Research Article

Metallic nutrients in enset (Ensete Ventricosum) corm and soil sample from some West Shoa Zone, Oromia Regional State, Ethiopia

Teressa Bedada* and Alemayehu Abebaw
Department of Chemistry, College of Natural and Computational Sciences Ambo University, P.O. Box 19, Ambo, Ethiopia

Abstract

The aim of this study was to investigate the essential and non-essential metals concentration in corm of Ensete ventricosum and soil samples from some West Showa Zone. Ca and Mg were determined complecometric titration method, K and Na were analyzed using Flame Absorption Emission spectroscopy and the other metals with flame atomic absorption spectroscopy after appropriate quality control measures were undertaken to verify and maintain the quality of the data generated. The optimized wet digestion method for corm and soil analysis was found effective for all of the minerals and as it was evaluated through the recovery experiment, a good percentage recovery 95% (Fe concentration in corm) to 112% (Pb concentration in soil) was obtained for the minerals identified. The results of this study showed that the average the metallic nutrient concentrations of elements in the soil sample were ranged in order of decreasing in (mg.kg\(^{-1}\)) 3521.11(K) >3497.85(Mg) > 3461.59(Fe) > 3294.93(Ca) >1096.89(Na) > (93.99(Mn) >16.74(Zn) > 3.77(Cu) > 0.26(Cd) but, the concentration of Pb was not detected and the metallic nutrient concentrations of elements in the corm were ranged in order of decreasing in (mg.kg\(^{-1}\)) 16425.13(Ca) >13813.33(K) > 1323.55(Mg) > 1131.11(Na)  > 76.78(Fe) > 16.16(Zn) >2.77(Cu) >2.77(Mn) but, Cd and Pb were not detected. The nutrient concentrations of the metals were also compared with recommended maximum permissible limits and some international reports; and found to be in a good agreement indicating no exposure risk of using the corm of Ensete ventricosum under the current situation. Statistical test of significance using ANOVA revealed that there were significant differences (P<0.05) between the values of metals in the corm and soil samples obtained from all the sampling sites except Zn concentration for corm and Ca concentration in soil) is not found.

Introduction

Root and tuber crops are widely cultivated in southern Ethiopia, which are supporting a considerable portion of the country’s population as source of food. Prominent among these are potato (Solanum tuberosum L.), sweet potato (Ipomoea batatas L.), Enset (E. ventricosum), Godere (Colocasia esculanta L.), Yams (Dioscorea spp.), Ethiopian dinch (Colesus parviflorus), koteharrie (Diaspora bulbiferous) and Anchote (Coccinia abyssinica). Among these, Enset, Anchote and some yams are endemic to Ethiopia [1]. Enset based farming systems play an important role in food security in Ethiopia [2] and Ensete ventricosum is one of the indigenous root crops widely cultivated in the Central, South and South Western parts of Ethiopia, but recurrent droughts have led to the expansion of Enset cultivation to other parts of the country [3,4].

Ensete ventricosum parts contained high percent of water (85 to 90%), which is beneficial when used as fodder during dry periods. Corm of Ensete ventricosum contained 17 of 20 amino acids. Leaves had 13% protein, among the highest available in Ethiopia, 20% crude fibre and 10% sugar. The pseudostem the main food source, was rich in 80% of soluble carbohydrates and 65% of starch, but has low protein content 4% [5].

Eating the right foods is an important part of maintaining a healthy lifestyle. A single day’s intake of nutrients may affect the body’s organs only slightly but over years and decades the effects of unhealthy diet compounds into disease, shortened
lifespan, and a less active lower productive life. The nutrients that we intake today will become part of us tomorrow. The nature and composition of what we eat as determines our future, Enset as a food would have its own influential impact on millions of peoples in Ethiopia. The corm of Enset has rich in essential nutrients and low in non-essential nutrients [6].

The human body requires a number of nutrition to preserve a good health that nutrition accumulated in different parts of plants (FAO, 2004). Thus plants are intermediate reservoirs through which trace elements from soil and partly from water and air, transfer to man and animal (Rogan, et al. 2009). The content of heavy metals is one of the criteria for the use of plant material as food or traditional medicines. Hence determination of mineral compositions in food and medicinal plant is essential for understanding their nutritive importance and health risk [7].

However, no literature report was found on comparative determination the concentration of essential and non-essential metals in corm of Ensete ventricosum with its supporting soil samples. Therefore, the aim of this study to compare the essential and non-essential metals concentration in corm of Ensete ventricosum and soil environment from some different locations of West Shoa Zone, Oromia Regional State, Ethiopia by using flame atomic absorption spectroscopy.

Materials and methods

Apparatus and instruments

Polyethylene plastic bags, Electronic analytical balance with 0.0001g sensitivity (AA-200DS Deriver Instrument Company, German), A 250 ml round bottomed flask fitted and reflux condensers, Whatman filter paper (No.42 150 mm, England), Digestive furnace (Model KDN-20C, China), Flame Photometer (ELICO, CL-378, India) and flame atomic absorption spectrophotometer FAAS (Buck Scientific Model 210 VGG, USA).

Chemicals and reagents

All reagents were analytical reagent grade. The reagents and chemicals used in this study were: HNO₃ (65–68%, Uni-Chem® Chemical Reagent, India), HClO₄ (70–72%, Uni-Chem® Chemical Reagent, India), H₂O₂ (30%, Uni-Chem®, India, EDTA-Na₂ (98.5–101%, Unic-Chem®, India) and Stock standard of metals.

Description of the study areas

The study was conducted in Oromia National Regional State, West Shoa administrative zone. It was found between 8°17’–9°56’N and 37°1’–38°45’E. The zone was bounded with Amhara Regional State in the Northern part, East Wollega and Horro Guduru Wollega in the West and North West, Jimma Zone in the South West, South West Shoa Zone in the South East, and North Shoa Zone in the North East. West Shoa Zone was 170Km long from North to South and 183Km wide from East to West. For administrative purpose the Zone is divided into eighteen districts Figure 1. Ambo was the capital city of zone and far 114Km to the West of Addis Ababa on the main way from Addis to Nekemte road.

The Zone has variable topography consisting of a high and rugged central plateau, flat and gentle slope, and the peripheral lowlands. The Mean annual temperature of the zone ranges from 10°C to 22°C in the highlands and from 22°C to 30°C in the lowlands. Thereby, Annual average rainfall in the area is ranging from 1200 to over 2000mm. The Zone has three agro climatic zones, namely Dega (57%), Woina dega (25%) and Kolla (18%) (Zone Office Agriculture and Rural Developing, 2015).

Figure 1: Map of study areas.
Sample collection and protocol

The corm of Ensete ventricosum sample was collected in January, 2016 from the three agricultural areas with its supporting soil. Each sample was collected purposely from four different sub-sites (farm lands) to provide replicate samples. From the three agricultural areas corm can be prepared according to the traditional method. The edible designated plant out of the land was cut into three parts for the separation of pseudostem and Corm with knife (Figure 2). The soil sample was collected from the surface 15cm-25cm depth of the same four sampling areas of Enset by spade. Finally three corm and soil samples one from each stated areas were collected and put into clean cooled.

Sample preparation

Corm and soil samples were collected from each sub-sites (kebeles) were air dried for three days to remove moisture and all clods and clumps. The samples were grinded with a mortar and pestle and then sieved through a 2mm mesh sieve. The four sub-samples were mixed equal proportion together to form a composite sample that represents each sampling areas. The powdered two samples were placed in pre-cleaned screw capped polyethylene container and stored in desiccators containing calcium chloride to keep constant dry weight till digestion [8].

Optimization of digestion procedure for corm and soil samples

0.5g of air dried and homogenized corm and soil samples were transferred into a 250ml round bottom flask. To this was added 3.5ml a mixture of HNO₃ (69-72%) and HClO₄ (70%) with a volume ratio of 2:1.5 for corm and 4.5ml a mixture of HNO₃ (69-72%), HClO₄ (70%) and H₂O₂ (30%) with a volume ratio of 2:1.5:1 for soil. The mixture were digested on a micro Kjeldahl digestion apparatus by setting the temperature 210 and 230 for 1:45hr and 2:30hr respectively. Then, after the digested solution were allowed to cool for 20min without dismantling the condenser. To the cooled solution 25ml of distilled water was added 3.5ml a mixture of HNO₃ (69-72%) and HClO₄ (70%) while to minimize dissolution of was added to dissolve the precipitate formed on cooling and the condenser. From the solution were allowed to cool for 20min without dismantling the condenser from the micro digestion apparatus by setting the temperature 210 and 230 for 1:45hr and 2:30hr respectively. Then, after the digested solution were allowed to cool for 20min without dismantling the condenser. To the cooled solution 25ml of distilled water was added and the solution was filtered with Whatman filter paper by the digest residue while filtering with Whatman filter paper. The round bottom flask was rinsed subsequently with 5ml distilled water until the total volume reached around 45ml. To this final solution, 1% lanthanum nitrate solution was added and the solution was filled to the mark 50ml with distilled water. The digested samples were kept in the refrigerator, until the level of all the metals in the sample solutions were determined by FAAS and FAES.

Method validation and quality control

Precision and accuracy: Precision and accuracy of the analytical method was assessed by repeatability and recovery studies of Matrix Spike (MS) and Laboratory Control Samples (LCS). Recovery study was performed by spiking three replicate of corm and soil samples with a known concentration of metal standard solution (mid-range calibration concentration). The spiked samples were then subjected to the same digestion procedure like the actual sample. Precision was calculated by

\[ \text{RSD} = \frac{\sum_{i=1}^{n} |X_i - \bar{X}|}{n} \times 100 \]

accuracy was calculated by this equation \[9\].

\[ \%R = \frac{C_{\text{Spike sample}} - C_{\text{Unspike sample}}}{C_{\text{added}}} \]

Instrument Detection Limit (IDL)

Instrument Detection Limits (IDLs) was estimated by taking seven replicate measurements of the calibration blank (Distilled water). The IDL is calculated to the concentration equal to three times the standard deviation of seven replicate measurements of blank \[10\].

\[ \text{IDL} = 3 \times S_b \]

Where, \( S_b \) is standard deviation of blank (\( n = 7 \)) and IDL is Instrument detection limit.

Method Detection Limit (MDL)

Method detection Limit is the minimum concentration of analyze that can be identified measured and reported with 99% confidence that the analyze concentration is greater than zero. MDL was based on seven replicate measurements of a series of calibration blanks (reagent blank) that are carried through the entire sample preparation scheme \[11\]. The MDL was calculated by:

\[ \text{MDL} = S \times \text{T-test} \]

Where, \( S \) is standard deviation of the replicated analysis with n-1 degree of freedom, \( t = 3.71 \) (\( T- \) test value for a 99% of confidence level for six degrees of freedom).

Method Quantification Limit (MQL)

Method quantification limit was obtained from analysis of seven reagents blanks which were digested in the same digestion procedure as actual samples. The method quantification limit was calculated by multiplying standard deviation of the reagent blank by ten plus the mean of the reagent blank signals \[11\]. It can be calculated by:

\[ \text{MQL} = \bar{X}_{\text{blank}} + 10 \times S_{\text{blank}} \]

Where, \( \bar{X}_{\text{blank}} \) is the mean of blank, \( S_{\text{blank}} \) is standard deviation of the blank.

Method blank

The method blank accounts for contamination that may occur during sample preparation and analysis. These could arise from the reagents, the glassware or the laboratory environment \[8\]. Sucrose was used as matrix since there was
no other plant and clear soil that can serve as the matrix for the corm and soil samples. The blank which was prepared from the sucrose and any reagents used for the digestion was taken through the entire measurement procedure to detect contamination from reagents, sample handling, and the entire measurement process [12].

Matrix spike

Matrix Spike (MS) is portion of a sample spiked with known concentration(s) of target analyte(s). The spiking occurs prior to sample preparation and analysis. The purpose of a matrix spike sample is to determine whether the sample matrix contributes bias to the analytical results [13]. In this study, Matrix spike was prepared for each sample item by spiking aliquots of 0.5g of each corm and soil samples with 2.5ml standards mixture solution giving concentrations of 1.0mgL-1 for K, Zn, Cd and Pb; 2.0mgL-1 for Na, Cu, Fe and Mn. They were all carried through the same digestion and analysis steps as an unspiked sample (Table 1). And the mean recovery values of Matrix Spikes were calculated by:

$$\%R = \frac{C_{\text{Spiked sample}} - C_{\text{Unspike sample}}}{C_{\text{Added}}}$$

**Transfer factor of metals from soil to Ensete Ventricosum**

Transfer factor is the ratio of the concentration of metals in a plant to the concentration of metals in soil. Transfer factor for each metals was computed based on the method Harrison and Chirigawi (1989) as described by [14] according to the following formula.

$$TF = \frac{p_m \text{ mg kg}^{-1}}{S_m \text{ µg g}^{-1}}$$

Where, $p_m$ is metals concentration in plant and $S_m$ is metals concentration in soil.

**Statistical analysis**

Analysis of variance (ANOVA) and F-test at $p<0.05$ are used to examine statically significant differences in the mean concentrations of metals among groups of soil and corm of *E. Ventricosum*. A probability level of $p<0.05$ is considered statistically significant. All statistical analysis was done by Microsoft Office Excel 2007 was used for data analysis and SPSS Version 16.0 Software Window was used for Analysis of variance (ANOVA) and correlation between metals in corm and soil samples [11].

**Result and discussion**

**Instrument detection, method detection and quantification limits**

As indicated Table 2 the method detection values ranged from 0.08mgkg$^{-1}$ (Cd in soil) to 2.441mg.kg$^{-1}$ (Zn in corm) and the MQL values lied in range from 0.3mg.kg$^{-1}$ (Pd in Corm) 4.21 mgkg$^{-1}$ (Mn in corm). The results revealed the both MDL and MQL values were greater than the IDL; hence, the results of the analysis could be reliable.

**Calibration**

The Calibration curves for the various concentrations were ranged between 0.9969 and 0.9999, which were all greater than the required limit (0.995) for trace element analysis. This showed that there was good correlation (relationship) between concentration and absorbance indicating good calibration of instrument (Table 2).

**Method precision and accuracy**

As it can be indicated Table 3, The mean percent recovery values ranged between 94.85% Fe in soil to 109% (Cu in corm), all lied in the acceptable range (80–120%) for metal analysis [12]. This showed that the analytical method provided results in the required level of accuracy. The RSD values of recovery was ranged between 0.05% (Mn in corm) to 12.22% (K in corm), all lied under the required limit ≤15% [15].

**Laboratory control samples result**

The percent recovery values of LCS measurements lied
in the range 94.0% (Na in soil) to 113.6% (Zn in bulla) and their relative standard deviations 0.09 (Cu in soil) to 11.37 (Cd in soil), and all the values were found under standard control limits 80–120% for LCS recovery, and 515% for RSD [15]. This showed that the method used for the study has provided the required level of accuracy and precision throughout the analytical process.

**Optimizations the digestion procedures of soil samples**

**Digestion of soil sample:** 0.5g of soil with a mixture of 2ml of concentration of HNO₃, 1.5ml concentration of HClO₄ and 1ml concentration of H₂O₂ digested at a temperature of 230 for 2:30 hours gave a clear colorless solution respectively. These optimum conditions were selected based on clarity of digest; minimum reagent consumption, minimum digestion time, and minimum temperature applied for complete digestion of samples. After digestion the samples were cooled and diluted to 50ml (Table 4).

**Metal concentration in soil used for Enset (Ensete ventricosum) cultivated**

The analyzed K concentration of the soil sample was ranged from 3403.33 to 3710mgkg⁻¹. The highest K concentration was observed in the Dire Enchini and the lowest in the Jeldu, Which is found within the permissible level 5000 to 30000 and 1000 to 15000mgkg⁻¹ of Ca and Mg in soil respectively (EPA, 2002). When, have compared the concentration of Ca and Mg in this study areas soil samples similar with Ca concentration but greater than in case of Mg concentration with the one reported by Wodaje and Alemayehu [16], which was ranged from 23866 to 32262 and 1751 to 4288mgkg⁻¹.

The highest Mg concentration was observed in te Jeldu and the lowest in the Dire Enchini, Which is found within the permissible level 5000 to 30000 and 1000 to 15000mgkg⁻¹ of Ca and Mg in soil respectively (EPA, 2002). When, have compared the concentration of Ca and Mg in this study areas soil samples similar with Ca concentration but greater than in case of Mg concentration with the one reported by Wodaje and Alemayehu [16], which was ranged from 23866 to 32262 and 1751 to 4288mgkg⁻¹.

The levels of six trace metals were analyzed in soil samples the results of total concentration of all metals of interest in the soil samples. The mean concentrations of Fe, Mn, Zn, Cu, Pb and Cd were presented in this study 102.87 to 184.1, 83.1 to 99.87, 11.67 to 20.13, 2.5 to 3.07, 0.49mgkg⁻¹ and not detect respectively in the soil samples. The concentrations of all metals in the analyzed soil samples of Enset environment were under the EPA maximum permissible limit of typical concentration in soil dry matter, Fe, Mn, Cu, Zn and Cd are 10–50, 20–30, 2–100, 10–200, 0.1–1mg kg⁻¹ respectively (EPA, 2002) Table 5.

**Optimizations the digestion procedures of corm sample**

**Digestion of corm of E. Ventricosum samples:** 0.5g of corm with a mixture of 2ml of concentration of HNO₃, 1.5ml concentration of HClO₄ digested at a temperature of 210 for 1:45 hours.
Metal levels in corm of *E. Ventricosum*

The mean concentrations of K was the second most accumulated corm next to calcium. The values of K concentration in corm sample were studied areas ranged from 12050.00 to 15013.33 mg kg\(^{-1}\). The highest K concentration was observed in the Jibat corm and lowest in the Jeldu corm, which is below the permissible limit of K concentration in plant dry matter was ranged from 15% [18]. While, as have compared the concentration of K in corm in this study less than in corm with the one reported by Ayalew, et al. [6] around welkete, which was ranged from 14100 to 32200 mg kg\(^{-1}\).

The analyzed sodium concentration of Corm was ranged from 1023.33 to 1326.67 mg kg\(^{-1}\). The lowest Na concentration was observed in Jeldu and highest in Jibat, This concentration was below the WHO recommendation on sodium maximum consumption for adults, which is 2 g sodium/day (WHO, 2012).

The analytical data of Mg concentration in corm of Enset was ranged from 1268.67 to 1370 mg kg\(^{-1}\). The highest Mg concentration was observed in the Jibat and lowest in the Jeldu, which is below the maximum allowed concentration of Mg 0.1 to 0.4% in dry matter of plant FAO [18]. The mean concentration of Mg in corm of Enset in this study less than in corm with the one reported by Ayalew, et al. [6] around welkete, which was ranged from 24900 to 26900 mg kg\(^{-1}\).

The average concentration of Calcium in corm was studied areas ranged from 15912.22 to 16875.78 mg kg\(^{-1}\). When compared the concentration of Ca, was observed highest in the Jibat lowest in the Jeldu. This concentration found within the range of the maximum permissible limit of Ca in plant dry matter was ranged from 15% [18]. While, as have compared the amount of Ca in corm of Enset in this study within the same range with the one reported by Ayalew, et al. [6] around welkete, which was ranged from 36100 to 39100 mg kg\(^{-1}\).

The concentration of iron was analyzed in corm of Enset sample was ranged from 53.67 to 62.0 mg kg\(^{-1}\). The lowest Fe concentration was observed in the Jibat Corm whereas highest in the Jeldu Corm. This concentration found within the range of the maximum permissible limit of iron between 50 to 250 mg kg\(^{-1}\) in plant dry matter [18]. When, as have compared the concentration of Fe in this study almost similar concentration in corm with the one reported by Ayalew, et al. [6], which was ranged from 18.2 to 54.4 mg kg\(^{-1}\).

The Concentration of Manganese in Corm of Enset was found between 1.13 to 2.40 mg kg\(^{-1}\), which was found below the FAO [18] maximum permissible limit of 20 to 300 mg kg\(^{-1}\). When as compared the manganese concentration in Corm from the three sites the lowest Mn concentration was observed in the Jeldu Corm but, the highest in the Dire Enchini Corm and the concentration of Mn in Corm of Enset in this study slightly less than with the one reported by Ayalew, et al. [6], around welkete, which was ranged from ND to 5.6 mg kg\(^{-1}\) and 2.0 to 5.0 mg kg\(^{-1}\).

The Concentration of Zn accumulated in corm next to iron among of micro nutrients from the study areas, which was ranged from 15.43 to 17.43 mg kg\(^{-1}\). While, the lowest Zn concentration is observed in the Jibat and the highest in the Dire Enchini, Which was found below the maximum permissible limit of Zn set FAO [18], was ranged from 50 to 250 mg kg\(^{-1}\) in plant dry matter. The concentration of Zn in this study with the one reported by Ayalew, et al. [6], which was ranged from 2.11 to 42.3 mg kg\(^{-1}\).

**Table 6**: Mean Concentration of Metals (mg kg\(^{-1}\)) in Soil Samples from each Sites.

| Metals | Dire Enchini | Jeldu | Jibat | Max. safe Limit in soil (mg kg\(^{-1}\)) |
|--------|-------------|-------|-------|----------------------------------------|
| Na     | 1047.33±15.53 | 1123.33±20.83 | 1120.00±51.96 | NA |
| K      | 3403.33±5.77  | 3710.00±78.10  | 3450.00±170   | 100 a |
| Mg     | 267.78±10.71  | 3934.67±5.36   | 3911.11±28.31 | NA |
| Ca     | 2647.33±10.08 | 3949.12±43.74  | 3788.83±50.08 | NA |
| Fe     | 183.90±0.17   | 102.87±2.11    | 184.10±0.85   | 5000 a |
| Mn     | 9.987±0.06    | 83.10±1.82     | 99.00±2.34    | 200 a |
| Zn     | 18.43±0.67    | 11.67±0.25     | 20.13±1.21    | 300 a |
| Cu     | 3.07±0.23     | 2.73±0.15      | 2.50±0.10     | 100 a |
| Cd     | 0.29±0.01     | ND             | 0.49±0.01     | 3 a |
| Pb     | ND            | ND             | ND            | 100 a |

ND is not detected; * source: FAO/WHO (Codex Alimentarium commission [17]).

**Table 5**: Optimization of digestion procedure for 0.5g the Corm sample.

| Trial | Amount of samples | Volume of reagents | Temperature (°C) | Times(Hr) | Observation |
|-------|-------------------|--------------------|-----------------|-----------|-------------|
|       |                   | HNO\(_3\) | HClO\(_4\) |               |             |
| 1     | 0.5               | 4       | 2      | 250         | 2.45        | Yellow      |
| 2     | 0.5               | 3.5     | 1.5    | 250         | 2.45        | Light yellow|
| 3     | 0.5               | 3       | 3      | 250         | 2.45        | Clear and colorless |
| 4     | 0.5               | 3       | 2      | 250         | 2.45        | Very Clear and colorless |
| 5     | 0.5               | 2.5     | 2.5    | 250         | 2.45        | Light yellow |
| 6     | 0.5               | 2       | 2      | 250         | 2.45        | Light yellow |
| 7     | 0.5               | 2       | 2      | 250         | 2.45        | Very Clear and colorless |
| 8     | 0.5               | 2*      | 1.5*   | 250         | 2.45        | Very Clear and colorless |
| 9     | 0.5               | 1.5     | 2      | 250         | 2.45        | Very Clear and colorless |

Optimization of Temperature

| Trial | Amount of samples | Volume of reagents | Temperature (°C) | Times(Hr) | Observation |
|-------|-------------------|--------------------|-----------------|-----------|-------------|
| 1     | 0.5               | 2.5                | 150             | 2.45      | Light yellow |
| 2     | 0.5               | 2                   | 170             | 2.45      | Clear light yellow |
| 3     | 0.5               | 2                   | 1.5             | 190       | Very Clear and colorless |
| 4     | 0.5               | 2                   | 210*            | 2.45      | Very Clear and colorless |
| 5     | 0.5               | 2                   | 230             | 2.45      | Very Clear and colorless |

Optimization of Time

| Trial | Amount of samples | Volume of reagents | Temperature (°C) | Times(Hr) | Observation |
|-------|-------------------|--------------------|-----------------|-----------|-------------|
| 1     | 0.5               | 2                   | 210             | 1.15      | Light yellow |
| 2     | 0.5               | 2                   | 210             | 1.30      | Light yellow |
| 3     | 0.5               | 2                   | 210             | 1.45*     | Very Clear and colorless |
| 4     | 0.5               | 2                   | 210             | 2.00      | Very Clear and colorless |
| 5     | 0.5               | 2                   | 210             | 2.15      | Very Clear and colorless |

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The mean concentrations of Cadmium and lead were not detected in corm sample, those were below the acceptable concentration for food stuff which is around 1 ppm [19–22], indicating no exposure risk to Cd and Pb. The lowest level of Cd which can cause yield reduction is 5–30 ppm [19–22] Table 7.

**Transfer factor of metals from soil to E. ventricosum**

The transfer factor of metals from soil to *E. ventricosum*, Ca, K, Na, Mg and Cu concentration were more accumulated. When compared the transfer factor among the different metals, Ca, K, Cu, Na and Zn concentration showed the maximum transfer factor value (Table 8), which ranged from 1.0 Mg in the Jibat to 6.0 Ca in the Dire Enchini and Mn and Fe concentration were minimum value which, ranging from 0.01 in the Jeldu to 0.6 in the Jeldu. This is indicated that corm is rich in essential metals. The high level of these metals in Ensete ventricosum high due to direct deposition a foliar absorption more than the translocation upper part of the plant to the root of the plants. This can be attributed to the redistribution of elements within the soil profile (Zubillaga, et al. 2008).

**Conclusion**

The level of essential and non-essential elements in corm of Enset and soil sample was determined by flame atomic absorption spectrometry, flame photometry and complexometric titration with EDTA. The distribution of the selected essential and non-essential metals over corm sample of *E. Ventricosum* and soil were observed, they were found to vary in the order of decreasing Ca > K > Mg > Na > Fe > Zn > Cu > Mn but, Cd and Pb could not be detected from Corm sample and K > Ca > Mg > Na > Fe > Mn > Zn > Cu > Cd > Pb (ND) in soil samples of the three areas. Based on the WHO recommended limit and FAO (2008) the maximum permissible limit for plant, K, Na, Mg, Ca, Fe, Mn, Cu, Zn and Pb, Cd were not found to cause any risk to the people by consuming E. Ventricosum plants grown in the area where the E. Ventricosum is planted (studied areas). Statistical test of significance using ANOVA revealed that there were significant differences (P<0.05) between the values of metals in the corm and soil samples obtained from all the sampling sites except Zn concentration in corm and Ca concentration in soil.

Statistical test of significance using ANOVA revealed that there were significant differences (P<0.05) between the values of metals in the corm and soil samples obtained from all the sampling sites except Zn concentration for corm and Ca concentration in soil) is not found.

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**Table 7: Mean concentration of metals (mgkg⁻¹) in Corm sample from Each sites (n = 3).**

| Metals | Dire Enchini | Jeldu | Jibat |
|--------|-------------|-------|-------|
| Na     | 1023.33±25.17 | 1043.33±15.27 | 1326.67±4163 |
| Ca     | 14376.67±137.48 | 12590.00±66.58 | 15013.33±325.17 |
| Mg     | 1332.00±8.76 | 1268.63±16.43 | 1370.00±18.00 |
| Mn     | 2.40±0.17 | 1.13±0.11 | 2.30±0.09 |
| Fe     | 84.33±1.15 | 62.00±1.73 | 84.00±7.51 |
| Zn     | 17.43±2.41 | 15.63±0.35 | 15.43±0.45 |
| Cu     | 1.87±0.23 | 2.93±0.11 | 3.30±0.55 |
| Cd     | ND | ND | ND |
| Pb     | ND | ND | ND |

ND is not detected NA is Not Available; *source: (FAO, 2008); *source: FAO/WHO/Codex Alimentarum commission [17].

**Table 8: Transfer (TF) of Metals from Soil to E. Ventricosum.**

| Sites       | Sample | Metals | Na | K | Mg | Ca | Fe | Mn | Zn | Cu |
|-------------|--------|--------|----|---|----|----|----|----|----|----|
| Dire Enchini| Corm   | 0.97   | 4.22 | 0.17 | 6.0 | 0.46 | 0.02 | 0.94 | 0.8 |
| Jeldu       | Corm   | 0.92   | 3.24 | 1.24 | 4.27 | 0.60 | 0.014 | 1.3 | 1.07 |
| Jibat       | Corm   | 1.18   | 4.35 | 1.0 | 4.35 | 0.29 | 0.02 | 0.76 | 1.4 |

Citation: Bedada T, Abebaw A (2021) Metallic nutrients in enset (Ensete ventricosum) corm and soil sample from some West Shoa Zone, Oromia Regional State, Ethiopia. J Agric Sc Food Technol 7(1): 073-080. DOI: https://dx.doi.org/10.17352/2455-815X.000091
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