ABOUT AJPAC

The African Journal of Pure and Applied Chemistry (AJPAC) is an open access journal that publishes research analysis and inquiry into issues of importance to the science community. Articles in AJPAC examine emerging trends and concerns in the areas of theoretical chemistry (quantum chemistry), supramolecular and macromolecular chemistry, relationships between chemistry and environment, and chemicals and medicine, organometallic compounds and complexes, chemical synthesis and properties, chemicals and biological matters, polymer synthesis and properties, nanomaterials and nanosystems, electrochemistry and biosensors, chemistry and industry, chemistry and biomaterials, advances in chemical analysis, instrumentation, speciation, bioavailability. The goal of AJPAC is to broaden the knowledge of scientists and academicians by promoting free access and provide valuable insight to chemistry-related information, research and ideas. AJPAC is a bimonthly publication and all articles are peer-reviewed.

African Journal of Pure and Applied Chemistry (AJPAC) is published twice a month (one volume per year) by Academic Journals.

Contact Us

Editorial Office:  ajpac@academicjournals.org
Help Desk:  helpdesk@academicjournals.org
Website:  http://www.academicjournals.org/journal/AJPAC
Submit manuscript online  http://ms.academicjournals.me/.
Editors

Prof. Tebello Nyokong
Acting Editor
Chemistry Department
Rhodes University
Grahamstown 6140,
South Africa.

Prof. F. Tafesse
Associate Editor
Associate professor
Inorganic chemistry
University of South Africa
South Africa.
Editorial Board

Dr. Fatima Ahmed Al-Qadri
Asst. Professor
Chemistry Department
Sana'a University
Republic of Yemen.

Dr. Aida El-Azzouny
National Research Center
(NRC, Pharmaceutical and Drug Industries Research Division)
Dokki-Cairo, 12622-Egypt.

Dr. Santosh Bahadur Singh
Department of Chemistry
University of Allahabad
Allahabad, India.

Dr. Gökhân Gece
Department of Chemistry
Bursa Technical University
Bursa, Turkey.

Dr. Francisco Torrens
Institute for Molecular Science
University of Valencia
Paterna Building Institutes
P. O. Box 22085
E-46071 Valencia
Spain.

Dr. Erum Shoeb
Asst. Professor
Department of Genetics
University of Karachi
Karachi-75270
Pakistan.

Dr. Ishaat Mohammad Khan
Physical Research Laboratory
Department of Chemistry
Aligarh Muslim University
Aligarh 202002, India.

Prof. Jean-Claude Bunzli
Department of Chemistry
Swiss Federal Institute of Technology Lausanne (EPFL)
Institute of Chemical Sciences and Engineering
BCH 1402
CH-1015 Lausanne (Switzerland).

Mrinmoy Chakrabarti
Department of Chemistry,
Texas A&M University
415 Nagle Street, College Station, TX 77840
USA.

Dr. Geoffrey Akien
430 Eisenhower Drive, Apartment B-2,
Lawrence, Kansas 66049,
United States.

Prof. Anil Srivastava
Jubilant Chemsys Ltd.,
B-34, Sector-58,
Noida 201301 (UP),
India.
ARTICLES

Research Articles

Phytochemical screening of the leaf extracts of *Hyptis spicigera* plant 83

*Colour of starch-iodine complex as index of retrogradability of starch pastes* 93
Louis M. Nwokocha and Gabriel B. Ogunmola
Phytochemical screening of the leaf extracts of *Hyptis spicigera* plant


1National Research Institute for Chemical Technology, Private Mail Bag 1052, Zaria, Nigeria.
2Department of Chemistry, Ahmadu Bello University, Zaria, Nigeria.
3Division of Agricultural Colleges, Ahmadu Bello University, Zaria, Nigeria.
4National Agricultural Extension and Research Liaison Services, Ahmadu Bello University, Zaria, Nigeria.

Received 4 March, 2014; Accepted 7 May, 2014

The present study reports the screening of phytochemical constituents of the leaf extracts of *Hyptis spicigera* using hexane, ethylacetate and methanol and the leaf powder of the plant. Qualitative analysis of phytochemical constituents showed the presence of the following secondary metabolites vitannins, carbohydrates, saponins, flavonoids, steroids, alkaloids, quinones, coumarin, terpenoids, resins and cardiac glycosides. The quantitative analysis of total phenolics, alkaloids, saponins, terpenoids and flavonoids carried out using standard protocols revealed the presence of flavonoids (8.82%), saponins (6.23%), terpenoids (16.10%), alkaloids (7.55%) and phenolics (20.75%) respectively. Phenolics showed the highest content (20.75%) while saponins (6.23%) gave the least content. The high content of phenolics in the plant showed that *H. spicigera* plant may contain antioxidant properties and could be a good source of natural antioxidants. Also, the richness in flavonoids, saponins, alkaloids and terpenoids in this plant can be correlated with its medicinal properties used by traditional herbal healers in Northern Nigeria.

Key words: *Hyptis spicigera*, phytochemical screening, secondary metabolites.

INTRODUCTION

Plants have been the subject of human curiosity and use for thousands of years (Ram et al., 2004) and have played important roles in many countries of the world for centuries by providing food, shelter, clothing, agrochemicals, flavours and fragrances and more importantly, medicines (Gurib-Fakim, 2006). Traditional people have relied on medicinal plants to combat various ailments caused by microorganisms such as bacteria, fungi and viruses that infect the body system. Plants have indeed formed the basis of sophisticated traditional medicine systems which will continue to provide mankind with new remedies for all forms of ailments (Gurib-Fakim, 2006). Bioactive natural products have enormous economic

*Corresponding author. E-mail: zakariladan@gmail.com
Author(s) agree that this article remain permanently open access under the terms of the Creative Commons Attribution License 4.0 International License
importance as specialty chemicals as they can be used as drugs, lead compounds, biological or pharmaceutical tools, feed stock products, excipients and nutraceuticals (Pieters and Vlietinck, 2005). In recent times, focus on plant research has increased all over the world and a large body of evidence and knowledge has accumulated in the literature to show immense potential of medicinal plants used in various medical, pharmaceutical, cosmetic, agrochemical applications. An advantage of natural bioactive molecule is that they have a milder side effect on the body in comparison to chemically synthesized drugs (Badisa et al., 2003). With the increasing acceptance of herbal medicines as alternative form of health care delivery, the screening of medicinal plants for bioactive compounds is imperative (Masoko et al., 2005; Cowan, 1999).

*Hyptis spicigera* belongs to the family Lamiaceae and is commonly known as Black beniseed, or Black sesame. It is an erect aromatic herb, up to 1 m in height, with a terminal inflorescence in which the seeds are packed in quadruplets or more in the flowers. The plant is found around Senegal to western Cameroon, possibly native to Brazil, now widely naturalized in tropical Africa and Asia as well as Nigeria. It grows naturally and commonly as a weed. It prefers roadsides, waste places, cultivated places and often damp places (Burkill, 1995). Generally, the whole plant is used in traditional stores to protect cowpea against damage by *Callosobruchus* species (Lambert et al., 1985). The Bajju and Attyapp people of Kaduna state, northern Nigeria, make use of the inflorescence (where the seeds are packed) to cure headaches by sniffing it and also crushing the leaves and applying to the head to relieve head colds and headaches (Dalziel, 1937).

This paper reports the phyto constituents of the leaf extracts of *H. spicigera* and their potential medicinal applications.

**MATERIALS AND METHODS**

**Collection of the plant**

About 500 g of the leaf part of *H. spicigera* was collected in Basawa Village, Zaria, Kaduna state, Nigeria on the 26th November, 2013. It was taxonomically identified and authenticated by Mallam U. S. Gallah of the Herbarium Section, Department of Biological Science, Ahmadu Bello University Zaria, Nigeria, and a sample Voucher No.528 was deposited at the Herbarium section of the Department of Biological sciences.

**Extraction and isolation**

The plant was dried in the shade for 14 days and pulverized to powder using pestle and mortar in the laboratory. Approximately 450 g of the powdered plant material was macerated sequentially with hexane (1 L), ethyl acetate (1 L) and methanol (1 L) at room temperature (27°C) and concentrated *in vacuo* to afford the various crude extracts which were stored in the refrigerator (4°C) until needed for further analysis.

**Phytochemical screening**

Phytochemical analysis of the crude extracts was carried out according to standard methods (Harborne, 1998; Sofowora, 1993; Fansworth, 1996; Rangari, 2002).

**Salkowski reaction test for phytosterols**

To 0.5 ml each of the extracts in a test tube was added 1.0 ml of concentrated H$_2$SO$_4$ (conc.) from the sides of the test tube and then 1.0 ml chloroform. Appearance of reddish brown colour in chloroform layer indicates the presence of phytosterols.

**Liebermann-Burchard’s test for triterpenoids**

Extracts were treated with few drops of acetic anhydride, boil and cool. Conc. sulfuric acid was added from the sides of the test tubes which showed a brown ring at the junction of two layers, and formation of deep red color indicated the presence of triterpenoids.

**Foam test for saponins**

Small amount (0.1 g) of the extracts were taken in test tubes with little quantity (1.0 ml) of water and shaken vigorously. Appearance of foam persisting for 10 min indicated presence of saponins.

**Dragendorff’s test for alkaloids**

About 0.5 g each of hexane, ethyl acetate and methanol extracts were dissolved in 1.0 ml chloroform and evaporated. The residue was acidified by adding few drops of Dragendorff’s reagent (Potassium bismuth iodide). Appearance of orange red precipitate indicated presence of alkaloids.

**Molisch’s test for carbohydrates**

About 0.5 g each of the extracts was mixed with Molisch reagent, and then added H$_2$SO$_4$ conc. along the sides of the test tube to form layers. Appearance of reddish violet ring the interference indicated the presence of carbohydrates.

**Lead acetate test for flavonoids**

To 0.1 g each of the extracts were dissolved in ethanol and few drops of 10% lead acetate solution were added. Appearance of yellow precipitate indicated presence of flavonoids.

**Legal’s test for lactones**

To 0.1 g each of the extracts 1.0 ml sodium nitroprusside and 1.0 ml pyridine were added in test-tubes. The mixtures were treated with 0.01 moldm$^{-3}$ NaOH. Appearance of deep red colour indicated the presence of lactones.

**Ferric chloride test for phenolic compounds and tannins**

About 2.0 ml of each extract was measured in a test tube and 0.01 mol dm$^{-3}$ Ferric chloride solution was added drop by drop. Appearance of bluish black precipitate indicated presence of phenolic compounds and tannins.
Ninhydrin test for proteins

Few drops of ninhydrin were added to the extracts. Appearance of blue colour indicated presence of amino acid. Proteins may rarely give positive result with this test.

Keller-Killiani test for glycosides

About 1 ml of glacial acetic acid, few drops of 0.01 mol dm\(^{-3}\)Ferric chloride solution and H\(_2\)SO\(_4\) (Conc) slowly through the sides of the test tube were added to the extracts. Appearance of reddish brown ring at the junction of the liquids indicated the presence of deoxysugars.

Quantitative determination of phytochemical constituents

**Determination of total phenolic compound (TPC)**

Total phenolic content of the hexane, ethylacetate and methanolic extracts was determined by standard method (Makkar et al., 1993) with little modifications, using tannic acid as a standard phenolic compound. The extracts were diluted with distilled water to a known concentration in order to obtain the readings within the standard curve range of 0.0 to 600 μg of tannic acid/ml. 250 μl of diluted extract or tannic acid solution was mixed with 1 ml of distilled water in a test tube followed by the addition of 250 μl of Folin-Ciocalteu reagent. The samples were mixed well and then allowed for 5 min at room temperature in order to allow complete reaction with the Folin-Ciocalteu reagent. Thereafter, 2.5 ml of 7% sodium carbonate aqueous solution was added and the final volume was made up to 6.0 ml with distilled water. The absorbance of the resulting blue colour solution was measured at 760 nm using spectrophotometer after incubating the samples for 90 min. All the experiment was conducted in three replicates.

**Determination of alkaloids**

About 5.0 g of the dried powdered plant was weighed into a 250 ml beaker and 200 ml of 10% acetic acid in ethanol was added. The mixture was covered and allowed to stand for 4 h. This was filtered and the extract concentrated on a water bath to one-quarter of the original volume. Concentrated ammonium hydroxide solution was added dropwise to the extract until the precipitation was complete. The solution was allowed to settle and the precipitate was collected and washed with dilute ammonium hydroxide and finally filtered, dried, weighed and the percentage alkaloid was calculated (Harborne, 1998).

**Determination of terpenoids**

About 2 g of the plant leaf powder was weighed and soaked in 50 ml of 95% ethanol for 24 h. The extract was filtered and the filtrate extracted with petroleum ether (60 to 80°C) and concentrated to dryness. The dried ether extract was treated as total terpenoids (Ferguson, 1956).

**Determination of saponins**

About 15 g of each sample was placed into a conical flask and 100 ml of 20% aqueous ethanol was added. The samples were heated over a hot water bath for 4 h with continuous stirring at 55°C. The mixture was filtered and the residue re-extracted with another 200 ml and 20% ethanol. The combined extracts were reduced to 40 ml over water bath at 90°C. The concentrate was transferred into a 250 ml separating funnel and 20 ml of diethyl ether was added and shaken vigorously. The aqueous layer was recovered while the ether layer was discarded. This purification process was repeated and 60 ml of n-butanol was added. The combined n-butanol extracts were washed twice with 10 ml of 5% aqueous sodium chloride. The remaining solution was heated in a water bath. After evaporation the samples were dried in the oven to a constant weight and the percentage saponin was calculated (Obdoni and Ochuko, 2001).

**Determination of flavonoids**

About 5.0 g of the plant sample was weighed and extracted repeatedly with 100 ml of 80% aqueous methanol at room temperature. The whole solution was filtered through whatman filter paper No 41. The filtrate was evaporated into dryness over a water bath and weighed to a constant weight. The percentage flavonoids was then calculated (Soni and Sosa, 2013).

**RESULTS AND DISCUSSION**

The extracts of the leaves of *H. spicigera* were screened for the presence of the following secondary metabolites: alkaloids, glycosides, flavonoids, carbohydrates, tannins, steroids, terpenoids and resins, coumarins, saponins and quinines. The results of the phytochemical screening showed the presence of all the secondary metabolites analyzed in ethylacetate and methanol extracts while hexane extract showed only the presence of alkaloids, glycosides, carbohydrates and resins. Other secondary metabolites such as flavonoids, tannins, steroids, terpenoids, coumarins, saponins and quinones were absent in the hexane extract.

The phytochemical content was found to be similar to that obtained by other authors (Onayade et al., 1991) with different extracts revealing the different partitioning abilities of the different solvents used. The presence of these phytochemicals in all the extracts is quite instructive as this lends credence of the use of the plant for medicinal purposes. A lot of the plants contain non-toxic glycosides that can be hydrolyzed to give phenolic compounds that are toxic to microbial pathogens (Abaoba and Efuwape, 2001). The saponin content in the ethylacetate and methanol extracts were found to be present in these extracts, respectively. Saponins possess the property of precipitating and coagulating red blood cells (Sodipo et al., 1991). It also foamed in aqueous solution and has hemolytic effect and can also bind on cholesterol sites. These properties make saponins present in the plant to exhibit medicinal properties (Sodipo et al., 1991) and this therefore supports the findings in this present study that extracts of the plants may be useful in chemotherapy of mycotic infections which the antimicrobial studies revealed (Ladan et al., 2009). Alkaloids were found present in hexane, ethylacetate and methanol extracts and this can be corroborated with literature reports which indicate that naturally occurring alkaloids and their synthetic derivatives.
have analgesic, antispasmodic and bactericidal activities (Okwu and Okwu, 2004). They exhibit marked physiological activity when administered to animals. Classes of alkaloids are among the major powerful poisons known and despite being poisonous, some of the alkaloids are known to be useful in correcting renal disorders (Konkwar, 1979). The use of some plants for medicinal purpose, in the traditional treatment of diseases is due to the presence of flavonoids and saponins (Zwadyk, 1992; Othira et al., 2009), hence the use of H. spicigera for the treatment of diarrhea, dysentery, colds and several other diseases by local herbalists or traditional healers is not surprising. The presence of flavonoid was evident in methanol and ethylacetate extracts, flavonoid containing plants have been used as diuretic, laxative, emollient and poultice (Baba-Mousa et al., 1999) therefore; the use of H. spicigera rich in saponins and other Hyptis species in traditional medicine lent credence to the medicinal potentials of the plant. Tannins in some medicinal plants have been found to be responsible for the antiviral and antibacterial activities exhibited by such plants (De-Ruiz et al., 2001; Elegani et al., 2002). Therefore, H. spicigera with high tannin content in ethylacetate and methanol extracts could probably be a source of phytochemicals for the treatment of bacterial infections. Phenolic compounds like tannins present in plant cells are inhibitors of many enzymes (proteolytic and hydrolytic) used by plant pathogens. Other compounds such as saponins have antifungal properties (Abaoba and Efuwape, 2001; Mohanta et al., 2007). Therefore, these phytochemicals detected in this study may be responsible for the antimicrobial potency of the leaf extracts of Hyptis spicigera and also lend credence to the claims of traditional application of the plant as remedies for various ailments.

**Quantitative phytochemical analysis**

Results of the quantitative analysis data of the plant material revealed significant levels of phytochemical constituents present in the leaf as evident in the qualitative analysis data (Table 1). Phenolic content (20.75%) is the highest followed by terpenoids content (16.10%) while flavonoid (8.82%), saponins (6.23%) and alkaloids (7.55%) followed respectively. Subhashini et al. (2013) and Soni and Sosa (2013) have reported various phyto constituents in the leaves of Ecboiium viride (Forks) Merrill plant and the methanolic and ethyl acetate extracts of the leaves of Anogeissus leiocarpus and found the following values: terpenoids (0.3034 w/w), saponins (0.1100 w/w), alkaloids (0.1340 w/w), flavonoids (0.0884 w/w) and phenols (0.03045) for the E. viride (Forks) while alkaloids (152.0 ± 0.1 mg/g ), phenolics 1294.81± 3.0 mg/g), flavonoids (330.7 ± 3.0 mg/g) in the methanol extract and alkaloids (80.20 ± 0.0 mg/g), phenolics (616.5 ± 4.4 mg/g), flavonoids (202.5 ± 4.0 mg/g) in the ethyl acetate extract of the A. leiocarpus plant have been reported. The quantitative values of these metabolites reported in the leaf part of H. spicigerae are higher (Table 2) than those reported for E. viride (Forks) Merrill and A. leiocarpus plants. Phenolic compounds are one of the most important constituents of plant secondary metabolites with marked physiological properties.

The phyto constituents found in the plant may be responsible for its biological properties such as antioxidative, anti-inflammatory, anti-carcinogenic, anti-hypertensive, anti-diabetic, anti-cancer, cardiovascular protection and improvement of endothelial function (Han et al., 2007). Several studies have described the antioxidant properties of different parts of various medicinal plants which are rich in phenolic compounds (Brown and Evans, 1998; Krings and Berger, 2001; Malencic et al., 2007). Natural anti-oxidants mainly come from plants in the form of phenolic compounds, such as flavonoids, phenolic acids, tocopherolsetc (Ali et al., 2008) and used for the treatment of degenerative diseases. The antioxidative treatment of degenerative diseases.
different mechanisms, such as scavenging of free radicals, chelation of metal ions, such as iron and copper and inhibition of enzymes responsible for free radical generation (Akinmoladun et al., 2007; Benavente-Garcia et al., 1997). This plant (H. spicigera) will provide the natural anti-oxidant needed to enhance good living by scavenging free radicals that cause ill health in humans.

Conclusion

Phytochemical screening of the leaf part of H. spicigera revealed the presence of tannins, carbohydrates, saponins, flavonoids, steroids, alkaloids, quinones, coumarin, terpenoids, resins and cardiac glycosides which are important secondary metabolites. The richness of the plant in phenolic contents and other secondary metabolites affirmed its medicinal efficacy and potentials. The finding from this study therefore suggests that the leaf could be a potential source of natural anti-oxidant that could have great importance as therapeutic agents in preventing or slowing ageing associated with oxidative stress and related degenerative diseases. It is recommended that further investigation on the isolation and characterization of the bioactive constituents of the leaf leading to structural elucidation is necessary.

Conflict of Interests

The author(s) have not declared any conflict of interests.

ACKNOWLEDGEMENT

The authors are grateful to the Management of National Research Institute for Chemical Technology (NARICT), Zaria, Nigeria for providing necessary chemicals and equipment for the conduct of this research work.

REFERENCES


Full Length Research Paper

Colour of starch-iodine complex as index of retrogradability of starch pastes

Louis M. Nwokocha1* and Gabriel B. Ogunmola2

1Department of Chemistry, University of Ibadan, Ibadan, Nigeria. 
2Institute of Genetic Chemistry and Laboratory Medicine, Bodija, Ibadan, Nigeria.

Received 23 April, 2014; Accepted 12 May, 2014

Starch retrogradation is mainly due to the amylose fraction of starch. Amylose quantification is usually done by iodine staining with which it forms a blue colour complex while starch retrogradability can be monitored by freeze-thaw stability measurements. In this work, spectroscopic determination of the blue value and visual monitoring of starch–iodine colour complex were used to study starch retrogradation. The results obtained were compared with that from freeze-thaw stability measurements. Native cassava starch, its carboxymethylated and cyanoethylated derivatives of different degrees of substitution (D.S) were used in the study. From the results, increase in starch derivatization reduced amylose ability to bind iodine, decreased the blue value with resultant decay in the blue colour of the starch-iodine complex. The blue black colour of the starch-iodine complex was lost in carboxymethyl starch at D.S of 0.145 and cyanoethyl starch at D.S of 0.141; at these degrees of substitution, the helical structure of amylose was no longer maintained by the starch molecules and retrogradation eliminated. Freeze-thaw stability study showed clear pastes with no evidence of retrogradation over 10 freeze-thaw cycles for derivatized starches above these degrees of substitutions. This showed a good agreement with that obtained from the colour of the starch-iodine complex. Hence derivatized starch products, which showed absence of blue black colour of starch-iodine complex would be freeze-thaw stable over a long period of cold storage. The disappearance of the blue colour of starch-iodine complex thus becomes an index of retrogradability and freeze-thaw stability of starch pastes on cold storage.

Key words: Starch pastes, retrogradability, blue value, starch-iodine colour.

INTRODUCTION

When pasted starch products are subjected to cold storage the starch macromolecules in the starch dispersions tend to re-associate and exude water, a process referred to as syneresis (Thomas and Atwell, 1999; Balagopolan et al., 1998). This re-association and subsequent crystallization from the dispersions (retrogradation) is a major problem limiting starch utilization. Retrogradation problem can minimized by derivationization of the starch hydroxyls through introducing groups which retard the side-by-side association of starch molecules thereby stabilizing the aqueous dispersion of the paste (Thomas and Atwell, 1999; Balagopolan et al., 1998; Yeh and Yeh, 1993). When starch is derivatized, there is a change in the molecular organization within the granules and in starch physicochemical properties (Ogunmola and Nwokocha, 2002); and since starch

*Corresponding author. E-mail: lm.nwokocha@mail.ui.edu.ng 
Author(s) agree that this article remain permanently open access under the terms of the Creative Commons Attribution License 4.0 International License.
retrogradation is a phenomenon primarily associated with amylose molecules (Collison, 1968) (amylopectin retrogrades very slowly), the effective changes must have been those associated with amylose units. Starch reacts with iodine to form a blue colour (Thomas and Atwell, 1999). Studies on starch fractions have shown that the blue colour complex observed with iodine is due to the colour formation of amylose units (Radley, 1968). Amylopectin reacts with iodine to form a violet-red colour. The ability to form blue colour complex with iodine is gradually destroyed with increasing derivatization until the blue colour disappears (Radley, 1968). Similarly with increasing degree of derivatization, retrogradation is found to decrease until a certain degree of derivatization when retrogradation is completely removed. Since the blue colour of starch-iodine complex and retrogradation in starch pastes are primarily associated with amylose, it is possible to use the disappearance of the blue colour to monitor the retrogradability of aqueous starch pastes and thus provide an assay method of retrogradability in starch products evaluation.

MATERIALS AND METHODS

Preparation of cassava starch derivatives

Cassava starch used in this study was isolated in Chemistry Department, University of Ibadan, Nigeria. All reagents used including iodine crystals, potassium iodide, acetic acid, ethanol and sodium hydroxide are products of Sigma-Aldrich Chemical Co., St Louis, MO, USA) and of reagent grade. Carboxymethylation (Reaction 1) was achieved by treating aqueous starch dispersions with acrylonitrile using sodium hydroxide as the alkalizing agent. The detailed procedures are reported elsewhere (Ogunmola and Nwokocha, 2002; Nwokocha and Ogunmola, 2008). Derivatization can occur at any of the three hydroxyl groups of the glucopyranosyl unit.

Measurement of freeze-thaw stability of native, carboxymethyl and cyanoethyl starches

The freeze-thaw stability of the starch samples was determined on 5% starch pastes. 0.5 g dry starch samples were weighed into centrifuge tubes and 10 ml distilled water added and sealed. The samples were shaken to disperse the starch and then pasted in a Clifton water bath at 95°C for 30 min with occasional shaking throughout the heating regime. The samples were removed, cooled and subjected to alternate freezing (freezer temperature) and thawing (25°C) for 18 and 3 h, respectively and centrifuged at 5000 rpm for 15 min and the amount of water separated after each freeze-thaw cycle measured. The freeze-thaw stability was determined as percentage of water exuded per weight of paste.

Determination of the blue value of native, carboxymethyl and cyanoethyl starch-iodine complexes

0.1 g dry starch sample was weighed into a boiling tube, 1 ml ethanol (95%) was added followed by 9 ml of 1 M NaOH solution and heated in a boiling water bath for 10 min to solubilize the starch. The starch solution was cooled and quantitatively transferred into a 100 ml standard volumetric flask and the volume made up to 100 ml mark with distilled water. 2.5 ml of starch solution was taken into 50 ml standard flask; 0.5 ml of 1 M acetic acid was added followed by 1 ml of stock iodine (0.2 g I₂/2.0 g KI/100 ml) and the solution made up to the 50 ml mark with distilled water. The resulting colour was left for 20 min to fully develop before the absorbance reading was monitored at 620 nm with a Perkin-Elmer Lambda 3B double beam UV/ visible spectrophotometer. Iodine solution of same concentration as above but without starch sample was used in the reference cell. The colour of the starch-iodine complex was monitored visually. The blue value was calculated according to the method of Gilbert and Spragg (1964) using the formula:

\[
\text{Blue value} = \frac{\text{Absorbance at } 620\text{nm} \times 4}{\text{Concentration (mg/dl)}}
\]

RESULTS AND DISCUSSION

Freeze-thaw stability of carboxymethyl and cyanoethyl starches

Tables 1 and 2 show the freeze-thaw stability of carboxymethyl and cyanoethyl starches respectively. Both modified starches exhibited unusual stability even at low degrees of substitution in comparison with native starch. In carboxymethyl starch, stability to syneresis was observed at D.S of 0.064 and above. At lower levels of substitution (D.S< 0.064), syneresis was observed from the 5th freeze-thaw cycle in D.S of 0.028 and 0.061 with maximum exudates achieved decreasing with increasing D.S. In cyanoethyl starch, maximum exudates of 11.50% were obtained after 10 freeze-thaw cycles in D.S of 0.038. However, D.S of 0.100 and higher levels were stable to freeze-thaw cycles. Native cassava starch (D.S. 0.0) could not survive the first freeze-thaw cycle and attained maximum exudates of 31.30% after seven freeze-thaw cycles. Thus derivatization introduced branching groups onto the amylose chains which
hindered reassociation of amylose and reduced their ability to retrograde.

Relationship between colour of starch-iodine complex and degree of substitution

From Tables 3 and 4, a general decrease in blue value was observed with increase in D.S. This implied a gradual derivatization of the amylose as the D.S increased. Also the helical structure of amylose in solution responsible for complex formation with iodine was gradually destroyed with increase in D.S. It was observed that the decrease in blue value did not vary uniformly with increase in D.S. This implied both amylose and amyllopectin were randomly derivatized. The blue colour of the starch-iodine complex was gradually destroyed as D.S increased until a green colour was observed. In carboxymethyl starch, the transition to green colour occurred at a D.S of 0.145. Higher D.S gave various shades of green colour. At D.S of 0.145, the helical structure of amylose was destroyed to the extent that retrogradation was completely removed. Hence, D.S of 0.145 and above should produce pastes that are stable to cold storage. In cyanoethyl starch, the transition to green colour occurred at a D.S of 0.141. Higher D.S gave

Table 1. Freeze–thaw stability of native and carboxymethyl starches at 5% (w/v) concentration.

<table>
<thead>
<tr>
<th>No. freeze-thaw cycles (days)</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>D.S</td>
<td>0.0</td>
<td>0.028</td>
<td>0.061</td>
<td>0.064</td>
<td>0.076</td>
<td>0.111</td>
<td>0.131</td>
<td>0.134</td>
<td>0.187</td>
<td>0.194</td>
<td>0.197</td>
</tr>
<tr>
<td>Percentage water separated</td>
<td>0.7</td>
<td>22.10</td>
<td>5.60</td>
<td>18.60</td>
<td>26.70</td>
<td>30.80</td>
<td>31.30</td>
<td>31.30</td>
<td>31.30</td>
<td>31.30</td>
<td>31.30</td>
</tr>
</tbody>
</table>

- clear paste with no water separation, D.S of native starch = 0.0.

Table 2. Freeze–thaw stability of native and cyanoethyl starches at 5% (w/v) concentration.

<table>
<thead>
<tr>
<th>No. freeze-thaw cycles (days)</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>D.S</td>
<td>0.0</td>
<td>0.038</td>
<td>0.100</td>
<td>0.141</td>
<td>0.168</td>
<td>0.199</td>
<td>0.255</td>
<td>0.303</td>
<td>0.368</td>
<td>0.603</td>
<td>0.644</td>
</tr>
<tr>
<td>Percentage water separated</td>
<td>0.7</td>
<td>6.90</td>
<td>8.14</td>
<td>8.56</td>
<td>10.15</td>
<td>10.51</td>
<td>11.03</td>
<td>11.08</td>
<td>11.50</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- clear paste with no water separation, D.S of native starch = 0.0.
Table 3. Relationship between the starch-iodine colour and degree of carboxymethylation.

<table>
<thead>
<tr>
<th>Degree of carboxymethylation</th>
<th>Blue value</th>
<th>Starch-iodine colour</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0.375±0.0005</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.014</td>
<td>0.374±0.0007</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.039</td>
<td>0.370±0.0010</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.055</td>
<td>0.342±0.0015</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.116</td>
<td>0.290±0.0004</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.128</td>
<td>0.300±0.0011</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.145</td>
<td>0.251±0.0013</td>
<td>Deep green</td>
</tr>
<tr>
<td>0.185</td>
<td>0.170±0.0006</td>
<td>Green</td>
</tr>
<tr>
<td>0.205</td>
<td>0.208±0.0004</td>
<td>Green</td>
</tr>
<tr>
<td>0.212</td>
<td>0.185±0.0004</td>
<td>Green</td>
</tr>
<tr>
<td>0.269</td>
<td>0.186±0.0002</td>
<td>Green</td>
</tr>
<tr>
<td>0.296</td>
<td>0.159±0.0004</td>
<td>Green</td>
</tr>
</tbody>
</table>

Table 4. Relationship between the starch-iodine colour complex and degree of cyanoethylation.

<table>
<thead>
<tr>
<th>Degree of cyanoethylation</th>
<th>Blue value</th>
<th>Starch-iodine colour</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0</td>
<td>0.375±0.0005</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.038</td>
<td>0.266±0.0009</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.100</td>
<td>0.266±0.0010</td>
<td>Blue black</td>
</tr>
<tr>
<td>0.141</td>
<td>0.266±0.0006</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.169</td>
<td>0.246±0.0005</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.199</td>
<td>0.114±0.0015</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.255</td>
<td>0.032±0.0006</td>
<td>Light green</td>
</tr>
<tr>
<td>0.274</td>
<td>0.081±0.0013</td>
<td>Light green</td>
</tr>
<tr>
<td>0.303</td>
<td>0.079±0.0013</td>
<td>Light green</td>
</tr>
<tr>
<td>0.368</td>
<td>0.214±0.0010</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.484</td>
<td>0.171±0.0012</td>
<td>Green</td>
</tr>
<tr>
<td>0.591</td>
<td>0.194±0.0008</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.603</td>
<td>0.075±0.0006</td>
<td>Light green</td>
</tr>
<tr>
<td>0.639</td>
<td>0.181±0.0005</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.644</td>
<td>0.149±0.0005</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.699</td>
<td>0.164±0.0010</td>
<td>Green-blue</td>
</tr>
<tr>
<td>0.963</td>
<td>0.127±0.0015</td>
<td>Green</td>
</tr>
</tbody>
</table>

various shades of green colour which in some cases had bluish tint. This has been attributed to crosslinking and surface derivatization which tends to occur when starch is cyanoethylated under high acrylonitrile concentration (Hefrieter, 1986). Due to surface derivatization and crosslinking some long segments amylose inside the granules were not accessible to the reagents and hence remained long enough to form blue complex with iodine, this was responsible for the blue tint observed at high cyanoethylation degree.

Comparison of results from freeze-thaw stability and colour of starch-iodine complex showed that stability was reached at slightly different points. The difference arose from the fact that each method monitored retrogradation by different parameters. Retrogradation by freeze-thaw stability measures re-association between adjacent starch molecules while retrogradation by colour of starch-iodine complex measures the extent of complexation of the amylose with iodine. At D.S less than 0.064 in carboxymethyl and 0.100 in cyanoethyl starches, re-association was significant enough to result in the crystallization of the starch molecules. At these degrees of substitution the amylose chains were long enough to enter into helical structure with iodine to give the blue colour complex. At higher D.S the associative forces were very weak to result in starch crystallization and at
D.S of 0.145 in carboxymethyl and 0.141 in cyanoethyl starches, the helical structure was no longer maintained resulting in destruction of the blue colour. The method based on complex formation of starch with iodine has the advantage of rapidity because the result can be obtained in less than an hour unlike the freeze-thaw stability method that requires at least a day to get result.

**Conclusion**

The stabilization of starch paste by derivatization was related to the colour of starch-iodine complex. The blue colour of starch-iodine complex decayed with increase in degree of substitution (D.S) of the derivatized starch until it completely disappeared. This corresponded to when freeze-thaw stability was achieved without any noticeable sign of retrogradation. The disappearance of the blue colour of starch-iodine complex thus becomes an index of retrogradability and freeze-thaw stability of starch on cold storage.

**Conflict of Interests**

The author(s) have not declared any conflict of interests.

**ACKNOWLEDGEMENT**

This study was supported by University of Ibadan Research Grant 94/95.
African Journal of Pure and Applied Chemistry

Related Journals Published by Academic Journals

- African Journal of Mathematics and Computer Science Research
- International Journal of the Physical Sciences
- Journal of Geology and Mining Research Technology
- Journal of Environmental Chemistry and Ecotoxicology
- Journal of Internet and Information Systems
- Journal of Oceanography and Marine Science
- Journal of Petroleum Technology and Alternative Fuels