Proximate Nutritional Analysis and Heavy Metal Composition of Dried *Moringa Oleifera* Leaves from Oshiri Onicha L.G.A, Ebonyi State, Nigeria

Offor, I.F.\(^1\), Ehiri, R.C.\(^1\), Njoku, C.N.\(^2\)

\(^1\) Department of Chemistry, Federal University Ndufu Alike, Ikwo, P.M.B. 1010, Abakaliki, Ebonyi State, Nigeria
\(^2\) Department of Industrial Chemistry, Ebonyi State University, P.M.B. 053, Abakaliki, Ebonyi State, Nigeria

**Abstract:** To carry out proximate analysis and heavy metal assessment of dried moringa leaves in order to ascertain their nutritional and metal content values. Fresh leaves of *moringa oleifera* lam were plucked from *moringa* trees growing at Egbooda, Oshiri in Onicha area of Ebonyi State, Nigeria. The period of sampling was for 2 weeks in the month of March, 2013. The leaves were air dried at room temperature and their proximate contents determined using standard analytical techniques. Ash and moisture contents were determined using the Association of Official Analytical Chemists (AOAC) method. Fat, crude fibre and protein content were determined using soxhlet fat extraction method, weende’s method and kjeldahl method respectively. In addition, carbohydrate content was determined using arithmetic difference method. Results show that the mean nutritional content of the samples were: 24.5% protein, 14.8% moisture, 17.3% crude fibre, 4.5% Fat, Ash 3.8% and 50.6% carbohydrate. Heavy metal composition was assessed using spectrophotometric and ethylenediaminetetraacetic acid (EDTA) titrimetric methods. The results indicated that Fe and Cd were not detected in the leaf samples. The metal contents in the leaves were Fe (505 mg/L), Zn (85,810.951 mg/L), Cu (1,016mg/L), Mg (386.555mg/L), Mn (79.563mg/L) and Pb (352.013 mg/L).

**Keywords:** leaves, *moringa oleifera*, nutritional value, proximate analysis

I. **INTRODUCTION**

*Moringa Oleifera* belongs to the monogeneric family of shrubs and trees, called *Moringaceae* [1]. The tree originated from Agra and Qudh in Northwestern region of India, South of the Himalayan Mountain. The tree has spread to almost all tropical belt because it is drought-resistant [2]. Also, the leaves, fruits, flowers and immature pods are edible and they form part of traditional diets in many countries of the tropics and sub-tropics [3]. The oil obtained from the seeds is pale yellow, sweet and edible. It is almost odourless and possesses an appreciable test [4].

*Moringa* is called the “Miracle Tree” for good reason. *Moringa* leaves, pods, flowers, fruits, roots, bark, and seeds can be utilized in water treatment, as food supplement and the extract can be used against bacterial or fungal skin infections. It is estimated that at least three hundred diseases can be cured by taking this supplement along with hundreds of other health benefits, it contain more than 90 nutrients, different antioxidants, and all eight essential amino acids [5]. *Moringa* is rich in many vitamins, including vitamin A, several forms of vitamin B, vitamin C, vitamin D and vitamin E. In fact, it has more of these vitamins than a variety of foods (such as carrots, oranges and milk) that claim to be excellent source of these vitamins [6]. The *Moringa* tree grows quickly in many types of soil environments. Much of the plant is edible by humans or by farm animals. The leaves are rich in protein, vitamin A, vitamin B, C and minerals [5]. *Moringa* leaves can be used in food preparation and also brewed as a tea, either alone or in combination with milk.

The term heavy metal refers to any metallic chemical element that has a relatively high density and may be toxic or poisonous even at low concentrations [7]. Heavy metals include zinc, lead, calcium, mercury etc. Heavy metals are natural components of the earth crust. They cannot be degraded or destroyed. To a small extent they enter our bodies via food, drinking water & air. As trace elements, some heavy metals (e.g. Cu, Zn, Fe) are essential for maintaining the human body metabolism. There are several sources of metal poisoning which include water contamination via lead pipes, food contamination and inhalation of contaminated dust.

Heavy metals are dangerous, because they tend to bio accumulate. Bio accumulation means an increase in the content of a chemical in a biological organism over time, compared to the chemicals content in the environment. Diverse amounts of heavy metals may be found everywhere in soil, water, air, sediments, plants etc. chemicals like heavy metals once introduced to the environment by one particular method may spread to various environmental components which may be caused by the nature of interactions occurring in this natural system. Heavy metals may chemically or physically interact with the natural compounds, which change their forms of existence in the environment. Soil variability and some environmental properties (e.g. climatic factors)
may change equilibrium found in the soil and cause leaching of trace toxic demerits like heavy metals tightly bound to soil particles [8]. The environment contains a wide range of heavy metals with varying concentration ranges depending on the surrounding geological environment and natural activities occurring or that has once occurred. These heavy metals can be Fe, Zn, Cr, Cd, Pb, Ni, Mn, Hg, etc. However, some heavy metals like Pb, Zn, Cd, Hg and Th are of great concern because of their potential effects on human health, agriculture and environment. In this study, proximate nutritional analysis and heavy metal accumulation in moringa oleifera leaves were investigated using standard analytical techniques.

II. Materials And Methods

2.1 Sampling
Fresh leaves of Moringa Oleifera were plucked from the Moringa tree growing at Egbooda Oshiri. The leaves was removed from leaf branches, spread on a clean plastic tray and air dried at room temperature for 3 weeks until no trace of water was found in it. This was done to avoid interference during analysis and also to reduce the level of water content.

2.2 Pretreatment
Moringa Oleifera leaves were ashed at 550°C overnight and the ash was dissolved in concentrated HNO₃, filtered and then diluted to 250ml with deionized water and the absorbance of the samples read directly on the spectrophotometer.

2.3 Proximate analysis
The proximate compositions of the dried Moringa leaves were determined using standard analytical methods. All measurements were done in duplicates and values presented in percentage.

2.3.1 Ash content determination
The ash content was determined using the method described in AOAC [9]. 5g of the sample was weighed into a crucible in a muffle furnace and heated at 550°C for six hours until it became gray ash. The dish was removed from the muffle furnace using crucible tong and placed in a desiccator to cool. When cooled it was re-weighed and the weight of ash was obtained by the difference.

2.3.2 Moisture content determination
The moisture content of the samples was determined using AOAC method [9]. The Petri-dish was washed thoroughly and placed in oven to dry. 5g of the sample was then placed in a pre-weighted Petri dish, and then placed in an oven to dry at 105°C for two hours. The dish and dry sample were transferred to a desiccator to cool at room temperature before being weighed again. The experiments were repeated until constant weight was obtained.

2.3.3 Fat content determination
Fat was determined using soxhlet fat extraction method [10]. 250ml boiling flask was washed thoroughly and dried in oven at 105°C for 30 minutes and then placed in a desiccator to cool. 2g of the dried sample was then weighed accurately into labeled thimbles. Cooled boiling flask was filled with 200ml of petroleum ether and boiled at 40-60°C. The extraction thimble was plugged lightly with a cotton wool and the boiling flask containing the petroleum ether was placed in the extraction thimble to boil and the soxhlet apparatus was allowed to reflux for six hours. The thimble was removed carefully, and the petroleum ether on top of the container was collected and drained into another container for reuse. When the flask was free of petroleum ether, it was removed and boiled for an hour at 105°C. It was finally transferred from the oven into a desiccator to cool before weighing.

2.3.4 Fibre content determination
Crude Fibre content was determined by Weende’s method [11]. 2g of the sample was weighed into a 250ml conical flask and 200ml of 1.25% H₂SO₄ was added and the mixture was boiled under reflux for 30minutes. The solution was filtered with whatman filter paper; the residue was rinsed thoroughly with hot water until it was no more acidic when tested using pH paper. The residue was transferred into a 250ml beaker and 200ml of 1.25% NaOH was added and boiled for 30minutes in a digestion apparatus after which it was filtered and rinsed with distilled water until the filtrate was neutral when tested with pH paper. The residue was transferred into a crucible and placed in electric oven at 100°C for eight hours to dry. It was then removed and placed in a desiccator to cool before weighing. After weighing, the sample was incinerated, cooled in a desiccator and reweighed.
2.3.5 Protein determination

Protein content of the sample was determined using the Kjeldahl method [12]. The total nitrogen was determined and multiplied by a conversion factor of 6.25 to obtain the protein content. 0.5g of the sample was weighed into a Kjeldahl digestion flask. A tablet of selenium catalyst was added to it. 20ml of H$_2$SO$_4$, 10g of Na$_2$SO$_4$, 1g of CuSO$_4$ were also added to the flask and digested by heating under a fume cupboard till the solution digested completely and changed to blue color. The solution was carefully removed and allowed to solidify for 24hrs until a white colour was obtained. The solution was dissolved with 100ml of distilled water in a 200ml volumetric flask. Then 60ml of 40% of NaOH and two pieces of zinc metal were added to the solution in the Kjeldahl distillation apparatus. The mixture was distilled until a total of 50ml distillate was collected into 250ml conical flask containing boric acid and was titrated with 0.1N H$_2$SO$_4$. The end point of the titration was observed when the color of the distillate changed to the initial color of the mixture of boric acid and screen methyl red indicator which was light pink.

2.3.6 Carbohydrate determination

The carbohydrate content of the test sample was determined by estimation using the arithmetic difference method [12].

\[
%\text{CHO} = 100 - (\% \text{fat.} + \% \text{ash} + \% \text{fiber} + \% \text{protein}) \quad (1)
\]

2.4 Heavy metal analysis

The heavy metals content of dried Moringa leaves was determined using AOAC method [13] and EDTA titrimetric method [9]. All determinations were done in duplicates and values of heavy metals were reported in Mg/L.

2.4.1 Determination of Zinc

This was determined by EDTA titrimetric method [9]. 2ml of sample solution was measured into a conical flask and 2ml of buffer solution was added to it. Then, 2 drops of Eriochrome Black T Indicator was added and the mixture titrated with 0.01 EDTA until the color changed from wine red to blue.

2.4.2 Determination of Lead

The Lead content in the sample was determined using spectrophotometric method [13]. 10ml of the sample solution was measured into a beaker, followed by addition of 5 drops of 10% KCN, 5ml of 1.2M NH$_3$ solution and 5ml of 10% NaSO$_4$. The resulting mixture was made up to the 50ml with distilled water. The spectrophotometer was set at wavelength of 430nm after calibration and the absorbance reading of the sample solution taken.

2.4.3 Determination of Iron

To determine the iron content, 5ml of the sample was pipetted into a 50ml of volumetric flask. 5ml of 17% CH$_3$COONa was then measured into the digested sample in a volumetric flask followed by addition of 0.5ml of 10% hydroxylamine hydrochloride and 5ml of 0.25% O-phenanthroline. Then the mixture was made up to 50ml with distilled water. The wavelength of spectrophotometer was set at 510nm for iron [13].

2.4.4 Determination of Copper

To determine the copper content, 10ml of the digested sample was measured into 50ml of volumetric flask. 10ml of 0.01N NH$_3$OH was added to the solution and the resultant mixture made up to 50ml using distilled water. It was allowed to develop for 30mins before the absorbance reading was taken with aid of the spectrophotometer at wavelength of 620nm [13].

2.4.5 Determination of Manganese

Manganese content was determined by using the spectrophotometer. 20ml of the digested sample was measured into 50ml volumetric flask, then 5ml of phosphorus and 0.3g of potassium periodate were added and the mixture shaken before boiling for 15mins. It was allowed to cool before diluting with distilled water to make up to the 50m mark on the standard flask. The resultant solution was then allowed to develop for 30mins. The absorbance of the solution was read using the spectrometer machine at wavelength of 520nm [13].

2.4.6 Determination of Cadmium

Cadmium determination was carried out by EDTA titrimetric method using Xylenol Orange as an indicator. 25ml of the sample was measured into a 250ml conical flask. Then 5ml of distilled water, 3 drops of xylenol orange indicator and 1 drop of dilute H$_2$SO$_4$ were added to the sample solution. Thereafter, the color
turned to yellow and hexamine powder was added until the color changed to deep red. It was then titrated with 0.05m EDTA until color change from deep red to initial yellow was observed.

2.4.7 Determination of Calcium
This was determined by ethylenediaminetetraaetic acid (EDTA) titrimetric method [14]. 10ml of the digested sample was measured into a 250ml conical flask. A pinch of potassium cyanide, a pinch of hydroxylamine hydrochloride, 5ml of 10% potassium hydroxide were added and shaken gently until the solids dissolve. Then a pinch of indicator (Pattons & Reader’s reagent) was added and the mixture titrated with the 0.01M EDTA solution until the color changed from wine red to blue which is the end point.

2.4.8 Determination of Magnesium
Magnesium was determined using EDTA titrimetric method [14]. 10ml of the digested sample was pipetted into 250ml conical flask, a pinch of KCN, a pinch of hydroxyl ammonium chloride solution and 5ml of 10% potassium hydroxide were added and shaken gently until the solid dissolved. A pinch of Eriochrome black T indicator was added and the mixture was titrated with 0.01M EDTA solution until the wine red color changes to blue color which is the end point.

III. Results And Discussions
Table 1 shows the results of proximate analysis of dried Moringa Oleifera. The results indicated that dried Moringa leaves contained appreciable amount of crude protein content (24.5 ± 0.9%) making it to be a good source of supplementary protein for man and livestock, The results also showed that Moringa leaves contain nutritious compounds.

Table 1: Proximate contents of the analyzed sample

<table>
<thead>
<tr>
<th>Parameter (%)</th>
<th>Mean ± S.E.M</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture</td>
<td>14.8 ± 0.2</td>
</tr>
<tr>
<td>Fat</td>
<td>4.5 ± 0.1</td>
</tr>
<tr>
<td>Crude Protein</td>
<td>24.2 ± 0.9</td>
</tr>
<tr>
<td>Crude Fibre</td>
<td>17.3 ± 0.2</td>
</tr>
<tr>
<td>Ash</td>
<td>3.8 ± 0.2</td>
</tr>
<tr>
<td>Carbohydrate</td>
<td>50.4 ± 0.2</td>
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</tbody>
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* S.E.M=Standard error of the Mean

Other similar studies have reported different values for the protein content of moringa leaves ranging from 16-40% [15-23]. Appreciable level of crude in moringa leaves is acceptable as it prevents the occurrence of diseases thereby promoting good health [24]. The carbohydrate content of dried moringa leaves was 50.4 ± 0.2%.This value however differs from similar findings which recorded carbohydrate contents of 45.43%, 43.88% and 63.11% respectively [20, 23, 25]. Carbohydrate deficiency causes depletion of body tissues [24]. Sufficiency of Carbohydrate is however necessary for optimum functioning of the brain, heart, nervous, digestive and immune systems [24]. High content of moisture (14.8±0.2%) in moringa leaves was recorded in this study which was relatively higher than that obtained in similar researches with moisture content values of 11.76%, 3.21%, 76.53% and 9.53% respectively [20, 23, 25-26]. Ash content of 3.82±0.2% was recorded in this study which was comparatively low when compared to values obtained in similar research with ash content values of 7.64%, 7.93%, 7.13% and 10.64% respectively [20, 23, 25-26]. The variation in nutritional makeup of the dried moringa leaves analyzed in this study and that of other similar researches could be attributed to the difference in genetic makeup of the plant and varying climatic and soil factors. Table 2 however shows that dried Moringa leaves contain high amounts of heavy metals. Iron content in the dried leaves of Moringa was 505 ± 53.0mg/L, indicating that the leaves could be a potential source of Fe supplement for human and livestock. Fe is an essential trace element for normal functioning of the central nervous system and in the oxidation of carbohydrates, proteins and fats [27]. Iron is also a necessary component of hemoglobin and myoglobin for oxygen transport and cellular processes of growth and division [28].
The mean Fe content in the sample was 505mg/L, which was comparable to similar studies done with reported Fe values of 103.75mg, 490mg, 28.2mg, 107ppm, 318.81ppm and 870mg of Fe respectively in dried Moringa leaves [20, 23, 25, 28-30]. Thus, Iron which is commonly deficient in many plant-based diets was found in abundance in these cited works.

The presence of Zn in high amounts is of special interest in view of the importance of Zn in the diet of animals and humans. Result from this analysis showed higher levels of zinc (85811mg/L ± 0.384) than a similar finding of 25.5mg in dried Moringa leaves [31]. Furthermore, related researches also reported the level of zinc in dried Moringa leaves as 148.54mg, 60ppm and 57.34ppm respectively [20, 23, 25]. Zinc is essential for the synthesis of DNA, RNA, insulin and function of several enzymes, zinc is also required for cell reproduction and growth especially the sperm cells [32].

The dried Moringa leaves contained Cu, which is considered to have strong effects on the immune system. Cu content found in the sample was 1.016mg. Other studies have reported variable Cu contents of 0.57mg, 0.143mg, 6.10mg, 5.73ppm, and 4.66mg respectively [5, 20, 23, 27, 33]. The Mg content of dried Moringa leaves, from the result was 386.6 ± 1.01mg/L. Mg is an essential element that the body required in relatively large quantities compared to other minerals as it plays an important role in the structure of skeleton and muscles. Other studies have reported variable contents of Mg in Moringa leave as 0.38ppm, 1.03ppm and 107.56mg respectively [23, 26, 33].

Calcium which is required for normal growth, strong muscles and skeletal development was surprisingly not detected in the leaves. It is suspected that Mg and Zn probably displaced the elemental Ca in the leaves and consequently plays the role of Ca in the plant. This view was further strengthened by the high amount of Mg found in the leaves (386.6mg/L). However other reports have it that the calcium content of dried Moringa leaves was 98.67mg, 2003mg, 3.65%, 2.47ppm, 1.91 and 3.4% [4-5, 20, 23, 26, 33].

Thus, Dried Moringa leaves are available as a good source of Ca to farm animals or humans. Cadmium was also not detected in the leave samples tested.

For the case of Mn, dried Moringa leaves had Mn content of 79.56 ± 1.188mg/L which was comparable to that obtained by other researchers in similar studies with Mn values of 13.55mg/L, 86.80mg/L, 81.65mg/L and 21.70mg/L respectively [26, 23, 33, 20]. Lead was also found in the leaves of Moringa at very high value of 352.01 ± 2.36mg/L. The elevated level of Pb observed was probably due to high large deposit of Galena (PbS) and other Pb ores in Oshiri, Ebonyi State where the leaves were harvested. In addition, a similar study reported a Pb level of 2.96mg/L in dried Moringa leaves investigated [23].

IV. Conclusion

Proximate analysis results show that dried moringa leaves is a good source of important nutrients and thus, the plant might be explored as a viable supplement in both animal and human food. Other nutritional contents in moringa leaves not covered in this study and the possible roles of edaphic factors on the nutritional makeup of moringa plants are areas for further investigation in future research. Furthermore, the phytochemical constituents of moringa oleifera leaves can be further explored in the search for further uses of moringa plants as herbal remedies.

In addition, the variation in the heavy metal contents of Moringa leaves found in this study when compared to earlier studies may be due to climatic and edaphic factors, solvents used for the analysis, the cultivation method used and age of the plant. Furthermore, the leaves have been known to serve as a herbal tonic and for the maintenance of tissues and cell membranes [34].

References


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