Production, Proximate Compositions and Dry Matters of Stored Achicha and Mpoto - Cocoyam Based Products

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ABSTRACT

The production, proximate compositions and dry matters of stored Achicha and Mpoto were conducted. Fresh cocoyam corms/cormels and leaves [ede ofe (NCE 002), coconoida (NCE 001) and ukpong/anampu (NCE 004)] were obtained from National Root Crop Research Institute, Umudike, Abia State, Nigeria. A 5 kg cocoyam corms/cormels of the samples was sorted, washed and boiled for 3 hours and was cooled, peeled and cut into small sizes of average of 2.0 cm by 1.5 cm dimension with a sharp kitchen knife. They were spread on a mat and dried under the sun for 5 days. The dried cocoyam corms/cormels (achicha) were pulverized before storage with a locally fabricated machine and stored in various plastic containers for 0, 1, 2, 3 months intervals. A sample of 300 g of cocoyam leaves were plucked, sorted, washed, spread on a mat and sun-dried for 3 days. The dried cocoyam leaves (mpoto) were pulverized before storage with a locally fabricated machine and stored in various plastic containers for 0, 1, 2, 3 months. The proximate compositions and dry matters of 3 different varieties of 4 samples of stored Achicha and Mpoto were determined. The results of stored Achicha showed that Edeofe had the least contents in moisture (12.33%) and Anampu had least contents in crude fibre (1.64%) and carbohydrate content (75.65%); Edeofe had the highest contents in ash (3.83%), crude protein (4.78%), crude fat (0.93%), and Coccoindia had the highest contents of dry matter (87.79% after 3 months storage. The proximate compositions of stored Mpoto showed that Edeofe had the least moisture content (10.16%), Anampu had the highest contents in ash (14.92%), Edeofe had the highest contents in crude protein (15.19%), crude fat (0.89%), crude fibre (8.74%), dry matters (89.85%); and Coccoindian had highest carbohydrate content (51.08%) after 3 months storage. This showed that stored Mpoto samples were richer in ash, crude protein, crude fibre and dry matters than the stored Achicha samples which were richer in moisture, crude fat and carbohydrate contents. It is recommendable to use the tuber and the leaves in food preparations for human consumption.

Keywords: Corm, Cormel, Protein, Fat, Fibre, Ash, Carbohydrate, Moisture, Edeofe, Coccoindia,

Introduction

Cocoyam is commonly referred to as Ede in Igbo land of Nigeria. Cocoyam’s Taro and Tannia have remained the two varieties mainly grown in Nigeria. The taro varied botanically known as Colosasia esculenta and commonly called Coco-India originated from Asia, while tannia (Xanthosoma sagittifolium) originated from America but were both introduced and grown in West Africa. These two species Colosasia esculenta (Taro) and Xanthosoma sagittifolium (Tannia) are the most widely accepted and cultivated varieties in Nigeria and other parts of the tropics and sub-tropic of Africa. Colosasia is thought to have originated into Indo–Malayan region, perhaps in Eastern India and Bangladesh, and spread eastward into South East Asia, Eastern Asia and the Pacific Island; Westward to Egypt and the Eastern Mediterranean, and then Southern and Westward from there into East Africa and West Asia, whence it spread to the Caribbean and Americans. It is known by many local names and often referred to as Elephant–ears when grown as an ornamental plant [1-5].

Taro plant is a perennial herb with clusters of long heart or arrow head-shaped leaves that point earthward. It is cultivated in the tropics, and the leaves are classified as large to very large, 20 to 150 cm (7.6 to 5.9 in) long, with a sagittate shape, growing on erect stems which may be green, red, black, or variegated. The new leaves and stems push out of the innermost stalk, unrolling as they emerge, with the stem several feet high [6]. Taro corms contains considerable amount of starch (70 to 80 g/100g dry Taro). Lebot (2009) did report taro corms to be rich in starch (61 to 88%DM) but little of protein (2.3 to 14.8% DM) [7]. The corm contains mainly starch and water together with small quantities of protein, fat, ash, vitamins B and C. The carbohydrate content of taro cultivated in different locations varied. The starch extracted from taro corms appears as fine granules in the 0.5 to 5 microns range, and thus offers smooth textured starch gel [1]. Moreover, the fine starch granule was reported to improve the binding and reduced breakage of snack products [7]. Meanwhile, taro leaves have been said to have a variable but generally high protein
content usually in the 16 to 27% DM range [5]. Even though lower values (13 to 16% DM) are also reported. Moreover, the leaves are a good source of thiamin, riboflavin, iron, phosphorus and zinc and a very good source of vitamin C, B6, niacin, potassium, copper and magnesium [8]. Cocoyam is most commonly grown for its starchy edible roots [6]. Containing several vitamins and minerals. Cocoyam also has appreciable content of crude fibre which aids in digestion and makes the elimination of stool very easy, as well as playing major role in preventing cancer [4, 2].

The cocoyam (Colocasia esculenta) is highly perishable root and leaves, as high as 40-60% post-harvest losses have been found [9]. The high perishability of the harvested and stored cocoyam roots and leaves is a major barrier to the wider utilization of the crop and there is need to diversify the uses to enhance demand and increase the rate of turn over or sale of the product [3]. Lack of adequate cocoyam processing technology inhibits production and processing. Over the years due to the high perishable nature of cocoyam local farmers had adopted sun-drying as a means of preserving the cocoyam. It becomes necessary to evaluate the effect of these processes and storage methods on the overall quality of the cocoyam products.

Cocoyam is an indigenous root crop that has not been utilized like other root crops such as cassava and yam. Despite the high nutritional value of cocoyam and soybean in relation to other root crops and legumes, lack of knowledge of their uses has limited their adoption, production and processing [10]. To bridge the gap, efforts are being made by research institutes, Non-Governmental Organizations (NGOs) and industries to promote the production, processing and utilization of cocoyam in Nigeria [2]. As there is a growing interest in the production of flours from locally available grains that can be used as substitutes for wheat in baked goods, this study was undertaken to produce sausage rolls of acceptable quality from cocoyam, soybean and wheat flour blends. It is mostly used as thickener in soup and few indigenous recipes. Therefore, its conversion into flour could be used efficiently in baking technology. Nigeria faces one of the most serious nutritional problems in protein-energy malnutrition. Nigeria has not been able to produce wheat in commercial quantity due to climatic and soil conditions [11, 3]. The main objectives of this study are to produce achicha (dried cocoyam corms/cormels) and mpoto (dried cocoyam leaves) and to evaluate the effect of storage periods (0-3 months intervals) on the proximate and dry matter compositions of cocoyam-based products.

Materials and Methods
Collection of Materials
Fresh cocoyam corms/cormels and leaves [ede ofe (NCE 002), cocoindia (NCE 001) and ukpong/anampu (NCE 004)] were obtained from National Root Crop Research Institute, Umudike, Abia State, Nigeria. The fresh samples were identified by the Agronomist, Cocoyam Unit, National Research Institute Umudike, Abia State. The cocoyam corms/cormels and cocoyam leaves are shown in Plates 1 and 2.

Processing of Corms/cormels into Achicha (Dried Cocoyam)
The cocoyam corms/cormels weighing 5kg for each of the samples was sorted, washed and boiled for 3hrs. It was cooled, peeled and cut into small sizes of average of 2.0 cm by 1.5 cm dimension with a sharp kitchen knife. They were spread on a mat and dried under the sun for 5 days between 9am- 6pm. The dried cocoyam corms/cormels (achicha) were pulverized before storage with a locally fabricated machine and stored in various plastic containers for a period of three months and analyzed at 0, 1, 2- and 3-months intervals. The pictures of the cocoyam plant, cocoyam corms/cormels and cocoyam leaves and the processed achicha are shown in Plates 1, 2, 3 and 4, respectively. Also, the flow diagrams of the production of achicha from cocoyam corms/cormels is shown in Figure 1.


**Plate 3: Raw achicha**

**Plate 4: Colocasia esculenta leaves**

**Proximate Analysis of Achicha and Mpoto Samples**

Proximate analysis of achicha and mpoto (dried cocoyam corms/ cornels and leaves) samples were evaluated for moisture, ash, crude protein, crude fibre, fat, and carbohydrate contents by the methods of [12, 13]. And were determined in triplicates.

**Determination of moisture content**

The moisture content was determined by the gravimetric methods as described by [14]. A measured weight of each sample (5 g) was put into a weighed moisture crucible. The weight of the crucible and the sample were taken. The crucible and its sample content were dried in the oven at 105 °C for 3 hours at first. They were cooled in desiccators and reweighed. The weights were recorded while the samples were returned to the oven for further drying. The drying, cooling and weighing continued repeatedly until a constant weight was obtained. The weight of the moisture lost was determined by differences and expressed as a percentage. It was calculated as shown in Equation 1:

\[
\text{% Moisture} = \frac{W_2 - W_4}{W_3 - W_4} \times 100 \quad \text{Equation 1}
\]

Where:
- \(W_1\) = Initial weight of empty crucible
- \(W_2\) = Weight of crucible and sample before drying
- \(W_3\) = Final weight of crucible and sample after drying at a constant weight.

**Determination of total ash content**

This was done using the furnace incineration gravimetric methods [12]. A measured weight (5 g) of each sample was put into a previously weighed porcelain crucible. The samples in crucible were put in a muffle furnace set at 500 °C and allowed to burn for 3 hours (until the sample became grey ash). The sample in crucible was carefully removed from the furnace (taking care not to allow air blow the ash away) and cooled in a desiccator. It was reweighed; weight of the ash was obtained by difference and expressed as a percentage given by the Equation 2:

\[
\text{% Ash (Dry basis)} = \frac{W_3 - W_1}{W_3 - W_2} \times 100 \quad \text{Equation 2}
\]

Where:
- \(W_1\) = Weight of empty crucible
- \(W_2\) = Weight of crucible + food before drying and/or ashing
- \(W_3\) = Weight of crucible + ash

**Determination of crude protein**

The crude protein was determined by Kjeldahl method described by [14]. The total nitrogen was determined and multiplied with the factor 6.25 to obtain the protein. For each sample, 5 g was mixed with 10 ml of concentrated sulphuric acid, AR grade (Analytical Reagent Grade) in a Kjeldahl digestion flask. A tablet of selenium catalyst was added to it and the mixture digested (heated) under a fume cupboard until a clear solution was obtained in a separate flask. The acid and other reagent were digested but without sample to form the blank (control).

All the digests were carefully transferred to 100 ml volumetric flask using distilled water and made up to a mark in the flask. A 100 ml portion of each digest was mixed with equal volume of 45 % NaOH solution in Kjeldahl distilling unit. The mixtures were distilled and the distillate collected into 10 ml of 4 % boric acid solution containing 3 drops of mixed indicator (bromocresol green and methyl red). A total of 50 ml distillate was obtained and titrated against 0.02 M H2SO4 solution. Titration was done from the initial green colour to a deep red end point. The Nitrogen content was calculated as shown in Equation 3:

\[
\% N_a = \frac{100}{W} \times \frac{N \times 14}{100} \times \frac{V_f}{V_a} \times T \quad \text{Equation 3}
\]

Where:
- \(W\) = Weight of sample analyzed
- \(N\) = Concentration of H2SO4 hydrant
- \(V_f\) = Total volume of digest
- \(V_a\) = Volume of digest distilled
- \(T\) = Titre value of blank.

**Determination of Crude Fibre**

This was determined by Wende methods [13]. A 5 g of each sample was defatted (during fat analysis). The defatted sample was boiled under reflux for 30 mins with 200 ml of a solution containing 1.25 g of H2SO4 per 100ml solution. After that the samples were washed with several portions of not boiling water using a two-folds muslin cloth to trap the particle, until the washings were no longer acidic. The washed samples were carefully transferred quantitatively back to the flask and 200ml of 1.25 g of NaOH solution was added to it. Again, the samples were boiled for 30 min. and washed as before with hot water. They were then carefully transferred to a weighed porcelain crucible and dried in the oven at 105 °C for 3 hours. After cooling in a desiccator, they were reweighed (W2), then put in a muffle furnace and burnt at 550 °C for 2 hours (until they became ash). Again, they were cooled in desiccators and reweighed. The crude fibre content was calculated gravimetrically as shown in Equation 4:

\[
\frac{W_2 - W_3}{W_1} \times 100 \quad \text{Equation 4}
\]

Where:
- \(W_1\) = Weight of sample.
- \(W_2\) = Weight of crucible and sample before incineration.
- \(W_3\) = Weight of crucible and sample ash.
Determination of fat content
Fat content of the sample was determined by the continuous solvent extraction methods using a Soxhlet apparatus. The methods as described by [12]. A measured weight of 5g of each sample was wrapped in a porous paper (Whatman No. 1 filter paper). The wrapped samples were put in a Soxhlet reflux flask containing 200ml of petroleum ether. The upper end of the reflux flask was connected to a condenser by heating the solvent in the flask through electro-thermal heater, it vaporized and condensed into the reflux flask. Soon, the wrapped samples were completely immersed in the solvent and remained in contact with the solvent until the flask is filled up and siphoned over them carrying oil extract from the samples down to the boiling flask. The process was allowed on repeatedly for 4 hours before the defatted samples were removed and reserved for crude fibre analysis. The solvent was recovered and the extracting flask with its oil content was dried in the oven at 60 °C for 3 minutes (to remove any residual solvent). After cooling in a desiccator, the flask was reweighed. By difference, the weight of fat (oil) extraction was determined and expressed as a percentage of the sample weights as shown in Equation 5:

\[ \% \text{ Fat} = \frac{\text{Weight of fat}}{\text{Weight of sample}} \times 100 \quad \text{Equation 5} \]

Determination of Carbohydrate
The carbohydrate content was calculated by difference as the nitrogen free extractive (NFE): a method separately described by [13]. as shown in Equation 6:

\[ \% \text{ NFE} = 100 - \% (a + b + c + d + e) \quad \text{Equation 6} \]

Where:
- \( a \) = Protein
- \( b \) = fat
- \( c \) = fibre
- \( d \) = ash
- \( e \) = moisture.

Determination of Dry Matter
The moisture content was determined by the gravimetric methods as described by Onwuka (2005; 2018). A measured weight of each sample (5 g) was put into a weighed moisture crucible. The weight of the crucible and the sample were taken. The crucible and its sample content were dried in the oven at 105 °C for 3 hours at first. They were cooled in desiccators and reweighed. The weights were recorded while the samples were returned to the oven for further drying. The drying, cooling and weighing continued repeatedly until a constant weight was obtained. The weight of the moisture lost was determined by differences and expressed as a percentage. It was calculated as shown in Equation 7:

\[ \% \text{ Moisture} = \frac{W_1 - W_3}{W_2 - W_1} \times 100 \quad \text{Equation 7} \]

Where:
- \( W_1 \) = Initial Weight of empty crucible
- \( W_2 \) = Weight of Crucible and sample before drying
- \( W_3 \) = Final weight of crucible and sample after drying at a constant weight.

% total solid (dry matter) = 100 - % moisture.

Statistical Analysis
All analysis was carried out in triplicates. The experiment was laid out in a completely randomized design (CRD). The data obtained were analyzed statistically using analysis of variance (ANOVA) at 5% level of significance while Least Significant Difference was used to separate the factor means [15].

Results and Discussion
Proximate and Dry Matter Compositions of Achicha during Three Months Storage
Table 1 showed the comparison of mean Proximate Composition of achicha processed from three different Colocasia varieties (edoefe, cocoindia, and anampu) during three months storage.

<table>
<thead>
<tr>
<th>Proximate</th>
<th>Edoefe (M)</th>
<th>Cocoindia (M)</th>
<th>Anampu (M)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Moisture</td>
<td>10.27±</td>
<td>10.75±</td>
<td>10.84±</td>
</tr>
<tr>
<td>±0.03</td>
<td>0.03</td>
<td>±0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Ash</td>
<td>4.31±</td>
<td>4.17±</td>
<td>3.94±</td>
</tr>
<tr>
<td>±0.03</td>
<td>0.03</td>
<td>±0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Crude Protein</td>
<td>5.77±</td>
<td>5.44±</td>
<td>4.81±</td>
</tr>
<tr>
<td>±0.03</td>
<td>0.03</td>
<td>±0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Crude Fat</td>
<td>1.05±</td>
<td>1.03±</td>
<td>0.95±</td>
</tr>
<tr>
<td>±0.03</td>
<td>0.03</td>
<td>±0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Crude Fibre</td>
<td>2.21±</td>
<td>2.16±</td>
<td>2.15±</td>
</tr>
<tr>
<td>±0.03</td>
<td>0.03</td>
<td>±0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>CHO</td>
<td>76.39±</td>
<td>76.45±</td>
<td>77.31±</td>
</tr>
<tr>
<td>±0.03</td>
<td>0.03</td>
<td>±0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Dry Matter</td>
<td>90.73±</td>
<td>90.25±</td>
<td>89.16±</td>
</tr>
<tr>
<td>±0.03</td>
<td>0.03</td>
<td>±0.03</td>
<td>0.03</td>
</tr>
</tbody>
</table>

Values are means of three independent determinations ±SD. Mean in the same row with the same superscript are not significantly (\( p > 0.05 \)) different.

Moisture content
Moisture contents of achicha varied significantly (\( p < 0.05 \)) from 9.82 to 13.57%. The least moisture content occurred in anampu at zero month (9.79%) compared with the other samples analyzed which showed that they may have longer storage lives if packaged well and stored. The highest moisture content was observed in anampu at three months (13.57%). The low moisture contents of the three samples make them easy to store at room temperature and less prone to fungal and bacterial infections because food spoilage...
micro-organisms thrive where moisture content is very high, making them more shelf stable products. These findings agree with those of [16]. Who reported the moisture content of cocoyam, yam and sweet potato to be 8.19, 10.51 and 7.07%, respectively [17]. Also reported yam and cocoyam flours to be 9.85% and 10.99%, respectively. The moisture contents of all the samples increased as the storage period progressed. The microorganisms must have produced some moisture for metabolic activities. [18]. reported that the lower the moisture content of a product to be stored, the better the shelf stability of such product.

**Ash content**

Ash is a reflection of the inorganic mineral elements present in the samples. Some of the samples investigated contained significant quantities of ash which differed significantly (p<0.05) from each other. The values varied from 3.71 to 4.31%. The highest ash content occurred in edeofe at zero month, while the lowest ash content was observed in coccoidia after 3 months storage. However, the ash low levels could be attributed to the solubilization and leaching of nutrients into processing (cooking) water. Ash content in cassava reported by [19]. Ranged from 1.44 to 2.35%.

**Crude Protein**

The mean values of crude protein varied significantly (p<0.05) from 3.93 to 5.77%. The highest value of crude protein was found in edeofe (5.77%) at zero month and the lowest value was observed in coccoidia (3.93%) at 3 months. This study showed that crude protein content decreased with increase in storage time. This could be attributed to some reactions which might have occurred during storage. [20]. reported crude protein of Colocasia esculenta (3.8g/100g) and Xanthosoma sagittifolium (4.75g/100g). Protein in diet helps primarily to build and maintain body cells. The protein content for all the samples tested was higher than those that have been reported in literature: 0.10-0.5% for yam starch [21, 22]. 0.9-1.3% for taro starch and 0.14-0.23% for sweet potato starch [23, 24].

**Crude Fat**

The mean values of crude fat in achicha samples differed significantly (p<0.05). Crude fat content varied from 0.86 to 1.05%. The least value occurred in anampu at three months, while the highest value was observed in edeofe at zero month. Fats are vital to the structure and biological functions of cells and are used as alternative energy source. [20]. reported that Xanthosoma and Colocasia varieties of cocoyam showed only low amount of fat expressed as ether extract (about 0.44 g/100g).

**Crude Fibre content**

This study revealed that crude fibre contents varied significantly (p<0.05) from 1.64 to 2.21%. The lowest value of crude fibre was observed in coccoidia at three months, while the highest value of crude fibre was found in edeofe at zero month. Previous studies showed that crude fibre content ranged from 1.53 to 2.31 for cocoyam varieties [20]. Crude fibre represents that portion of food not used up by the body but mainly made up of cellulose together with a little lignin and is known to increase bulk stool [19]. Crude fibre consists largely of cellulose and lignin (97%) plus some mineral matter. It represents only 60-80% of the cellulose and 4-6% of the lignin. The crude fibre content is commonly used as a measure of the nutritive value of poultry and livestock feeds and also in the analysis of various foods and food products to detect adulteration, quality and quantity.

**Carbohydrate Content**

The carbohydrate contents of the samples differed significantly (p<0.05). The values ranged from 75.65 to 79.46%. The highest value of carbohydrate occurred in coccoidia at two months, while the least value of carbohydrate was observed in anampu at three months. Reported carbohydrate in yam flour (71.70%) and cocoyam (73.48%). The high content of carbohydrate in the corms agrees with the fact that tuber and root crops are generally rich in carbohydrate [17]. The reason for the observed difference in the carbohydrate may be partly attributed to the differences in their moisture content.

**Dry Matter Content**

The result of the dry matter contents varied significantly (p<0.05) from 86.44 to 90.73%. The lowest value was observed in anampu at three months while the highest value was found in edeofe at zero month. Storage of these samples resulted to decrease in dry matter content. The microorganisms must have used up some dry matter for metabolic activities.

**Proximate and Dry Matter Compositions of Mpoto during Three Months Storage**

The results in Table 2 showed the comparison of Proximate Composition of mpoto processed from three different Colocasia varieties (edeofe, coccoidia, and anampu) stored from zero to three months interval.
### Table 2: Proximate and Dry Matter Compositions (%) of Mpoto during Three Months Storage

<table>
<thead>
<tr>
<th></th>
<th>Edoefe (M)</th>
<th>Cocioindia (M)</th>
<th>Anampu (M)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>0.01</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Ash</td>
<td>15.34±</td>
<td>15.17±</td>
<td>15.22±</td>
</tr>
<tr>
<td></td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Crude Protein</td>
<td>16.83±</td>
<td>16.34±</td>
<td>15.28±</td>
</tr>
<tr>
<td></td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Crude Fat</td>
<td>0.68±</td>
<td>0.62±</td>
<td>0.94±</td>
</tr>
<tr>
<td></td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Crude Fibre</td>
<td>8.88±</td>
<td>8.37±</td>
<td>8.92±</td>
</tr>
<tr>
<td></td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>CHO</td>
<td>50.04±</td>
<td>50.69±</td>
<td>50.38±</td>
</tr>
<tr>
<td></td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
</tr>
<tr>
<td>Dry Matter</td>
<td>91.77±</td>
<td>91.17±</td>
<td>90.74±</td>
</tr>
<tr>
<td></td>
<td>0.03</td>
<td>0.03</td>
<td>0.03</td>
</tr>
</tbody>
</table>

Values are means of three independent determinations ±SD. Mean in the same row with the same superscript are not significantly p>0.05 different.

### Moisture content

Moisture content of mpoto varied significantly (p<0.05) from 8.23 to 11.36%. The moisture content of the samples was significantly (p<0.05) different. The least moisture content occurred in edoefe (8.23%) at zero and first months, while the highest moisture content was observed in anampu (11.36%) at three months.

### Ash content

The samples investigated contained significant quantities of ash which differed significantly (p<0.05) from each other. The values varied from 14.31 to 16.17%. The highest ash content occurred in cocioindia (16.17%) at zero month, while the lowest ash content was observed in edoefe (14.31%) at three months. Ash is a reflection of the inorganic mineral elements present in the samples. These values indicate that these vegetables species may be considered as good sources of minerals when compared to values (2 – 10 %) obtained for cereals and tubers (FAO).

### Crude Protein content

The mean values of crude protein ranged from 14.76 to 16.83%. The mean value was significantly (p<0.05) different. The highest value of crude protein was found in edoefe (16.83%) at zero month and the least value was observed in cocioindia (14.76%) at three months. This study showed that crude protein content decreased with increase in storage time. This could be attributed to changes in other proximate constants. Protein in diet helps primarily to build and maintain body cells. The protein content of V. unguculata (21.96 ± 0.30 %) was higher than that reported for some high value leafy vegetables such as Momordica balsamina (11.29 %) and Moringa oleifera (20.72 %) [25]. It’s worth emphasizing that plant foods which provide more than 12 % of their calorific value from proteins have been shown to be good source of proteins [26]. This suggests that mpoto leaves investigated are good sources of proteins and could play a significant role in providing cheap and available proteins for rural communities.

### Crude Fat content

The mean values of crude fat in mpoto samples differed significantly (p<0.05). Crude fat varied from 0.64 to 0.94%. The least value occurred in anampu (0.64%) at zero and first months, while the highest value was observed in edoefe (0.94%) at two months. Fats are vital to the structure and biological functions of cells and are used as alternative energy source [19].

### Carbohydrate Content

The carbohydrate contents of the samples differed significantly (p<0.05). The values ranged from 48.94 to 51.08%. The highest value of carbohydrate occurred in cocioindia (51.08%) at three months, while the least value of carbohydrate was observed in cocioindia (48.94%) at zero month. The reason for the observed difference in the carbohydrate may be partly attributed to the differences in their moisture content. The carbohydrate contents in this study were higher than 20, 23.7 and 39.05 % reported for Senna obtusifolia, Amaranthus incurvus and Momordica balsamina leaves, respectively. These values are however; lower than those reported for Corchorus tridens (75 %) and sweet potato leaves (82.8 %).

### Dry Matter content

The result of the dry matter content varied significantly (p<0.05) from 88.67% to 91.77%. The lowest value was found in edoefe (88.67%) at three months, while the highest value was found in cocoindia (91.77%) at zero month. There were no significant (p>0.05) difference among the samples. Storage of these samples resulted to decrease in dry matter content. This could be attributed to some reactions which might have occurred during storage. The microorganisms must have used up some dry matter for metabolic activities.

### Conclusion and Recommendations

#### Conclusion

This study showed the effect of processing and storage on the Cocoyam based products (achicha and mpoto). Achicha is nutritionally rich in carbohydrates. However, its composition...
varies according to the variety. Mpoto leaves examined have high contents of ash, crude protein, crude fibre with low fat content and crude fibre. All these results suggest that the studied leaves if consumed in sufficient amount would contribute greatly to the human nutritional requirement for normal growth and adequate protection against diseases arising from malnutrition.

**Recommendation**

Efforts should be geared towards determining and perfecting proper food processing techniques to encourage the full inclusion of mpoto leaves in the list of vegetables in recipes for traditional cuisines. Both achicha meal and mpoto soup can contribute significantly to the nutrient requirements of humans and could be recommended as cheap sources of nutrients. Further studies should be carried out on the effect of storage of these cocoyam-based products.

**References**


_Citation_: David-Chukwu NP, Aji RU, Ndukwe KO, Odom TC, Chukwu MN (2021) Production, Proximate Compositions and Dry Matters of Stored Achicha and Mpoto - Cocoyam Based Products. Journal of Food Technology & Nutrition Sciences. SRC/JFTNS/144.