

ABSTRACTS

AIM:

This research investigates the pollution level of heavy metals and their variation in five selected areas in Kano state, Nigeria. The heavy metals investigated are Cadmium (Cd), Chromium (Cr), Manganese (Mn), Zinc (Zn), Lead (Pb), Copper (Cu), Iron (Fe), and Nickel (Ni).

PLACE AND DURATION OF STUDY:

The area under investigation is found to be associated with various activities (e.g. Industrialization, blacksmithing, metal scrap dump site, agriculture etc) for the past 40 years but due to increase in population, it is now a compact (Nucleated) settlement. In agricultural areas, some of the farmers use the polluted water released for their source of irrigation activities. The study covers a period of six months (November 2015 to April 2016), based on the period of activities in the selected sites (e.g. cultivation by the farmers, Industrial activities, Melting, Metal scraps etc).

METHODOLOGY:

Sample Preparation, Preservation and Digestion

The soil samples are collected for Three Months (February, March and April, 2016) after the sites were prepared for three months (November, December 2015 and January, 2016) for the experiment. Each time the sample was collected it was shade-dried for seven days on the plastic trays to avoid metal contact. The dried samples are grinded using ceramic coating, then sieved into refined powder and leveled into polythene bags, for storage under the ambient temperature (Isa *et al.*, 2017).

PROCEDURE

A beaker containing 1gm of soil sample and 30ml of Aqua regia (HNO₃ +HCl) at 3:1 ratio was placed into mixer (vibrator) for one hour thirty minutes. Filter paper (Whiteman No.42) was used to filter the solution (suspension) on a separate beaker and distilled water was added to marked 50ml. Atomic absorption spectroscopy ((ASS)_{-Model 210 VGP}) was used to determine the presence and concentration of; Pb, Cd, Ni, Cu, Cr, Zn, Mn and Fe with the corresponding wavelength of each metal; 248.3, 213.9, 232, 357.9, 228.8, 217, 279.5 and 324.8nm respectively. The result obtained was further analyzed using SPSS 20.0.

RESULTS:

It is found that in all the five (5) sites (locations) of the study, there exist all the eight heavy metals (HMs) in varying concentrations. The slopes are deduced with the values as; Cd (0.109), Cr (0.119), Cu (0.022), Fe (0.026), Ni (0.013), Mn (0.02), Pb (0.022) and Zn (0.017). These values are used to compute the concentration of the eight metals identified, which gave the order of concentration s as: Zn>Ni>Mn>Fe>Cu>Pb>Cr>Cd (for February and March, 2016) but Ni>Cu>Pb>Mn>Fe>Cd>Cr>Zn (For April, 2016). The pollution load index for the five locations is obtained as: 1.2927 (BUK), 1.6249 (Naibawa), 1.6783 (Kofar Ruwa), 1.4197 (BUK Screen) and 1.559 (Sharada).

Conclusion

The results obtained reveals that eight (8) HMs are determined - (Cr, Cd, Cu, Fe, Zn, Ni, Pb, and Mn). These HMs recorded different/varying concentrations (within the soil). The correlation matrix generated from the concentrations of samples obtained shows that in each site, there is group of HMs that originate from the same source(s) and others that emanate from another source (s). In Naibawa, Cd, Cr. Cu. Fe. Ni, Mn, and Pbhave high probability of originating from the same source while Zn might have originated from a different source But in Kofar Ruwa site, Fe and Zn recorded high probability of originating from the same source while Cd, Cr, Cu, Ni, Mn and Pb are from other source(s). In BUK - E; Cd, Cr, Cu, Fe and Pb are probably from the same source, while Ni, Mn and Znare from different source. In the control area (BUK C site), Cd, Ni, Mn, Pb recorded values have probabilities, indicating they are from the same source while Cr, Cu, Fe and Zn are contrary from the latter. In the overall sites, the data generated reveals that Cr and Cu are from the same source while Cd, Cr, Cu, Ni, Mn and Pb are from another source. From the soil pollution load index computed (before, during and after planting), the study indicates decrease in the level of contamination in all the sites.

Keywords: AAS, HeavyMetals, Phythoremediator, Sunflower and Vegetable.

1. INTRODUCTION

Several efforts are made towards safeguarding the health of the society by conducting researches on the composition of samples using various techniques. These researchers range from identification, determination, study and evaluation of samples (Biological and geological). Natasaet al., (2015) reports that; Melting operation, sludge dumping, intensive agriculture, traffic activities, power transmission, cement - pollution and smelting are possible ways of heavy metal accumulation. Metal Contamination in agricultural soil is of increasing concern, due to food safety issues and potential health risk (Yeasminet al, 2013). Heavy Metal (HMs) pollution has pervaded many parts of the developing countries and affects humans because of their longevity and accumulation in their organs via different ways (Li et al., 2014 and Zhang et al, 2010). The non-biodegradability of HMs and their potential to cause inappropriate effect made them the most noxious material (Seydou and Timoty, 2016). It is widely reported that they have both positive and negative role in human life. The elements play important role in the biological process, but at high concentrations they may be toxic to biota, disturb the biochemical process and cause hazards. Excessive content of HMs beyond maximum permissible level (MPL) leads to number of nervous, cardiovascular, renal, neurological impairment as well as bone diseases, which significantly contribute to decrease human life expectancy (9-10 years), within the affected area and several other health disorders (Yeasminet al, 2013). Khan et al., (2008), reports that National Research Council (NRC) has outlined four steps (processes) in estimating health risk agent, which are hazard identification, exposure assessment, dose/response assessment, and risk 31 32 characterization. This problem is not an exception in Nigeria as Ahmed et al., (2016) reports that the risk level Nigerians 33 and other African countries are exposed to. There search scope is restricted to Kano State, Nigeria (within five locations). Kano is a state in Nigeria, located between the latitude 12°15'S and 12°35'N of equator and the longitude 8°20'W and 34 35 8°27'E of meridian, as presented in figure 1. 36

The study areas are found to be an industrial area for the past 40 years but due to the increase in population, the areas is 37 now a compacted (Nucleated) settlement. Also some of the peopleuse the water released from the industries for their 38 39 irrigation activities.

The study is aims at determining the level of concentration of HMs (as Pollutants) in some selected area in Kano state 40 due to the increased in population, industrial activities (effluent), metal scraps, agricultural activities, provided possible 41 solution and the to call the attention of the authority to come to the aid of the residents. The specific objective of the 42 43 study is identifying the HMs in these areas, finding out whether the metals comes from the same source or not and at what level of concentration are they placed and determining the level of contamination in the selected areas. 44



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46 Figure 1: The five (5) selected sample site: Sharada, KofaRuwa, Naibawa and Bayero University (two locations) 47 Kano.

48 **1.1 THEORETICAL BACKGROUND** 49

One of the governing equations that gives a relationship between, α (the analyte's absorptivity with units of cm⁻¹conc⁻¹); 50 51 Concentration, C; Absorbance, A; and width, b; is the Beer's law (some time called Beer - Lambert Law), as presented in equation (1): 52

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 $A = \alpha b C$

(1)

When expressing the concentration using molarity, then α will be replaced with the molar absorptivity, ε , which has unit of 54 $cm^{-1} M^{-1}$. Hence: 55 56

 $A = \varepsilon bC$

(2)

The concentration of HMs is directly related to the absorbance of the metals by a substance. In this research work we are 57 interested in the Soil Samples Concentration (C sample), and Pollution Load Index (PLIs). In order to have the 58 concentrations of these metals, the equations used by Udoet al. (2009) and Khan et al., (2008) were employed. 59 60

$$C_1 V_1 = C_2 V_2 \tag{3}$$

62 Where C_n is the concentration of solution and V_n is the volume (for n=1,2,3,...,n).

63 Concentration of sample (C sample)

$$C_{sample} = \left(\frac{Abs.}{Standard/Slope}\right) \times \frac{Volume}{WeightofSample}$$
(4)

66 where Abs. is Reading of absorbance (with respect to Heavy Metals)

Pollution Load Index Soil (PLIs) 68

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Ahmed *et al.*, (2013), reported methods used in indicating the level of contamination of soil ranging from low, moderate and severe contamination. The equations are given as:

$$C_f = \frac{C_n}{C_r}$$

72 Where C_f is the contamination factor, C_n is the soil concentration and C_r is the background level of the study area. The 73 PLIs is a dimensionless quantity, which depends on C_f . The expression for PLIs is given as:

$$PLIs = \sqrt[n]{C_{f1} + C_{f2} + C_{f3} + \dots + C_{fn}}$$
(6)

(5)

75 **2.0 Materials and Method**

Five (5) experimental sites are set up within Kano State, Nigeria. These are: (a) Bayero University, Kano Screen House (BUK-C) $- 8^{\circ}28'0" \in \& 11^{\circ}59'0" N$, (b) Bayero University, Kano Environment (BUK-E) $- 8^{\circ}28'0" \in \& 11^{\circ}59'0" N$ (c) Kofar Ruwa (K) $- 8^{\circ}29'5" \in \& 12^{\circ}1'5" N$,(d) Naibawa (N) $- 8^{\circ}35'0" \in \& 11^{\circ}58'0"N$ and (e) Sharada (S)- $8^{\circ}29'5" \in \& 11^{\circ}58'0"N$. as shown in Figure 1.

81 **2.1 Sample Preparation, Preservation and Digestion**

The soil samples are collected for Three Months (February, March and April, 2016) after the sites were prepared (before, during and after plantations) for three months (November, December and January, 2015).Each time the sample were collected it was shade-dried for seven days on the plastic trays to avoid metal contact. The dried samples are grinded using ceramic coating, then sieved into refined powder and leveled into polythene bags, for storage under the ambient temperature (Isa *et al.*, 2017).

87 2.2 Procedure

A beaker containing 1gm of soil sample and 30ml of Aqua regia (HNO₃ +HCl) at 3:1 ratio is placed into mixer (vibrator) for one hour thirty minutes. Filter paper (Whiteman No.42) is used to filter the solution (suspension) on a separate beaker and distilled water is added to marked 50ml. Atomic absorption spectroscopy ((ASS)_{-Model 210 VGP}) is used to determine the presence and concentration of; Pb, Cd, Ni, Cu, Cr, Zn, Mn and Fe with the corresponding wavelength of each metal; 248.3, 213.9, 232, 357.9, 228.8, 217, 279.5 and 324.8nm respectively. The result obtained was further analyzed using SPSS 20.0.

94 2.2 Statistical Method

95 SPSS 20.0 version was employed to analyze the concentrations of the eight heavy metals determined. Regression 96 analysis is also used to obtain the slope values that are used to compute the concentrations. The correlation matrix was 97 equally generated and the heavy metals are identified and discussed to be from the same or different source(s). The 98 correlation is in term of probabilities with a heavy metal selected as it reference base on the activities in the area. 99

100 3.0 Results and Discussion

101 **3.1 Samples Concentrations**

The concentration of heavy metals is directly related to the absorbance of metals by the samples, equation (3) was used to calculate the concentrations of metals in the sample. The standard/slope was computed using equation (4). Different volumes of solutions at different concentrations are prepared and analyzed using AAS machine to obtain the absorbance. The concentration and absorbance of each metal are given in **Table 1**

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112 Table 1: Cd, Cr, Cu Fe, Ni, Mn, Pb, and Zn Concentration (mg/kg) and Absorbance Values

Cadmium (Cd) 01.00 00.80 00.60 00.40 00.20 00.00 Concentration 00.111 00.087 00.063 00.044 00.023 00.00 Absorbance Chromium(Cr) 01.00 00.80 00.60 00.40 00.20 00.00 Concentration Absorbance 00.118 00.097 00.071 00.049 00.026 00.00 Copper(Cu) Concentration 05.00 04.00 03.00 02.00 01.00 00.00 Absorbance 00.111 880.00 00.066 00.043 00.022 00.00 Iron(Fe) Concentration 10.00 08.00 06.00 04.00 02.00 00.00

Absorbance	00.262	00.212	00.164	00.112	00.054	00.00			
Nickle (Ni)									
Concentration	10.00	08.00	06.00	04.00	02.00	00.00			
Absorbance	00.131	00.112	00.084	00.053	00.027	00.00			
Manganese(Mn)									
Concentration	10.00	08.00	06.00	04.00	02.00	00.00			
Absorbance	00.202	00.162	00.122	00.081	00.042	00.00			
Lead(Pb)									
Concentration	10.00	08.00	06.00	04.00	02.00	00.00			
Absorbance	00.223	00.174	00.129	00.086	00.045	00.00			
Zinc(Zc)									
Concentration	10.00	08.00	06.00	04.00	02.00	00.00			
Absorbance	00.171	00.137	00.102	00.067	00.031	00.00			

The values of the concentration for these heavy metals (HMs), in the soil samples were analyze in five different sites. In determining the concentration (in the soil samples) of the HMs, various solution (with different volume) and varying concentration and its equivalent absorbance was produced using Atomic Absorbance Spectroscopy (AAS), The values of absorbance and concentrations are tabulated inTable 1.Using the same table 1, slopes of these HMs were deduced with the values as; Cd (0.109), Cr (0.119), Cu (0.022), Fe (0.026), Ni (0.013), Mn (0.02), Pb (0.022) and Zn (0.017). The computed values of the slope reveal that the concentrations is directly proportional to the absorbance. Using equation (4) the concentrations were generated and presented in **Figures 2**.



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122 Figure 2a: Comparison of Heavy metals from different sites base on their concentrations in February 2016

Figure 2a reported the concentration of each HMs with respect to their sites. In February 2016, all the HMs studied Zn and Ni recorded the highest values, followed by Fe, Mn, Cu, Pb, Cr and Cd.



126 Figure 2b: Comparison of Heavy metals from different sites base on their concentrations in March

In Figure 2b, the same HMs were presented for the month of March 2016, where it was observed that there are general decrease in the concentrations of the HMs when compared with the concentrations of these metals in the months of February 2016. By extension there are changes in the conditions of the sites (soils).

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132 Figure 2c: Comparison of Heavy metals from different sites base on their concentrations in April

As for the month of April 2016, (Given in Figure 2c), similar behavior as recorded in the previous month (March, 2016) was significantly seen, this is connected to the common activities in the sites (farming) as reported by Isa *et al.*, (2017).

However Cu and Zn appear to have the highest concentrations when compared with the other HMs.

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Figure 3.0: Total concentration of the Heavy Metals for Three Months (February, March and April,

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2016).

Figure 3.0, gave all the concentrations of the eight (8) HMs in different sites and their comparison base on Months and metals. From it, **Ni** records the highest concentration and **Cd** has the least. Overall the concentration base on monthly basis ascends from February, March, and then April 2016.

144 **3.2 Correlation of the Eight Heavy Metals.**

To investigate the correlations between the metals, SPSS 20.0 was used and the result obtained was tabulated in Tables 2, 3, 4, 5, and 6.

147 Table 2: Correlation Matrix of the Heavy Metals from Naibawa site.

	Cd	Cr	Cu	Fe	Ni	Mn	Pb	Zn
Cd	1.000							
Cr	0.952	1.000						
Cu	0.990	0.900	1.000					
Fe	0.947	0.804	0.983	1.000				
Ni	0.980	0.873	0.998	0.992	1.000			
Mn	0.996	0.923	0.998	0.971	0.993	1.000		
Pb	0.982	0.78	0.999	0.990	1.000	0.995	1.000	
Zn	-0.993	-0.908	-1.000	-0.979	-0.997	-0.999	-0.998	1.000

From Table 2, it was observe that there is highest probability that Cd, Cr, Cu Fe, Ni, Mn, and Pb are from the same source(s), while Zinc originate from a different source(s). This was expected considering the nature of the site (Dump site).

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153 **Table 3: Correlation Matrix of the Heavy Metals from KofarRuwa site.**

	Cd	Cr	Cu	Fe	Ni	Mn	Pb	Zn
Cd	1.000							
Cr	0.977	1.000						
Cu	0.985	0.925	1.000					
Fe	-0.376	-0.170	-0.531	1.000				
Ni	0.979	1.000	0.929	-0.178	1.000			
Mn	0.947	0.994	0.876	0.057	0.993	1.000		
Pb	0.943	0.992	0.871	0.047	0.991	1.000	1.000	
Zn	-0.996	-0.996	-0.967	0.296	-0.993	-0.971	-0.968	1.000

Table 3, reported the correlation probabilities of the HMs at Kofar Ruwa market (Iron scraps). It was found that Cd, Cr and Cu originate from the same source(s). Fe, Ni, Mn, Pb and Znstands alone with a unitaryprobability but variable probabilities when compared to the other HMs in the sites. However looking at the nature the site this results is expected.

157 Table 4: Correlation Matrix of the Heavy Metals for BUK Environs site.

	Cd	Cr	Cu	Fe	Ni	Mn	Pb	Zn
Cd	1.000							
Cr	0.873	1.000						
Cu	0.943	0.662	1.000					
Fe	0.753	-0.978	0.492	1.000				
Ni	-0.237	-0.680	0.99	-0.818	1.000			
Mn	-0.339	0.162	0.632	0.364	-0.834	1.000		
Pb	0.412	-0.085	0.691	-0.290	0.788	-0.997	1.000	
Zn	-0.026	0.465	-0.356	0.638	-0.965	-0.949	-0.922	1.000

158 In table 4, the probability shows that Cd, Cr, Cu, Fe and Pb are produced from the same sources while Ni, Mn, and Zn 159 were produced from a different source(s).

160 Table 5: Correlation Matrix of the Heavy Metals for BUK Screen House site.

	Cd	Cr	Cu	Fe	Ni	Mn	Pb	Zn
Cd	1.000							
Cr	0.353	1.000						
Cu	-0.875	0.144	1.000					
Fe	-0.975	-0.137	0.961	1.000				
Ni	1.000	0.351	-0.876	-0.976	1.000			
Mn	0.986	0.501	-0.784	-0.926	0.986	1.000		
Pb	0.866	-0.162	-1.000	0.955	0.867	0.773	1.000	
Zn	-0.996	-0.433	0.830	0.952	-0.996	-0.997	-0.820	1.000

161 The probabilities in Table 5, report that Cd, Ni, Mn and Pb are produced from the same source while Cr, Cu, Fe, and Zn 162 have been from another source(s).

163 **Table 6: Correlation Matrix of the Heavy Metals for all the sites**.

	Cd	Cr	Cu	Fe	Ni	Mn	Pb	Zn
Cd	1.000							
Cr	0.993	1.000						
Cu	0.989	0.993	1.000					
Fe	-0.963	0.989	0.992	1.000				
Ni	0.984	-0.963	0.946	-0.989	1.000			
Mn	0.985	0.985	0.949	-0.902	1.000	1.000		
Pb	0.943	0.990	0.958	-0.915	0.999	1.000	1.000	
Zn	-0.975	-0.975	-0.931	0.879	-0.999	-0.999	-0.997	1.000

Table 6 shows the overall summary of the correlation between the metals studies. It was found that Cu and Cr are produced from the same source in all the locations while Cd, Ni, Mn, Zn and Pb are produced from different sources.

167 **3.3 Pollution Load Index (PLIs)**

Pollution load Index (PLIs) is another way used to determine the level of pollution in a given sample (soil). Three (3) factors were studied, using the concentrations of the eight HMs (geological samples)computed using equations (4). These factors are Concentration of soil (C_n), Background Concentration (C_r) and Contamination factor (C_f) using equations 5 and 6. These factors were then employed in computing the PLIs and presented in **Figures4a**, **4b**, **4c**, **4d**, **and 4e**.



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173 Figure 4a: Barchart indicating BUK site contamination level and the value of PLIs (1.2927)

Figure 4a shows that there is high contamination factor of the HMs, with Ni, recording the highest, their by decreasing in the following sequence Fe,Mn,Zn,Cd,Pb, Cu, and Cr in BUK site with PLIs value of 1.2927.



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177 Figure 4b: Barchart indicating Naibawa site contamination level and the value of PLIs (1.6249)

178 In Naibawa it was found that Zn recorded the highest contamination value in the soil followed by Fe, Cd, Mn, Cr, Ni, Cu 179 and Pb with PLIs value of 1.6249 as presented in figure 4b.



183 Figure 4c: Barchart indicating Kofar Ruwa site contamination level and the value of PLIs (1.6783)

184 Similarly Zn, Fe, Mn, Ni, Pb, Cd, Ni, and Cu are the order of the level of contamination (by HMs) in Kofar Ruwa site. The 185 PLIs computed in this site is 1.6783 as presented in Figure 4c.



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187 Figure 4d: Barchart indicating BUK Screen House site contamination level and the value of PLIs (1.4197)

188 In the control site (BUK Screen House) the contamination is relatively low compare to the background and concentration

189 of the HMs. However, the PLIs was obtained to be 1.4197. The HMs contamination factor level decrease in sequence Zn,

- 190 Cd, Mn, Ni, Cr, Pb, and Cu, this is presented in Figure 4d.
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193 Figure 4e: Barchart indicating Sharada site contamination level and the value of PLIs (1.559)

The PLIs value is 1.559 in Sharada with Fe recording the highest contamination factor then followed by Mn, Cd, Zn, Pb, Ni, Cr, and Cu.

196 Considering figures 4a to 4e, the computed contamination factors are all greater than 1, this means that the sites are 197 contaminated.

198	Table 7: Pollution Load Index of Soil (I	PLIs) Site

PLIs	Before Planting of the	During Planting of the	After Planting of the
	Samples	Samples	Samples
BUKS	1.2927	1.2444	1.2318
Naibawa	1.6249	1.6067	1.5098
KofarRuwa	1.5783	1.5386	1.4372
BUKN	1.4197	1.4029	1.3028
Sharada	1.5590	1.5253	1.4449

199 The level of contamination of the soil (sites) was analyze in three phase. The first phase is before plantation (farming), i.e. 200 the three months preparation for cultivation, then the second phase is during the plantations and t east phase is after the plantations. In each period the samples were collected and the PLIs was determined. Table 7 gave the tabulated readings 201 for the three periods for each sites. The values compute in relation to the concentrations (C_n , C_r , and C_f), are used to 202 compute the level of contamination. PLIs is use to indicate at what level is our site place base on the values obtained. 203 According to Ahmed et al., (2014), if the C_f 1, indicates low contamination, $1 \le C_f \le 3$; Moderate Contamination, $3 \le C_f \le 3$ 204 205 6 and C_f> 6; Severe Contamination. While for PLIs: when PLIs < 1; absence of Contamination, PLIs = 1; Low 206 contamination, and PLIs >1; High contamination. Figure 5 show the representation of table 7 in form of a bar chart.



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Figure 5: Bar chart representing contamination levels from the five sites.

209 It can be deduced that the five sites are contaminated with HMs. However looking at the different periods in which 210 pollution level varies, one can say that the pollution reduces with a time relative to the plantation of the samples. As reported by Isa *et al.* 2017, the declining (decreasing) values in this report indicate that the PLIs decreases as the plants grow in the five sites as a result of absorption of the metals by the plants.

213 **4.0 CONCLUSION**

- The concentrations of eight (8) HMs (Cr, Cd, Cu, Fe, Zn, Ni, Pb, and Mn) are determined. These heavy metals recorded
- different/varying concentrations, within the soil and the plant's samples.
- 216 The correlation matrix generated from the concentrations of samples obtained reveals that in each site, there are group
- of HMs that originate from the same source(s) and others that emanate from the other source(s).
- The Pollution Load Index computed (PLI) in each site is greater than 1, hence the sites are considered to be
- 219 contaminated. However the pollution Load Index computed, before, during, after planting of the two samples, its hows
- that there is significant decrease in the level of contamination which could be attributed to some amount of the HMs
- absorbed by the samples during plantation of the samples, and if more are planted, the metal level in the soil would be
- reduced drastically.
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