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Modification of African breadfruit (*Treculia africana*, Decne) kernel starch: Physicochemical, morphological, pasting, and thermal properties

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ABSTRACT

The need for starch to meet the ever-increasing industrial applications and its inherent problems in the native state have led to researches into unconventional starch sources and starch modifications, respectively. In this study, starch was isolated from African breadfruit kernels and modified to produce acetylated, AC, oxidized, OX, and acid-thinned, AT, starches. The effects of modifications on the native starch, NA, were examined through physicochemical, pasting, morphological and thermal properties. Modifications caused significant (p < .05) reductions in the NA starch moisture, ash, crude protein, crude fat, and crude fibre compositions. Swelling power had direct relationship with increase in temperature. Following modifications, OX starch showed the most significant increase in oil absorption capacity while NA had better water absorption capacity than the modified starches. Acid-thinning significantly improved the gelation and reduced the pasting properties of NA starch. Morphological study using scanning electron micrograph, revealed oblong and oval granules with rough surfaces for all the starches with no significant differences. Thermogravimetry of the starches showed single-step decomposition with increased thermal stability of African breadfruit kernel starch following modifications. DSC study showed that gelatinisation transition temperature of NA reduced following modification. The study showed

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1. Introduction

Starch, a naturally abundant polymer of plant origin, is an important raw material with versatile industrial applications. It shows different functionalities and it is the most regularly applied hydrocolloid. Its use in its native form however, has limitations [1] such as inability to dissolve in cold water, changing gelling power and viscosity after cooking, syneresis, a consequence of retrogradation [2], thermal decomposition and low shear resistance among others. To overcome these limitations, starch modifications are usually done to introduce desirable alterations in the starch structure with a resultant ability to predict and control starch behavior [3].

Starch modifications are achieved by physical, chemical and biotechnological means. Derivatives obtained have varied uses in pharmaceutical/biomedical, food and non-food industries as binders, disintegrants, lubricants [4], blood plasma expanders, cryoprotective agents for erythrocytes, fat replacers, flavor stabilisers, thickeners, gelling agents, salad dressing, water purification scavengers, flocculants, sizing agents and improvers of printing dyes in textile industry, sanitary product manufacture, binders in insulation of fibreglass, resins of adhesive in plywood, fluid loss control in subterranean drilling [5], floatation and sedimentation agents in ore mining [6], etc.

The diverse use of starch derivatives and the attendant overdependence on conventional sources such as corn, wheat, rice, sorghum, potato, tapioca, etc., which serve as major staple foods in developing countries, necessitated the search for alternative sources of starch to meet the ever-increasing demands of the insatiable starch industry. These resulted in researches into new sources of starch from unconventional/under-utilised sources such as mucuna beans [7], Cyperus sedge [8], cocoyam [9], bambara groundnut [10].

African Breadfruit is a tropical plant that is rich in carbohydrate (73%) [11]. It grows well in the Western and Eastern regions of Nigeria. It is a rain-forest plant, about 40 m high and with 30–50 fruits produced annually with weights in the range of 30 and 40 kg. The use of its kernel is limited to thickening of soup, making cookies, breadfruit cakes and various snacks [12,13]. However, the use of this kernel as

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source of starch is an unacknowledged potential. With the dearth of report on its use as starch source, it is reasonable to explore this virgin starch source with the view to maximizing its use as raw material particularly, as alternative source of starch for industrial applications. The optimum employment of African breadfruit starch is however, hinged on its modifications by various methods aforementioned, knowing full well that its potentials in the native states, like every other native starch, are limited. The objective of this work was to determine the effects of acetylation, oxidation and acid-thinning on the physicochemical, pasting, morphological and thermal properties of African breadfruit kernel starch with the view to improving its functional properties and maximizing its potential for industrial applications.

2. Materials and methods

2.1. Materials

Unprocessed African breadfruit was bought from Afigwe Main Market, Anambra State, Nigeria. The seeds were sun-dried for 4 days and the defective ones screened by hand-picking. The seeds were manually dehulled, dried, milled in a grinder (Marlex Excella, KIL, Daman, India) and kept in LDPE bag until required. Chemicals used were of analytical grade apart from sodium hypochlorite which was reagent grade.

2.2. Isolation and purification of starch

Isolation and purification of starch from African breadfruit kernel was done by the method of Lawal and Adebowale [14] with modifications. A 1000 g weight of the African breadfruit flour was suspended in 4000 mL of distilled water and the pH was adjusted to 8.0 using NaOH solution (0.2% w/v) at room temperature for 4 h with continuous stirring. The suspension obtained was centrifuged at 4500 rpm and the supernatant discarded. The pellet was suspended in 4 L distilled water, screened using muslin cloth and centrifuged for 30 min at 4500 rpm. (ROTANTA 460 R, Hettich GmbH & Co. KG, Tuttlingen, Germany). The starch obtained was washed twice before drying in the air for 48 h at 30 ± 2 °C. The native starch was labelled as NA.

2.3. Starch modification

2.3.1. Oxidation of starch

Oxidation of NA was done by the Forssel et al. method as reported by Lawal [9] with modifications. A 250 g weight of starch was suspended in 2500 mL of distilled water. The suspension pH was made 9.5 with 8% w/ v NaOH. NaOCl (10 g, 4% w/w active chlorine) was added to the suspension at the rate of 0.33 g/min at a pH range of 9–9.5 while stirring continuously at room temperature. Additional 10 min was allowed for the reaction after adding the NaOCl. The reaction medium pH was adjusted to 7 using 1.0 M HCl solution. The starch suspension was centrifuged, repeatedly washed with distilled water (four times) and centrifuged after each washing. The recovered oxidized starch was air-dried at room temperature for 2 days and labelled OX.

2.3.2. Acetylation of starch

Native starch was acetylated using the method of Sathe and Salunkhe described by Lawal [9]. A 20% w/v starch suspension in 0.5 L was made in distilled water. The pH of the suspension was adjusted to 8 using 4% w/v NaOH after magnetically stirring for 20 min. Acetic anhydride ($(CH_3CO)_2$, 10.2 g) was added at the rate of 0.17 g/min at a pH of 8–8.5. Additional 5 min was allowed for the reaction after adding the ($CH_3CO)_2$. The reaction medium pH was adjusted to 4.5 using 1.0 M HCl solution. The starch suspension was centrifuged, repeatedly washed with distilled water four times and centrifuged after each washing. The recovered acetylated starch was air-dried at room temperature for 2 days and labelled AC.

2.3.3. Acid-thinned starch

Starch thinning was done by Lawal method [9]. A 100 g weight of NA was suspended in 0.5 L of HCl (0.15 M). The starch suspension was magnetically stirred for 8 h at 50 °C. The modified starch was centrifuged at 4500 rpm (Rotanta 460 R, Hettich GmbH & Co. KG, Tuttlingen, Germany) for 10 min. The pellet recovered was washed repeatedly with distilled water four times and centrifuged after each washing. The recovered acid-thinned starch was air-dried at room temperature for 2 days and labelled AT.

2.4. Physicochemical, pasting, morphological and thermal properties

2.4.1. Physicochemical properties

Nwinuka et al. [15] method was used for determining the moisture, ash, crude fibre, crude protein and crude fat contents of the native and derivatised starches.

The carboxyl and carbonyl contents of OX were determined by the method of Lawal [9]. For the carboxyl content, about 8% w/v starch sample in 25 mL of 0.1 M HCl was prepared. The starch slurry was occasionally swirled for 30 min and then filtered using suction through a sintered glass funnel of medium porosity. It was washed with 0.4 L of distilled water and the starch was transferred into a 0.5 L beaker. The volume of the starch slurry was adjusted to 0.3 L with distilled water and it was heated with constant stirring over a period of 15 min for gelatinisation in a boiling water bath (Memmert W270, MEMMERT GmbH+Co.KG, Schwabach, Germany). The volume of the suspension of hot starch was then made up to 0.45 L using distilled water and titrated with 0.01 M NaOH to pH 8.3. NA was used for blank test. Eq. (1) was used to calculate the carboxyl group content.

Percent carboxyl

$$=\frac{(ST-BT)mL \times NaOH \ concentration \times 0.045 \times 100}{S \ (g)}$$
(1)

where ST = Titre value of oxidized sample; BT = Titre value of NA; S = sample mass.

To determine the carbonyl content of oxidized starch, 25% w/v hydroxylamine hydrochloride in 0.5 M NaOH was prepared. The solution was then diluted to 0.5 L with distilled water. 100 mL of 4% w/v starch sample was put in a 0.5 mL conical flask. The starch dispersion was heated for 20 min in a thermostated water bath (Memmert W270, MEMMERT GmbH+Co.KG, Schwabach, Germany) to attain gelatinisation. It was cooled to 40 °C, and the pH adjusted to 3.2 using 0.1 M HCl. This was followed by the addition of 15 mL of hydroxylamine reagent. A rubber bung was used to stopper the flask and heated at 40 °C for 4 h in a thermostated water bath stirring slowly. Excess hydroxylamine content of the suspension was determined by swiftly titrating the mixture with standard 0.1 M HCl to a pH value of 3.2. Hydroxylamine reagent only was used to calculate the carbonyl content of OX.

$$Carbonyl (C = 0) content (\%) = \frac{(BT - ST) mL \times HC1 concentration (M) \times 0.028 \times 100}{S (g)}$$
(2)

where BT = Titre value of NA, ST = Titre value of oxidized sample; S = sample mass.

The percentage acetyl group content of AC and the degree of substitution were determined using the method of Smith as described by Lawal [9]. A 5 g weight of AC was added to 50 mL distilled water in a 0.25 L conical flask and thoroughly mixed. The suspension was titrated with 0.1 M NaOH to obtain a permanent phenolphthalein pink endpoint. 0.45 M NaOH (25 mL) was added and the conical flask tightly stoppered with a rubber bung. It was vigorously shaken for a period of 30 min after which the bung was removed, rinsed into the flask together with the flask walls using distilled water. The mixture, with its excess NaOH, was titrated with 0.2 M HCl solution to a colourless phenolphthalein indicator colour. NA was subjected to the same treatment for a blank titre. Acetyl content (%) and the degree of acetyl substitution were calculated using Eqs. (3) and (4), respectively.

$$=\frac{(B1-S1)ML \times HC1 Concentration (M) \times 0.043 \times 100}{S(g)}$$
(3)

·· (10) 0.042 100

where BT = Titre value of NA, ST = Titre value of acetylaed sample; S = sample mass.

Degree of substitution (DS) =
$$\frac{162A}{4300-42A}$$
 (4)

where A = acetyl content (%).

The effect of temperature variation on swelling power and solubility was carried out, at 55–95 °C, as reported by Lawal [9] with modification. 1 g of starch sample was weighed and transferred quantitatively into a clean dry test tube and weighed. 10 mL of distilled water was added to the test tube and the starch suspension was thoroughly mixed for 30 s and heated at the chosen temperatures (55–95 °C) for 30 min on a thermostated water bath (Memmert W270, MEMMERT GmbH+Co. KG, Schwabach, Germany). The mixture was cooled to 27 ± 2 °C and centrifuged for 15 min at 4500 rpm (Centrifuge 80-3 Union Laboratories, England). The starch pellet after centrifugation and the test tube was weighed. Swelling power (g/100 g) at the chosen temperature was calculated using Eq. (5)

Swelling of Starch =
$$\frac{W_2 - W_1}{Weight of Starch}$$
 (5)

where $W_1 = mass$ of dry test tube + dry starch; $W_2 = mass$ of dry test tube + wet starch; S = mass of dry starch sample.

A 5 mL aliquot of the centrifugation supernatant was oven-dried (OV/125, Genlab Limited, Cheshire, England) at 110 °C to a constant weight. The residue obtained (%) was quantified and represents the extent of solubilisation of starch in water per 100 g of starch sample at the chosen temperature.

The method of Beuchat as reported by Lawal [16] was used for the determination of starch water and oil absorption capacities. Starch sample (1 g) in 10 mL water or oil (Power oil, Nigeria) was prepared and thoroughly mixed for 30 s. It was then allowed to stand for 30 min. The volume of the water or oil absorbed was then calculated as the difference between the initial volume and the final volume after allowing 30 min. The mass of oil or water absorbed was expressed as g/100 g starch on a dry weight basis.

Gelation studies were performed by the method of Lawal [16]. Starch samples (0.1, 0.2, 0.3, 0.4, 0.5, 0.6, 0.7, 0.8, and 0.9 g) were suspended in 5 mL of distilled water in test tubes. The starch suspensions were mixed thoroughly for 5 min and heated at 90 °C for 30 min in thermostated water bath. That was followed by cooling rapidly at the cold water tap and refrigeration for 2 h at 4 °C. The lowest concentration at which the sample in the inverted tube did not slip was taken as the least gelation concentration.

Fourier transform infrared (FTIR) spectroscopy was done using a Spectrum 100TM FTIR instrument (Perkin-Elmer, USA) equipped with an ATR accessory. All the spectra were the average of 16 scans from 4000 cm⁻¹ to 650 cm⁻¹ and were acquired at a resolution of 4 cm⁻¹. Tablets were prepared from a mixture of the sample with KBr at a ratio of 1:100 (starch:KBr)

2.4.2. Pasting properties

Pasting properties of starches were determined using Rapid Visco-Analyser (RVA-4 Newport Scientific, Australia). A 10% w/v starch suspension was prepared in distilled water. The slurry was subjected to measured heating-and-cooling cycle at constant shear where it was maintained for 1 min at 50 °C, heated from 50 to 95 °C with temperature variation of 12 °C/min and the sample was then held at 95 °C for 2.5 min, followed by 50 °C for 5 min. The final viscosity, maximum gelatinisation temperature, peak viscosity, breakdown, set back and holding strength were determined.

2.4.3. Morphological characteristic

The morphologies of the starch granules were examined using a scanning electron microscope (Phenom Pro Desktop SEM, Thermo Fisher Scientific). All of the samples were observed at an acceleration voltage of 25 kV.

2.4.4. Thermogravimetric properties

Starch sample thermal decomposition was determined using a Pyris-1 TGA apparatus (Perkin–Elmer, Shelton, USA). Samples were heated between 30 and 650 °C at a rate of 10 °C/min in an inert atmosphere of nitrogen flowing at 20 mL/min.

2.4.5. Differential scanning calorimetry

Gelatinisation properties of native and modified starches were determined by differential scanning calorimeter (DSC3, Mettler Toledo Co., Switzerland) using the method of Xu et al. [17] with modifications. Starch suspension was prepared by mixing starch and distilled water at a ratio of 1:3 (g/mL), which was sealed in an aluminium pan and placed at room temperature for 24 h to equilibrate. Samples were heated from 30 to 120 °C at a heating rate of 10 °C/min in a nitrogen flow rate of 50 mL/ min. An empty pan was used as reference. The onset temperature (To), peak temperature (Tp), conclusion temperature (Tc) and gelatinisation enthalpy (Δ H) were obtained by data analysis software STARE Software 16.10.

3. Results and discussion

3.1. Physicochemical properties

The results of chemical compositions of native and modified African breadfruit kernel starches are presented in Table 1. Significant (p < .05) reductions were observed in the percent moisture, ash, crude protein, crude fat and crude fibre following starch modifications. These reductions may be attributed to the degradative and eroding abilities of the oxidant and acid used for hydrolysis and loss of degraded molecules with wash water. Similar observations were made by other workers for hybrid maize [18], new cocoyam [9], jack bean [19] and white

Table 1

Native (NA), acetylated (AC), oxidized (OX) and acid-thinned (AT) starches of African breadfruit kernel chemical compositions*.

Sample	Yield (%)	Moisture (%)	Ash (%)	Crude protein (%)	Crude fat (%)	Crude fibre (%)
NA AC OX AT	$\begin{array}{r} 69.56 \pm 1.32 \\ 94.71 \pm 0.54 \\ 96.75 \pm 0.98 \\ 93.41 \pm 0.05 \end{array}$	$\begin{array}{l} 11.44 \pm 0.26^{a} \\ 9.52 \pm 0.48^{b} \\ 10.45 \pm 0.04^{c} \\ 9.05 \pm 0.11^{b} \end{array}$	$\begin{array}{l} 0.38 \pm 0.31^{a} \\ 0.24 \pm 0.07^{b} \\ 0.31 \pm 0.38^{c} \\ 0.29 \pm 0.29^{c} \end{array}$	$\begin{array}{c} 0.97 \pm 0.44^{ m a} \\ 0.55 \pm 0.38^{ m b} \\ 0.34 \pm 0.23^{ m c} \\ 0.52 \pm 0.26^{ m b} \end{array}$	$\begin{array}{l} 0.72 \pm 0.08^{a} \\ 0.58 \pm 0.57^{b} \\ 0.48 \pm 0.47^{c} \\ 0.42 \pm 0.20^{d} \end{array}$	$\begin{array}{c} 0.76 \pm 0.37^{a} \\ 0.46 \pm 0.41^{b} \\ 0.49 \pm 0.19^{b} \\ 0.55 \pm 0.28^{c} \end{array}$

* Means \pm SD of triplicate determinations. Yield of native starch, NA, was based on dry matter of African breadfruit kernel. Yields of derivatives are based on recovery after modification. Means within columns with different superscripts are significantly different at 95% confidence interval. OX: COOH = 0.35 per 100 anhydrous glucose unit; CHO = 0.19 per 100 anhydrous glucose unit; AC: percent acetyl substituent = 1.84%.



Fig. 1. FTIR spectra of native (NA), acetylated (AC), acid-thinned (AT), and oxidized (OX) African breadfruit kernel starches.

sorghum [20] starches. The range of values of moisture contents in native and modified starches are desirable for prolonged shelf life and are similar to those obtained by Olayinka et al. [20]. Low fat and protein contents establish the level of purity of the starch isolates. Similar values were obtained by [21] for breadfruit starch.

Fig. 1 presents the FTIR spectra of native, acetylated, oxidized and acid-thinned African breadfruit kernel starches. Native and modified starches showed strong absorption bands at various regions, depending on the starch type. Significant bands included those at 3277–3287 cm⁻¹, 2927–2932 cm⁻¹, 1733–1739 cm⁻¹, 1635–1642 cm⁻¹, 1150–1076 cm⁻¹, 1100–990 cm⁻¹, and 928–861 cm⁻¹ regions. The broad absorption bands in the 3277–3287 cm⁻¹ region are characteristic of hydrogen bonded —OH stretching vibration [22]. The intensities of absorption however decreased following modification, especially in AC and OX starches, as a

result of introduction of acetyl and carbonyl/carboxyl functional groups, respectively. This observation is similar to the findings of Rahim et al. [23] and Dao et al. [24]. Absorption peaks at 2927–2932 cm⁻¹ are attributable to $-CH_2$ stretching vibration while the presence of C=O symmetric perturbation are evident in OX and AC starches as shown by the sharp bands at 1733–1739 cm⁻¹ which are absent in the native starch [22–24]. 1635–1642 cm⁻¹ bands are characteristic of H₂O bending vibration as a result of water absorbed from bonding of hydrogen to the starch molecule -OH groups. The absorbance at 1150 and 1076 cm⁻¹ are the coupling of C–C, C–O and O–H bond bending, stretching, and asymmetric stretching of the C–O–C glycosidic linkage and absorbance at 928 and 861 cm⁻¹ are assigned both for C–H bending [23].

Figs. 2 and 3 show how temperature variation affects swelling power and solubility of native and derivatised African breadfruit kernel



Fig. 2. Mean swelling power of native (NA), acetylated (AC), acid-thinned (AT), and oxidized (OX) African breadfruit kernel starches with temperature variation from 55 to 95 °C. Standard deviations of triplicate determinations are used for error bars.



Fig. 3. Mean solubility of native (NA), acetylated (AC), acid-thinned (AT), and oxidized (OX) African breadfruit kernel starches with temperature variation from 55 to 95 °C. Standard deviations of triplicate determinations are used for error bars.

starches, respectively. Swelling power and solubility of all starches are directly related to temperature and increased with increase in temperature. Similar trends of gradual increase in swelling power and starches solubility with temperature were reported by other workers for Canavalia ensiformis [19,25], Kyllinga nemoralis [8], Araucaria brasiliensis [26] Artocarpus artilis [21], Artocarpus heterophyllus cultivars [27], cocoyam and cassava [28] and corn [29]. Increase in swelling power was gradual from 55 to 65 °C (1.93–4.18 g/g). However, a sharp rise in swelling power was observed between 65 and 75 °C for all starches except AT starch. Solubility increase with temperature, on the other hand, was gradual for all starches. Luo et al. [30] reported a similar jump in swelling power between 65 and 74 °C for crosslinked waxy potato starch while Adebowale et al. [10] reported noticeably high swelling power between 75 and 95 °C for unmodified bambara groundnut starch. NA starch had a significantly (p < .05) higher increase in swelling with temperature, reaching 11.60 \pm 0.13 g/g at 95 °C. That was followed by AC and OX starches with 9.98 \pm 0.27 and 9.15 \pm 1.79 g/g, respectively. The least swelling power was exhibited by AT starch. High swelling power with increase in temperature has been attributed to weakening of the bindings in intra-granular starches, which facilitated unrestricted swelling as temperature increased. Higher swelling power by native starches over modified starches were also reported by Remya et al. [31], Adebowale and Lawal [7] and Liu et al. [32] for modified lentil and banana, mucuna and common buckwheat starches. On the contrary, Singh and Adedeji [33] and Adebowale et al. [21] reported increase in swelling power of acid-modified and acetylated, oxidized and heat-moisture-treated breadfruit starches over their native counterparts, respectively. Increase in swelling power of AC starch could be attribued to introduction of bulky functional groups in the starch chain which facilitated entrance of water into the granules of starch. Oxidation of starch has been reported to reduce swelling power due to possible chain scission which disrupts structure of granules and hence ability to hold water. AT starch had lower swelling power similar to the reports of previous workers. Sandhu et al. [34] reported that acidthinning reduced swelling power of normal and waxy maize starches. Similar reports were given by Lawal [9] after oxidation and acidthinning of new cocoyam starch at varying temperatures and Xiao et al. [35] for oxidized rice starch. All the modifications, but acidthinning, reduced African breadfruit native starch solubility significantly (p < .05). Significant increase in solubility of starch following acid-thinning was observed ($10.33 \pm 0.58-56.67 \pm 1.16 \text{ g}/100 \text{ g}$). Dey and Sit [36], Sandhu et al. [34], and Lawal et al. [18] all reported increase in solubility of starches following acid-thinning. This increased solubility can be attributed to erosion of the amorphous amylose content of starch which is leached into the medium of suspension.

Table 2 shows the results of gelation properties of native and modified African breadfruit starches. A measure of gelation ability, the least gelation concentration. LGC, represents the minimum concentration of starch suspension in water that produced gel. From the results obtained. gelation tendency increased with increase in the concentration of starch suspension in water. LGC ranged from 6 to 10% w/v with acid-thinning reducing LGC (6% w/v) of native starch among the modifications while oxidation increased gelation concentration (10% w/v). Similar reduction of LGC by acid-thinning has been reported for corn, potato and rice and jack bean [25,37]. Reduced LGC can be attributed to molecular reordering of starch granules after imbibition of water and swelling due to bridging of intergranular binding forces among starch molecules [18]. Increase in LGC of OX may be attributed to fragmentation and molecular disintegration [38] and the introduction of carbonyl and carboxyl groups on the starch chain which reduced hydrogen bonding required for gelation [25]. NA and AC had 8% w/v LGC, a value similar to that reported for jack bean by Lawal and Adebowale [25].

Table 2

Gelation properties of native (NA), acetylated (AC), acid-thinned (AT) and oxidized (OX) starches of African breadfruit kernel.

Concentration (% w/v)	NA	AC	AT	OX
2	L	L	L	L
4	L	L	V	L
6	V	V	G	V
8	G	G	G	V
10	G	G	G	G
12	G	G	G	G
14	G	G	G	G
16	G	G	G	G
LGC	8	8	6	10

LGC = Least Gelation Concentration; L = liquid; V = viscous; G = gel.



Fig. 4. Oil and water absorption capacities of native (NA), acetylated (AC), acid-thinned (AT), and oxidized (OX) starches of African breadfruit kernel.

Results of oil and water absorption capacities of native and modified African breadfruit starches are presented in Fig. 4. Oil and water absorption capacities ranged from 39.20 \pm 0.43–117.60 \pm 0.09 g/100 g and 163.33 \pm 0.58–220.00 \pm 0.20 g/100 g, respectively. Modifications led to increase in oil absorption capacity (OAC) of native African breadfruit kernel starch with significant increase (p < .05) recorded by OX. Similar observation of increase in OAC following oxidation has been reported for bambara groundnut starch [10] and jack bean starch [25]. OAC of AT (68.6 \pm 0.32 g/100 g) was higher than that observed for NA $(39.20 \pm 0.43 \text{ g}/100 \text{ g})$. This was contrary to the observation of Alimi and Workneh [39] who reported decrease in OAC of citric acid modified acha and iburu starches after modification. Native starch however, had a better water absorption capacity (WAC) which was significantly greater than that of AT. Low WAC by AT is attributable to eroded amylose content following acid treatment. This gives credence to its low swelling power.

3.2. Pasting properties

Significant differences in pasting properties of native and modified African breadfruit starches were observed (Table 3). Pasting properties are affected by starch molecular structure and composition [40]. Peak viscosities of the modified starches, except for OX (2962.5 \pm 181.73 cP at 87.3 \pm 0.0 °C in 5.3 \pm 0.19 min), were lower than that obtained for NA (1677 \pm 85.56 cP at 85.7 \pm 1.2 °C in 4.9 \pm 0.09 min) with significant reduction observed in AT. High peak viscosity for oxidized new cocoyam starch has been reported by Lawal [9]. High peak viscosity reflects the extent to which starch is damaged [41]. Final viscosities of AC and OX were greater than the viscosity obtained for NA with OX having the highest (5092.5 \pm 197.28 cP) of the two. It is obvious that thinning greatly decreased the pasting properties of NA.

observations have been made by Zhang et al. [42] and Khan et al. [43]. The significantly reduced pasting ability of AT might be attributed to stiff granules not breaking up easily after erosion of the amorphous amylose layer making the movement of molecules of water into the starch granules slow, consequently preventing swelling of granules [44]. Also, acid hydrolysis caused an apparent breaking of the α -1,4-glycosidic linkages of the amylopectin chains, with consequential lowering of viscosity. Apart from AT, AC had the utmost thermal and mechanical shearing stability at 95 °C holding temperature as shown by reduced breakdown viscosity (234 ± 1.41 cP) because higher breakdown viscosity implies lower ability in withstanding stress from heating and shearing [41]. Similar reduction in breakdown viscosity following modification has been observed by Xiao et al. [45] for cross-linked oxidized potato starch. Breakdown reduction might be credited to the rearrangement in the altered starch granules [29]. The pasting temperature of OX was highest (87.3 \pm 00 °C). On the contrary, Xiao [45] reported reduction in pasting temperature following oxidation of rice, maize and potato starches. Reduced pasting temperature of AC is similar to that reviewed by Golachowski et al. [46]. Retrogradation tendencies of African breadfruit starch increased following oxidation and acetylation. OX has higher tendency to retrogradation with a setback viscosity of 2705.0 ± 24.04 cP. Acid-thinning significantly increased resistance to retrogradation as a result of shorter amylose chain following acidthinning and debranching of amylopectin chains [47]. Adebowale and Lawal [38] reported lowered setback viscosity following starch modifications.

3.3. Morphological characteristics

Fig. 5 presents the SEM of native and modified African breadfruit kernel starches. All the starches have diameters of about 10 µm similar

Table	3
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Pasting properties of native (NA), acetylated (AC), oxidized (OX) and acid-thinned (AT), starches of African breadfruit kernel.

Sample	PeV (cP)	TrV (cP)	BrV (cP)	FnV (cP)	SeV (cP)	PeT (min)	PaT (°C)
NA AC AT OX	$\begin{array}{r} 1677.5\pm85.56^a\\ 1490.0\pm29.70^a\\ -16.5\pm2.12^b\\ 2962.5\pm181.73^c\end{array}$	$\begin{array}{c} 1136.5 \pm 102.53^a \\ 1256 \pm 28.28^a \\ -24 \pm 1.41^b \\ 2387.5 \pm 173.24^c \end{array}$	$\begin{array}{c} 541 \pm 16.97^{a} \\ 234 \pm 1.41^{b} \\ 7.5 \pm 0.71^{c} \\ 575 \pm 8.49^{a} \end{array}$	$\begin{array}{c} 1872.5 \pm 101.12^a \\ 2626.5 \pm 12.02^b \\ -17.0 \pm 1.41^c \\ 5092.5 \pm 197.28^d \end{array}$	$\begin{array}{l} 736.0 \pm 1.41^a \\ 1370.5 \pm 40.31^b \\ 7.0 \pm 00^c \\ 2705.0 \pm 24.04^d \end{array}$	$\begin{array}{c} 4.9 \pm 0.09^{a} \\ 5.1 \pm 0.19^{a} \\ 5.6 \pm 0.47^{b} \\ 5.3 \pm 0.19^{a} \end{array}$	$\begin{array}{c} 85.7 \pm 1.20^{a} \\ 85.2 \pm 0.60^{a} \\ 0.0 \pm 00^{b} \\ 87.3 \pm 00^{a} \end{array}$

Means \pm SD of triplicate determinations. Columns with different alphabets are significantly different (p < .05); PeV = Peak Viscosity; TrV = Trough viscosity; BrV = Breakdown viscosity; FnV = Final viscosity; SeV = Setback viscosity; PeT = Peak Time; PaT = Pasting Temperature.



Fig. 5. Scanning Electron Micrograph of native (NA), acetylated (AC), oxidized (OX) and acid-thinned (AT) African breadfruit kernel starches.

to those reported by Zabot et al. [48] for annatto seeds. Oblong and oval particles with rough surfaces characterised all the starches. No significant morphological difference appeared on starch particle surfaces following oxidation, acetylation and acid-thinning. Unchanged surface morphology of OX is similar to the reports of Sangseethong et al. [49] for hypochlorite oxidized cassava starch, Kuakpetoon and Wang [50] for potato, corn and rice starches and Halal et al. [51] for barley starches but contrary to the report of Sukhija et al. [52] for elephant foot yam starch and Vanier et al. [53] for bean starch. Vanier et al. [53] also reported that acid hydrolysis at a mineral acid concentration of 3.16 M resulted in *exo*-erosion of starch granule surface.

3.4. Thermal properties

The thermograms of the TGA of native and modified African breadfruit kernel starches are shown in Fig. 6. All the starches showed single step decomposition with about 291.90 °C, 298.60 °C, 291.53 °C and 291.90 °C on-set decomposition for NA, AC, OX and AT, respectively. The initial loss in weight is attributable to desorption of bonded water molecules from starch granules. This occurred around 86–118 °C. Fifty percent weight loss for the starches occurred around 336.33 °C, 346.00 °C, 344.50 °C and 337.17 °C for NA, AC, OX and AT, respectively. Comparing the modified starches with the native, complete decomposition of AC, OX and AT occurred at 540.09 °C, 566.50 °C and 538.17 °C, respectively. These correspond to increase in thermal stability of African breadfruit kernel starch following modification. NA had significantly lower complete decomposition temperature of 369.50 °C. Aziz et al. [54] and Emeje et al. [55], reported increase in thermal stability of acetylated corn, pea and sweet potato, and acha starches, respectively. Reduction in thermal stability following oxidation was reported by Zhang et al. [22] and was attributed to starch polymer chain scission following oxidation while Tanetrungroj and Prachayawarakorn [56] reported lowered thermal stability for oxidized casava starch.

The results of the gelatinisation properties of the native and modified African breadfruit kernel starches are presented in Table 4. The gelatinisation onset temperature, peak temperature, conclusion temperature and enthalpy ranged from 73.7–79.3 °C, 80.2–84.7 °C, 83.8–89.4 °C, and 11.8–15.7 J/g, respectively. The gelatinisation transition temperatures of all the starch derivatives reduced than that of the native starch. These reductions can be attributed to structural weakening following starch modification as a result of acetyl and carbonyl/carboxyl groups introduced into the AC and OX, respectively, which enhanced water absorption, leaching of amylose, and hydration [49]. Other workers have reported similar decrease in gelatinisation temperatures following acetylation [38,57] and oxidation [49,57]. As a result of weakening caused by the introduced functional groups, early rupture of amylopectin double helices and consequent reductions in the gelatinisation enthalpies, Δ H, of acetylated and oxidized starches were



Fig. 6. TGA curves of Native (NA), acetylated (AC), oxidized (OX) and acid-thinned (AT) African breadfruit kernel starches.

observed [38] in a manner similar to those observed by Adebowale and Lawal [38] for mucuna bean, Lawal and Adebowale [25] for jack bean, and Nur and Purwiyatno [58] for corn starches. Reduced onset gelatinisation temperature in AT on the other hand can be adduced to reduced amylopectin chain length [59]. Increased enthalpy of gelatinisation was observed in AT due to the rigid crystalline amylopectin constituent left after erosion of the amorphous amylose layer by acid hydrolysis [60].

4. Conclusion

Acetylated, oxidized and acid-thinned starches were produced from African breadfruit kernel native starch. The physicochemical properties of native starch were significantly affected by modification. The FTIR spectroscopy revealed that hydroxyl contents of native starch reduced following acetylation and oxidation and new functional groups were introduced in the acetylated and oxidized starches. Improved solubility and gelation ability and reduced pasting properties were observed in acid-thinned starch. Starch morphology remained unchanged following modifications. Modification improved starch thermal stability. The native starch is a potentially good thickener for reducing water contents of preparations owing to its high, water absorption capacity while the oxidized African breadfruit kernel starch can function as flavor retention agent. The acid-thinned starch has the potential of usage in gelling, in quick-cook preparations with low viscosity, and frozen products with low tendency to syneresis.

Table 4

Gelatinisation properties of native (NA), acetylated (AC), oxidized (OX) and acid-thinned (AT), starches of African breadfruit kernel.

Sample	To (°C)	Tp (°C)	Tc (°C)	Tc – To (°C)	$\Delta H (J/g)$
NA	79.3	84.7	89.4	10.1	14.1
AC	76.5	81.9	85.6	9.1	11.8
AT	73.7	80.2	87.6	13.9	15.7
OX	74.3	82.4	83.8	9.5	12.9

To = onset temperature; Tp = peak temperature; Tc = conclusion temperature; Tc – To = gelatinisation temperature range; ΔH = enthalpy of gelatinisation.

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CRediT authorship contribution statement

Oderinde Abdulazeez Adewale:Conceptualization, Methodology, Validation, Formal analysis, Investigation, Resources, Writing - original draft, Writing - review & editing.**Ibikunle Ahmed Adeola:**Conceptualization, Methodology, Validation, Resources, Writing - original draft, Writing - review & editing.**Bakre Lateef Gbenga:**Conceptualization, Validation, Formal analysis, Writing - review & editing.**Babarinde Najeem Abiola Adesola:**Conceptualization, Supervision, Validation, Resources, Writing - original draft, Writing - review & editing.

Declaration of competing interest

The authors declare no conflict of interest.

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