Phytotherapeutic Potentials of Synedrella Nodiflora: In-Vitro Quantification of Phytochemical Constituents, Antioxidant Capacities and Skin Enzymes Inhibiting Activities

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Abstract

Synedrella nodiflora is a useful medicinal plant that has been evaluated for its application in treatment of several ailments in sub-Saharan Africa and Bangladesh since time immemorial. In the present study, the methanol extract of the plant was investigated in-vitro for the presence of some essential phytochemicals, its antioxidant capacities, activity inhibition of skin degenerating enzymes of elastase and tyrosinase as well as its anti-lipid peroxidation activity. Determination of the total antioxidant capacities of the extract was achieved through evaluation of oxygen radical absorbance capacity (ORAC), ferric ion reducing antioxidant potential (FRAP) and iron-II induced inhibition of lipid peroxidation (LPO) assays. The quantitative analysis outcome of this study showed the relative abundance and concentration of flavanols, alkaloids, flavonols, phenolics, tannins, proanthocyanidins and saponin in Synedrella nodiflora methanol extract with the relative abundance (%) of the phytochemicals is in the order of polyphenols (19.09)>Saponin (18.61)>flavonol (18.00)> proanthocyanidins (14.52) > flavanol (10.35)>Alkaloid (10.07)>tannins (9.36).

The outcome of this study showed the ability of the plant extracts to scavenge free radicals and to inhibit degradation of lipids due to oxidative damage. Thus, S. nodiflora may be used in the treatment and management of oxidative stress and its related diseases. It was also observed that the plant extract possess a mild in-vitro tyrosinase and elastase inhibiting activities which implies that the plant may find applications in cosmetic preparations for skin depigmentation and anti-wrinkle agents in pharmaceutical formulations.

Keywords: Tyrosinase, Elastase, Synedrella Nodiflora, Phytochemicals; Phytotherapy, Antioxidant Phytochemicals, Phytotherapy, Antioxidant

Introduction

Many plants in nature have been assessed for their medicinal values. The background knowledge of the chemical constituents of a plant is highly desirable not only for the discovery of therapeutic agents but also because such information may be of value in disclosing new sources of economic materials like tannins, polyphenols, gum, oils, and other precursors for the synthesis of complex chemical substances [1]. Phytochemical screening enables tracing of the medicinal value of plants constituent and some biochemical substances that generate a definite physiological action on the human body [2,3]. Some important of the bioactive compounds in plants are the alkaloids, flavanol, flavonoids, tannins, alkaloids saponins, glycosides, cardenolides, proanthocyanidins and polyphenolic compounds [4-7].

Medicinal plants have found applications in treatments of several ailments and disorders like rheumatism, muscle swelling, inflammation, pains, diabetes, high blood pressure, oxidative stress,
cancer etc. Such plants include *Citrus coloecynthis*, *Acacia arabica*, *Ocimum gratissimum*, *Azadirachta indica*, *Phyllanthus amarus*, *Eclipta alba* and *Vernonia amygdalina* [8-13]. The ethnopharmaceutical and ethnomedicinal applications of these plants are due to their varying degree of bioactive phytochemical constituents as well as their characteristic antioxidant properties [14-17]. The enzymes inhibiting properties of medicinal plants have been found exciting, adding more value to phytotherapy and making plants an excellent alternative to conventional synthetic chemicals in biomedicine. The alpha amylase and alpha glucosidase inhibition activities of some plants have been reported [18, 19]. Recent independent researches by [20,21,22] have shown the inhibitory effects of some plant materials on the enzyme tyrosinase present in the skin and they confirmed the possible application of such plant material in solving skin hyperpigmentation and premature skin aging problems.

*Synedrella nodiflora* is from the kingdom Plantae, infrakingdom of Streptophyta, subkingdom of Viridiplantae, division of Tracheophyta, superdivision of Embryophyta, subdivision of Spermatophyta, class of Magnoliopsida, order of Asterales, superorder of Asteranae, family of Asteraceae, genus of Synedrella Gaertn., and species of *Synedrella nodiflora* [23]. *Synedrella nodiflora* is a branched ephemeral herb that can grow up to 80 cm tall. It is characterized with a shallow root system which is usually strongly branched. The lower part of the stem may root at the nodes, especially in moist conditions while the leaves exist in opposite pairs and are usually 4-9 cm long. The presence of various bioactive constituents in the leaf extract of *Synedrella nodiflora* was reported after screening studies by [24,25]. Solvent extracts of *Synedrella nodiflora* have been shown to possess flavonoids, alkaloids, glycosides, steroids, tannins, saponins and phytosterols, triterpenoids gums and reducing sugars [26-29]. The plant is essential for the treatment of various diseases and its leaves are eaten as a vegetable by some livestock and human. It also possesses sex hormone activity [30]. The leaves can be used as Pregnant Mare Serum Gonadotrophin supplier in animal husbandry and to improve reproductive parameters in female animals [31]. Literatures on the skin enzymes inhibitory activities of the plant are still limited according to SciFinder and Dictionary of Natural products. This research is therefore directed at investigating the inhibitory actions of *Synedrella nodiflora* on the degenerating actions of enzymes present in the surface of the skin (tyrosinase and elastase) as well as its phytochemical constituents and antioxidant properties to support and validate its acclaimed ethnopharmacological application.

**Materials and Methods**

**Chemicals and Reagent**

The reagents and standards used for this work were all of analytical grades with high percentage purity, secured from Sigma-Alchdrich, Inc (St. Louis, MO, USA) and Pascal scientific Ltd (Alaure, Nigeria).

**Methods**

**Samples Collection and Preparation**

Aerial part of *S. nodiflora* was sourced from a local farm in Ikere-Ekiti (7.4991° N, 5.2319° E), Ekiti State Nigeria. The plant was identified by the Herbarium curator at the Department of Biological Science Koladaisi University, Ibadan, Nigeria, where a voucher specimen number Kdu/IB/034 was assigned to the *S. nodiflora*. The obtained plant materials were prepared by washing them with distilled water and drying at room temperature for two weeks. The plant parts were crushed separately using pestle and mortar. Thereafter, the crushed parts were mixed together, pulverized by an electric blender into a homogenized powder, weighed and stored in different airtight sterile sample bottles pending analysis.

**Extraction**

The powdered plant material was extracted with methanol. Approximately 50g of the powdered material was soaked in 1000 mL of the solvent in a vial for 72 hours for cold extraction. The extract was filtered and concentrated at 50 °C using rotary evaporator. The concentrate was stored in an air tight sample vial pending analysis.

**Quantitative Phytochemical Analysis**

**Determination of Total Phenol**

The total phenolic content of the aerial part of the plant was determined using the Folin-Ciocalteu method [32]. The procedure involved adding both distilled water and Folin-Ciocalteu reagent to a 125 μL of the solvent extract. The mixture was allowed to stand for 6 min before the adding sodium carbonate solution (7.0% w/v). Thereafter, the mixture was allowed to stand for 90 min after which the absorbance was read at 760 nm on a SpectrumLab70 spectrophotometer and the result were expressed in terms of Gallic acid in mg/mL of extract.

**Determination of Saponin**

The saponin concentration of the plant was determined using spectrophotometric method as described by [33]. In this method, 2.0 g of the extract was weighed into a beaker containing isobutyl alcohol (butan-2-ol) was added. The mixture was stirred and filtered through No 1 Whatman filter paper into a beaker containing 40% magnesium carbonate solution. Thereafter, 1 mL of the solution was transferred into a 500 mL volumetric flask 2 mL iron (III) chloride (FeCl3) solution was added to the solution and made up to mark with distilled water. The resulting solution was allowed to stand for 40 min for color development and absorbance was read at 380 nm on a SpectrumLab70 spectrophotometer.
Determination of Tannin

The total tannin content was assessed by the experimental protocol of [18] with slight modifications. The tannin determination was carried out as follows. Approximately 0.5 g of the extract was diluted with 90% ethanol. 0.1 mL of the diluted sample was measured and added to 2 mL of Folin Ciocalteu reagent. After 10 min, 7.5 mL of sodium carbonate (7%) solution was added and the mixture incubated for 2 hours. The absorbance of this mixture was measured at 760 nm and the tannin content was estimated using tannic acid (TA) curve as the standard.

Determination of Alkaloid

Alkaloid content of the plant’s solvent extract was determined by weighing 5.0 g portion of the solvent extract into a beaker and 250 mL of 10% acetic acid in ethanol was added and allowed to stand for 5 min. The mixture was filtered and the extract was concentrated on a water bath to one fifth of the original volume. Ammonium hydroxide solution was then added in drops to the concentrated extract until the precipitation was completed [19]. The precipitate was collected, washed several times with dilute ammonium hydroxide and filtered. The residue was dried in a desiccator and weighed.

Flavanol Content

The flavanol content of the Solanum nodiflora was determined spectrophotometrically in accordance to the modified method of Popoola et al. [7]. A 25 mL 0.05% 4 dimethylaminocinnam-aldehyde (DMACA) solution prepared by dissolving DMACA in 8% HCl prepared in methanol was added to 50 mg of the solvent extract. The mixture was allowed to stand for 30 min and the absorbance was read on a SpectrumLab70 spectrophotometer at 640 nm. The result was expressed in terms of catechin equivalents in mg CE/mg of extract. The determination was carried out in triplicate.

Determination of Flavonol

The flavonol content of the plant extract was determined by the method described (2008) with slight modifications. 2 mL of AlCl3 prepared in ethanol and 3 mL of 50 g/L sodium acetate solution were added to 2 mL of the extracted sample in a test tube and mixed. The mixture was incubated for three hours (3 hrs) at 20°C. Series of stock solution of 20, 40, 60, 80, and 100 μg/mL were thereafter prepared. The absorbance of the solutions was measured at 440 nm against a blank at 593 nm using a SpectrumLab70 spectrophotometer at 640 nm. The resulting mixture was pre incubated for 5 min at 37°C. Fifty micro liters (50μL) of peroxide radical AAPH solution was added immediately and decaying of fluorescence was scanned for 35 min at 1 min interval at 37°C [35]. The decay in fluorescence was

Determination of Proanthocyanidins

The total proanthocyanidin content of the plant material was vortex mixed with 3 mL of vanillin (4%) prepared in methanol and 1.5 mL of hydrochloric acid. The mixture was allowed to stand for 15 min at ambient temperature. Thereafter, the absorbance was read at 500 nm. The total proanthocyanidin content was expressed in terms of catechin (CE mg/g).

Antioxidant Activities Determination

Ferric Reducing Antioxidants Potential (FRAP) Assay

The FRAP assay was performed according to a modified method of [19]. 1 mL of the extracted sample (1.0 mg/mL) was mixed with 5 mL FRAP reagent in a test tube and vortex mixed. Blank samples were prepared for the methanol extract. The sample and blank were both incubated in water bath for 30 minutes at 37°C. The absorbance of the samples was measured against the blank at 593 nm. Series of stock solution with concentration of 20, 40, 60, 80, and 100 μg/mL were prepared and using aqueous solution of FeSO₄·7H₂O as standard curve.

Inhibition Lipid Peroxidation Assay (LPO)

For the determination of the lipid peroxidation inhibition assay, a little modification was made on the adopted method of Snijman [34]. In this method, both the plant extract and positive standards were prepared by incubating a mixture of 50μL anti-oxidative solutions (in different test tubes) corresponding to the extract or the positive standard and 300μL microsomes at 37°C for 30 min in a shaking water bath. 100 μl of FeSO₄ solution and KCl-buffer were added to each resulting mixture. Each of the mixtures was incubated at 37°C for 1 hour in a shaking water bath. After the incubation, 1 mL trichloroacetic acid (TCA) reagent was added to each test tube containing each mixture, vortex mixed and centrifuged at 2000 rpm for 15 min. Thereafter, each supernant was decanted; 1mL of each supernatant was removed and added to new test tubes containing approximately 1 mL 0.67% TBA solution. The samples were further vortex mixed and heated for 20 min at 90°C in a water-bath. Absorbance of each sample and standard was measured at 532 nm. The percentage inhibition was measured according to Equation (I).

% inhibition = \frac{A_{standard} - A_{sample}}{A_{standard}} \times 100 \quad \text{(I)}

Oxygen radical absorbance capacity (ORAC) assay

The methods of [22,35] were adopted for the ORAC assay. The reaction was carried out in a 75 mM phosphate buffer (pH 7.4) medium that yielded a final reaction mixture of 200μL. The Plant extract (10μL, varying from 1 to 50μg/mL) and fluorescein (100μL; 1.6μg/mL) solutions were added to the reaction mixture and vortex. The resulting mixture was pre incubated for 5 min at 37°C. Fifty micro liters (50μL) of peroxide radical AAPH solution was added immediately and decaying of fluorescence was scanned for 35 min (at 1 min interval) at 37°C [35]. The decay in fluorescence was
measured with a fluorescence detector at 486 nm excitation and 538 nm emission wavelengths. The ORAC value was calculated by dividing the sample curve-area by the trolox curve area [22]. The ORAC value of the extract was expressed as mg/mL Trolox equivalent (TE) of dry extract.

**Enzymes Inhibition Activities**

**Tyrosinase Inhibition Assay**

The aerial part of the *S. nodiflora* was also assayed for its tyrosinase inhibition activity in a 96-well reader using a modified procedure of [36]. The anti tyrosinase activity of the plant was carried out at room temperature. A reaction mixture of 30 µL potassium phosphate buffer (50 Mm, pH 6.8) and 343 Units/mL mushroom tyrosinase and 110 µL L-tyrosine 2 mM was pre-incubated for 15 mins before adding L-tyrosine. The resulting mixture upon addition of L-tyrosine was further incubated for 15 mins before measuring the absorbance at 490 nm. A mixture without L-tyrosine was used as blank while the reaction mixture with the corresponding solvents (without plant material) was used as standard; kojic acid was served as the positive standard. The percentage inhibition of tyrosinase was calculated using Equation (II) while the IC50 value obtained for extract was from the dose-effect curves.

\[
\text{Tyrosinase inhibition (\%)} = \frac{(A-B)-(C-D)}{A-B} \times 100 \quad \text{... (II)}
\]

Where A = Absorbance of standard with enzyme; B=Absorbance of the standard without enzyme; C = Absorbance of sample with enzyme; D = Absorbance of sample without enzyme.

**Elastase Inhibition Assay**

The Elastase inhibition activity of the plant material was assayed by modifying the method of [37]. In this method, N-succ-(Ala)3-nitroanilide (SANA) was used as the substrate the release of p-nitronilide was monitored. The assay was performed in 0.2 M Tris-HCl buffer (pH 8.0). Porcine pancreatic elastase (PPE) was dissolved to make a 100 Units/mL stock solution in sterile water. The prepared substrate SANA was dissolved in the tris-HCl buffer without the plant extract. The plant extract (10µL) of varying concentrations (20-100µg/mL) was pre-incubated with 10µL enzyme (0.01 Units/mL) and 125µL of Tris-HCl for 10 min; thereafter, the substrate was added while the reaction proceeded further for 10 min at 25°C. The absorbance was measured at 410 nm wavelength immediately after incubation. The percentage inhibitions were calculated using Equation (III). IC50 values were obtained for from the dose-effect curves.

\[
\text{Elastase inhibition (\%)} = \frac{(A-B)}{A} \times 100 \quad \text{... (III)}
\]

Where A is the absorbance at 410 nm without plant extract and B is the absorbance at 410 nm with plant extract.

**Statistical Analysis**

Values shown in tables are the means and the standard deviations of three parallel measurements. The IC50 values of the enzymes (elastase and tyrosinase) inhibition activities of the plant’s extract were obtained from linear regression curve of enzymes inhibitory activities (%) versus extract’s concentrations (µg/mL) using Microsoft Excel (2010).

**Results and Discussion**

The phytochemical analyses of the *S. nodiflora* extract revealed the relative abundance (Figure 1) and concentration (Figure 2) of individual component present as saponin, alkaloids, polyphenols, proanthocyanidins, flavanol, tannin and flavonol (Figure 1). The phytochemical constituents of medicinal plants have been reported to be directly associated with their biological activities [10, 25, 38]. Saponins, alkaloids, glycoside, polyphenols, proanthocyanidins, flavanol, tannin and flavonol content are some of the phytochemicals known to be directly associated with antioxidant activities. These compounds are often referred to as powerful chain-breaking antioxidants [39-41]. The presence of these bioactive phytochemicals in the plant showed that the plant is a useful antioxidant material that may exert protective effects against inflammation, cancer diabetes and cardiovascular disease; they may also play a vital role in microbial inhibition.
Further, the results of the present study demonstrate that the plant exhibit variations in its phytochemical constituents (Table 1). Polyphenols (44.97 ± 1.33 mg of GAE/mL) was the most abundant (19.09 %) of the plant’s evaluated phytochemicals while tannin (22.00 ± 0.16 mg/g) was estimated to be the least abundant (9.36 %). The relative abundance (%) of the phytochemicals is in the order of polyphenols (19.09) > Saponin (18.61) > flavanol (18.00) > proanthocyanidins (14.52) > flavanol (10.35) > Alkaloid (10.07) > tannins (9.36) (Table 1). The mechanism of lipid peroxidation has been suggested to proceed via a free radical chain reaction Zhiyong and Yuanzong [42] which has been associated with cell damage in biomembranes [22,43]. The damage has been shown to initiate the development cancerous cells, cardiovascular diseases and diabetes according to [44,45]. The potency of the S. nodiflora extract to inhibit the process of lipid peroxidation was investigated in this study. The methanol extract of the plant was able inhibit lipid peroxidation by 55.60% well below that of the standard EGCG (90.95%). However, the value (Table 2) recorded for this plant showed its mild ability to inhibit lipid peroxidation processes (Table 2). The reducing power of a compound is related to its electron transfer ability and may therefore serve as a significant indicator of its potential antioxidant activity [7]. To estimate the reductive ability of the plant’s extract, the ferric ion reducing antioxidant potential of the extract was assayed. The result of this study showed Fe$^{3+}$ to Fe$^{2+}$ transformation potential of the plant’s extract with FRAP value of 681.10±0.13 mg/mL which is higher than that of the standard 3.20±0.01 mg/mL as shown in Table 2. The higher reducing ability of the plant’s extract implies a higher electron donating ability of the extract.

Table 1: Concentrations and the relative abundance of the evaluated phytochemicals in S. nodiflora.

| Phytochemicals | Concentrations ± SD | Relative abundance (%) |
|----------------|---------------------|------------------------|
| Flavonol       | 24.38 ± 0.03 mg CE/g| 10.35                  |
| Flavonol       | 42.41 ± 1.53 mg QE/mL| 18                     |
Table 2: Antioxidant activities of the S. nodiflora.

| Antioxidants Assay | FRAP (mg/mL) | LPO inhibition (%) | ORAC (μM TE/g × 10^3) |
|--------------------|--------------|--------------------|------------------------|
| Extract            | 6.81 ± 0.13  | 55.6               | 7.85                   |
| EGCG               | 3.20.50 ± 0.01 | 90.95             | 6.1                    |

Table 3: Enzyme inhibition activities of S. nodiflora.

| Enzymes inhibition Assay | Tyrosinase IC_{50} (μg/mL) | Elastase IC_{50} (μg/mL) |
|--------------------------|-----------------------------|--------------------------|
| Extract                  | 11.29                       | 71.20μg/mL               |
| Kojic acid               | 11.25                       | -                        |
| Xanthone                 | -                           | 20.2                     |

The Oxygen Radical Absorbance Capacity (ORAC)

The Oxygen Radical Absorbance Capacity of the plant’s extract is presented in Table 2. The ORAC values were determined in terms of Trolox equivalents (TE). The S. nodiflora extract exhibited a higher capacity to absorb oxygen radicals than the standard. The value ORAC value recorded for the plant’s extract is 7.85μM TE/g × 10^3 while that of EGCG is 6.10μM TE/g × 10^3. The higher oxygen radical absorbance capacity of the plant’s extract may imply that there are more electron withdrawing groups in the extract than in the standard. The antioxidant capacities and lipid peroxidation inhibition activity of the plant is mainly due to the presence and abundance of the bioactive compounds like Tannins, alkaloids, polyphenols and flavonol in the extract. The presence of phenolic compounds has been shown to correlate to the antioxidant activity of natural plant products [7,27,28] (Table 3).

The inhibitory effect of S. nodiflora on tyrosinase activity was investigated in-vitro and the result is presented in Table 3. The assay was carried out to show the possibility of using the plant to solve problems relating skin pigmentation. The potential of the plant’s extract and kojic acid (standard) to inhibit this enzyme increased from 20 to 100μg/mL with respect to the results obtained from this study. This shows that the inhibition activity of the materials increased with increasing concentration of the extract (Figure 3,4). The S. nodiflora extract and kojic acid showed optimum inhibition (39.75 and 95.9 % respectively) at the concentration of 100 μg/mL (Table 3). The IC50 value for the plant’s extract was 112.90μg/mL while that of the standard was 9.25 μg/mL. The enzyme tyrosinase was susceptible and mildly inhibited by the plant as a result of strong tyrosinase inhibitors like flavanol, flavonol and polyphenols present in the plant’s extract. South African Rooibos (Aspalathus linearis), Aspilia africana (Compositae) and Melanthera scandens are plants evaluated to have contained these phytochemicals and shown to have inhibited the enzyme tyrosinase [7,22].

The tyrosinase inhibition activity of the plant showed its value as a promising material for solving the problems of hyperpigmentation or may be used for depigmentation purpose. Elastase inhibition activity of the S. nodiflora was also investigated to evaluate its applicability in skin treatment and problems solving involving skin wrinkling and dry skin. The performance efficiency of the plant’s extract to inhibit Elastase at optimum concentration (100μg/mL) was 44.4%. The percentage by which both the extract and the standard (xanthone) inhibited the enzyme increase with their increasing concentrations respectively. It is possible that increasing the concentration of the extract beyond 100μg/mL could increase its inhibition activities. However, the plant material was able to demonstrate ability to maintain healthy skin and anti-wrinkle activities even at lower concentrations.

Conclusion

The presence of free radicals and reactive oxygen species is a major cause of skin aging and cellular oxidative stress related problems. Medicinal plants have found applications in solving deleterious effects of free radicals and reactive oxygen species in biological
system making them excellent natural alternatives to conventional synthetic chemicals. This study has shown the promising phytotherapeutic value of Synedrella nodiflora. The plant has demonstrated good free radical scavenging ability, reactive oxygen absorbing capacity and anti-lipid peroxidation ability and hence, a good antioxidant. In addition S. nodiflora possess the ability to inhibit the enzymes tyrosinase and elastase making it an economic important plant for skin treatment and healthy skin maintenance. To meet skin-whitening and anti-wrinkle’s properties demand for cosmetic preparation in the global market, S. nodiflora as well as some other plant’s extract capable of inhibiting tyrosinase and elastase activities may be good choice. The outcome of this study confirms the acclaimed ethnopharmacological applications of S. nodiflora and its biological roles in solving oxidative stress related diseases.

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