

**EFFECTS OF BOILING AND FERMENTATION ON
PHYSICOCHEMICAL PROPERTIES, FATTY ACID AND
MICRONUTRIENTS COMPOSITION OF *Hibiscus
sabdariffa* SEEDS**

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13 **ABSTRACT**

Aim: To ascertain the effects of boiling and fermentation on the physicochemical properties, fatty acid, mineral and vitamin composition of *Hibiscus sabdariffa* (HS) seeds.

Study design: Comparison-wise.

Place and Duration of Study: Rivers and Anambra states, Nigeria, between February and September, 2019.

Methodology: Two portions of 200 grams of HS seeds each were subjected to boiling and fermentation. The three samples were designated HSR, HSB and HSF for raw, boiled and fermented HS seeds respectively. Standard methods were used in determining the physicochemical properties and micronutrient composition, while fatty acid constituents were identified using a gas chromatography.

Results: The acid, free fatty acid, peroxidase values and specific gravity were significantly increased ($p < 0.05$), while iodine value was significantly reduced ($p < 0.05$) after boiling and fermentation. Saponification value showed a mixed trend, while refractive index was not significantly ($p > 0.05$) altered. Lauric (5.51–33.79%), palmitic (27.23–30.87%) and myristic (12.69–35.00%) acids were the predominant saturated fatty acids in HSR, HSB and HSF samples respectively. Oleic, linoleic, alpha-linolenic and arachidonic acids were the unsaturated fatty acids present in the samples. Boiling increased oleic acid level, while fermentation caused a drastic reduction (>90%) in its amount. Linoleic acid level improved up to 43% after fermentation. Magnesium, iron and sodium amounts significantly ($p < 0.05$) reduced after boiling and fermentation, while zinc, calcium and molybdenum levels were significantly ($p < 0.05$) improved after boiling. Na/K ratios for all the samples were greater than 0.60, while Ca/Mg values ranged between 0.82 and 3.46, below the recommended value (1.0). Vitamins B1, B3, B12 and D were significantly reduced ($p < 0.05$) after boiling and fermentation, while fermentation significantly increased ($p < 0.05$) vitamins B2, A, E and K levels.

Conclusion: Physicochemical properties of HS seeds suggest its favourable industrial applicability, whereas boiling and fermentation produced wide-ranging alterations in the micronutrient composition of HS seeds, most of which maximizes its usefulness as quality nutritional plant.

14
15 **Keywords:** [*Hibiscus sabdariffa* seeds, processing, physicochemical properties, fatty acid, minerals,
16 vitamins]

17 1. INTRODUCTION

18
19 *Hibiscus sabdariffa* Linn. is part of Malvaceae family believed to be from East Africa, Asia (India to
20 Malaysia) or Tropical Africa. *Hibiscus sabdariffa* (HS) Linn seeds are cultivated in many countries such as
21 Egypt, India, Mali, Malaysia, Nigeria, and Sudan have been found to contain high amount of protein,
22 dietary fiber, vitamins, lipids, and minerals [1 – 5]. Seeds of HS have already been noted as prolific and
23 were reported early in the century among African food grains, as being consumed in Northern Nigeria
24 after grinding into a coarse meal. They are highly regarded as a nourishing food [6]. They are crushed
25 and boiled in water to the consistency of a thin porridge and eaten as a sauce with staple foods among
26 the Banyoro of Uganda. In the Sudan, HS seeds are used as a seasoning after fermentation, and in the
27 South of Sudan the seeds are ground into flour [6].

28



29

30 Figure 1: *Hibiscus sabdariffa* seeds

31 In the northern regions of Cameroon, HS seeds are used to make “Mbuja” a condiment produced by
32 Fermentation. Mbuja is also known as Bikalga (Burkina Faso), Dawadawa botso (Niger), Datou (Mali),
33 Furundu (Sudan) [7].

34 According to Anioara-Arleziana *et al.* [8], physicochemical properties are imperative in determining the
35 overall stability and quality of food materials. Some of the important physicochemical properties are acid
36 value, specific gravity, iodine value, saponification value, peroxide value and refractive index. They are
37 used to monitor the compositional quality of oils. Fatty acids are inherent in plant oils and the property of
38 such oil is usually a function of the constituent fatty acids, which may either be a non-essential fatty acid
39 (omega – 9) or the essential fatty acids (omega – 3 and 6) gotten from the diet [9]. Great proportions of
40 unsaturated fatty acids are predominant in triglycerides from plant sources of oils, and the extent of
41 unsaturation is related to the extent of oxidative deterioration. Therefore, determination of fatty acid
42 composition of oils highlights the characteristics and stability of the oil.

43 Micronutrients are useful properties of food substances that enhance quality nutrition [10]. Minerals are
44 very important in human nutrition for proper metabolic activities and enzymatic actions in the body.
45 Magnesium is involved in regulating the acid-base balance in the body, utilization of iron and enzyme
46 activity, while calcium and magnesium play major roles in carbohydrate metabolism, nucleic acids and
47 binding agents of cell walls. Potassium is essential in synthesis of amino acids and proteins. Calcium
48 helps in teeth development. Iron is very essential in formation of haemoglobin in red blood cells; hence it

49 can help in stimulation of erythropoiesis. Vitamins can contribute to normal growth of body cells and skin,
50 proper immune function, normal vision, cell development, gene expression and maintenance of epithelial
51 cell functions [11].

52 Processing of seeds (such as boiling and fermentation) or other plant parts can either adversely affect or
53 improve their nutrient composition. Also, bioavailability, usefulness and utilization of nutrients in food
54 sources are seriously affected by the degree, nature or extent of processing they pass through. Boiling and
55 fermentation have been shown to significantly alter the quantities of nutrients and anti-nutrients in seeds
56 [12]. The impact of temperature on the stability, viscosity, peroxide value, iodine value to assess the
57 quality and functionality of the oil have been studied by Farhoosh *et al.* [13] and Li *et al.* [14].

58 **2. MATERIAL AND METHODS**

59 **2.1 Sample Collection and preparation**

62 *Hibiscus sabdariffa* seeds were collected from Mangu Local Government Area, Plateau State, Nigeria.
63 They were properly cleaned and sorted, ensuring that no debris was found in the sample. The cleaned
64 seeds of HS were pulverized into a fine powder with an electric blender and stored in a lid-tight container
65 for further analyses in the laboratory.

67 **2.2 Processing of *Hibiscus sabdariffa* Seeds**

68 **2.2.1 Boiling**

69 This was done according to the method modified from Mariod *et al.* [15]. *Hibiscus sabdariffa* (3 x 200 g) of
70 raw seeds of HS was boiled in 500 ml distilled water for forty (40) minutes till they become softened when
71 squeezed between the fingers. The cooked seeds were drained, dried, pulverized into fine powders and
72 stored in a tight-lid container for further analyses.

74 **2.2.2 Fermentation**

75 This was carried out using a modified method of Parkouda *et al.* [7]. After boiling and draining off water
76 from boiled seeds, the seeds were covered in a container and allowed to ferment for 3 – 4 days. They
77 were dried, ground into powder with an electric blender and stored in a tight container in a refrigerator.

79 **2.3 Determination of physicochemical parameters**

80 Standard methods were used in determining the physicochemical properties. Acid, saponification,
81 peroxide and iodine values were determined using the methods of A.O.A.C. [16]. Refractive indices were
82 analyzed using Abbe refractometer at 25 °C according to Oderinde and Ajayi [17]. pH was measured
83 electrometrically according to APHA [18] using an electric pH meter. Thiobarbituric acid value was
84 determined as mg malondialdehyde per kg sample.

85 **2.4 Determination of Fatty Acid Composition**

86 The fatty acid constituents were identified on a gas chromatography (Agilent 6890N) equipped with Flame
87 Ionization Detector and a 30 x 0.32m DB-225 silica capillary column (J and W Scientifics, USA). The split
88 injector (1 ml) and detector were operated at a temperature of 230 °C and 25 °C respectively, while the
89 oven temperature of 160 °C/2min was increased to 230 °C on a scale of 4 °C/min. Nitrogen was the
90 carrier gas at a flow rate of 1.5 ml/min. The peaks were compared with standard methyl esters while the
91 percentage area was recorded with standard Chemstation system.

93 **2.5 Determination of Mineral Composition**

94 Mineral composition was determined using Agilent FS240AA Atomic Absorption Spectrophotometer
95 (AAS) according to the method of American Public Health Association [19].

96 2.6 Determination of Vitamins

97 Retinol and tocopherol (vitamins A and E) were determined calorimetrically using the method of Kirk and
98 Sawyer [20]. Determination of thiamine, riboflavin, niacin and cobalamin (vitamins B1, B2, B3 and B12
99 respectively) and vitamin K were by spectrophotometric method while pyridoxine and ascorbic acid
100 (vitamins B6 and C respectively) were determined by titrimetric method according to Kirk and Sawyer
101 [20]. These methods are as described by AOAC [16].
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103 3. RESULTS AND DISCUSSION

104 3.1 Physicochemical Properties

106 The mean values of the physicochemical properties of the oils of raw (HSR), boiled (HSB) and fermented
107 (HSF) *Hibiscus sabdariffa* seed oils are shown in Table 1. Generally, acid value can be related to the
108 quality of the fatty acids in oil in terms of stability and shelf life. The acid values in this study were higher
109 than the acid value for raw, sun-dried and roasted groundnut oil (2.35, 1.79 and 2.52 mgKOH/g
110 respectively) as reported by Ayoola and Adeyeye [21]. It was however, lower than that found in *Duranta*
111 *repens* seed oil (21.01 mgKOH/g) as reported by Agomuo *et al.* [10] and *plukenetia aconophora* (11.5mg
112 KOH/g) as shown by Akintayo and Bayer [22]. The low acid values of the raw, boiled and fermented
113 seeds of HS (2.51, 4.02 and 4.66 mgKOH/g respectively) strongly suggest that the oil may be very
114 suitable for manufacture of soap, cooking, manufacture of margarine, mayonnaise, salad oils and
115 cosmetics. The free fatty acids (FFA) for HSR sample is similar to that of raw groundnut oil (1.18%) [21].
116 The FFA values were all below the maximum limit of 5.0% reported for Nigerian palm oil [23]. An increase
117 in the level of FFA in the samples may be as a result of hydrolysis of triglycerides which may occur by the
118 action of lipase enzyme, an indicator of processing and storage conditions (i.e., high temperature and
119 relative humidity, tissue damage) [24]. FFAs are sources of flavours and aromas. Samples with lower FFA
120 values tend to be soluble in water and volatile with characteristic smell, while samples with higher FFA
121 values are more prone to oxidation in their free form and their breakdown products (aldehydes, ketones,
122 alcohols, and organic acids) provide characteristic flavors and aromas [24]. The low levels of percentage
123 FFA in the three samples (1.26, 2.01 and 2.05% respectively) indicate that the oils from them may be
124 useful edible oils that may be stored for a long time without spoilage via oxidative rancidity. The peroxide
125 value of the samples were higher than that reported for water melon seed (3.24 mEq/kg) by Gladvin *et al.*
126 [25] and different refined groundnut species (1.30 – 1.73 mEq/kg) by Nkafamiya *et al.* [26] but lower than
127 those of *Opuntia dillenii* (15.60 mEq/kg) by Njoku *et al.* [27], crude groundnut oils (22.06 - 25.03 mEq/kg)
128 [26], *Duranta repens* leaf oil (20.00 mEq/kg) and *Duranta repens* seed oil (12.29 mEq/kg) by Agomuo *et*
129 *al.* [10]. The peroxide values of the samples of *Hibiscus sabdariffa* seeds increased possibly as a result of
130 boiling and fermentation. The peroxide value has been identified as the most common indicator for lipid
131 oxidation [28] and [29] demonstrated that peroxide values greater than 10 mEq/kg is an indication that
132 the oils are highly prone to auto-oxidation as a result of presence of trace elements or moisture. Such oils
133 can be unstable and may easily go rancid. The peroxide values of all the oil samples were less than the
134 standard peroxide value (10 mEq/kg) for vegetable oil deterioration and thus, suggest that they can be
135 put on storage for an elongated time without becoming rancid or deteriorating. The lower level of the
136 peroxide value of the raw sample suggests that it may have higher shelf life than the processed samples.
137 Fresh oils have value less than 10 mEq/kg and value between 20 and 40 mEq/kg leads to rancid taste
138 [30]. The low peroxide value indicated slow oxidation of these oils as suggested by Demian [31]. The
139 iodine value for raw seeds of HS (72.07 g₂/100g) was higher than 52.0 g₂/100g for palm oil [32], *Opuntia*
140 *dinelli* (63.33 g₂/100 g) by Njoku *et al.* [27] and similar to the iodine value of *Duranta repens* leaf oil (72.65
141 g₂/100 g) by Akubugwo *et al.* [33], while those of boiled (46.47 g₂/100g) and fermented (40.07 g₂/100g)
142 were lower. The values of the boiled and fermented samples were higher than that in *Cocos nucifera* (9.60
143 g₂/100g), *Pentaclethra macrophylla* (20.50 g₂/100g) and *Treculia africana* (27.50 g₂/100g) as reported
144 by Akubugwo *et al.* [33], whereas the iodine value of the raw sample was higher than them all. The iodine
145 value of the boiled sample is similar to that of crude Kampala Michika oil (46.88 g₂/100 g) [26] and *Duranta*
146 *repens* seed oil (44.84) [10], while that of the fermented sample was similar to that of *Citrullus vulgaris*
147 with 38.1% [34] and Hausa melon seed with 38.50% [35]. The iodine values of all the samples were lower
148 than that found in both crude and refined gargarjiya oil (81.94 and 97.13 g₂/100 g respectively) by
149 Nkafamiya *et al.* [26] and water melon seed oil (112 g₂/100 g) [25]. The reduction in iodine value after

150 processing was similar to the trend found in raw and heat-processed groundnut oil where the iodine value
151 reduced from 110.7 $\text{gI}_2/100 \text{ g}$ for raw groundnut to 100.7 $\text{gI}_2/100 \text{ g}$ for roasted groundnut oil as reported
152 by Ayoola and Adeyeye [21]. With the classification of Duel [36] for oils and fats, (drying oils: IV 200-130,
153 Semi drying: IV 130-100 and Non-drying: IV lower than 100), the samples all had iodine values less than
154 100 and therefore can be classified as non-drying oils in terms of industrial importance and also, as
155 classified by Aremu *et al.* [37]. The iodine value is the generally accepted parameter used in showing the
156 degree of unsaturation and number of carbon-carbon double bonds in fats or oils [38]. This value may be
157 useful in determining the amount of double bonds present in the oil which in turn reflects the susceptibility
158 of the oil to oxidation. The lower iodine values in the boiled and fermented samples in this study may
159 imply few unsaturated bonds found in them and hence low susceptibility to oxidative rancidity [39]. The
160 decrease in iodine value after processing (boiling and fermentation) may suggest lipid oxidation, which
161 could be as a result of presence of metal ions and other factors, which enhances or promotes oxidation
162 after the formation of hydroperoxide [40,41]. The SV of the raw sample of HS seed oil was similar to that
163 of Winsor orange-coloured cashew nut seed oil (212.00) by Aremu and Akinwumi [42], *Jatropha curcas*
164 seed oil (208.50) by Igwenyi [43] and yellow melon seed oil (210.00) by Egbebi [44]. The SV of the boiled
165 sample of HS seeds was similar to that of melon seed oil (148.50) reported by [45] and Almond seed oil
166 (151.55) as reported by Ogunsuyi and Daramola [46], while that of fermented sample was similar to
167 coconut oil (248-265) [47] and *C. nucifera* (246.00) as reported by Amoo *et al.* [48]. The saponification
168 value of oils is of interest when considering using the oil for industrial purposes [49]. Saponification value
169 is applicable in tracking adulteration [50]. The larger the saponification values of oil, the better their soap-
170 making abilities [51]. The saponification values greater than 200 mgKOH/g may indicate high proportion
171 of unsaturated short chain fatty acids in the samples and may promote stability of the oil. This shows that
172 they have a very high potential use in soap making and food industries. Denniston *et al.* [52] reported that
173 high saponification value indicated the presence of greater ester bonds, suggesting that the fat molecules
174 were intact. These properties make it useful in soap making industry. Furthermore, the high saponification
175 values indicate oxidation and its decrease suggest the onset of oxidation. Rossel [53] reported similar
176 observation. TBA values are used in assessing the level of oxidation of fats and oil (lipid oxidation) in
177 terms of the amount of malondialdehyde (secondary product of oxidation of fats and oil) present in a
178 sample. The presence of thiobarbituric acid in the samples suggests that some forms of oxidation had
179 taken place as suggested by Lukaszewicz *et al.* [54]. These values may be useful in carrying out sensory
180 tests aimed at ascertaining rancidity in food systems as suggested by these authors [55 – 57]. The raw
181 and fermented samples had higher TBA values than the boiled sample in this study. The refractive index
182 (RI) of the samples were 1.40, 1.42 and 1.40 for raw, boiled and fermented samples respectively. These
183 values were less than the standard values for refined and virgin oils (1.4677–1.4707) according to
184 CODEX-STAN [32]. However, they were higher than the RI of melon seed oil (1.35) as ascertained by
185 Edidiong and Ubong [58], while the RI of the boiled sample was found to be same as that of cashew nut
186 seed oil (1.420) by Aremu and Akinwumi [42]. The RI of an oil denotes the ratio of speed of light to its
187 speed in the oil/fat itself, at a particular wavelength. The RI is important during quality control by indicating
188 isomerization and hydrogenation which are necessary when ascertaining the purity of a substance [10].
189 The pH ranged from 4.67 – 6.17, with the fermented sample being the most acidic (4.67). The pH of the
190 raw sample is similar to the pH of *Duranta repens* seed oil (6.16) as determined by [10]. The decrease in
191 the pH may be attributed to the effect of microorganisms, which produces carbon dioxide during
192 fermentation, thereby making the samples more acidic. This can be influenced by the duration of the
193 fermentation process. The pH and acid values are used to assess the quantity of free fatty acids present
194 in oils and can as well, determine their shelf life and stability [10]. The SG of the raw sample was similar
195 to those of Koto/Pteryogota seed oil (0.930), *Pteryogota macrocarpa* (0.928) and Luffa gourd seed
196 (0.930) as reported by Amoo and Agunbiade [59] and Oluba *et al.* [60] respectively. The boiled and
197 fermented samples had SG values similar to Castor seed oil (0.959) and Cashew nut seed (0.964) as
198 determined by Akpan *et al.* [61] and Aremu *et al.* [62]. The SG in the current study were higher than found
199 in Melon seed oil (0.850) by Edidiong and Ubong [58], groundnut seed oil (0.914) by Musa *et al.*, [63] and
200 pumpkin seed oil (0.830) by Akubugwo *et al.* [33]. Whereas, they were found to be lower than SG of
201 *Duranta repens* seed and leaf oils (1.64 and 1.02) [10] and Almond seed oil (1.71) by Akpambang *et al.*
202 [64]. The result showed that oils of the sample in the present study are less dense than water (1 g/cm^3)
203 and therefore may find application in cream production, because it could make the oils flow and can
204 easily be spread on the skin [45]. SG can be used alongside other figures in assessing the purity of oil
205 [65].

206 **Table 1: Physicochemical analysis of the oil of raw, boiled and fermented *Hibiscus sabdariffa***
 207 **seeds**

Parameters	HSR	HSB	HSF
Acid value (mgKOH/g)	2.51 ± 0.01 ^a	4.02 ± 0.04 ^b	4.66 ± 0.12 ^c
Free fatty acid (%)	1.26 ± 0.01 ^a	2.01 ± 0.21 ^b	2.05 ± 0.01 ^b
Iodine value (gI ₂ /100g)	72.07 ± 2.04 ^c	46.47 ± 4.01 ^b	40.07 ± 3.10 ^a
Peroxide value (mleq/kg)	4.40 ± 0.20 ^a	9.6 ± 0.50 ^c	8.25 ± 1.45 ^b
Saponification value (mgKOH/g)	210.10 ± 8.57 ^b	148.72 ± 7.11 ^a	256.68 ± 10.20 ^c
Thiobarbituric acid (mg.mal/kg)	3.58 ± 0.06 ^b	2.63 ± 0.30 ^a	3.58 ± 0.10 ^b
pH	6.17 ± 0.01 ^c	5.20 ± 0.03 ^b	4.67 ± 0.07 ^a
Refractive index	1.40 ± 0.01 ^a	1.42 ± 0.00 ^a	1.40 ± 0.02 ^a
Specific gravity	0.93±0.02 ^a	0.99±0.05 ^b	0.97±0.01 ^b

208 Values are means of three determinations ± standard deviation (SD). At ($P < 0.05$), means with different
 209 superscripts in a row are significantly different from each other.

211 3.2 Fatty acid profile

212 The fatty acid profile of the samples (HSR, HSB and HSF) is presented in table 2. The results showed
 213 that Lauric (5.51 – 33.79%), palmitic (27.23 – 30.87%) and myristic (12.69 – 35.00%) acids were the
 214 predominant saturated fatty acids in HSR, HSB and HSF samples respectively. Oleic, linoleic, alpha
 215 linolenic and arachidonic acids were the unsaturated fatty acids present in the samples. Oleic acid was
 216 found in all the samples (HSR – 15.85%, HSB – 21.50% and HSF – 0.79%), linoleic in HSB (19.85%) and
 217 HSF (34.84%), alpha linoleic in HSF (2.18%) and arachidonic acid only in HSR (2.36%). Generally, the
 218 levels of lauric, palmitic and myristic acids in this study were higher than those reported by Rao [2] for
 219 mesta (*Hibiscus sabdariffa*) seeds and *Duranta repens* leaf oil [10]. Kostik *et al.* [66] reported higher
 220 amounts of lauric acids in coconut (48%) and palm kernel (41%) oils, and lower amounts of myristic acid
 221 in corn oil (0.6%), cottonseed (0.4%) and Safflower (0.5%). However, Ahmad *et al.* [67] reported similar
 222 amount of palmitic acid in HSB (30.87%) for *Hibiscus sabdariffa* seed oil; Agomuo *et al.* [10] in HSR and
 223 HSB in myristic acid for *Duranta repens* seed oil. Also, Al-Wandawi *et al.* [68] and Ahmed and Hudson
 224 [69] reported similar palmitic acid levels in Iraqi karkade cultivars (17.85–28.46) and crude karkade seed
 225 oil (20.5%) respectively. The stearic acid levels in HSR and HSB were lower than that of *Duranta repens*
 226 leaf (6.78%) and seed (8.05%) oils [10], Canola type 2 oil (6.9%) [66] and mature stems of *Opuntia dillenii*
 227 [27], but similar to soybean (4.00%), peanut (4.50%), and Canola type 1 (5.2%) oils [66] and crude
 228 Karkade seed oil (5.8%) by Abu-Tarboush [70]. The stearic acid level of HSF (19.23%) was much higher
 229 than all the oils mentioned above. HSR, HSB and HSF had stearic acid levels higher than mesta seed
 230 (2.4%), sunflower seed (2.0%), linseed (3.5%), cotton seed (2.0%), palm kernel (2.0%) and coconut
 231 (2.0%) oils reported by Kostik *et al.* [66]. The ratio of unsaturated to saturated fatty acids (SFAs) was
 232 found to be low, compared to another study by Soheir and Deba [71]. This may be as a result of
 233 geographical factors, growing conditions, degree of maturation etc. This implies that the samples had
 234 more SFAs and short chain FA may be used in chemical industries for soap and cosmetic production [72].
 235 However, many studies have reported the harmful impacts small chain fatty acids on the human body by
 236 mainly lowering HDL cholesterol and increasing LDL cholesterol [73]. The oleic acid content in HSR and
 237 HSB were higher than that in coconut oil (8.8%) [66], egusi melon oil [74] and *Duranta repens* seed oil
 238 (11.47%) [10]; but lower than in crude Karkade seed oil [70], and groundnut oil (44.90%), cashew seed oil
 239 (34.47%) and pumpkin seed oil (36.10%) [74]. The oleic acid composition in this study is comparable to

240 that of Safflower (16.6%) and Linseed (22.5%) [66] and rubber seed oil (23.74%) [74]. In comparison to
 241 the findings of Kostik *et al.* [66], the linoleic acid contents of the samples in this study were higher than
 242 those in coconut (0.5%), palm kernel (1.25%) and olive (7.0%) oils, much lower than those in corn
 243 (48.0%), soybean (49.5%), sunflower (59.5%) oils and similar to those in linseed (20.5%), peanut (20.0%)
 244 and canola variety 1 (18.8%) oil. Bello and Anjorin [74] also reported linoleic acid content in groundnut oil
 245 (32%) and cashew seed oil (34.47%) similar to HSF (34.84%) in this study. Also, Okra seeds contain
 246 31.48% linoleic acids [75]. From the results of this study, the samples had lower amounts of unsaturated
 247 fatty acids. However, the unsaturated fatty acids were more concentrated in HSB (41.35%) and HSF
 248 (37.81%) samples, but they were all lower than the saturated fatty acids. Polyunsaturated fatty acids are
 249 essentially fatty acids needed for normal growth, physiological functioning and maintenance of the body.
 250 Linolenic acid is an omega – 3 polyunsaturated fatty acid (PUFA) involved in the regulation of biological
 251 functions and management of a many human diseases like hearth and inflammatory diseases [76].
 252 However, further increase in PUFA may predispose the oil to oxidation [77]. The presence of oleic acid,
 253 linoleic, alpha linolenic and arachidonic acids suggests that the samples may find industrial applicability
 254 for pharmaceuticals, soaps, shampoo and cosmetics productions. Unsaturated fatty acid improves lipid
 255 profile, whereas excess consumption of SFAs may cause obesity and elevated cholesterol levels [78].
 256 Boiling and fermentation increased the levels of the SFAs – magaric, myristic and stearic acids, while the
 257 level of lauric acid reduced after boiling and fermentation. Varied effects of boiling and fermentation on
 258 the unsaturated fatty acids were observed in the study. While boiling increased the amount of oleic acid,
 259 fermentation caused a drastic reduction (> 90%) in its amount. However, the amount of linoleic acid
 260 improved by up to 43% after fermentation. These alterations may be as a result of the breakage of the
 261 fatty acid bonds or their complete degradation. Fermenting microorganisms may also contribute to the
 262 breakdown of fatty acids.

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Table 2: Fatty acid profile of raw, boiled and fermented *Hibiscus sabdariffa* seed

Fatty acid	% Composition		
	HSR	HSB	HSF
SATURATED FATTY ACIDS			
C8 = Caprylic acid	0.91	ND	ND
C12 = Lauric acid	33.79	5.51	10.14
C14 = Myristic acid	13.36	12.69	35.00
C16 = Palmitic acid	27.23	30.87	ND
C17 = Magaric acid	1.47	4.49	ND
C18 = Stearic acid	4.17	5.10	19.23
C20 = Arachidic acid	1.14	ND	ND
UNSATURATED FATTY ACIDS			
C18:1 = Oleic acid	15.58	21.50	0.79
C18:2 = Linoleic acid	ND	19.85	34.84

C18:3 = Alpha linolenic acid	ND	ND	2.18
C20:4 = Arachidonic acid	2.36	ND	ND

ND = Not detected

3.3 Mineral Composition and Mineral ratios

The results of the mineral composition of HSR, HSB and HSF are presented in table 3. Magnesium, iron and sodium amounts (21.35, 24.08 and 17.24 mg/kg respectively) in HSR were significantly higher ($p < 0.05$) than in HSB (5.95, 10.00 and 8.40 mg/kg respectively), which was also significantly higher than in HSF (3.74, 5.28 and 2.79 mg/kg respectively). Zinc and calcium contents of HSR (11.06 and 20.60 mg/kg respectively) were significantly higher ($p < 0.05$) than in HSB and HSF, while lead and molybdenum contents were significantly higher in HSB (0.40 and 0.19 mg/kg) than in HSR and HSF. Lead (0.07 – 0.1 mg/kg), cobalt (0.09 mg/kg) and molybdenum (0.02 – 0.19 mg/kg) were found to be in least amounts. The variations in the levels of the macronutrients may be as a result of different geographical locations, methods of cultivation, soil types, processing methods etc, which they were subjected to. During boiling, there are tendencies of these nutrients to be leached into the boiling water, thereby causing their loss. The decrease in magnesium after boiling is in consonance with the observation of Tounkara *et al.* [79] which investigated the effect of boiling of the physicochemical properties of Roselle seeds in Mali. However, there was a disagreement in the results for other elements. All the Na/K values were greater than 0.60 (4.03, 2.29 and 1.09) for HSR, HSB and HSF respectively. This is the ratio that favours none enhancement of high blood pressure disease in man [80]. This denotes that the samples may not be suitable for managing high blood pressure. To bring this ratio low, consumption of foods rich in potassium is highly encouraged. The Ca/Mg values ranged between 0.82 and 3.46 whereas the recommended value is 1.0 [80]. Therefore, only HSB (3.46) and HSF (1.51) had the recommended Ca/Mg level. Both Ca and Mg would need adjustment for good health.

Table 3: Mineral Composition of raw, boiled and fermented *Hibiscus sabdariffa* seeds

Parameters	Concentration (mg/kg)		
	HSR	HSB	HSF
Magnesium	21.35 ± 0.68 ^c	5.95 ± 0.05 ^b	3.74 ± 0.58 ^a
Lead	0.12 ± 0.02 ^a	0.40 ± 0.10 ^b	0.07 ± 0.02 ^a
Manganese	2.45 ± 0.07 ^b	ND	0.08 ± 0.02 ^a
Copper	1.02 ± 0.06 ^b	0.22 ± 0.01 ^a	0.18 ± 0.01 ^a
Iron	17.24 ± 0.63 ^c	8.40 ± 0.10 ^b	2.79 ± 0.12 ^a
Zinc	8.25 ± 0.09 ^b	11.06 ± 0.59 ^c	0.90 ± 0.10 ^a
Cadmium	0.43 ± 0.03 ^b	0.47 ± 0.09 ^b	0.10 ± 0.00 ^a
Molybdenum	0.02 ± 0.02 ^a	0.19 ± 0.01 ^b	ND
Sodium	24.08 ± 1.21 ^c	10.00 ± 0.26 ^b	5.28 ± 0.19 ^a
Potassium	5.98 ± 0.21 ^b	4.37 ± 0.46 ^a	4.84 ± 0.07 ^a
Calcium	17.46 ± 0.13 ^b	20.60 ± 0.20 ^c	5.64 ± 0.09 ^a

Aluminum	0.96 ± 0.68 ^b	ND	0.01 ± 0.01 ^a
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290 Values are means of three determinations ± standard deviation (SD). At ($P < 0.05$), means with different
 291 superscripts in a row are significantly different from each other. HSR = Raw *Hibiscus sabdariffa* seeds;
 292 HSB = Boiled *Hibiscus sabdariffa* seeds and HSF = Fermented *Hibiscus sabdariffa* seeds. ND = Not
 293 detected

294 **Table 4: Mineral ratios of raw, boiled and fermented *Hibiscus sabdariffa* seeds**

Mineral Ratios	HSR	HSB	HSF
Na/K	4.03	2.29	1.09
Ca/K	3.07	4.71	1.17
Ca/Mg	0.82	3.46	1.51
Zn/Cu	8.09	50.27	5.00
Fe/Cu	16.9	38.18	15.50

296 Values are calculated from means of mineral concentrations

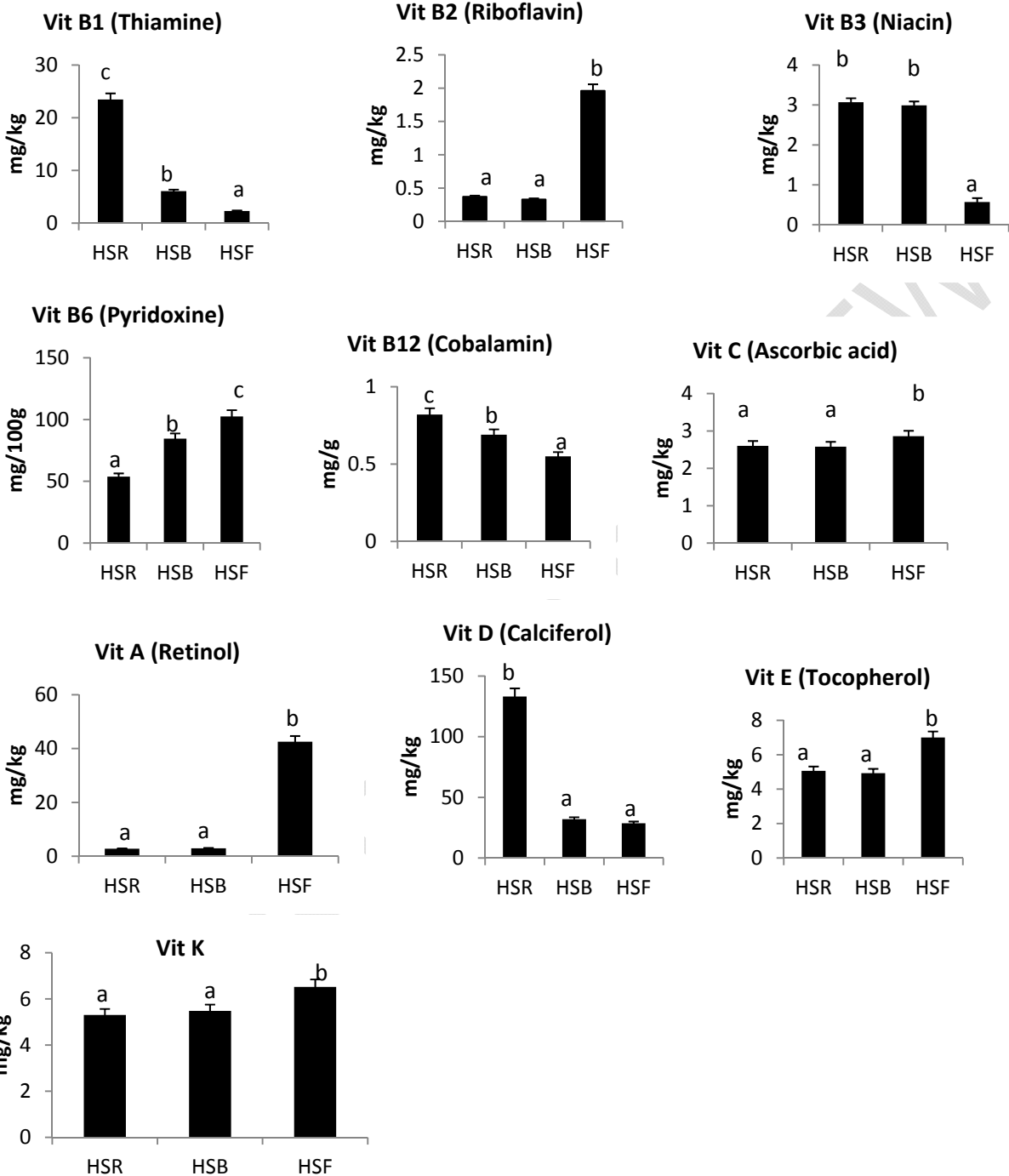
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300 3.4 Vitamin composition

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 302 The results of the vitamin composition of HSR, HSB and HSF are presented in figure 1. Vitamin B1
 303 (thiamine) (23.45 mg/kg) and vitamin D (calciferol) (133.17 mg/kg) contents in HSR were significantly
 304 higher ($p < 0.05$) than in HSB and HSF which did not differ significantly ($p > 0.05$) from each other. Vitamins
 305 B2 (riboflavin), B6 (pyridoxine), E (tocopherol), A (retinol), C (ascorbic acid) and K contents in HSF (1.96
 306 mg/kg, 102.50 mg/100g, 7.00 mg/kg, 42.53 mg/kg, 2.86 mg/kg and 6.52 mg/kg respectively) were
 307 significantly higher ($p < 0.05$) than in HSR and HSB. Vitamin B12 (cobalamin) was significantly lower
 308 ($p < 0.05$) in HSF than in HSB and HSF. Vitamins B2 (riboflavin), E (tocopherol), A (retinol), C (ascorbic
 309 acid) and K contents in HSR and HSB were not significantly different ($p > 0.05$) from each other. Vitamin
 310 B6 content in HSR (53.74 mg/100g) was significantly lower than in HSB and HSF. The vitamin
 311 compositions of the seeds of HS show that they are good sources of vitamins and the presence of these
 312 vitamins can contribute to normal growth of body cells and skin, proper immune function, normal vision,
 313 cell development, gene expression and maintenance of epithelial cell functions [11]. Vitamin B6 was
 314 present in a reasonable amount and it helps in formation of red blood cells and maintenance of brain
 315 function. This vitamin also plays an important role in the proteins that are part of many chemical reactions
 316 in the body. Vitamin B12 is involved in formation of red blood cells and vitamin K aids in blood clotting
 317 [81]. Vitamin C is important for proper body function and its deficiency may interfere with the normal
 318 formation of intracellular substances which could lead to impaired growth and development in the body. It
 319 is also crucial in the maintenance and repair of tissues such as bones, skin and teeth. The antioxidant
 320 vitamins (A, C and E) were present in the raw, boiled and fermented seeds of HS and they neutralize free
 321 radicals that can accumulate in the body which in turn, leads to aging and some diseases. Therefore, the
 322 seeds of HS may possess ameliorative potentials if supplemented with other anti-oxidant rich plants
 323 against diseases linked with oxidative stress. The reduction in vitamin B1 (thiamine) and B3 (Niacin)
 324 contents after boiling is in agreement with the earlier reports of Fadahunsi [82] on Bambara groundnut
 325 flour, Prinyawiwatkul *et al.* [83] on cowpeas and Barampama and Simard [84] on beans. Further decrease
 326 in the amount of thiamine after fermentation is also in line with Fadahunsi [82] on Bambara groundnut
 327 flour, Philips *et al.* [85] on fermented cowpea and Wang and Hesseltine [86]; Murata *et al.* [87] on
 328 fermented soybeans; Van Veen *et al.* [88] and Keuth and Bispring [89] on fermented wheat. This

329 decrease in the amount of vitamin B1 may be due to rapid utilization of vitamin B1 for optimum growth
330 and other functions at a higher rate than its synthesis by the fermenting organisms [86]. For Vitamin B2
331 (Riboflavin), there was a reduction in its amount after boiling (though not significant). Similar report was
332 given by Fadahunsi [82] for Bambara groundnut flour, Philips *et al.* [85] and Uzogara *et al.* [90] for boiled
333 cowpea, whereas Deosthale [91] reported such in chicken peas and green peas. The significant ($p < 0.05$)
334 increase in vitamin B2 after fermentation is in tandem with Fadahunsi [82] on Bambara groundnut flour
335 and Philips *et al.* [85] on fermented cowpea. Fermentation also significantly increased the amount of the
336 antioxidant vitamins (A, C and E) and K. This increase in vitamins was also reported by Akinyele and
337 Akinlosotu [92] on fermented cowpeas and Eka [39] on fermented locust bean. However, it disagrees with
338 Barampama and Simard [84] which reported a decrease in the vitamin content of beans after
339 fermentation. The variations in the levels of the vitamins may be as a result of different geographical
340 locations, methods of cultivation, type of soil, processing methods e.t.c., which they were subjected to.

UNDER PEER REVIEW



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Figure 1: Vitamin composition of HSR, HSB and HSF. Values are means of three determinations \pm standard deviation (SD). At $P < 0.05$, bars with different superscripts in a chart are significantly different from each other

4 CONCLUSION

The results of the physicochemical analysis showed that raw and processed *Hibiscus sabdariffa* seed oils are suitable for industrial applications in terms of shelf life, stability, density and resistance to auto-oxidation. The seeds are majorly composed of saturated fatty acids and also some polyunsaturated fatty acids. Minerals and vitamins were detected in reasonable amounts. Processing caused varied alterations

354 in the micronutrient composition of HS seeds, most of which maximizes its usefulness as quality
355 nutritional plant.

356

357 **COMPETING INTERESTS**

358

359 Authors have declared that no competing interests exist.

360

361 **AUTHORS' CONTRIBUTIONS**

362 This work was carried out in collaboration among all authors.

363 **CONSENT (WHERE EVER APPLICABLE)**

364

365 None

366

367

368 **ETHICAL APPROVAL (WHERE EVER APPLICABLE)**

369

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595
596

597 APPENDIX

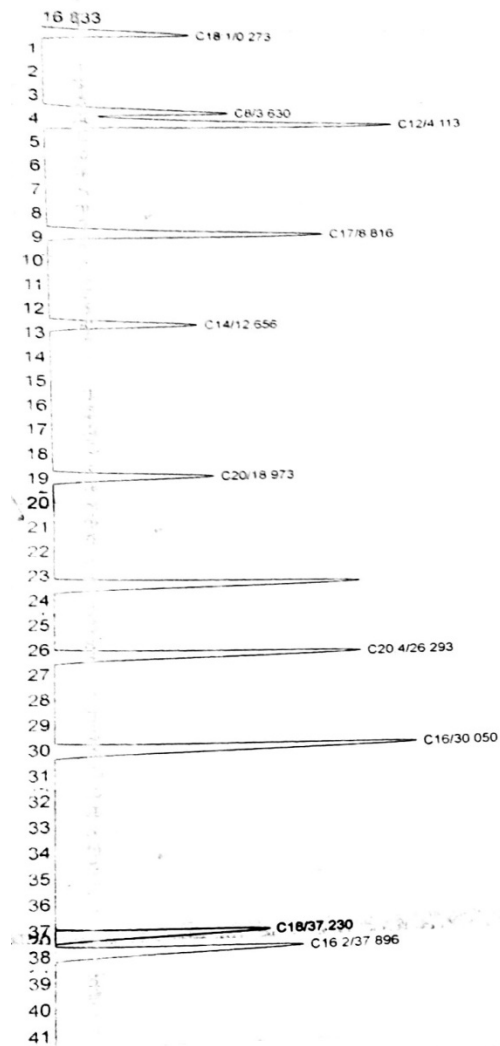
598

599 A Chromatogram of fatty acid profile of raw *Hibiscus sabdariffa* seed

Client: Charles
Client ID: DA134
Collected: 24/01/18
Description: FID
Column: RESTEK 15METER MXT-1 HSR
Carrier: HELIUM AT 5 PSI
Components: fame standard cpt
Data file: Charles fatty acid profile ()
Sample: fatty acid profile
Comments: TYPE YOUR COMMENTS HERE

Events

Time Event



2408

Component	Retention	Area	Height	External	Units
C18 1	0 273	4636 9515	273 847	12 6256	ppm
C8	3 630	15478 1068	339 145	0 7386	ppm
C12	4 113	11295 9556	632 310	27 3788	ppm
C17	8 816	8838 2628	500 548	1 1903	ppm
C14	12 656	4803 1439	272 736	10 8266	ppm
C20	18 973	5160 5610	293 146	0 9214	ppm
C20 4	26 293	9544 9712	539 008	1 9148	ppm
C16	30 050	11160 0115	630 662	22 0624	ppm
C18	37 230	6529 9332	371 001	3 3792	ppm

600

B Chromatogram of fatty acid profile of boiled *Hibiscus sabdariffa* seed

Lab name: Springboard Lab
Client: Charles
Client ID: DA134
Collected: 25/09/18
Method: Syringe Injection
Description: FID
Column: RESTEK 15METER MXT-1
Carrier: HELIUM AT 5 PSI
Control filename: DEFAULT CON
Data file: Charles Fatty acid profile ()
Sample: fatty acid profile
Comments: TYPE YOUR COMMENTS HERE

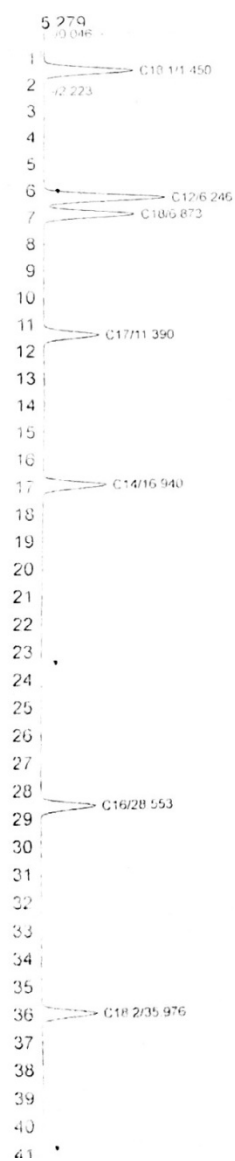
HSB

Temperature program

Init temp	Hold	Ramp	Final temp
70.00	10.000	10.000	220.00
220.00	5.000	5.000	280.00

Events

Time	Event
0.000	ZERO
0.000	SOUND



C Chromatogram of fatty acid profile of fermented *Hibiscus sabdariffa* seed

Lab name: Springboard Lab
 Client: Charles
 Date: 29/07/18
 Method: Syringe Injection
 Description: FID
 Column: RES1EK 15METER MXT-1
 Carrier: HELIUM AT 5 PSI
 Data file: Charles HSF fatty acid profile ()
 Comments: TYPE YOUR COMMENTS HERE

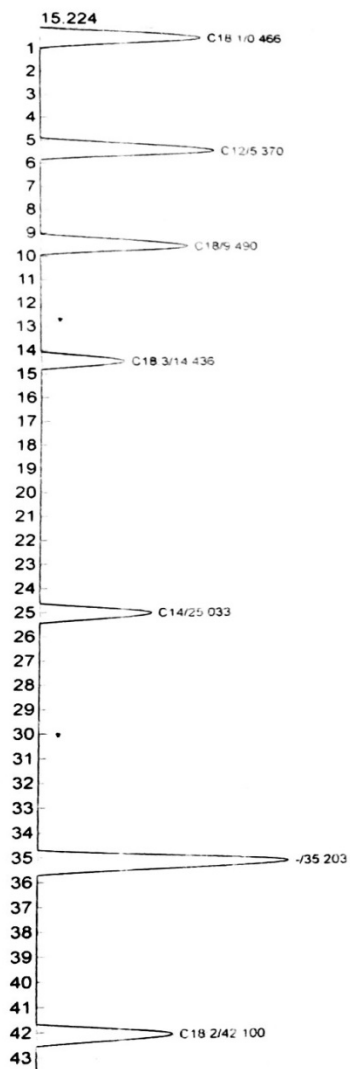
HSF

Temperature program

Init temp	Hold	Ramp	Final temp
80.00	5.000	5.000	220.00
220.00	2.000	5.000	300.00

Events

Time	Event
0.000	ZERO
0.000	SOUND



977

Component	Retention	Area	Height	External	Units
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