IN VITRO ANTIOXIDANT ACTIVITY OF N-BUTANOL EXTRACT OF CURCUMA LONGA AND ITS POTENTIAL TO PROTECT ERYTHROCYTES MEMBRANE AGAINST OSMOTIC-INDUCED HAEMOLYSIS

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ABSTRACT
Antioxidant activity of Curcuma longa rhizomes (Turmeric) has been widely investigated in Asia. This work evaluated the antioxidant potential of Nigerian variety of Curcuma longa with the main hypothesis that variation in plant phytochemicals in type and/or amount is a function of geographical location/soil type. The experimental models used for the antioxidant studies include: iron (III) reducing capacity, total antioxidant capacity, 1,1-diphenyl-2-picrylhydrazyl (DPPH) scavenging assay and in vitro inhibition of osmotic-induced haemolysis. The extract was found to reduce osmotic pressure-induced haemolysis of mammalian erythrocytes in a dose-dependent manner and also gave high total antioxidant capacity. At 1000 µg/mL concentration, its DPPH radical scavenging ability compared favourably well with those of the standard compounds - ascorbic and garlic acids. The implications of these findings in the usage of Curcuma longa as food additives and ethnomedicinal practice in Nigeria are discussed.

Keywords: Curcuma longa, In vitro, Antioxidant potential, mammalian erythrocytes.

INTRODUCTION
Oxidative stress occurs in cells and tissues of organisms wherever oxidation and reduction processes abound. This has been attributed to production of reactive oxygen species (ROS) beyond the ability of the system to neutralize them (Halliwell, 1996). Oxidative stress has been incriminated in the pathogenesis of several disease conditions such as cancer, cardiovascular, liver injury, myocardial infarction, Alzheimer’s disease, Parkinsonism as well as in the ageing process (Grzegorczyk et al., 2007). Continuous exposure to synthetic compounds, which is characteristic of orthodox medicine, could lead to increase in the amount of free radicals in the body and hence cause different toxicities. This, among other factors of toxicity and cost, has caused significant increase in research interests towards medicinal plant research in recent years.

A lot of scientific researches has focused on screening plants and herbs for their potential antioxidant properties (Motterlini, 2000) because active principles from these plants are thought to play important roles in preventing diseases caused by oxidative stress. Again, since about 80% of the world’s population is thought to rely on traditional medicines, which is mainly the use of plant extracts, for some aspect of their primary health care needs (WHO, 1993) and that traditional medicine may be the surest means to achieving total health care coverage in the world (WIPO, 1998), it seems justifiable that the search for antioxidants potentials of plants should continue.

Curcuma longa which is also known as ‘turmeric’, is a herbaceous perennial plant belonging to the ginger family, Zingiberaceae (Figures 1). It is extensively cultivated in China, India and other countries with tropical climate (Paramasivam et al., 2009) including Nigeria where it is commonly called ‘Zabibi’ or ‘Magina’ among Hausa speaking Nigerians. It is extensively used as spice, food preservative and colouring material in India, China and South-East Asia (Neeta et al., 2007). It has been used in traditional medicine as a household remedy for various diseases, including biliary disorders, anorexia, cough, diabetic wounds, hepatic disorders, rheumatism and sinusitis (Araujo and Leon, 2001). Literature review revealed that curcumin (diferuloylmethane), the main yellow bioactive component of turmeric, possesses wide spectrum of biological properties including anti-inflammatory (Motterlini et al., 2000; Jureka, 2009), antioxidant (Motterlini et al., 2000) , anticarcinogenic and antimutagenic (Ammon and Wahl, 1991;Regunathan and Panneerselvam, 2007), antilulcer (Natchaya et al., 2004), and antiprotozoal (Mazumdar et al., 1995; Reddy et al., 2005), hypotensive and hypocholesteremic activities (Chattopadhyay et al., 2004). Of all these antioxidant studies, almost all were conducted with curcumin/curcumin-rich ethanolic extracts obtained from Curcuma longa cultivated in Asian countries with virtually few studies on extracts from Nigerian cultivars. This is important when one considers that variation in phytochemicals such as phenols and flavonoids has been established among same plant cultivated in different geographical locations (Ndhlala et al, 2014). There is therefore, the need to investigate the antioxidant potentials of Turmeric cultivated in Nigeria where there is paucity of literature on the antioxidant potentials of this rhizome and where a lot of emphasis has been given to its role in dyeing of various local fabrics. This paper investigated the antioxidant properties of n-butanol extract of Nigerian variety of Curcuma longa and determined the ability of the extract to protect, in vitro, mammalian erythrocytes against lysis induced by osmotic pressure. In Nigeria, the plant is locally cultivated in the Northern parts of the country and the colour pigment is used mainly in traditional dyeing industry.

In Vitro Antioxidant Activity of N-Butanol Extract Of Curcuma Longa and Its Potential to Protect Erythrocytes Membrane Against Osmotic-Induced Haemolysis
**Materials and Methods**

**Plant Material**
Rhizomes of turmeric (*Curcuma longa*) were obtained from Kaura village in Kaura Local Government Area of Kaduna State, Nigeria and were subsequently dried in a water bath. The crude extract, a mixture of n-butanol and water, was used for the experiments.

**Blood Sample**
Fresh blood samples were obtained from healthy sheep, cattle and goat animal after slaughter in Kakuri Abattoir, Kaduna State, Nigeria.

**Chemicals**
DPPH (1,1-diphenyl-2-picrylhydrazyl), Trichloroacetic acid (TCA), Ammonium Molybdate, Ascorbic acid, Gallic acid were purchased from Sigma Co. U.S.A. All other chemicals used were of analytical grade.

**Methods**

**Extraction**
Fresh rhizomes (200g) of turmeric were pounded to puree and extracted with 200ml of petroleum ether at 40°C using soxhlet apparatus, after which residual puree was re-extracted with 200ml of n-butanol at 60°C. The extract was concentrated by removing the solvent with the aid of a rotary evaporator (NYC R-205D) and subsequently dried in a water bath. The crude extract, a brownish gummy exudates, was then stored in the refrigerator until required for the experiment.

**Preparation of erythrocyte suspension**
Fresh blood samples from sheep, cattle and goat were collected in separate EDTA bottles from Kakuri abattoir (in Kaduna metropolis, Nigeria). After swirling to mix, 2ml of each blood sample was transferred into centrifuge tubes using a sterile glass pipette. Five (5) ml of phosphate buffer saline (PBS), pH 7.4, was thereafter added to each tube and centrifuged at 2000 r.p.m. for 10 min. The supernatant was discarded and 5 ml of the buffer (PBS) was added to the washed erythrocytes pellet. This process was repeated twice (the supernatant for the third washing is usually free of haemoglobin). The washed cells (RBCs) were finally made ready for use in 4 ml of PBS buffer. The cell density of the suspension was determined using haemocytometer.

**Fragility Test**
The method described by Tizard et al. (1978) was adopted. Briefly, to each set of test tubes containing 5 ml of different concentrations (0.00-0.85%) of saline, 10⁶ washed RBCs was added. The mixture was incubated in water bath at 37°C for 30 minutes, centrifuged at 2000 r.p.m. for 10 minutes, the supernatant fluid removed and its haemoglobin content estimated spectrophotometrically at 540 nm against 0.85% buffered saline (the blank). Appropriate positive and negative controls were employed at the same time. Percentage haemolysis (mean ± standard deviation) was calculated as shown below:

\[
\% \text{ Haemolysis} = \left( \frac{\text{Absorbance of individual fluid} - \text{Absorbance with 100% haemolysis}}{\text{Absorbance with 100% haemolysis}} \right) \times 100
\]

**Study on the effect of aqueous butanol extract of *C. longa* on haemolysis induced by osmotic-pressure**

From the fragility curve, the effective concentration that will cause 50% haemolysis (EC₅₀) was extrapolated. This EC₅₀ was used to incubate the washed RBCs of each animal with varying concentrations of aqueous butanol extract of *C. longa*. The test tubes were set as follows:

i. **Positive controls**: tubes contained 10⁵ washed RBCs in distilled water.

ii. **Negative controls**: tubes contained 10⁵ washed RBCs in phosphate buffer, pH 7.4.

iii. **Extract-treated groups**: tubes contained 1 ml of the extract solution with 10⁴ of each washed RBCs. The total volume was made up to 5ml by adding phosphate buffer, pH 7.4, to give different active concentrations of 100, 200, 400, 800 and 1600 µg/mL. The extract solutions were prepared with the same buffer, and each experiment was carried out in triplicate.

The incubation was done at 37°C for 60mins at the end of which the tubes were centrifuged at 2000 r.p.m. for 10 mins and the haemoglobin content of the supernatant was determined spectrophotometrically (540 nm) as described above. The ameliorative potential of the extract was expressed as Percent retardation (mean ± standard deviation) and was calculated using the formula below:

\[
\% \text{ Retardation} = \left( \frac{A - B}{A} \right) \times 100
\]

Where A is haemolysis by EC₅₀ saline concentration while B is haemolysis caused by the concurrent addition of extract.

**Determination of Total antioxidant capacity**
The antioxidant activity of butanol extract of *C. longa* was evaluated by the phosphomolybdate method of Prieto et al. (1999). The assay measures the reduction of Mo(VI) – Mo (V) by the extract and the subsequent formation of a green phosphate/Mo (V) complex at acid pH. Briefly, a 0.3 ml extract solution was combined with 3 ml of reagent solution (0.6 M sulfuric acid, 4 mM ammonium molybdate and 28mM sodium phosphate). The tubes containing the reaction solution were incubated at 95°C for 90 min. The absorbance of the solution at 695 nm was measured after cooling to room temperature using spectrophotometer (Jenway UV/Vis 6505) against blank (0.3 ml butanol extract in phosphate buffer). The antioxidant activity is expressed as the number of gram equivalents of ascorbic acid.

**Reducing Power assay**
Different concentrations aqueous butanol extract of *C. longa* (31.25-1000 µg/mL) in 1ml distilled water with 2.5ml phosphate buffer.
buffer (pH 6.6, 0.2M) and 2.5ml of potassium ferricyanide [1% K₃Fe(CN)₆] were incubated at 50°C for 20mins. A portion (2.5 ml) of 10 % trichloroacetic acid (TCA) was added to the mixture and then centrifuged at 3000 r.p.m. for 10 min. The upper layer of the solution (2.5 ml) was mixed with 2.5 ml distilled water and 0.5 ml of iron (II) chloride (1% FeCl₃). The absorbance was measured at 700 nm using spectrophotometer (Jenway UV/Vis 6505), increased absorbance of the reaction mixture indicated increased reducing power. Ascorbic acid and garlic acid were used as standards. Phosphate buffer, pH 6.6 was used as blank solution. Each test was carried out on triplicate and absorbances of the final reaction mixtures were expressed as mean ± standard deviation. This procedure was described by Saba et al., (2008).

**DPPH radical scavenging activity**

The determination of the free radical scavenging activity of butanol extract of *C. longa* using DPPH (1,1- diphenyl - 2-picylhydrazy) as assay as described Blois (1958) with slight modification.

Various concentrations (1000- 31.25µg/mL) of butanol extract in methanol were prepared. 1ml of 0.3mM DPPH in methanol was added to 2ml solution of the extract and standards, and allowed to stand at room temperature in a temperature in a dark chamber for 30mins. The change from deep violet to yellow was then measured at 518nm on a spectrophotometer (Jenway UV/Vis 6505). The decrease in absorbance was then converted to percentage scavenging activity (% SA) using the formula;

\[
\text{% Scavening Activity} = \left(\frac{\text{Absorbance of sample} - \text{Absorbance of blank}}{\text{Absorbance of control}}\right) \times 100
\]

Blank is methanol (1ml) plus sample solution (2.0ml). Negative control is DPPH solution (1ml, 0.25mM) and methanol (2ml), ascorbic acid and garlic acid were used as standards. The scavenging reaction between (DPPH) and an antioxidant (H- A) can be written as;

\[
(DPPH) + (H-A) \rightarrow DPPH- H + (A) \rightarrow \text{yellow}
\]

**Results and Discussion**

Herbs and plants have been used traditionally as food and as sources of medicine and thus providing strong link between nutrition and chemotherapy. In *C. longa*, such connection is easily seen as the rhizome of the plant is use by local dwellers as part of their staple food and also in the management of local ailments (Narasinga, 2003; Ishita et al., 2004). The aim of this study was to investigate the potential antioxidant properties of Nigerian turmeric and to assess the ability of the extract to protect mammalian erythrocytes from haemolysis caused by osmotic pressure.

The n-butanol extract showed high DPPH scavenging activity at 1000 µg/ml comparable to standard Ascorbic and Gallic acids. It means that the extract exhibited free radical scavenging activity but it is not known, by this work, if the antioxidant potential in Nigerian Extract will surpass that seen the Asian Turmeric (Fig. 3). The reducing capacity of compounds could serve as indicator of potential antioxidant property (Meir et al., 1995). However, the n-butanol extract of *C. longa* revealed poor reducing power, which indicates low ability to reduce FeCl₃ to FeCl₂ (Fig. 4). This could be attributed to poor solubility of the extract’s active components in aqueous solvent. The total antioxidant capacity of the extract based on the phosphomolybdate protocol exhibits high values with increasing concentration in Ascorbic acid equivalents (Table 1). This corroborate the DPPH scavenging activity in this study and agrees with observation made on turmeric extract from Asia in which curcumin has been identified as the active principle (Narasinga, 2003; Ishita et al., 2004). The high antioxidant activity may be due to the presence of phytochemicals such as phenols, flavonoids and anthocyanins (Aderogba et al., 2004). The antioxidant mechanism of curcumin is attributed to its unique conjugated structure, which includes two methoxylated phenols and enol form of β- diketone; the structure shows typical radical-trapping ability as a chain-breaking antioxidant (Ishita et al., 2004).

Fragility test conducted showed that the n-butanol extract of Curcuma longa lowered haemolysis of sheep, goat and bovine erythrocytes under saline-induced stress, thus suggesting a degree of protection of the erythrocytes against damage of their membrane integrity. Bovine erythrocyte was most protected from this stressor. This could be partly attributed to the variation in composition among various erythrocytes membranes an observation that has already been made (Lanham and Godfrey, 1970). This implies that the extract may possess principles that could protect the erythrocytes from damages due to stressors, at least in vitro. Similar observation has been made by Neeta and Ramtej (2007) using aflatoxin-induced haemolysis model.

Our results showed that Nigerian turmeric also showed tremendous antioxidant properties but whether these properties are due to curcumin or other cucurminoids is yet to be ascertained. This finding will be useful to both food and drug industries. Curcumin is known to be a component ingredient of curry powder. But the component curcuminoids may vary in chemical structure and is possible that the chemical and colour characteristics as well as the functional properties will vary (Paramasivam et al., 2009). If this study is consolidated with further researches, it may provide strong basis to discontinue importation of food condiment so that production and local utilization will be encouraged.
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Fig. 4. Reducing power assay of butanol of Curcuma longa

Fig. 5. Retardation of osmotic induced haemolysis of sheep erythrocytes by aqueous butanol extract of C. longa

Table 1: Total antioxidant capacity (TAC) of n- butanol extract of C. longa.

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<thead>
<tr>
<th>Concentration (µg/mL)</th>
<th>TAC of extract (Equivalent to ascorbic acid)</th>
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<tr>
<td>1000.00</td>
<td>370</td>
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<td>500.00</td>
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<td>31.25</td>
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Fig. 6. Retardation of osmotic induced haemolysis of aqueous butanol extract on bovine erythrocytes

Fig. 7. Retardation of osmotic induced haemolysis on goat erythrocytes by aqueous butanol extract

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REFERENCES


