Comparative Influence of Dehulling on the Composition, Antioxidative and Functional Properties of Sorrel (Hibiscus sabdariffa L.) Seed

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Abstract

Although animal proteins provide indispensable amino acids that the body requires for normal growth, maintenance and function, their expensiveness makes them unaffordable especially for most families in the developing countries. This has given impetus to extensive research into under-utilized protein-rich oilseeds such as sorrel as possible alternate sources of good quality protein for tackling the challenge of protein-energy malnutrition which is fast becoming a global challenge. Sorrel seed may hold great potentials as a source of good quality protein, however the presence of hard seed coat, bitter after-taste and associated antinutritional factors have limited its use as protein supplement for humans and food ingredient. This study therefore compared the effect of dehulling sorrel seed to boiling, germination and roasting. This was with the aim of enhancing its utilization as protein source for human nutrition and functional ingredient in food product development. Flours obtained were analyzed for their proximate, mineral, antinutrient, amino and fatty acids composition; in vitro starch and protein digestibility, and functional and antioxidative properties. Protein content (ranged from 24.93% - 32.91%) significantly increased due to processing; dehulling alone accounted for a percentage increase of 32.01%. Similarly, dehulling increased all essential amino acids (except isoleucine and valine) at percentage which ranged from 3.63% - 61.17% whereas other processing methods caused significant reductions. Lysine, leucine, valine, arginine and phenylalanine were the most abundant essential amino acids, while methionine and cystine were the first and second limiting amino acids. Palmitic, linoleic, oleic and stearic acids were the most abundant fatty acids. Mineral composition was K > Ca > Mg > Na > Fe > Zn > Mn. Dehulled seed flour had highest in vitro protein digestibility.
(75.87%). Improved amino acid composition, antioxidative and functional properties of sorrel seed flour due to dehulling may indicate the potential of this flour to serve as a protein supplement and functional ingredient for food product development.

**Keywords**

Dehulling, Amino Acid Composition, Functional Properties, *In Vitro* Protein Digestibility, Human Nutrition, Processing, Protein-Energy Malnutrition, Under-Utilized Sorrel Seed

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1. **Introduction**

Malnutrition, currently one of the most devastating challenges being faced globally, is the most important risk factor for illness and death worldwide; with hundreds of millions of pregnant women and young children particularly affected [1] [2]. Particularly in developing countries (Asia and Africa), protein-energy malnutrition (PEM) is the major health challenge and leading nutrition problem most prevalent. This situation is further worsened by the increasing level of poverty which has been reported to be the major cause of malnutrition [3]. Hence, diets of most households in these regions are frequently deficient in macronutrients (protein, carbohydrates and fat, leading to protein-energy malnutrition), micronutrients (electrolytes, minerals and vitamins, leading to specific micronutrient deficiencies) or both [2] [4]. Although animal proteins provide indispensable (essential) amino acids that the body requires for synthesis of tissue and organ proteins and other nitrogen-containing compounds necessary for normal growth, maintenance and function [5]; their relative expensiveness makes them largely unaffordable for most families especially in the developing countries. Thus, most staple foods in these regions are largely composed of starch.

The major nutrition challenge for governments of both developing and industrialized societies is being able to provide their populations with adequate amounts of food proteins to meet physiological and nutritional requirements [6]. Direct human consumption of oilseeds has the potential to improve the protein content and quality of diets of over half of the world’s population. Hence, Müller and Krawinkel [2] have suggested the use of locally available protein- and micronutrient-rich leguminous plants as an effective and sustainable intervention needed to tackle the problems of protein-energy malnutrition and micronutrient deficiency in developing countries. This is because in contrast to low protein content of cereal grains (6% - 14%), oilseeds and legumes have high protein content (20% - 25%), are adapted to grow under a wide variety of climatic conditions, are relatively cheap and since they are already a part of the diet in many parts of the world, efforts to increase their consumption is greatly simplified [7]. Usually, legumes (rich in lysine but poor in methionine and cystine)
are complemented with starchy foods such as cereals (deficient in lysine but rich in methionine and cystine), roots and tubers like rice, millet, maize, yam, cassava, potatoes and sorghum to provide an alternative source of dietary protein of vegetable origin [8]. Extensive research on a number of these legumes (including soybean, cottonseed, sunflower and peanut) has resulted in their wide acceptability and utilization in human nutrition at both domestic and industrial levels. This has made them almost as expensive as animal protein and in some cases not readily available since quantities produced are never enough to meet the high demands for production of a wide range of products, isolates, concentrates, oil and flour [9]. This has necessitated exploiting new protein sources from other under-utilized protein-rich oilseeds; as such attention has been recently shifted to under-utilized protein-rich legumes and oilseeds that can be used as alternative protein sources in food ingredients and functional foods formulation [2] [10].

Sorrel (*Hibiscus sabdariffa* L.) seed is one of the under-utilized protein-rich seeds with promising nutritional potentials and is currently attracting research interest as a source of good quality protein [11] [12]. A member of the *Malvaceae* family, sorrel plant also known as Roselle is one of the over 300 species of hibiscus around the world and one of the most common flower plants grown worldwide. This is because it is relatively easy to grow, can be grown as part of multi-cropping systems and is usually grown as a home garden crop due to its ease to maintain [13] [14]. In comparison with other seeds such as black seed, sunflower seed, melon seed, chickpea, pigeon peas, cowpea, soybean and groundnut, sorrel seed contains higher protein content (32.28% - 34%) and its essential amino acid profile is comparable to soybeans. It also contains high amounts of cellulose, dietary fibre, minerals such as phosphorus, calcium and magnesium, vitamins C and E; a good source of cholesterol-free vegetable oil rich in unsaturated and essential fatty acids like linoleic acid. The high dietary fibre (39% - 42%) of the seed contributes to its strength when compared with other common sources such as wheat, rice bran, oat and fibre [13]-[18]. Hence, this highly-nutritious seed possesses features that can make it serve as a potential good quality protein source that can supplement the often high starch staples frequently consumed in developing countries.

Despite these, sorrel seed is still grossly under-utilized both as protein supplement for humans or functional ingredient in food product development; as such large quantities are usually discarded as by-products after removal of the calyces in most producing areas. Its major use is as animal feed, although the fermented extract of the seed has been reportedly used in some countries as condiment for soup preparation while the residue is mostly thrown away or used as animal feed [19]. This process avoids the ingestion of the whole seed kernel, thus depriving humans of essential nutrients that could be derived from actual consumption of the seed kernel. This is due to the bitter after-taste associated with its hard seed coat which is also high in antinutritional factors and food toxicants [14] [20]. This has also limited the use of sorrel seed in food product de-
velopment and till date no study has reported any final product application for sorrel seed or any use as protein supplement for human nutrition. According to Carter et al. [21], processing technology is one of the factors that limit the maximum utilization of oilseeds; appropriate processing techniques are very vital to harnessing the abundant nutrients in these plant proteins for human nutrition while inefficient processing techniques limit maximum utilization of legumes and oilseeds. Hence, processes such as drying, sundrying, boiling, roasting, fermentation and germination which have been extensively reported to reduce antinutrient contents and improve nutritional value and digestibility of sorrel seed [14] [15] [16] [17] [18] have not transformed to wide acceptance and maximum utilization of the seed for human nutrition. Hence, to explore its potential as a protein source and food ingredient and make its nutrients more easily accessible, the seed coat has to be removed (dehulled). Apart from seed coat/hull removal, dehulling also enhances texture, appearance, cooking time, digestibility of protein, palatability, nutritive value and colour of food due to removal of antinutritional and toxic factors which abound in the seed coat [22].

Presently, information is lacking on the effect of dehulling on sorrel seed composition and this is vital to effectively maximize it as a protein supplement and functional ingredient for formulation of new food products. Information is also lacking on the functional properties of both raw and processed sorrel seed flours, while reports on the antioxidative potential and bioactive composition of the processed and raw seed flours (which provide information on the health-promoting potential of the seed) are scarce. Information on functional properties is relevant to determine the level of utilization of flours in ingredient formulation and food product development [23]. This present study has therefore compared the effect of dehulling to two heat processing methods (boiling and roasting) and germination on the nutrient and antinutrient composition, amino and fatty acid profiles, in-vitro protein and starch digestibility, antioxidative and functional properties of sorrel seed. It is expected that results of this study will provide baseline information that will stimulate extensive research into optimization and mechanization of the dehulling process of sorrel seed. This will consequently take out the drudgery associated with the manual dehulling process and encourage production and utilization of the dehulled seed flour. Also, this study has proposed utilization possibilities of dehulled sorrel seed flour as a potential functional ingredient in food product development and protein supplement.

2. Materials and Methods

2.1. Sample Acquisition and Pretreatment

Red sorrel seeds (10 kg) were obtained from a local farm in Sokoto State, Nigeria and transported to the Federal University of Technology, Akure, Ondo State, Nigeria where this study was carried out from May to December, 2017. All reagents and chemicals used were of analytical grade.
2.2. Processing of Sorrel Seed and Production of Processed Flours

The seeds were sorted, washed severally with clean tap water to remove dirt and foreign materials, drained and divided into 5 equal portions of 2 kg each for preparation of flours.

2.2.1. Raw Oven-Dried Sorrel Seed Flour
Raw oven-dried sorrel seed flour which served as the control was prepared by oven drying the washed seeds at 60˚C until a constant weight was obtained [24].

2.2.2. Dehulled Sorrel Seed Flour
This was prepared by boiling the washed seeds in an aluminium pot at 100˚C until the seed coat was soft; drained, cooled and dehulled manually by rubbing between the palms to extract the kernels from the hulls. Clean tap water was thereafter poured in and the hulls separated from the kernels by repeated sieving.

2.2.3. Boiled Sorrel Seed Flour
Washed seeds were boiled with distilled water at 100˚C for 1 h in an aluminium pot and drained [12].

2.2.4. Roasted Sorrel Seed Flour
The method of Duwa et al. [25] was adopted for preparation of roasted sorrel seed flour. Washed seeds were roasted for 40 min at 100˚C in a frying pan placed on an electric cooker and allowed to cool to ambient temperature (25˚C ± 2˚C).

2.2.5. Germinated Sorrel Seed Flour
Washed seeds were steeped in 1 litre of distilled water at room temperature for 24 h; thereafter drained, spread on a clean moistened jute bag and allowed to germinate at room temperature (25˚C ± 2˚C) to a mean sprout height of 0.5 cm [12].

All processed seeds, apart from the roasted seeds, were oven dried at 60˚C until a constant weight was obtained, milled using a Waring Commercial Blender (Model 24CB10, USA), sieved using a mesh aperture of 0.4 mm, packaged in different airtight containers, labeled appropriately and stored at 4˚C for further analyses.

2.3. Determination of Chemical and Amino and Fatty Acid Compositions of Sorrel Seed Flours

Proximate composition of the seed flours was determined using the standard methods of Association of Official Analytical Chemists [26]. Crude protein content was determined by the micro Kjeldahl nitrogen method and a conversion factor of 6.25 was used to convert the nitrogen content to protein. Carbohydrate content was calculated by difference (100 – [moisture + total ash + crude fat + crude fiber + protein]).

Mineral element composition was determined using the methods described by
Isaac and Johnson [27] and Jones and Case [28]. Briefly, triplicate samples (1 g each) were ashed using a muffle furnace at 500°C in clean ceramic crucibles for the first 2 h and thereafter held for another 2 h to cool to room temperature. The ash were digested using distilled water, 400 ml conc. HCl and 133 ml 70% Nitric acid and diluted with distilled water to 2 litre mark in a volumetric flask to produce the Aqua Regia solution. The mixture was vortexed and thereafter centrifuged at 3000 rpm for 10 min. The clear supernatant was decanted into vials for mineral determination using the Unicam 919 atomic absorption spectrophotometer (Unicam Ltd, Cambridge, UK) for Ca, Mg, Mn, Zn and Fe. K and Na were determined using flame photometer (Corning EEL), and phosphorus by Phospho-vanadomolybdate method [26].

The amino acid profile of the seed flours was determined using the Ion Exchange chromatography (IEC). Briefly, triplicate samples were defatted using chloroform/methanol mixture of 2:1, hydrolyzed, evaporated in a rotary evaporator and injected into the Technicon sequential multisampling Amino Acid Analyzer (TSM). Fatty acid profile was determined by the method of Oh [29]. Fatty acids were extracted using a CHCl₃-MeOH (2:1 v/v) solution. The samples were centrifuged at 3000 rpm, supernatant collected were mixed with 0.9% NaCl solution and centrifuged again at 3000 rpm. The CHCl₃ phase was evaporated under nitrogen gas and treated with 14% boron trifluoride methanol solution (BF₃-MeOH) for 10 min at 100°C. After cooling to room temperature, 1 ml water and 2 ml pentane were added. The pentane phase was evaporated under nitrogen gas and dissolved in n-hexane. Fatty acid composition was analyzed using a gas chromatography (GC) (Acme 6000, Young-Lin Co.), which was equipped with a flame ionization detector (FID) and SPMTM-fused silica capillary column (130 mm × 0.25 mm, 0.25 µm, Supelco Co. USA). Nitrogen gas was used as carrier. Individual fatty acid methyl esters (FAME) were quantified as a percentage of total FAME analyzed. Free fatty acid was determined by Akintayo and Bayer [30] method.

Total carotenoid content of samples was determined by methods described by Rodríguez-Amaya and Kimura [31] with slight modifications. Briefly, 2.5 g of each sample was weighed into a conical flask. 30 ml hexane, 20 ml ethanol and 2 ml 2% NaCl solution were added and mixed thoroughly. The solution was transferred into a separating funnel and allowed to stand for 10 min to aid extraction of carotenoid. The lower content of the solution was run off and the upper layer was collected and absorbance was measured at 454 nm. Sample preparation and readings were done under less intense light in the laboratory because of the sensitivity of carotenoid to light. Total carotenoid content (mg/100g) was calculated and expressed on dry weight basis using the following equation:

\[
\text{Total carotenoid content (µg/g)} = \frac{A \times \text{Volume (mL)} \times 10^4}{A_{1cm}^\% \times \text{sample weight}}
\]

where \(A\) = absorbance; volume = total volume of extract (25 mL); \(A_{1cm}^\%\) = ab-
sorption coefficient of β-carotene in PE (2592), PE-Petroleum ether.

2.4. Determination of In-Vitro Starch and Protein Digestibility

In vitro protein digestibility (IVPD) of the seed flours was determined using the multienzyme procedure of Hsu et al. [32]. Enzymes used were porcine pancreatic trypsin (ZF.93615.0025), bovine pancreatic chymotrypsin (ZF.27270) and porcine intestinal peptidases (Z.F.77163.0500, Zefa Lab service, GMBH Germany). The activity of the enzymes was initially determined before use by using them to digest casein. One hundred milligrams (100 mg) of each of the seed flour was dispersed in 1 ml. Each sample suspension was adjusted to pH 8.0 and incubated in water bath at 3˚C with constant stirring. Fresh Multienzyme solution was prepared to contain 1.6 mg trypsin, 3.1 mg chymotrypsin and 1.4 mg peptidase dissolved in 1 ml distilled water. The pH of enzyme solution was maintained at 8.0. Five millimeter (5 ml) of the multienzyme solution was added to each sample suspension with constant stirring at 37˚C. The pH of each sample suspension was recorded at 10 min and 15 min respectively after adding the enzyme solution. IVPD was calculated using the equation of Hsu et al. [32]. In vitro starch digestibility was determined using pancreatic amylase [33]. Briefly, 50 mg of each sample was dispersed in 1 ml of 0.2 M phosphate buffer (pH 6.9). Twenty milligrams of the enzyme was dissolved in 50 ml of the same buffer and 0.2 ml of both the sample and enzyme were added. The mixture was heated for 5 min in a boiling water bath, cooled thereafter and the absorbance was read at 540 nm against a blank containing buffer while maltose was used as a standard.

2.5. Determination of Anti-Oxidant Potentials and Free Radical Scavenging Activity

2.5.1. DPPH Free Radical Scavenging Ability

DPPH free radical-scavenging ability of the samples was measured using spectrometric assay of Butrits and Bucar [34] which measures hydrogen atom or electrons-donating ability from the bleaching of purple-coloured methanolic DPPH solution. Aqueous extract of 100 µl, 200 µl, 300 µl and 400 µl containing 2 mg, 4 mg, 6 mg and 8 mg of the sample respectively was dispensed into test tubes and made up to 500 µl with distilled water followed by addition of 600 µl of methanolic DPPH and incubated in the dark at room temperature for 15 min and absorbance read at 517 nm using a JENWAY UV-Visible spectrophotometer (JENWAY Inc.). The DPPH radical-scavenging capacity (%) was calculated as:

$$DPPH(\%) = \frac{Abs_{reference} - (Abs_{sample} - Abs_{negative})}{Abs_{reference}} \times 100$$

2.5.2. Total Phenolic Content

Total phenolic content (TPC) of the samples was determined as described by Singleton et al. [35] with slight modification using gallic acid as standard. Fifty microliters (50 µl) of the aqueous extract containing 0.5 mg of aqueous extract
was dispensed into a test tube, 50 µl of distilled water and 500 µl of Folin-Ciocalteu reagent were added and shaken thoroughly. After 3 min at room temperature, 400 µl of 7.5% sodium carbonate solution was added and the mixture was incubated in the dark at 45°C in a water bath for 40 min and the absorbance read thereafter at 750 nm in a JENWAY UV-Visible spectrophotometer. Total phenolic content was expressed as gallic acid equivalent per gram of sample (mg of GAE/g sample) through the calibration curve of gallic acid and calculated as follows:

\[
\text{Total Phenolic Content (mg/GAE/g)} = \frac{\text{Abs}_{\text{sample}} \times \text{Conc. Standard (mg/ml)}}{\text{Abs}_{\text{standard}} \times \text{Conc. Sample (g/ml)}}
\]

### 2.5.3. Flavonoid Content

Flavonoid content was determined using the aluminum chloride colorimetric assay [36] with slight modifications. Briefly, 500 µl of aqueous extract of the samples was diluted with 500 µl methanol in a 10 ml flask. To this 500 µl, 10% AlCl₃, 50 µl of 1 M potassium acetate and water added to a total volume of 2.5 mL. The solution was incubated at room temperature for 40 min and absorbance read against blank at 415 nm. Total flavonoid content was calculated thus:

\[
\text{Total Flavonoid Content (mg/QE/G/g)} = \frac{\text{Abs}_{\text{sample}} \times \text{Conc. Standard (mg/ml)}}{\text{Abs}_{\text{standard}} \times \text{Conc. Sample (g/ml)}}
\]

### 2.5.4. Ferric Reducing Antioxidant Property (FRAP)

The ferric reducing antioxidant property of the flours was determined using the method of Pulido et al. [37]. 0.25 ml of sample extract (2.5 mL) was mixed with equal volumes of 0.2 M phosphate buffer (pH 6.6) and 1% potassium ferriycyanide [K₃Fe(CN)₆] and the resulting solution was incubated for 20 min at 50°C. Afterwards, 2.5 mL of freshly prepared 10% trichloroacetic acid was added and the mixture centrifuged at 600 rpm for 10 min. The supernatant (5 mL) was mixed with equal volume of distilled water and 1 mL of 0.1% FeCl₃ and the absorbance was immediately read at 700 nm using a UV-visible spectrophotometer. Ascorbic acid was used as standard and the ferric reducing power was determined as ascorbic acid equivalent per milliliter of the sample extract.

### 2.6. Determination of Some Functional Properties

Water and oil absorption capacities (WAC and OAC) were determined according to the methods of Rodriguez-Ambriz et al. [38] as described by Omowayne-Taiwo et al. [39]. Each sample (1 g) was weighed into a 15 ml already weighed centrifuge tube and 10 ml of distilled water (for WAC) or 10 ml of soybean oil (with density 0.92 g/ml) (for OAC) was added stepwise with continuous stirring at room temperature for 10 min. Thereafter, the tubes were centrifuged at 2500 xg for 20 min and volume of the supernatant measured. The WAC or OAC was calculated as the difference between the initial volume of water or oil used and the final volume of the decanted supernatant and calculated in percen-
tages (with consideration of the density of the oil). Swelling index was determined by Ukpabi and Ndimele [40] method. A sample size of 25 g was added to water in a measuring cylinder and left to swell for 4 h at room temperature. The procedure was replicated thrice and the swelling index was calculated as follows:

\[
\text{Swelling Index (v/v)} = \frac{\text{Final Volume}}{\text{Initial Volume}}
\]

Foaming capacity (FC) and foaming stability (FS) were carried out as described by Sze-Tao and Sathe [41]. The sample (0.5 g) was dispersed in 50 ml of distilled water in a 100 ml graduated cylinder and the solutions homogenized at a speed of 1600 ×g for 5 min. The volume was recorded before and after whipping. FC was expressed as the volume (%) increase due to whipping. This was then stored for 1 hr and the foam-volume changes in the graduated cylinder were recorded as FS. Both were calculated in percentages as shown below:

Foaming capacity

\[
\text{Foaming capacity} = \frac{\text{Volume after homogenization} - \text{Volume before homogenization}}{\text{Volume before homogenization}} \times 100
\]

Foaming stability

\[
\text{Foaming stability} = \frac{\text{Volume of foam after set time}}{\text{Initial volume of foam}} \times 100
\]

Emulsion capacity (EC) was determined according to the method of Chavan et al. [10]. One gram of the sample in 25 ml distilled water was homogenized at a speed of 5000 ×g for 1 min at 27˚C. The protein solution was then mixed with 25 ml of soybean oil followed by homogenization at 10,000 ×g for 1 min. The emulsion volume was then used in calculating the EC as shown below:

\[
\text{Emulsion capacity} = \frac{\text{Height of emulsified layer}}{\text{Height of the contents of the tube}} \times 100
\]

The effect of pH on protein solubility of the seed flours was determined by the method of Palić et al. [42].

2.7. Determination of Antinutrient Composition

Phytate was determined according to the method of Wheeler and Ferrel [43] and tannin as described by Makkar [44]. Oxalate was determined according to the method of Day and Underwood [45], while saponin content of the samples was determined using AOAC [46] methods.

2.8. Statistical Analysis

All determinations were carried out in triplicate on three independent batches of samples. Data obtained were subjected to analysis of variance (ANOVA) expressed as mean ± standard deviation using SPSS 16.0 for windows computer software package. The difference in means was compared using Duncan’s new Multiple Range test and significant level was established at P < 0.05.
3. Results and Discussion

3.1. Proximate and Mineral Element Compositions and in Vitro Digestibility of Raw and Processed Sorrel Seed Flours

Crude protein content of the seed flours showed significant (p < 0.05) increase due to processing from 24.93% to a range of 27.65% - 32.91% (% DW); accounting for a 10.91% - 32.01% increase in which dehulling caused the highest increase and roasting the least (Table 1). Values reported in the present study compare well with previous reports [15] [16] [47]; although Emmy Hainida et al. [47] reported that sun-drying and boiling caused crude protein reduction in sorrel seed. However, the increase here corroborates increase reported by Yagoub et al. [18] and Duwa et al. [25] in processed sorrel seed. This increase may be attributed to heat treatment which destroys heat-sensitive anti-nutritional factors such as protein inhibitors that bind to proteins and inactivate enzymes that speed up nutrient damage, hence improving the availability of nutrients [14]. Furthermore, these higher values as compared to those previously reported for some common seeds and grain legumes, such as sunflower seeds, melon seeds, cowpeas, soybeans and groundnuts [48], may make sorrel seed a potential source of cheap and available protein supplement. Dehulling may have resulted in the highest increase due to synergistic activities of seed coat removal and heat treatment applied to the seed before dehulling since hulls have been reported to

Table 1. Proximate composition (% dry weight), mineral composition (mg/100g), in vitro protein and starch digestibility of processed sorrel seed flours.

<table>
<thead>
<tr>
<th>Parameters</th>
<th>RDSF (Control)</th>
<th>DDSF</th>
<th>BDSF</th>
<th>ROSF</th>
<th>GDSF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture (g/100g)</td>
<td>6.40 ± 0.07</td>
<td>5.94 ± 0.07</td>
<td>6.72 ± 0.07</td>
<td>4.36 ± 1.00</td>
<td>5.67 ± 0.08</td>
</tr>
<tr>
<td>Crude protein (g/100g)</td>
<td>24.93 ± 1.00</td>
<td>32.91 ± 0.08</td>
<td>29.69 ± 0.08</td>
<td>27.89 ± 0.08</td>
<td>27.65 ± 0.08</td>
</tr>
<tr>
<td>Crude fat (g/100g)</td>
<td>26.24 ± 1.00</td>
<td>25.60 ± 1.00</td>
<td>28.66 ± 0.12</td>
<td>24.44 ± 0.12</td>
<td>22.24 ± 0.12</td>
</tr>
<tr>
<td>Crude fibre (g/100g)</td>
<td>2.47 ± 0.13</td>
<td>2.04 ± 0.07</td>
<td>4.69 ± 0.13</td>
<td>4.41 ± 0.13</td>
<td>4.09 ± 0.13</td>
</tr>
<tr>
<td>Total ash (g/100g)</td>
<td>4.36 ± 0.45</td>
<td>3.16 ± 0.05</td>
<td>3.38 ± 0.15</td>
<td>3.26 ± 0.01</td>
<td>3.41 ± 0.04</td>
</tr>
<tr>
<td>Carbohydrate (g/100g)</td>
<td>42.00 ± 0.06</td>
<td>36.29 ± 1.00</td>
<td>33.58 ± 0.06</td>
<td>40.01 ±1.00</td>
<td>42.61 ± 0.06</td>
</tr>
<tr>
<td>Sodium (mg/100g)</td>
<td>0.66 ± 0.01</td>
<td>1.11 ± 0.30</td>
<td>1.10 ± 0.30</td>
<td>0.76 ± 0.02</td>
<td>0.88 ± 1.00</td>
</tr>
<tr>
<td>Calcium (mg/100g)</td>
<td>20.86 ± 0.10</td>
<td>23.86 ± 0.01</td>
<td>25.75 ± 0.20</td>
<td>24.17 ± 0.10</td>
<td>23.02 ± 0.20</td>
</tr>
<tr>
<td>Potassium (mg/100g)</td>
<td>90.00 ± 0.20</td>
<td>91.86 ± 0.45</td>
<td>100.21 ± 0.09</td>
<td>111.41 ± 0.13</td>
<td>110.02 ± 0.20</td>
</tr>
<tr>
<td>Magnesium (mg/100g)</td>
<td>20.51 ± 0.02</td>
<td>21.61 ± 0.01</td>
<td>21.54 ± 0.10</td>
<td>20.49 ± 0.01</td>
<td>20.51 ± 0.01</td>
</tr>
<tr>
<td>Manganese (mg/100g)</td>
<td>0.02 ± 0.01</td>
<td>0.05 ± 0.01</td>
<td>0.11 ± 0.09</td>
<td>0.10 ± 0.09</td>
<td>0.11 ± 0.09</td>
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<tr>
<td>Zinc (mg/100g)</td>
<td>0.11 ± 0.98</td>
<td>0.12 ± 0.98</td>
<td>0.12 ± 0.98</td>
<td>0.14 ± 0.98</td>
<td>0.14 ± 0.98</td>
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<tr>
<td>Iron (mg/100g)</td>
<td>0.17 ± 0.23</td>
<td>0.19 ± 0.23</td>
<td>0.19 ± 0.23</td>
<td>0.12 ± 0.51</td>
<td>0.04 ± 0.13</td>
</tr>
<tr>
<td>IVPD (%)</td>
<td>78.32 ± 1.00</td>
<td>75.87 ± 1.00</td>
<td>72.36 ± 1.00</td>
<td>73.64 ± 1.00</td>
<td>74.40 ± 1.00</td>
</tr>
<tr>
<td>IVSD (%)</td>
<td>59.69 ± 1.00</td>
<td>61.68 ± 1.00</td>
<td>69.86 ± 1.00</td>
<td>67.18 ± 1.00</td>
<td>63.25 ± 1.00</td>
</tr>
</tbody>
</table>

Means ± standard deviation for at least 3 determinations; Means with different superscripts on the same row are significantly different at p < 0.05; Legends: RDSF (Control)—Raw oven-Dried Sorrel Seed Flour; DDSF—Dehulled Dried Sorrel Seed Flour; BDSF—Boiled Dried Sorrel Seed Flour; ROSF—Roasted Sorrel Seed Flour; GDSF—Germinated Dried Sorrel Seed Flour; IVPD—in vitro protein digestibility; ISPD—in vitro starch digestibility.
contain significant amounts of anti-nutritional factors [14] [20]. Dehulling may therefore serve as a better processing method for increasing protein content of sorrel seed and the dehulled seed flour may provide a higher source of nutritional protein supplement. Crude fat (ranging from 22.24% - 28.66%) reduced due to germination, dehulling and roasting, while boiling significantly (p < 0.05) increased it. Whereas dehulling decreased crude fibre possibly due to removal of hulls; germination, roasting and boiling increased it. Lower values in the present study (2.04% - 4.69%) as compared to those (13.10 - 15.50) previously reported by Duwa et al. [25] may be due to sieving of the flours after milling; although comparable values were reported by Kwari et al. [12]. Based on the high protein and moderate crude fibre contents of the dehulled sorrel seed flour (DDSF), this sample may be useful as a cheap, available protein supplement for enriching protein content of cereal-based complementary foods which serve as major weaning staples in Africa including Nigeria. The total ash and carbohydrate (except germinated sorrel seed flour—GDSF) contents of the processed flours were significantly lower than the raw sorrel seed flour (Table 1).

Results obtained showed that potassium (which ranged from 90.0 - 111.41 mg/kg) was the most abundant mineral in both raw and processed sorrel seed flours; the trend in abundance being K > Ca > Mg > Na > Fe > Zn > Mn (Table 1). The predominance of K corroborates previous reports that potassium is usually the most abundant mineral element in tropical plants and previous workers have similarly reported predominance of K, followed by Ca and Mg in sorrel seed [15] [16] [47]. However, processing caused significant variations in the mineral composition of sorrel seed. All processing methods increased K with the highest increase recorded in the roasted sorrel seed flour (ROSF) and lowest in DDSF, Ca [highest in boiled seed flour (BDSF) and lowest in GDSF], Na, Mg and Mn (increase not significant in DDSF); while Zn and Fe were not significantly affected. Although increased mineral contents due to processing may be unusual, Emmy Hainida et al. [47] also reported similar increase in Na, Ca and K due to boiling as compared to sun-drying and freeze-drying. This increase may be due to destruction of anti-nutritional factors (ANFs) which bind to mineral elements and reduce their contents and availability [14]. This is further corroborated by reports of Yagoub et al. [18] that processing improves HCl extractability of minerals; thus explaining the increase observed in this study. Overall, sample DDSF had the lowest mineral contents and may be linked to the lowest total ash recorded in DDSF. This may indicate that the seed’s hulls contain more of the inorganic component of the seed; and thus the dehulled seed flour may require micronutrient fortification when being used for protein enrichment if cereal-based complementary foods.

In vitro protein digestibility (IVPD) of sorrel seed flours ranged from 72.36% - 78.32%, comparing favourably with a range of 79% - 82% previously reported for sorrel seed and even higher than defatted soybean flour [15] [17] [18]. These high values indicate that the protein of these flours is highly digestible and as
such can serve as a nutritional source of amino acid, since digestibility is an important factor for determining the nutritional value of proteins [14]. However, there was a reduction of 3.13% - 7.61% due to processing, the highest recorded in the boiled seed flour (BDSF) (72.36%) and the lowest in the dehulled seed flour (DDSF) (75.87%) (Table 1). This indicates that the dehulled sorrel seed flour may serve as the best nutritional source of amino acid as compared to other flours, thus further emphasizing the potential of dehulling as the best processing method for production of sorrel seed flour that will serve as an important source of amino acid for human nutrition. Similar reductions have been previously reported and attributed to unfolding of the seed proteins which increases surface contact of embedded hydrophobic amino acids with water molecules, thus reducing solubility and consequently digestibility. Also, it has been reported that during heat treatment, changes occur in proteins resulting in reduced protein digestibility due to formation of isopeptides and high polymer protein fractions [18]. On the other hand, in vitro starch digestibility (IVSD) (which ranged from 59.69% - 69.86%) significantly increased due to processing; the highest increase recorded in BDSF and the lowest in DDSF. Heating improves the digestibility of starch through gelatinization and destruction of antinutrients and this may account for the increase in this study since all the processing methods except germination involved heating, whereas increased starch digestibility during germination of legumes may be attributed to activation of the amylolytic enzymes during germination [49].

3.2. Amino and Fatty Acid Profiles of Raw and Processed Sorrel Seed Flours

Table 2 shows the effect of processing on the amino acid composition of sorrel seed. Lysine, leucine, valine, arginine and phenylalanine generally had the highest amounts in the samples as compared to other essential amino acids and compared favourably with the reference; while for the non-essential amino acids, higher values were obtained for aspartic and glutamic acids. This is in line with previous reports that leucine, lysine, arginine, phenylalanine, valine and glutamic acid are abundant amino acids in sorrel seed [11] [16] [47]. On the other hand, methionine was the most limiting essential amino acid in the seed and previous workers had reported similarly [16] [17] [47]. Similar results have been reported in other oil-rich protein seeds [50] [51]. Significant reductions occurred due to other processing methods, however, dehulling significantly (p < 0.05) increased all essential and non-essential amino acids except isoleucine and valine where percentage reduction were 0.88% and 47.26%, respectively. Highest increase in the essential amino acids in sample DDSF occurred in leucine, methionine, lysine, cystine, threonine, tyrosine, histidine and phenylalanine with percentage increase of 61.17%; 52.27%, 12.35%, 27.75%, 16.09%, 13.7%, 20.65%, 11.11%, respectively. Similar reductions in amino acids due to heat treatment, germination and fermentation have been reported in sorrel seeds and other
Table 2. Essential and non-essential amino acid composition of raw and processed sorrel seed flours, whole egg protein and FAO/WHO recommended pattern of human requirement.

<table>
<thead>
<tr>
<th>Amino Acid (g/100g protein)</th>
<th>Samples</th>
<th>Reference</th>
<th>Human requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RDSF</td>
<td>DDSF</td>
<td>BDSF</td>
</tr>
<tr>
<td>Essential amino acids</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leucine</td>
<td>5.28 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>8.51 ± 0.02&lt;sup&gt;a&lt;/sup&gt; (&gt;61.17)</td>
<td>5.10 ± 0.05&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>3.39 ± 0.11&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.36 ± 0.01&lt;sup&gt;b&lt;/sup&gt; (&lt;0.88)</td>
<td>3.85 ± 0.04&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Lysine</td>
<td>5.91 ± 1.12&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.64 ± 0.01&lt;sup&gt;a&lt;/sup&gt; (&gt;12.35)</td>
<td>4.95 ± 0.11&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Threonine</td>
<td>2.64 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.06 ± 0.03&lt;sup&gt;a&lt;/sup&gt; (&gt;16.09)</td>
<td>2.58 ± 0.04&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Methionine</td>
<td>0.88 ± 0.02&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.34 ± 0.01&lt;sup&gt;a&lt;/sup&gt; (&gt;52.27)</td>
<td>0.80 ± 0.01&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Valine</td>
<td>4.19 ± 0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.21 ± 0.02&lt;sup&gt;b&lt;/sup&gt; (&lt;2.76)</td>
<td>4.01 ± 0.14&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>3.52 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>3.96 ± 0.02&lt;sup&gt;a&lt;/sup&gt; (&gt;11.11)</td>
<td>3.34 ± 0.01&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Histidine</td>
<td>1.84 ± 0.10&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.22 ± 0.01&lt;sup&gt;a&lt;/sup&gt; (&gt;20.65)</td>
<td>1.71 ± 0.02&lt;sup&gt;c&lt;/sup&gt;</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>2.65 ± 0.03&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.98 ± 0.01&lt;sup&gt;a&lt;/sup&gt; (&gt;13.70)</td>
<td>2.31 ± 0.03&lt;sup&gt;c&lt;/sup&gt;</td>
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<tr>
<td>Cystine</td>
<td>1.73 ± 0.03&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.21 ± 0.12&lt;sup&gt;a&lt;/sup&gt; (&gt;27.75)</td>
<td>1.66 ± 0.01&lt;sup&gt;c&lt;/sup&gt;</td>
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<tr>
<td>Arginine</td>
<td>4.68 ± 0.03</td>
<td>4.85 ± 0.04 (&lt;3.63)</td>
<td>4.68 ± 0.01</td>
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<tr>
<td>Total essential amino acids</td>
<td>36.71</td>
<td>41.34 (&lt;12.61)</td>
<td>34.99 (&lt;&lt;4.69)</td>
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</tbody>
</table>

Non-essential amino acids

<table>
<thead>
<tr>
<th>Amino Acid</th>
<th>Samples</th>
<th>Reference</th>
<th>Human requirement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine</td>
<td>4.80 ± 0.02</td>
<td>5.18 ± 0.32 (&lt;7.92)</td>
<td>4.47 ± 0.10</td>
</tr>
<tr>
<td>Serine</td>
<td>4.20 ± 0.01</td>
<td>3.06 ± 0.05 (&lt;27.14)</td>
<td>4.08 ± 0.13</td>
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<tr>
<td>Proline</td>
<td>4.29 ± 0.14</td>
<td>3.71 ± 0.32 (&lt;15.63)</td>
<td>3.94 ± 0.10</td>
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<tr>
<td>Aspartic acid</td>
<td>22.31 ± 0.51</td>
<td>23.26 ± 0.74 (&lt;4.26)</td>
<td>21.21 ± 0.09</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>14.85 ± 0.05</td>
<td>15.21 ± 1.01 (&lt;2.42)</td>
<td>14.28 ± 0.15</td>
</tr>
<tr>
<td>Glycine</td>
<td>3.63 ± 0.01</td>
<td>4.19 ± 0.12 (&lt;15.43)</td>
<td>3.40 ± 0.01</td>
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</table>
Total non-essential amino acids

<table>
<thead>
<tr>
<th>Value</th>
<th>Mean ± SEM</th>
<th>Percentage Increase/Decrease</th>
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<tbody>
<tr>
<td>54.08</td>
<td>54.61 (&gt;0.98)</td>
<td>51.38 (&lt;4.99)</td>
</tr>
<tr>
<td>47.37</td>
<td>&lt;12.41</td>
<td>42.95 (&lt;20.58)</td>
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<td>42.6</td>
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</table>

Ratio of essential to non-essential amino acids

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<th>Value</th>
<th>Mean ± SEM</th>
<th>Percentage Increase/Decrease</th>
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<tr>
<td>0.68</td>
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<tr>
<td>0.69</td>
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<td>0.72</td>
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<td>1.29</td>
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Arg/Lys

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<th>Percentage Increase/Decrease</th>
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<td>0.79</td>
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<tr>
<td>0.73</td>
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<td>0.95</td>
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<tr>
<td>0.94</td>
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<td></td>
</tr>
<tr>
<td>0.96</td>
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1st limiting amino acid

<table>
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<tr>
<th>Value</th>
<th>Mean ± SEM</th>
<th>Percentage Increase/Decrease</th>
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<td>Met</td>
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2nd limiting amino acid

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<th>Value</th>
<th>Mean ± SEM</th>
<th>Percentage Increase/Decrease</th>
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<td>Cys</td>
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<tr>
<td>Cys</td>
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</tr>
</tbody>
</table>

Values are mean ± SEM (n = 3). Values in parenthesis show percentage increase or decrease in the amino acid composition due to processing; aadopted from Iyenagbe et al. [53]. bAmino acid requirement/kg divided by safe level of reference protein/kg. cValues adopted from Emmy Hainida et al. [47]. dAmino acid composition of human milk. eIndividual aged 2 - 5 years. fIndividual aged 10 - 12 years.

Continued

However, increase due to dehulling is an indication that the kernel of sorrel seed contains the largest deposit of protein and amino acids in the seed, thus indicating the usefulness of dehulling as the best method for obtaining sorrel seed flour with enhanced protein quality. Thus, dehulled sorrel seed flour may serve as a better and more adequate protein supplement for human nutrition. Also, the highest lysine content of the dehulled seed flour which was higher than the FAO/WHO [53] human requirement may make this flour useful as a supplement food mixture for poor lysine food sources such as cereals used as weaning foods and a major staple in developing countries. Hence, it may contribute significantly to lysine content of complementary foods when combined with cereals (which are poor in lysine) and thus be useful for combating PEM in young children since values in this flour are higher than FAO/WHO [53] requirements for infants, pre-school and school age children.

Apart from the control sample which had 36.71% total essential amino acids, the dehulled seed flour (DDSF) was the only sample among the processed seed flours that met and exceeded (41.34%) the 36% recommended for an ideal protein [54]. Overall, roasting had the worst effect on the amino acids. The Arg/Lys ratios reported in this study (from 0.73 - 0.96) are lower than that of soybean (1.40) but higher than that for casein (0.44) [50]. Malomo et al. [55] reported that high ratio of Arg/Lys in the diet produces beneficial hypocholesterolemic effects that may improve cardiovascular health and help in regulation of hypertension. Hence, results obtained in the present study show potential of sorrel seed flours in impacting positive effects on the cardiovascular system.

Fatty acid composition of sorrel seed presented in Table 3 showed that sorrel seed is abundant in palmitic, linoleic, oleic and stearic acids as have been previously reported [15] [16], hence a rich source of these fatty acids. In most cases, processing had no significant (p < 0.05) effect on the fatty acids. Oleic and linoleic acids observed to be most abundant unsaturated fatty acids have also been
Table 3. Fatty acid composition and total carotenoid content of sorrel seed flours.

<table>
<thead>
<tr>
<th>Fatty Acids (%)</th>
<th>RDSF (Control)</th>
<th>DDSF</th>
<th>BDSF</th>
<th>ROSF</th>
<th>GDSF</th>
<th>Codex Alimentarius Commission (2009)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myristic acid (C14:0)</td>
<td>2.23 ± 0.11&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.26 ± 0.11&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.21 ± 0.11&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.51 ± 0.11&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.27 ± 0.11&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-</td>
</tr>
<tr>
<td>Palmitic acid (C16:0)</td>
<td>18.29 ± 0.06&lt;sup&gt;b&lt;/sup&gt;</td>
<td>18.36 ± 0.06&lt;sup&gt;b&lt;/sup&gt;</td>
<td>18.40 ± 0.06&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>18.66 ± 1.0&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>18.22 ± 0.1&lt;sup&gt;b&lt;/sup&gt;</td>
<td>8.0 - 13.5</td>
</tr>
<tr>
<td>Stearic acid (C18:0)</td>
<td>6.29 ± 0.72&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.22 ± 0.72&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.35 ± 0.71&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>6.27 ± 0.71&lt;sup&gt;b&lt;/sup&gt;</td>
<td>6.20 ± 0.71&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2.0 - 5.4</td>
</tr>
<tr>
<td>Oleic acid (C18:1)</td>
<td>22.28 ± 0.09&lt;sup&gt;b&lt;/sup&gt;</td>
<td>22.02 ± 1.00&lt;sup&gt;bc&lt;/sup&gt;</td>
<td>23.26 ± 1.0&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>23.17 ± 1.0&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>22.35 ± 0.0&lt;sup&gt;b&lt;/sup&gt;</td>
<td>17.0 - 30.0</td>
</tr>
<tr>
<td>Linoleic acid (C18:2)</td>
<td>50.09 ± 0.82&lt;sup&gt;b&lt;/sup&gt;</td>
<td>50.14 ± 1.00&lt;sup&gt;b&lt;/sup&gt;</td>
<td>50.17 ± 1.0&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>50.24 ± 1.0&lt;sup&gt;b&lt;/sup&gt;</td>
<td>49.86 ± 0.8&lt;sup&gt;b&lt;/sup&gt;</td>
<td>48.0 - 59.0</td>
</tr>
<tr>
<td>Linolenic acid (C18:3)</td>
<td>1.51 ± 0.16&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.59 ± 1.00&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.55 ± 0.15&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.50 ± 0.16&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.53 ± 0.16&lt;sup&gt;a&lt;/sup&gt;</td>
<td>4.5 - 11.0</td>
</tr>
<tr>
<td>US: S</td>
<td>3:1</td>
<td>3:1</td>
<td>3:1</td>
<td>3:1</td>
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</tr>
<tr>
<td>FFA</td>
<td>0.72 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.84 ± 0.02&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.56 ± 0.01&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.42 ± 0.01&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.61 ± 0.02&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
</tr>
<tr>
<td>Total Carotenoid (mg/100g)</td>
<td>0.04 ± 0.01&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.02 ± 0.01&lt;sup&gt;abc&lt;/sup&gt;</td>
<td>0.03 ± 0.01&lt;sup&gt;ab&lt;/sup&gt;</td>
<td>0.01 ± 0.01&lt;sup&gt;d&lt;/sup&gt;</td>
<td>0.03 ± 0.01&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
</tr>
</tbody>
</table>

Means ± standard deviation for at least 3 determinations. Means with different superscripts on the same row are significantly different at P < 0.05. Legends: RDSF (Control), Raw Dried Sorrel Seed Flour; DDSF, Dehulled Dried Sorrel Seed Flour; BDSF, Boiled Dried Sorrel Seed Flour; ROSF, Roasted Sorrel Seed Flour; GDSF, Germinated Dried Sorrel Seed Flour; US: S, Ratio of Unsaturated to Saturated Fatty Acid; FFA, Free Fatty Acid.

similarly reported and have been linked to its high level of antioxidants, particularly tocopherols [56]. This accounts for the high ratio of saturated to unsaturated fatty acids of 1:3 observed for all the samples. This is in agreement with the range of 1:2-1:3 reported by El-Adawy and Khalil [15]. Since intake of adequate amounts of unsaturated fatty acids has been linked to physiological benefits on blood pressure, heart rate, endothelial function, cardiac diastolic function and reduced risk of fatal coronary heart disease [14], sorrel seed may be termed a functional food which may impart these physiological benefits and contribute significantly to maintaining consumers’ health. All processing methods except dehulling reduced free fatty acid (FFA) from 0.72 to a range of 0.42 - 0.61. Low FFA (0.42 - 0.84) reported in this present study is an indication that the oil is less prone to enzymatic hydrolysis since free fatty acid is an important index for oil quality. Thus, foods manufactured using sorrel seed and/or its oil will have a long shelf life and be free from off-flavor development during storage. Although insignificant (except in ROSF), total carotenoid content reduced in all cases and this may be due to its sensitivity to oxidation and high temperature during heat processing since heat treatment destroys heat-sensitive nutrients including vitamins [14].

3.3. Anti-Oxidative Potentials and Free Radical Scavenging Activity of Raw and Processed Sorrel Seed Flour

Total phenolic content of sorrel flours ranged between 7.36 and 10.24 mg GAE/g; processing significantly (p < 0.05) increased the phenolic content. De-
hulling caused the highest increase from 7.36 mg GAE/g to 10.24 mg GAE/g and germination the lowest from 7.36 mg GAE/g to 7.70 mg GAE/g (Table 4). Oxidative stress ensues when there is an imbalance between the antioxidant system of the body and the formation of reactive oxygen species and phenolics act as antioxidants and have the ability to scavenge free radicals, which would otherwise build up in the body and cause harm [57]. Hence, increase in the phenolic content of sorrel seed due to processing and inclusion of these flours, especially DDSF into diets may provide protective roles against oxidative stress that can lead to the initiation phase of degenerative and cardiovascular diseases in human [58]. On the other hand, flavonoid content was significantly reduced by processing. Values ranging between 0.16 mg rutin Eqv/mL and 0.96 mg rutin Eqv/mL further shows that sample DDSF had the highest content as compared to other samples, hence the least reduction (0.57 mg rutin Eqv/mL) while no significant difference existed among samples BDSF (0.27 mg rutin Eqv/mL), ROSF (0.16 mg rutin Eqv/mL) and GDSF (0.33 mg rutin Eqv/mL).

DPPH (2,2-diphenyl-1-picrylhydrazyl) is one of the assays used to determine free radical-scavenging activities. DPPH possesses a proton-free radical with a characteristic absorption which decreases significantly on exposure to proton radical scavengers. During the assay, a change from the characteristic purple to yellow indicates that the radicals in the reaction medium are scavenged resulting in decreased absorbance and the degree of decolourization is an indication of the scavenging ability of the samples [14] [59]. The DPPH radical-scavenging activity of the flours ranged between 56.42% and 88.83% as shown in Table 4. These high values are an indication of the high radical-scavenging ability of sorrel seed. Processing significantly increased the DPPH of sorrel seed; RDSF (control sample) had the lowest value of 56.42%, and sample DDSF (dehulled sorrel seed flour) had the highest activity (88.83%). Thus, indicating that dehulling had the highest positive effect on the radical-scavenging ability of sorrel seed. DPPH of BDSF (67.43%) and ROSF (66.74%) were not significantly (p < 0.05) different, whereas GDSF had the second highest value (80.74%) (Table 4). Ferric reducing

### Table 4. Antioxidant properties of sorrel seed flours.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Total Phenolic content (mg/g)</th>
<th>Total Flavonoid (mg rutin Eqv/mL)</th>
<th>DPPH (%)</th>
<th>FRAP (μg Vit C E/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDSF (Control)</td>
<td>7.36 ± 0.15d</td>
<td>0.96 ± 1.00a</td>
<td>56.42 ± 0.05e</td>
<td>2.69 ± 1.00a</td>
</tr>
<tr>
<td>DDSF</td>
<td>10.24 ± 0.15a</td>
<td>0.57 ± 1.00b</td>
<td>88.83 ± 1.00a</td>
<td>2.37 ± 1.00b</td>
</tr>
<tr>
<td>BDSF</td>
<td>7.86 ± 0.15c</td>
<td>0.27 ± 1.00c</td>
<td>67.43 ± 0.05c</td>
<td>1.68 ± 0.63c</td>
</tr>
<tr>
<td>ROSF</td>
<td>9.21 ± 0.15b</td>
<td>0.16 ± 1.00c</td>
<td>66.74 ± 1.00c</td>
<td>2.15 ± 1.00c</td>
</tr>
<tr>
<td>GDSF</td>
<td>7.70 ± 0.15c</td>
<td>0.33 ± 1.00c</td>
<td>80.74 ± 1.00b</td>
<td>1.66 ± 0.63c</td>
</tr>
</tbody>
</table>

Means ± standard deviation for at least 3 determinations. Means with different superscripts on the same column are significantly different at p < 0.05. Legends: RDSF (Control), Raw Dried Sorrel Seed Flour; DDSF, Dehulled Dried Sorrel Seed Flour; BDSF, Boiled Dried Sorrel Seed Flour; ROSF, Roasted Sorrel Seed Flour; GDSF, Germinated Dried Sorrel Seed Flour.
antioxidant power (FRAP) assay measures antioxidant power by the reduction of ferric iron (Fe³⁺) to ferrous iron (Fe²⁺). FRAP values ranged from 1.66 - 2.69 mg Vitamin C Eqv/mL (Table 4). The control sample (RDSF) had the strongest reducing ability but was decreased by processing. Boiling (BDSF) and germination (GDSF) had the strongest negative effect on FRAP, while dehulling had the least negative effect with the highest value of 2.37 mg Vitamin C Eqv/mL. This may indicate that dehulling will ensure the production of sorrel seed flour with higher antioxidative and scavenging activity than other processing methods.

3.4. Functional Properties of Raw and Processed Sorrel Seed Flour

Functional properties are non-nutritional, intrinsic characteristics of a food or food additive which affect the behaviour of proteins in food systems during processing, manufacturing, storage, preparation and consumption and are usually affected by processing treatments and the environment. These properties are important in determining the level of utilization of any flour in ingredient formulation and food product development and reflect the composition and conformation of proteins, and their interactions with other food components [23] [60]. During heat treatment, denaturation of protein occurs thereby influencing its functional properties and bringing about significant modification of physicochemical characteristics including dissociation into constituent subunits, unfolding and surface exposure of hydrophobic side groups [14]. This may explain changes observed in this present study.

3.5. Water and Oil Absorption Capacities and Swelling Power

Results in Table 5 showed that all processing methods significantly (p < 0.05) increased water absorption of sorrel seed flour from 115% to a range of 165% - 266%; with highest increase in roasted seed flour (ROSF) and the lowest in DDSF. Iyenagbe et al. [50] reported similar higher increase of WAC in toasted sorrel seed flours.

Table 5. Functional properties of sorrel seed flours.

<table>
<thead>
<tr>
<th>Samples</th>
<th>WAC (%)</th>
<th>OAC (%)</th>
<th>Swelling power (%)</th>
<th>Foaming Capacity (%)</th>
<th>Foaming Stability (%)</th>
<th>Emulsifying Capacity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RDSF (Control)</td>
<td>115 ± 1.00d</td>
<td>156 ± 1.00d</td>
<td>74.0 ± 1.00b</td>
<td>16.66 ± 1.00a</td>
<td>50.00 ± 1.00a</td>
<td>66.67 ± 1.00a</td>
</tr>
<tr>
<td>DDSF</td>
<td>165 ± 1.00c</td>
<td>184 ± 1.00b</td>
<td>93.3 ± 1.00a</td>
<td>8.33 ± 1.00b</td>
<td>20.02 ± 0.57d</td>
<td>53.27 ± 0.91b</td>
</tr>
<tr>
<td>BDSF</td>
<td>264 ± 1.00a</td>
<td>166 ± 0.13c</td>
<td>49.1 ± 1.00d</td>
<td>4.84 ± 1.00c</td>
<td>33.24 ± 1.00b</td>
<td>53.77 ± 1.00d</td>
</tr>
<tr>
<td>ROSF</td>
<td>266 ± 0.66a</td>
<td>217 ± 1.00a</td>
<td>66.0 ± 1.00b</td>
<td>8.35 ± 1.00b</td>
<td>20.75 ± 0.55c</td>
<td>53.31 ± 0.91b</td>
</tr>
<tr>
<td>GDSF</td>
<td>206 ± 1.00b</td>
<td>173 ± 0.13c</td>
<td>19.5 ± 1.00a</td>
<td>3.35 ± 1.00a</td>
<td>20.18 ± 0.57d</td>
<td>50.15 ± 1.00c</td>
</tr>
</tbody>
</table>

Means ± standard deviation for at least 3 determinations. Means with different superscripts on the same column are significantly different at p ≤ 0.05. Legends: RDSF (Control), Raw Dried Sorrel Seed Flour; DDSF, Dried Dried Sorrel Seed Flour; BDSF, Boiled Dried Sorrel Seed Flour; ROSF, Roasted Sorrel Seed Flour; GDSF, Germinated Dried Sorrel Seed Flour; WAC, Water Absorption Capacity; OAC, Oil Absorption Capacity.
conophor seed flour. This increase may be due to protein denaturation which occurs during heat processing, resulting in increased WAC [50]. Also, increased protein content reported earlier in this study (Subsection 3.1) may account for increased WAC since proteins have the ability to absorb and retain water [61]. Values reported in this study (115% - 266%) are consistent with those previously reported for most defatted oilseed flours (100% - 260%), lupin seed (120%), soy (130%) and pigeon pea (138%) flours [62]. Hence, both raw and processed sorrel seed flours may serve as functional ingredients in liquid and semi liquid foods like soups, gravies and baked products where water absorption/thickening is a critical factor; and may serve as good replacement for some legumes and oil seeds currently being used as thickeners since they will be relatively cheaper. Similarly, oil absorption increased (from 156% to 166% - 217%) due to processing and again roasting caused the highest increase. Values reported in this study (156% - 217%) compare favourably with those for some oil seed flours (140%, 142%, 193% and 142% for chickpea, yam bean, soy and fluted pumpkin seed flours respectively) [62] [63] [64], but are higher than those reported for pigeon pea (89.7%), jack bean (105.6%) and gourd seed (96%) flours [62] [65]. This further suggests their possible applicability in baked foods because good oil absorption capacity of flour samples is an indication of their usefulness in food preparations that involve oil mixing like in bakery products, where oil is an important ingredient [65]. Their use as composite in baked foods will not only improve protein quality of these foods whose major ingredient is wheat flour, but will also reduce amount of wheat in the product and the associated problems of gluten intolerance. The water/fat binding capacity of proteins which is an index of its ability to absorb and retain oil, influences flavour retention, texture and mouth feel of food products like ground meat formulations, doughnuts, pancakes, baked goods and soups [63]. Hence, sorrel seed flours can be used as flavour retainer and to improve the mouth feels of food.

On the other hand, significant reduction occurred in swelling power due to boiling (49.1%), roasting (66.0%) and germination (19.5); while only dehulling increased it (Table 5). The highest reduction caused by germination which may be due to activation of amylolytic enzymes during germination which degrade starch for use as energy source since swelling power of flours is associated with granule structure and chemical composition, particularly amylose and lipid content [66]. Also, heat processing has been reported to reduce swelling power of starch. Increase in swelling power due to dehulling implies that the flour may find wide application in food systems where retention of water is desirable especially in baked foods.

3.6. Foaming and Emulsification Capacities

Foaming capacity and stability were significantly reduced due to processing. Low foaming capacity and stability of both raw and processed sorrel seed flours (ranging from 3.35% - 8.35% and 20.02% - 33.24% respectively) indicate that the
flours may find little or no application in foods such as ice-cream and alcoholic beverages where foaming is desirable, hence may not be useful as aerating agents in foods such as ice cream. This is because foams are very vital in foods like whipped toppings and beverages where the proteins unfold forming a layer that keeps air bubbles in suspension and prevents them from collapsing [50]. However, the low FS of the flours may indicate that very slow film formation at the air-water interfaces coupled with poor film visco-elasticity [55].

Similarly, emulsion capacity was significantly reduced by all processing methods from 66.67% in sample RDSF to 33.77%, 50.15%, 53.27% and 53.31% in samples BDSF, GDSF, DDSF and ROSF, respectively. While boiling caused the highest reduction, dehulling and roasting [which were not significantly (p < 0.05) different] resulted in the lowest decrease. Similar reduction has been reported in blanched fluted pumpkin seed flour and has been attributed to heat effect [64]. Values reported in this present study are higher than those previously reported for soya bean (15.0%), breadnut (18%), wheat (7% - 11%), calabash seed (23.2%) and pigeon pea (49.1%) flours [62] [65] [67] but are lower than 75.1% for sunflower flour [68]. Despite the reductions, sorrel seed flours can still find use as emulsifiers since relatively high values (ranging from 33.77% - 66.67%) were obtained. Emulsion capacity indicates the maximum amount of oil that can be emulsified by protein dispersion, hence, based on McWatters and Cherry’s [68] model for describing thickness or emulsion consistency of flours; samples DDSF, ROSF and GDSF would have a thick mayonnaise-like emulsion (50 - 59 ml/g), while the control flour (RDSF) with a higher value (66.67%) would have a very thick mayonnaise-like (>60 ml/g) emulsion. However, sample BDSF having the lowest value (33.77%) would have a very thick salad dressing-like emulsion (30 - 39 ml/g). Hence, the flours can find use as emulsifying agents and meat additive in sausage production, salad dressing preparation, pie fillings, ice creams and mayonnaise.

3.7. Protein Solubility

There was an inconsistent reduction in the protein solubility of sorrel seed flour; at pH 3, solubility increased from 4.78 in the control (RDSF) to 7.14 in BDSF; and at pH 6 from 6.1 in RDSF to 6.22 and 6.82 in DDSF and BDSF respectively (Figure 1). Maximum protein solubility of 10.01 and 9.63 were obtained at pH 9 for RDSF and GDSF, while minimum protein solubility of 2.64 was obtained at pH 8 in DDSF. Dehulled and roasted seed flours had maximum values of 6.22 and 5.43 at pH 6, while BDSF had maximum solubility of 7.14 at pH 3. Minimum protein solubility for BDSF (4.27) and GDSF (3.52) was reached at pH 2, while the control had minimum value of 4.78 at pH 3. Decreased protein solubility which has been linked to reduced in vitro protein digestibility in this study (Subsection 3.1) may be attributed to denaturation of proteins and increase in amounts of insoluble protein aggregates; carbohydrates and polyphenols during heating [34]. Thus, the control (RDSF) and sprouted (GDSF) seed flours with
high protein solubility at pH 9 may be useful in vegetable milk production where emulsification is important [62].

3.8. Effect of Processing on the Antinutrient Contents of Sorrel Seed Flour

Figure 2 which presents the effects of different processing methods on the antinutrient content of sorrel seed showed that of the antinutrients determined, saponin was most abundant in sorrel seed, with values ranging from 6.89 - 30.26 mg/g. It was followed by phytate (3.93 - 9.12 mg/g), tannin (2.47 - 6.04 mg/g) and oxalate (1.30 - 3.23 mg/g). Processing significantly (p < 0.05) reduced saponin; roasting caused the highest reduction from 30.26 mg/g to 6.89 mg/g. The trend in decreasing order was DDSF > GDSF > BDSF > ROSF. A similar trend was observed for oxalate and tannin contents, while for phytate the trend was slightly different DDSF > GDSF > ROSF > BDSF. These results indicate that heating had the most significant effect on the antinutrients as compared to germination and dehulling. Hence, irrespective of the processing method used, sorrel seed meant for human consumption has to be subjected to some form of heating to reduce the antinutrient content and enhance safety from toxigenic components.

4. Conclusion

Results of the present study have shown that dehulling has the potential of
improving protein quality and food functionality of the under-utilized sorrel seed and make it an important source of nutritional amino acid for human nutrition and a functional ingredient in food product development. Dehulling enhanced the nutritional, antioxidative and functional properties of sorrel seed as compared to other methods that have been previously used. Hence, the dehulled seed flour may provide a higher source of nutritional protein supplement and a better source of essential amino acid. High in vitro digestibility values especially in the dehulled seed flour indicate that the protein of these flours is highly digestible and as such can serve as a nutritional source of amino acid. Also, the dehulled seed flour (DDSF) exhibited the highest antioxidative and radical-scavenging activities as compared to other processed seed flours thus indicating that greater amounts of the bioactive compounds of sorrel seed are present in the kernel than in the hulls/seed coat. Thus, removal of the hull aided in concentrating the antioxidants in the kernels. Good thickening and emulsifying properties of the flours indicate they can find use as functional food ingredients for food product development. However, the lowest amounts of ash and minerals in the dehulled seed flour may necessitate micronutrient fortification of this flour when being used as a protein-enhancer in cereal-based complementary foods. Further in vivo studies are required to ascertain true biological values and health benefits of dehulled sorrel seed flour. Further studies are also required to verify the food product applicability of the flour and to optimize the traditional dehu-
ling process for mechanical dehulling so as to remove the drudgery associated with the manual dehulling process. This will encourage increased production, widespread acceptance and consumption of this flour.

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Conflicts of Interest

The authors declare no conflict of interest (financial or non-financial).

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