blood boosting. The high calcium content found in the leaves of
181 *AM* may be essential for blood clotting (hence coping with internal haemorrhage), bone
182 formation, contraction of muscles, normal functioning of the respiratory and nervous systems as
183 well as a vital co-factor for the process of erythropoiesis. The high copper content in the roots of
184 *FZ* shows that it can aid proper absorption of iron from the gastrointestinal tract, which leads to
185 increase in iron concentration (boosting iron stores). Zinc also observed in the roots and leaves of
186 *FZ* is known to play a pivotal role as essential components of several enzyme systems such as
187 carbonic anhydrase, alanine peptidase, carboxypeptidase, carbonic anhydride. The impact of zinc
188 is a salient one, but on the average about 20% of children in Nigeria is at risk of inadequate zinc
189 intake with values of micronutrient deficiency in south-southern Nigeria increasing by the day
190 (Dioxin and Harris, 2004). Thus, its presence in these plants may imply the benefit of the plants
191 to protein synthesis, cell differentiation and replication as well as increased immunity as immune
192 cells require iron, copper and zinc for their continuous generation in the bone marrow. The
193 minerals in these plants may thus be used to combat micronutrient deficiency.

194 Analysis of vitamins content revealed varying concentrations of vitamins with significantly high
195 (p≤0.05) concentrations of folate and ascorbic acid observed in both *AM* and *FZ*. The leaves of

196 *Annona muricata* were also observed to contain significantly higher ($p \leq 0.05$) concentrations of
 197 vitamins D, E and K. The vitamin components of these plants may also prove their relevance in
 198 several nutritional deficiency disorders.

199 **Table 3: Vitamin content in the leaves and roots of *Annona muricata* and *Fagara***
 200 ***zanthoxyloide***

	<i>Annona muricata</i>		<i>Fagara zanthoxyloide</i>	
	Leaves	Roots	Leaves	Roots
Retinol ($\mu\text{g}/100\text{g}$)	3.81 \pm 0.14 ^{a,c}	1.97 \pm 0.09 ^{a,c,d}	0.16 \pm 0.17 ^b	ND
Niacin (mg/Kg)	4.86 \pm 0.19 ^a	4.23 \pm 0.32 ^{c,d}	9.18 \pm 0.19 ^{d,e}	8.23 \pm 0.81 ^{d,e}
Riboflavin (mg/kg)	9.72 \pm 0.29 ^{c,e}	7.89 \pm 0.11 ^{a,d,e}	6.28 \pm 0.02 ^{c,e}	10.21 \pm 0.27 ^{c,d}
Folate (mg/Kg)	26.82 \pm 0.48 ^{a,b}	23.47 \pm 0.03 ^{b,c}	15.82 \pm 0.18 ^{a,c}	20.63 \pm 0.91 ^{d,e}
Ascorbic acid (mg/Kg)	31.97 \pm 0.03 ^{a,b}	26.89 \pm 0.19 ^{d,e}	13.86 \pm 0.13 ^{c,e}	30.21 \pm 0.01 ^{c,d}
Vitamin D (mg/Kg)	4.21 \pm 0.21 ^{c,d,e}	0.91 \pm 0.16 ^{c,e}	1.11 \pm 0.26 ^{b,d}	3.21 \pm 0.49 ^{b,d,e}
Vitamin E (mg/Kg)	5.82 \pm 0.01 ^{a,d}	0.18 \pm 0.19 ^{c,d}	0.27 \pm 0.48 ^{a,c,e}	5.08 \pm 0.04 ^c

201 Values expressed as Mean \pm SEM of triplicate determinants. Values with same superscript are
 202 statistically significant ($p \leq 0.05$). *ND = Not detected

203
 204 Ascorbic acid present in significantly high ($p \leq 0.05$) concentrations in all samples has been
 205 known to be an antioxidant which reduces the concentration of reactive oxygen species in the
 206 body and as result increases immunity and decrease peroxidation (Gora *et al.*, 2006). Scurvy
 207 haemorrhage a condition common to this region may be alleviated by these plants due to the
 208 ascorbic acid content of which its deficiency results in weakening of the endothelial wall of the
 209 capillaries around the gums. The presence of vitamin E and riboflavin in the leaves of *AM* and
 210 roots of *FZ* have also been known to induce antioxidant properties when consumed thereby
 211 protecting cells of the body against free radical-induced oxidative damage. A diet rich in
 212 riboflavin as seen in all samples have also been linked to the proper maintenance of the
 213 connective tissues thus facilitating wound healing. Niacin and riboflavin common to both
 214 samples may aid co-enzyme formation leading to increased oxidative phosphorylation and thus
 215 energy production through the electron transport chain. Retinol although observed in small
 216 quantities in the leaves and roots of *AM* used in this study may in conjunction with ascorbic acid
 217 lead to an increase in the iron absorption from the gastrointestinal tract and its release from iron

218 stores. This would thus promote the proliferation of the red blood cells in the bone marrow and
 219 reduce anaemic related condition observed among young women and geriatric individuals in this
 220 region.

221 Phytochemical analysis of the plants revealed the presence of alkaloids, flavonoid, tannins,
 222 phenols and steroids in the leaves and roots of both plants used in the study. Saponin was
 223 however not observed in the roots of *Annona muricata* but present in *Fagara zanthoxyloide* as as
 224 illustrated in Table 4. Quantitative phytochemical examination revealed significantly high
 225 ($p \leq 0.05$) concentrations of phytochemicals in the leaves of *Annona muricata* than in the roots.

226 **Table 4: Phytochemicals in leaves and roots of *Annona muricata* and *Fagara zanthoxyloide***

Phytochemical	<i>Annona muricata</i>		<i>Fagara zanthoxyloide</i>	
	Leaves	Roots	Leaves	Roots
Alkaloids	+	+	+	+
Flavonoids	+	+	+	+
Tannins	+	+	+	+
Terpernoids	+	+	+	+
Saponins	+	-	+	+
Phenols	+	-	+	+

227 **Present: +, Absent: -**

228 The result of quantitative phytochemical analysis on *Fagara zanthoxyloide* revealed a high
 229 concentration of phytochemicals in the roots on comparison with the leaves. The roots were
 230 observed to be rich in alkaloids, tannins and terpenoid which were also significantly higher
 231 ($p \leq 0.05$) in roots than in the leaves (Table. 5). The leaves of *AM* and roots of *FZ* were observed
 232 to posses significantly high ($p \leq 0.05$) concentrations of all phytochemicals investigated when
 233 compared with the roots. The presence of a significant array of phytochemicals in the leaves of
 234 *Annona muricata* and the roots of *Fagara zanthoxyloide* may be the reason for their preferred
 235 use by traditional medicine dispensers within South-southern Nigeria than the other plant part
 236 investigated.

237 **Table 5: Quantitative phytochemicals in the leaves and roots of *Annona muricata* and**
 238 ***Fagara zanthoxyloide***

Phytochemical (mg/100g)	<i>Annona muricata</i>		<i>Fagara zanthoxyloide</i>	
	Leaves	Roots	Leaves	Roots
Alkaloids	27.34 ± 0.15 ^a	12.98 ± 0.98 ^{a,b}	35.55 ± 0.95 ^{a,c}	50.90 ± 0.83 ^{b,c}
Flavonoids	19.66 ± 0.04 ^{c,d,e}	3.71 ± 0.46 ^{b,c}	3.27 ± 0.34 ^{c,e}	8.63 ± 0.27 ^{a,c}
Tannins	11.24 ± 0.05 ^{a,c}	3.86 ± 0.22 ^{c,d}	28.70 ± 0.19 ^{a,e}	55.37 ± 0.47 ^{b,c,e}
Terpernoids	8.19 ± 0.11 ^{b,d}	5.21 ± 0.19 ^{b,c}	18.23 ± 0.08 ^{c,d,e}	41.21 ± 0.16 ^c
Saponins	6.32 ± 0.14 ^{a,e}	1.25 ± 0.07 ^{a,d,e}	7.43 ± 0.41 ^{a,d}	19.44 ± 0.59 ^{a,e}
Phenols	15.10 ± 0.11 ^{a,c}	0.07 ± 0.42 ^{b,c}	2.17 ± 0.2 ^{a,d}	13.23 ± 0.17 ^{c,d}

239 Values expressed as Mean ± SEM of triplicate determinants. Values with same superscript are
240 statistically significant (p≤0.05).
241

242 The phytochemicals seen in the roots and leaves of *Annona muricata* and *Fagara zanthoxyloide*
243 have been suggested in several studies to elicit several physiological properties. The high
244 alkaloids content in all samples which offer repellent properties to plants against predators and
245 parasites have been known to also be resourceful in intestinal infections which accompany
246 immunodeficiency disorders. The tannin content significantly high in the leaves and roots of FZ
247 has been implicated in the treatment of inflamed tissues. Generally, the presence of terpenoid **as**
248 and leaves of *AM* has been known to elicit stimulation of the immune system (Llauradó *et al.*,
249 2012). As such these plant extracts may be applied in the management of secondary
250 immunodeficiency conditions such as; HIV/AIDs, graft *vs* host diseases, leukaemia and
251 lymphoma. Notably, the high phenol content as seen in the leaves of *AM* and roots of *FZ* may
252 induce haematopoietic responses as well as confer antioxidant properties which have been
253 implicated in treatment and management of haemolytic anaemia (Oboh and Akindahunsi, 2004).
254 This possibly may be one of the reasons for the proposed use of the leaves of *Annona muricata*
255 and the roots of *Fagara zanthoxyloide* by traditional medicine dispensers within this region for
256 the management of symptoms of sickle-cell anaemia (Dubost *et al.*, 2007). All of these may
257 contribute synergistically to the use of these plants in herbal medications within this region.

258 ***Acute toxicity (LD₅₀)***

259 Result of the acute toxicity study on the extract of *Annona muricata* leaves and *Fagara*
260 *zanthoxyloide* roots are illustrated in Tables 6 and 7. The results revealed that administration as
261 from 8g/ml/Kg bw for *Annona muricata* leaves and 6 g/ml/Kg bw for *Fagara zanthoxyloide*

262 roots resulted in signs of toxicity and oral administration below this levels was well tolerated in
 263 mice even beyond 7 days. The result for the toxicity of *Annona muricata* was however slightly
 264 higher than the findings by Abdul-Wahab *et al.*, (2018) in which he observed kidney toxicity
 265 above 5g/Kg bw and Bertin *et al.*, (2017) who observed 100% mortality at 5g/Kg bw with
 266 *Annona muricata* from Benin was investigated.

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 268

269 **Table 6: Acute toxicity test on extracts of *Annona muricata* leaves**

Dosages (g/ml/Kg bw)	No. of rats used	No. of mortality	Remarks
-	5	0	
2	5	0	
4	5	0	
6	5	0	
8	5	1	Salivation, weight loss.
10	5	2	Lacrimation, reduced irritability

270
 271
 272

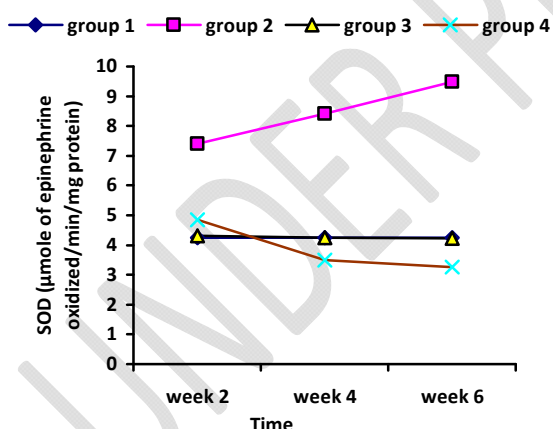
Table 7: Acute toxicity test on extracts of *Fagara zanthoxyloide* roots

Dosages (g/ml/Kg bw)	No. of rats used	No. of mortality	Remarks
-	5	0	
2	5	0	
4	5	0	
6	5	1	Reduced irritability, fur coat changes observed
8	5	2	Muscle paralysis, weight loss.
10	5	2	Weakness and salivation

273

274 *Fagara zanthoxyloide* acute toxicity test resulted in signs of toxicity as from 6g/Kg bw with
 275 animals witnessing weight loss, lacrimation and reduced irritability (Table 7). The median acute
 276 toxicity value (LD₅₀) which was estimated suggests that the extract possessed no short time
 277 toxicity. This value was also similar with that observed for *Fagara zanthoxyloide* root-peels by
 278 Ogwal-Okeng *et al.*, (2003) at 5g/ml/Kg bw; however the reduced irritability at dosages higher
 279 than the LD₅₀ may possibly culminate in the findings of Ogwal-Okeng *et al.*, (2003), suggesting
 280 a direct effect of extracts on nervous system.

281 Results on oxidative stress makers are shown in Figures 1-5 below. The CAT activity was
 282 observed to reduce in groups 3-4 when compared with group 2 with the values at week 6
 283 observed to approach the activity observed in the normal control (Figure 2). GSH concentration
 284 was observed to decrease in groups 3 and 4 when compared with group 2 at week 2. The
 285 concentration in group 4 was also observed to decrease below the value observed for group 1 at
 286 weeks 4 and 6. No significant change was observed in the H₂O₂ concentration when the values in
 287 group 3 and 4 were compared with the normal control (Figure 4). The level of lipid peroxidation
 288 was observed to be significantly higher in groups 3 and 4 in comparison with the normal control
 289 but lower than that observed for the negative control (Figure 5), with the values in group 4
 290 observed to significantly decrease at week 4 and 6. The administration of zidovudine resulted in
 291 an increase in superoxide dismutase (SOD) and catalase (CAT) activities, glutathione (GSH),
 292 hydrogen peroxide (H₂O₂) and lipid peroxidation (LPO) concentrations at week 2-6 (Figure 1-5).
 293 Treatment with extracts of *Annona muricata* and *Fagara zanthoxyloide* resulted in a significant
 294 decrease ($p \leq 0.05$) in these bio-markers which continued as time progressed. Treatment with
 295 *Fagara zanthoxyloide* roots caused a significant decrease ($p \leq 0.05$) in the SOD activity at weeks
 296 4 and 6 when compared to all groups (Figure 1).



297 Figure 1. Superoxide dismutase activities in wistar albino rats treated with extracts of *Annona muricata* leaves and *Fagara zanthoxyloide* roots

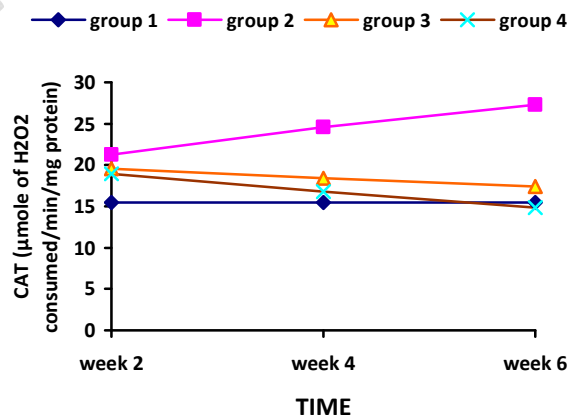
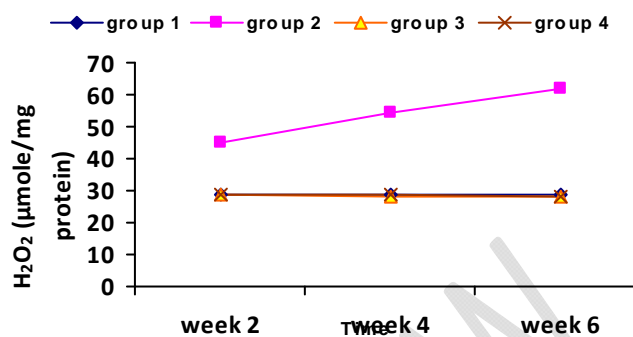
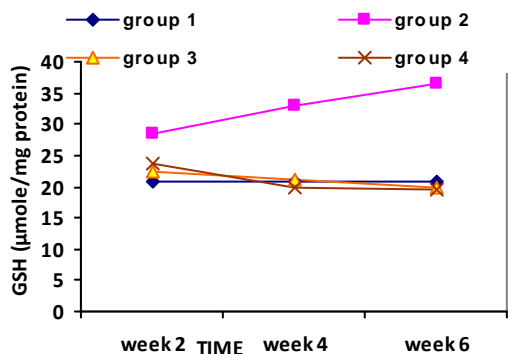
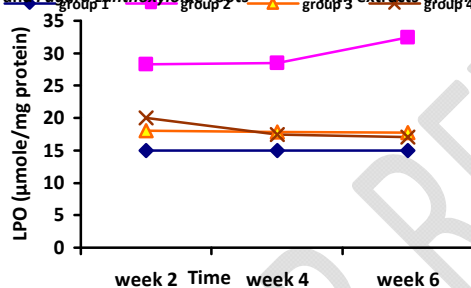


Figure 2. Catalase activities in wistar albino rats treated with extracts of *Annona muricata* leaves and *Fagara zanthoxyloide* roots



298

Figure 3. Glutathione concentrations in wistar albino rats treated with extracts of *Annona muricata* leaves and *Fagara zanthoxyloide* roots. Figure 4. Hydrogen peroxide concentrations in wistar albino rats treated with extracts of *Annona muricata* leaves and *Fagara zanthoxyloide* roots



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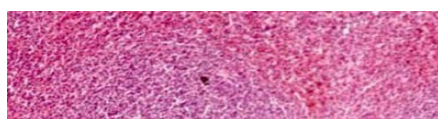
Figure 5. Lipid peroxidation concentrations in wistar albino rats treated with extracts of *Annona muricata* leaves and *Fagara zanthoxyloide* roots

300

301 The reduction in the levels of these biomarkers on treatment with extract may suggest that the plants
 302 may serve as good sources of antioxidants which aid in alleviating cytotoxic effects of reactive
 303 oxygen species which results in damages to biological molecules, DNA, membrane function and
 304 ultimately ageing.

305 The reduction in the activities of the enzymatic antioxidants by the extracts of *Annona muricata*
 306 leaves and *Fagara zanthoxyloide* roots together with the presence of non-enzymatic antioxidants
 307 (phenols, riboflavin, ascorbic acid, vitamins D and E) observed in both plants may buttress the
 308 impact of these plants in oxidative stress related scenarios. These non-enzymatic antioxidants may
 309 serve as the reason for the reduction in the concentration of biomarkers of oxidative stress in the
 310 spleens of rats used in the study and the proposed antioxidant effects of these plants.

311 The result of the splenic histopathology showed a reduction in the pore size of the splenic
 312 sinusoids seen in the negative control group as well as atrophy and necrosis but no case of
 313 splenomegaly (Plate 2-4). Treatment with extract of *Annona muricata* and also *Fagara*
 314 *zanthoxyloide* resulted in the preservation of the architecture of the spleen with normalization of
 315 sinusoid space and absence of atrophy and necrosis as observed in the negative control. There



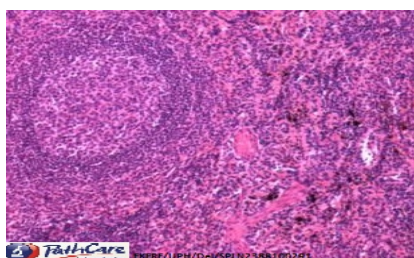
316 was however an observed increase in red blood cells and β -lymphocyte aggregation (Plate 9-10)
 317 in the group treated with *Fagara zanthoxyloide*.

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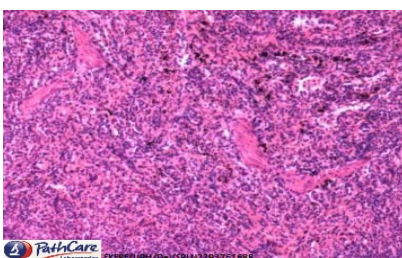
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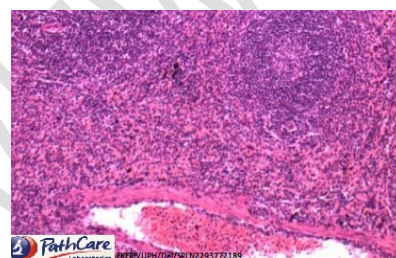
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323
324 **Plate 2.** Spleen of group 2 at week 2
325 showing scattered red and white pulp
regions with parenchyma cells.

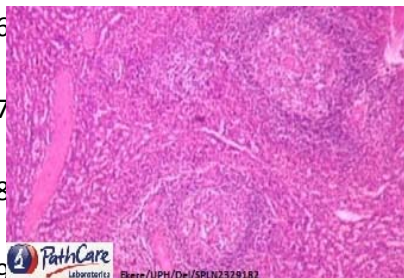


326
327 **Plate 3.** Spleen of group 2 at week 4
328 showing red and white pulp regions
329 with infiltration of neutrophils and
lymphocytes

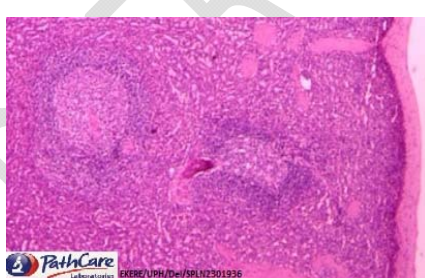


330 **Plate 4.** Spleen of group 2 at week 6
331 showing a normal central artery
332 constrictions in red pulp with cyanocilic
333 fibrosis and necrotic stroma.

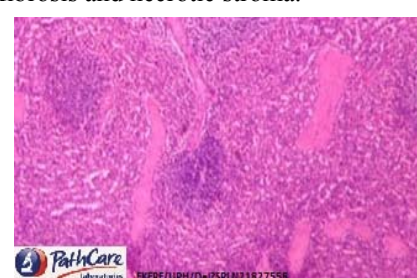
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329 **Plate 5.** Spleen of group 3 at week 2
330 showing white pulp with stream of
331 myeloblast

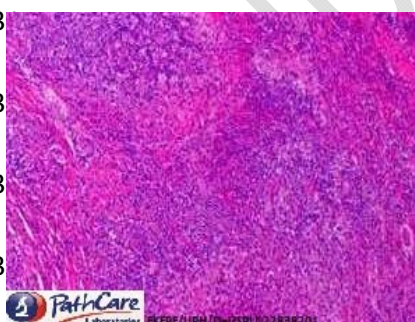


332
333 **Plate 6.** Spleen of group 3 at week 4
334 showing splenic vacuolation and
335 sinusoidal space.

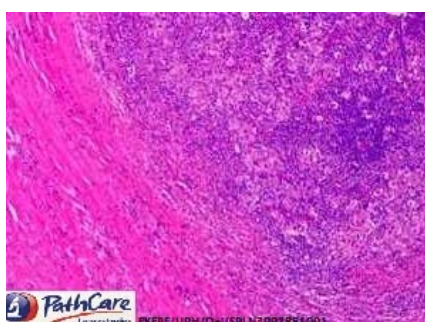


336 **Plate 7.** Spleen of group 3 at week 6
337 showing lymphoid sheath and
338 malpighian follicles

33



33
33
33
33 **Plate 8.** Spleen of group 4 at week 2
335 showing red blood sinusoids



336 **Plate 9.** Spleen of group 4 at week 4
337 showing region of β -lymphocyte
338 aggregation and lymph nodes



Plate 10. Spleen of group 4 at week 6
339 showing a lymphocyte around the
340 splenic capsule

336

337 The results from the treatment groups (Plates 5-10) suggest that the extracts may confer
338 protection against possibly oxidative stress induced necrosis and atrophy as observed in the
339 negative control group (Plates 2-4), thus preserving the architecture of the white and red pulps
340 and enabling the spleen perform its quality control function on the haematopoietic system. The
341 restoration of sinusoidal pore sizes also buttress the protective effect of the extracts on membrane
342 of splenic sinusoids against oxidation as proposed by Chapman and Azevedo, (2018) and may be
343 attributed to the presence of phytochemicals such as tannins, phenols and flavonoids as well as
344 other mineral and vitamins which confer anti-oxidative properties to the extracts. The increase in
345 red blood cells and β -lymphocyte aggregation in the group 4 may indicate normalization in the
346 splenic functionality on treatment with extracts of *Fagara zanthoxyloide* (Belonwu *et al.*, 2013a
347 &b).

348 CONCLUSION

349 Plants are a great source of food and medicine for humans. The proposed acclaimed effect of *AM*
350 and *FZ* by traditional healers may be due to the activity of several biochemical compounds in
351 them. An analysis of *Annona muricata* and *Fagara zanthoxyloide* has revealed that these plants
352 accumulate a high amount of phytochemicals and possess vitamins and minerals which can help
353 in cases of micronutrient deficiency and alleviating symptoms observed in several physiological
354 conditions. These bioactive components as seen from *in vivo* studies may also serve as potential
355 antioxidants and aid in reducing oxidative stress derived from toxicants, heavy metals and free
356 radicals present in the ecosystem.

358 Ethical Approval:

359
360 As per international standard or university standard ethical approval has been collected and
361 preserved by the authors.

363 CONFLICT OF INTERESTS

364 The authors declare that they have no conflicting interests.

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