PROXIMATE COMPOSITION AND FUNCTIONAL PROPERTIES OF SANDBOX SEEDS AS INFLUENCED BY PROCESSING METHODS

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ABSTRACT

Sandbox (Hura crepitans) seeds were processed as untreated (raw) sandbox flour (USF), cooked fermented flour (CFS), and soaked fermented flour (SFS). The three samples were defatted (with acetone) to obtain defatted untreated sandbox (DUS) flour, defatted cooked fermented (DCF) sandbox flour, and defatted soaked fermented (DSF) sandbox flour. The DSF was used to prepare sandbox protein concentration (SPC) and sandbox protein isolates (SPI). The physicochemical (pH and bulk density), functional properties (water absorption capacity, WAC), oil absorption capacity (OAC), least gelation concentration (LGFC), and in-vitro protein digestibility (IVPD) of defatted untreated and soaked fermented flours and protein concentrate and isolate were determined using standard procedures. The results showed that the moisture content varied between 5.13 – 8.23% and that processing treatments, such as fermentation and defatting significantly (at p < 0.05) increased protein, ash and carbohydrate contents of sandbox seed flour. Sandbox protein isolates exhibited highest protein content (87.49%), but the lowest crude fibre (0.02%), fat (1.31%) and ash (0.47%) contents. All the samples except SPC were acidic in aqueous solution and the bulk density values ranged between 0.45 g/mL – 0.67 g/mL. Defatted untreated sandbox (DUS) seed flour and sandbox protein isolate (SPI) had the lowest (1.06 mL/g) and highest (3.38 mL/g) significantly different (p < 0.05) WAC values, respectively, while DUS and SPI had the lowest (1.18 mL/g) and highest (2.30 mL/g) OAC, respectively. The foaming capacities for DUS, DSF, SPC and SPI were 42.00, 36.84, 20.00 and 12.00%, respectively, while emulsifying activity index ranged between 12.38 – 21.85m g⁻¹. All processing treatments were found to increase the in-vitro protein digestibility with values between 50.90 and 87.21%.

Key Words: Sandbox seeds, Processing techniques, Nutrient composition, Functional properties.

INTRODUCTION

Scarcity of protein-rich food is a major problem faced by many tropical developing countries, including Nigeria. The dependency of cereal-based diets by children and pregnant women for protein and energy deprives them of essential amino acids (EAAs). In recent years, plant proteins have been playing significant roles in the same developing countries where average protein intake is less than required (Khadi et al., 2003). Due to inadequate supplies of food proteins, there has been a constant search for unconventional legumes or oil seeds as new protein sources for use as both functional supplements (Onweluzo and Nwabugwu, 2009).

Modern research has thus focused more on oil seed crops as largely unexploited sources of food crops. Sandbox (Hura crepitans) seed falls into this group of underutilized species of plants. Sandbox as an underutilized plant in Nigeria is often grown as an ornamental plant in the tropics (Allen, 2000). It belongs to the “spurge” family (Euphorbiaceae) and is often planted in towns and villages as a cover tree. It has short densely crowned spines on the trunk and branches, the long-stalked leaves with prominent closely parallel pinnate nerves, the purple flower spikes and the large fluted flattened fruits are highly distinctive.

This tree flowers usually at the beginning of and at the end of raining season. One nut is a flattened and fluted disc with 5 – 20 lobes about 2.5 cm deep and 7.5 cm wide on a stout stalk. The capsule splits explosively releasing one flattened circular seed bout 18 mm across from each chamber (Fowomola and Akindahunsi, 2005). Seeds of sandbox are a potential source of dietary protein (37.62%) in West Africa and Nigeria (Abdulkadir et al., 2013).

In view of the high level of protein in sandbox seed, processing the whole seed to protein rich products such as defatted flour, protein concentrate and isolate could enhance its utilization as food ingredients. However, its
utilization as food ingredient could be limited due to the presence of anti-nutrients, such as alkaloids, oxalate, saponins, tannins, phytate and cyanide (Fowomola and Akindahunsi, 2005). Fasoyiro et al. (2006) revealed that soaking, cooking and fermentation are capable of reducing the anti-nutritional factors and organoleptic acceptability of the seeds (Christiana and Marcel, 2008; Omafurbe et al., 2004).

Seed proteins are required to possess the essential requisite functional properties for successful utilization in various food products or systems (Koladoye and Akanbi, 2015). These functional properties are intrinsic physicochemical characteristics which affect the behaviour of properties in food system during processing, manufacturing, storage and preparations. These properties include emulsion capacity and stability. It also includes foam capacity and stability. Other prominent properties are protein solubility, water and fat absorption capacity, bulk density (Aremu et al., 2007).

The purpose of the present investigation was to assess the effect of processing methods, such as soaking, cooking, and fermentation among others on the aforementioned properties of sandbox seeds.

MATERIALS AND METHODS
Dried seeds of sandbox (Hura crepitans) pods were collected at different locations in Obafemi Awolowo University, Ilé-Ifé, Osun State, Nigeria. The seeds were carefully removed from the pods and were washed under tap water to remove adhering dirt. The cleaned seeds were air-dried for about 2 hours and decorticated. The creamy white seeds of sandbox were then sun-dried for a week, packaged in polythene bags and stored in a freezer until use.

Preparation of whole and fermented sandbox flour samples
The sandbox seeds were divided into three portions and subjected to different processing treatments as shown in Figure 1. The first portion was oven-dried in a Gallenkamp oven (OVB 305, United Kingdom) at 50°C for 12 hours and the dried seeds were milled using hammer mill, sieved through a 300 µm sieve (Endecotts sieve, United Kingdom), then packaged using polythene bags. The sample obtained was referred to as untreated sandbox flour (USF). The second portion was prepared using a method described by Ayanwale and Kolo (2001). The seeds were cooked at atmospheric pressure for 2 hours, drained and fermented in a calabash lined with clean plantain leaves for 72 hours in the incubator (Memmert, IN30, Germany), and the fermentation was terminated by drying the cooked fermented seeds at 50°C (Gallenkamp oven, United Kingdom) for 12 hours. It was milled for 12 hours and then hammer milled, sieved through a 300 µm sieve (Endecotts sieve, United Kingdom) after which it was packaged in zip lock polythene bags. The sample obtained was referred to as cooked fermented sandbox flour (CFS).

The third portion was prepared employing same method as the second portion except that the seeds were soaked in lieu of cooking before fermenting. The seeds were soaked for 24 hours as described in the method of Nwosu (2010), drained and fermented in calabash lined with clean plantain leaves for 72 hours in incubator (Memmert, IN30, Germany), and the fermentation was terminated by drying the soaked fermented seeds at 50°C (Gallenkamp oven, United Kingdom) for 12 hours and then hammer milled, sieved through a 300 µm sieve (Endecotts sieve, United Kingdom). This was packaged using a zip lock polythene bags. The sample obtained was referred to as soaked fermented sandbox flour (SFS). All samples were stored in a freezer (at -20°C) pending further analysis.
Preparation of Defatted Sandbox Flour
Defatted sandbox seed flours were prepared from untreated and fermented flour sample using the method of Sathe (1994) as modified by Gbadamosi et al. (2011) as shown in Figure 2. The flour sample was defatted with cold (4°C) acetone using flour to solvent ratio of 1:5 w/v. The mixture was stirred over a magnetic stirrer for 4 h. The slurry was then filtered through a Whatman No. 1 filter paper. The residue was re-extracted twice in a similar fashion. The defatted flour was desolventized by drying in a fume hood at room temperature and the dried flour was finally ground in a blender (Binatone BLG-450, China) set at high speed to obtain homogeneous defatted flour. The defatted flour was stored in an air-tight plastic bottle and kept in a freezer until later used.

Preparation of Protein Concentrate
Sandbox protein concentrate was prepared from fermented defatted flour samples by a modification of the method described by Cheftel et al. (1985). A known weight (200 g) of the defatted flour was dispersed in 2 L of distilled water to give a final flour to water ratio of 1:10. The dispersion was gently stirred on a magnetic stirrer for 10 minutes. The pH of the resultant slurry was adjusted with 0.1 N HCl to pH 4 at which the protein was least soluble. This was determined from the solubility profile of the defatted flour during preliminary investigation.
The extraction was allowed to proceed with gentle stirring for 4 hours, keeping the pH constant. Soluble carbohydrates (oligosaccharides) and minerals were removed by centrifugation at 3,500 rpm for 30 minutes using MSE Harrier 15/80 Centrifuge (United Kingdom). The precipitate (concentrate) was collected and dried in an oven at 45°C for 8 hours. The flow chat of the process is shown in Figure 3.

Preparation of Protein Isolate

Sandbox protein isolate was prepared from fermented defatted flour by a method described by Chavan et al. (2001). A 200-g sample of the defatted flour was dispersed in 2 L of distilled water to give a final flour to liquid ratio of 1:10. The dispersion was gently stirred on a magnetic stirrer for 10 minutes. The pH of the resultant slurry was adjusted with 0.1 N NaOH to pH 10 at which the protein was most soluble. The extraction proceeded with gentle stirring for 4 hours keeping the pH constant. The non-solubilized materials were removed by centrifugation at 3,500 rpm for 10 minutes. The proteins in the extract were precipitated by drop wise addition of 0.1 N HCl with constant stirring until the pH was adjusted to pH 4.0. The mixture was centrifuged at 3,500 rpm for 10 minutes using a centrifuge (MSE Harrier 15/80, United Kingdom) in order to recover the protein. After separation of proteins by centrifugation, the precipitate was washed twice with distilled water. The precipitated protein was re-suspended in distilled water and the pH was adjusted to 7.0 with 0.1 M NOH, centrifuged and then freeze-dried. The freeze-dried protein was stored in air-tight glass containers at room temperature for further use. The flow chart used for the preparation of sandbox protein isolate is in Figure 4.
Figure 3: Scheme for the production of Sandbox protein concentrate
Adapted from Cheftel et al. (1985) as modified by Gbadamosi et al. (2011)

Figure 4: Scheme for the production of Sandbox protein isolate (Adapted from Chavan et al. (2001))
Proximate Composition of Whole, Defatted Sandbox Flour, Sandbox Protein Concentrate and Isolate

Moisture contents, total ash, crude fibre and crude fat, crude protein using Kjedahl apparatus were determined for all the samples according to AOAC (2000) methods.

Physicochemical and Functional Properties Determination

Bulk Density

Bulk density was determined by the method of Okezie and Bello (1988). A 10 mL graduated cylinder, previously tared, was gently filled with the samples (protein concentrates and isolates). The bottom of the cylinder was gently tapped on a laboratory bench several times until there was no further diminution of the sample level after filling to the 10 mL mark. Bulk density was calculated as weight of sample per unit volume of sample (g/mL).

\[ \text{Bulk density} = \frac{\text{weight of sample (g)}}{\text{volume of sample after tapping (mL)}} \] (1)

\[ \text{pH} \]

The pH was measured by making a 10% w/w suspension of the sample in distilled water. The suspension was mixed thoroughly and the pH (after standardizing the pH meter with buffer solutions of pH 7 and pH 4) was measured with a Hanna Checker pH meter (Model M11270).

Water Absorption Capacity (WAC)

The WAC was determined at room temperature and at temperatures ranging between 60 – 90°C using a combination of the AACC (1995) method and those of Soldolski (1962) and Rutkowski and Kozlowska (1981). A 2 g (W₁) sample (protein concentrate and isolate) was weighed separately into a known weight (W₂) centrifuge tube (2 cm in diameter) and 10 mL of pure Gino oil was dispersed into the sample. The content was mixed with 10 mL of pure Gino oil for 60 seconds and the mixture was allowed to stand for 10 minutes at room temperature, centrifuged at 1,788 rpm for 30 minutes using the centrifuge (0502-1 Hospibrand, USA) and the oil that separated was carefully decanted and the tubes were allowed to drain at a 45° angle for 10 minutes and then weighed (W₃). Oil absorption was expressed as percentage of the volume of oil absorbed by the weight of the sample as shown in equation (3) below:

\[ \text{Oil absorption capacity (\%)} = \frac{W₃ - W₂}{W₁} \times 100 \] (3)

where \( W₁ \) = weight of tube + sample before centrifuging and decanting

\( W₂ \) = weight of tube + sample after centrifuging and decanting

\( W₃ \) = weight of sample.

Least Gelation Concentration (LGC)

The method of Sathe and Salunkhe (1981) was employed for the determination of gelling concentration. Sample suspensions of 1, 3, 5, 7, 9, 11, 13, 15, 17 and 20% (w/v) were prepared in 5 mL distilled water separately into test tubes which were then heated in a boiling water bath for 1 hour followed by rapid cooling under running cold tap water. The test tubes were further cooled for 2 hours at 4°C. Least gelling concentration was determined as the concentration when the sample form the inverted test tube did not fall down or
Emulsifying Activity Index (EAI) and Emulsion Stability Index (ESI) of Samples

The EAI was determined by the method described earlier (Gbadamosi et al., 2011) with some modifications. About 500 mg of the samples was dispersed in 100 mL of distilled water NaCl solution. The protein solution was mixed with 50 mL of pure Gino oil and the mixture was homogenized using a blender (O’Qlink SN2200, China) set at high speed for 60 seconds. Fifty microlitres of the aliquot of the emulsion was transferred from the bottom of the blender after homogenization, and mixed with 5 mL of 0.1% sodium odecyl sulphate (SDS) solution. The absorbance of the diluted solution was then measured at 500 nm using spectrophotometer (722-2000 Spectronic, England). The absorbance obtained was used to calculate the EAI as shown in equation (4) below using the method suggested by Pearce and Kinsella (1978).

\[
\text{Emulsifying Activity Index (m}^2/\text{g)} = \frac{2 \times 2.303 \times A}{0.25 \times \text{protein weight (g)}} \tag{4}
\]

where \( A \) = absorbance at 0 minutes after homogenization.

To determine the ESI, the emulsions was allowed to stand for 10 minutes at room temperature and the ESI was determined as described below and it was expressed based on the absorbance at 0, 10 minutes and the time difference as shown in the formula:

\[
\text{Emulsion stability index} = \frac{AA \times \Delta t}{AA} \tag{5}
\]

where \( AA \) = absorption at 10 minutes
\( A \) = absorbance at 0 minute
\( \Delta t \) = change in time = 10 minutes.

Foaming capacity and stability of samples

Foam capacity and foam stability was determined by a modification of the method described by Chavan et al. (2001). Approximately 2 g of sample was dispersed in 100 mL of distilled water. The solution was then homogenised for 2 minutes using a blender (O’Qlink Blender, China) at high speed and then transferred into 250 mL measuring cylinder. The percentage ratio of the volume increase to that of the original volume of protein solution in the measuring cylinder was calculated and expressed as foam capacity or whippability (Ogunwolu et al., 2009). Foam stability was expressed as percentage of the volume of foam remaining in the measuring cylinder to that of the original volume after 30 minutes of quiescent period.

\[
\text{Foaming capacity (\%)} = \frac{v_2 - v_1}{v_1} \times 100 \tag{6}
\]

\[
\text{Foaming stability (\%)} = \frac{v_2 - v_3}{v_1} \times 100 \tag{7}
\]

where \( v_1 \) = volume before whipping (mL)

\( v_2 \) = volume after whipping (mL)

\( v_3 \) = volume after standing for 30 minutes (mL)

In vitro Protein Digestibility Determination

In vitro protein digestibility of samples was measured according to the combined methods of Saunders et al. (1973) and as modified by Chavan et al. (2001). Two hundred and fifty milligrams of the sample was suspended in 15 mL of 0.1 M HCl containing 1.5 mg pepsin, followed by gentle shaking for 1 hour at room temperature. The resultant suspension was neutralized with 0.5 M NaOH and treated with 4.0 mg pancreatin in 7.5 mL of phosphate buffer (0.2 M, pH 8.0). The mixture was shaker bath for 24 hours at room temperature. The mixture was then filtered using Whatman No. 1 filter paper and the residue washed with distilled water, air-dried and used for protein determination using Kjedhal procedure (AOAC, 2000) as described earlier. Protein digestibility was obtained by using the equation:

\[
\text{In vitro protein digestibility (\%)} = \frac{I}{F} \times 100 \tag{8}
\]

where \( I \) = protein content of sample before digestion

\( F \) = protein content of sample after digestion.

RESULTS AND DISCUSSION

Proximate Composition

The proximate composition of untreated sandbox flour (USF), defatted untreated sandbox flour
(DUS), cooked fermented sandbox flour (CFS), defatted cooked fermented sandbox flour (DCF), soaked fermented sandbox flour (SFS), defatted soaked fermented sandbox flour (DSF), sandbox protein concentrate (SPC) and isolate (SPI) is shown in Table 1.

The moisture content of samples varied from 5.123 – 8.23%. Defatted cooked fermented sandbox flour (DCF) exhibited the lowest (5.13%) and USF the highest (8.23%). The value of 8.23% obtained for USF was higher than the value reported by Abdukadir et al. (2013) for sandbox seed (3.10%). Sandbox protein isolate (SPI) moisture content (5.52%) was observed to be higher when compared to the moisture content of walnut protein isolates (4.50%) (Hua and Mao, 2012). Generally, foods high in moisture are susceptible to microbial attack. Therefore, the low moisture exhibited in sandbox flour samples may make it stable to microbial growth and thus conferred on the products extended shelf life.

Protein contents of the sandbox flour samples varied between 31.08 and 87.49%. Untreated sandbox flour (USF) exhibited the least protein content (31.08%) while SPI the highest (87.49%). It was revealed statistically that the values of CFS (43.78%) and SFS (41.59%) were not significant (p ≤ 0.05). The increase in protein value with fermentation time could be attributed to net synthesis of protein by fermenting organisms which might resulted in the production of some amino acids during protein synthesis (Uwagbute et al., 2000). The value of USF compare favourably with the report of Olatidoye et al. (2010) on crude protein of sandbox seed flour. The protein content of USF was higher than the values reported by Okolie et al. (2012) and Oyeleke et al. (2012) on sandbox seed flour protein (22.36% and 25.76% respectively). The protein content of SPI compared well with bambara protein isolates (85.97%) (Eltayeb et al., 2011), safflower protein isolates (90.1%) (Ulloa et al., 2011), but was higher than conophor nut protein isolate (80.00%) (Gbadamosi et al., 2011).

The crude fat content of the samples ranged from 1.31 – 48.89%. Sandbox protein isolate (SPI) had the lowest value (1.31%) while USF exhibited the highest value (48.89%). The fat content in USF compared well with the report of Olatidoye et al. (2010) on the crude fat of sandbox seed (43.52%), and was lower that the crude fat contents of 53.61% and 53.81% reported for sandbox seeds by Okolie et al. (2012) and Abdulkadir et al. (2013), respectively. The decrease in fat contents of fermented samples; CFS (43.41%) and SFS (43.67%) when compared with the whole untreated sample observed in this study may be attributed to the increased activities of the lipolytic enzymes during fermentation which hydrolysed fat components into fatty acid and glycerol (Chinma et al., 2009).

The crude fibre content of samples varied from 0.02 – 3.05%. SPI had the lowest while DUS exhibited the highest with significant difference (p ≤ 0.05). The observed decrease in crude fibre of fermented samples in this study could be as a result of degradation of the fibre by fermenting microbes (Babalola and Giwa, 2012). The crude fibre value of USF (1.25%) for this study compared well with the values for sandbox seed (1.45%) by Okolie et al. (2012) and 1.21% reported by Abdulkadir et al. (2013). Fibre in foods generally offers a variety of health benefits and it is essential in reducing the risk of chronic diseases (Food Science Avenue, 2008). The ash content varied significantly (p ≤ 0.05) between 0.42 – 3.72% with SPI exhibiting the lowest (0.42%) while DUS exhibited the highest value (3.72%). The value of USF (2.42%) agreed with the value reported by Olatidoye et al. (2010) for sandbox seed four (2.60%). The ash content of USF was lower compared to the sandbox seed ash content reported by Okolie et al. (2012), Oyeleke et al. (2012) and Abdulkadir et al., (2013) (6.42, 3.54 and 3.55% respectively). However, high ash content implies high mineral contents which helps retard the growth of certain micro organisms and some minerals are necessary in diets for health benefits (McClements, 2003).

The carbohydrate contents of the samples varied between 2.15 and 31.55% with significant difference (p ≤ 0.05). Fermentation significantly decreased total carbohydrate content of sandbox samples as observed in USF (8.13%), CFS (2.15%) and SFS (4.91%) due to enzymatic activities. The observed changes in carbohydrates with fermentation agreed with the report of
Achinewhu and Isichei (1990), Nnam (1995), and Onweluzo and Nwagbugwu (2009) on fermented fluted pumpkin seeds, fermented cowpea and fermented millet, respectively. The apparent decrease may be attributed to increased activity of amylolytic enzymes which hydrolyze starch and other complex carbohydrates to simpler sugars. The simpler sugars then probably provided energy for the fermenting micro organism as carbon source for possible synthesis of other compounds (Kazanas and Fields, 1981).

Functional Properties of Sandbox Flours and its Protein Concentrate and Isolate

Bulk density

The bulk densities of DUS, DSF, SPC and SPI are shown in Table 2. There was no significant difference in the BD of DUS (0.067 g/mL) and SPI (0.64 g/mL), but the values were significantly higher (p ≤ 0.05) than those of DSF (0.54 g/mL) compared favourably with the report of Appiah et al. (2011) on bulk density of fermented breadfruit pulp flour (0.57 g/mL). The reduction in bulk density as a result of fermentation was similar to the observation of Onimawo et al. (2003) and Elkhalalifa et al. (2005) on fermented pumpkin seed and sorghum flour respectively, where it was noticed that the fermented samples were less dense than the raw samples. Bulk density is a vital parameter that determines the suitability of flours for the ease of packaging and transportation of particulate foods as well as for infant formulations (Shittu et al., 2005).

Table 1: Proximate composition of sandbox seed whole, defatted and its protein concentrate and isolate

<table>
<thead>
<tr>
<th>Sample</th>
<th>USF</th>
<th>DUS</th>
<th>CFS</th>
<th>DCF</th>
<th>SFS</th>
<th>DSF</th>
<th>SPC</th>
<th>SPI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture (%)</td>
<td>8.23±0.04</td>
<td>6.83±0.11</td>
<td>7.62±0.02</td>
<td>5.13±0.04</td>
<td>7.74±0.05</td>
<td>5.83±0.03</td>
<td>8.03±0.04</td>
<td>5.52±0.15</td>
</tr>
<tr>
<td>Protein (%)</td>
<td>31.08±0.62</td>
<td>51.22±0.62</td>
<td>43.78±1.24</td>
<td>61.29±1.02</td>
<td>41.59±0.62</td>
<td>59.10±0.61</td>
<td>75.11±2.04</td>
<td>87.49±0.60</td>
</tr>
<tr>
<td>Fibre (%)</td>
<td>1.25±0.07</td>
<td>0.92±0.02</td>
<td>0.92±0.02</td>
<td>2.86±0.02</td>
<td>0.88±0.01</td>
<td>2.69±0.01</td>
<td>0.14±0.01</td>
<td>0.02±0.01</td>
</tr>
<tr>
<td>Fat (%)</td>
<td>48.89±0.01</td>
<td>3.63±0.01</td>
<td>43.41±0.01</td>
<td>3.11±0.01</td>
<td>43.67±0.01</td>
<td>3.34±0.01</td>
<td>3.94±0.06</td>
<td>1.31±0.01</td>
</tr>
<tr>
<td>Ash (%)</td>
<td>4.72±0.07</td>
<td>3.72±0.02</td>
<td>2.12±0.01</td>
<td>3.17±0.06</td>
<td>1.22±0.01</td>
<td>2.12±0.02</td>
<td>0.77±0.05</td>
<td>0.47±0.02</td>
</tr>
<tr>
<td>Carbohydrate (%)</td>
<td>8.13±0.78</td>
<td>31.55±0.81</td>
<td>2.15±1.26</td>
<td>24.45±1.19</td>
<td>4.91±0.57</td>
<td>26.93±0.63</td>
<td>12.03±2.11</td>
<td>5.25±0.50</td>
</tr>
</tbody>
</table>

* Values reported are means ± standard deviation of triplicate determinations. Means values with different superscript within the same row are significantly (P ≤ 0.05) different.

USF: Untreated sandbox flour; DUS: Defatted untreated sandbox flour; CFS: Cooked fermented sandbox flour; DCF: Defatted cooked fermented sandbox flour; SFS: Soaked fermented sandbox flour; DSF: Defatted soaked fermented sandbox flour; SPC: Sandbox protein concentrate; SPI: Protein sandbox isolate

The bulk density of the SPC therefore suggests usefulness in infant formulations, since low bulk density flours are desirable in infant food preparation (Nelson-Quartey et al., 2007) and nutritionally it promotes digestibility off food products, particularly among children with immature digestive system (Osundahunsi and Awoh, 2002).
The aqueous solutions of DUS, DSF, SPC and SPI had pH values of 5.80, 5.97, 7.20 and 6.67 respectively as shown in Table 2. The pH values obtained were acidic and slightly neutral, which indicated the presence of some organic acids. The increase in the pH of DSF over DUS agreed with the report of Omafuvbe et al. (2000) that the activities of microorganisms involved in the fermentation resulted in the release of ammonia by the organism, hence causing increase in the pH of the medium.

The pH of flour suspension is important since it affects functional properties, such as solubility, emulsifying activity and foaming properties (Chavan et al., 2001; Odoemelam, 2003; Khalid et al., 2003; Gbadamosi et al., 2011).

Water Absorption Capacity (WAC)

The values of water absorption capacities of the samples were significantly different (p ≤ 0.05) from one another and varied from 1.06 – 3.38 mL/g (Table 2). It was observed that SPI exhibited the highest water absorption capacity (3.38 mL/g) while DUS (1.06 mL/g) had the lowest. The water absorption capacities of SPC and SPI were higher than that of breadfruit flour (2.19 mL/g) (Nelson-Quartey et al., 2007). The WAC of SPI was higher than that of protein isolate from the seeds of fenugreek (1.68 mL/g) (El-El-Nasir and El-Tinay, 2007), while the WAC of DSF (1.34 mL/g) compared well with the 1.40 mL/g obtained for defatted white bean flour (Adebowale et al., 2005), and WAC value of DUS (1.06 mL/g) compared favourably with the 1.12 mL/g of soybean flour (Alfaro et al., 2004).

Adebowale et al. (2005) reported that defatted flours have higher water absorption capacities when compared to full fat samples. The absorption capacity of water by flour is a useful functionality for protein utility in aqueous food formulations, especially those involving high handling (Osungbaro et al., 2010). The matrix of protein with water is advantageous to properties, such as hydration, swelling power, solubility and gelation (Etudaiye et al., 2009).

Effect of Temperature on WAC

The effect of temperature on water absorption capacity is shown in Figure 5. The WAC values of samples with respect to temperature increase from ambient temperature to 90°C were DUS (1.06 – 1.45 mL/g), DSF (1.34 – 1.45 mL/g), SPC (3.13 – 3.39 mL/g) and SPI (3.38 – 3.82 mL/g). There were minimal increase in WAC of samples at lower temperatures (30 – 70°C), but as temperature increased above 70°C, water absorption capacity of the samples significantly increased (p ≤ 0.05). Increase in water absorption capacity of flour as a result of increase in temperature is in conformity with the increase in pH values observed in Table 2.
with earlier report for fermented maize flour (Fasasi et al., 2007). The presence in higher amounts of carbohydrate materials and higher protein contents in defatted flour samples and protein products respectively may be attributed to the high WAC exhibited with respect to change in temperatures. This high WAC demonstrated by SPC and SPI is an advantage as the samples could be used as ingredients in soups, gravy, baked products, and as thickeners in liquid and semi-liquid foods, since the samples have the ability to absorb water for improved consistency in food particularly at high temperatures.

Figure 5: Effects of temperature on water absorption capacity of defatted sandbox flours and proteins

Oil absorption capacity (OAC)
The oil absorption capacity of samples had significantly different (p ≤ 0.05) variation between 1.18 and 2.30 mL/g as shown in Table 2. The oil absorption capacity of SPI compared favourably with that of raw breadfruit flour (2.30 mL/g) (Odoemelam, 2005), and was within the range of OAC values reported by Adebowale et al. (2005) on mucuna bean flours (2.00 – 2.60 mL/g). Fat absorption is an important property in food formulations because fats improve the flavour retention and give soft texture to mouth-feel of foods (Odoemelam, 2003; Aremu et al., 2006; Ubbor and Akobundu, 2009). The value of DUS (1.18 mL/g) which was less to DSF (1.21 mL/g) as a result of fermentation treatment in this study, agreed with the report of Yadav et al. (2012) on the OAC of some selected cereal flour samples; sorghum (7.03 ml/g), pearl millet (6.7 mL/g) and maize (6.9 mL/g), that fermentation significantly (p ≤ 0.05) increased their oil holding capacity by 15.0, 22.0 and 23.0% respectively after 36 hours of fermentation, and 7% increase in oil holding capacity after 8 hours of fermentation for sorghum flour (Elkhalalifa et al., 2005). The oil absorption capacity observed in SPI suggests the presence of good lipophilic constituents and therefore may be suitable as recipes for sausage, soups, and cakes (Aremu et al., 2006).

Least gelation capacity (LGC)
The results of the least gelation capacity of DUS, DSF, SPC and SPI are presented in Table 2. DUS and DSF had the lowest LGC (3%) while SPC exhibited the highest LGC (9%). Lawal et al. (2007) reported on a study on pumpkin, that lower least gelation capacity implies better gelation capacity implying that DUS and DSF had better
gelling power. The ability of proteins to form gels and provide structural matrix for holding water, flavours, sugars, and food ingredients makes them useful application and future product developments (Aremu et al., 2006). The values of LGC obtained in this study for defatted flour samples and protein products were lower than the value (10%) reported for soybean flour by Alfaro et al. (2004) but were within the range of values reported for African bread fruit (6 – 12%) (Fasasi et al., 2007). SPC value compared favourably with LGC value pumpkin seed protein concentrate (8.00%) reported by Atuonnwu and Akobundu (2010).

In vitro protein digestibility (IVPD)
The result of in vitro digestibility of samples is presented in Table 2. The digestibility values of samples were significantly different ($p \leq 0.05$) and varied between 50.90 and 87.21%. The results of protein digestibility of samples compared favourably with and were within the range of values reported by Yadav et al. (2012) for the IVPD of fermented and unfermented sorghum (65.0 – 83.0%), pearl millet (68.0 – 84.0%) and maize (63.0 – 81.0%). The results obtained were also observed to be higher than the values obtained for uncooked soybean flour (30.5%) (Maha et al., 2009) and conophor protein concentrate (35.00%) (Gbadamosi et al., 2011). High digestibility in this study especially for the protein products may be attributed to decrease in the non-protein compounds especially polysaccharides as well as increase in the availability of the protein for enzymatic activities. Fermentation in this study also improved the IVPD of the defatted flour samples. This was attributed to the partial degradation of complex storage proteins into more simple and soluble products (Mohiedeen et al., 2010).

Emulsifying properties
The results of emulsifying activity index (EAI) and emulsion stability index (ESI) for sandbox flour samples are shown in Table 2. The values of EAI for the samples varied from 12.38 – 21.85 m$^2$/g and were significantly different ($p \leq 0.05$). DSF (21.85 m$^2$/g) and DUS (12.38 m$^2$/g) had the highest and lowest EAI respectively. The emulsion stability index (ESI) with significantly different ($p \leq 0.05$) values varied between 11.97 and 71.45%. The highest stability occurred in SPC (71.45%), while the least was observed in SPI (11.97%). This study revealed that both SPC and SPI had lower emulsion capacity compared to DSF. This result was in agreement with earlier observation by Atuonnwu and Akobundu (2010) who observed that the emulsion capacities of defatted pumpkin seed flour (22.0%) was higher compared to its protein concentrate ad isolate (5.0% and 1.0% respectively). According to Ogunwoli et al. (2009) and Hua and Mao (2012), emulsifying capacity of defatted cashew nut powder and defatted walnut flour (24.63% and 53.28% respectively) exhibited the highest values, while cashew nut protein concentrate and isolate (12.48% and 50.01% respectively) were lower as compared to the defatted flour. The preparation conditions of the samples can also affect their emulsifying properties as reported by Atuonnwu and Akobundu (2010), as difference in residual lipid content and composition can conditionally result from the defatting processes which are responsible for the modified protein functionality (Wu and Wang, 2003).

Foaming properties
The foam capacity of the samples varied between 12.00 – 42.00% with significant difference ($p < 0.05$) as shown in Table 2. It was observed that DUS (42.00%) had the highest FC while SPI (12.00%) had the lowest FC. The foam stability of the samples were between 4.00 – 35.00% with SPI having the lowest FS while DUS exhibited the highest value and were significantly different. Adebowale et al. (2005) explained that the low foam capacity may be attributed to high levels of globular proteins, which resist surface deaturation. Foaming result obtained for this study revealed that DUS and DSF had better capacities and stabilities than the protein products. This agrees with the observations of Atuonnwu and Akobundu (2010) on pumpkin seed protein concentrate and isolate which exhibited poor foamability, while pumpkin seed flour had a better foaming capacity and stability. This could be attributed to the protein solubility which has an important influence on the foaming behaviour of proteins (L'Hocine et al., 2006). Also, DUS (42.00%) and DSF (36.84%) had higher foaming capabilities than praseed flour (32.6%) reported by Choonhahirun (2010) and conophor defatted
flour (24.17%) as reported by Gbadamosi et al. (2011). The foam capacity of sandbox seed concentrate (20.00%) and isolate (12.00%) were lower than conophor protein concentrate (35.00%) and isolate 50.00%) reported by Gbadamosi et al. (2011), and walnut protein concentrate (38.78%) and isolate (46.34%) reported by Hua and Mao (2012). The results suggest that sandbox seed protein might not be suitable as whipping agents in food formulations, while the defatted flours could perform well as aerating agents in whipped toppings, frozen desserts and sponge cakes.

CONCLUSION
This study examined the effect of processing on nutrient composition, physicochemical and functional properties of sandbox seed flour as well as its in vitro protein digestibility (IVPD). The result of proximate composition revealed that fermenting and defatting significantly increased the protein contents of sandbox flour samples. Sandbox protein concentrates and isolates demonstrated high water and oil absorption capacities for possible use as flavour retainers and in bakery mixes, while defatted untreated and soaked fermented sandbox flours exhibited high gelling powers. Defatted soaked fermented sandbox flour showed good emulsifying and foaming capabilities for possible use as emulsifying and foaming agents especially in ice-cream and whipped toppings. The high in-vitro protein digestibility of the sandbox seed proteins means that a high percentage of the sandbox seed proteins can be digested by the proteolytic enzymes of the digestive system.

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