#### 1 Original Research Article

Evaluation of functional properties of spontaneous and starter culture fermentedsweet potato flour.

### 4 ABSTRACT

Sweet potato tubers obtained from a local market were sorted, washed processed 5 into fermented and unfermented sweet potato flour. The samples obtained were 6 analysed for their functional properties (swelling power, solubility index, water 7 absorption capacity, dispersibility and bulk density) using standard laboratory 8 procedures. A significant difference (p < 0.05) was observed in the loose and packed 9 bulk density with values ranging from 0.488 to 0.607 g/mL and 0.701 to 0.801 10 g/mL respectively but there was no significant difference (p>0.05) in the water 11 absorption capacity, oil absorption capacity and dispersibility. There was no 12 significant difference (p>0.05) in the swelling power but numerically the swelling 13 power increased with increase in temperature. A significant difference was 14 observed in the solubility index above 75°C and increase in solubility with increase 15 16 in temperature was observed. The result of this study showed that fermentation had no significant effect on the functional properties of the sweet potato flour except its 17 effect on the porosity of the granules as shown in the result of the bulk density. The 18 functional properties of these flours showed their uniqueness in each parameter 19 measured and on be useful for food application processes. 20

21 Keywords: Sweet potato flour, starter culture, fermentation, functional properties.

# 22 INTRODUCTION

Functional properties are properties that are used to predict the application of a food material as well as the end use for various food produce and the behavior of the functional properties depend on the source of raw material, presence of various ingredients, and processing conditions (Akinwale *et al.*, 2017). They interact with other food components directly or indirectly affecting processing applications,food quality and ultimate acceptance.

Sweet potato [Ipomoea batatas (L.) Lam.] is one of the major staple crops and 29 food the most important security promoting root crops in the world, 30 al.. especially in sub Saharan Africa (Low et 2009). Well adapted to the 31 tropical and subtropical regions, sweet potato has nutritional advantage for the 32 rural and urban dwellers (Ingabire and Hilda, 2011). Sweet potato (Ipomoea 33 batatas [L.] Lam.) is a dicotyledonous plant from the family Convolvulaceae 34 that grows in tropical and subtropical areas and even in some temperate 35 zones of the developing world (Ahn, 1993). In developing countries, sweet 36 potato ranks fifth economically after rice, wheat, maize, and cassava, sixth in 37 dry matter production, seventh in digestible energy production, and ninth in 38 production (Starters at al., 2005; Thottappilly and Loebenstein, protein 39 is about 131 million tonnes 1986).World production 40 per year, on approximately 9 million ha with mean estimated yields of 13.7 tonnes ha-1 41 (FAO, 2009). China is the world's leading producer of sweet potato, accounting 42 for about 80% of the total production worldwide. Nigeria is the most abundant 43 sweet potato producer in Africa and second to China in world production (FAO, 44 2014). 45

Sweet potato is an excellent source of energy (438 kJ/100 g edible portion) 46 and can produce more edible energy per hectare per day than cereals, such 47 as wheat and rice (Abu et al., 2000) and has other advantages, such as 48 versatility, high yield, hardiness, and wide ecological adaptability (Laurie et 49 al., 2012). Sweet potato roots are rich in starch, sugar, vitamin C,  $\beta$ -carotene, 50 iron, and several other minerals (Laurie et al., 2012; Oloo et al., 2014). 51 Despite its high carbohydrate content, sweet potato has a low glycemic index 52 due to low digestibility of the starch making it suitable for diabetic or 53

overweighed people (Ellong et al., 2014; Fetuga et al., 2014; ILSI, 2008; 54 Ooi and Loke, 2013). In addition, some varieties of sweet potatoes contain 55 colored pigments, such as  $\beta$ -carotene, anthocyanin, and phenolic compounds. 56 These pigments form the basis for classifying the foods as nutraceuticals (Oloo 57 et al., 2014). Nowadays, several research programmes are focusing on orange-58 fleshed or vitamin A sweet potato with great potential to prevent and 59 60 combat vitamin A deficiency for the crops value chain upgrading within the West African sub-region (Inaghe and Hilda, 2011). In addition, cassava is also 61 largely consumed after processing into garri, traditional flour, lafun, and 62 improved flour. However, the potential benefits of crop such as sweet potato are 63 marginalized and are underutilized despite their technological potential which 64 is well recognized and exploited elsewhere. 65

Fermentation is the conversion of carbohydrates to alcohol and carbon 66 dioxide or organic acids using yeasts, bacteria or a combination under 67 anaerobic conditions. The primary benefit of fermentation is the conversion 68 of sugars and other carbohydrates to usable end products. According to 69 Steinkraus (1995), the fermentation of foods improve flavour, aroma, and texture 70 in food substrates, preservation and shelf-life extension through lactic acid, 71 alcohol, acetic acid and alkaline fermentation, enhancement of food quality 72 with protein, essential amino acids, essential fatty acids and vitamins, 73 improving digestibility and nutrient availability, detoxification of anti-nutrient 74 through food fermentation processes. Starter cultures are living microorganisms 75 of defined combination used for fermentation purposes. They help to ellicit 76 specific changes in the chemical composition, nutritional value and sensorial 77 properties of the substrate (Opere et al., 2012) and they are generally 78 recognised as safe (Augirre and Collins, 1993). Moreover, their properties are 79

as follows: They are harmless, initiate and control the fermentation process,
typical for product, help in rapid acid formation, and help protect against
spoilage organisms.

Starter cultures are cheaply reproducible in large amount, they also help 83 provide desirable sensory properties and also assists in reducing fermentation 84 period.Lactic acid bacteria (LAB) are Gram positive acid tolerant, generally 85 non-sporulating, either rod or cocci shaped bacteria that produce lactic acid 86 as the major metabolic end product of carbohydrate fermentation. Lactic acid 87 bacteria have been reported to be predominant microorganisms in most of 88 89 the African indigenous fermented foods (Nout, 1991; Halm et al., 1993; Hounhouigan et al., 1993; Sanni, 1993; Steinkraus, 1996; Olasupo et al., 90 1997; Nago et al., 1998, Kunene et al., 2000; Duhan et al., 2013). 91 Yeasts are eukaryotic, single-celled microorganisms classified as members of the 92 fungus kingdom. The first yeast originated hundreds of millions of years ago, and 93 1,500 species are currently identified (Kurtzman and Fell, 2006; Hoffman et al., 94 2015). By fermentation, the yeast species Saccharomyces cerevisiae converts 95 carbohydrates to carbon dioxide and alcohols.For thousands of years the carbon 96 dioxide has been used in baking and the alcohol in alcoholic beverages (Legras et 97 al.,2007). The objectives of this study was to determine the application and use of 98 fermented sweet potato flour for various food products. 99

# 100 MATERIALS & METHODS

### 101 Sample Collection

102 Fresh raw sweet potato samples used for this work were purchased from Arena103 market, Bolade, Oshodi, Lagos state Nigeria. The samples were brought to

Biotechnology Department of Federal Institute of Industrial Research, Oshodi,(FIIRO) Lagos State.

#### **Preparation of Samples**

The sweet potato tubers were thoroughly sorted to remove bad ones from the lot. The sorted tubers were washed to remove adhering soil particles, weighed accordingly into four different portions. The tubers after weighing were thereafter peeled and sliced into small pieces, transferred into sterile fermentation bowls, appropriate volume of clean water was added to the sweet potato samples.

### 112 **Preparation of Inoculum**

Starter cultures (Lactobacillus brevis and Debaromyces polymorphous) used for 113 this study were isolated from fermenting sweet potato broth, after isolation the 114 organisms were subcultured by streaking on MRS agar (Oxoid) for Lactic acid 115 bacteria and incubated anaerobically at 37° C for 24 hours. Pure culture of yeast 116 isolates was cultivated by streaking on potato dextrose agar PDA (Oxoid) and 117 incubated at 25°C for 3 days. A colony was picked from each pure culture plates of 118 MRS and PDA plates and inoculated aseptically into MRS broth and YEPD ( 119 Yeast Extract Potato Dextrose) respectively then incubated. After incubation, the 120 organisms were harvested from the broth media by centrifuging at 5000 rpm for 15 121 minutes. The supernatants were decanted and the cell biomass disloged using 122 sterile distilled water 123

# 124 Preparation of starter culture fermented sweet potato flour

The sweet potatoes were washed to remove adhering soil particles and peeled. The peeled tubers were chipped into slices (4 to 5 mm) and soaked in potable water and inoculated with the starter cultures (Appropriate volume of sterile distilled water was used to wash the organisms into the various fermentation bowls containing the sweet potato samples). (Ajayi et al., 2016) for a period of 48 h and 72 h. After this period has elapsed, the fermented chips were drained and dried in a cabinet drier (Mitchel, Model SM220H) at 55°C for 9 h and milled into flour ( $\leq$ 250 µm) (Ajayi et al., 2016).

### 133 Preparation of spontaneous fermented sweet potato flour

The sweet potatoes were washed to remove adhering soil particle, the sweet potato was peeled and the peeled tubers were chipped into slices (4 to 5 mm) and soaked in potable water for a period of two days (48 h). After this period has elapsed, the fermented chips were drained and dried in a cabinet drier (Mitchel, Model SM220H) at 55°C for 9 h and milled into flour ( $\leq 250 \mu m$ ) (Oluwole et al., 2012).

# **139 Preparation of unfermented sweet potato flour**

The sweet potatoes were washed to remove adhering soil particles. Sweet potato was peeled the peeled tubers were chipped into slices (4 to 5 mm). After this period has elapsed, the fermented chips were drained and dried in a cabinet drier (Mitchel, Model SM220H) at 55°C for 9 h and milled into flour ( $\leq$ 250 µm) (Oluwole et al., 2012).

# 145 Functional properties determination

# 146 Bulk density determination

Bulk density was determined using standard methods (Ashraf *et al.*, 2012). Sample of 10g was measured into a 50 ml graduated measuring cylinder and gently tapped on the bench 10 times to attain a constant height. The volume of sample was recorded and expressed as grams per millilitre. Hausner ratio was determined as a ratio of the packed bulk density to the loose bulk density of the flour (Dossou *et al.*, 2014).

# 153 Water absorption capacity determination

The method described by Adebowale *et al.*, (2012) was used for determining the water absorption capacity (WAC). Sample of 1g was weighed into clean preweighed dried centrifuge tube and mixed with 10 ml distilled water with occasional stirring for 1 h. The dispersion was centrifuged at 3000 rpm for 15 min. After centrifuging, the supernatant was decanted and the tube with the sediment was weighed after removal of the adhering drops of water. The weight of water (g) retained in the sample was reported as WAC.

### 161 **Dispersibility determination**

Standard method was used for determining dispersibility (Kulkarni *et al.*, 1991). Sample of 10g was dispersed in distilled water in a 100 ml measuring cylinder and distilled water was added up to 50 ml mark. The mixture was stirred vigorously and allowed to settle for 3 h. The volume of settled particles was noted and percentage dispersibility was calculated as follows;

167 Dispersibility (%) =  $(50 - volume of settled particle) \times 100$ 

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# 169 Oil absorption capacity (OAC)

One gram of sample was mixed with vegetable oil (1:10). The mixture was stirred for 30 min at room temperature. After samples were centrifuged (2500g, 30 min), the supernatant was transferred to a graduated cylinder of 10 mL, where the volume was measured. The OAC was expressed as milliliters of vegetable oil held per gram of sample (Chau and Huang, 2003).

### 175 Swelling power and solubility index determination

The method described by Hirsch (2002) was used for swelling power and solubilityindex determination. One gram of sample was poured into pre-weighed graduated

centrifuge tube appropriately labeled. Then, 10 ml of distilled water was added to 178 the weighed sample in the centrifuge tube and the solution was stirred and placed 179 in a water bath heated at different temperature range (55, 65, 75, 85, 95 °C) for 1 h 180 while shaking the sample gently to ensure that the starch granules remained in 181 suspension until gelatinization occurred. The samples were cooled to room 182 temperature under running water and centrifuged for 15 min at 3000 rpm. After 183 184 centrifuging, the supernatant was decanted from the sediment into a pre-weighed petri-dish; the supernatant in the petri-dish was weighed and dried at 105 °C for 1 185 h. The sediment in the tube was weighed and the reading recorded. The starch 186 swelling power and solubility was determined according to the equations below; 187

188 Swelling power = weight of swollen sediment  $\times 100$ 

- 189 Weight of dry starch
- 190 Solubility = weight of dry supernatant  $\times 100$
- 191 weight of starch sample

#### 192 Statistical Analysis

All experimental data obtained were subjected to analysis of variance (ANOVA)

194 procedure of SPSS version 15.0 (SPSS Inc., 2006) at 5% significant level.

# 195 **RESULTS**

196 Table 1: Functional properties of various sweet potato flour.

Sample	LBD (g/ml)	PBD (g/ml)	Hausner ratio	WAC (g/g)	(OAC) (g/ml)	Dispersibility (%)
A	0.500 <sup>b</sup>	0.660 <sup>b</sup>	1.32	1.515 <sup>a</sup>	9.100 <sup> a</sup>	0.355 <sup>a</sup>
B	0.607 <sup>a</sup>	0.801 <sup>a</sup>	1.32	1.480 <sup>a</sup>	9.050 <sup>a</sup>	0.385 <sup>a</sup>

С	0.488 <sup>b</sup>	0.702 <sup>b</sup>	1.44	1.540 <sup>a</sup>	9.100 <sup>a</sup>	0.410 <sup>a</sup>
D	0.519 <sup>b</sup>	0.701 <sup>b</sup>	1.35	1.610 <sup>a</sup>	8.550 <sup>a</sup>	0.355 <sup>a</sup>

197 Values are mean. Mean values (n = 2) having different superscript alphabets in the same column are significantly

different (p < 0.05) A: Spontaneously fermented sweet potato flour; B: Unfermented sweet potato flour; C: 48 hours

- starter culture fermented sweet potato flour; D: 72 hours starter culture fermented sweet potato flour; LBD: Loose
  bulk density; PBD: Packed bulk density; WAC: water absorption capacity; OAC: Oil absorption capacity
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Figure 1:Swelling power of different sweet potato flour samples at different temperatures.Sample A: Spontaneously
 fermented sweet potato flour, Sample B: Unfermented sweet potato flour; Sample C: 48 hours starter culture
 fermented sweet potato flour; Sample D: 72 hours starter culture fermented sweet potato flour.

Table 2 : Solubility index of various sweet potato flour.

VISIA.	AUD AU				
s/n	Solubility Index	Solubility	Solubility	Solubility	Solubility
	(%) 55 ° C	Index (%)	Index (%)	Index	Index (%)
		65 ° C	75 ° C	(%)85 ° C	95 ° C
A.	349.50 <sup>a</sup>	335.00 <sup>b</sup>	2.50 <sup>b</sup>	6.50 <sup>b</sup>	5.00 <sup>a</sup>
B.	441.00 <sup>a</sup>	212.00 <sup>b</sup>	3.00 <sup>ab</sup>	$4.50^{ab}$	6.50 <sup>a</sup>
C.	333.50 <sup> a</sup>	259.50 <sup> a</sup>	$4.00^{ab}$	$4.50^{ab}$	7.00 <sup> a</sup>
D.	347.00 <sup>a</sup>	338.00 <sup>a</sup>	6.50 <sup>a</sup>	$2.00^{a}$	4.50 <sup>a</sup>

Sample A: Spontaneously fermented sweet potato flour, Sample B: Unfermented sweet potato flour; Sample C: 48
 hours starter culture fermented sweet potato flour; Sample D: 72 hours starter culture fermented sweet potato flour.

209 Values Are Average Of Two Determinations. Values in the same column not followed by the Same Superscript are

significantly different (P < 0.05).

#### 211 **DISCUSSION**

The functional property of sweet potato flour at different fermentation time is depicted in Table 1. A significant difference (p<0.05) was observed in the loose bulk density (LBD), packed bulk density (PBD) but there was no significant difference (p>0.05) in the water absorption capacity (WAC), dispersibility and oil absorption capacity (OAC).

Loose bulk density of the flour samples ranged between 0.488 and 0.607 g/ml with sample C having the lowest value while sample B had the highest value. Loose bulk density reveals the ability of a flour sample to occupy larger storage space per weight (Dossou *et al.*, 2014). This implies that sample C will occupy larger storage space while sample B will occupy less storage space.

Packed bulk density of the samples ranged between 0.701 and 0.801 g/ml, sample 222 D had the least value while B had the highest value. Packed bulk density is a 223 functional property that predicts the ease of transportation and packaging of 224 powdery products (Akinwale et al., 2017). Increase in fermentation time caused a 225 decrease in the packed bulk density although there was no significant difference in 226 the 48 and 72 hours fermentation period, this is in agreement with the findings of 227 Oloyede et al. (2016) who also reported a decrease in bulk density of the sample as 228 fermentation increased. This could be due to the effect of fermenting organisms on 229 the porosity of the flours and this probably implies that fermented flours up to 48 230 hours will occupy less space during packaging and more flour can be transported. 231

Hausner ratio of the flour samples ranged between 1.32 and 1.44. Hausner ratio is the ratio of packed bulk density to loose bulk density and this predicts the flow properties of food flour or powd and it has been reported that hausner ratio less

than 1.4 will facilitate conveying, blending and packaging of the flour/powder 235 which encourages its use in industrial food manufacture (Barbosa-Canovas et al., 236 2005; Ogunsina et al., 2010). Thus, due to the low hausner ratio of sample B and 237 D, this suggests that these samples may be the best applicable flour with the best 238 conveying and blending ability suitable for industrial use. There was no significant 239 difference (p>0.05) in the water absorption capacity of the fermented flours. Water 240 241 absorption capacity is a measure of the amount of water held by the protein matrix at room temperature. The values ranged from 1.48 to 1.61 g/ml, B had the least 242 while D had the highest value. This finding deviates from the findings of Oloyede 243 et al. (2016) who reported significant increase in water absorption capacity of 244 defatted Moringa oleifera seed flour as fermentation increased. This probably 245 implies that there was almost equal modification of the macromolecules during 246 fermentation exposing the hydrophilic bond of the sweet potato flours causing each 247 sample to have almost the same water affinity rate. There was no significant 248 difference (p>0.05) in the oil absorption capacity of the fermented sweet potato 249 flour. The oil absorption capacity ranged from 8.55 to 9.10 g/ml. All the samples 250 absorbed oil almost equally. This corroborates with the work of Fagberni, (1999) 251 which reported that good absorption of oil suggest its usefulness in preparation of 252 food products such as baked foods. 253

Significant difference was not observed (p>0.05) in the dispersibility of the samples. Dispersibility is a property that indicates the rate of reconstitution of flour sample in water.(Oluwole *et al.*, 2016).This probably indicates that the samples will disperse similarly in water.

The swelling power is depicted in Table 2. There was no significant difference (p>0.05) between the samples but numerically, slight increase in swelling power was observed with increased period of fermentation. This is in support with the findings of Oloyede *et al.*, (2016) who reported increase in swelling power of moringa seed flour with increase in fermentation time. Yuliana *et al.*, (2018) also reported increase in swelling power of sweet potato flour with increase in fermentation time. The increase in swelling power as the length of fermentation increased could be due to the modification of the sample's starch granules which disengaged the bonds allowing more water uptake and swelling effect to occur but this effect was not significant enough to indicate this modification.

The solubility index of fermented sweet potato flour is as shown in Table 3. There was no significant difference in the fermented sweet potato flour at 55 and 65 °C but significant difference was observed above 65 °C. All the samples at temperature range of 55 and 65°C had very high solubility values; this could be due to the gelatinization temperature of sweet potato that has not been attained. The gelatinization temperature of sweet potato has been reported to be between 64.6 to 84.6°C (Hoover, 2001).

The solubility above 65 °C increased with increase in fermentation time and it also increased as the temperature increased. The increase in the solubility observed above 65 °C may be due to the impact of fermentation period on the granular structure of the sweet potato as reported by Sobowale *et al.*, (2007) that when starch is heated in excess water above its gelatinization temperature, disruption of granular structure occurs causing molecules to disperse in solution.

Relating the swelling power and solubility values above 65 °C, the swelling power were high while the solubility index were low and it has been reported that high swelling power and low solubility are required for formation of highly viscous and elastic gels or dough (Baah *et al.*, 2005).This probably suggest that the fermented sweet potato flours will be beneficiary for viscous and elastic food products suchas dough in bread baking.

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