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# Chapter

# Antioxidants from Nigerian Medicinal Plants: What Are the Evidence?

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#### **Abstract**

The search for natural antioxidants from plants would continue to be a dominant research interest for many years. This is because of the increasing understanding on the role of oxidative stress in damaging cell structures such as DNA, due to over production of free radicals and reactive oxygen species (ROS) in human systems, which are linked to inflammation, cancer and diabetes. However, phenolic compounds especially from phytochemicals or vegetable foods play important roles in reducing the risk of these diseases and reinforces the importance of natural antioxidants in human health. These antioxidant molecules neutralize or quench the ROS by either hydrogen atom transfer or single electron transfer mechanisms. Thus, the capacity to scavenge ROS and free radicals or inhibits lipid peroxidation is measured quantitatively as the strength of antioxidant activity. Several chemical and biochemical protocols have been used in the evaluation of plant extracts as antioxidants. Overwhelming literature reports have indicated varying degrees of antioxidant efficacies of extracts from Nigerian medicinal plants in comparison to synthetic antioxidants. These efficacies were analyzed to provide insight into the strength of antioxidant activity. This chapter reviewed 250 Nigerian medicinal plants in search of evidence for effective antioxidants.

Keywords: Nigerian medicinal plants, antioxidants, DPPH, ROS, free radicals

#### 1. Introduction

1

Since the discovery of enzyme superoxide dismutase (SOD) and the evidence that emerged in support of the role of free radicals in biological systems, human understanding of free radical biochemistry in health and disease continue to advance [1]. This provided the basis for continuous search on natural antioxidants from foods and phytomedicines. Overwhelming reports on Google search engine has indicated 92,800 hits for "antioxidant activity" of medicinal plants in the last 10 years (2008–2018). This is due to growing interest on the antioxidant properties of medicinal plants. Several chemical and biochemical protocols have been used in the

evaluation of antioxidant activity including the oxygen radical absorbance capacity (ORAC), total radical-trapping antioxidant potential (TRAP), total oxidant scavenging capacity (TOSC), chemiluminescence (CL), croton bleaching, low density lipoprotein (LDL) oxidation, ferric reducing antioxidant power (FRAP), copper reduction assay (CUPRAC), 2,2' azinobis(3-ethylbenzothiazoline-6-sulfonic acid (ABTS) assay, 2,2-diphenyl-1-picrylhydrazyl (DPPH), nitric oxide (NO), hydroxyl radical (OH) hydrogen peroxide ( $H_2O_2$ ) and total phenolic assay among others [2]. Biochemical protocols are based on animal models for *in vivo* evaluations of oxidative stress biomarkers. However, this study is focused on in vitro evaluations of antioxidants from plants based on hydrogen atom transfer or single electron transfer mechanisms [2]. The strength of antioxidant activity measured from a combination of different methodologies was used to evaluating antioxidant effectiveness [3]. This review provides fundamental background on free radical and ROS in human health and disease with a view to understand the roles of natural antioxidants. We reviewed 250 Nigerian medicinal plants evaluated for antioxidant activity in search of evidence for effective antioxidants.

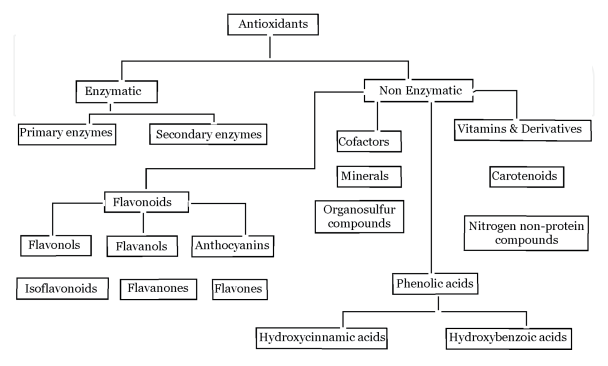
# 2. Reactive oxygen species (ROS) in human health and disease

Human system uses oxidation for normal metabolic activities in the transformation of nutrients into energy. During oxidation, reactive oxygen species (ROS) are also produced at low levels in normal physiological conditions, which are necessary for maintaining normal cell functions such as signaling immunity and homeostasis [4]. These activities are maintained by endogenous antioxidant (enzymatic) defense systems produced by the body for protection against harmful effects. These include superoxide dismutase (SOD), glutathione peroxidase (GSH-Px), glutathione reductase (GSH-Rx) and catalase [5]. Excessive production of ROS beyond the body defense mechanisms can be extremely harmful to cellular functions by damaging nucleic acids, oxidizing proteins, and causing lipid peroxidation [6]. The resultant cell damage by free radicals and ROS appeared as major contributor to aging and degenerative diseases of aging such as cancer, cardiovascular disease, immune system decline, liver diseases, diabetes mellitus, inflammation and brain dysfunction among others [7, 8]. These ROS and reactive nitrogen species (RNS) including superoxide anion O<sub>2</sub>-, hydroxide ion OH-, hydroxyl radical OH, peroxyl radical ROO and nitric oxide NO as well as  $H_2O_2$ , lipid peroxides ROOH, and singlet  $O_2$  are very reactive and can initiates free radical reactions or lipid peroxidation in living cells.

ROS can be produced either by external sources (e.g., tobacco smoke, stress, etc.), as by-products during the mitochondrial electron transport of aerobic respiration or by oxidoreductase enzymes and metal catalyzed oxidation [9]. But the biological effects of ROS depend on the types of cell or tissue in relation to enzyme production, signal transduction and DNA repair [10]. ROS are harmful when excessive productions are not balanced by body antioxidant mechanism. This imbalance between ROS production and enzymatic antioxidant defense systems is called oxidative stress [11]. Antioxidants counteract oxidative stress by neutralizing free radicals because they are reducing agents that react with and buffer ROS as a form of defense against oxidative stress [12].

# 3. Phytochemicals as sources of natural antioxidants

Antioxidants are molecules that prevent oxidation or inactivates the reactive oxygen species and thus prevent oxidative damage to the cells and body tissues [13]. Antioxidants can also inhibit, quench or scavenge free radicals converting them into new and stable chemical compounds [14]. Broadly, antioxidants are classified as enzymatic and non-enzymatic with each class providing complementary role of protection against free radicals in human systems. Previous work has concisely discussed on antioxidants classification [3] as summarily reproduced in **Figure 1**. But our focus is the non-enzymatic antioxidant involving flavonoids, phenolic acids, vitamins, carotenoids, minerals and cofactors. They are exogenous sources of protection through diet. Plants foods contain a variety of nutrients and non-nutrients chemicals which are good antioxidant agents. These sources of natural antioxidants including Vitamin A (retinol) obtained from  $\beta$ -carotene, vitamin C (ascorbic acid), Vitamin E ( $\alpha$ -tocopherol), lycopene and carotenoids occur naturally in fruits, vegetables, legumes and grains which are commonly consumed and play important role in the defense against free radicals [3, 15]. Medicinal plants are rich source of phenolic compounds such as flavonoids, phenolic acids and coumarins [16, 17]. Flavonoids are antioxidants compounds composed of anthocyanins, flavanones, flavonols, flavones, isoflavonoids and flavanones, while hydroxycinnamic and hydroxybenzoic acids such as gallic acid are components of phenolic acids widely distributed secondary metabolites in plants with antioxidant and antiradical properties [18]. They are important as chelators and free radical scavengers of hydroxyl and peroxyl radicals, superoxide anions and peroxynitrites [19]. Carotenoids natural pigments are important phytochemical antioxidants obtained from plants. They are structurally grouped into carotenes and xanthophyll based on the degree of oxygenation of carotenoid hydrocarbons and exert antioxidant effect by singlet oxygen quenching ability [3]. Several studies on the antioxidant activities of various herbal plants have indicated their enormous medicinal values as inhibitors of free radical and ROS [20].

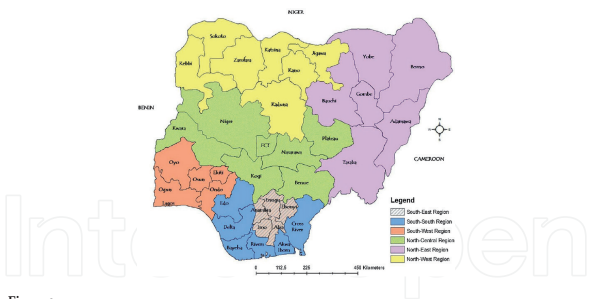


**Figure 1.**Board classification of antioxidants adapted from Carocho and Ferreira [3].

## 4. Antioxidants from Nigerian medicinal plants

Nigeria is a west African country with an area of 923,769 km² having a population of 198 million with 250 ethnic groups [21]. The country shares border with republic of Cameroun to the east, Niger and Chad republics to the north, Benin republic to the west and Gulf of Guinea to the south. Nigeria has favorable climate conditions with enormous diversity of plant species, which are distributed across geographical contrast of the mangrove swamps in South-South (SS), to the tropical rain forests covering South-West (SW) and South-East (SE) and to the grassland vegetation of North-Central (NC) up to the Sahel savannah of semi-arid North-East (NE) (**Figure 2**). Many of these plants are used as medicines for treatment of illness or management of human and animal health among rural and urban dwellers.

The application of herbal recipes especially in the management of human metabolic diseases such as diabetes and cancer is common knowledge among Nigerians. This prompted research interest in academia on the potentials of phytomedicines as complimentary or alternative treatment agents, and consequent research efforts to validate their pharmacological properties. The number of Nigerian medicinal plants reported for antioxidants is enormous. However, 250 medicinal plants evaluated for antioxidant activity were studied in addition to the 28 compounds isolated from 44 plants. But antioxidant evaluations on crude extracts rather than on pure compounds largely dominated the literature. Thus, effective activity based on concentrations required to inhibit 50% free radicals ( $IC_{50}$ ) for selected extracts are presented (**Table 1**) together with concentrations of various standard antioxidants used.



Map of Nigeria showing the six geopolitical zones.

S. No	Plant name	Family	Part used	IC <sub>50</sub> sample	IC <sub>50</sub> standard	Ref.
1	Abrus precatorius	Leguminosae	Seed	1.92 <sup>*</sup> 2.10 <sup>*</sup>	AA = 1.83 AA = 1.20	[37]
2	Acalypha ornata	Euphorbiaceae	Leaf	20.50 <sup>*</sup>	TC = 15.4	[68]
3	Acalypha wilkesiana	Euphorbiaceae	Leaf	15.25 <sup>*</sup>	AA = 7.26	[26]
4	Acanthospermum hispidum	Asteraceae	Aerial	28.9 <sup>*</sup>	AA = 1.41	[39]
5	Aframomum melegueta	Zingiberaceae	Fruit Leaf	0.04** 0.07**	AA = 0.03	[69]

S. No	Plant name	Family	Part used	IC <sub>50</sub> sample	IC <sub>50</sub> standard	Ref
6	Ageratum conyzoides	Asteraceae	Leaf	31.25*	AA = 7.26	[26]
7	Allamanda cathartica	Apocynaceae	Leaf	0.46**	VE = 0.25	[70
8	Allanblackia floribunda	Guttiferae	Leaf	0.02** 0.1**	VE = 0.01	[71
9	Alstonia boonei	Apocynaceae	Stem	0.12**	AA = 0.06	[72
10	Alstonia congensis	Apocynaceae	Root	19.7*	AA = 4.9	[73
11	Alternanthera dentata	Amaranthaceae	Leaf	35*	AA = 125	[46
12	Amaranth caudatus	Amaranthaceae	Leaf Stem	15.81**	TC = 13.2	[27
13	Anacardium occidentale	Anacardiaceae	Bark Leaf	5.66 <sup>*</sup> 7.77 <sup>*</sup>	AA = 4.57	[74
14	Annona senegalensis	Annonaceae	Leaf	45.72 <sup>*</sup> 49.0 <sup>*</sup>	GA = 48.77 TX = 72.9	[35
15	Aspilia africana	Asteraceae	Leaf	160 <sup>*</sup>	AA = 120	[75
16	Asystasia gangetica	Acanthaceae	Leaf	100 <sup>*</sup>	AA = 150	[75
17	Bauhinia galpinii	Caesalpiniaceae	Leaf	20.52 <sup>*</sup>	AA = 19.8	[76
18	Bauhinia monandra	Caesalpiniaceae	Leaf	5.56 <sup>*</sup>	AA = 30.0	[45
19	Bixa orellana	Bixaceae	Leaf	0.45*	VE = 0.25	[70
20	Borreria ocymoides	Rubiaceae	Aerial	1.85**	AA = 0.05	[77
21	Borreria verticillata	Rubiaceae	Leaf	2.98*	AA = 1.05	[78
22	Bridelia ferruginea	Euphorbiaceae	Leaf	12.5*	AA = 7.26	[26
23	Bridelia micrantha	Euphorbiaceae	Leaf	0.1 μΜ	AA = 2.0 μM	[38
24	Bryophyllum pinnatum	Crassulaceae	Leaf	0.41**	VC = 0.067	[79
25	Calliandra surinamensis	Mimosaeae	Flower	28**	VE = 38	[80
26	Calyptrochilum christyanum	Orchidaceae	Whole	50.6 <sup>*</sup>	AA = 1.41	[39
27	Canthium subcordatum	Rubiaceae	Leaf	23.9*	AA = 4.9	[74
28	Capsicum annuum	Solanaceae	Fruit	1.15**	BHA = 0.96	[25
29	Capsicum frutescens	Solanaceae	Fruit	0.67**	BHA = 0.96	[25
30	Carica papaya	Caricaceae	Seed	0.227**	AA = 0.109	[67
31	Cassia sieberiana	Leguminosae	Leaf	24.1* 46.6*	AA = 4.9 TC = 38.9	[73
32	Cassia singuaena	Leguminosae	Leaf	1.20*	AA = 2.56	[48
33	Celosia trigyna	Amaranthaceae	Leaf	120 <sup>*</sup>	AA = 120	[75
34	Cissampelos owariensis	Menispermaceae	Leaf	2.77*	AA = 0.067	[25
35	Citrus aurantifolia	Rutaceae	Peel	12.1**	VC = 0.067	[25
36	Commiphora kerstingii	Burseraceae	Leaf	0.33 <sup>**</sup> 0.54 <sup>**</sup>	AA = 0.49	[23
37	Corchorus olitorius	Malvaceae	Leaf	11.8 <sup>**</sup> 27.52 <sup>*</sup>	TC = 13.2 AA = 188.3	[27 [33
38	Crassocephalum rubens	Asteraceae	Leaf	2.91 <sup>**</sup> 1.73 <sup>**</sup>	VC = 1.18 VC = 0.56	[62
39	Cucumis sativus	Cucurbitaceae	Leaf	1.68 <sup>**</sup> 71.1 <sup>**</sup>	BHA = 0.96 TC = 13.2	[27 [25

S. No	Plant name	Family	Part used	IC <sub>50</sub> sample	IC <sub>50</sub> standard	Ref
40	Cucurbita moschata	Cucurbitaceae	Leaf	150 <sup>*</sup>	AA = 120 TC = 50	[75]
41	Cymbopogon citratus	Poaceae	Leaf	1.35*	VE = 0.25	[70]
42	Daniellia oliveri	Leguminosae	Leaf	15.5 <sup>*</sup>	TC = 0.25	[50]
43	Daucus carota	Apiaceae	Aerial	4.61**	BHA = 0.96	[25
44	Ehretia cymosa	Boraginaceae	Leaf	0.47**	GA = 2.09	[53]
45	Emilia coccinea	Asteraceae	Leaf	120 <sup>*</sup>	AA = 120	[75
46	Eugenia caryophyllata	Myrtaceae	Leaf Bud	0.03 <sup>*</sup> 0.02 <sup>*</sup>	AA = 0.03	[47]
47	Eupatorium adenophorum	Asteraceae	Root	22.4 <sup>*</sup> 53.7 <sup>*</sup>	AA = 4.9 RT = 3.3	[73
48	Eupatorium odoratum	Asteraceae	Leaf	0.07**	AA = 0.06	[72
49	Euphorbia hirta	Euphorbiaceae	Leaf	2.5**	VC = 4.5	[81
50	Feretia apodanthera	Rubiaceae	Root	0.053**	VC = 0.048	[43
51	Ficus exasperata	Moraceae	Leaf	0.86*	VE = 0.25	[70
52	Ficus gnaphalocarpa	Moraceae	Leaf	45.3 <sup>*</sup> 44.6 <sup>*</sup>	GA = 48.8 TX = 72.9	[34
53	Ficus sycomorus	Moraceae	Stem	42.0*	VC = 25.0	[82
54	Globimetula oreophila	Loranthaceae	Leaf	0.38**	VC = 0.06	[79
55	Gongronema latifolia	Asclepiadaceae	Leaf	70.0*	VC = 50	[83
56	Grewia carpinifolia	Tiliaceae	Leaf Stem	0.32** 0.39**	AA = 0.31 AA = 0.18	[32
57	Harungana madagascariensis	Hypericaceae	Stem	37.5 <sup>*</sup>	BHT = 16.2	[36
58	Heliotropium indicum	Boraginaceae	Aerial	48.4*	AA = 1.41	[39
59	Hibiscus sabdariffa	Malvaceae	Leaf	0.14*	AA = 0.02	[84
60	Holarrhena floribunda	Apocynaceae	Leaf	7.2*	QT = 2.95	[85
61	Ipomoea asarifolia	Convulvulaceae	Leaf	24.3*	AA = 1.41	[39
62	Irvingia gabonensis	Irvingiaceae	Root Stem	12.4 <sup>*</sup> 25.5 <sup>*</sup>	AA = 4.9 TC = 38.9	[73
63	Justicia secunda	Acanthaceae	Leaf	1.58 μΜ	AA = 2.52	[86
64	Kalanchoe pinnata	Crassulaceae	Leaf	180 <sup>*</sup>	AA = 120	[75
65	Lactuca sativa	Asteraceae	Whole	0.26**	QT = 0.83	[25
66	Landolphia owariensis	Apocynaceae	Root	8.8 <sup>*</sup> 49.1 <sup>*</sup>	AA = 4.9 TC = 38.9	[73
67	Laportea ovalifolia	Urticaceae	Leaf	100*	AA = 150	[75
68	Lasianthera africana	Icacinaceae	Leaf Root	0.30** 0.27**	RT = 0.26	[28
69	Launaea taraxacifolia	Asteraceae	Shoot Leaf	1.94 <sup>**</sup> 1.59 <sup>**</sup>	VC = 1.18 VC = 0.56	[62
70	Lawsonia inermis	Lythraceae	Leaf	3.80*	AA = 7.26	[26
71	Leptadenia hastata	Asclepiadaceae	Leaf	42.3*	GA = 48.8	[35
72	Lycopersicon esculentum	Solanaceae	Fruit	1.16 <sup>**</sup> 1.47 <sup>**</sup>	QT = 0.83	[25

S. No	Plant name	Family	Part used	IC <sub>50</sub> sample	IC <sub>50</sub> standard	Ref
73	Massularia acuminata	Rubiaceae	Leaf	70.0*	VC = 7.59	[87]
74	Mondia whitei	Apocynaceae	Leaf	6.1*	AA = 3.4	[66]
75	Moringa oleifera	Moringaceae	Leaf	0.16*	AA = 0.02	[84]
76	Murraya koenigii	Rutaceae	Leaf	7.35 <sup>*</sup>	TC = 13.2	[27]
77	Nauclea diderrichii	Rubiaceae	Stem	18.12*	AA = 1.41	[39]
78	Nauclea latifolia	Rubiaceae	Leaf	12.9*	AA = 4.9	[73
79	Ocimum basilicum	Lamiaceae	Leaf	1.0*	AA = 9.0	[55]
80	Ocimum gratissimum	Lamiaceae	Leaf Stem	0.14 <sup>*</sup> 8.67 <sup>*</sup>	AA = 0.02 BHA = 3.36	[33] [27]
81	Parinari curatellifolia	Chrysobalanaceae	Leaf	13.5*	VC = 1.98	[88]
82	Parkia biglobosa	Leguminosae	Stem	15.65 <sup>*</sup>	AA = 7.26	[26
83	Phragmanthera capitata	Loranthaceae	Leaf	1.9 <sup>*</sup> 1.0 <sup>*</sup>	BHT = 4.6 VC = 10	[41
84	Piliostigma reticulatum	Fabaceae	Leaf	10.3*	AA = 3.9	[22
85	Piliostigma thonningii	Fabaceae	Leaf	14.7*	AA = 3.9	[22
86	Piper guineense	Piperaceae	Seed	74*	AA = 31.7	[89
87	Psidium guajava	Myrtaceae	Leaf	0.04**	BHA = 0.05	[24
88	Sapium ellipticum	Euphorbiaceae	Stem	0.19**	BHT = 0.11	[90
89	Senna alata	Fabaceae	Leaf	0.59**	VC = 0.067	[79
90	Simarouba glauca	Simaroubaceae	Stem	4.7*	BHT = 5.0	[42
91	Solanum macrocarpon	Solanaceae	Leaf	6.21**	TC = 13.2	[27
92	Spinacia oleracea	Amaranthaceae	Leaf	12.6 <sup>*</sup>	TC = 13.2	[27
93	Spondias purpurea	Anacardiaceae	Stem	8.3*	AA = 11.5	[52
94	Stachytarpheta jamaicensis	Verbenaceae	Leaf	5.0*	AA = 9.0	[51
95	Strophanthus hispidus	Apocynaceae	Root	1.18**	VC = 0.067	[79
96	Telfairia occidentalis	Cucurbitaceae	Leaf	0.16**	AA = 0.02	[84
97	Trichilia catigua	Meliaceae	Stem	30.28*	AA = 20.72	[64
98	Vernonia amygdalina	Asteraceae	Leaf	31.25*	AA = 7.26	[26
99	Vernonia calvoana	Asteraceae	Leaf	1.90 μΜ	AA = 2.0 μM	[49
100	Vernonia cinerea	Asteraceae	Leaf	6.50 <sup>*</sup> 8.0 <sup>*</sup>	GA = 0.62	[30
101	Vernonia migeodii	Asteraceae	Leaf	20.0*	AA = 18.0	[91
102	Vitex doniana	Verbenaceae	Leaf	53.23 <sup>*</sup>	GA = 48.8	[34
103	Xylopia aethiopica	Annonaceae	Fruit	1.04**	VC = 0.067	[79
104	Zingiber officinale	Zingiberaceae	Rhizome	47.0 <sup>*</sup>	AA = 36.4	[92

AA, ascorbic acid; QT, quercetin; RT, rutin; GA, gallic acid; VC, vitamin C; VE, vitamin E; TX, trolox; BHT, butylated hydroxy toluene; BHA, butylated hydroxy anisole; TC, tocopherol.

**Table 1.**Antioxidant activities of selected Nigerian plants.

 $<sup>^*</sup>IC_{50} = \mu gmL^{-1}$ .

 $<sup>^{**}</sup>IC_{50} = mgmL^{-1}$ .

#### 5. Antioxidant activities of crude extracts

The antioxidant efficacies of Nigerian plants were largely evaluated using protocols involving DPPH, ABTS, FRAP, TAC, NO, OH and or H<sub>2</sub>O<sub>2</sub> targets. The DPPH radical scavenging assay is one of the commonly used techniques for quick evaluation of antioxidant capacity. Plant extracts tested for DPPH inhibition have demonstrated interesting efficacies for instance, crude extracts of P. reticulatum (40.10  $\mu$ gmL<sup>-1</sup>) and P. thoninngii (50.94  $\mu gmL^{-1}$ ) showed comparable activity with Ginkgo biloba (EC<sub>50</sub>  $40.72 \,\mu\text{gmL}^{-1}$ ) [22]. Nigerian plants evaluated for antioxidants activity between 2008 and 2012 were reported in 40 publications representing over 166 extracts from 119 plants. These studies showed 29 extracts with effective activity on various free radical targets. However, 15 extracts have comparable antioxidant efficacies to standard antioxidants, while 14 have higher percent (%) inhibition or lower IC<sub>50</sub> values than the standards used. These include stem methanol extract of C. kerstingii (IC<sub>50</sub> 26.27  $\mu$ gmL<sup>-1</sup>, ascorbic acid 33.59  $\mu$ gmL<sup>-1</sup>) [23] and leaf methanol extract of *P*. guajava (IC<sub>50</sub> 0.037 mgmL<sup>-1</sup>, BHA 0.049 mgmL<sup>-1</sup>) [24]. But DPPH inhibition studies on selected vegetable plants showed better effective activity for L. sativa (IC<sub>50</sub> 0.26 mgmL<sup>-1</sup>), Z. officinale (IC<sub>50</sub> 0.29 mgmL<sup>-1</sup>) and C. frutescens (IC<sub>50</sub> 0.67 mgmL<sup>-1</sup>) respectively compared to BHA ( $IC_{50}$  0.96 mgmL<sup>-1</sup>) and quercetin ( $IC_{50}$  0.83 mgmL<sup>-1</sup>) [25]. The activity of *L. inermis* was most profound of the 36 medicinal plants surveyed in Southwestern Nigeria, with lower  $IC_{50}$  of 3.80  $\mu$ gm $L^{-1}$  than ascorbic acid (7.26  $\mu$ gmL<sup>-1</sup>) [26]. Similar evaluations of DPPH inhibition on 15 medicinal plants showed S. oleracea extract with lower IC<sub>50</sub> of 12.6 mgmL<sup>-1</sup>. But S. macrocarpon extract was most effective with IC<sub>50</sub> 6.21 mgmL<sup>-1</sup> lower than  $\alpha$ -tocopherol (13.20 mgmL<sup>-1</sup>) [27].

The analysis of antioxidant efficacies on medicinal plants reported from 2013 to 2017 in 55 publications, involving 211 extracts from 144 plants was carried out. We observed that 70 extracts from 50 plants have exhibited good antioxidant efficacies on various free radical targets with 51 extracts from 53 plants having comparable efficacies to standard antioxidants. However, lower IC<sub>50</sub> or higher percent (%) inhibitions compared to standards were observed with 20 extracts from 17 medicinal plants. The NO inhibition on root extract of L. africana (IC<sub>50</sub> 0.27 mgmL<sup>-1</sup>) compared very well with rutin (IC<sub>50</sub> 0.28 mgmL<sup>-1</sup>) [28]. The DPPH inhibition on *P. guajava* (IC<sub>50</sub> 1.564  $\mu$ gmL<sup>-1</sup>) extract also indicated effective activity compared to ascorbic acid (IC<sub>50</sub> 5.950  $\mu$ gmL<sup>-1</sup>) [29]. Other plant extracts including *V. cinerea* (IC<sub>50</sub> 6.50  $\mu$ gmL<sup>-1</sup>) compared to gallic acid (IC<sub>50</sub> 0.62  $\mu$ gmL<sup>-1</sup>) [30] and K. senegalensis stem bark (IC<sub>50</sub> 95.76  $\mu$ gmL<sup>-1</sup>) with ascorbic acid (223.35  $\mu$ gmL<sup>-1</sup>) indicated effective activity [31]. The inhibition of lipid peroxidation using thiobarbituric acid reactive substances (TBARS) assay on leaf extract of G. carpinifolia was very effective with  $IC_{50}$  of 0.24 mgmL<sup>-1</sup> compared to ascorbic acid ( $IC_{50}$  0.38 mgmL<sup>-1</sup>). Moreover, the ABTS assay indicated 100% inhibitions for both extracts and ascorbic acid [32]. The antioxidant evaluations on two of the most locally utilized vegetable plants such as V. amygdalina and O. gratissimum showed effective inhibitions compared to the standard ascorbic acid [33].

Furthermore, DPPH inhibitions on *S. occidentalis* (IC<sub>50</sub> 42.80 μgmL<sup>-1</sup>) compared to gallic acid (48.77 μgmL<sup>-1</sup>) was effective, but ABTS assay on *F. gnaphalocarpa* (44.63 μgmL<sup>-1</sup>) was more effective than Trolox (72.92 μgmL<sup>-1</sup>) [34]. Similarly, *L. hastata* (IC<sub>50</sub> 42.32 μgmL<sup>-1</sup>) when compared to gallic acid (48.77 μgmL<sup>-1</sup>) and ABTS on *A. senegalensis* (IC<sub>50</sub> 48.98 μgmL<sup>-1</sup>) with Trolox (72.92 μgmL<sup>-1</sup>) have interesting lower IC<sub>50</sub>values [35]. But *H. madagascariensis* exhibited moderate activity [36] while *A. precatorius* [37] and *B. micrantha* [38] have demonstrated effective inhibitions (**Table 1**). The analyses of Nigerian plants in 2018 showed interesting activities with 15 plant extracts from 32 published reports. Plants with moderate DPPH inhibition include *A. hispidum*, *A. laxiflora*, *C. christyanum*, *H. indicum* and

*I. asarifolia* [39]. However, effective inhibitions were observed on root extracts of *D. tripetala* (IC<sub>50</sub> 0.631 μgmL<sup>-1</sup>) and *M. excelsa* (IC<sub>50</sub> 0.194 μgmL<sup>-1</sup>) compared to 4.60 μgmL<sup>-1</sup> ascorbic acid [40]. Similarly, *P. capitata* (27.4 μgmL<sup>-1</sup>) was effective than BHT (56.0 μgmL<sup>-1</sup>) [41], and the evaluation of *S. glauca* stem bark on FRAP (4.70 μgmL<sup>-1</sup>) and NO $^{\bullet}$  (11.90 μgmL<sup>-1</sup>) were effective than 5.0 μgmL<sup>-1</sup> and 18.0 μgmL<sup>-1</sup> of BHT respectively [42]. Lastly, plant crude extracts have demonstrated varying but strong efficacies on different free radical targets which in many cases surpassed standard antioxidants. The report on DPPH inhibition of *F. apodanthera* root bark ethanol extract represents effective activity with IC<sub>50</sub> of 0.053 μgmL<sup>-1</sup> in comparison to vitamin C (0.048 μgmL<sup>-1</sup>) standard [43].

# 6. Chemical composition and antioxidant activity

# 6.1 GC-MS analysis of extracts and evaluation of antioxidant activity

The antioxidant evaluations of Nigerian medicinal plants with determination of chemical composition using gas chromatography-mass spectrometry (GC-MS) have become routine studies. The GC-MS is intended to give insight on the probable chemical entities of volatile components present in the sample extract. Several plants constituents have been analyzed using GC-MS by comparison of compounds' retention times with library of standard chemical entities provided by the National Institute of Standards and Technology (NIST) database imbedded in the instrument. The chemical constituents with low molecular weights such as terpenoids, long chain alkanes, phenolics and fatty acid methyl esters (FAME) are separated and detected by GC-MS. This is perhaps one reason that FAME are prevalent from among plant extracts, but sharp contrast between lipophilic and hydrophilic components are determined by solvent polarity or method of extraction [44].

The GC-MS analyses and evaluation of antioxidants on *B. monandra* hexane extract showed 4-hydroxy-5-methyl-3-propyl-2-hexanone (42.7%) and oleic acid (20%) as major compounds. The DPPH inhibition (IC<sub>50</sub> 5.56  $\mu$ gmL<sup>-1</sup>) with ascorbic acid (IC<sub>50</sub> 30.0  $\mu$ gmL<sup>-1</sup>) showed interesting efficacy, but ethyl acetate extract containing largely oleic acid (40.76%) and hexadecanoic acid (21.75%) was more effective (IC<sub>50</sub> 0.01  $\mu$ gmL<sup>-1</sup>) [45]. The evaluation on A. dentata methanol extract containing hexadecanoic acid (31.6%), phytol (24.6%) and octadecanoic acid (10.56%) was found to be poor. However, the FRAP inhibition showed optimum activity (0.65  $\mu$ molL<sup>-1</sup>) compared to ascorbic acid (2.00  $\mu$ molL<sup>-1</sup>) [46]. The DPPH screening on buds, leaf, root and stem of commonly used spice, E. caryophyllata was reported. The various ethanol extracts showed effective activities IC50 of 0.02, 0.03, 3.66 and 0.99  $\mu$ gmL<sup>-1</sup> respectively, compared to ascorbic acid (IC<sub>50</sub>  $0.03 \, \mu gmL^{-1}$ ) and gallic acid (IC<sub>50</sub>  $0.05 \, \mu gmL^{-1}$ ) standards. This indicated an important response to the DPPH scavenging capacity which have been largely attributed to aromatic phenols, caryophyllene, aromatic esters and ethers [47]. Similar comprehensive study on the leaf, stem bark and root of *C. singueana* was reported. The DPPH, OH and NO<sup>•</sup> showed IC<sub>50</sub> of 1.20, 2.58 and 35.99 μgmL<sup>-1</sup> for DPPH inhibition of stem bark ethanol, root aqueous and leaf ethanol extracts respectively. But the response on OH showed IC<sub>50</sub> of 1.58, 2.05 and 6.47  $\mu$ gmL<sup>-1</sup> respectively, for stem bark ethyl acetate, root aqueous and leaf ethanol extracts. The NO results however, was interesting on leaf aqueous extract (IC<sub>50</sub> 2.81  $\mu gmL^{-1})$  better than the ascorbic acid (IC50 26.28  $\mu gmL^{-1})$  and Trolox (IC50 599.21  $\mu$ gmL<sup>-1</sup>) standards used. The chemical components such as resorcinol (54%) and phytol (23.7%) were largely detected from ethanol extracts of stem bark and leaf respectively [48].

The leaf ethyl acetate extract of *V. calvoana* harvested from the South–South Nigeria contains largely aromatic compounds such as ethyl benzene (22%) and 1,2,3-trimethyl benzene (12.5%). FRAP inhibitions on extract (1.98  $\mu$ M) and ascorbic acid (2.0 μM) were more effective than on DPPH [49]. Although the inhibition on DPPH by plant extracts have been promising but chloroform extract of D. oliveri exudate showed rather poor (IC<sub>50</sub> of 15.5  $\mu$ gmL<sup>-1</sup>) when compared to  $\alpha$ -Tocopherol (0.25  $\mu$ gmL<sup>-1</sup>) [50]. But *S. jamaicensis* methanol extract (IC<sub>50</sub> 5.0  $\mu gmL^{-1}$ ) was more effective than ascorbic acid (IC<sub>50</sub> 9.0  $\mu gmL^{-1}$ ). Compounds such as 3,5-dihydroxy-6-methyl-2,3-dihydro-4H-pyran-4-one (13.7%) and D-arabinitol (13.5%) have been largely identified [51]. It was interesting to note that S. purpurea hexane extract showed effective DPPH inhibition (IC<sub>50</sub> 8.3  $\mu$ gmL<sup>-1</sup>) than ascorbic acid (IC<sub>50</sub> 11.5  $\mu$ gmL<sup>-1</sup>) [52]. The evaluation of *E. cymosa* leaf extracts on ethyl acetate (IC<sub>50</sub> 0.56 mgmL<sup>-1</sup>) and methanol (IC<sub>50</sub> 0.60 mgmL<sup>-1</sup>) extracts justifies effective activity compared to gallic acid (IC<sub>50</sub> 0.47 mgmL $^{-1}$ ). The 2-hexadecycloxirane (34.2%) and methyl linoleate (28.9%) were detected as major components of methanol extract [53]. Lastly, the GC-MS analyses of various plant extracts and antioxidant evaluation have revealed similar pattern of contents and composition in addition to contrasting influences of solvent polarity to radical inhibition efficacies. Nevertheless, interesting antioxidant efficacies were observed.

### 6.2 GC-MS analysis of essential oils and evaluation of antioxidant activity

The essential oils (EO) from Nigerian medicinal plants have been analyzed using the GC-MS and evaluated for antioxidants activity. Because they are mixtures of several constituents containing largely low molecular weights compounds, EO are rapidly analyzed using GC-MS to ascertain their chemical composition. The essential oils (EO) from P. guajava showed that 3, 6-dioxa-2,4,5,7-tetraoctane-2,2,4,4,5,5,7,7-octamethyl (11.7%) and cyclononane (10.7%) are largely identified. DPPH inhibition showed 71.83% comparable to 68.7% of ascorbic acid [54]. The antioxidant efficacy of O. basilicum EO was also interesting (IC<sub>50</sub> of 1.0 μgmL<sup>-1</sup>) probably due to phenolic constituents such as methyl eugenol (15.5%), o-nitrocumene (14.0%) and 2-phenyl-1-hexanol (14.0%) [55]. The EO of common spices such as A. melegueta leaf, C. crepidioides stem bark, O. gratissimum leaf and M. myristica stem bark showed various chemical constituents with interesting antioxidant activity. The EO components largely identified from the four plants are myrtenyl acetate (29.1%), thymol (44%), γ-terpinene (53%) and γ-cardinene (31.1%) respectively. The highest antioxidant activity was found from EO of O. gratissimum (96.4%) which compared to BHA (96.7%). However, other EOs have demonstrated radical scavenging of >50% inhibition [56]. The leaf EO of C. portoricensis was found to contain thymol (9.64%) and  $\beta$ -caryophyllene (9.15%) as main compounds which might have resulted to 75% DPPH inhibition compared to BHT (95%) [57]. Similarly, analysis on M. alternifolius EO that yielded largely tricosane (19.45%) and z-14-nonanosene (13.4%) with interesting efficacy (97.95%) compared to ascorbic acid (97.88%) [58]. This trend of activity demonstrated by the EO was observed in *E. maculata* which contained  $\alpha$ -pinene (8.0%),  $\beta$ -trans-ocimene (8.0%), 1S-α-pinene (7.0%) and cyclofenchene (7.0%) as main components, with the DPPH (IC<sub>50</sub> 8.0  $\mu$ gmL<sup>-1</sup>) and FRAP (10.0  $\mu$ gmL<sup>-1</sup>) inhibition efficacies in comparison to 9.0 and 20.0 μgmL<sup>-1</sup> ascorbic acid respectively [59]. Lastly, GC-MS analyses of EO from Nigerian plants have revealed interesting but similar chemical compounds with some degree of antioxidant efficacies. EO composition containing phenolics moieties and terpenoids have indicated evidence of effective radical inhibitions.

#### 6.3 HPLC analysis of extracts and evaluation of antioxidant activity

The high-performance liquid chromatography (HPLC) has been reported in the analysis of major chemical constituents of plant extracts alongside with the antioxidant activity. The HPLC technique uses reverse phase chromatography because of simplicity, versatility and sensitivity towards separation, purification, quantification and identification of diverse natural products such as plant phenolics, steroids, alkaloids and flavonoids [60]. Hence, the combination of HPLC methods with antioxidants evaluations may provide the needed understanding of antioxidant efficacies of plant extracts. Previous HPLC profiling of ethanol extract of Z. zanthoxyloide showed quercetin, kaempferol and caffeic acid largely quantified. DPPH inhibition (IC<sub>50</sub> 38.58  $\mu$ gmL<sup>-1</sup>) in comparison to ascorbic acid (6.63  $\mu$ gmL<sup>-1</sup>) was poor [61]. Similarly, aqueous extracts of *L. taraxacifolia* (IC<sub>50</sub> 6.59 μgmL<sup>-1</sup>) and C. rubens (IC<sub>50</sub> 6.21  $\mu$ gmL<sup>-1</sup>) were less effective than Trolox (IC<sub>50</sub> 0.51  $\mu$ gmL<sup>-1</sup>). Although methanol extracts of both plants contain gallic acid, caffeic acid, quercetin, rutin, isoquercetin and kaempferol as the main compounds identified, yet the activity was not interesting. But the OH inhibition on aqueous extracts showed rather interesting results with IC<sub>50</sub> 1.94 and 1.09  $\mu$ gmL<sup>-1</sup> in comparison to IC<sub>50</sub> 1.18  $\mu$ gmL<sup>-1</sup> of vitamin C [62].

Although antioxidant activities of plant extracts using DPPH have been established to correlate with phenolics and flavonoids contents [63]. However, many of the plants evaluated for antioxidants activity have no correlation with the number and amounts of phenolics and flavonoids quantified by HPLC. The report on T. catigua ethanol, ethyl acetate, dichloromethane and butanol extracts showed DPPH inhibition with IC<sub>50</sub> of 9.17, 30.28, 42.42 and 76.35  $\mu$ gmL<sup>-1</sup> respectively. These, in comparison to ascorbic acid (20.72  $\mu$ gmL<sup>-1</sup>) indicated poor activity except the ethanol extract with lower IC<sub>50</sub> than the standard. The extract was quantified to be rich in gallic acid, chlorogenic acid rutin and quercetin [64]. Similarly, S. dulcificum contains phenolic acids and flavonoids but demonstrated poor efficacies on DPPH  $(IC_{50} 139.45 \,\mu gmL^{-1})$ , ABTS  $(IC_{50} 135.83 \,\mu gmL^{-1})$ , NO $^{\bullet}$   $(IC_{50} 119.17 \,\mu gmL^{-1})$  and OH (IC<sub>50</sub> 147.65  $\mu$ gmL<sup>-1</sup>) [65]. However, *M. whitei* contains largely caffeic acid with interesting efficacies on NO $^{\bullet}$  (IC<sub>50</sub> 6.1  $\mu$ gmL $^{-1}$ ) and FRAP (IC<sub>50</sub> 5.7  $\mu$ gmL $^{-1}$ ) compared to ascorbic acid (3.4 and 7.0  $\mu$ gmL<sup>-1</sup>) respectively [66]. Similarly, C. papaya seeds protein analyzed using the LC-ESI-DAD-MS with largely ferulic acid in addition to flavonoid sugars, justifies the antioxidant efficacies on DPPH (IC<sub>50</sub> 0.227 mgmL<sup>-1</sup>) and Fe<sup>2+</sup> chelating (IC<sub>50</sub> 0.157 mgmL<sup>-1</sup>) in comparison to ascorbic acid (IC<sub>50</sub>  $0.109 \text{ mgmL}^{-1}$ ) and EDTA (IC<sub>50</sub>  $0.091 \text{ mgmL}^{-1}$ ) respectively [67]. The HPLC quantification of plant extracts have shown similar classes of compounds such as chlorogenic acid, ellagic acid, caffeic acid, gallic acid, p-coumaric acid, apigenin, quercetin, rutin and kaempferol which have been repeatedly found in Nigerian plants. But the antioxidant efficacies observed were not reflective of HPLC quantification. This may indicate that phenolic compounds are quantified at miniature level which can only serve as evidence of qualitative presence in plant extracts.

# 7. Antioxidant activities of isolated compounds

The antioxidant evaluations on isolated compounds from Nigerian medicinal plants are rarely reported. This is probably due to funding problems associated to plant chemistry research in Nigeria, coupled with dysfunctional analytical instruments such as the NMR spectrometer. Most of the published research on isolation and characterization of compounds were carried out abroad. Of the 250 plants

analyzed for antioxidant evaluations, only 28 compounds were isolated from 44 plants together with full spectral characterization. The antioxidant activities of quercetin and quercetin-3-O-rutinoside from *B. monandra* were probably the first report on pure compounds [93]. Since then several isolated compounds were evaluated for antioxidant efficacies and in most cases compared with standard antioxidants. Thus, compounds' efficacy only with IC<sub>50</sub> values of standards are presented in **Table 2**. The analysis of isolated compounds showed that flavonoids and

S. No	Chemical name	Plant	Model	Compd. (IC <sub>50</sub> )	Stand. (IC <sub>50</sub> )	Ref
1	Quercetin	Bauhinia monandra	DPPH	10.64	AA = 12.52	[93]
2	Quercetin-3-O-rutinoside	Bauhinia monandra	DPPH	16.11	AA = 12.52	[93]
3	Isovitexin	Croton zambesicus	DPPH	189.1^	QT = 5.31	[98]
4	Trans-ethyl-3-(3, 4-dihydroxyphenyl acrylate	Aspilia africana	DPPH	14.49	AA = 13.18	[99]
5	p-hydroxy benzaldehyde	Aspilia africana	DPPH	73.50 <sup>*</sup>	VC = 37.5	[102]
6	Tiliroside	Croton gratissimus	DPPH	360.1 <sup>*</sup>	AA = 70.12	[100]
7	Isovitexin	Croton gratissimus	DPPH	211.6*	AA = 70.12	[100]
8	Helichrysoside-3'-methyl ether	Croton zambesicus	DPPH	183.4 <sup>*</sup>	AA = 70.12	[100]
9	Betulin	Parinari curatellifolia	DPPH	>100*	VC = 1.98	[88]
10	β-sitosterol	Parinari curatellifolia	DPPH	>50*	VC = 1.98	[88]
11	Betulinic acid	Parinari curatellifolia	DPPH	>100*	VC = 1.98	[88]
12	4-(3',3-dihydroxyl- 1-mercaptopropyl) phenyl-glucosylpyranoside	Massularia acuminata	DPPH	75 <sup>*</sup>	VC = 7.59	[87]
13	Agathisflavone	Anacardium occidentale	DPPH	366.4 <sup>*</sup>	AA = 4.57	[74]
14	Quercetin-3-O-rutinoside/ rhamnoside	Anacardium occidentale	DPPH	0.96*	AA = 4.57	[74]
15	Rosmarinic acid	Solenostemon monostachyus	DPPH	4.99*	QT = 2.32	[97]
16	Methyl rosmarinate	Solenostemon monostachyus	DPPH	5.97 <sup>*</sup>	QT = 2.32	[97]
17	Caffeic acid	Solenostemon monostachyus	DPPH	3.03*	QT = 2.32	[97]
18	Methyl caffeate	Solenostemon monostachyus	DPPH	13.41*	QT = 2.32	[97]
19	Apigenin	Solenostemon monostachyus	DPPH	26.67 <sup>*</sup>	QT = 2.32	[97]
20	Luteolin	Solenostemon monostachyus	DPPH	5.35 <sup>*</sup>	QT = 2.32	[97]
21	Apigenin glucuronide	Solenostemon monostachyus	DPPH	185.89 <sup>*</sup>	QT = 2.32	[97]
22	Epicatechin	Chrysophyllum albidum	DPPH	19.02 <sup>^</sup>	GA = 12.82	[101]

S. No	Chemical name	Plant	Model	Compd. (IC <sub>50</sub> )	Stand. (IC <sub>50</sub> )	Ref.
23	Epigallocatechin	Chrysophyllum albidum	DPPH	15.88 <sup>^</sup>	GA = 12.82	[101]
24	Procyanidin B5	Chrysophyllum albidum	DPPH	8.80^	GA = 12.82	[101]
25	Kaempferol-3-O- rutinoside	Holarrhena floribunda	FRAP	394.8 <sup>*</sup>	QT = 2.95	[85]
26	Quercetin-3-O-glucoside	Holarrhena floribunda	LPI FRAP	10.4 <sup>*</sup> 1649.4 <sup>*</sup>	QT = 2.95	[85]
27	Kaempferol-3-O-glucoside	Holarrhena floribunda	FRAP	337.5*	QT = 2.95	[85]
28	Quercetin-3-O-glucoside/ galactoside mixture (1: 1)	Holarrhena floribunda	LPI FRAP	9.8 <sup>*</sup> 1589.9 <sup>*</sup>	QT = 2.95	[85]
29	Quercetin	Cassia sieberiana	DPPH ABTS	1.58 <sup>#</sup> 0.81 <sup>#</sup>	AA = 2.44 TX = 0.81	[96]
30	Kaempferol	Cassia sieberiana	DPPH	7.75 <sup>#</sup>	AA = 2.44	[96]
31	Dihydrokaempferol	Cassia sieberiana	DPPH	82.93#	AA = 2.44	[96]
32	Piceatannol	Cassia sieberiana	DPPH	3.96#	AA = 2.44	[96]
33	(–)-Catechin	Alchornea floribunda	DPPH H <sub>2</sub> O <sub>2</sub>	88 <sup>*</sup> 13 <sup>*</sup>	AA = 6 AA = 8	[95]
34	(+)-epicatechin	Alchornea floribunda	DPPH H <sub>2</sub> O <sub>2</sub>	40 <sup>*</sup> 10 <sup>*</sup>	AA = 6 AA = 8	[95]
35	(–)-epicatechin	Alchornea floribunda	DPPH H <sub>2</sub> O <sub>2</sub>	10 <sup>*</sup> 8 <sup>*</sup>	AA = 6 AA = 8	[95]
36	(2R,3R)-dihydroquercetin	Alchornea floribunda	DPPH H <sub>2</sub> O <sub>2</sub>	46 <sup>*</sup> 18 <sup>*</sup>	AA = 6 AA = 8	[95]
37	Catechin	Annona senegalensis	DPPH Fe(II)	0.03** 1.29**	AA = 0.01 EDTA = 0.05	[15]

DPPH, 2,2-diphenyl-1-picrylhydrazyl radical; ABTS, 2,2'-azinobis (3-ethylbenzothiazoline-6-sulfonic acid; FRAP, ferric reducing antioxidant power; TAC, total antioxidant capacity; LPI, lipid peroxidation inhibition; NO, nitric oxide assay;  $H_2O_2$ , hydrogen peroxide assay; AA, ascorbic acid; QT, quercetin; RT, rutin; GA, gallic acid; VC, vitamin C; VE, vitamin E; TX, trolox; EDTA: ethylenediaminetetraacetic acid.

Table 2.

Antioxidant activity of isolated compounds of Nigerian plants.

flavonoids glycosides constitute major classes of antioxidants reported. Catechin isolated from A. senegalensis had effective DPPH inhibition (IC<sub>50</sub> 0.03 mgmL<sup>-1</sup>) and Fe<sup>2+</sup> chelating activity (1.29 mgmL<sup>-1</sup>) when compared to ascorbic acid (0.01 mgmL<sup>-1</sup>) and EDTA (IC<sub>50</sub> 0.05 mgmL<sup>-1</sup>) respectively [94]. The evaluation on H<sub>2</sub>O<sub>2</sub> inhibition by (–)-epicatechin isolated from A. floribunda showed effective activity with equal strength as standard ascorbic acid (IC<sub>50</sub> 8.0 µgmL<sup>-1</sup>) [95]. Similarly, the ABTS inhibition by quercetin isolated from C. sieberiana has resulted to effective activity of equal strength to Tocopherol (0.81 mM) [96]. The DPPH inhibition by caffeic acid (IC<sub>50</sub> 3.03 µgmL<sup>-1</sup>) from S. monostachys is another effective activity comparable to quercetin standard (IC<sub>50</sub> 2.32 µgmL<sup>-1</sup>) [97]. However, the most outstanding DPPH inhibition was recorded on quercetin-3-O-rutinoside/rhamnoside isolated from A. occidentalis. The 1:1 mixture of flavonoid glycoside exhibited

 $<sup>^{\</sup>hat{}}IC_{50}=\mu M.$ 

 $<sup>^{\#}</sup>IC_{50} = mM.$  $^{*}IC_{50} = \mu gmL^{-1}.$  $^{**}IC_{50} = mgmL^{-1}.$ 

 $IC_{50}$  0.96 μgm $L^{-1}$  less than ascorbic acid ( $IC_{50}$  4.57 μgm $L^{-1}$ ). [74]. But generally, the antioxidant efficacies of isolated compounds from Nigerian plants are not interesting. Out of the 28 compounds isolated from 44 plants only 7 compounds from 6 plants exhibited the efficacies with strength of standard antioxidants.

#### 8. Conclusion

Analysis of antioxidant efficacies of Nigerian medicinal plants reported from 1998 to 2018 was carried out. The aim was to provide evidence for effective antioxidants. Our findings have shown the enormous potentials of Nigerian plants as sources of natural antioxidants. We have observed various crude extracts obtained mainly from polar solvents with antioxidant efficacies better than standard compounds. Such preponderance of evidence indicated by broad spectrum of free radical and non-free radical inhibitions has defined the comparable strength of plant extracts to standard antioxidants. Nigerian plants have the capacity to protect or inhibit damage induced by free radical species. This study attempts to provide insights on the strength of antioxidant efficacies of plant extracts comparable to standard antioxidants. However, it is recommended that comprehensive approach to plant bioactive research must be adopted in search of antioxidants to avoid replication of studies especially on certain species. There is need for collaboration among Nigerian scientist working in related areas to enhance on the scope of research questions and improve on the quality of research output.

# Acknowledgements

Authors acknowledged the plant taxonomist, Umar Shehu Gallah of National Research Institute for Chemical Technology (NARICT) Zaria, for providing the local (Hausa) names of plants. We are grateful to Ahmadu Bello University, Zaria for providing some of the facilities used during the study.



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