Secondary Metabolites from the Leaf of Millettia Aboensis Against Streptococcus Mutans Isolated from Carious Lesions


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ABSTRACT

Objectives: The use of traditional medicine as a source of relief from illness is a long-lasting practice. This is because of the possible adverse effects associated with synthetic drugs. The investigation of more plants with antimicrobial activities has been fuelled by the increase of multidrug resistivity of most clinical isolates. Although the leaf of Millettia aboensis has shown antimicrobial potentials against some clinical isolates, there is no report against Streptococcus mutans and detection of the phytoconstituents present.

Methods: Extraction was done according to the maceration process using methanol and further purification via liquid-liquid partitioning. The clinical samples used were cultured anaerobically and aerobically within 72 hr and 48 hr with the blood agar and brain heart infusion broth respectively. The isolates were identified using appropriate tests and methods. The fractions were further analysed for their Phyto-constituents with the help of High pressure liquid chromatography – diode array (HPLC-DAD).

Results: Methanol extraction using 50 g of plant material produced 12.72% wet weight, while the fractions using 90 g of methanol extract produced 30.38% for water, 20.57% for hexane, 6.65% for ethyl acetate, and 4.19% for butanol as percentage yield. The compounds detected from the fractions of the leaf extract of M. aboensis includes septicine - 1, cyclopenol - 2, psammaplin A - 3, 3-phenyl chromen-4-one - 4, hyperin - 5, rikuzenol - 6, hyperoside – 7 and N N-dimethyl tryptophan methyl ester - 8. There was zero antimicrobial activity recorded at 6.25, 12.5, 25 and 50 mg/ml against Strep. mutans.

Conclusion: The phytoconstituents present in the fractions are promising lead compounds for subsequent investigation in search of new bioactive agents.

Key Words: Millettia aboensis, Antimicrobial, Extract, fractions, Streptococcus mutans.

INTRODUCTION

Medicinal plants are plants with antimicrobial, antibacterial, anticancer, antitubercular, antimalaria activities. Traditional medicine is the main source of medical care for a great proportion of the population of the developing world. In Africa, indigenous plants play an important role in the treatment of a variety of diseases. As medicinal plants are a suitable alternative for synthetic and chemical drugs; they contain secondary metabolites which have medicinal and physiological potentials.

Millettia aboensis are small trees of 30-40 feet high and up to 2feet in girth, but usually 12feet high with reddish-brown pubescence on the petioles, branches and fruits. The flowers are purple in erect woody racemes, up to 18 inches long. Almost all the parts of M. aboensis have medicinal properties. The leaf is used by traditional herbalist for general healing, including ulcer and laxatives, while the root is used in treating gastrointestinal disturbances and liver diseases. Also, a decoction of the leaf, stem and root with other plant species is used to cure venereal diseases such as gonorrhoea, syphilis,etc. However, the plant parts of M. aboensis are known to be a rich source of steroids, phenolic acids, flavonoids, and alkaloids, hence with medicinal and physiological potentials. Although the leaf of M. aboensis has shown some antimicrobial potential against some clinical isolates,
no report against Streptococcus mutans and detection of the phytoconstituents present. The present study was therefore designed to provide information on the antimicrobial potentials of the extract/fractions and or the secondary metabolites from the leaf of M. Aboensis against S. mutans isolated from carious lesions and detect the phytoconstituents using High-pressure liquid chromatography – diode array (HPLC-DAD).

MATERIALS AND METHODS

Collection, identification and extraction of plant materials

Fresh leaves of M. aboensis were identified and collected from their natural habitat in Amudo-Awka, Anambra State of Nigeria, in January 2015. The plants were identified by Mr. Alfred Ozieko of Bioresources, Development, and Conservation Program (BDCP), Nsukka. Voucher specimens have been deposited in the herbarium section of the Department of Pharmacognosy and Traditional Medicine, Faculty of Pharmaceutical Sciences, Nnamdi Azikiwe University, Awka, with specimen number PCG/474/A/021. The leaves were thoroughly washed using running water and rinsed with distilled water and air-dried to a constant weight at room temperature of 25-27°C for 2 weeks, after which it was milled to a fine powder with the aid of a Marlex Exceller grinder.

Solvent extraction

The extraction process was carried out according to the methods of Ajaegbu et al. and Onoja et al. with little modification. Extraction was done according to the maceration process in methanol and further purification via liquid-liquid partitioning. The powdered plant materials of the leaf (750g) were extracted exhaustively for two days using the cold maceration method in methanol with intermittent shaking. The maceration process was then repeated for 24 hours in methanol for maximal extraction. The methanol extract was then collected and concentrated almost to dryness under vacuum at 40°C using a rotary evaporator. The concentrated methanol leaf extract (MLE) was reconstituted with 200 ml of methanol: water (2:8) and fractionated successively with hexane, ethyl acetate and butanol and water to give the hexane (MLHF), ethyl acetate (MLEF), butanol (MLBF) and aqueous (MLWF) fractions.

Preparation of test organism

Clinical isolates of S. mutans that were used for this study were taken from carious lesions of patients that attended the clinic at Federal College of Dental Technology and Therapy, Trans-Ekulu, Enugu. Firstly, all reagents that were to be used for the studies were prepared and stored in a refrigerator. Subsequently, brain heart infusion broth and blood agar were prepared and autoclaved at 121°C for 15 mins before antibacterial testing.

Subculturing

After 48hr of incubation for aerobic and 72 hr of incubation for anaerobic, the plates were brought out of the incubator. Colonies that developed were respectively subculturing into a freshly prepared blood agar using the streaking method for isolation of pure colonies. The plates were incubated aerobically and anaerobic culture condition. Pure colonies were identified by macroscopic examination of the colony growth on agar plates, morphological characteristics, motility test and other biochemical tests according to standard microbiological procedures. All the organisms that were isolated and identified were stabbed in nutrient agar slant, labelled appropriately and kept in the refrigerator.

Bacterial identification

Gram stain: Smears of the isolates were prepared and heat-fixed on clean grease-free slides. The smears were stained for 1 minute with crystal violet. This was washed out with gentle running tap water. The slides were flooded with dilute gram’s iodine solution for 1 minute. This was washed off with water and the smears were decolourized using 95% alcohol for 30 sec. The smears were then counterstained with Safranine solution for 30 sec. Finally, the slides were washed with running tap water, air-dried and observed under oil immersion objective lens.

Motility test: Motility test is usually used to differentiate motile organisms from non-motile ones. For this test, the hanging drop technique was employed and the technique was carried out as described by Cheesbrough and Adeoye et al. A little Vaseline jelly was rubbed around the cavity of hanging drop slip. A drop of peptone water containing the pure culture was then placed on a coverslip. The hanging drop slide was then placed over the drop of peptone water in such a way that the centre of the depression lies over the drop. The slide was quickly inverted and viewed under the microscope, using the 40X objective lens.

Catalase test: This test is usually used as an aid to differentiate other catalase-positive organisms from catalase negative. A loop full of the pure colony was transferred onto a plane clean glass test tube containing 2ml of 3%v/v hydrogen peroxide. The reaction was observed immediately for gas production indicated by the production of gas bubbles confirmed the presence of catalase.

Coagulase test: In slide test, a loop full of the isolates was mixed with human plasma at one end of the slide and a loop full of the isolate was mixed with water at the other end to serve as the negative control. The slide was rocked gently for 30 seconds and was allowed to stand. Particles indicating aggregation was used as an indication of coagulase reaction.

Sugar fermentation test: Each of the isolates were tested for its ability to ferment a given sugar with the product of acid and gas or acid only. Since most bacteria especially
gram-negative bacteria utilize different sugars as a source of carbon and energy with the production of either acid and gas or acid only. The test is used as an aid in their differentiation. The growth medium used was peptone water and the method used was that described by Kirk et al\(^1\). Peptone water was prepared in a conical flask and the indicator, bromoresol purple was added. The mixture was dispensed into test tubes containing Durham tubes. The test tubes with their content were sterilized by autoclaving at 121°C for 15min. 1% solution of the sugar was prepared and sterilized separately at 115°C for 10 min. This was then aseptically dispensed in 5 ml aliquots volume into the test tubes containing the peptone water and indicator. The tubes were inoculated with a young culture of the isolates and incubated at 37°C. Acid and gas production or acid only were observed after about 24 hr of incubation. The control tubes were not incubated\(^1\). Acid production was indicated by the change of the medium from light green to yellow color while gas production was indicated by the presence of gas in the Durham tubes\(^1\).

**Indole test:** The test is usually used as an aid in the differentiation of gram-negative, bacilli especially those of the Enterobacteriaceae. Tubes of peptone water were inoculated with a young culture of the isolates. The tubes were incubated at 37°C for 48hr. About 4 drops of Kovac reagents were added into 1ml of each of the culture tubes. The positive test was indicated by a red colour that occurs immediately at the upper part of the test tube\(^1\).

**Haemolytic test:** Nutrient agar was prepared and allowed to cool at 45°C and 5% sterile sheep blood was added and poured on sterile plates. The plates were then inoculated with the test organism and incubated at 37°C for 24 hr. After incubation, the plates were checked for zones of haemolysis\(^1\).

**Oxidase test:** This test was used to identify the organism that produces the enzyme cytochrome oxidase by oxidizing phenylenediamine. A piece of filter paper was soaked with few drops of oxidase reagent, using a glass rod, an inoculum was picked and was smeared on the filter paper. This was observed for blue-purple colour after 60 sec\(^1\).

**RESULTS**

**Extraction**

The yields of the extract and fractions of the extraction process of *M. aboensis* leaf are shown in Table 1, with MLWF having the highest yield (30.38%), followed by MLHF (20.57%) and MLEF (6.65%) while MLBF gave the lowest yield (4.19%).

**Bacterial Isolate**

From the 36 samples collected, *S. mutans* were present in 32 samples, 10 from male and 22 from females.

The morphological examination indicates a circular raised cream colony which test positive to gram stain, appears as short rods in shape and is immotile. The test isolate tested negative to catalase, oxidase and indole test. It is coagulase-positive and can ferment mannitol, glucose and lactose.

**Antimicrobial evaluation**

The methanol extract and the fractions of the leaf of *M. aboensis* showed no zone of inhibition against *S. mutans* with varying concentration of 6.25 to 50 µg/ml on the culture plates after 24 h, while the conventional antibiotic showed significant inhibition zone against *S. mutans* even at the lowest concentration of 10 µg/ml as shown in Table 2.

**HPLC-DAD analysis**

HPLC-DAD analysis was suggestive of eight phytoconstituents in the fractions of the methanol extract of the leaf of *M. aboensis*. The structures, Ultraviolet (Uv) spectra and chromatograph of the compounds are shown in Figure 1 – 4.

**DISCUSSION**

The advent of multi-drug resistance in human and animal pathogenic bacteria, as well as undesirable side effects of certain antibiotics, has triggered immense interest in the search of new antimicrobial drugs of plant origin\(^1\). Nature is a unique source of structures of high phytochemical diversity, many of them possessing interesting biological activities and medicinal properties. Previous studies show that the phytochemical component of *M. aboensis* is rich in flavonoids, tannins, saponins, and cardiac glycosides\(^1\). The use of plant extracts with known antimicrobial properties can be of great significance in treatments but several studies have also reported various types of contamination of herbal medicines which include microorganisms and toxins produced by microorganisms, pesticides and toxic heavy metals\(^1\). Oral diseases are the major problems in public health and one of the chronic diseases that affect humans drastically. The use of plant extract with known antimicrobial activities for the regulation of oral diseases and also an alternative to synthetic antimicrobials can be of great importance in dental caries treatment\(^1\). Previous preliminary phytochemical screening indicates that *M. aboensis* is a rich source of flavonoids, phenolic acids, tannins, saponins, alkaloids and steroids\(^1\). The methanol maceration process from 750 g of plant material of *M. aboensis* gave a yield of 12.72% w/w. From the results reported, MLWF had more yield than all other fractions, while MLBF gave the lowest yield. *S. mutans* are predominant in the patient under the age group of 18 – 25 years, and this may be as a result of frequent consumption of sugary beverages and food without restrictions\(^1\). The clinical samples indicate that the females
are more prone to *S. mutans* than male. This can be attributed to the difference in the eating habits of male and female. The consumption of candy cake chocolate, gum, soft drinks ice cream is higher in female than in male.22

The antimicrobial screening of the extract/fractions of the leaf of *M. aboensis* on the clinical isolates of *S. mutans* despite their varying concentrations at 6.25mg/ml-50mg/ml showed no inhibition. This can also be seen in the report given by Onyegeme-Okerenta and Okafor23, indicating no antimicrobial effect of *M. aboensis* extract against *E. coli*, but showed positive antimicrobial effects on *S. aureus, P. aeruginosa* and *K. pneumoniae* at 12.5 mg/ml – 400 mg/ml concentrations. The observed result with the methanol extract/fractions may be due to solubility of active compounds in methanol or inhibitors to the antimicrobial component.24

Okigbo et al and Okigbo & Igwe25-26 reported also inactivity of plant extracts which may be as a result of the age of the plant, type of extracting solvents, the period of harvesting of plant material and extraction method.

The HPLC screening suggested the presence of septicine - 1 in MLHF (Figure 1); cyclopenol – 2, psammaplin A – 3, 3,3-phenylchromen-4-one – 4 and hyperin – 5 in MLEF (Figure 2); rikuzenol – 6 and hyperoside – 7 in MLBF (Figure 3); and N,N-dimethyl tryptophan methyl ester in MLWF (Figure 4). Septicine is a toxic amine resulting from bacterial proteolysis which has potential as a cytotoxic (anticancer) agent.27 Cyclopenol is a naturally occurring 7-membered 2,5-dioxopiperazine alkaloid isolated from the extract of fungus penicillium cyclopium, penicillium sclerotiorum etc. Cyclopenol can be converted into the quinoline viridicatol, by the enzyme cyclopenase present in the conidia. Cyclopenol has phytotoxic and antimicrobial activities.28

Psammaplin is a marine natural product which inhibits aminopeptidase N and suppresses angiogenesis in vitro. It is a phenolic natural product isolated from a marine sponge, which showed potent cytotoxicity against several cancer cell lines.29 3-phenylchromen-4-one is an isoflavone. Isoflavones are reported to have a variety of bioprotective effects, including antioxidant, antimutagenic, anticarcinogenic and antiinflammatory activities. They protect the body from hormone-related cancers, like breast, endometrial (uterine) and prostatic.30,31,32,33,34 Hyper in belongs to the class of organic compounds known as flavonoid-3-o-glycosides. It has shown anti-inflammatory, anticarcinogenic and antiangiogenic properties. It helps in the reduction of depression, heart disease, fatigue, anxiety and cancer.35

Rikuzenol is a phenyl ether that was first isolated from cultured lichen mycobionts of *Graphis rikuzensis*.36 When screened for its cytotoxicity against murine lymphoma cell line (L5178Y) using the MTT method, it exhibited no activity.37 Hyperoside also is known as quercetin-3-O-D-galactoside is a flavonol glycoside present in a variety of vegetables and fruits.38,39 It has been isolated from various medicinal plants such as *Hypericum perforatum*40, *Ligularia fischeri*,41 *Crataegus davisi*,42 *Divaricate saposhnikovia*43, and *Hypericum mysorens*e44. Hyperoside is associated with several potent pharmacological activities which include anti-inflammatory, anti-thrombotic, anti-diabetic, anti-viral, anti-fungal, hepatoprotective, and antioxidant protective effects.45,46,47,48,49 N,N-dimethyl tryptophan methyl ester belong to the class of tryptophan esters. Tryptophan esters have shown to bypass the absorptive defect of Hartnup disease in young children50, and antitumor properties in various animal models.51

CONCLUSION

From this study, HPLC-DAD techniques revealed the presence of eight compounds and it can be concluded that the antimicrobial screening of the methanol extract/fractions of *M. aboensis* showed no activity against *S. mutans*, but with promising phytoconstituents present, there is need for purification and identification of these compounds present in the extract/fractions which could be exploited for pharmaceutical use.

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Conflict of interest

The authors declare no conflict of interest.

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